I. Introduction

A. Purpose and Overview

The Tobacco Control Act was enacted on June 22, 2009, amending the Federal Food, Drug, and Cosmetic Act (FD&C Act) and FCLAA, and providing FDA with the authority to regulate tobacco products (Pub. L. 111–31; 123 Stat. 3109–21). This rule is the result of a request for regulations under the authority provided by the act to implement a requirement for health warnings on cigarette packages and in cigarette advertisements. This rule amends the Federal Cigarette Labeling and Advertising Act (Tobacco Control Act) to require each cigarette package and advertisement to bear one of nine new textural warning statements. The final rule specifies the color graphic images that must accompany each of the nine new textural warning statements.

II. Proposed Rule

The tobacco industry has challenged the requirement for large graphic warnings on cigarette packages and in cigarette advertisements. This final rule amends the Federal Cigarette Labeling and Advertising Act (Tobacco Control Act) to require each cigarette package and advertisement to bear one of nine new textural warning statements. This final rule specifies the color graphic images that must accompany each of the nine new textural warning statements.

III. Final Rule

The Food and Drug Administration (FDA) is amending its regulations to add a new requirement for the display of health warnings on cigarette packages and in cigarette advertisements. This rule implements a provision of the Family Smoking Prevention and Tobacco Control Act (Tobacco Control Act) that requires FDA to issue regulations requiring color graphics, depicting the negative health consequences of smoking, to accompany the nine new textural warning statements required under the Tobacco Control Act. The Tobacco Control Act amends the Federal Cigarette Labeling and Advertising Act (FCLAA) to require each cigarette package and advertisement to bear one of nine new textural warning statements. This final rule specifies the color graphic images that must accompany each of the nine new textural warning statements.

DATES:

This rule is effective September 22, 2012. See section VIII of this document, Implementation Date, for additional information. The incorporation by reference of a certain publication listed in the rule is approved by the Director of the Federal Register as of September 22, 2012.

FURTHER INFORMATION CONTACT:

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II. Need for the Rule and Responses to Comments

A. Cigarette Use in the United States and the Resulting Health Consequences

1. Smoking Prevalence and Initiation in the United States

In explaining the need for the proposed rule, we provided information in the NPRM on smoking prevalence and initiation rates among adults and children in the United States. As stated in the NPRM (75 FR 69524 at 69526), approximately 46.6 million U.S. adults (or 20.6 percent of the adult population) are cigarette smokers (Ref. 4). Moreover, almost half (46.3 percent) of youth in grades 9 through 12 in the United States have tried cigarette smoking, and 19.5 percent of youth in grades 9 through 12 are current cigarette smokers (Ref. 5 at p. 7). Smoking rates among U.S. adults have shown virtually no change during the 5-year period from 2005 to 2009 (Ref. 4), and smoking rates among U.S. youth have not decreased from 2006 to 2009 (Ref. 6).

Furthermore, each year millions of U.S. adults and children become new smokers. Data from the 2008 National Survey on Drug Use and Health indicate that 2.4 million persons aged 12 or older in the United States smoked cigarettes for the first time in the past 12 months (Ref. 7 at p. 59). In addition, these data indicate that almost 1 million Americans aged 12 or older started smoking cigarettes daily within the past 12 months (Ref. 7 at p. 60).

In other words, approximately 6,600 people aged 12 or older in the United States become new cigarette smokers every day, and more than 2,500 individuals become new daily cigarette smokers every day (Ref. 7 at pp. 59–60). Moreover, nearly 4,000 of the people who become new cigarette smokers every day and nearly 1,000 of the individuals who become new daily cigarette smokers every day are children under the age of 18 (Ref. 7 at pp. 59–60). These statistics for youth smokers are particularly concerning, as studies suggest that the age people start smoking can greatly influence how much they smoke per day and how long they smoke, which in turn influences their risk of tobacco-related disease and death (Refs. 8, 9, and 10).

FDA received many comments that were strongly supportive of the proposed rule, some of which provided data and information consistent with that in the NPRM regarding cigarette use prevalence and initiation in the United States (75 FR 69524 at 69526 through 69527). Many of these comments also stated that smokers would be more
likely to quit smoking and that nonsmokers would be less likely to start smoking if cigarette advertisements and packages display, visually and graphically, the health effects of cigarettes. Most of these comments expressed a belief that the required warnings would help reduce the existing and future use of cigarettes. Some comments that were supportive of the proposed rule discussed the smoking prevalence and initiation rates in the United States in particular populations. These comments, and FDA’s responses, are summarized in the following paragraphs.

(Comment 1) Multiple comments indicated that people with less education and lower incomes have higher smoking prevalence rates in general. One comment from a health care association indicated that women of low educational background have higher smoking prevalence rates and that many of these women still are not aware of cigarettes’ impact on life expectancy, heart disease, and pregnancy.

(Response) We agree that adults with low education levels have higher than average smoking prevalence rates. For example, as discussed in the NPRM (75 FR 69524 at 69526), 49.1 percent of adults with a General Education Development certificate (GED) and 28.5 percent of adults with less than a high school diploma were current smokers in 2009, compared with 5.6 percent of adults with a graduate degree (Ref. 4). We also agree that graphic health warnings may be particularly important communication tools for these smokers, as there is evidence suggesting that countries with graphic health warnings demonstrate fewer disparities in health knowledge across educational levels (Ref. 11 at p. 18 and Ref. 3 at p. 295).

(Comment 2) Multiple comments noted that smoking rates vary by race and ethnicity, with American Indians/Alaska Natives having the highest rates. One comment also noted that the health and economic costs of smoking vary by race and ethnicity. For example, the comment stated that African-American smokers suffer disproportionately from smoking-related diseases, including lung cancer, heart disease, and strokes (citing Ref. 12), and called for measures to address these disparities.

One comment from a State public health agency indicated that racial minority populations and economically disadvantaged populations have smoking prevalence rates that are two to three times higher than the general population.

(Response) We agree that smoking rates vary by race and ethnicity and socioeconomic status. For example, prevalence data from 2009 for current U.S. adult cigarette smokers indicate that, among racial/ethnic groups, adults reporting multiple races had the highest smoking prevalence (29.5 percent), followed by American Indians/Alaska Natives (23.2 percent) (Ref. 4). We also agree that economically disadvantaged populations have higher smoking prevalence rates. For example, data from 2009 indicate that the prevalence of current smoking was higher among U.S. adults living below the Federal poverty level (31.1 percent) than among those at or above this level (19.4 percent) (Id.).

We have selected required warnings that will help effectively convey the negative health consequences of smoking to a wide range of population groups, including different racial and ethnic groups and different socioeconomic groups, and that can help both to discourage nonsmokers from initiating cigarette use and to encourage current smokers to consider quitting. For additional information regarding our selection of required warnings to reach a broad range of population groups, see section III of this document regarding our selection of the final images.

(Comment 3) Multiple comments stated that tobacco use disparities exist among lesbian, gay, bisexual, and transgender individuals. One comment from a community organization stated that lesbian, gay, bisexual, and transgender individuals smoke at rates almost 50 percent to 200 percent higher than the rest of the population and strongly supported the proposed rule.

(Response) We agree that evidence suggests that gay, lesbian, bisexual, and transgender populations have higher smoking rates than their heterosexual counterparts (Ref. 13). The required warnings will help convey information about various health risks of smoking to individuals from a wide range of demographic groups and will help encourage smoking cessation and discourage smoking initiation.

(Comment 4) One comment from a nonprofit research organization indicated that members of the U.S. military have rates of smoking that are unacceptably high, particularly among younger members. The comment detailed the negative outcomes of smoking to military personnel, including lower physical performance, an increased risk of injury during physical tasks, a greater number of days sick and unable to report for duty, poorer job performance, and a higher likelihood of premature discharge from active duty, and stated that smoking and its negative effects among active duty personnel costs the military an estimated $1 billion annually in health care and lost productivity (Ref. 14). The comment also referred to evidence suggesting the tobacco industry has targeted military members and fought efforts to reduce tobacco product consumption by military personnel, and indicated that the proposed rule is an important step in protecting military members from the health harms of cigarette use and will likely decrease cigarette use among military personnel.

(Response) We agree that members of the U.S. military have higher smoking prevalence rates than the general population; approximately 20.6 percent of the U.S. adult population smoke cigarettes, while data from 2008 indicate that 31 percent of active duty military personnel smoke cigarettes (Ref. 15). We agree that the required warnings will help convey information about various health risks of smoking to a wide range of individuals, including members of the U.S. military and veterans who began smoking while in military service, and that the required warnings will encourage smoking cessation and discourage smoking initiation in these individuals.

2. Health Consequences of Smoking

Smoking is responsible for at least 443,000 premature deaths per year in the United States, and each year cigarettes are responsible for approximately 5.1 million years of potential life lost (Ref. 1). Annual direct health care expenses due to smoking total approximately $96 billion, and annual productivity losses due to premature deaths alone from cigarette smoking total approximately $96.8 billion (Id.).

The Agency received many comments that were supportive of the proposed rule, some of which reiterated the health risks of smoking described in the NPRM (75 FR 69524 at 69527 through 69529) and stressed the need for measures, such as graphic health warnings, to curb smoking in the United States in order to improve health and to reduce the massive health care costs attributable to tobacco-related illnesses. Some of these comments cited data demonstrating that smoking is the leading cause or most powerful risk factor for particular diseases, such as chronic obstructive pulmonary disease (COPD), bladder cancer, and atherosclerosis.

However, FDA also received multiple comments disputing the health risks of smoking. These comments and FDA’s responses are summarized in the following paragraphs.

(Comment 5) One comment from an individual expressed a belief that addiction to nicotine is 99 percent...
psychological and only 1 percent pharmacological, and that nicotine is no more addictive than caffeine.

(Response) We disagree with the assertion that nicotine addiction does not have a substantial physiologic component. While we acknowledge that behavioral processes play a role in initiation and maintenance of nicotine addiction, nicotine is a powerful pharmacologic agent that acts in a variety of ways at different sites in the body. As stated in the NPRM, nicotine causes physical dependence characterized by withdrawal symptoms that usually accompany nicotine abstinence (75 FR 69524 at 69528).

Regarding the relative addictiveness of nicotine and caffeine, caffeine is distinct from nicotine in its abuse liability, which includes a consideration of multiple factors, including the dependence potential of a substance and the degree to which it produces adverse effects (see Ref. 16 at p. 304). Caffeine produces only minimal disruptive physiological effects and, unlike nicotine from tobacco products, caffeine is generally not used in ways that are considered to be of significant adverse health effect (see Id. at pp. 285 and 304).

(Comment 6) One comment stated that nicotine withdrawal is the only medical condition that is irreversibly caused by cigarettes.

(Response) We disagree with this comment. While nicotine addiction is one negative health effect of cigarette smoking, it is not the only medical condition that is irreversibly caused by cigarettes. As detailed in the 2004 report of the Surgeon General, “The Health Consequences of Smoking,” which summarizes thousands of peer-reviewed scientific studies and was itself peer-reviewed, cigarettes have been shown to cause an ever-expanding number of diseases and conditions, including lung cancer, laryngeal cancer, oral cavity and pharyngeal cancers, esophageal cancer, bladder cancer, pancreatic cancer, kidney cancer, stomach cancer, cervical cancer, acute myeloid leukemia, all the major clinical cardiovascular diseases, COPD, and a range of acute respiratory illnesses (Ref. 2).

Maternal smoking during pregnancy causes a reduction in lung function in infants, and women who smoke during pregnancy are more likely to experience premature rupture of the membranes, placenta previa, and placental abruption (Id. at pp. 508 and 576). Smoking also increases rates of preterm delivery and shortened gestation, and women who smoke as likely as nonsmokers to have low birth weight infants; smoking also increases the risk of sudden infant death syndrome (SIDS) (Id. at pp. 569, 576, 587 and 601). Children who smoke experience impaired lung growth and an early onset of lung function decline (Id. at pp. 508–509, 2004 SG). Smoking during adulthood also leads to a premature onset of accelerated age-related decline in lung function (Id. at p. 509). Smoking also results in poor asthma control and causes a range of respiratory symptoms in children, adolescents, and adults, including coughing, phlegm, wheezing, and shortness of breath (Id.).

Furthermore, cigarette smokers have poorer overall health status compared to nonsmokers, and an increased risk of adverse surgical outcomes related to wound healing and respiratory complications compared to nonsmokers. Smokers are also at an increased risk for hip fractures, and smoking increases the risk for periodontitis, cataract, and the occurrence of peptic ulcer disease in persons who are *Helicobacter pylori* positive (Id. at pp. 717–719, 736, 777, 780, and 1418).

In addition, exposure to secondhand smoke has been shown to cause a variety of negative health effects in nonsmokers, including lung cancer, cardiovascular disease, and respiratory symptoms (see Ref. 17).

(Comment 7) Some comments were submitted by individuals disputing the negative health consequences of smoking that are described in the graphic warnings. These comments generally indicated that the individuals submitting the comments were smokers, and that they and/or their family members (who were exposed to secondhand smoke) had not experienced negative health effects from smoking.

(Response) We disagree with these comments. Cigarette smoking has been shown to cause a wide range of negative health consequences, as detailed in the previous response. Furthermore, it can be years before some of the negative health consequences of smoking clinically manifest. Thus, the personal health status of the individuals submitting these comments could change in the future. A scientific determination that a product causes a particular negative health consequence is based on data from large groups of individuals, and the fact that an individual product user has not experienced (or has not yet experienced) a particular negative health consequence does not mean the product does not cause that harm.

Moreover, to the extent these comments indicate that many smokers do not fully understand the serious health risks of cigarettes or do not believe that these risks apply to them, they illustrate the need for health warnings that effectively communicate the negative health consequences of smoking to consumers. For additional information regarding consumers’ lack of knowledge of smoking risks, see section II.C of this document.

(Comment 8) One comment stated that cigarettes are a minor public health concern compared to obesity and alcohol, and that cigarette use results in less health care costs than medical treatment for the obese.

(Response) As discussed in the NPRM, cigarette smoking is the leading cause of preventable death and disease in the United States (Ref. 4). Furthermore, cigarettes are responsible for health care expenditures and productivity losses resulting in a combined economic burden of approximately $193 billion per year (Ref. 1). The total costs of smoking to society are much higher, as the estimate for productivity losses does not include costs associated with smoking-related disability, employee absenteeism, or costs associated with secondhand-smoke attributable disease morbidity and mortality (Id.).

We disagree that cigarettes are a minor public health concern, even as compared to other public health issues, and also disagree with the implication that the public health issue of smoking should not be addressed because other public health issues exist. The required warnings will have a significant, positive impact on public health (75 FR 69524 at 69526), and as a result will help mitigate the single largest cause of preventable death and disease in the United States.

B. Inadequacy of Existing Warnings

In the preamble to the proposed rule, FDA explained how cigarette packages and advertisements can be effective channels for communication of important health information, particularly given that part-time smokers are potentially exposed to warnings more than 7,000 times per year (75 FR 69524 at 69529). However, the existing warnings have suffered from three crucial problems: (1) They have not changed in more than 25 years, (2) they often go unnoticed, and (3) they fail to convey relevant information in an effective manner. FDA also explained that larger, graphic warnings communicate the health risks of smoking more effectively. The preamble to the proposed rule presented extensive evidence from other countries’ experiences with graphic warnings as well as information from the 2007 IOM Report (75 FR 69524 at 69531). On the
basis of the available scientific evidence, the IOM concluded that larger, graphic warnings would promote greater public knowledge of the health risks of using tobacco and would help reduce consumption (Ref. 3).

We received numerous comments regarding the adequacy of the existing warnings that appear on cigarette packages and advertisements. The large majority of these comments supported our analysis of the existing warnings, but a few comments disagreed with this analysis. These comments, and our responses, are summarized in the following paragraphs.

(Comment 9) A substantial number of comments, including those from health institutions, nonprofit organizations, academics, and consumers, agreed with FDA’s conclusion that the existing warnings that appear on cigarette packages and advertisements are ineffective at conveying the health risks of smoking (75 FR 69524 at 69529 through 69531).

However, one comment stated that the current warnings were “fine.” Two comments expressed the belief that the existing warnings have worked successfully in the current information environment.

(Response) We disagree with the comments stating that the existing warnings that appear on cigarette packages and advertisements are effective. As several other comments noted, the Surgeon General has long recognized that the cigarette warnings are deficient when evaluated in terms of proper public health criteria” (Ref. 18). That same year, the IOM concluded that the warnings were “inadequate * * * and woefully deficient when evaluated in terms of proper public health criteria” (Ref. 19 at p. 237). Yet those same warnings are still in use more than 16 years after the Surgeon General’s report and 26 years after their inception. Accordingly, we conclude that the existing warnings for cigarettes do not adequately communicate the health risks of smoking.

C. Consumers’ Lack of Knowledge of the Health Risks

In the preamble to the proposed rule, FDA described how the existing warnings that currently appear on cigarette packages and advertisements have largely gone unnoticed by both smokers and nonsmokers (75 FR 69524 at 69530). FDA also provided clear evidence that the warnings have failed to convey appropriately crucial information such as the nature and extent of the health risks associated with smoking cigarettes (75 FR 69524 at 69530 through 69531).

FDA received many comments regarding the level of consumers’ knowledge regarding the health risks of smoking. Several comments stated that consumers are adequately informed about the risks of smoking or even overestimate the risks of smoking, while many other comments explained that consumers lack knowledge about a wide variety of smoking risks. A summary of these comments, and our responses, is included in the following paragraphs.

(Comment 10) Several comments, including comments from tobacco product manufacturers and individual consumers, objected to the new required warnings, in part because they claimed that consumers already know the health risks associated with smoking. The submitters expressed the belief that the new warnings are unnecessary, because the new warnings provide information that the public has been aware of for many years.

(Response) We disagree. Many comments provided significant evidence to support the notion that consumers, including those in communities with low literacy rates and military personnel, actually lack knowledge or underestimate the risks associated with smoking. As discussed in this document, this lack of knowledge may involve either an incomplete understanding of the statistical risks or a failure to understand the personal (as opposed to the statistical) risks (see also section X.B.2 of this document). There is also a possibility that the risks are not considered at the time of purchase, even if they are understood—a special problem for those who are deciding whether to start to smoke. The requirements adopted here should help to counteract all of these problems.

While most smokers understand that smoking poses certain statistical risks to their health, many fail to appreciate the severity and magnitude of those risks (Refs. 20 and 21), and there is evidence that even when smokers appreciate the statistical risk, they underestimate the personal risk that they face (Ref. 22). A 2007 survey found that two in three smokers underestimate the chance of a smoker developing lung cancer compared to a nonsmoker (Ref. 23). The survey also found that up to a third of smokers erroneously believe that certain activities, such as exercise and taking vitamins, could “undo” most of the effects of smoking (Id.).

Other research also highlights how smokers misunderstand the health effects of smoking. For example, in a 2008 survey, more than one-quarter of current smokers did not agree that smoking increases a person’s chances of getting cancer “a lot” (Ref. 24). Furthermore, one study, involving smokers’ perception of their personal risk, found that only 40 percent of current smokers believed they had a higher-than-average risk of cancer and only 29 percent believed they had a higher-than-average risk of heart disease (Ref. 25). Even among heavy smokers (those who smoke at least 40 cigarettes per day), less than half believed they were at increased risk for these diseases (Id.). In another demonstration of underestimation of personal risk, a study found that adolescent smokers underestimated their personal risk, even if they had an accurate sense of the statistical risk (Ref. 22).

A 2005 study of smokers in the United States and three other countries found that there were significant gaps in smokers’ knowledge about the risks of smoking and that smokers living in countries where health warnings referred to specific diseases and the consequences of smoking were much more likely to be aware of those consequences (Ref. 26). The study concluded that smokers are not fully informed about the risks of smoking, and that warnings that are graphic, larger, and more comprehensive in content are more effective in communicating the health risks of smoking (Id.).

Thus, even if consumers are aware of certain negative health consequences of smoking, such as lung cancer and emphysema, and even if they are aware of certain statistical risks, many smokers underestimate their personal risks, and many Americans are under-informed about other health risks associated with smoking. For example, while nearly all daily smokers in one study correctly identified that smoking caused lung cancer (99 percent) and emphysema (97 percent), a lower percentage of respondents correctly identified smoking as causing low birth weight babies (88 percent), worsened asthma (85 percent), miscarriages (76 percent), other cancers (69 percent), head and neck cancers (68 percent), cervical cancer (48 percent), stomach ulcers (46 percent), reproductive difficulties (44 percent), osteoporosis (41 percent), and SIDS (40 percent) (Ref. 27). In fact, research indicates that most people know only one or two of the many diseases causes by smoking. One survey found that while a majority of people knew that smoking caused life-threatening illnesses, more than half of the respondents were unable to name a smoking-related illness other than lung cancer (Ref. 28). Similarly, researchers
found that when asked about health risks of smoking, 39 percent of respondents either answered incorrectly or said they did not know (Ref. 29). Americans also lack adequate understanding of the addictive nature of cigarettes. Although one comment provided local surveys showing that adults already know that cigarettes are addictive, there is also evidence that many adolescents do not appreciate the addictive nature of cigarettes. The 2007 IOM Report explained that “adolescents misperceive the magnitude of smoking harms and the addictive properties of tobacco and fail to appreciate the long-term dangers of smoking, especially when they apply the dangers to their own behavior” (Ref. 3 at p. 93). In addition, one survey found that fewer than 5 percent of daily smokers in high school think that they will still be smoking at all in 5 years, yet more than 60 percent of high school smokers are regular daily smokers 7 to 9 years later (Ref. 30). Another survey found that only 7.4 percent of adult smokers and 4.8 percent of young smokers expected to smoke longer than 5 years when they started, but 87 percent of these adults and 76 percent of these youth reported that they had been smoking for more than 5 years (Ref. 31).

There is also evidence that certain demographic groups are even less aware of the negative health consequences of smoking, which is particularly concerning in light of the evidence that these groups also have some of the highest smoking prevalence rates (see section II.A.3 of this document). For example, research shows that knowledge of smoking risks is lower among people with lower incomes and fewer years of education (Refs. 32 33 and 24). Smokers in the military also underestimate the actual risk of serious disease and substantially underestimate their own risks (a point that fits well with the evidence of underestimation of personal risks) (Refs. 34 35 and 36). In addition to underestimating the risks smoking poses to their own health, Americans understate the health effects of secondhand smoke on others. In the 2010 Report, “How Tobacco Smoke Causes Disease: The Biology and Behavioral Basis for Smoking-Attributable Disease,” the Surgeon General concluded that “many of the effects from active smoking can be observed in persons involuntarily exposed to cigarette smoke” (Ref. 37). In addition, individual studies have shown that secondhand smoke triggers childhood asthma and is associated with bone and cancer (Ref. 17). Yet, most parents believe that smoke exposure has little or no negative impact on children’s asthma (Ref. 38), and a 2009 study found that nearly one-fifth of Americans do not believe that secondhand smoke is dangerous to nonsmokers (Ref. 39).

There is a final point. Even if many people do have an accurate understanding of the statistical risk, and even if, in the abstract, many smokers also have an accurate understanding of their personal risk, that understanding may be too abstract to be thought of at the time of purchase, especially (but not only) for those who are starting to smoke. Efforts to make the relevant risks salient are justified and indeed required under the Tobacco Control Act.

(Comment 11) A few comments claimed that adults actually overestimate the risks of smoking-related disease, and stated that this further underscores the lack of a need for graphic health warnings. In particular, one comment referred to a Montana survey in which adults believed that smoking caused colon cancer.

(Response) We disagree with these comments. While the Montana survey referred to in one of the comments indicates that some consumers are not aware of the precise relationship between smoking and certain diseases (for example, the 2004 Surgeon General’s report notes that the evidence is suggestive but not sufficient to infer a causal relationship between smoking and colorectal cancer (Ref. 2 at p. 26)), we are aware of significant research indicating that many consumers are not sufficiently aware of the risks associated with smoking, as discussed in the previous response. We find that the weight of evidence clearly demonstrates that many consumers lack adequate knowledge about the health risks of smoking—especially the personal risks. In addition, the comments claiming that adults overestimate smoking’s risks fail to take into account consumers’ lack of knowledge of other health risks due to smoking, like the dangers of secondhand smoke, reproductive difficulties, and miscarriages, as described in the previous response.

D. Larger, Graphic Warnings Communicate More Effectively

Since Canada first introduced graphic health warnings for cigarettes in 2001, an extensive evidence base has been developed to examine the effects of graphic health warnings in Canada and in the more than 30 other countries that have adopted similar requirements for graphic health warnings on cigarettes. As FDA noted in the NPRM (75 FR 69524 at 69531 through 69533), the research literature indicates that large graphic health warnings, such as those being required in this rule, are more likely than text-only warnings to (1) get consumers’ attention, (2) influence consumers’ awareness of cigarette-related health risks, and (3) affect smoking intentions and behaviors. FDA received many comments on the efficacy of large, graphic warnings, as well as comments regarding the potential for any rebound effect from the use of graphic warnings. Those comments, and FDA’s responses, are summarized in the following paragraphs.

(Comment 12) A wide variety of comments, including those from health institutions, nonprofit organizations, and academics, agreed with FDA’s findings in the NPRM that larger, graphic warnings are effective.

However, several comments stated that the changes in the format and placement of the warnings being proposed, including the use of graphic images, will not result in reductions in cigarette use given the experiences in other countries. For example, one comment noted that Health Canada’s own data found, among other things, that there was no statistically significant decline in smoking incidence consumption for adolescents or adults after the introduction of graphic warnings. This comment expressed the belief that Canada’s warnings have been ineffective and that FDA’s graphic health warnings will be similarly ineffective.

(Response) For the reasons stated in the NPRM, we conclude that larger, graphic warnings are effective in conveying the health risks of smoking, influencing consumer awareness of these risks, and affecting smoking intentions. We disagree with comments stating that the change in format and placement of the warnings will not be effective. The set of required warnings we have selected will satisfy our primary goal, which is to effectively convey the negative health consequences of smoking on cigarette packages and in advertisements, and this effective communication can help both to discourage nonsmokers, including minor children, from initiating cigarette use and to encourage current smokers to consider cessation to greatly reduce the serious risks that smoking poses to their health.

The research literature clearly indicates that larger, graphic warnings are effective at communicating the health risks associated with smoking, encouraging users to quit smoking, and discouraging nonsmokers from beginning to smoke. We already included significant research to
substantiate this conclusion in the preamble to the proposed rule, and the comments did not specifically dispute this analysis (see 75 FR 69524 at 69531 through 69532). In addition, as we noted in the NPRM, the available evidence demonstrates that graphic health warnings are (1) more likely to be noticed than text-only warnings, (2) more effective for educating smokers about the health risks of smoking and for increasing the time smokers spend thinking about the health risks, and (3) associated with increased motivation to quit smoking (Id.). As several comments noted, evidence from countries with graphic health warnings also indicates that such warnings are an important information source for younger smokers, and that pictures are effective in conveying messages to children (Ref. 40 at pp. 3, 20, and 24–26). These important effects of graphic warnings are sustained longer than any impact from text-only warnings (Ref. 41).

Further, the data from Health Canada does not indicate that the warnings have been ineffective at conveying the health risks of smoking and impacting smoking intentions. We cited several studies in the preamble (including data from Health Canada) that illustrated the effectiveness of the Canadian graphic health warnings, which have been found effective at providing youth and adult smokers with health information, making consumers think about the health effects of smoking, and increasing smokers’ motivations to quit smoking, among other things (see 75 FR 69521 at 69532). For example, national surveys conducted on behalf of Health Canada indicate that approximately 95 percent of youth smokers and 75 percent of adult smokers report that the Canadian pictorial warnings have been effective in providing them with important health information (Ref. 3 at p. 294).

(Comment 13) One comment suggested that the new required warnings will have a greater impact on nonsmokers who inadvertently view cigarette packages than on smokers and, therefore, will not be effective in achieving FDA’s goals.

(Response) We are not aware of any evidence to substantiate this comment. Further, our required warnings are intended to have an impact on both smokers and nonsmokers. As stated in the preamble to the proposed rule, “the new required warnings are designed to clearly and effectively convey the negative health consequences of smoking on cigarette packages and in cigarette advertisements, which would help both to discourage nonsmokers, including minor children, from initiating cigarette use and to encourage current smokers to consider cessation to greatly reduce the serious risks that smoking poses to their health” (75 FR 69524 at 69526). Therefore, the warnings are intended to have an impact on nonsmokers as well as smokers, and the required warnings will effectively communicate the negative health consequences of smoking to both of these important audiences.

(Comment 14) Several comments, including comments from cigarette manufacturers and individual consumers, expressed concerns that the new required warnings on cigarette packages and advertisements would cause people not to look at packages or cause them to hold their cigarettes in decorative cases. The comments also indicated that some of the proposed images would induce youth to purchase cigarettes rather than deter them from smoking, because the new images would be striking to youth. These comments stated that this “rebound effect” would undermine the intent of the warnings and decrease their effectiveness.

(Response) We disagree. Comments expressing concerns about a potential rebound effect did not provide persuasive scientific evidence to demonstrate such an effect is likely to occur (or that it would have sufficient magnitude to be a significant concern). The comments referenced older studies that did not specifically address graphic warnings on cigarette packages and advertisements, and also referred to a qualitative study conducted on the European Union’s graphic warnings, in which some focus group participants commented that some warnings were humorous or that they were not persuasive in educating consumers about dental diseases associated with smoking (Ref. 42). When weighing this qualitative information against the quantitative research available, including evidence from countries with graphic health warning requirements, as well as the findings of the expert panel of the IOM in its 2007 report (see Ref. 3), the information referenced in the comments is not persuasive. (While focus groups can provide useful information, it is well known that they are not as reliable as real-world evidence for drawing conclusions about causal relationships and generalizing results to the population as a whole (Ref. 43).)

Furthermore, we note that in the European Union qualitative study referenced in the comments, the researchers concluded that pictures have the potential for adding a powerful element to health warning messages and that the old text-only messages were not working (Ref. 42 at p. 43). They also noted that some of the warning messages the comments referred to, including the referenced dental disease image, provoked a highly emotional response in all the countries surveyed despite the comments from certain focus group participants (Id. at p. 35). The research literature suggests that images that evoke emotional responses can increase the likelihood smokers will reduce their smoking, make an attempt to quit, or quit altogether (Ref. 44). While one comment said that the failure of fear-inducing messages based on health effects is “well-known in areas outside of smoking prevention,” the comment did not provide sufficient evidence of such failure in the area of smoking prevention. In fact, as some comments discussed, there is scientific evidence relating to cigarette graphic health warnings illustrating the success of fear-inducing messages (see, e.g., Ref. 44). For example, one comment referred to research that found that smokers exposed to Canada’s graphic health warnings generally did not try to avoid the fear-inducing messages, and that any avoidance engaged in by smokers does not appear to undermine quitting intentions or attempts (citing Ref. 45). Similarly, researchers analyzing data related to graphic warnings found that:

[T]here is no evidence that pictorial warnings lead to boomerang effects. An analysis of data from the ITC Four Country Survey found that the Australian pictorial warnings, introduced in 2005, led to greater avoidant behaviours (e.g. covering up the pack, keeping it out of sight, or avoiding particular labels), compared to Canada, the United Kingdom, and the USA. Importantly, those smokers who engaged in avoidant behaviours were no less likely to intend to quit or to attempt to quit replicating the findings of a study of the Canadian warnings. Thus, although pictorial warnings can lead to avoidance and defensive reactions, such reactions are actually indicators of positive impact.

(Ref. 46. citing Refs. 20 and 44). To the extent that smokers engage in any defensive avoidance with respect to the new required warnings, we are adding a reference to a cessation resource to give smokers an immediate way to act upon this impulse and access cessation assistance. The research literature suggests that such a reference is effective in diminishing potential avoidance effects in response to messages that arouse fear (see Ref. 40 at pp. 39–41). See section V.B.6 of this document for additional information regarding our rationale and authority for including a reference to a cessation resource in the required warnings.

(Comment 15) Several expressed concern about the potential
effectiveness of the new required warnings, particularly those that are fear-based, with certain portions of the population. These comments expressed the following concerns: (1) Many youths and young adults are rebellious and will be attracted to what they perceive as the “forbidden fruit;” (2) fear-based warnings fail with groups that have low self-esteem; (3) fear-based warnings fail with adolescents, because they tend not to be influenced by health-based deterrents; and (4) the new required warnings are “high fear messages” that may actually inhibit reductions in smoking, because they decrease a person’s perceived ability to quit smoking. These comments expressed the belief that the new required warnings would be ineffective.

(Response) While acknowledging the concerns, we disagree. It is true that messages that induce fear, pointing to a risk, may not be effective when people are unaware of how to reduce the risk, but in this case, the best way to reduce the risk is clear. We have chosen a balanced set of images, including those that may arouse fear and those that may generate other emotional responses in certain individuals in order to reach a diverse population of smokers and nonsmokers, as well as youth, young adults, and adults. Furthermore, as is explained in more detail in section III.B of this document, we conducted a research study to quantitatively evaluate the relative efficacy of the proposed required warnings in communicating the health harms of smoking to adults (aged 25 or older), young adults (aged 18 to 24), and youth (aged 13 to 17). The nine selected required warnings showed positive effects on important study measures in all study populations, including youth, relative to the text-only control. In particular, as is discussed in more detail in section III of this document, the selected required warnings showed strong impacts on the salience measures in our research study, including emotional and cognitive measures.

The research literature suggests that these measures are likely to be related to behavior change. For example, the literature suggests that risk information is most readily communicated by messages that arouse emotional reactions (see Ref. 45), and that smokers who report greater negative emotional reactions in response to cigarette warnings are significantly more likely to have read and thought about the warnings and more likely to reduce the amount they smoke and to quit or make an attempt to quit (Ref. 44). The research literature also suggests that warnings that generate an immediate emotional response from viewers can confer negative feelings about smoking and undermine the appeal and attractiveness of smoking (Ref. 45 and Ref. 40 at pp. 37–38). In addition, research has shown that younger adolescents are more likely to notice and think about health warnings that include graphic images (Ref. 47).

The required warnings will effectively communicate the negative health consequences of smoking, and we do not agree that they will have unintended negative effects among younger population groups.

(Comment 16) One comment expressed concern that the new graphic images on cigarette packages and advertisements would actually make cigarette smokers sicker, as the images would increase smokers’ anxiety and damage their self-esteem. (Response) We disagree. We are not aware of any scientific evidence to support this claim. In fact, as discussed in the preamble to the proposed rule, the available research suggests that graphic health warnings can benefit the public health by increasing smokers’ intentions to quit and reducing the likelihood of initiation by nonsmokers (75 FR 69524 at 69532).

(Comment 17) A few comments stated that fear-based warnings fail to work when the message being conveyed is already clearly understood and does not provide new information. These comments expressed the view that, because consumers already understand the risks associated with smoking, the new required warnings would not be effective in achieving FDA’s goals. (Response) We disagree. As explained in section II.C of this document, there is substantial evidence demonstrating that the premise of these comments is not correct and that many consumers do not adequately understand the personal risks associated with smoking.

E. Need To Refresh Required Warnings

As amended by the Tobacco Control Act, FCLAA includes provisions that can help prevent or delay the wear out of the new required warnings. For example, section 4(c)(1) of FCLAA (15 U.S.C. 1333(c)(1)) indicates that the required warnings on cigarette packages must be randomly displayed in each 12-month period, in as equal a number of times as is possible on each brand of the product, and be randomly distributed throughout the United States, in accordance with a warning plan approved by FDA. Section 4(c)(2) of FCLAA requires the warnings to be rotated quarterly in cigarette advertisements, also in accordance with a warning plan approved by FDA.

Nevertheless, as stated in the NPRM, we intend to monitor the effects of the new required warnings once they are put into use. We will conduct research and keep abreast of scientific developments regarding the efficacy of various required warnings and the types and elements of various warnings that improve efficacy. As stated in the NPRM, we will use the results of our monitoring and such research to help determine whether any of the textual warning statements or accompanying graphic images should be revised in a future rulemaking (75 FR 69524 at 69534). This commitment to continued empirical testing is consistent with Executive Order 13563, section 1, which states that our regulatory system “must measure, and seek to improve, the actual results of regulatory requirements.”

FDA received numerous comments regarding the need periodically to refresh the warnings to minimize wear out, which we have summarized and responded to in the following paragraphs.

(Comment 18) Many comments, including comments from health institutions, nonprofit organizations, and academics, suggested that FDA should refresh the graphic warnings on a regular basis because consumers can become habituated to and ignore warnings. The comments referred to scientific research on the effectiveness of graphic warnings for cigarette packages and advertisements, which strongly recommends that warnings be periodically refreshed to maintain their effectiveness and impact on consumers (Refs. 18, 42, 44, and 26). The comments suggested a wide range of timeframes as to when FDA should refresh the graphic warnings. One comment suggested that FDA track the effectiveness of the required warnings on a quarterly basis and that the results of any testing be made publicly available. One comment suggested that FDA establish a conclusion that new graphic warnings for cigarette packages and advertisements will be required at no more than a 2-year interval. A few comments also suggested that FDA establish a target schedule for reconsideration and revision of the warnings, which would include ongoing consumer research and re-examination of the effectiveness of the required warnings.

(Response) We agree that refreshing the required warnings on a periodic basis can help maintain their effectiveness. Researchers have found that graphic images and text messages are likely to have greater impact at the time they are introduced and that
In all, we proposed 36 potential section 201 of the Tobacco Control Act. These 36 proposed required warnings were made available as electronic files in portable document format (.pdf) and displayed in the document entitled “Proposed Required Warning Images,” which was included in the docket for the proposed rule. The proposed required warnings were also made available on FDA’s Web site. Consistent with section 4 of FCLAA, 2 versions of each of the 36 proposed required warnings were developed: one with the textual warning statement in black font on a white background, and one with the textual warning statement in white font on a black background.

As explained in the preamble to the proposed rule (75 FR 69524 at 69534), in considering and developing appropriate color graphic images to accompany the nine textual warning statements set forth in section 201 of the Tobacco Control Act, FDA assessed the graphic warnings that other countries have required, and worked with various experts in the fields of health communications, marketing research, graphic design, and advertising to develop 36 proposed required warnings. Each of the proposed color graphic images depicted the negative health consequences of smoking, and the themes and subjects depicted in each image illustrated the message conveyed by the accompanying textual warning statement.

The NPRM explained that we planned to select 9 final required warnings from among the 36 proposed required warnings. We sought comments on what color graphic images to require in this final rule, including comments on the 36 proposed color graphic images included with the NPRM.

In addition, as described in more detail in section III.B of this document, we conducted research on the 36 proposed required warnings to evaluate the relative effectiveness of the proposed color graphic images and their accompanying textual warning statements at conveying information about various health risks of smoking, and additionally, at encouraging smoking cessation and discouraging smoking initiation.

In order to determine which color graphic images to require in the final rule, we considered a number of factors. First, we considered the relative effectiveness of the proposed required warnings based on the strength of effect the different color graphic images had across the various populations included in our study (see section III.B of this document for more detailed description of the research study).

In addition, we considered the substantive public comments received in the docket related to the 36 proposed required warnings (see section III.C of this document for more information on the comments received; the comments relating to each image are summarized and responded to in sections III.D and III.E of this document). We also considered the comments received in the docket that suggested that we use other images in the required warnings, including images that have been used in other countries’ graphic health warnings. However, as discussed in more detail in the following comment summaries and in section III.B of this document, we selected images for the nine required warnings from among the images we developed and proposed. Our research study, among other information, indicated these required warnings will effectively communicate the negative health consequences of smoking to a wide range of population groups. As explained in the comment responses throughout this section III, the comments submitted to the docket did not persuade us that other images, including images used in other countries’ graphic health warnings, were more appropriate for use in the required warnings than the images we selected.

Furthermore, we considered the relevant scientific literature in the docket, and in particular the extent to which the literature supported or refuted aspects of the images and the extent to which the literature helped determine the appropriate weight to give to other information (including the appropriate weight to give to the various endpoints considered in our research study).

We also considered the variety and diversity reflected in the images in making selection decisions in order to ensure that the final set of required warnings effectively communicates risk information to a diverse range of audiences, including audiences that have been targeted by tobacco industry marketing efforts. We took into account the importance of selecting a set of required warnings that includes a diversity of styles (e.g., photographic versus illustrative), themes, and human images (e.g., race, gender, age). This is consistent with the evidence base for graphic health warnings from countries that have already implemented such warnings, which indicates that variety is important in enhancing the noticability and salience of warnings and broadening their relevance for target groups (Ref. 40 at p. 46 and Ref. 48 at

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Section 202(b) of the Tobacco Control Act amends section 4 of FCLAA (15 U.S.C. 1333) to add a new subsection (d), “Change in Required Statements.” However, section 201 of the Tobacco Control Act also amends section 4 of FCLAA to add a new subsection (d), “Graphic Label Statements.”
p. 9), and which suggests that warnings that include pictures of people should broadly represent the ethnic/racial profile of the relevant country (Ref. 11).

We also considered whether to have one image accompany each of the textual warning statements set forth in section 201 of the Tobacco Control Act.

We received multiple comments regarding our proposal to select 9 final required warnings and our proposal to select them from among the 36 proposed color graphic images that were made available with the NPRM. We have summarized and responded to these comments in the following paragraphs (we also received a number of comments on the proposed color graphic images themselves; these comments are summarized in sections III.D and III.E of this document. In addition, we received a number of comments regarding our research study, which assessed the relative effectiveness of the 36 proposed color graphic images; these comments are summarized in section III.F of this document).

(Comment 19) Several comments suggested that FDA select more than one graphic image for each new textual warning statement. The comments reasoned that by limiting the warnings to one graphic image per textual statement, the health warnings would effectively communicate to fewer segments of the smoking and nonsmoking populations. Some comments also suggested that selecting more than one image per warning statement would counteract wear out of the required warnings. One comment suggested that FDA develop multiple series of images and require that each series be used one at a time to delay wear out.

(Response) We decline to select more than one image for each warning statement as suggested in these comments. We believe that the set of nine required warnings we selected will be sufficient at this time to achieve our goal of effectively communicating the negative health consequences of smoking and to prevent wear out of the required warnings for several years. Furthermore, the nine selected required warnings will appeal to a diverse range of audiences, and, as discussed in section III.D of this document, the images selected showed significant effects on important measures in our research study across the three study populations (adults, young adults, and youth).

We intend to monitor the effects of these required warnings once they are put in use. We will conduct research and keep abreast of scientific developments regarding the efficacy of various required warnings and the types and elements of various warnings that improve efficacy. Given the significant changes being made to the text, format, and placement of the existing warnings by this rule, it will be valuable to obtain relevant data on the effects of the complete set of required warnings as soon as possible. If we were to expand the number of required warnings, it could delay an assessment of efficacy of the warnings under conditions of real-world use. We intend to use the results of our monitoring and of research conducted on the required warnings once they are in public use to determine whether changes should be made to the required warnings in a future rulemaking, including changes to add new images or to modify the existing required warnings. Accordingly, at this time we decline to select more than nine images.

(Comment 20) Multiple comments suggested that FDA use graphic warning images that have been tested or used in other countries instead of or in addition to one or more of the images that FDA proposed. Some of these comments indicated that images that are in use in other countries would be more effective and educational than some or all of FDA’s proposed images.

(Response) We decline to follow this suggestion. FDA’s research study evaluated the 36 proposed required warnings. The results from this research study suggest that the nine selected required warnings will effectively communicate negative health consequences of smoking to a diverse range of audiences. Moreover, if we were to select images that were not evaluated in our study, it would be difficult to objectively assess the relative efficacy of such images compared to the 36 proposed images. Compared to the information provided by our research study, the supporting information in the comments did not convince us that the images suggested by those comments would more effectively communicate the negative health consequences of smoking than the images we have selected in this final rule.

(Comment 21) A number of comments suggested that FDA use other images than those published with the proposed rule. For example, some comments recommended that FDA use images that depict real people with real diseases and not models. A few recommended that FDA include images that show negative cosmetic effects of smoking, such as stained fingers and bad breath, in order to impact adolescents concerning the image. One comment suggested that FDA portray a picture of an arbitrary, while another recommended the use of an image depicting the amount of money smokers spend to purchase cigarettes every year.

(Response) We decline to select the images suggested in these comments. Each of the required warnings selected by FDA was quantitatively tested to assess its relative effectiveness in communicating the negative health consequences of smoking. In selecting the set of nine required warnings, we considered the results of our research study and a number of other factors and have concluded that the nine selected required warnings effectively communicate the negative health consequences of smoking. In addition, we are adopting the nine textual warning statements mandated by Congress in section 4(a)(1) of FCLAA. The images selected were designed to correlate with those warning statements; the available evidence base highlights the value of the text and images in graphic health warnings relating to one another in a meaningful way (see Ref. 40 at p. 41). Including images inconsistent with the textual warning statements could confuse consumers and detract from the effectiveness of the warnings. Furthermore, some of our selected images do show the negative cosmetic effects that can occur as a result of the health consequences of smoking. Moreover, some of the images proposed for use in the comments, such as an image showing the amount of money smokers spend to purchase cigarettes, would not be consistent with the statutory requirement that the required warnings depict the negative health consequences of smoking.

B. FDA’s Research Study

As explained in the NPRM (75 FR 69524 at 69535), we conducted research on the 36 proposed required warnings. Specifically, we conducted an Internet-based consumer research study with over 18,000 participants that quantitatively examined the relative efficacy of the 36 proposed color graphic images in communicating the harms of smoking to 3 target groups: Adult smokers (age 25 or older), young adult smokers (aged 18 to 24), and youth (aged 13 to 17) who currently smoke or who are susceptible to smoking.

The purpose of the study was to: (1) Measure consumer attitudes, beliefs, and intended behaviors related to cigarette smoking in response to the proposed color graphic images and their accompanying textual statements; (2) determine whether consumer responses to the proposed color graphic images and their accompanying textual statements differed across various groups based on age, smoking status, or...
other demographic variables; and (3) evaluate the relative effectiveness of the proposed color graphic images and their accompanying textual warnings statements at conveying information about various health risks of smoking, and additionally, at encouraging smoking cessation and discouraging smoking initiation.

We placed a report (Ref. 49; see also Ref. 50) that described the research study and presented the results of the analyses from the research study in the docket for the proposed rule and announced the report's availability by a notice in the Federal Register on December 7, 2010 (see 75 FR 75936 at 75936 through 75937) so that the public had an opportunity to comment on the results.

This section briefly describes the design of FDA’s research study and key endpoints examined in the research study; a full description of the study and the several hundred pages of data and data analyses are contained in the study report and accompanying appendices (Ref. 49) that was placed in the docket for the proposed rule. This section also describes how the results from this research study informed the selection of the final required warnings.

FDA received numerous comments in the docket related to the research study; this section also includes a summary of the substantive comments received about the research study and FDA’s responses to these comments.

1. Study Design

FDA’s research study evaluated the required warnings proposed for each of the nine warning statements against a text-only control (which contained the warning statement without any accompanying color graphic image). Study participants were randomly assigned to be exposed to either one of the 36 proposed required warnings (treatment groups) or one of the nine textual warning statements (control groups). Treatment groups for each target population (adults, young adults, and youth) viewed a hypothetical pack of cigarettes that included one of the proposed required warnings, which appeared on the upper 50 percent of the pack, while the control group viewed a hypothetical pack of cigarettes with a warning statement (but no warning image), which appeared on the side of the pack. Furthermore, among adults, an additional treatment group viewed a hypothetical advertisement with a warning statement in the same location (but without a warning image) that was presented using the size and format currently required by FCLAA. The study tested the relative efficacy of each proposed required warning relative to the text-only control for that warning statement for the various outcomes measured.

Each respondent viewed either a single cigarette package or advertisement that displayed one of the proposed required warnings or a text-only warning. Respondents answered questions about their immediate reactions to the cigarette package or advertisement, related attitudes and beliefs about smoking, as well as intentions to quit or start smoking. At the end of the survey, subjects were asked to recall which warning statement and image they viewed earlier in the survey to assess the accuracy of recall. In addition, 1 week after completing the survey, subjects were re-contacted and asked to recall the warning statement and image to which they were exposed. Overall, the following key outcomes were measured after exposure to one of the required warnings or the text-only control, and/or at 1 week follow-up:

- **Salience**—The study examined emotional and cognitive responses to the cigarette packages and advertisements that bore health warnings. Participants provided ratings of their responses to the packages and advertisements. The ratings were aggregated to create two scales: (A) An emotional reaction scale, which included ratings on how the warning made the respondent feel, such as “depressed,” “discouraged,” and “afraid”; and (B) a cognitive reaction scale, which included ratings on what the respondent thought about the warning, such as “believable,” “meaningful,” and “convincing”.3

2 While the numerical results reported in the study report (Ref. 49) were correct, and while all of the results discussed in this rule are accurately described, some of the descriptors contained in the study report were in error. An errata sheet for the study report (Ref. 50), which lists all the errors and the corrections, has been prepared and is being placed in the docket. These errors did not adversely impact commenters’ ability to convey their assessment of the images and the study results in their comments. To the extent some comments included inaccurate statements about the study results in their significant comments as a result of the errors, we recognized the inaccuracy and were able to discern the material points in the comment and evaluate them appropriately, as is reflected in the comment summaries and responses.

3 Some additional cognitive measures, including the reaction item “the pack was difficult to look at” (or, for the adult sample viewing the print ad, “the ad was difficult to look at”) were also evaluated but were not reported as part of the composite cognitive reaction scale. These items were not sufficiently correlated with the other cognitive measures to include in the composite measure.
changes in beliefs and intentions and ultimately to behavior change.

2. Use of FDA’s Research Study Results in Selection of Images

As described in section III.A of this document, in order to determine which color graphic images to require in the final rule, we considered a number of factors, including the results from our research study. We carefully examined the research results for the 36 proposed required warnings on all the different outcomes in determining which images to require in this final rule. However, the responses on the salience measures served as a primary basis for distinguishing among the 36 proposed required warnings for a number of reasons.

First, many of the proposed required warnings elicited significant impacts on the salience measures (emotional and cognitive measures), which the research literature suggests are likely to be related to behavior change (Ref. 51). For example, the literature suggests that risk information is most readily communicated by messages that arouse emotional reactions (see Ref. 45), and that smokers who report greater negative emotional reactions in response to cigarette warnings are significantly more likely to have read and thought about the warnings and more likely to reduce the amount they smoke and to quit or make an attempt to quit (Ref. 44). The research literature also suggests that warnings that generate an immediate emotional response from viewers can result in viewers attaching a negative affect to smoking (i.e., feel bad about smoking), thus undermining the appeal and attractiveness of smoking (Ref. 45 and Ref. 40 at pp. 37–38).

In comparison to the salience measures, fewer of the proposed required warnings elicited significant impacts on the beliefs measures in our research study, and on the whole the proposed required warnings did not elicit strong responses on the intentions measures. Given the design of our research study, where participants had only a single exposure to one proposed required warning, it is not surprising that the proposed required warnings did not consistently show effects on these beliefs and intentions measures, which are more eventual outcomes in the behavior change process than the salience responses, which occur more immediately. However, this does limit the utility of these longer-term measures in discriminating across the proposed required warnings. Thus, given the design of the study, the results on the salience measures, which the research literature indicates are predictors of more eventual behavioral outcomes, were considered to be more meaningful than the results on the beliefs and intentions measures in discriminating between the images.

In addition, we gave greater weight to outcomes on the salience measures than to outcomes on the statement recall measures for several reasons. First, there is evidence to suggest that, while recall of associated warning message statements may be reduced in the short term by moderately or highly graphic pictorial warnings versus text-only controls or less graphic pictorial warnings, these warnings still increase intentions to quit through evoked emotional responses (Ref. 52). Second, as described previously, participants in the research study were exposed to a single viewing of the proposed required warnings, which does not allow for assessment of the effect that repetitive viewing of the required warnings may have on recall. Recall can be expected to increase in real world settings where consumers will be exposed to the warnings multiple times. Third, recall was generally high for all the proposed required warnings, even where there was not a significant difference compared to the text-only control or where recall was significantly lower for the proposed required warning than the text-only control. For example, for the nine required warnings that we selected for use in this final rule, the research study shows that recall of both the textual warning statements and the color graphic images was high at both baseline and at 1-week follow-up, exceeding 50 percent on all measures, and, in many cases, exceeding 80 percent.

3. Comments on FDA’s Research Study

FDA received a number of comments related to its research study in the docket for the proposed rule, which are summarized and responded to in the following paragraphs.

a. Study design. Several comments addressed the cross-sectional design of the study.

(Comment 22) Several comments, including comments from cancer researchers, nonprofit organizations, and academics noted that participants in the study were exposed to a proposed required warning only once in a controlled environment. These comments stated that the single exposure study design makes it impossible to assess long term or actual effects of the proposed required warnings. We responded that comments recommended that FDA conduct longitudinal research or post-market surveillance to assess actual long-term effects.

(Comment 23) A comment from tobacco product manufacturers stated that a longitudinal study demonstrating that the required warnings would have actual effects on smoking prevalence was necessary to support the final regulation.

(Comment 24) A comment from cancer researchers, nonprofit organizations, and academics noted that participants in the study were exposed to a proposed required warning only once in a controlled environment. These comments stated that the single exposure study design makes it impossible to assess long term or actual effects of the proposed required warnings. We responded that comments recommended that FDA conduct longitudinal research or post-market surveillance to assess actual long-term effects.
(Comment 24) Several comments discussed behavioral models similar to that described in FDA’s research study (see Ref. 49) and explained how those models provide a rationale for how health warnings can effectively communicate risk information about the harmful effects of tobacco use. For example, one comment from a researcher working on an international project to evaluate the impact of graphic health warnings for tobacco products stated that the primary objectives of health warnings are to educate and inform smokers and nonsmokers about the many negative health consequences of smoking and to provide information that can enhance their efficacy for quitting. The comment noted that effective health warnings increase knowledge and thoughts about the harms of cigarettes, the extent to which the smoker could personally experience a smoking-related disease, and as a result, increase motivation to quit smoking. Another academic who also is conducting research on graphic health warnings commented that a wide variety of research suggests that health warnings with pictures are significantly more likely to draw attention, result in greater information processing, and improve memory for warnings than text-only warnings. A comment from a researcher with expertise in risk perceptions and decisionmaking stated that changes in smoking behavior based on warning labels appear to require four steps: (1) Immediate, negative affective reactions to the potential consequences of smoking; (2) associations of these emotional reactions to smoking cues; (3) increases in perceptions of the risks of smoking, and finally (4) increases in quit contemplation and reductions in smoking behaviors.

(Comment 25) Several comments presented a detailed discussion of the scientific literature to substantiate our conclusion that graphic health warnings can effectively communicate the negative health consequences of cigarette use to smokers and nonsmokers, which is critical given the seriousness of these consequences. Greater understanding of those health effects will motivate some smokers to stop smoking and prevent some nonsmokers from starting to smoke. The preamble to the proposed rule presented a detailed discussion of the scientific literature to substantiate our conclusion that graphic health warnings can be an effective means of communicating important health information about the risks of smoking (see 75 FR 69524 at 69531 through 69533). These comments provide additional support for that conclusion.

b. Study results. Several comments discussed the results from FDA’s research study.

(Comment 25) Several comments, including comments from academics, nonprofit organizations, and health professional organizations, stated that FDA’s research study provides data consistent with the overall literature demonstrating the effectiveness of graphic health warnings. For example, one comment stated that in general the study results are consistent with prior findings that the addition of graphic images to health warnings is beneficial in comparison to text-only warnings. Another comment stated that, based upon the FDA study and the existing scientific literature, it is possible to conclude that the proposed graphic warnings are likely to be effective.

Other comments, including comments from tobacco product manufacturers, advertising industry associations, and a public policy organization, asserted that FDA’s research study fails to provide evidence of efficacy. These comments stated that the study did not show evidence that the proposed required warnings would actually affect prevalence of smoking, and failed to demonstrate sufficient evidence that the proposed required warnings would significantly affect consumer knowledge of the risks of smoking or actual behavior change.

(Response) We agree that the study is generally consistent with the existing scientific evidence demonstrating that graphic health warnings can effectively communicate the negative consequences of cigarette smoking, and by doing so, can encourage smoking cessation and discourage smoking initiation. We disagree that the study results do not support the efficacy of the warnings. We presented substantial research in the preamble to the proposed rule supporting the efficacy of graphic health warnings (75 FR 69524 at 69531 through 69533), and our research study is consistent with that research.

c. Study outcome measures.

Numerous comments discussed the key outcomes measured in FDA’s research study.

(Comment 26) FDA received a wide variety of comments concerning the use of emotional reactions to assess the relative effectiveness of the proposed graphic warnings. A number of comments, including those from academics, medical institutions, and public health groups, supported the inclusion of emotional reaction measures. These comments stated that graphic health warnings that elicit strong emotional reactions, especially negative feelings, are more effective in communicating the negative health consequences of smoking and in motivating healthier behaviors than warnings that do not elicit emotional reactions, and indicate that these effects are well established in the scientific literature.

For example, one comment stated that the scientific literature shows that graphic depictions of the negative health effects of smoking arouse reasonable fears and are associated with greater consideration of health risks, increases in motivations to quit, and ultimately with attempts at cessation. Another comment stated that theoretical models and studies in communications and social psychology suggest that graphic health warnings can be effective because they elicit greater emotional engagement with the information provided and it is that engagement that drives behavior change. Another comment from an academic researcher stated that considerable psychological research suggests that risk is more readily communicated by information that arouses emotional associations with the activity. Emotional reactions can be readily accessed from memory by mere presentation of the stimulus, and appear to be powerful predictors of smoking behavior. Yet another comment stated that growing evidence from controlled experiments and survey research indicates that, compared to text-only warnings, graphic health warnings evoke stronger emotional reactions and increase motivations to quit or not start smoking. The comment indicated that these studies are consistent with cognition and neuroscience research demonstrating that relative to linguistic or text information, imagery-based information can be processed more rapidly, evoke stronger emotional responses, induce greater cognitive processing and attitude change and can be recalled more easily.

However, other comments stated that reliance on emotional measures for assessing graphic health warnings is inappropriate. A joint comment from tobacco product manufacturers stated that the study measured only the effect of eliciting strong emotional and cognitive reactions, which confirms that the warnings were intended not to inform consumers with purely factual and uncontroversial information, but rather to shock consumers into adopting the Government’s preferred course of conduct. Another tobacco product manufacturer commented that, to the extent FDA selected images based on emotional or cognitive reactions and not
on ability to inform consumers about the health risks of smoking, the regulations would not pass constitutional muster. A comment from a public policy organization commented that emotional and cognitive responses are irrelevant measures of effectiveness if there is no behavior response.

(Response) On the basis of our review of the relevant scientific literature and the feedback received in the docket, we conclude that our inclusion of emotional reaction measures to evaluate the relative effects of the 36 proposed required warnings was appropriate and is consistent with well-established principles in the scientific literature. As discussed in the study report that was placed in the docket (Ref. 49) and in other comments summarized in previously in this document, eliciting strong emotional and cognitive reactions to graphic warnings enhances recall and information processing, which helps to ensure that the warning is better processed, understood, and remembered. Thus, these responses can enhance the effective communication of the health warning message. These responses in turn influence short-term outcomes, such as later recall of the message and changes in knowledge, attitudes, and beliefs related to the dangers of tobacco use and exposure to secondhand smoke. As attitudes and beliefs change, they eventually lead to changes in intentions to quit or to start smoking and then later can lead to lower likelihood of smoking initiation and greater likelihood of successful cessation.

We disagree that use of emotional reaction measurements demonstrates the Agency’s intent to advocate a preferred position or course of conduct. Each of the nine graphic warnings required by the final regulations communicates negative health consequences of smoking that are well-established in the scientific literature. Consistent with the Tobacco Control Act, the purpose of these required warnings is to communicate effectively and graphically the very real, scientifically established adverse health consequences of smoking. The overall body of scientific evidence indicates that health warnings that evoke strong emotional responses enhance an individual’s ability to process the warning information, leading to increased knowledge and thoughts about the harms of cigarettes and the extent to which the individual could personally experience a smoking-related disease. Increased knowledge and thoughts about the negative consequences of smoking, in turn, are reasonably likely to result in more informed and healthier behaviors, such as trying to quit smoking or deciding not to start.

(Comment 27) We also received two comments concerning the cognitive measure used in the study. A comment filed by tobacco product manufacturers observed that “looks cool” was one of the measured cognitive reactions. The comment stated that the study analysis omits responses on whether the warnings “looked cool,” and contended that if a substantial number of participants viewed a warning as “looks cool,” the warning would be unlikely to have the intended effect. The comment concluded that the ratings for the “looks cool” measure do not appear to have been neutral; the comment stated that regression results for the “looks cool” measure indicates that this measure elicited one of the strongest estimated effects of the study and the results go in the opposite direction of effectively communicating health risk information. (Response) We disagree that data concerning the “looks cool” outcome was omitted or that the results for this outcome go in the opposite direction of the intended effect of communicating the negative health consequences of smoking. Although the “looks cool” outcome was not included in the reported composite cognitive measure, the study report (Ref. 49) includes the results for this measure in its appendices. The measure was reverse coded, so that a higher value corresponded with the intended directionality for other measures. Thus, a high value for “looks cool” corresponds to a response of “strongly disagree” from the respondent. The data presented in the appendices demonstrate that for each of the nine selected required warnings, significantly more participants disagreed that the warning “looked cool” than participants who viewed the text-only control warning. Eight of the nine required warnings elicited significantly higher ratings than the text-only control warning across all target audiences. Ratings for the ninth required warning, which includes the textual statement “WARNING: Quitting smoking now greatly reduces serious risks to your health,” show that significantly more adults disagreed that the selected required warning “looked cool.” Responses for young adults and youth were in the appropriate direction, but the responses were not significantly different from the text-only control warning.

(Comment 28) We also received a comment concerning the believability measure. This comment raised a concern that some of the 36 proposed required warnings may be perceived as unrealistic because they did not vividly portray immediate health risks, which could lead some smokers to discount the warning. The comment recognized that a believability measure was included in the study as part of the cognitive reaction scale, but stated that specific results for believability were not reported, and recommended that FDA examine the mean scores of the specific believability items in conjunction with other important measures included in the study. (Response) We agree with the comment that believability is a helpful measure for assessing the relative effectiveness of warning images. All of the selected images scored significantly higher than the controls on the cognition measures, which included ratings on how meaningful the warning was, whether it was informative, and whether it was believable. While the results do not include mean scores for believability and other individual measures, the appendices include the parameter estimates from regression analyses on these individual measures. The results show that, in most cases, the images selected for the nine required warnings scored significantly better than the control with respect to believability.

(Comment 29) One comment stated that the statement recall measure is less important and less relevant to decisions about smoking than negative affective reactions because the warning statements are now believed by smokers and nonsmokers. (Response) Statement recall was appropriately included as part of the assessment of the relative effectiveness of the 36 proposed required warnings. As discussed in section II.C of this document, while both smokers and nonsmokers have some understanding about some of the risks of smoking, there are significant gaps in their knowledge, including about the magnitude and severity of the risks of smoking. We also note that, as explained in section III.B.2 of this document, although we carefully examined the research results on all the study measures for the 36 proposed required warnings, including recall, the responses on the salience measures served as a more important basis than recall for distinguishing among the 36 proposed required warnings.

(Comment 30) A joint comment submitted by tobacco product manufacturers asserted that the study fails to demonstrate that the published graphic warnings will have any discernible effects on smoking risk beliefs.
(Response) We disagree with this comment. Four of the nine selected required warnings did show a significant impact on beliefs about the health risks of smoking relative to the text-only control among at least one study population. In addition, there is substantial evidence in the scientific literature showing that graphic health warnings effectively increase consumer understanding of the health risks of smoking. In the preamble to the proposed rule (75 FR 69524 at 69531 through 69533), we presented substantial research showing that graphic health warnings significantly increase consumer thoughts about and understanding of the health risks of smoking after they were introduced in other countries. In addition, as discussed previously in this document, considerable scientific evidence shows that health warnings that elicit strong emotional and cognitive reactions are better processed and more effectively communicate information about the negative health consequences of smoking. Each of the nine required warnings elicited strong effects on the emotional and cognitive reaction scales, which indicates that these warning will effectively communicate information about the negative health consequences of smoking.

Based on the results of our research study and the existing scientific literature, we conclude that graphic health warnings, including the nine selected required warnings, are likely to increase consumer knowledge and understanding of the health risks of smoking.

(Comment 31) A comment submitted by tobacco product manufacturers criticized the study’s use of intentions to measure behavioral change and stated that FDA should have presented data showing actual effects on behavior.

(Response) We disagree that intentions are an inappropriate variable for assessing potential behavioral changes. While measures of intended behavioral outcomes do not perfectly predict a future behavior outcome, it is a necessary precursor. The scientific literature indicates that one’s intentions to quit smoking must be increased before one makes the actual quit attempt. Thus, we conclude that it was appropriate in our research study to assess quit intentions as a proxy for behavior change. In accordance with Executive Order 13563, after the rule is in effect we will be undertaking analysis to better understand the behavioral effects of the warnings.

(Comment 32) Several comments raised concerns that the lack of strong statistically significant results concerning intentions in FDA’s research study is an indication that the required warnings will not be effective. For example, a comment submitted by tobacco product manufacturers stated that the results of FDA’s research study show that graphic health warnings will not result in a statistically significant reduction in youth initiation or overall prevalence of smoking, and thus, confirms that the warnings will not be effective.

(Response) We disagree that our study results indicate that the required warnings will not be effective. It is important to recognize that FDA’s research study was not designed or intended to produce evidence demonstrating actual effects on behavior. Rather, the study was designed to provide data concerning the relative effects of the graphic health warnings in order to provide a more objective and scientific basis for our selection of the set of nine required warnings in the final regulation. There is considerable evidence in the scientific literature demonstrating that graphic health warnings effectively increase awareness of the health risks of smoking, which is the principal purpose of the warnings, and that this awareness in turn can influence smoking intentions and behaviors. We included significant research to substantiate this conclusion in the preamble to the proposed rule (see 75 FR 69524 at 69531 through 69533). For example, as discussed in the proposed rule, a 2007 report from an expert IOM panel that evaluated the existing scientific evidence on health warnings concludes that the available scientific evidence indicates that larger, graphic health warnings would promote greater public understanding of the health risks of using tobacco and would help to reduce consumption (Ref. 3).

FDA’s research study cannot be viewed in isolation from the overall body of scientific evidence evaluating the efficacy of graphic health warnings. While the research study itself did not provide evidence of strong effects on intentions (which, as noted in section III.B.2 of this document, is not surprising given the single-exposure design of the study), the overall body of scientific literature does provide sufficient evidence that the required warnings, by increasing public understanding of and thoughts about the health risks of smoking, will be effective in encouraging smoking cessation and discouraging smoking initiation.

A number of comments provide additional support for our conclusion. For example, a comment from a researcher conducting an international longitudinal study on graphic health warnings states that studies show that graphic depictions of smoking’s adverse effects on the body are associated with greater consideration of health risks, increases in motivations to quit smoking, and ultimately, attempts at cessation. A comment by a researcher with expertise in risk perceptions and decisionmaking concludes that emotional associations to smoking appear to be powerful predictors of smoking behavior and may well be causally implicated in efforts to either stop or start smoking.

(Comment 33) A comment from tobacco product manufacturers stated that the responses to the “smoking urges” questions included in the study would provide a better measure for assessing whether the proposed required warnings affected smoking behavior and, referring to the responses regarding these questions, the comment asserts that, on balance, seeing the proposed required warnings increased the desire to have a cigarette. The “smoking urges” measures were reverse coded, so that a higher value corresponds with the intended directionality for other measures in the study. Thus, a high value corresponds to a response of “strongly disagree” from the respondent. The data presented in the study report appendices (Ref. 49, study report) show that, for three of the nine selected required warnings, significantly more participants in at least one target group disagreed with the statement that they wanted a cigarette than participants exposed to the text-only control warning. For one of the selected required warnings, significantly more adult participants who viewed the warning on a cigarette pack disagreed that they wanted a cigarette, but significantly more adults who viewed the warning in a cigarette advertisement agreed. For one of the selected required warnings, significantly more participants in one target audience agreed that they wanted a cigarette than participants exposed to the text-only control warning. Results for the remaining selected required warnings and sample groups were not significantly different from the text-only control warning. Thus, on balance, the study does not show that exposure to the final set of nine images increased the desire to smoke a cigarette among study participants. As discussed in the previous response, the overall body of
scientific literature provides ample evidence that the required warnings, by increasing public understanding of and thoughts about the health risks of smoking, are likely to encourage smoking cessation and discourage smoking initiation. Data from our research study regarding “smoking urges” provide no basis for calling into question that evidence.

d. Study limitations and issues regarding methodology. A number of comments discussed a wide variety of issues concerning limitations of FDA’s research study and raised various issues concerning the study methodology.

(Comment 34) Several comments, including comments from health institutions, nonprofit organizations, and academics, raised concerns that the demographics of FDA’s research study did not include adequate sample sizes for minority populations and persons of lower income or lower education status. These comments noted that the findings of the study therefore may not be relevant to persons with high smoking prevalence and to those consumers who might be most impacted by graphic health warnings. Some of the comments recommended further testing in these populations.

(Comment 35) A comment from tobacco product manufacturers criticized the study methodology because it did not include a nationally representative sample of participants and claimed that this failure biased the study results. The comment stated that the study report (Ref. 49, study report) fails to disclose basic sampling information and provides no indication that those conducting the study adjusted for the effect of choosing participants by soliciting volunteers. The comment concluded that this failure was significant because the participants in the study may not reflect the population of interest and may bias the statistical estimates.

(Response) We disagree that the study results are invalid due to the demographic composition of the sample. The research study was not intended to be a survey of the national population, but rather a study using random assignment to study conditions. The study included individuals from certain target groups, particularly current smokers and youth who may be susceptible to initiation of smoking. Statistical methods were used to assess the relative impact of each of the proposed required warnings on various outcomes, rather than to assess the absolute impact one would expect to observe in the U.S. population as a whole.

(Comment 36) One comment raised a concern that lack of adequate pretesting of the proposed required warnings evaluated in FDA’s research study could compromise the overall effectiveness of the pool of images tested. The comment stated that it would have been more helpful to conduct pilot testing with a very large group of images (at least 20 per textual warning statement) to ensure testing and selection of the most effective graphic warnings.

(Response) We agree that more extensive pretesting may have been useful. However, we disagree with the suggestion that the overall effectiveness of the required warnings could be compromised by the inability to conduct additional pretesting prior to the research study. The results of the research study as well as research submitted by others during this rulemaking proceeding indicate that the overall efficacy of the pool of proposed required warnings is quite strong. Based on those data, as well as the overall scientific literature, we conclude that the required warnings will effectively communicate the negative health consequences of smoking to smokers and nonsmokers.

(Comment 37) A comment from tobacco product manufacturers asserted that selection bias is a serious methodological flaw of the study. The comment stated that participants were recruited from an Internet panel and offered the opportunity to participate in the research study, creating a selection bias that was compounded by the fact that the invitation to participate stated that the study was funded by FDA. The comment noted that there is no indication that the study corrected for the selection bias and opines that one would not expect the selection bias to be neutral given the identification of FDA as the sponsor of the study.

(Response) We disagree that the study design itself introduced a potential bias. While it is true that biased sampling can affect the results of a study, we believe that the design and implementation of the study were robust. The study used random assignment to study conditions, which helps to control for selection bias. Additionally, the study included a large number of participants, which increases the generalizability of the results. The study results suggest that the overall effectiveness of the proposed required warnings is strong.

(Comment 38) A comment from tobacco product manufacturers stated that FDA’s research study is seriously flawed because 32 percent of the participants dropped out of the study before completing the questionnaire. The comment stated that the survey was not likely to be a random event and may have been a result of smokers who are not receptive to graphic health warnings dropping out. If so, the comment suggested that this would have significantly overstated the results of the study.

(Response) We disagree that the dropout rate observed in the study undermines the validity of the results of the study. Table 3–1 from the methodology report displays the total number of individuals entering the study. However, these values represent the total number of individuals who entered the study’s “landing page,” which is the site to which invitees link from the e-mail invitation. The invitation from e-Rewards, as well as the landing page, refers to the study as a “Study about Consumer Products.” There were no references to FDA, smoking, or tobacco in either the invitation or the landing page. Though it is true that a number of invitees chose not to continue after seeing the invitation or the landing page, their decision not to participate cannot be attributed to a bias for or against FDA or the implementation of graphic health warnings on cigarettes.

In addition, the number of individuals identified as “Quits” in table 3–1 of the methodology report includes individuals who quit after viewing the landing page and those who quit after having been informed of FDA’s involvement and that the survey concerned smoking or tobacco. Of those individuals identified as “Quits,” only a very small number were in the latter category (i.e., quit after being informed of FDA’s involvement and that the survey concerned smoking or tobacco). For
example, of the 13,673 respondents who entered the adult pack survey, 2.179 chose at some point to discontinue. Of these, only 148 individuals, or about 1.1 percent of those entering the study, chose to discontinue the survey after being informed of FDA’s involvement and that the survey concerned smoking or tobacco. A similar pattern exists for all of the study samples: In the adult pack follow-up sample 23 individuals, or 0.6 percent, chose to discontinue after being informed; in the adult ad study sample 193 individuals, or 2.1 percent, chose to discontinue after being informed; in the adult ad follow-up sample 26 individuals, or 0.7 percent, chose to discontinue after being informed; in the young adult study sample 152 individuals, or 1.3 percent, chose to discontinue after being informed; in the youth study sample 104 individuals, or 0.3 percent, chose to discontinue after being informed; and in the youth follow-up sample 13 individuals, or 0.5 percent, chose to discontinue after being informed. The drop-out rate, as calculated here, varies across the study samples but never exceeds 2.1 percent. Therefore, we do not agree that the drop-out rate invalidates the results of the study.

(Comment 39) A comment from tobacco product manufacturers stated that the youth component of FDA’s research study is subject to a response bias. The comment stated that the study failed to address the risk that the youth participants might alter their responses due to a concern that their parents might see the results.

(Response) We disagree that the youth sample is likely subject to a response bias. Youth participants were told at the outset of the study that their responses would be kept confidential. Once the study was complete, other household members could not retrieve those responses. Moreover, if youth participants were concerned about parental awareness of their participation, it would likely have resulted in a decision not to participate rather than a decision to alter their responses.

(Comment 40) A comment from tobacco product manufacturers raised a concern that the youth sample is subject to a selection bias because participants were derived from families whose parents also participated in the study.

(Response) We disagree. As discussed in section 2.2.3 of the methodology report (included in the docket as part of the study report (Ref. 49, study report)), most of the youth were sampled from a separate youth panel, which was independent of the adult panel. Some of the youth were sampled from the households of the adult panel. However, those in the latter group were sampled independently and randomly from the adults that participated in the study. Although possible, it is unlikely that both a parent and child from a single household received an invitation for the study and completed the study.

(Comment 41) A comment from tobacco product manufacturers objects to the manner in which the study assessed emotional and cognitive reactions. The comment states that the study weighted the responses to multiple questions, but fails to disclose the weights used and the justification for those weights, and states that without information on the weighting system, one cannot assess these measures for bias.

(Response) We disagree with this comment. Section 4.2 of the methodology report for our research study (included in the docket as part of the study report (Ref. 49, study report)) indicates that a factor analysis was used to determine the appropriate items to include within each scale. A weighting scheme was used. Rather, items were combined using a simple summative scale. Use of a simple summative scale is a widely-used method of analyzing these data.

(Comment 42) A comment from tobacco product manufacturers states that the study used an inappropriate methodology by measuring risk awareness and smoking intentions on a scale. The comment states that evaluating these measures on a scale is inappropriate for testing awareness of a fact and also resulted in the authors making subjective and undisclosed decisions about how to weight those values.

(Response) We disagree. It is appropriate to measure the impact of a warning on the strength of an individual’s awareness, beliefs, and intentions. To do this, one must use a scaled response, rather than a dichotomous response, to each question. In the research study, items were not weighted within each scale. Rather, they were combined using a simple summation of ratings. This is a widely-used methodology for this type of study.

(Comment 43) A comment attached to the comment from tobacco product manufacturers criticizes FDA’s research study for failing to assess baseline knowledge among participants to determine whether the proposed required warnings increased awareness of the health effects of smoking.

(Response) The lack of an assessment of baseline knowledge does not make the study results less reliable or invalid. In a study such as FDA’s research study, responses to the control conditions serve as proxies for baseline knowledge, awareness, beliefs, and intentions. Comparing the treatment responses to those of the control allow for an assessment of the potential impact the treatment has on baseline measures.

C. Comments to the Docket

FDA received hundreds of comments on the 36 proposed required warnings; the comments relating to each proposed required warning are discussed in sections III.D and III.E of this document. Some comments discussed the 36 proposed required warnings generally or discussed different styles or themes used in the set of proposed required warnings. These comments are summarized and responded to in this section.

As explained in section III.A of this document, we considered the comments submitted to the docket as we determined which color graphic images to require to accompany the nine textual warning statements in the final rule. We did not simply count the number of comments received supporting or opposing the use of a particular image as a way to measure the relative effectiveness of our proposed images or of images recommended by comments, but rather evaluated the substantive input contained in the comments to help inform our decisions in selecting or not selecting a particular image and to obtain other relevant information related to research on the images. Many of the comments contained information about the submitter’s personal opinions, beliefs, and attitudes related to various images. While this information is helpful in understanding how people might interpret various images and in raising issues for further exploration, this type of qualitative information is not as useful as quantitative assessments of the relative effectiveness of the 36 proposed required warnings at conveying information about the negative health consequences of smoking, such as the assessment provided in FDA’s research study.

Furthermore, as described in more detail in the comment summaries and responses in sections III.D and III.E of this document, some of the information contained in comments that criticized or opposed the use of various proposed images suggested through the language of comments negative emotional reactions in the viewer. The research literature,
however, suggests that warnings that evoke these reactions can increase the likelihood smokers will reduce their smoking, make an attempt to quit, or quit altogether (Ref. 44).

1. Comments Submitting Research on FDA’s Proposed Required Warnings

We received several comments, including comments from academics, a nonprofit organization, and a prevention specialist, that described the results of scientific investigations that the submitter had conducted to examine the potential effectiveness of FDA’s proposed required warnings on various outcomes. We address that research and our responses to these comments in the comment summaries and responses in this section. The information contained in these comments about particular proposed required warnings is also discussed as applicable in sections III.D and III.E of this document.

As is discussed in the summaries in this section, the nine required warnings we have selected for use on cigarette packages and in cigarette advertisements generally performed well in the studies discussed in these comments. These comments indicate that the findings from our own research study are robust, as they have generally been confirmed under the various different study designs utilized in the research discussed in these comments.

However, in contrast to our own research study, we did not have access to the raw data or to all the statistical analyses for the studies discussed in these comments. In addition, the design of some of these studies did not allow for an assessment of the relative effectiveness of FDA’s 36 proposed required warnings. This limited the utility of the information provided in the submissions.

Thus, while we carefully considered the information provided in these submissions, the results of our own study were more helpful in making research-based selection choices. (Comment 44) One study was submitted by a group from a medical institution and by a collaborating academic who has conducted research on graphic health warnings. Participants were recruited from an Internet panel of adults, young adults, and youth. The report for the study states that it was intended to assess the potential effectiveness of FDA’s 36 proposed required warnings. Among other things, participants were asked to provide certain demographic information as well as information concerning their smoking status and attitudes and beliefs about smoking. In addition, the study tested nine “sets” of warnings, one for each of the textual warning statements required by the Tobacco Control Act. Each set included each of the proposed required warnings published with the proposed rule for use with the specific textual warning statement as well as at least one alternative warning. Each participant was randomly assigned to view and rate two sets of health warnings.

Warnings within each set were first rated individually on a scale of 1 to 10 and then participants were asked to rank order the entire set for perceived effectiveness for discouraging smoking. The comment presented the rating and ranking scores for the health warnings. The comment also presented preliminary statistical analyses for the overall ranking scores; statistical data were not presented for individual ratings for the individual measures assessed. The comment concludes that preliminary results from the study show that warnings that were more explicit about the health risks of smoking were rated as being more effective among both adults and youth. The academic who conducted the study similarly concluded that health warnings that were more explicit and that elicited greater emotional reactions were rated as being most effective, and the researcher recommended that FDA select certain graphic warnings that received high rating and ranking scores in the study (including required warnings proposed by FDA as well as graphic warnings that have been used in other countries).

(Response) The results of this study are generally consistent with the results of the scientific literature and the study sponsored by FDA. This study shows that the existing cigarette warnings are not salient among either adults or youth. Among other responses, 50.3 percent of adults responded that they never or rarely noticed the health warnings on cigarette packs, while 23.7 percent stated that they often or very often noticed the warnings. Among youth, 63.3 percent responded that they never or rarely noticed the warnings. The graphic warnings selected for inclusion in the final regulation generally performed relatively well in both this study and in FDA’s research study. It is difficult to assess the results of this study more specifically, however, without additional information concerning the study protocol, methods, and statistical analyses.

(Comment 45) A study was submitted by a researcher with expertise in risk perceptions and decisionmaking. Participants were young adult college students, including smokers, nonsmokers, and “vulnerable” nonsmokers. The study assessed emotional reactions, risk perceptions, and smoking aversion. Participants were randomized into four conditions, with each viewing 18 graphic warnings. Two conditions viewed graphic warnings being used in other countries, one condition viewed 18 graphic warnings published with the proposed rule, and the fourth condition viewed the proposed FDA graphic warnings plus three graphic warnings from other jurisdictions. According to the comment, warnings “that were perceived as more graphic, more intense, less good, and more fearful produced more thoughts about not wanting to smoke.” The comment concludes that, compared to the viewed warnings being used in other countries, the FDA proposed required warnings did not maximize thoughts of health risk perceptions or smoking aversion, although the differences between the warnings from other jurisdictions and FDA’s proposed required warnings were marginal.

(Response) The nine required warnings that we have selected performed relatively well in this study. Many performed as well as the warnings from other jurisdictions and some performed better. It is difficult to assess the results of this study more specifically, however, without additional information concerning the study protocol, methods, and statistical analyses.

(Comment 46) A study was submitted by a group of behavioral scientists whose research focuses on cognitive, emotional, and imagery processes that influence how people respond to messages about health risks. Their experimental study evaluated the 36 proposed required warnings published with the proposed rule. Participants were young adults ages 18 to 25, and included smokers and nonsmokers. Each participant viewed 18 of the 36 proposed required warnings and was asked to rate each on the following measures: Perceived comprehension, worry about the health risks of smoking, and the extent to which the warning discouraged the participant from wanting to smoke a cigarette. The comment states that the study provides strong support that most of the graphic warnings proposed by FDA are perceived by young adult smokers as easy to understand, as enhancing worry about the health risks of smoking, and as discouraging young adult smokers from wanting to smoke. The comment states that the results of the study are consistent with the growing body of
evidence showing that, compared to text-only warnings, graphic warnings can evoke stronger emotional responses and reduce motivations to smoke.

(Comment 48) An organization of high school students submitted the results of a study they conducted to assess the efficacy of the 36 proposed required warnings published with the proposed rule. Organization members recruited participants from their high schools and communities. Each participant viewed 18 of the proposed required warnings and was asked to rate each warning for perceived effectiveness in stopping someone from smoking. Findings were reported as arithmetic means and modes. The comment concludes that study respondents generally believed that the most effective images were the more graphic images.

(Comment 47) A study was submitted by two researchers at a university-based public policy center. The comment states that the study, of young adult and adult smokers, was conducted to assess limitations of the FDA study and to identify ways to increase the impact of the warnings. The study used the same online survey firm as that used in the FDA study, although respondents who participated in the FDA study were not eligible to participate in this study. The study was limited to four of the nine warning statements required by the Tobacco Control Act. The graphic warnings assessed for each of these four statements included some of the proposed FDA warnings, these same proposed warnings with additional text or color added, and some graphic warnings used in Canada. Graphic warnings were compared against a text-only control warning that appeared on the side of a cigarette pack. The study used two indices to assess efficacy. The first assessment was perceived effectiveness in discouraging someone from smoking. For the second assessment, participants were asked to imagine themselves smoking a cigarette and then to report how good or bad they would feel smoking a cigarette. The comment states that in three of the four warning messages required by the Tobacco Control Act, a single exposure to a large graphic warning was more effective in creating immediate negative emotional associations with the act of smoking than exposure to the text-only warning. The comment states that the study did not show that the single exposure affected immediate plans to quit smoking; the authors of the comment note that a brief test following a single exposure is unlikely to detect this effect, and that they would expect quit intentions to increase through repeated exposures to the warnings.

(Comment 49) One comment contained the results of a study conducted by two individuals among college students at a U.S. university. In this study, 63 college students, apparently including both smokers and nonsmokers, were shown the 36 proposed required warnings and asked to rate them on a scale of 1 to 7 on their perceived effectiveness in helping smokers’ intent to quit. According to the comment, certain demographic information also was obtained from participants. The comment identifies the five proposed required warnings that were ranked as being the most effective warnings and the five proposed required warnings that were ranked as being the least effective. According to the comment, demographic factors did not affect the rating scores. The only factor identified as having an impact on rating was smoking status, with participants who had a history of smoking more likely to rate the graphic warnings as being effective than subjects who did not have any history of smoking.

In another comment, submitted by a self-identified prevention specialist from a U.S. public school district, 1,339 high school students viewed the 36 proposed required warnings and were asked “which image would change your mind about smoking.” The comment identified the “top three” proposed required warnings.

(Comment 46) A study was conducted to assess whether the FDA’s graphic health warning statements were ranked as being the most effective images in helping smokers consider the health risks of smoking. The study respondents generally believed that graphic health warnings that elicit strong emotional responses help people process the warning information. This leads to increased knowledge and thoughts about the health risks of smoking and the extent to which an individual could personally experience smoking-related disease, which can increase motivation to quit smoking. For example, the literature suggests that risk information is most readily communicated by messages that arouse emotional reactions (see Ref. 45), and that smokers who report greater negative emotional reactions in response to cigarette warnings are significantly more likely to have read and thought about the warnings and more likely to reduce the amount they smoke and to quit or make an attempt to quit (Ref. 37–38).

2. Other Comments

FDA also received a number of other comments that discussed the proposed required warnings generally or highlighted issues that applied to some or all of the proposed required warnings. These comments are summarized and responded to in the following paragraphs.

(Comment 50) Many comments stated that graphic health warnings that elicit strong emotional responses are most effective in communicating the negative health consequences of smoking and in encouraging smoking cessation and discouraging smoking initiation. Most of these comments recommended that FDA select the warnings that evoke the strongest emotional responses. Some of these comments cited graphic warnings used in other countries or international research showing that images that trigger emotional responses promote greater awareness and better recollection of the health risks of smoking. Some of these comments also stated that warnings that trigger these responses retain their effectiveness longer. Some of these comments recommended that FDA select graphic warnings that portray graphically disturbing images or images that evoke fear or disgust.

(Comment 51) We agree that eliciting strong emotional responses helps communicate health information. The overall body of scientific literature indicates that health warnings that evoke strong emotional reactions enhance an individual’s ability to process the warning information. This leads to increased knowledge and thoughts about the health risks of smoking and the extent to which an individual could personally experience a smoking-related disease, which can in turn motivate positive behaviors. For example, the literature suggests that risk information is most readily communicated by messages that arouse emotional reactions (see Ref. 45), and that smokers who report greater negative emotional reactions in response to cigarette warnings are significantly more likely to have read and thought about the warnings and more likely to reduce the amount they smoke and to quit or make an attempt to quit (Ref. 37–38). In FDA’s study, eight of the nine selected required warnings elicited strong emotional reactions across all target audiences. As is further discussed in section III.D of this document, the ninth selected required warning, which, unlike the other eight required warnings, contains a warning statement that is framed in a positive manner, also
showed significant effects on the emotional reaction scale in one study population. Given the manner in which this ninth warning is framed, it is not expected to arouse the same level of response on the emotional reaction scale used in FDA’s research study as the other eight warning messages (see section III.D of this document).

Some of the required warnings we selected include images that may be more emotionally disturbing to certain individuals than others. As we discussed in the preamble to the proposed rule, the use of health warnings with disturbing tonal qualities appears to be effective (75 FR 69524 at 69534). But research also indicates that other types of graphic images, including some that individuals do not find frightening or disturbing, can also be effective in communicating the health risks of smoking (I.d.). The set of nine graphic warnings we selected includes a balanced set of images in order to reach the broadest target audience of smokers and potential smokers. (Comment 52) Some comments raised concerns about the quality of the proposed required warnings published by FDA. Some believed that the proposed required warnings were weaker than those used in other countries, and thus, would be less impactful than those in use in other countries. A few comments said the images were overdone and insulting, and a few indicated that the submitters believed that the visuals were poorly crafted.

(Response) We disagree with these comments. We have chosen a balanced set of images for use with the required warnings, and these warnings are generally consistent with the graphic health warnings used in other countries. The results from our research study and the overall body of scientific literature on graphic warnings provide a strong basis for concluding that the nine selected required warnings will effectively communicate the negative health risks of smoking to smokers and potential smokers.

(Comment 52) Some comments raised concerns that the proposed required warnings were too explicit and too visually disturbing. Some of these comments raised concerns that the images were too disturbing for children to see, and others indicated that nonsmokers should not have to be subjected to “gross” images when they go into retail establishments. Two comments raised concerns that images that showed humans in distress or human remains were disrespectful and degrading. One comment stated that the proposed warnings crossed the line and were an effort to manipulate people to stop smoking or not to start.

(Response) We disagree. The set of nine required warnings we selected include a balanced set of images. Some individuals may find certain images more visually disturbing than others. The images are not intended to shock or disturb, but rather to effectively educate and inform smokers and potential smokers about the serious health consequences of smoking. Each of the nine graphic warnings communicates negative health consequences of smoking that are well-documented in the scientific literature. By appropriately conveying the serious health consequences in a truthful, forthright manner, the images contain information that may disturb some viewers because the severe, life-threatening and sometimes disfiguring health effects of smoking are disturbing. The overall body of scientific evidence indicates that larger, graphic health warnings will effectively communicate these risks. We do not agree that these warnings are disrespectful or degrading.

(Comment 53) A number of comments advocated for the selection of a set of images that could communicate with the diverse U.S. population, and emphasized the importance of human diversity in the images, in part to help ensure the images reach people of low socioeconomic status that are more likely to be smokers and/or to have lower literacy. The comments stated that graphic health warnings are an especially important communication tool for these population groups. A few comments also raised concerns that not enough of the 36 proposed required warnings depicted younger people, and indicated this could reduce their impact among youth.

(Response) We agree that it is important to select a set of images that can communicate with the diverse U.S. population. As discussed in section III.A of this document, we considered the need for diversity when making image selections, and the images selected include a diversity of human images (e.g., race, gender, age), as well as a diversity of styles (e.g., photographic versus illustrative) and themes. This is consistent with the evidence base for graphic health warnings from countries that have already implemented such warnings (see Ref. 40 at p. 46 and Ref. 11).

(Comment 54) A number of comments raised concerns that some of the proposed graphic warnings included graphic illustration or “cartoon-style” images. Some of these comments stated that these warnings might trivialize the serious health risks of smoking or diminish the importance of the warnings, with some asserting that this style is contradictory to the serious messages being conveyed. One comment believed that these warnings would soften the message, while another believed the graphic illustration warnings were “harsh.” Some comments stated that these warnings would negatively affect the believability of the warnings and would not be taken seriously by youth. One comment expressed concern that the graphic illustration style images might resonate with youth, but would not be effective with young adults or adults. It was also noted in the comments that the images presented in this style may inadvertently suggest approval of tobacco use to low-literacy populations that do not comprehend the accompanying textual statement, and that these images could allow smokers to deny the health consequences that are presented. Another comment stated that the research suggests “cartoon-style” images and overly conceptual images are easily dismissed by smokers.

(Response) We disagree with the contention that the use of graphic illustration style images is categorically inappropriate. One of the required warnings we selected is presented in this style. As discussed in section III.B of this document, our research study shows that the selected required warnings, including the required warning that includes a graphic illustration style image, showed strong effects in terms of emotional reaction scale, cognitive reaction scale (including believability), and the “difficult to look at” measure. Given these results, we concluded that the graphic illustration style can be an effective style for communicating the negative health risks of smoking, including to a diverse range of viewers. In addition, it is important to include a variety of different styles in the final set of warnings. As discussed in the preamble to the proposed rule, a varied set of warnings is consistent with the scientific literature, facilitates better targeting of specific groups whose interests may vary, and has been shown to be effective in delaying or counteracting wear out of the warnings (75 FR 69524 at 69534).

(Comment 55) A number of comments advocated that FDA select only required warnings with photographic images. Some of these comments stated that the use of photographic images was important to realistically portray the negative health consequences of smoking and to provide a real-life similarity to the warnings. A comment stated that photographic images were needed to ensure that smokers and
potential smokers understood that the depicted health consequence could really happen and to provide a more physical connection. One comment stated that photographic images would be more engaging and remembered than images presented in other styles. One comment stated that warnings with abstract imagery that require individuals to “connect the dots” and draw inferences present an unnecessary and counterproductive hurdle for viewers, and are unlikely to have an effect on smokers.

(Response) We agree that graphic warnings with photographic images can effectively communicate the negative health consequences of smoking, and most of the required warnings we selected include photographic images. The existing scientific literature, the experience of other countries, and the results of our research study show that graphic warnings using photographic images can effectively communicate the negative health consequences of smoking. At the same time, we do not agree that photographic images are the only style of imagery capable of effectively communicating these health risks. A balanced set of warnings with a variety of image styles is more likely to effectively reach a broad group of target audiences, and we note that graphic warnings used in many other countries include a mix of imagery, including photographic and other styles.

(Comment 56) Some comments stated that graphic warnings will not be effective in deterring smoking. One comment stated that smokers already know the health risks of smoking and are very brand loyal, so graphic images will not affect their smoking decisions. Another comment stated that youth will not be deterred by pictures and the graphic warnings could instead make smoking more enticing to youth. One comment stated that smokers are addicted to cigarettes and “flashy” pictures will not stop them from smoking but instead will only encourage them to cover the pictures. On the other hand, other comments concluded that graphic health warnings are likely to affect smoking decisions. One comment stated that graphic warnings will deter initiation, and another stated that the warnings will lead to a decrease in cigarette sales. One comment stated that graphic warnings will reach people who otherwise would not read text-only warnings.

(Response) As previously discussed, we concluded that large graphic warnings are effective in conveying the health risks associated with smoking, influencing consumer awareness and knowledge of those risks and having an impact on smoking intentions. We disagree with comments stating that required warnings will not be effective. We have determined that the set of required warnings we have selected will effectively convey the negative health consequences of smoking, which will help discourage non-smokers, including children and adolescents, from starting to smoke cigarettes, and help encourage current smokers to consider cessation to greatly reduce the serious risks that smoking poses to their health.

D. Selected Images

This section discusses the nine color graphic images that we selected for use with the textual warning statements set forth in section 201 of the Tobacco Control Act and the factors that influenced each selection decision, including the results from our research study, the substantive comments received in the docket, the relevant scientific literature, and any other considerations weighed, such as the diversity a particular image contributes to the overall set of required warnings.

The document entitled “Proposed Required Warning Images” that was included in the docket for the proposed rule displayed each of the 36 proposed required warnings (consisting of the proposed images and accompanying warning statements) on two consecutive pages, with one display showing the warning statement accompanying the image in black text on a white background and one display showing it in white text on a black background. The images are referred to in this section by the pages on which they appear in the “Proposed Required Warning Images” document and by the descriptive names used for each image in the study report (Ref. 49) summarizing the results of our research study.

In this section’s discussion of the results from our research study for each selected image, the endpoints that the images showed a statistically significant effect on in one or more of the study populations (adult smokers aged 25 or older, young adult smokers aged 18 to 24, and youth who currently smoke or who are susceptible to smoking aged 13 to 17) are described. This discussion also notes the level of significance of the effects by providing p-values: (p<0.05), (p<0.01), and (p<0.001). The p-value is reflective of the percent chance the finding could have happened by coincidence. For example, for a finding that is significant at 0.1 percent (p<0.001), there is less than one chance in a thousand that the finding happened by coincidence. The full description of our research study and the analyses are contained in the study report (Ref. 49, study report) that was placed in the docket for the proposed rule.
The required warnings, consisting of the nine color graphic images we selected and the textual warning statements, are contained in a document titled “Cigarette Required Warnings,” as is further discussed in section V of this document.

1. “WARNING: Cigarettes are Addictive”

We selected the image which appears on pages one and two of the document “Proposed Required Warning Images,” referred to as “hole in throat,” for use with this warning statement.

In our research study, this image had a significant effect (p<0.001) on all salience measures (emotional reaction scale, cognitive reaction scale, and difficult to look at measure) in all three study populations (adults, young adults, and youth). The image had the numerically largest effects of the images proposed for use with this warning statement on the emotional reaction scale and the difficult to look at measure in all three study populations, as well as on the cognitive reaction scale in adults. As discussed in section III.B of this document, these salience impacts are important, as the research literature suggests that they are likely to be related to behavior change.

The image also had a significant impact (p<0.05) on adult 4 beliefs about the health risks of smoking for smokers, and a significant impact (p<0.05) on adult beliefs about the health risks of secondhand smoke exposure for nonsmokers, relative to the text-only control.

However, young adults viewing the image had significantly lower statement recall at one week follow-up than those who viewed the text-only control (55.9 percent versus 74.3 percent), as did adults viewing a hypothetical advertisement containing the proposed required warning (64.1 percent versus 87.7 percent). However, recall of the statement was generally high for the image (ranging from 55.9 percent to 86.3 percent), even where it was significantly lower than for the text-only control, and we conclude that repetitive viewing of the required warning is likely to increase recall. As explained in section III.C of this document, we gave greater weight to outcomes on the salience measures than to outcomes on the recall measures.

We received a number of comments on this image, which we have summarized and responded to in the following paragraphs.

(Comment 59) FDA received a large number of comments supporting the use of the image “hole in throat,” including comments from individuals (including former smokers), public health advocacy groups, academics, State and local public health agencies, and health care professionals. Many comments stated that this image is the best image for use with this warning statement. Some comments indicated that the image was appropriately compelling and effectively communicates the risks of smoking. Other comments stated that the image will be an effective deterrent to smoking by making a smoker think twice before buying cigarettes and/or by making children think twice before starting to smoke. Several comments also indicated that the image concretely conveys the health harms of smoking.

(Response) We selected this image for use with this warning statement.

(Comment 60) One comment supported use of this image in part because of the diversity reflected in the image, and noted that it could be a Latino smoker or a man of color, which could make it more relevant than other proposed images with low socioeconomic status smokers. Another comment noted that the image targets a critical demographic group by portraying an image of a man.

(Response) We agree that it is beneficial to have a diverse set of images that communicates with a wide range of audiences, including population subgroups with higher smoking prevalence rates. In light of this, we selected a set of nine required warnings (including the image “hole in throat,” which portrays a man of color) that includes a variety of human images that are broadly representative of the overall population.

(Comment 61) As mentioned in section III.C of this document, some comments submitted to the docket described the results of scientific investigations that the submitters had conducted to examine the potential effectiveness of FDA’s proposed images on various outcomes. This image was discussed in some of these comments. For example, in one submitter’s study, participants rated this image highly on its ease of comprehension. It also induced relatively greater worry and feelings of discouragement from wanting to smoke than a text-only control. The submitter concluded that this image was the most effective of the images proposed for use with this warning statement. Additionally, this image was one of two images deemed effective in another submitter’s survey of comparative effectiveness of the 36 proposed required warnings at stopping someone from smoking, and it received the highest overall rating of the images examined for use with this statement in another submitter’s study of the potential effectiveness of the images.

(Response) As discussed in section III.C of this document, we carefully considered the comments submitted to the docket that described the results of studies conducted by the submitters on our proposed required warnings. The results summarized in these comments are generally supportive of our image selection decisions.

(Comment 62) FDA also received some comments that opposed the use of the image “hole in throat.” One comment noted that the image was “too gross to be effective,” while another comment stated that it “offends against human dignity.” In addition, one comment stated that the image would only have a one-time shock value, and another comment indicated that the image was too vague in nature.

(Response) We disagree with these comments. The image effectively and concretely communicates the negative health consequences of smoking. The image clearly portrays the addictive nature of cigarettes, depicting a man who is still smoking despite prior evidence (a stoma in his neck) of surgery for cancer. As discussed, this image had a highly significant effect (p<0.001) on all salience measures (emotional reaction scale, cognitive reaction scale, and difficult to look at measure) in all three study populations (adults, young adults, and youth). The research literature indicates that images that evoke emotional reactions can promote greater awareness and better recollection of the health risks of smoking, and can increase the likelihood smokers will reduce their smoking, make an attempt to quit, or quit altogether (Ref. 20, 44, and 45).

Furthermore, contrary to the assertion that the image will only have a one-time shock value, the research literature suggests that more vivid warnings are more likely to retain their salience over time (Ref. 3 at p. C–4 and Ref. 41).

2. “WARNING: Tobacco Smoke Can Harm Your Children”

We selected the image which appears on pages 9 and 10 of the document “Proposed Required Warning Images,” referred to as “smoke approaching baby,” for use with this warning statement.

In our research study, this image had a significant effect (p<0.001) on all the salience measures (emotional reaction scale, cognitive reaction scale, and...
difficult to look at measure) in the adult and youth samples. In young adults, the image also had a significant effect on all the salience measures (emotional reaction scale (p<0.01), cognitive reaction scale (p<0.001), and difficult to look at measure (p<0.05)).

The image had a significant effect (p<0.05) on recall of the warning statement at baseline compared to the control for adults and youth. The image also had a significant effect (p<0.05) on statement recall at 1 week follow-up in young adults. The image also showed some of the largest effect sizes for image recall (at baseline and at 1 week follow-up) in adults and young adults across the images proposed for use with this warning statement.

The image had a statistically significant effect (p<0.05) on youth intentions to not smoke in the next year, with 71.6 percent of youth viewing the image reporting that they would not be likely to smoke in the next year compared to 56.9 percent of youth viewing the text-only control.

As is discussed in further detail in section III.E of this document, three other images proposed for use with this warning statement, “smoke at toddler,” “girl crying,” and “girl in oxygen mask,” also had significant effects on all the salience measures (emotional reaction scale, cognitive reaction scale, and difficult to look at measure) in all three study populations (adults, young adults, and youth). While several of the images proposed for use with this warning statement could effectively convey the negative health consequences of tobacco smoke exposure for nonsmokers (and in particular, children), we ultimately considered “smoke approaching baby” to have the strongest overall research results of the images proposed for use with this warning statement for multiple reasons.

First, two of the images that also showed significant effects on all the salience measures across the study populations, “girl crying” and “girl in oxygen mask,” were negatively associated with beliefs about the health risks of secondhand smoke exposure for nonsmokers in the adult sample. In other words, adults who viewed these images were less likely to believe that nonsmokers will suffer from negative health effects related to secondhand smoke exposure than adults who viewed the text-only control.

As described in section III.B of this document, we determined that the salience results from our research study are the fundamental basis for making distinctions among the images given the design limitations of the research study, which exposed each participant to each image only once, and thus may not be able to accurately distinguish the relative effects of the images on more eventual outcomes, such as changes in beliefs, as reliably as their effects on more immediate emotional and cognitive reactions. However, the negative results observed on the secondhand smoke beliefs measure for the images “girl crying” and “girl in oxygen mask” were of concern, particularly given that the subject of the warning statement is the health risks of secondhand smoke exposure for children. Thus, “smoke approaching baby” was considered a preferable alternative to these two images.

Furthermore, “smoke approaching baby” was associated with youth reporting that they would be less likely to be smoking 1 year from now. We received a number of comments on this image, which we have summarized and responded to in the following paragraphs.

(Comment 65) As mentioned in section III.C of this document, some comments submitted to the docket described the results of scientific investigations that the submitters had conducted to examine the potential effectiveness of FDA’s proposed images on various outcomes. This image was discussed in some of these comments. For example, it was rated highly on its ease of comprehension and induced relatively greater worry and feelings of discouragement from wanting to smoke than a text-only control in one submitter’s study.

(Response) As discussed in section III.L of this document, we carefully considered the comments submitted to the docket that described the results of studies conducted by the submitters on our proposed required warnings. The results summarized in these comments are generally supportive of our image selection decisions.

(Comment 66) FDA also received some comments critical of the image “smoke approaching baby.” These comments suggested that the child does not appear to be suffering harms to his health and/or looks too healthy. One of these comments also stated that the image was associated with youth reporting that they would be more likely to be smoking 1 year from now, and advised against its use.

(Response) We do not agree that the image does not depict the health hazards of secondhand smoke. Graphic depictions of the visible effects of disease are not the only way of communicating the health risks of secondhand smoke for children (see Ref. 11), some of which (such as impaired lung growth), are not necessarily externally visible in a photograph of a child exposed to secondhand smoke. Furthermore, it is important to keep in mind that the image is not used in isolation, but accompanies the textual warning statement, which provides additional context for what is shown. As evidenced by the significant effects the image had on the salience measures compared to the text-only control across the populations participating in FDA’s research study, the required warning depicts the health consequences of
secondhand smoke exposure in a manner that has an impact on both smokers and potential smokers. Thus, we conclude that the required warning effectively conveys the message that exposure to tobacco smoke is harmful for children.

We also note that the comment stating that the image was associated with youth reporting that they would be more likely to be smoking 1 year from now is incorrect. In fact, the image had a statistically significant effect on decreasing youth intentions to smoke (see Ref. 49 at p. 4–4; see also Ref. 50).

As stated previously, 71.6 percent of youth viewing this image reported that they would not be likely to smoke in the next year, compared to 56.9 percent of youth viewing the text-only control.

3. “WARNING: Cigarettes Cause Fatal Lung Disease”

We selected the image which appears on pages 25 and 26 of the document “Proposed Required Warning Images,” referred to as “healthy/diseased lungs,” for use with this warning statement.

In our research study, this image had a significant effect (p<0.001) on all the salience measures (emotional reaction scale, cognitive reaction scale, and difficult to look at measure) in all three study populations (adults, young adults, and youth). The image had the numerically largest effects of the images proposed for use with this warning statement on the salience measures. As discussed in section III.B of this document, these salience impacts are important, as the research literature suggests that they are likely to be related to behavior change.

The image also showed some of the largest effect sizes for image recall (at baseline and at 1 week follow-up) in adults and youth across the images proposed for use with this warning statement.

We received a number of comments on this image, which we have summarized and responded to in the following paragraphs.

(Comment 67) FDA received a large number of comments supporting the use of the image “healthy/diseased lungs,” including comments from individuals, public health advocacy groups, medical organizations, academics, State and local public health agencies, and health care professionals. Many comments indicated that this image is the best image for use with this warning statement, with one stating that the image dramatically depicts a health consequence of smoking, and another noting that it was appropriately gripping and compelling.

Several comments noted that, based on FDA’s research results, this image is the clear choice among the four images proposed by FDA for use with this warning statement. Some comments noted that similar images have been used effectively in other countries that require graphic health warnings on cigarette packages. One comment noted that this image could reach a younger audience, and hopefully prevent them from starting to smoke.

(Response) We selected this image for use with this warning statement.

(Comment 68) As mentioned in section III.C of this document, some comments submitted to the docket described the results of scientific investigations that the submitters had conducted to examine the potential effectiveness of FDA’s proposed images on various outcomes. This image was discussed in some of these comments. For example, in one submitter’s study, participants rated this image highly on its ease of comprehension. It also induced relatively greater worry and feelings of discouragement from wanting to smoke than a text-only control. The submitter concluded that this image was the most effective of the images proposed for use with this warning statement.

Another comment also submitted research suggesting that this image was the highest rated for potential effectiveness among the set of images proposed for use with this warning statement. Another submitter showed that, in a survey, respondents rated this image as one of the most effective of the 36 proposed images for encouraging smokers to quit smoking. The image was also identified in a survey of high school students as one of the “top three” proposed required warnings (out of 36) in another submitter’s study.

(Response) As discussed in section III.C of this document, we carefully considered the comments submitted to the docket that described the results of studies conducted by the submitters on our proposed required warnings. The results summarized in these comments are generally supportive of our image selection decisions.

(Comment 69) FDA also received some comments critical of the image “healthy/diseased lungs.” One comment noted that the image was “too gross to be effective,” while several comments expressed the opposite belief, with some suggesting that the diseased pair of lungs should be more damaged.

(Response) The image “healthy/diseased lungs” is an appropriate image that effectively conveys the negative health consequences of smoking. While, as reflected in the above summary, some comments expressed a belief that the image of the diseased lung is “too gross” and some expressed a belief that the image is too healthy in appearance, the image effectively evoked emotional and cognitive reactions in reviewers in FDA’s research study, which in turn suggests that the image has the potential to promote greater awareness of the health risks of smoking and motivate positive behavioral outcomes, including an increased likelihood that smokers will reduce their smoking, make an attempt to quit, or quit altogether (Refs. 20, 44, and 45).

4. “WARNING: Cigarettes Cause Cancer”

We selected the image which appears on pages 33 and 34 of the document “Proposed Required Warning Images,” referred to as “cancerous lesion on lip,” for use with this warning statement.

In our research study, this image had a significant effect (p<0.001) on all the salience measures (emotional reaction scale, cognitive reaction scale, and difficult to look at measure) in all three study populations (adults, young adults, and youth). The image had the numerically largest effects of the images proposed for use with this warning statement on the emotional reaction scale and had the numerically largest effects on the cognitive reaction scale in young adults and youth. As discussed in section III.B of this document, these salience impacts are important, as the research literature suggests that they are related to behavior change.

The image also had a significant impact (p<0.05) on beliefs about the health risks of smoking for smokers, and a significant impact (p<0.01) on beliefs about the health risks of secondhand smoke exposure for nonsmokers relative to the text-only control in the adult sample that viewed a hypothetical advertisement containing the proposed required warning.

The image also showed some of the largest effect sizes for image recall (at baseline and 1 week follow-up) in adults and youth across the images proposed for use with this warning statement, though it showed lower recall among study participants who viewed this image (ranging from 86.3 percent to 85.1 percent). However, recall of the warning statement was generally high at 1 week follow-up among study participants who viewed this image (ranging from 68.3 percent to 77 percent), and, based on the scientific literature, we conclude that repetitive viewing of the required warning is likely to increase recall. As explained in section III.C of this document, we gave greater weight to
outcomes on the salience measures than to outcomes on the recall measures. As is discussed in further detail in section III.E of this document, another image proposed for use with this warning statement, “deathly ill woman,” also had significant effects on all the salience measures (emotional reaction scale, cognitive reaction scale, and difficult to look at measure) in all three samples (adults, young adults, and youth). While we agree that this image, similar to the selected image of “cancerous lesion on lip,” is a very strong image that effectively conveys the negative health consequences of smoking, we ultimately chose “cancerous lesion on lip” for use with this warning statement for several reasons.

First, “cancerous lesion on lip” was the only image among the images proposed for use with this warning statement that had a positive impact on beliefs about the health risks of smoking and secondhand smoke exposure in one of the study samples (adults viewing a hypothetical advertisement).

Furthermore, as is stated in several comments (see the following paragraphs), the selected image, “cancerous lesion on lip,” is likely to have particular relevance for youth. As explained in some of these comments, the research literature suggests that youth are likely to relate to and be susceptible to cigarette warnings depicting the negative short-term impacts of smoking on their personal appearance, including their lips and teeth (Ref. 53).

We received a number of comments on this image, which we have summarized and responded to in the following paragraphs.

(Comment 70) FDA received a large number of comments supporting the use of the image “cancerous lesion on lip,” including comments from individuals, public health advocacy groups, a medical organization, academics, State and local public health agencies, and health care professionals. Several comments suggested that FDA should use this image because it has a very high potential to reach consumers and positively influence their behavior.

A few comments also specifically addressed the benefits of using an image that shows the public that cigarettes cause oral cancers, noting that public awareness of this negative health consequence is low, and that many smokers and nonsmokers only relate cigarettes to lung cancer (see also section II.C of this document regarding consumers’ lack of knowledge regarding the health risks of smoking).

Multiple comments also noted that, based on FDA’s research results, this image was the best choice among the four images proposed for use with this warning statement, significantly outperforming “white cigarette burning” and “red cigarette burning,” and slightly outperforming “deathly ill woman.”

(Response) We selected this image for use with this warning statement.

(Comment 71) Several comments noted that the image could be especially effective with younger audiences and could positively influence such audiences by illustrating how the health effects caused by smoking negatively affect their physical appearance. The comments indicated that adolescents can relate to and will be susceptible to this message.

(Response) We agree with these comments. It is important to include content in the required warnings that is relevant to youth. The image “cancerous lesion on lip” has the potential to positively impact youth behavior, in addition to adult and young adult behavior.

(Comment 72) As mentioned in section III.C of this document, some comments submitted to the docket described the results of scientific investigations that the submitters had conducted to examine the potential effectiveness of FDA’s proposed images on various outcomes. This image was discussed in some of these comments. For example, in one submitter’s study, participants rated this image highly on its ease of comprehension. It also induced relatively greater worry and feelings of discouragement from wanting to smoke than a text-only control. The submitter concluded that this image, along with “deathly ill woman,” was one of the most effective of the images proposed for use with this warning statement. In addition, this image was rated as the most effective of the 36 proposed images in another submitter’s survey of comparative effectiveness of the images in helping smokers quit. It was also the highest rated image among the set of images proposed by FDA for use with this warning statement in another submitter’s study of the potential effectiveness of the images, and was identified by high school students as one of the “top three” proposed required warnings (out of 36) in another submitter’s study.

(Response) As discussed in section III.C of this document, we carefully considered the comments submitted to the docket that described the results of studies conducted by the submitters on the docket that described the results of scientific investigations that the submitters had conducted to examine the potential effectiveness of FDA’s proposed images on various outcomes. This image was discussed in some of these comments. For example, in one submitter’s study, participants rated this image highly on its ease of comprehension. It also induced relatively greater worry and feelings of discouragement from wanting to smoke than a text-only control. The submitter concluded that this image, along with “deathly ill woman,” was one of the most effective of the images proposed for use with this warning statement. In addition, this image was rated as the most effective of the 36 proposed images in another submitter’s survey of comparative effectiveness of the images in helping smokers quit. It was also the highest rated image among the set of images proposed by FDA for use with this warning statement in another submitter’s study of the potential effectiveness of the images, and was identified by high school students as one of the “top three” proposed required warnings (out of 36) in another submitter’s study.

(Comment 73) FDA also received some comments critical of the image “cancerous lesion on lip.” Two comments indicated that the image was “too gross” to be effective, while another comment stated that it borders on the offensive. In contrast, some comments suggested that the image should be more graphic. Another comment suggested that oral cancer was an odd choice of cancers to depict in the graphic warning.

(Response) We disagree with these comments. With respect to the comments stating that the image was “too gross” or that it was offensive, the research literature indicates that images that evoke strong emotional reactions can promote greater awareness and better recollection of the health risks of smoking and can increase the likelihood smokers will reduce their smoking, make an attempt to quit, or quit altogether (Refs. 20, 44, and 45).

With respect to the suggestion that the image is not graphic enough, as discussed previously, this image had a highly significant effect (p<0.001) on all the salience measures (emotional reaction scale, cognitive reaction scale, and difficult to look at measure) in all three study populations (adults, young adults, and youth), which in turn suggests that the image has the potential to motivate positive behavior change (Id.).

Furthermore, the choice of cancers depicted in the required warning is appropriate, and will help inform the public that cigarettes cause oral cancers, and thus increase public awareness of the negative health consequences of smoking.

5. “WARNING: Cigarettes Cause Strokes and Heart Disease”

We selected the image which appears on pages 39 and 40 of the document “Proposed Required Warning Images,” referred to as “oxygen mask on man’s face,” for use with this warning statement.

In our research study, this image had a significant effect (p<0.001) on all the salience measures (emotional reaction scale, cognitive reaction scale, and difficult to look at measure) in all three study populations (adults, young adults, and youth). The image had the numerically largest effects of the images proposed for use with this warning statement on the emotional reaction scale and the difficult to look at measure in all the study populations. These images are important, as the research literature suggests that graphic warnings that evoke responses of this kind are...
likely to increase awareness of the health risks of smoking and increase the likelihood that smokers will reduce their smoking, make an attempt to quit, or quit altogether (Refs. 20, 44, and 45).

The image also showed some of the largest effect sizes for image recall (at baseline and 1 week follow-up) in adults and youth across the images proposed for use with this warning statement.

We received a number of comments on this image, which we have summarized and responded to in the following paragraphs.

(Comment 74) FDA received a large number of comments supporting the use of the image “oxygen mask on man’s face,” including comments from individuals, medical organizations, public health advocacy groups, health care professionals, State public health agencies, and academics. Many of these comments indicated that this image is the best image for use with this warning statement. Some also noted that the image will make smokers think twice about continuing to smoke. Some comments also noted that the image is beneficial in that it will inform the public of negative consequences of smoking aside from lung disease.

Some comments also noted that, based on FDA’s research results, this image was the best choice for use with this warning statement, noting that it elicited the highest scores on the emotional reaction scale of the images tested for use with this statement in FDA’s research study.

(Response) We selected this image for use with this warning statement.

(Comment 75) As described in section III.C of this document, some comments submitted to the docket described the results of scientific investigations that the submitters had conducted to examine the potential effectiveness of FDA’s proposed images on various outcomes. This image was discussed in some of these comments. For example, in one submitter’s study, participants rated this image highly on its ease of comprehension. It also induced relatively greater worry and feelings of discouragement from wanting to smoke than a text-only control. The submitter concluded that this image was the most effective of the images proposed for use with this warning statement. In another submitter’s study, this image was the highest-rated of the FDA-proposed images for use with this warning statement; however, this study also evaluated two images used with similar warning statements in other countries (one of open heart surgery, one of a bloody brain), and noted that they rated higher than FDA’s proposed images.

(Response) As discussed in section III.C of this document, we carefully considered the comments submitted to the docket that described the results of studies conducted by the submitters on our proposed required warnings. The results summarized in these comments are generally supportive of our image selection decisions.

(Comment 76) FDA also received some comments critical of the image “oxygen mask on man’s face.” One comment noted that the image was “too gross to be effective,” and one comment stated that the image should feature a younger person to highlight the fact that heart attacks and stroke can occur in young smokers as well as in older smokers.

(Response) The image “oxygen mask on man’s face” is an appropriate image that effectively conveys the negative health consequences of smoking. We do not agree with the statement that the image is “too gross to be effective;” the image effectively elicited emotional and cognitive reactions in viewers in our research study, which in turn suggests that the image has the potential to promote greater awareness of the health risks of smoking and motivate positive behavioral outcomes, including an increased likelihood that smokers will reduce their smoking, make an attempt to quit, or quit altogether (Refs. 20, 44, and 45).

While we agree with the statement that heart disease and strokes can occur in young smokers as well as in older smokers, the selected required warning will effectively communicate with a range of audiences, including consumers of different ages. As described previously, “oxygen mask on man’s face” had a significant effect (p<0.001) on all the salience measures (emotion measures, cognition measures, and difficult to look at measure) in all three study populations (adults, young adults, and youth). We considered the variety and diversity reflected in the images in making selection decisions, and took into account the importance of selecting a set of required warnings that includes a diversity of styles (e.g., photographic versus illustrative), themes, and human images (e.g., race, gender, age). While the person shown in this image is an older man, some of the images show younger people. Overall, the nine selected required warnings will effectively communicate with a wide range of consumers, including both young and older smokers.


We selected the image which appears on pages 45 and 46 of the document “Proposed Required Warning Images,” referred to as “baby in incubator,” for use with this warning statement.

In our research study, this image had a significant effect (p<0.001) on all the salience measures (emotional reaction scale, cognitive reaction scale, and difficult to look at measure) in all three study populations (adults, young adults, and youth). The image had the numerically largest effects of the images proposed for use with this warning statement on the salience measures. As discussed in section III.B of this document, these salience impacts are important, as the research literature suggests that they are likely to be related to behavior change.

The image had a significant effect (p<0.01) on recall of the warning statement at baseline compared to the text-only control in youth. The image also had a significant effect (p<0.05) on statement recall at follow-up in young adults, and showed the largest effect sizes for image recall (at baseline and 1 week follow-up) in adults and youth across the images proposed for use with this warning statement.

The image had a significant impact (p<0.05) on beliefs about the health risks of smoking for smokers in adults, although it had a negative significant impact (p<0.05) on beliefs about the health risks of smoking for smokers in youth. Thus, the results on this beliefs measure were mixed for “baby in incubator.” However, given the strength of the effects observed for this image on the salience measures, the required warning that includes the “baby in incubator” image is likely to increase awareness of the health risks of smoking and increase the likelihood that smokers will reduce their smoking, make an attempt to quit, or quit altogether (Refs. 20, 44, and 45).

We received a number of comments on this image, which we have summarized and responded to in the following paragraphs.

(Comment 77) FDA received a number of comments supporting the use of the image “baby in incubator,” including comments from individuals, a community organization, a public health advocacy group, health care professionals, a State public health agency, and academics. Several of these comments indicated that this image is the best image for use with this warning statement, with some noting that the image effectively shows how smoking during pregnancy can damage a baby’s health. One comment noted that the image could stimulate discussion about how smoking affects pregnancy among youth.
One comment also noted that the image “baby in incubator” outperformed the other image proposed for use with this warning statement in FDA’s research study on the key criteria that have proven most meaningful.

(Response) We selected this image for use with this warning statement.

(Comment 78) As described in section III.C of this document, some comments submitted to the docket described the results of scientific investigations that the submitters had conducted to examine the potential effectiveness of FDA’s proposed images on various outcomes. This image was discussed in some of these comments. For example, in one submitter’s study, participants rated this image highly on its ease of comprehension. It also induced relatively greater worry and feelings of discouragement from wanting to smoke than a text-only control. The submitter concluded that this image was the most effective of the images proposed for use with this warning statement. However, in another submitter’s study, this image was evaluated against images used in other countries, one of which was very similar in composition to “baby in incubator” but which was a photograph rather than a graphic illustration. In that submitter’s study, the photographic image was rated significantly higher than “baby in incubator.”

(Response) As discussed in section III.C of this document, we carefully considered the comments submitted to the docket that described the results of studies conducted by the submitters on our proposed required warnings. The results summarized in these comments are generally supportive of our image selection decisions.

(Comment 79) FDA also received a number of comments critical of the image “baby in incubator.” The majority of these comments objected to the graphic illustration style used for the image, with some submitters approving of the concept but stating that a photograph would be more impactful, and some indicating that the style is inappropriate, either because it downplays the seriousness of the risk described in the required warning or because it would inappropriately appeal to youth without discouraging them from smoking.

Some comments indicated that the lettering style used in the image was difficult to read, and one comment stated that the results from FDA’s research study for this image, while better than the results for the other images proposed for use with this warning statement (“pacifier & ashtray”), were not compelling.

One comment stated that the image bordered on the offensive.

(Response) The image “baby in incubator” is an appropriate image that effectively conveys the negative health consequences of smoking. As discussed in section III.C of this document, we are aware that many comments received in the docket expressed concern about the use of graphic illustration style images and expressed a belief that this style was not strong enough to elicit appropriate reactions. However, as discussed in section III.C of this document, we disagree with the contention that the use of graphic illustration style images is categorically inappropriate. As the results from our research study demonstrate, the “baby in incubator” image effectively elicited emotional and cognitive reactions, showing a highly significant effect (p<0.001) on these measures in all study populations, which in turn suggests that the image has the potential to promote greater awareness of the health risks of smoking and motivate positive behavioral outcomes, including an increased likelihood that smokers will reduce their smoking, make an attempt to quit, or quit altogether (Refs. 20, 44, and 45).

In addition, based on the study results, we also do not agree that the image is inappropriately offensive or that our research results for this image are not compelling. Based on the overall feedback received, we also disagree that the text in the proposed warning is difficult to read.

7. “WARNING: Smoking Can Kill you”

We selected the image which appears on pages 49 and 50 of the document “Proposed Required Warning Images,” referred to as “man with chest staples,” for use with this warning statement. In our research study, this image had a significant effect (p<0.001) on all the salience measures (emotional reaction scale, cognitive reaction scale, and difficult to look at measure) in all three study populations (adults, young adults, and youth). The image had the numerically largest effects of the images proposed for use with this warning statement on the salience measures. As discussed in section III.B of this document, these salience impacts are important, as the research literature suggests that they are likely to be related to behavior change.

The image was also associated with higher intentions to quit smoking compared to the text-only control (p<0.05) in adults.

The proposed required warning featuring the “man with chest staples” image showed some of the largest effect sizes for image recall among the images proposed for this warning statement at baseline in all study populations and at 1 week follow-up in young adults and youth.

Young adults viewing the image had significantly lower recall of the warning statement than those viewing the text-only control at baseline (76.2 percent versus 92.3 percent) and 1 week follow-up (78.9 percent versus 91.3 percent). However, recall of the statement was generally high at baseline and follow-up among study participants who viewed this image (ranging from 76.2 percent to 90.4 percent), and repetitive viewing of the required warning is likely to increase recall. As explained in section III.C of this document, we gave greater weight to outcomes on the salience measures than to outcomes on the recall measures.

We received a number of comments on this image, which we have summarized and responded to in the following paragraphs.

(Comment 80) FDA received a large number of comments supporting the use of the image “man with chest staples,” including comments from individuals (including former smokers), public health advocacy groups, medical organizations, health care professionals, State and local public health agencies, and academics. Many of these comments indicated that this image is the best image for use with this warning statement, while some also noted that the image is appropriately attention-grabbing or powerful and that it will make smokers think twice about continuing to smoke, or help them smoke less. Some comments also noted that the image is an excellent way of driving home the message that smoking can kill you. One comment stated that the image is a strong, solid concept that has been used effectively in other countries that require graphic health warnings on cigarette packages.

Some comments stated that, based on FDA’s research results, this image is the best choice for use with this warning statement, noting that it ranked the highest scores on the emotional reaction scale of the images tested for use with this statement in FDA’s research study, and had other positive results.

(Response) We selected this image for use with this warning statement.

(Comment 81) As described in section III.C of this document, some comments submitted to the docket described the results of scientific investigations that the submitters had conducted to examine the potential effectiveness of FDA’s proposed images on various outcomes. This image was discussed in some of these comments. For example,
in one submitter’s study, participants rated this image highly on its ease of comprehension. It also induced relatively greater worry and feelings of discouragement from wanting to smoke than a text-only control. In another submitter’s study, it was noted that, based on respondents’ rating and ranking of this image’s effectiveness, the image clearly stands out as the highest rated of the images FDA proposed for use with this warning statement.

(Response) As discussed in section III.C of this document, we carefully considered the comments submitted to the docket that described the results of studies conducted by the submitters on our proposed required warnings. The results summarized in these comments are generally supportive of our image selection decisions.

(Comment 82) FDA also received some comments critical of the image “man with chest staples.” One comment stated that the image was “too gross to be effective,” while another stated the image “offends against human dignity.” A few comments suggested that the person in the image should look worse (e.g., paler, weaker, thinner, like he had suffered more), and some comments suggested the person’s death should be more clearly tied to smoking by the image. One comment indicated that persons unfamiliar with an autopsy may not understand the image.

(Response) The image “man with chest staples” is an appropriate image that effectively conveys the negative health consequences of smoking. We do not agree that the image “is too gross to be effective” or that it “offends against human dignity”; the image shows a realistic outcome of the negative health consequences caused by smoking, and effectively elicited emotional and cognitive reactions in viewers in our research study. This in turn suggests that the image has the potential to promote greater awareness of the health risks of smoking and motivate positive behavioral outcomes, including an increased likelihood that smokers will reduce their smoking, make an attempt to quit, or quit altogether (Refs. 20, 44, and 45).

Viewers will understand that the image shows someone who has died from a smoking-related cause. Although we agree that not all viewers will necessarily be familiar with an autopsy scar, it is important to keep in mind that the image is not used in isolation, but accompanies the textual warning statement, which provides additional context for what is shown. The results of our research study suggest that viewers from all age groups understood and reacted to this image in desirable ways. The figure shown is appropriate; although some of the negative health consequences of smoking may lead to the effects on appearance suggested in the comments (e.g., significant disease-related weight loss), other consequences, such as heart attacks, can kill smokers without first causing these effects.

8. “WARNING: Tobacco Smoke Causes Fatal Lung Disease in Nonsmokers”

We selected the image which appears on pages 57 and 58 of the document “Proposed Required Warning Images,” referred to as “woman crying,” for use with this warning statement.

In our research study, this image had a significant effect (p<0.001) on the emotional reaction scale in all three study populations (adults, young adults, and youth). It also showed significant effects on the difficult to look at measure in all study populations (adults (p<0.01), young adults (p<0.001), and youth (p<0.001)). The image was the only image proposed for use with this warning statement that showed significant effects on all the salience measures in our research study.

The image also had a significant impact (p<0.05) on beliefs about the health risks of smoking for smokers in young adults.

The proposed required warning that included this image also showed the largest effect sizes for image recall (at baseline and 1 week follow-up) in adults, young adults, and youth across the images proposed for this warning statement. Youth viewing the image had significantly lower recall of the warning statement than those viewing the text-only control at baseline (52.4 percent versus 68.9 percent). However, recall of the statement was generally high among study participants who viewed this image, and repetitive viewing of the required warning is likely to increase recall. As explained in section III.C of this document, we gave greater weight to outcomes on the salience measures than to outcomes on the recall measures.

FDA received a number of comments on this image, which the Agency has summarized and responded to in the following paragraphs.

(Comment 83) FDA received several comments supporting the use of the image “woman crying,” including comments from individuals (including former smokers) and public health advocacy groups. Some of these comments indicated that this image is the best image of the ones proposed for use with this warning statement. One comment stated that the image stood out as particularly effective among the proposed required warnings because it shows the devastating effects secondhand smoke can have on people who have tried to protect themselves by not smoking, and indicated that the image will remind smokers that they are harming their loved ones and others around them as well as themselves. Others noted that the image sends a powerful message.

One comment indicated that the image outperformed the other images proposed for use with this warning statement on the emotional reaction scale and the difficult to look at measure in FDA’s research, and noted that it appears to be a cut above the other images.

(Comment 84) One comment approved of the diversity reflected in the image (which shows an African-American woman).

(Response) We agree that it is beneficial to have a diverse set of images that communicate with a wide range of audiences, including a variety of population subgroups. In order to ensure that the final set of required warnings effectively communicates risk information to a diverse range of audiences, we selected a set of nine required warnings, including the image “woman crying,” that includes a variety of human images that are broadly representative of the overall population.

(Comment 85) As described in section III.C of this document, some comments submitted to the docket described the results of scientific investigations that the submitters had conducted to examine the potential effectiveness of FDA’s proposed images on various outcomes. This image was discussed in some of these comments. For example, this image induced relatively greater worry and led to higher ratings of feeling discouraged from wanting to smoke than a text-only control in one submitter’s study.

(Response) As discussed in section III.C of this document, we carefully considered the comments submitted to the docket that described the results of studies conducted by the submitters on our proposed required warnings. The results summarized in these comments are generally supportive of our image selection decisions.

(Comment 86) FDA also received some comments critical of the image “woman crying.” One comment indicated that the image borders on the
offensive, while another stated it is too sensational to be effective.

Other comments suggested that the image did not directly portray a health consequence of secondhand smoke, or that the image is not clearly tied to secondhand smoke. One comment also suggested that the image should not be used because it did not have an impact on beliefs about the health harms of secondhand smoke or on quit intentions in FDA’s research study.

(Response) We disagree with these comments. The image “woman crying” is an appropriate image that effectively conveys the negative health consequences of smoking. We do not agree that the image is offensive or too sensational; the image is a realistic portrayal of the negative health consequences caused by exposure to secondhand smoke that affect people. It effectively elicited emotional and cognitive reactions in those who viewed it in our research study, which in turn suggests that the image has the potential to promote greater awareness of the health risks of smoking and motivate positive behavioral outcomes, including an increased likelihood that smokers will reduce their smoking, make an attempt to quit, or quit altogether (Refs. 20, 44, and 45).

We do not agree that the image does not depict a health consequence of secondhand smoke. Graphic depictions of the visible effects of disease are not the only way of communicating the health risks of secondhand smoke exposure (see Ref. 11). The negative health consequences caused by secondhand smoke exposure, including fatal lung disease, have many dimensions, including emotional suffering. This image highlights that dimension. Furthermore, it is important to keep in mind that the image is not used in isolation, but accompanies the textual warning statement, which provides additional context for what is shown. As evidenced by the image’s significant impact on the salience measures across the populations participating in our research study, the proposed required warning effectively depicts the health consequences of secondhand smoke exposure, including the suffering endured by those experiencing these health consequences.

9. “WARNING: Quitting Smoking Now Greatly Reduces Serious Risks to Your Health”

We selected the image which appears on pages 67 and 68 of the document “Proposed Required Warning Images,” referred to as “man I Quit t-shirt,” for use with this warning statement.

In our research study, the image had a statistically significant effect on the emotional reaction scale in young adults (p<0.05), and on the cognitive reaction scale in adults (p<0.05), young adults (p<0.01), and youth (p<0.001).

The proposed required warning that included this image also showed the largest effect sizes for image recall (at baseline and 1 week follow-up) in adults, young adults, and youth across the images proposed for this warning statement.

Although this image, along with the other images proposed for use with this warning statement, did not elicit the magnitude of reactions on the salience measures (emotional reaction scale, cognitive reaction scale, difficult to look at measure) that some of the images proposed for use with other warning statements did, this is likely a result of the information being conveyed in the warning statement, which emphasizes the positive health benefits of quitting smoking. The content of this required warning is not expected to arouse the same level of response on some of the salience measures as the other messages.

However, the research literature suggests that warnings that focus on the benefits of quitting are effective at encouraging cessation, and suggests that positive, self-efficacy messages can be used effectively as one component of graphic health warnings to increase smokers’ motivations and confidence about quitting (Ref. 40 at pp. 35, 39–41). The research literature also highlights the importance of including one or more warnings that provide solutions, such as the “man I Quit t-shirt” required warning, in a set of warnings conveying the negative health consequences of smoking. Specifically, the literature recommends that, in addition to communicating the health risks of smoking, some warnings should also provide information on how to avoid these risks (i.e., by quitting), in order to optimize the effectiveness of the overall set of warning messages (see Ref. 48 and Ref. 40 at p. 37).

As is discussed in further detail in section III.E of this document, another image proposed for use with this warning statement, “cigarettes in toilet bowl,” also had significant effects on the emotional reaction scale in some study populations and on the cognitive reaction scale, as well as showing positive effects on other study measures. While this image, similar to the selected image (“man I Quit t-shirt”), could be effectively used with this warning statement, we ultimately selected “man I Quit t-shirt” with this warning statement based on a consideration of multiple factors, including the feedback received in the docket, which is discussed in the comment summaries in the following paragraphs and in section III.E of this document.

Furthermore, as noted in section III.A of this document, in order to ensure that the final set of required warnings effectively communicates risk information to a diverse range of audiences, we selected a set of nine required warnings, including the image “man I Quit t-shirt,” that includes a variety of human images that are broadly representative of the overall population. The image “man I Quit t-shirt” contributes to the variety seen in the final set of images by picturing a man who is younger than the men in the other required warning images.

Additionally, as reflected in the comment summary, the man shown in the image is perceived by many viewers as strong and “macho,” suggesting that the image has the potential to reach and effectively communicate with a demographic group that has been heavily targeted by tobacco industry cigarette advertising (see Ref. 54 at p. 151). The depiction of men as strong, powerful, macho, rugged, and independent, and the association of these characteristics with cigarette brands, has long been a prominent theme in tobacco industry advertising (Id. at p. 151), and targeted marketing efforts by the tobacco industry have led to greater smoking uptake and lower cessation rates in targeted subgroups (Id. at p. 211).

We received a number of comments on this image, which we have summarized and responded to in the following paragraphs.

(Comment 87) FDA received a number of comments supporting the use of the image “man I Quit t-shirt,” including comments from individuals, public health advocacy groups, medical organizations, and State and local public health agencies. Many of these comments indicated that this image is the best image of the ones proposed for use with this warning statement. Several of the comments discussed specific favorable aspects of the image or potential effects of the image, including that the image models a positive behavior, is compelling, and that it will encourage others to quit. Several comments believed that the image could reach a critical demographic group by showing a younger, “cool,” “macho” man and suggesting that it is manly and/or cool to quit smoking.

Some comments also suggested that the image is positive in that it shows that quitting is a heroic decision.

(Response) We selected this image for use with this warning statement.
positive manner, discussing the health benefits of ceasing to smoke, and the image is consistent with this text. This required warning, particularly as part of the overall set of required warnings, will help educate consumers about the negative health consequences of smoking and help encourage positive behavior (see Ref. 40 at pp. 35 and 40).

Based on the overall feedback received and the results from our research study, we also disagree that the text in the proposed warning is difficult to read or that the image is banal.

10. Image for Advertisements With a Small Surface Area

In addition to proposing 36 required warnings for use on cigarette packages and in cigarette advertisements in the NPRM, we also proposed two other color graphics for use solely in advertisements with a small surface area of less than 12 square inches (75 FR 69524 at 69539). As we explained in the NPRM, these two proposed color graphics differ in their composition from the other proposed images in that the details of these two color graphics should be clear, conspicuous, and legible even when the image is reduced in size to occupy 20 percent of a surface with an area of less than 12 square inches (75 FR 69524 at 69535). We proposed that whichever of these options was selected would be used in combination with one of the nine textual statements only in advertisements with a small surface area (i.e., less than 12 square inches).

However, as we noted in the NPRM, even an advertisement with a relatively small surface area would need to be large enough so that the required graphic and accompanying textual warning statement are clear, conspicuous, and legible (75 FR 69524 at 69539). We selected the image which appears on page 75 of the document entitled “Proposed Required Warning Images” for use with the textual warning statements solely in advertisements with a small surface area (defined as less than 12 square inches). This image depicts a black exclamation mark enclosed within a red equilateral triangle.

As stated previously, FDA proposed two images for use solely with the textual warning statements in advertisements with a small surface area; the selected image described in the previous paragraph and an image of a burning cigarette enclosed in a red circle with a red bar across it. We did not receive any comments on either of the proposed images.

Versions of both of these images have been used in other contexts. For example, the image of an exclamation mark enclosed within a triangle is often used to draw attention to a warning of danger or hazards that could result in personal injury or a threat to health (see, e.g., 16 CFR 1211.15, 16 CFR 1407.3; 16 CFR 1500.19; and Ref. 56). The image of a burning cigarette enclosed in a red circle with a red bar across it is the international “No Smoking” symbol (Ref. 56) and is often used on signs and placards to denote an area where smoking is prohibited (see, e.g., 14 CFR 23.853, 49 CFR 374.201).

In light of the other contexts in which the two proposed images are used, we selected the image of the exclamation mark enclosed within a red equilateral triangle, as we believe this image is more appropriate than the other proposed image for use in the required warnings. As stated, this image is commonly used to draw attention to a warning of danger which could result in personal injury or a threat to health, which is consistent with its purpose in cigarette advertisements with a small surface area. Many consumers have likely been exposed to similar symbols in other contexts and, as a result, are likely to recognize and understand that the image is drawing attention to a warning of a threat to health.

E. Non-Selected Images

This section discusses the 27 color graphic images that we proposed but have not selected for use at this time, and the factors that influenced the decision not to use each image, including the research results for the images, the comments received in the docket, and the relevant scientific literature.

Consistent with the discussion of selected images in section III.D of this document, the images are referred to in this section by the pages on which they appear in the “Proposed Required Warning Images” document and by the descriptive names used in the study report (Ref. 49, study report) summarizing the results of FDA’s research study.

1. “WARNING: Cigarettes Are Addictive”

As discussed in section III.D of this document, we selected the image “haze in throat” for use with the statement, “WARNING: Cigarettes are addictive.” We proposed three other images for use with this statement: “cigarette injection,” which appears on pages 3 and 4 of the document “Proposed Required Warning Images”; “red puppet,” which appears on pages 5 and
that, although cigarettes are legal products, they are just as addictive as illegal drugs like heroin. One comment indicated that the image would be particularly effective with underage smokers.

FDA also received several comments that opposed the use of the image “cigarette injection.” Many of these comments objected to the graphic illustration style used in the image, with some stating it would be ineffective or less effective than a photographic image, and some indicating it would detract from the seriousness of the message being conveyed. Some comments also expressed concern that the style would inappropriately appeal to youth without deterring them from smoking.

A few comments also objected to the comparison of legal cigarette products with illegal drugs, with one comment indicating this downplayed the seriousness of intravenous drug use, and another comment noting that the analogy of cigarette use to heroin use could cause consumers to discount the message if they believe that cigarette and heroin use are not comparable.

Some comments also stated that the image could be misunderstood or was too abstract, and one comment stated that the image does not illustrate adverse health effects.

One comment noted that the proposed required warning featuring the “cigarette injection” image was not rated highly on its ease of comprehension in a research study the submitter conducted on the 36 proposed required warnings, though it did show a significant effect on worry and feeling discouraged from wanting to smoke relative to a text-only control.

(Response) We are not selecting this image for use in a required warning and instead have selected the image “hole in throat” for the reasons given in section III.D of this document.

Red puppet. In FDA’s research study, the image “red puppet” had significant effects on the emotional and cognitive reaction scales in all three study populations. It also showed higher correct recall of the warning statement compared to the control in young adults at 1 week follow-up.

However, the selected image, “hole in throat,” had numerically larger effects than this image on the salience measures (emotional and cognitive reaction scales, difficult to look at measure) in all three study populations. In addition, looking across the different measures used in the research study, both the image “hole in throat” and “cigarette injection” had stronger overall research results than this image.

FDA received a number of comments on this image, which the Agency has summarized and responded to in the following paragraphs.

(Comment 90) FDA received several comments that supported the use of the image “cigarette injection,” including comments from individuals, public health advocacy groups, and a State public health agency. Some of the comments stated that the image would be an effective smoking deterrent. Several of the comments noted that the image would help smokers understand
Looking across the different measures used in FDA’s research study, this image was relatively less effective than other images proposed for this warning statement, including the image selected for use in the required warnings “hole in throat.”

FDA received a number of comments on this image, which the Agency has summarized and responded to in the following paragraphs.

(Comment 92) FDA received multiple comments that supported the use of the image “woman in rain,” including comments from individuals, a community organization, and a State public health agency. Some of the comments stated that the image is likely to be effective, and one stated that smokers would be able to relate to the image.

FDA also received a number of comments that opposed the use of the image “woman in rain.” Some of these comments stated that the image would not be effective and is not emotionally arousing, while some stated that it shows a very weak harm (i.e., standing in the rain). Another comment stated that the image makes smoking seem like a normal behavior.

Several comments expressed concern that the image would not be understood by consumers, indicating it was too vague in nature and requires a high analytical ability to understand.

Several comments state the image does not convey a health consequence of smoking, while three comments stated that the results from FDA’s research study for this image did not support its selection from among the images proposed for use with this warning statement.

Two comments noted that the proposed required warning featuring the “woman in rain” image was not highly rated in research studies conducted by the submitters. One comment noted that the image was not rated highly on its ease of comprehension and did not increase worry relative to a text-only label or discourage respondents from smoking relative to a text-only label in the submitter’s study, while another noted that the image was ranked as one of the least effective of the 36 proposed images by respondents in the submitter’s study.

(Comment 93) FDA received multiple comments that supported the use of the image “smoke approaching baby,” including comments from individuals, a medical organization, public health advocacy groups, academics, and State and local public health agencies. Some of these comments indicated that the image would cause people to reconsider smoking due to the harm it can cause to others, especially a child or a baby.

Three comments noted that the image showed positive impacts in research studies conducted by the submitters. Specifically, in one submitters’ study this image had the relatively greatest impact in discouraging respondents from wanting to smoke of the images proposed for use with this warning statement. In another submitter’s study of the potential effectiveness of the images, this image received the highest overall rating of the images proposed for use with this warning statement. In addition, it was one of the two highest rated images of the FDA images proposed for use with this warning statement in another submitter’s study.

FDA also received several comments that opposed use of the image “smoke at toddler.” Multiple comments stated that the image would be perceived as demeaning to smokers by suggesting they blow smoke directly at their children, and one comment cited the image as an unreal portrayal. Another comment expressed concern that the image would prompt denial among smokers, who would interpret the image to mean that their children are not at risk if they do not blow smoke directly at them. One comment said the image does not depict a negative health consequence of smoking, while another comment stated the image was too positive, in that the child looked too happy. Finally, another comment stated that other images tested in FDA’s research study for use with this warning statement elicited higher scores on the emotional and cognitive reaction scales than this image.

(Comment 94) FDA received comments that supported the use of the image “smoke at baby” and significant impacts on all the salience measures in all three study populations, and also showed significant impacts on recall and behavioral intentions in some populations.

FDA received comments that supported the use of the image “smoke at toddler” and significant impacts on all the salience measures in all three study populations, and also showed significant impacts on recall and behavioral intentions in some populations.

FDA received comments that supported the use of the image “smoke approaching baby” and significant impacts on all the salience measures in all three study populations, and also showed significant impacts on recall and behavioral intentions in some populations.

FDA received comments that supported the use of the image “smoke at baby,” including comments from individuals, a community organization, a medical organization, academics, and a State public health agency. Some of these comments indicated that the image would cause people to reconsider smoking due to the harm it can cause to others, especially a child or a baby.

FDA received comments that supported the use of the image “smoke at baby,” including comments from individuals, a community organization, a medical organization, academics, and a State public health agency. Some of these comments indicated that the image would cause people to reconsider smoking due to the harm it can cause to others, especially a child or a baby.
studies conducted by the submitters. Specifically, this image had a significant impact in discouraging respondents from wanting to smoke in one submitter’s study, and it was one of the two highest-rated images of the FDA images proposed for use with this warning statement in another submitter’s study.

FDA also received several comments that opposed the use of the image “smoke at baby.” Many of these comments objected to the graphic illustration style used in the image, with some stating it would be ineffective or less effective than a photographic image, and some indicating it would detract from the seriousness of the message being conveyed. Some comments also expressed concern that the style would inappropriately appeal to youth without deterring them from smoking.

Multiple comments stated that the image would be perceived as demeaning to smokers by suggesting they blow smoke directly at their children, and one commented the image as an unreal portrayal. Another comment expressed concern that the image would prompt denial among smokers, who would interpret the image to mean that their children are not at risk if they do not blow smoke directly at them.

A couple of comments stated that other images tested in FDA’s research study for use with this warning statement outperformed this image, with one noting that other images elicited higher scores on the emotional reaction scale and difficult to look at measure than this image, and another noting that other images had higher scores on the quit intentions and recall measures than this image.

One comment expressed concern that the image could be perceived to mean that mothers who smoke should not breastfeed their children. Another comment stated that the text used in the proposed required warning was difficult to read.

(Response) We are not selecting this image for use in a required warning and instead have selected the image “smoke approaching baby” for the reasons given in section III.D of this document.

Girl crying. In FDA’s research study, the image “girl crying” had significant effects on all the salience measures (emotional reaction scale, cognitive reaction scale, and difficult to look at measure) in all three study populations (adults, young adults, and youth). It also showed higher correct recall of the warning statement compared to the control in adults at baseline, and higher correctly recalled warning statement at 1 week follow-up compared to the text-only control for adults and young adults. Youth who viewed the image also reported that they would be significantly less likely to be smoking 1 year from now compared to youth who viewed the control.

However, the image had a significant negative impact on adult beliefs about the health risks of secondhand smoke exposure for nonsmokers, i.e., adults who viewed the image were less likely to believe that nonsmokers will suffer from negative health effects due to secondhand smoke exposure than adults who viewed the text-only control.

As discussed in section III.D of this document, the selected image, “smoke approaching baby,” had significant impacts on all the salience measures in all three study populations, and also showed significant impacts on recall and behavioral intentions in some populations. Thus, while “girl crying” showed positive effects on several important measures in FDA’s research study, the selected image was considered to be a stronger choice, as it also showed positive effects on several important measures and did not show any negative effects.

FDA received a number of comments on this image, which the Agency has summarized and responded to in the following paragraphs.

(Comment 95) FDA received several comments that supported the use of the image “girl crying,” including comments from individuals and from a State public health agency. Some comments noted that the submitter found this image to be the most effective of the images proposed for use with this warning statement, and others noted it would appropriately elicit negative emotions in viewers.

FDA also received several comments that opposed use of the image “girl crying.” Multiple comments stated that it was not clear why the girl was crying, and one comment stated that the image does not depict a health consequence of secondhand smoke exposure. One comment indicated that the image was too sensational to be effective, and another comment cited the image as an unreal portrayal, stating that young children do not know they are being harmed when they are exposed to smoke and thus would not cry as a result of such exposure, and noted that this is what makes secondhand smoke exposure so insidious. One comment indicated that other images tested in FDA’s research study for use with this warning statement had superior overall results to this image.

(Response) We are not selecting this image for use in a required warning and instead have selected the image “smoke approaching baby” for the reasons given in section III.D of this document.

Warning in child lettering. In FDA’s research study, the image “warning in child lettering” had significant effects on the emotional and cognitive reaction scales in all three study populations (adults, young adults, and youth). It also showed higher correct recall of the warning statement compared to the control in adults and young adults at baseline, and higher correct recall of the warning statement at 1 week follow-up compared to the control for adults, young adults, and youth. However, “warning in child lettering” showed lower correct recall of the image at baseline and follow-up for adults, young adults, and youth compared to the other images.

Looking across the different measures used in FDA’s research study, this image was relatively less effective than other images proposed for use with this warning statement, including the image selected for use in the required warnings, “smoke approaching baby.”

FDA received a number of comments on this image, which the Agency has summarized and responded to in the following paragraphs.

(Comment 96) FDA received several comments that supported the use of the image “warning in child lettering,” including comments from individuals, a public health advocacy group, a medical organization, and a State public health agency. Some comments felt the use of child’s handwriting in the image would be especially impactful with parents, and one comment noted that this image would have wide appeal, resonating with parents of any race or ethnicity.

FDA also received several comments that opposed use of the image “warning in child lettering.” Multiple comments objected to the image style, indicating that a photographic depiction would be more effective at deterring people from smoking, with one comment noting that the image style would be inappropriately appealing to youth without discouraging them from smoking. One comment indicated that the image does not depict a negative health consequence of smoking, and another indicated that the image was not eye-catching.

Two comments noted that other images proposed for use with this warning statement had superior overall results compared to this image in FDA’s research study and stated that FDA should not select this image for use in the required warning. In addition, two comments noted that the image was not highly rated in studies conducted by the submitters. One comment noted that the image was
ranked as the least effective of the 36 proposed images by respondents in the submitter’s study, while another noted that the image was ranked the lowest by a considerable margin of the images proposed for use with this warning statement in the submitter’s study.

(Comment 97) FDA received a number of comments on this image, which the Agency has summarized and responded to in the following paragraphs.


FDA also received several comments that opposed use of the image “girl in oxygen mask.” Some comments suggested that the image should show more severe disease or more clear association between the girl’s illness and smoke exposure.

(Comment 98) FDA received a number of comments on this image, which the Agency has summarized and responded to in the following paragraphs.

(Comment 99) FDA received some comments that supported the use of the image “lungs full of cigarettes,” including comments from individuals and State and local public health agencies. Some of these comments indicated that the image is the best choice for use with this warning statement, while some also noted that

to higher ratings of feeling discouraged from wanting to smoke than a text-only control in one submitter’s study. The image was also one of the five images rated most effective among the images used in FDA’s 36 proposed required warnings in another submitter’s study of the potential effectiveness of the images.

FDA also received several comments that opposed use of the image “toe tag,” with some submitters indicating that consumers, and in particular minority populations, might not understand what the image of a toe tag signifies. Some comments stated that the image “offend[s] against human dignity” or is “too sensational to be effective,” while it was alternatively stated that the image should be more graphic or show more suffering. It was also noted in the comments that the image did not test as well as other images proposed for use with this warning statement in FDA’s research study.

(Comment 98) FDA received a number of comments on this image, which the Agency has summarized and responded to in the following paragraphs.

As discussed in section III.D of this document, the selected image, “smoke approaching baby,” had significant impacts on all the salience measures in all three study populations, and also showed significant impacts on recall and behavioral intentions in some populations. Thus, the selected image was considered to be a stronger choice than “girl in oxygen mask,” as it showed positive effects on several important measures, but did not show any negative effects.

FDA also received several comments that opposed use of the image “girl in oxygen mask.” Some comments noted that it was unclear that the person portrayed in the image was a child, and suggested that the image would be more persuasive if the person shown were younger. One comment expressed concern that persons of low socioeconomic status would not understand the image, and a few comments suggested that the image should show more severe disease or more clear association between the girl’s illness and smoke exposure.

(Comment 99) FDA received some comments that supported the use of the image “lungs full of cigarettes,” including comments from individuals and State and local public health agencies. Some of these comments indicated that the image is the best choice for use with this warning statement, while some also noted that

Some comments noted that the image showed positive effects in research studies conducted by the submitters. Specifically, this image was rated highly on its ease of comprehension and induced relatively greater worry and led
the image is particularly appropriate for use with the warning statement.

As discussed in section III.C of this document, some comments submitted to the docket described the results of research conducted by the submitters to examine the potential effectiveness of FDA’s proposed images. This image was discussed in some of these comments. Specifically, in one submitter’s study, participants rated this image highly on its ease of comprehension. It also induced relatively greater worry and feelings of discouragement from wanting to smoke than a text-only control. However, the image was rated as one of the least effective of the images proposed by FDA for use with this warning statement in another submitter’s study of the potential effectiveness of the images.

FDA also received several comments that opposed use of the image “lungs full of cigarettes,” with some submitters indicating that consumers might not understand the image, and some comments that the image should show the consequences of lung disease on a real person or on real lungs and suggesting that the proposed image did not depict health consequences in an understandable, hard-hitting manner. One comment noted that the secondary message highlighted by the use of bold face emphasis in this proposed required warning (“I cause disease”), could be interpreted as blaming smokers for their addiction, and expressed concern that this could undermine the proposed required warning’s ability to communicate effectively with smokers. One comment also stated that the image did not show desirable effects on some measures in FDA’s research study.

(Response) We are not selecting this image for use in a required warning and instead have selected the image “healthy/diseased lungs” for the reasons given in section III.D of this document.

Dr. with X-ray. In FDA’s research study, the image “Dr. [doctor] with X-ray” had significant effects on the emotional and cognitive reaction scales in all three study populations (adults, young adults, and youth). It also had significant effects on the difficult to look at measure. As discussed in section III.D of this document, the selected image, “healthy/diseased lungs,” had significant effects on all the salience measures in all study populations, and had the largest numerical effects of the images proposed for use with this warning statement on the salience measures.

Among young adults, the image “Dr. with X-ray” prompted higher correct recall of the warning statement at baseline and at 1 week follow-up than the text-only control, as well as higher correct recall of the warning statement at follow-up among youth and the adult sample that viewed a hypothetical advertisement featuring this proposed required warning.

However, among young adults, as well as among the adult sample who viewed a hypothetical advertisement featuring this image, “Dr. with X-ray” was negatively associated with beliefs about the health risks of secondhand smoke exposure to nonsmokers (i.e., participants viewing this image were less likely to believe that nonsmokers will suffer health consequences related to secondhand smoke exposure than participants viewing the text-only control).

FDA received a number of comments on this image, which the Agency has summarized and responded to in the following paragraphs.

(Comment 100) FDA received some comments that supported the use of the image “Dr. with X-ray,” including comments from individuals, a public health advocacy group, a community organization, and a State public health agency. These comments noted that the “Dr. with X-ray” image is particularly appropriate for use with the warning statement, or expressed the view that the image is the best choice for use with this warning statement.

As discussed in section III.C of this document, some comments submitted to the docket described the results of research conducted by the submitters to examine the potential effectiveness of FDA’s proposed required warnings. This image was discussed in some of these comments. Specifically, this image was rated highly on its ease of comprehension in one submitter’s study, but failed to show an effect on other study measures (worry, discouragement from smoking). The image was one of the five images rated least effective among the images used in FDA’s 36 proposed required warnings in another submitter’s study of the potential effectiveness of the images, and it was also rated as the least effective of the images proposed by FDA for use with this warning statement in another submitter’s study of the potential effectiveness of the images.

FDA also received several comments that opposed use of the image “Dr. with X-ray,” with some submitters indicating that the X-ray shown in the image is unclear and that the image would not be understood by consumers, and some indicating that it was too vague or clinical in nature and did not effectively convey the full impact of lung disease. It was also noted in the comments that the image failed to show desirable effects on some measures in FDA’s research study, and that it showed negative effects on the beliefs measure among some of the study participants.

(Response) We are not selecting this image for use in a required warning and instead have selected the image “healthy/diseased lungs” for the reasons given in section III.D of this document.

4. “WARNING: Cigarettes Cause Cancer”

As discussed in section III.D of this document, FDA selected the image “cancerous lesion on lip” for use with the statement, “WARNING: Cigarettes cause cancer.” FDA proposed three other images for use with this statement: “Deathly ill woman,” which appears on pages 29 and 30 of the document “Proposed Required Warning Images;” “white cigarette burning,” which appears on pages 31 and 32 of the document “Proposed Required Warning Images;” and “red cigarette burning,” which appears on pages 35 and 36 of the document “Proposed Required Warning Images.”

Deathly ill woman. The image “deathly ill woman” had strong overall research results in FDA’s research study, including significant effects on all the salience measures (emotional reaction scale, cognitive reaction scale, and difficult to look at measure) in all three study populations (adults, young adults, and youth).

However, overall the selected image, “cancerous lesion on lip,” had slightly higher numerical scores on the emotional and cognitive reaction scales than this image.

Among adults, the image “deathly ill woman” prompted lower correct recall of the warning statement at baseline and at 1 week follow-up. However, the image showed some of the largest effect sizes for image recall (baseline and follow-up) across the images proposed for use with this warning statement.

FDA received a number of comments on this image, which the Agency has summarized and responded to in the following paragraphs.

(Comment 101) FDA received a large number of comments that supported the use of the image “deathly ill woman,” including comments from individuals, public health advocacy groups, medical organizations, academics, and State and local public health agencies. Many of these comments indicated that this image is the best image for use with this warning statement, with some stating that the image would communicate effectively to women and other comments approving the image’s accurate portrayal of the effects cancer can have on personal appearance.
Some comments noted that the image showed positive impacts in research studies conducted by the submitter. Specifically, in one submitter’s study, participants rated this image highly on its ease of comprehension. It also induced relatively greater worry and feelings of discouragement from wanting to smoke than a text-only control. The submitter concludes that this image, along with “cancerous lesion on lip,” was the most effective of the images proposed for use with this warning statement. The image was also one of the five images rated most effective among the images used in FDA’s 36 proposed required warnings in another submitter’s study of the potential effectiveness of the images. It was also one of two images rated effective among FDA’s 36 proposed color graphic in another submitter’s study of the effectiveness of the images at stopping someone from smoking, and it was identified by high school students as one of the “top three” proposed required warnings in another submitter’s study.

FDA also received comments that opposed the use of the image “deathly ill woman.” Some comments noted that the image “offend[s] against human dignity,” while one stated it was “too sensational to be effective.” Conversely, some comments indicated that the image should show more obvious signs of illness. It was also noted in the comments that the image did not show desirable effects on all the measures in FDA’s research study.

(Response) We are not selecting this image for use in a required warning and instead have selected the image “cancerous lesion on lip” for the reasons given in section III.D of this document.

White cigarette burning. In FDA’s research study, the image “white cigarette burning” had significant effects on the emotional and cognitive reaction scales in all three study populations (adults, young adults, and youth). It also had significant effects on the difficult to look at measure and local public health agencies. These comments noted that the “white cigarette burning” image is particularly appropriate for use with the warning statement, or expressed the submitter’s preference that the image be used with this warning statement. As discussed in section IIIC of this document, some comments submitted to the docket described the results of research conducted by the submitters to examine the potential effectiveness of FDA’s proposed images. This image was discussed in some of these comments. Specifically, this image was rated highly on its ease of comprehension in one submitter’s study, but failed to show an effect on other study measures (worry, discouragement from smoking). The image was rated as the least effective of the images proposed by FDA for use with this warning statement in another submitter’s study of the potential effectiveness of the images.

FDA also received several comments that opposed use of the image “white cigarette burning,” with some submitters indicating that the image does not depict the negative health consequences of smoking or that the image is not appropriately evocative of cancer, and some noting that the image is unclear and will not be understood by consumers. Some comments also criticized the design of the image, and one stated that the image is not appropriately evocative of the Tobacco Control Act. Some comments also noted that this image of a burning cigarette could trigger cravings in smokers. It was also noted in the comments that the image failed to show desirable effects on some measures in FDA’s research study. One comment noted that the secondary message highlighted by the use of bold face emphasis in this proposed required warning (“I cause cancer”) could be interpreted as blaming smokers, and expressed concern that this could undermine the proposed required warning’s ability to communicate effectively with smokers.

(Response) We are not selecting this image for use in a required warning and instead have selected the image “cancerous lesion on lip” for the reasons given in section III.D of this document.

Red cigarette burning. In FDA’s research study, the image “red cigarette burning” had significant effects on all the salience measures (emotional reaction scale, cognitive reaction scale, and difficult to look at measure) in all three study populations (adults, young adults, and youth).

However, the selected image, “cancerous lesion on lip,” generally had numerically larger effects than this image on the salience measures.

Among adults, young adults, and youth, the image “red cigarette burning” prompted lower correct recall of the warning statement at baseline and at 1 week follow-up. The proposed required warning featuring this image also prompted relatively lower recall of the image at baseline and at 1 week follow-up among adults, young adults, and youth than “cancerous lesion on lip.”

Youth viewing the image “red cigarette burning” reported being more likely to be smoking 1 year from now than youth viewing the text-only control.

FDA received a number of comments on this image, which the Agency has summarized and responded to in the following paragraphs.

(Comment 102) FDA received some comments that supported the use of the image “white cigarette burning,” including comments from individuals and from State and local public health agencies. These comments noted that the “white cigarette burning” image is particularly appropriate for use with the warning statement, or expressed the submitter’s preference that the image be used with this warning statement. As discussed in section IIIC of this document, some comments submitted to the docket described the results of research conducted by the submitters to examine the potential effectiveness of FDA’s proposed images. This image was discussed in some of these comments. Specifically, this image was rated highly on its ease of comprehension in one submitter’s study, but failed to show an effect on other study measures (worry, discouragement from smoking). The image was rated as the least effective of the images proposed by FDA for use with this warning statement in another submitter’s study of the potential effectiveness of the images.

FDA also received several comments that opposed use of the image “white cigarette burning,” with some submitters indicating that the image does not depict the negative health consequences of smoking or that the image is not appropriately evocative of cancer, and some noting that the image is unclear and will not be understood by consumers. Some comments also criticized the design of the image, and one stated that the image is not presented in color as required by the Tobacco Control Act. Some comments also noted that this image of a burning cigarette could trigger cravings in smokers. It was also noted in the comments that the image failed to show desirable effects on some measures in FDA’s research study. One comment noted that the secondary message highlighted by the use of bold face emphasis in this proposed required warning (“I cause cancer”) could be interpreted as blaming smokers, and expressed concern that this could undermine the proposed required warning’s ability to communicate effectively with smokers.

(Comment 103) FDA received some comments that supported the use of the image “red cigarette burning,” including comments from individuals, a public health advocacy group, and from State and local public health agencies. These comments noted that the “red cigarette burning” image is particularly appropriate for use with the warning statement, or expressed the submitter’s preference that the image be used with this warning statement.

As discussed in section IIIC of this document, some comments submitted to the docket described the results of research conducted by the submitters to examine the potential effectiveness of FDA’s proposed images. This image was discussed in some of these comments. Specifically, in one submitter’s study, participants rated this image highly on its ease of comprehension. It also induced relatively greater worry and feelings of discouragement from wanting to smoke than a text-only control. In another submitter’s study, particular aspects of the image were evaluated, and the submitter reported that the use of the color red to accentuate the warning content in “red cigarette burning” was effective.

However, the image was rated as one of the least effective of the images proposed by FDA for use with this warning statement in another submitter’s study of the potential effectiveness of the images, and the image was rated as one of the five least effective images used in FDA’s 36 proposed required warnings in another submitter’s study of the potential effectiveness of the images.
FDA also received several comments that opposed use of the image “red cigarette burning,” with some submitters indicating that the image does not depict the negative health consequences of smoking or that the image is not appropriately evocative of cancer. Some comments also criticized the design of the image, with one stating that it looked like an image from a cigarette advertisement. Some comments also noted that this image of a burning cigarette could trigger cravings in smokers. It was also noted in the comments that the image failed to show desirable effects on some measures in FDA’s research study and showed some undesirable effects. Some comments also suggested that other cancers, including bladder cancer, should be added to the cancers listed in the image.

(Response) We are not selecting this image for use in a required warning and instead have selected the image “cancerous lesion on lip” for the reasons given in section III.D of this document.

5. “WARNING: Cigarettes Cause Strokes and Heart Disease”

As discussed in section III.D of this document, FDA selected the image “oxygen mask on man’s face” for use with the statement, “WARNING: Cigarettes cause strokes and heart disease.” FDA proposed three other images for use with this statement: “hand with oxygen mask,” which appears on pages 37 and 38 of the document “Proposed Required Warning Images;” “red lightning with heart,” which appears on pages 41 and 42 of the document “Proposed Required Warning Images;” and “man in pain with hand on chest,” which appears on pages 43 and 44 of the document “Proposed Required Warning Images.”

Hand with oxygen mask. In FDA’s research study, the image “hand with oxygen mask” had significant effects on all the salience measures (emotional reaction scale, cognitive reaction scale, and difficult to look at measure) in all three study populations (adults, young adults, and youth).

However, the selected image, “oxygen mask on man’s face,” also had significant effects on all the salience measures, and generally had numerically larger effects than this image on the emotional reaction scale and the difficult to look at measure.

Adults viewing the image “hand with oxygen mask” reported being less likely to quit smoking within the next month than adults viewing the text-only control.

FDA received a number of comments on this image, which the Agency has summarized and responded to in the following paragraphs.

(Comment 104) FDA received some comments that supported the use of the image “hand with oxygen mask,” including comments from individuals, a community organization, and State public health agencies. These comments noted that the “hand with oxygen mask” image is the best image for use with the warning statement or stated that the image was appropriate for use with this warning statement.

As discussed in section III.C of this document, some comments submitted to the docket described results of research conducted by the submitters to examine the potential effectiveness of FDA’s proposed images. This image was discussed in some of these comments. Specifically, this image was rated highly on its ease of comprehension and induced relatively greater worry and led to higher ratings of feeling discouraged from wanting to smoke than a text-only control in one submitter’s study. However, the image was rated as the least effective of the images proposed by FDA for use with this warning statement in another submitter’s study of the potential effectiveness of the images.

FDA also received several comments that opposed use of the image “hand with oxygen mask,” with some submitters indicating that the image is hard to understand or not appropriately compelling. Some comments also stated that the image would be more appropriate for use with a statement about lung-related health consequences (such as COPD). It was also noted in the comments that the image failed to show desirable effects on some measures in FDA’s research study and showed some undesirable effects.

(Response) We are not selecting this image for use in a required warning and instead have selected the image “oxygen mask on man’s face” for the reasons given in section III.D of this document.

Red lightning with heart. In FDA’s research study, the image “red lightning with heart” had significant effects on the emotional and cognitive reaction scales in all three study populations (adults, young adults, and youth). The image also had significant effects on the difficult to look at measure in adults and young adults.

However, the selected image, “oxygen mask on man’s face,” had significant effects on all the salience measures in all the study populations, and it generally had numerically larger effects than this image on the salience measures.

Among adults, young adults, and youth, the image “red lightning with heart” prompted higher correct recall of the warning statement at 1 week follow-up than the text-only control. However, the proposed required warning featuring this image prompted relatively lower recall of the image at baseline and at 1 week follow-up among youth than the selected image, “oxygen mask on man’s face.”

FDA received several comments on this image, which the Agency has summarized and responded to in the following paragraphs.

(Comment 105) FDA received a few comments that supported the use of the image “red lightning with heart,” including comments from State and local public health agencies, which noted that this image is appropriate for use with the warning statement.

As discussed in section III.C of this document, some comments submitted to the docket described results of research conducted by the submitter to examine the potential effectiveness of FDA’s proposed images. This image was discussed in some of these comments. Specifically, this image was rated highly on its ease of comprehension in one submitter’s study, but failed to show an effect on other study measures (worry, discouragement from smoking). The image was rated as one of the least effective of the images proposed by FDA for use with this warning statement in another submitter’s study of the potential effectiveness of the images.

FDA also received several comments that opposed use of the image “red lightning with heart,” with some submitters criticizing the design of the image, which was characterized as too conceptual and not easily understandable. Some comments also criticized the illustration style, stating that it does not have the impact a photograph would have, and would not compel or move viewers, and may inappropriately appeal to youth without discouraging them from smoking. It was also noted in the comments that the image failed to show desirable effects on some measures in FDA’s research study. Some comments also noted in the comments that the photoograph would have, and would not compel or move viewers, and may inappropriately appeal to youth without discouraging them from smoking. It was also noted in the comments that the image failed to show desirable effects on some measures in FDA’s research study.

(Response) We are not selecting this image for use in a required warning and instead have selected the image “oxygen mask on man’s face” for the reasons given in section III.D of this document.

Man in pain with hand on chest. In FDA’s research study, the image “man in pain with hand on chest” had significant effects on the emotional reaction scale in all three study populations (adults, young adults, and youth). The image also had significant effects on the difficult to look at measure in adults and young adults.

However, the selected image, “oxygen mask on man’s face,” had significant effects on all the salience measures in all the study populations, and it generally had numerically larger effects than this image on the salience measures.
adults viewing a hypothetical advertisement containing “man in pain with hand on chest.” The image also had significant effects on the difficult to look at measure in adults and youth. However, the selected image, “oxygen mask on man’s face,” had significant effects on all the salience measures in all the study populations, and had numerically larger effects than this image on the salience measures.

Among youth, the image “man in pain with hand on chest” prompted higher correct recall of the warning statement at 1 week follow-up than the text-only control. However, the proposed required warning featuring this image prompted relatively lower recall of the image at baseline among adults than “oxygen mask on man’s face.”

FDA received a number of comments on this image, which the Agency has summarized and responded to in the following paragraphs.

(Comment 106) FDA received several comments that supported the use of the image “man in pain with hand on chest,” including comments from individuals, public health advocacy groups, a health care professional, and a State public health agency. Several of these comments indicated that this image is the best choice for use with this warning statement, with some comments noting that the image appropriately shows how painful heart attacks can be.

As discussed in section III.C of this document, some comments submitted to the docket described results of research conducted by the submitters to examine the potential effectiveness of FDA’s proposed images. This image was discussed in some of these comments. Specifically, in one submitter’s study of the potential effectiveness of FDA’s proposed images, this image was rated highly on its ease of comprehension compared to a text-only control in one submitter’s study, but failed to show an effect on other study measures (worry, discouragement from smoking). The image was also rated as the most effective of the images proposed by FDA for use with this warning statement in another submitter’s study of the potential effectiveness of the images, but an image used in another country was rated significantly higher than either of FDA’s proposed images in this study (however, as discussed in section III.A of this document, at this time FDA does not believe it is necessary or appropriate to use graphic warnings used in other countries in place of the images developed by FDA).

FDA also received several comments that opposed use of the image “pacifier & ashtray,” with some submitters criticizing the design of the image, which was characterized as too symbolic and abstract to be understood, and as lacking in emotional impact. Some comments stated that the image does not show a health consequence of smoking, and some indicated the image is not graphic enough. A few comments also noted that the image would be more appropriate for a warning related to post-partum secondhand smoke-related risks, rather than a pregnancy warning, because pacifiers are used post-rather than pre-partum. One comment stated that the background used for the textual warning statement in the image looks unprofessional. It was also stated in the comments that the image failed to show large effects on the salience measures or to show desirable effects on some other measures in FDA’s research study.

(Comment 107) FDA received several comments that supported the use of the image “pacifier & ashtray,” including comments from individuals, public health advocacy groups, and State and local public health agencies. In general, these comments indicated that this image is the best choice for use with this warning statement, with some noting that the image is compelling and powerful.

As discussed in section III.C of this document, some comments submitted to the docket described the results of research conducted by the submitters to examine the potential effectiveness of FDA’s proposed images. This image was discussed in some of these comments. Specifically, this image was rated highly on its ease of comprehension compared to a text-only control in one submitter’s study, but failed to show an effect on other study measures (worry, discouragement from smoking). The image was also rated as the most effective of the images proposed by FDA for use with this warning statement in another submitter’s study of the potential effectiveness of the images, but an image used in another country was rated significantly higher than either of FDA’s proposed images in this study (however, as discussed in section III.A of this document, at this time FDA does not believe it is necessary or appropriate to use graphic warnings used in other countries in place of the images developed by FDA).

As discussed in section III.D of this document, FDA selected the image “baby in incubator” for use with the statement, “WARNING: Smoking during pregnancy can harm your baby.” FDA proposed one other image for use with this statement: “pacifier & ashtray,” which appears on pages 47 and 48 of the document “Proposed Required Warning Images.”

Pacifier & ashtray. In FDA’s research study, the image “pacifier & ashtray” had significant effects on the emotional and cognitive reaction scales in all three study populations (adults, young adults, and youth). The image also had significant effects on the difficult to look at measure in adults and youth. However, the selected image, “baby in incubator,” had significant effects on all the salience measures in all the study populations, and had numerically larger effects than this image on all the salience measures.

Among young adults, the image “pacifier & ashtray” prompted higher correct recall of the warning statement at baseline and at 1 week follow-up than the text-only control. However, the proposed required warning featuring this image prompted relatively lower recall of the image at baseline and at 1 week follow-up among adults, young adults, and youth than pre-partum. One comment stated that the background used for the textual warning statement in the image looks unprofessional. It was also stated in the comments that the image failed to show large effects on the salience measures or to show desirable effects on some other measures in FDA’s research study.

(Comment 107) FDA received several comments that supported the use of the image “pacifier & ashtray,” including comments from individuals, public health advocacy groups, and State and local public health agencies. In general, these comments indicated that this image is the best choice for use with this warning statement, with some noting that the image is compelling and powerful.

As discussed in section III.C of this document, some comments submitted to the docket described the results of research conducted by the submitters to examine the potential effectiveness of FDA’s proposed images. This image was discussed in some of these comments. Specifically, this image was rated highly on its ease of comprehension compared to a text-only control in one submitter’s study, but failed to show an effect on other study measures (worry, discouragement from smoking). The image was also rated as the most effective of the images proposed by FDA for use with this warning statement in another submitter’s study of the potential effectiveness of the images, but an image used in another country was rated significantly higher than either of FDA’s proposed images in this study (however, as discussed in section III.A of this document, at this time FDA does not believe it is necessary or appropriate to use graphic warnings used in other countries in place of the images developed by FDA).

As discussed in section III.D of this document, FDA selected the image “baby in incubator” for use with the statement, “WARNING: Smoking during pregnancy can harm your baby.”

FDA also received several comments that opposed use of the image “pacifier & ashtray,” with some submitters criticizing the design of the image, which was characterized as too symbolic and abstract to be understood, and as lacking in emotional impact. Some comments stated that the image does not show a health consequence of smoking, and some indicated the image is not graphic enough. A few comments also noted that the image would be more appropriate for a warning related to post-partum secondhand smoke-related risks, rather than a pregnancy warning, because pacifiers are used post-rather than pre-partum. One comment stated that the background used for the textual warning statement in the image looks unprofessional. It was also stated in the comments that the image failed to show large effects on the salience measures or to show desirable effects on some other measures in FDA’s research study.

(Comment 107) FDA received several comments that supported the use of the image “pacifier & ashtray,” including comments from individuals, public health advocacy groups, and State and local public health agencies. In general, these comments indicated that this image is the best choice for use with this warning statement, with some noting that the image is compelling and powerful.

As discussed in section III.C of this document, some comments submitted to the docket described the results of research conducted by the submitters to examine the potential effectiveness of FDA’s proposed images. This image was discussed in some of these comments. Specifically, this image was rated highly on its ease of comprehension compared to a text-only control in one submitter’s study, but failed to show an effect on other study measures (worry, discouragement from smoking). The image was also rated as the most effective of the images proposed by FDA for use with this warning statement in another submitter’s study of the potential effectiveness of the images, but an image used in another country was rated significantly higher than either of FDA’s proposed images in this study (however, as discussed in section III.A of this document, at this time FDA does not believe it is necessary or appropriate to use graphic warnings used in other countries in place of the images developed by FDA).
on pages 55 and 56 of the document “Proposed Required Warning Images.”

Red coffin with body. In FDA’s research study, the image “red coffin with body” had significant effects on all the salience measures (emotional reaction scale, cognitive reaction scale, and difficult to look at measure) in adults and youth. It also had a significant effect on the cognitive reaction scale in young adults.

However, the selected image, “man with chest staples,” had a significant effect on all the salience measures in all study populations, and had numerically larger effects than this image on these measures.

Among adults, the image “red coffin with body” prompted higher correct recall of the warning statement at baseline than the text-only control.

The image also had a significant impact on beliefs about the health risks of smoking for smokers relative to the text-only control in the adult sample that viewed a hypothetical advertisement containing the proposed required warning.

FDA received a number of comments on this image, which the Agency has summarized and responded to in the following paragraphs.

(Comment 108) FDA received several comments that supported the use of the image “red coffin with body,” including comments from individuals and a community organization. Several of these comments indicated that this image is the best choice for use with this warning statement, with some approving of the colors used in the image and some noting that the image gets the message across in a straightforward manner.

As discussed in section III.C of this document, some comments submitted to the docket described the results of research conducted by the submitters to examine the potential effectiveness of FDA’s proposed images. This image was discussed in some of these comments. Specifically, this image was rated highly on its ease of comprehension compared to a text-only control in one submitter’s study, but failed to show an effect on other study measures (worry, discouragement from smoking). The image was rated as one of the least effective of the images proposed by FDA for use with this warning statement in another submitter’s study of the potential effectiveness of the images.

FDA also received several comments that opposed use of the image “red coffin with body,” with some submitters stating that the image is too conceptual and not easily understandable. Several comments stated that the image is not impactful and is unlikely to be effective, with some indicating the image would be more effective if it were a photograph of an actual person. It was also suggested in the comments that the image style may inappropriately appeal to youth without discouraging them from smoking. Some comments noted that the image failed to show desirable effects on some measures in FDA’s research study.

(Response) We are not selecting this image for use in a required warning and instead have selected the image “man with chest staples” for the reasons given in section III.D of this document.

Man in casket. In FDA’s research study, the image “man in casket” had significant effects on all the salience measures (emotional reaction scale, cognitive reaction scale, and difficult to look at measure) in adults and youth. It also had a significant effect on the cognitive reaction scale in young adults.

However, the selected image, “man with chest staples,” had significant effects on all the salience measures, and generally had numerically larger effects than this image on these measures.

Among youth, the image “man in casket” prompted higher correct recall of the warning statement at baseline than the text-only control. However, among young adults, the image “man in casket” prompted lower correct recall of the warning statement at baseline than the text-only control.

The image also had a significant impact on beliefs about the health risks of smoking for smokers relative to the text-only control in the adult sample that viewed a hypothetical advertisement containing the proposed required warning.

FDA received a number of comments on this image, which the Agency has summarized and responded to in the following paragraphs.

(Comment 109) FDA received several comments that supported the use of the image “man in casket,” including comments from individuals, a public health advocacy group, and a State public health agency. Several of these comments indicated that this image is the best choice for use with this warning statement, with some noting that the image grabs viewers’ attention and clearly depicts death.

As discussed in section III.C of this document, some comments submitted to the docket described the results of research conducted by the submitters to examine the potential effectiveness of FDA’s proposed images. This image was discussed in some of these comments. Specifically, in one submitter’s study, participating image highly on its ease of comprehension. It also induced relatively greater worry and feelings of discouragement from wanting to smoke than a text-only control. In another submitter’s study, particular aspects of the image were evaluated, and the proposed required warning containing the image “man in casket” was found to be significantly more effective at discouraging others from smoking than a text-only statement on the side of a cigarette package. However, the image was rated as less effective than the selected image, “man with chest staples,” in another submitter’s study of the potential effectiveness of the images.

FDA also received several comments that opposed use of the image “man in casket.” Multiple comments stated the image looks staged because the man pictured does not look like he is dead or like he suffered from smoking-related disease. It was also suggested in the comments that the image may not be understood by all cultures. The image was also criticized as lacking a clear association to smoking. It was also noted in the comments that the image failed to show desirable effects on some measures in FDA’s research study.

(Response) We are not selecting this image for use in a required warning and instead have selected the image “man with chest staples” for the reasons given in section III.D of this document.

Cigarettes = RIP. In FDA’s research study, the image “cigarettes = RIP” had significant effects on all the salience measures (emotional reaction scale, cognitive reaction scale, and difficult to look at measure) in adults and youth. It also had a significant effect on the emotional and cognitive reaction scales in young adults.

However, the selected image, “man with chest staples,” had significant effects on all the salience measures in all the study populations, and generally had numerically larger effects than this image on these measures.

Among adults, the image “cigarettes = RIP” prompted higher correct recall of the warning statement at baseline than the text-only control. However, the proposed required warning featuring this image prompted relatively lower recall of the image at baseline and at 1 week follow-up than the selected image, “man with chest staples.”

The image had a significant impact on beliefs about the health risks of smoking for smokers relative to the text-only control in the adult sample that viewed a hypothetical advertisement containing the proposed required warning.

FDA received a number of comments on this image, which the Agency has summarized and responded to in the following paragraphs.

Graveyard. In FDA’s research study, the image “graveyard” had significant effects on the emotional reaction scale in all three study populations (adults, young adults, and youth). The image also had significant effects on the cognitive reaction scale in young adults and youth, and on the difficult to look at measure in youth.

However, the selected image, “woman crying,” had significant effects on the salience measures in all study populations, and it generally had numerically larger effects than this image on all the salience measures. Among adults and youth, the image “graveyard” prompted lower correct recall of the warning statement at baseline than the text-only control. Among young adults, the image prompted lower correct recall of the warning statement at 1 week follow-up than the text-only control.

The image “graveyard” had a significant impact on beliefs about the health risks of smoking for smokers in young adults.

FDA received a number of comments on this image, which the Agency has summarized and responded to in the following paragraphs.

(Comment 111) FDA received several comments that supported the use of the image “graveyard,” including comments from individuals, a community organization, and a State public health agency. Several of these comments indicated that this image is the best choice for use with this warning statement, with some noting that the image gets the message across in a straightforward manner, and some noting the image could deter people from starting to smoke.

As discussed in section III.C of this document, some comments submitted to the docket described the results of research conducted by the submitters to examine the potential effectiveness of FDA’s proposed images. This image was discussed in some of these comments. Specifically, in one submitter’s study, the image was highly effective on its ease of comprehension. It also induced relatively greater worry and feelings of discouragement from wanting to smoke than a text-only control. This image was also rated as the most effective of the images proposed by FDA for use with this warning statement in another submitter’s study of the potential effectiveness of the images, although an image used in another country was rated more highly than this image.

FDA also received several comments that opposed use of the image “graveyard.” Some comments indicated that the image would not be effective, noting that it is easy to disregard or, alternatively, too sensational to be effective. It was also stated in the comments that the image did not show large impacts on the emotional reaction scale and failed to show desirable effects on some other measures in FDA’s research study.

(Comment 112) FDA received several comments that supported the use of the image “woman crying,” including comments from individuals and State public health agencies. Several of these comments indicated that this image is the best choice for use with this warning statement, with some noting that the image would make smokers think about how their habit may cause others to avoid them. It was also noted that the image effectively shows how innocent bystanders are affected by smokers.
As discussed in section III.C of this document, some comments submitted to the docket described the results of research conducted by the submitters to examine the potential effectiveness of FDA’s proposed images. This image was discussed in some of these comments. Specifically, in one submitter’s study, participants rated this image highly on its ease of comprehension. It also induced relatively greater worry and feelings of discouragement from wanting to smoke than a text-only control. The submitter also concluded that the image was the most effective of the images proposed for use with this warning statement. However, the image was rated as one of the less effective images proposed by FDA for use with this warning statement in another submitter’s study of the potential effectiveness of the images.

FDA also received several comments that opposed use of the image “man smoke at woman.” Some comments indicated that the image is not realistic, stating that smokers do not blow smoke at their friends. One comment indicated that the image failed to portray an obvious health consequence of secondhand smoke, and multiple comments indicated that the image conveyed a bad message by showing the nonsmoker covering her nose and mouth, stating that these actions do not protect you from secondhand smoke. It was also noted in the comments that the image failed to show desirable effects on some measures in FDA’s research study.

(Response) We are not selecting this image for use with this required warning and instead have selected the image “woman crying” for the reasons given in section III.D of this document.

“Woman smoke at man.” In FDA’s research study, the image “woman smoke at man” had significant effects on the emotional reaction scale in adults, young adults, and youth. The image also had significant effects on the cognitive reaction scale in young adults and youth, and on the difficult to look at measure in adults and youth.

However, the selected image, “woman crying,” had significant effects on the salience measures in all study populations, and it had numerically larger effects than this image on the emotional reaction scale in all study populations (adults, young adults, and youth) and on the cognitive reaction scale in young adults and youth.

However, the selected image, “woman crying,” had significant effects on all the salience measures in all study populations, and it had numerically larger effects than this image on all these measures.

The proposed required warning featuring the image “man hands up & smoke” also prompted relatively lower correct recall of the image at baseline and at 1 week follow-up than the selected image, “woman crying.”

The image “woman smoke at man” had a significant impact on young adult’s intentions to quit smoking in the next month compared to the text-only control.

FDA received a number of comments on this image, which the Agency has summarized and responded to in the following paragraphs.

(Comment 113) FDA received several comments that supported the use of the image “woman smoke at man,” including comments from individuals, a public health advocacy group, a medical organization, and State and local public health agencies. Several of these comments indicated that this image is the best choice for use with this warning statement, with some noting that the image will make smokers think about how their actions negatively affect social situations.

As discussed in section III.C of this document, some comments submitted to the docket described the results of research conducted by the submitters to examine the potential effectiveness of FDA’s proposed images. This image was discussed in some of these comments. Specifically, this image was rated highly on its ease of comprehension compared to a text-only control in one submitter’s study but failed to show an effect on other study measures (worry, discouragement from smoking). The image was rated as one of the least effective of the images proposed by FDA for use with this warning statement in another submitter’s study of the potential effectiveness of the images.

FDA also received several comments that opposed use of the image “woman smoke at man.” Some comments indicated that the image would not be effective, suggesting that it is not impactful and probably would not stop people from smoking. One comment indicated that the image fails to portray an obvious health consequence of secondhand smoke, and another was critical of the actions of the nonsmoker in the image, noting that covering your nose and mouth does not protect you from secondhand smoke. It was also stated in the comments that the image failed to show desirable effects on some measures in FDA’s research study.

(Comment 114) FDA received some comments that supported the use of the image “man hands up & smoke,” including comments from individuals and a State public agency. These comments generally indicated that this image would be the best choice for use with this warning statement.

As discussed in section III.C of this document, some comments submitted to the docket described the results of research conducted by the submitters to examine the potential effectiveness of FDA’s proposed images. This image was discussed in some of these comments. Specifically, this image was rated highly on its ease of comprehension compared to a text-only control in one submitter’s study but failed to show an effect on other study measures (worry, discouragement from smoking). The image was rated as the least effective of the images proposed by FDA for use with this warning statement in another submitter’s study of the potential effectiveness of the images.

FDA also received several comments that opposed use of the image “man hands up & smoke.” Some comments indicated that the image is unrealistic in that it looks like the man is in fog or a house fire as opposed to being affected by secondhand smoke. One comment indicated that the image does not portray a health consequence of secondhand smoke; it was also stated in the comments the image is ineffective and unintentionally humorous. One comment stated that the image failed to show large effects on salience measures or to show desirable effects on other measures in FDA’s research study and indicated it should not be selected.

(Comment 14) FDA received several comments that supported use of the image “woman smoke at man,” including comments from individuals and a medical organization. These comments generally indicated that this image would be the best choice for use with this warning statement.

As discussed in section III.C of this document, some comments submitted to the docket described the results of research conducted by the submitters to examine the potential effectiveness of FDA’s proposed images. This image was discussed in some of these comments. Specifically, this image was rated highly on its ease of comprehension compared to a text-only control in one submitter’s study but failed to show an effect on other study measures (worry, discouragement from smoking). The image was rated as the least effective of the images proposed by FDA for use with this warning statement in another submitter’s study of the potential effectiveness of the images.

FDA also received several comments that supported use of the image “woman smoke at man,” including comments from individuals and a State public agency. These comments generally indicated that this image would be the best choice for use with this warning statement.

As discussed in section III.C of this document, some comments submitted to the docket described the results of research conducted by the submitters to examine the potential effectiveness of FDA’s proposed images. This image was discussed in some of these comments. Specifically, this image was rated highly on its ease of comprehension compared to a text-only control in one submitter’s study but failed to show an effect on other study measures (worry, discouragement from smoking). The image was rated as the least effective of the images proposed by FDA for use with this warning statement in another submitter’s study of the potential effectiveness of the images.

FDA also received several comments that opposed use of the image “woman smoke at man.” Some comments indicated that the image would not be effective, suggesting that it is not impactful and probably would not stop people from smoking. One comment indicated that the image fails to portray an obvious health consequence of secondhand smoke, and another was critical of the actions of the nonsmoker in the image, noting that covering your nose and mouth does not protect you from secondhand smoke. It was also stated in the comments that the image failed to show desirable effects on some measures in FDA’s research study.
9. “WARNING: Quitting Smoking Now Greatly Reduces Serious Risks to Your Health”

As discussed in section III.D of this document, FDA selected the image “man I Quit t-shirt” for use with the statement, “WARNING: Quitting smoking now greatly reduces serious risks to your health.” FDA proposed two other images for use with this statement: “cigarettes in toilet bowl,” which appears on pages 69 and 70 of the document “Proposed Required Warning Images;” and “woman blowing bubble,” which appears on pages 71 and 72 of the document “Proposed Required Warning Images.”

Cigarettes in toilet bowl. In FDA’s research study, the image “cigarettes in toilet bowl” had significant effects on the emotional reaction scale in adults and young adults and significant effect on the cognitive reaction scale in all study populations (adults, young adults, and youth).

Among youth, the image “cigarettes in toilet bowl” prompted higher correct recall of the warning statement at 1 week follow-up than the text-only control. However, the proposed required warning featuring this image prompted relatively lower recall of the image at baseline and at 1 week follow-up than the selected image, “man I Quit t-shirt.”

The image “cigarettes in toilet bowl” had a significant impact on beliefs about the health risks of smoking for smokers in young adults.

FDA received a number of comments on this image, which the Agency has summarized and responded to in the following paragraphs.

(Comment 115) FDA received some comments that supported the use of the image “cigarettes in toilet bowl,” including comments from individuals, a community organization, and a local public health agency. Some comments noted that this image is the best choice for use with this warning statement, and it was also noted in the comments that the image is effective because it creates an association between cigarettes and other undesirable things that belong in a toilet bowl.

As discussed in section III.C of this document, some comments submitted to the docket described the results of research conducted by the submitters to examine the potential effectiveness of FDA’s proposed images. This image was discussed in some of these comments. Specifically, this image failed to show any significant effects in one submitter’s study on measures of ease of comprehension, worry, and feeling discouraged from smoking compared to a text-only control. In addition, the image was rated as less effective than the selected image, “man I Quit t-shirt,” in another submitter’s study of the potential effectiveness of the images.

FDA also received several comments that opposed use of the image “cigarettes in toilet bowl.” These comments noted that the image is not clear or does not convey a health consequence of smoking. It was also noted that the image is not easily understood, or alternatively, that it is banal. Multiple comments expressed concern about what is shown in the image, stating that it recommends a bad or unhealthy action (i.e., flushing cigarettes down a toilet, which the comments stated could clog the toilet and pollute the environment). Some comments also stated that the statement was difficult to read in the “cigarettes in toilet bowl” image. It was also stated in the comments that the image did not show large effects on the emotional and cognitive reaction scales in FDA’s research study and failed to show desirable effects on other measures. (Response) We are not selecting this image for use in a required warning and instead have selected the image “man I Quit t-shirt” for the reasons given in section III.D of this document.

Woman blowing bubble. In FDA’s research study, the image “woman blowing bubble” had a significant effect on the cognitive reaction scale in youth. The image “woman blowing bubble” had a negative impact on youth beliefs about the health risks of smoking for smokers and for nonsmokers (i.e., youth who viewed this image were less likely to believe that smokers will suffer negative health consequences or that nonsmokers exposed to secondhand smoke will suffer negative health consequences than youth who viewed the text-only control). Furthermore, the adult sample that viewed a hypothetical advertisement containing the proposed required warning reported that they were less likely to quit smoking in the next 30 days compared to adults who viewed the text-only control.

(Comment 116) FDA received some comments that supported the use of the image “woman blowing bubble,” including comments from individuals, a public health advocacy group, and a State public health agency. Multiple comments noted that the image appropriately shows how quitting smoking allows for a better lung capacity or noted that it effectively conveys the idea that there are beneficial effects of quitting.

As discussed in section III.C of this document, some comments submitted to the docket described the results of research conducted by the submitters to examine the potential effectiveness of FDA’s proposed images. This image was discussed in some of these comments. Specifically, this image led to lower levels of worry and lower reports of feeling discouraged from smoking relative to a text-only control in one submitter’s study. In addition, the image was rated as the least effective of the images proposed by FDA for use with this warning statement in another submitter’s study of the potential effectiveness of the images.

FDA also received a number of comments that opposed use of the image “woman blowing bubble.” Multiple comments stated that the image is confusing and is not appropriately compelling and would not be effective in encouraging smokers to quit. Some comments indicated that the image does not effectively convey the message contained in the warning statement, and some noted that the image is banal or, alternatively, too positive. Multiple comments also stated the image is hard to understand, and that smokers may not comprehend the association between the image and the warning statement. It was also stated that the image would inappropriately appeal to youth without discouraging them from smoking, and that the image is inappropriate because it is sexually suggestive. It was also noted in the comments that the image showed negative results on some measures in FDA’s research study, and failed to show desirable effects on other measures. (Response) We are not selecting this image for use in a required warning and have instead selected the image “man I Quit t-shirt” for the reasons given in section III.D of this document.

10. Image for Advertisements With a Small Surface Area

As discussed in section III.D of this document, FDA selected the image which appears on page 75 of the document entitled “Proposed Required Warning Images” for use with the textual warning statements solely in advertisements with a small surface area (defined as less than 12 square inches). We also proposed one other image for use with this statement, which appears on page 74 of the document entitled “Proposed Required Warning Images.” The proposed image on page 74 depicts a burning cigarette enclosed by a red circle with a red bar across the image. We did not receive any comments on either of the proposed images.

As explained in section III.D of this document, we have selected the image of a black exclamation mark enclosed within a red equilateral triangle for use...
in advertisements with a small surface area because we have concluded that the common purpose of this image, to denote a warning of a threat to health or of a hazard which could result in personal injury, makes it the most appropriate for use in the required warning context.

IV. Comments Regarding Textual Warning Statements

A. Changes to Textual Warning Statements

As we explained in the proposed rule, section 202(b) of the Tobacco Control Act, amending section 4 of FCLAA (15 U.S.C. 1333), gives us the authority to adjust the format, type size, color graphics, and text of any of the required warning statements if such a change “would promote greater public understanding of the risks associated with the use of tobacco products.” In addition, under section 4(d) of FCLAA, FDA may adjust the type size, text, and format of the warning statements as the Agency determines appropriate “so that both the graphics and the accompanying label statements are clear, conspicuous, legible and appear within the specified area.” Such adjustments, including adjustments to the text and format of some of the warning statements, were included with some of the proposed warnings (75 FR 69524 at 69534). We did not receive comments about these adjustments. Two of the warning statements we have selected for this final rule are presented in all uppercase letters, as they were in the proposal. In addition, one of the proposed required warnings, “baby in incubator,” was presented without the signal word “WARNING.” The research literature on graphic health warnings indicates that signal words, such as “Warning,” have been found to enhance the noticeability of safety warnings and convey the degree of risk (see Ref. 40 at p. 33). In the final rule, we are thus not removing the word “WARNING” from this required warning, such that the text in this required warning is the same as the text presented in section 201 of the Tobacco Control Act (“WARNING: Smoking during pregnancy can harm your baby”).

Moreover, section 906(d) of the FD&C Act (21 U.S.C. 387f(d)) authorizes FDA to issue regulations restricting the sale or distribution of cigarettes and other tobacco products. As is discussed in more detail in section V.B.6 of this document, a reference to a cessation resource has been included in the final required warnings. Although we did not receive any comments about the adjustments we made to the text of some of the warning statements in the 36 proposed required warnings, we received numerous comments requesting other changes to the textual statements for the new required warnings, including requests to strengthen the text, to add additional information to the text or to otherwise modify the text of the warnings statements. We also received requests to substitute alternative warning statements for some or all of the textual statements and to expand the warning statements by adding additional statements regarding smoking-related risks. The comments, and our responses, are summarized in the following paragraphs. We also received numerous comments about our proposal to include a reference to a cessation resource in the required warnings; these comments and our responses are summarized in section V.B.6 of this document.

(Comment 117) Several comments suggested that some of the textual warning statements should be changed to include language asserted to be stronger and more direct. For example, multiple comments suggested that the statement, “WARNING: Tobacco smoke can harm your children,” should be reworded to be more assertive, for example, to state “Tobacco smoke harms your children.” One comment referenced the conclusion from the 2010 Surgeon General’s report that there is no risk-free level of exposure to secondhand smoke as support for this modification (Ref. 37). Similarly, multiple comments recommended that FDA change the warning statement, “WARNING: Smoking during pregnancy can harm your baby,” to be more strongly worded. For instance, comments suggested this statement could instead be worded as “WARNING: Smoking during pregnancy harms your baby” or “WARNING: Smoking when pregnant harms your baby” or “WARNING: Smoking harms your baby” or “WARNING: Smoking harms the fetus and babies.” Multiple comments also suggested the warning statement “WARNING: Smoking can kill you” should not be worded in a conditional manner. One comment suggested that the text could instead state “Smoking kills.”

(Comment 118) Many comments recommended that FDA include additional textual information to give further context for the health warnings. For example, comments requested that FDA add information such as research statistics, factual testimonials, or other explanatory text to further enhance the effectiveness of the new required warnings. Several of the comments suggested specific text for particular warning statements; for example, one comment suggested the warning

HIGHLY addictive,” while another comment suggested the statement read “Cigarettes are addictive and shorten your life.” Similarly, a comment from a health care professional suggested the warning should state “Cigarettes are addictive and deadly.” Another comment from a nonprofit foundation suggested that the statement “Cigarettes cause strokes and heart disease” be modified to state “Cigarettes cause strokes, heart disease, and amputations.”
statement related to addiction be accompanied by the following explanatory text: “Studies have shown that tobacco can be harder to quit than heroin or cocaine.” Other comments suggested that the statement “WARNING: Cigarettes cause cancer” be modified to add explanatory text about specific cancers caused by cigarettes, including cancers of the mouth, throat, esophagus, lungs, kidney, bladder, pancreas, stomach, cervix, and bone marrow. Another comment suggested that the statement “Cigarettes cause strokes and heart disease” be accompanied by explanatory text stating “Cigarette smoking doubles your chances of strokes and can cause heart attacks” and that the statement “Cigarettes cause fatal lung disease” be accompanied by explanatory text stating that “Every cigarette you smoke increases your chances of dying from lung disease.” In addition, the comment suggested that the statement “Tobacco smoke causes fatal lung disease in nonsmokers” be accompanied by explanatory text stating “You’re not the only one smoking cigarettes. The smoke is not just inhaled by smokers, it becomes second-hand smoke, which contains more than 50 cancer agents.” Another comment suggested adding information to the required warnings that state alternatives to smoking, such as exercise and healthy eating.

(Response) We decline to make such changes at this time. As stated previously, the nine new textual warning statements mandated by Congress in section 4(a)(1) of FCLAA objectively communicate some of the major health risks associated with smoking in an effective manner. In addition, research has shown that warning statements that are short and to the point and that are presented in larger fonts sizes are likely to be more effective (Ref. 40 at p. 33). If the additional requested information were added to the required warnings, the resulting warning statements would be longer, and the font size of the warning statements would likely decrease in order for them to fit within the specified area. This could undercut the effectiveness of the warnings (see, e.g., Ref. 57). If research later indicates that adding such information to the new required warnings will promote a greater understanding of the risks associated with smoking, we will consider making these changes using our authority under section 202(b) of the Tobacco Control Act.

(Comment 119) One comment suggested that the warning statements that reference “tobacco smoke” should be modified to instead reference “cigarette smoke” to apply more directly to the target audience.

(Response) We disagree that this change is warranted. The statements in section 4(a)(1) of FCLAA, including those that reference “tobacco smoke,” are scientifically accurate, and we do not believe that consumers will fail to understand that the warning statements referencing “tobacco smoke” apply to the products on which they appear (i.e., cigarettes), which are tobacco products.

(Comment 120) FDA received a number of comments suggesting that some of the negative health effects that are the subject of individual warning statements be replaced with other warnings. For example, one comment from a medical organization suggested that the statement “WARNING: Tobacco smoke causes fatal lung disease in nonsmokers” should instead focus on heart attacks, stating that the magnitude of fatal heart disease caused by secondhand smoke exposure is greater than the magnitude of fatal lung disease caused by smoke exposure. One comment from an individual suggested that FDA use other warnings about the health harms of smoking instead of the warning about addiction.

Another comment suggested that there should be fewer warnings regarding the health risks of secondhand smoke to babies and children and more warnings directed at young teens and pre-teens. One comment stated that the warnings about smoking during pregnancy and about the harms of tobacco smoke to children are only relevant to those who are pregnant or who have children and suggested that these warnings are thus less impactful than the other warning statements.

However, other comments stated that the warnings about the risks of smoking during pregnancy and about the health risks of secondhand smoke to children address important health issues, will help make smokers aware that they are harming innocent people around them, and will help smokers appreciate the severity and magnitude of some of the lesser-known risks of smoking. One comment from an individual noted that secondhand smoke kills an estimated 45,000 nonsmokers who live with smokers from heart disease each year, as well as increasing the risk of SIDS, acute respiratory infections, ear problems, and severe asthma in children, and causing respiratory symptoms and slowing lung growth in children.

(Response) We decline to amend the warning statements as suggested by the comments. As stated previously, the nine text warnings provided by Congress in section 4(a)(1) of FCLAA appropriately communicate important health risks of smoking. Furthermore, we disagree with the suggestion that there should be fewer warnings about the health risks of smoking during pregnancy and of secondhand smoke to children. These warnings comprise two of the nine warning statements, and we agree with the comments indicating that these warnings communicate information about important health issues and will help smokers understand some of the significant health harms caused by cigarettes. In addition, while these warnings may be especially impactful with parents and expectant parents, using a variety of messages, including messages that may particularly impact certain audiences, will strengthen the overall impact of the required warnings (Ref. 40 at pp. 7–8).

Similarly, we disagree with the suggestion that the warning about addiction should be replaced by a warning about other health hazards. As discussed in the preamble to the proposed rule (75 FR 69524 at 69528 through 69529), the magnitude of public health harm caused by cigarettes is inextricably linked to the addictive nature of these products (Ref. 16 at p. 14 and Ref. 3 at p. xi), and many people, particularly adolescents, have a poor understanding of how difficult it is to quit smoking due to the addictive nature of cigarettes (Ref. 3 at p. 91). Thus, we conclude this is an important and appropriate health warning.

(Comment 121) One comment suggested that graphic health warnings on cigarette packages and advertisements should have one broad warning that states: “Cigarette smoking may cause cancer, death, and other serious life-threatening health hazards.” Another comment suggested one broad warning that states: “Smoking Can Kill You.”

(Response) We disagree. We are not aware of any scientific evidence that one broad warning statement would be more effective in communicating the multitude of health risks to smokers and nonsmokers in all age categories than the nine specific textual warnings specified in section 4(a) of FCLAA. As noted in the proposed rule, evidence shows that warnings about specific health risks, such as cancer, heart disease, and stroke, are more effective than general warnings (75 FR 69524 at 69533 through 69534).

Utilizing a single broad statement like the ones proposed in the comments would also fail to communicate important information about the detrimental effects associated with secondhand smoke—and messages about secondhand smoke have been effective in moving smokers to consider
the health risks associating with smoking (75 FR 69524 at 69534). For example, the new set of warnings includes the following statement: “WARNING: Tobacco smoke causes fatal lung disease in non-smokers.” This important warning would be lost if we chose to use just one of the suggested broad warning statements. In addition, one of the new required warnings clearly notifies smokers that if they quit smoking, they can greatly reduce serious risks to their health. Again, that important message would be lost if we were to use just one of the suggested broad statements.

[Comment 122] One comment stated that the ninth warning statement provided by Congress in the Tobacco Control Act. “WARNING: Quitting smoking now greatly reduces serious risks to your health,” should appear on all packages after one of the other eight warning statements.  

(Response) We disagree that such a change is warranted. As discussed in section V.B.6 of this document, we have included a reference to a cessation resource in the required warnings, which we conclude is more appropriate than including the ninth warning statement in all the required warnings.

[Comment 123] Many comments suggested that FDA add additional warning statements to state that cigarette smoking may increase the risk of other diseases such as bladder cancer, impotence, blindness, or COPD. One comment stated that medical studies have shown that women who smoke a pack of cigarettes a day double the risk of orofacial cleft birth defects in their children, and suggested that a warning be added to include this risk and pictures of children with this birth defect (citing, e.g., Ref. 58). One comment also suggested that the required warnings indicate that smoking may increase the risk of breast cancer. Another comment suggested including messages about short-term effects of smoking, such as nutritional deficiencies.

(Response) We decline to add additional warning statements, as suggested in these comments. At this point, we have determined the nine textual statements mandated by Congress in section 4(a)(1) of FCLAA appropriately communicate major health risks of smoking. As stated previously, we intend to monitor the effects of these required warnings once they are put into use. We will conduct research and keep abreast of scientific developments regarding the efficacy of various required warnings and the types and elements of various warnings that improve efficacy. We intend to use the results of our monitoring and such research to determine whether changes should be made to the nine textual statements in a future rulemaking. We recognize that cigarettes cause negative health consequences in both smokers and non-smokers beyond those addressed in the nine warning statements provided by Congress, and will take this into account in making future determinations as to whether the textual statements should be revised by rulemaking.

[Comment 124] A few comments also suggested that when FDA initiates a new rulemaking to establish its next set of graphic warnings, the Agency should consider adding health warnings that refer to other smoking-related diseases that are not specifically mentioned in this first set of required warnings.

(Response) We intend to periodically review the required warnings to assess their effectiveness and determine whether the warnings are suffering from wear out. During this review, we will re-examine the scientific literature and possibly conduct our own research to determine if additional textual warnings about the scientifically documented negative health consequences of smoking are appropriate.

[Comment 125] One comment suggested that FDA utilize different warnings with featured messages targeted to specific audiences based on their different attitudes and beliefs. As an example, this comment pointed to the Canadian health warning directed at young males, which stresses that tobacco can make the smoker impotent (Ref. 55).

(Response) We conclude that the nine textual statements required by Congress in section 4(a)(1) of FCLAA are appropriate. In addition, we have selected color graphics to accompany the new warning statements that use a variety of different fonts, typography, and layouts; depict a variety of human subjects; and use a variety of styles, including photographic and graphic illustrations. The required warnings will reach a wide variety of audiences, including youth, young adult, and adult smokers and non-smokers. For information on FDA’s selection of images, see section III of this document. As previously stated, we intend to monitor the effectiveness of these required warnings once they are put into use. If our monitoring finds that the messages are not reaching an appropriately broad population and that targeted messages would be more effective, we will consider revising the textual statements in a future rulemaking.

[Comment 126] One comment suggested that FDA require a standard pack size and shape, which would help to ensure the readability of warnings. (Response) We do not believe it is necessary to adopt a standard pack size and shape. We have taken steps to ensure that the required warnings will be conspicuous and legible on cigarette packages and in advertisements.

B. Attribution to the Surgeon General

Section 4(a)(1) of FCLAA contains the nine new textual warning statements that, when combined with the Surgeon General’s image, comprise the required warning. Congress did not include an attribution to the Surgeon General in the new textual warning statements, as it has done in past laws on cigarette health warnings. Accordingly, when we issued our proposed rule and released the 36 proposed required warnings, the textual warning statements did not include a reference to the Surgeon General. A number of comments, including those from former Surgeons General and Commissioned Public Health Service Officers, questioned why the new health warnings no longer contain any attribution to the “Surgeon General.” A summary of the comments and our response regarding this issue is included in the following paragraphs.

[Comment 127] The comments noted that, since Surgeon General Luther Terry’s 1964 report highlighting the adverse health effects of tobacco use, the Office of the Surgeon General has been inextricably linked to smoking prevention and that the reduction in smoking rates since the initial report and the advent of the first Surgeon General’s warning is due to the public confidence associated with the Surgeon General’s recommendations. In addition, they claimed that the new warnings would be less effective without the Surgeon General attribution. Two other comments also suggested that FDA include “the federal government logo” on the health warnings to communicate that the Department of Health and Human Services (HHS) endorses the health message. Another comment from a public health advocacy group suggested that the warning statements add a reference to FDA and/or the U.S. Government to legitimize the warnings. In contrast, one comment stated that it did not support continued use of the Surgeon General attribution, but if FDA decides to include the attribution, it should be placed on the side of the package where it does not detract from the new health warnings.

(Response) We agree with comments highlighting the benefits of the Surgeon General’s work in tobacco prevention, but we decline to add the “Surgeon General” attribution to the
required warnings at this time. Congress did not include an attribution to the Surgeon General as it has done in the past. In addition, there is inconsistency among the limited scientific literature as to whether the attribution of health warnings to government sources enhances their credibility (see, e.g., Refs. 42, 36, 57, and 59). Attribution to a government resource may increase believability of the information; however, if the government is generally disliked or mistrusted, a government source attribution may result in rejection of the health warning (Ref. 11).

One 1997 study found that the attribution to a government source, including the U.S. Surgeon General, did increase the credibility and viewers’ intentions to comply with the warnings for cigarettes (Ref. 57). Similarly, in a study conducted prior to Israel’s decision to require new cigarette warnings on packages, researchers found that consumers preferred warnings with attribution to a government source or medical research rather than warnings without attribution (Ref. 59).

However, in a developmental study assessing appropriate attributes for new cigarette warnings in Australia, researchers found that the mention of “government” in an attribution reminded smokers that the government collects tax revenue from cigarettes and led smokers to challenge the sincerity of the government in issuing cigarette health warnings (Ref. 48). Similarly, researchers for the European Commission in the European Union looked at respondents’ reactions to three potential attributions for cigarette warnings: (1) Government/regulatory bodies; (2) health authorities/cancer charities; and (3) tobacco industry (Ref. 42). They found smokers did not respond well to regulatory bodies as a potential source for cigarette warning messages, believing that government bodies did not care about their smoking behavior or were motivated by self-interest (Id.).

Moreover, even though the 1997 study did find benefits associated with government source attribution, researchers also noted the potential trade-offs associated with government attribution (Ref. 57). They noted the surface area restrictions associated with warnings and that the amount of information that one can give without losing readers is limited (Id.). They also noted that the addition of attribution information may require the use of smaller font size, which may impact legibility and noticability of the warning (Id.).

In fact, as we noted in the preamble to the proposed rule, the length and font size of the existing warnings contribute to their ineffectiveness, and larger font sizes enhance the noticeability of cigarette warnings (75 FR 69524 at 69530 and 69534; Ref. 40 at 30–31). Therefore, given the inconsistency in the available research and the potential tradeoffs associated with including a government source attribution in the required warnings, we conclude that there is insufficient evidence to support addition of an attribution at this time.

We will continue to work in partnership with other components within HHS to educate consumers about the risks of smoking. FDA and others also will continue to conduct research regarding the efficacy of required warnings. If such research indicates that adding the Surgeon General attribution to the cigarette required warnings will improve their efficacy, we will consider adding a government attribution as part of a future rulemaking to update the warnings.

### C. Foreign Language Translations

As we explained in the preamble to the proposed rule, consistent with section 4(b) of FCLAA, proposed §1141.10(b)(2) would mandate that the textual component of the required warning appear in the English language in cigarette advertisements with two exceptions. First, per proposed §1141.10(b)(2)(i), if an advertisement appears in a non-English language publication, the textual portion of the required warning would need to appear in the predominant language of the publication. Second, per proposed §1141.10(b)(2)(ii), if an advertisement is in an English language publication but the advertisement itself is presented in a language other than English, the textual portion of the required warning would need to be presented in the same foreign language principally used in the advertisement. To accommodate the potential need for Spanish language translations of the textual warning statements, we included Spanish translations with the proposed rule. We received several comments regarding foreign language translations in advertisements and one comment requesting the use of foreign language translations on packages. We have summarized and responded to these comments in the following paragraphs.

(Comment 128) One comment indicated that the submitter was pleased to see Spanish translations of the warnings, but asked that FDA continue to work with as many languages as possible.

(Response) We understand the importance of ensuring that the textual portion of the required warnings is translated accurately so that the message is appropriately communicated to foreign language speakers. As indicated in the NPRM, we included Spanish language translations in recognition of the fact that Spanish is the foreign language most commonly used for cigarette advertisements in the United States (75 FR 69524 at 69537 through 69538). We also will work with any advertiser who plans to advertise cigarettes in any non-language advertisement, or who plans to utilize a non-English advertisement in an English-language publication in accordance with §1141.10(b)(2)(ii).

Specifically, upon request, we will assist advertisers in generating a true and accurate translation of the textual statements for the nine new required warnings for use in advertisements that are subject to §1141.10(b)(2).

(Comment 129) One comment expressed concerns that foreign language translations sometimes can be “too literal” and could inappropriately impact the meaning of the warning.

(Response) We are sensitive to this concern, and the final rule requires that any translation of the required warning statements results in a true and accurate foreign language version of the warning statements. As stated in the previous response, we will assist any advertiser who plans to advertise cigarettes with a foreign language translation of the required warnings.

(Comment 130) One comment stated that all cigarette advertisements in predominantly Spanish speaking areas, such as Puerto Rico, and in Spanish language publications should include warnings in Spanish. Another comment recommended that the required warnings in advertisements be in the language of the publication or advertisement.

(Response) We agree in certain circumstances. As stated in the proposed rule and required in §1141.10(b)(2), any advertisement that appears in a Spanish language publication must present the textual portion of the required warning in Spanish (see §1141.10(b)(2)(ii)). In addition, for advertisements in English language publications, if the advertisement itself is presented in Spanish, the required warning in the advertisement also must be in Spanish (see §1141.10(b)(2)(ii)). However, if an English language publication that includes English language advertisements is sold in predominantly Spanish speaking areas, the textual component of the required warnings will still be required to appear in
English, as specified by section 4 of FCLAA. We conclude that these requirements will appropriately ensure that the target audience of any advertisement is able to read and understand both the promotional content of the advertisement and the important warning information.

(Comment 131) One comment requested that the required warnings on all cigarette packages exported to Puerto Rico and Latin America be in Spanish.

(Response) We decline to adopt this request. Section 4(b)(2) of FCLAA and § 1141.10(b)(2) require translation of required warnings for certain advertisements only. Neither FCLAA nor the Tobacco Control Act requires foreign language warnings on cigarette packages sold or distributed within the United States, including within the Commonwealth of Puerto Rico. Furthermore, with limited exceptions, FCLAA does not apply to packages of cigarettes for export from the United States.

V. Description of the Final Rule

A. Overview of the Final Rule

This final rule adds new part 1141 to Title 21 of the Code of Federal Regulations, requiring new warnings on cigarette packages and in cigarette advertisements. These new required warnings consist of one of the nine textual warning statements set forth in section 201 of the Tobacco Control Act accompanied by color graphic images depicting the negative health consequences of smoking. We have selected nine images, such that each required warning consists of one of the nine textual warning statements and an accompanying color graphic.

As required by section 201 of the Tobacco Control Act, the rule requires the new warnings to appear prominently on cigarette packages and in advertisements, occupying at least 50 percent of the area of the front and rear panels of cigarette packages and the top 20 percent of the area of advertisements. We also have exercised our authority under sections 201 and 202 of the Tobacco Control Act, which allow FDA to adjust the type size, text, and format of the textual warning statements. For example, under section 4(d) of FCLAA (as amended by section 201 of the Tobacco Control Act), FDA may adjust the type size, text, and format as we determine appropriate so that both the textual warning statements and the accompanying graphics are clear, conspicuous, legible, and appear within the specified area. Such adjustments, including adjustments to the type size and the addition of information regarding a cessation resource, are included for the required warnings in this final rule. In addition, we are requiring a reference to 1–800–QUIT–NOW as part of the required warnings in accordance with section 906(d) of the FD&C Act as appropriate for the protection of the public health.

B. Description of Final Regulations and Responses to Comments

1. Section 1141.1—Scope

In the proposed rule, proposed § 1141.1 set forth the scope of the proposed regulations. In particular, proposed § 1141.1(b) limited the applicability of the proposed requirements by clarifying that they would not apply to manufacturers or distributors of cigarettes that do not manufacture, package, or import cigarettes for sale or distribution in the United States. Proposed § 1141(c) described situations where a cigarette retailer would not be in violation of the proposed rule for displaying or selling cigarette packages that do not comply with the rule, so long as certain conditions were met (75 FR 695112 at 695355). We received several comments regarding the scope of the regulation, which we have summarized and responded to in the following paragraphs.

(Comment 132) One comment requested that all imported cigarettes and tobacco products have required warnings to come into U.S. ports and be sold in the United States and its territories, including Puerto Rico.

(Response) We agree that imported cigarette packages must bear a required warning in accordance with section 4 of FCLAA and part 1141. Section 1141.10 provides that it is unlawful for any person to import for sale or distribution within the United States any cigarettes the package of which fails to bear one of the required warnings on both the front and rear panels. Section 1141.3 defines United States to include specified U.S. territories, including Puerto Rico. In addition, as explained in section V.B.2 of this document, we are revising the definition of importer to clarify that the term importer includes any person who imports any cigarette, regardless of where it was manufactured. With respect to whether other tobacco products should have required warnings, we have determined that issue is outside the scope of this rulemaking.

(Comment 133) One comment supported the imposition of the required warnings on all cigarette packages manufactured in the United States, including all exported cigarette packages. The comment said that it would be unconscionable for FDA to protect residents in the United States and not the rest of the world when they are smoking U.S.-made products. According to this comment, cigarettes that are being exported are essentially bought in the United States and these products are under the FDA’s jurisdiction.

(Response) We disagree that it is appropriate to impose a requirement that cigarettes that are manufactured in the United States for export bear a required warning. Section 4(a) of FCLAA applies to cigarette packages that are “for sale or distribution within the United States.” Section 12 of FCLAA provides:

Packages of cigarettes manufactured, imported, or packaged (1) for export from the United States or (2) for delivery to a vessel or aircraft, as supplies, for consumption outside of the United States, shall be exempt from the requirements of this Act, but such exceptions shall not apply to cigarettes manufactured, imported, or packaged for sale or distribution to members or units of the Armed Forces of the United States located outside of the United States.

(Comment 134) One comment requested that FDA exercise enforcement discretion for retailers and distributors selling cigarettes that do not bear a specified warning label because retailers do not control the labeling of the products supplied by manufacturers. The comment claimed that if a product is provided by a licensed supplier, and not altered by the distributor, the distributor should likewise be relieved of liability.

(Response) FCLAA provides a very limited exemption for retailers and we do not agree that it is appropriate to broaden the exemption to distributors. Nor do we agree that it is appropriate to adopt a broad enforcement discretion policy for retailers and distributors. By choosing to distribute and sell cigarettes, distributors are under an obligation to make sure that the products they receive from manufacturers, importers, and other distributors and subsequently distribute or sell comply with the law, including checking to see whether the packages include a required warning on the front and rear panel. Retailers, however, are not in violation if they display or sell a cigarette package that includes a health warning, even if it is not one of the nine required warnings, as long as other...
statutory requirements are met (see 15 U.S.C. 1333(a)(4)). The preamble to the proposed rule made clear that manufacturers, importers, and distributors have the primary responsibility for ensuring that the required warnings on cigarette packages comply with all the provisions of part 1141.

(Comment 135) One comment expressed concern regarding the exemption of retailers from an obligation to ensure packages depict required warnings. This comment claimed that the exemption hampers enforcement, because an inspector needs to be able to seize noncompliant packaging at retail.

(Comment 136) One comment requested that FDA revise its 2010 Regulations Restricting the Sale and Distribution of Cigarettes and Smokeless Tobacco to Protect Children and Adolescents (75 FR 13225, March 19, 2010) ("reissued 1996 rule") to ensure that the Agency does not exceed the scope of the Tobacco Control Act by imposing liability on retailers and distributors for labeling or advertising in specific situations. This comment contended that the Tobacco Control Act provides specific situations in which retailers should not be held liable for labeling or advertising and those situations are not recognized in the reissued 1996 rule.

(Comment 137) Multiple comments advocated for the placement of graphic warnings on all tobacco products, including smokeless tobacco products.

(Comment 138) One comment asked that FDA expand the definition of importer to include persons who introduce into commerce cigarettes manufactured in the United States, exported from the United States, and subsequently imported. According to this comment, legislation in 2000 substantially curtailed this practice, but it is still possible.

(Comment 139) With respect to the definition of retailer, one comment requested that FDA revise the definition to clarify that Internet sellers are included in this definition. The comment noted that it appears the retailer definition is broad enough to cover Internet sellers, but clarification would avoid any arguments to the contrary.

As explained in the preamble to the proposed rule, proposed § 1141.3 defined ‘‘importer,’’ for purposes of part 1141, as any person who introduces into commerce any cigarette that: (1) Was not manufactured in the United States and (2) is intended for sale or distribution to consumers in the United States. Proposed § 1141.3 defined ‘‘retailer’’ as any person who sells cigarettes to individuals for personal consumption, or who operates a facility where vending machines or self-service displays of cigarettes are permitted (75 FR 69524 at 69536).

2. Section 1141.3—Definitions

Proposed § 1141.3 included definitions for the following terms:
- Cigarette
- Commerce
- Distributor
- Front panel and rear panel
- Importer
- Manufacturer
- Package
- Person
- Required warning
- Retailer
- United States

We received only a few comments regarding definitions described in the proposed rule. In light of the few comments, we are revising the definition of ‘‘importer.’’

3. Section 1141.10—Required Warnings

The Tobacco Control Act directs FDA to require that color graphic images depicting the negative health
consequences of smoking accompany each of the textual warning statements that must be randomly displayed on cigarette packages (i.e., in each 12-month period, all of the different warnings must appear in equal numbers of times on each brand of the product and be randomly distributed in all areas of the United States in which the product is marketed) and rotated quarterly in cigarette advertisements under FCLAA.

Accordingly, in proposed § 1141.10, we proposed that cigarette packages and advertisements contain such a combination graphic-textual warning. Proposed § 1141.10 provided that the warnings required by this section be obtained from two documents entitled “Cigarette Required Warnings—English and Spanish” and “Cigarette Required Warnings—Other Foreign Language Advertisements.” “Cigarette Required Warnings—English and Spanish” was proposed to contain the required warnings that must be included on all cigarette packages, and in cigarette advertisements in which the text of the required warning must be set forth in the English language or the Spanish language. “Cigarette Required Warnings—Other Foreign Language Advertisements” was proposed to contain the electronic files that were to be used to generate the required warnings for advertisements in which the text of the required warning must be set forth in a foreign language (other than Spanish).

The material that was proposed to be contained in the two documents entitled “Cigarette Required Warnings—English and Spanish” and “Cigarette Required Warnings—Other Foreign Language Advertisements” is now contained in a single document entitled “Cigarette Required Warnings.” We have provided this information in a single document because each of the electronic files for use in advertisements contained in “Cigarette Required Warnings” allows users to select an English or Spanish textual warning statement or to remove the textual warning statement and insert a true and accurate foreign language (other than Spanish) translation of the warning statement into the file. It is thus unnecessary to provide separate documents with electronic files for English and Spanish language advertisements and for advertisements in which the text of the required warning must be set forth in a foreign language (other than Spanish). Section 1141.10 has been updated to reference this single document, “Cigarette Required Warnings,” rather than the two proposed documents (“Cigarette Required Warnings—English and Spanish” and “Cigarette Required Warnings—Other Foreign Language Advertisements”).

Section 1141.10(a) sets forth the requirement specific to cigarette packages, explaining that the new required warning must comprise at least the top 50 percent of the front and rear panels of the package, except for cartons where the warnings shall comprise 50 percent of the left side of the front and rear panels. This regulation implements section 4(a)(2) of FCLAA and is in line with the provisions of the Framework Convention on Tobacco Control (FCTC) (Ref. 60). Section 1141.10(a)(3) specifically provides that the “required warning shall appear directly on the package and shall be clearly visible underneath the celophane or other clear wrapping.” Section 1141.10(b) sets forth the requirements for advertisements, including the requirement that the warnings comprise at least 20 percent of the area of the advertisements. Section 1141.10(c) provides that the required warnings shall be indelibly printed on or permanently affixed to the package or advertisement. For the final rule, we have deleted the language from § 1141.10(a)(2) and (b)(3) that specified that the electronic images must be adapted as necessary to meet the requirements of section 4 of FCLAA and part 1141. As explained in the NPRM (75 FR 69524 at 69536 through 69538), this language was used to indicate that regulated entities should modify the size of the required warnings to ensure they are visible and occupy the required area of the cigarette package or advertisement. However, § 1141.10(a)(4) and (b)(5) set forth the size and placement requirements for required warnings on packages and advertisements, so this language in proposed § 1141.10(a)(2) and (b)(3) was not necessary. In addition, § 1141.10(a)(1) and (b)(1) make clear that the required warnings on cigarette packages and in cigarette advertisements must be “in accordance with section 4 of the Federal Cigarette Labeling and Advertising Act.”

We also have made minimal changes to § 1141.10(b)(4), which used similar language. Specifically, proposed § 1141.10(b)(4) indicated that the required warnings for foreign language advertisements (other than Spanish) must be adapted as necessary to meet the requirements of section 4 of FCLAA and part 1141. For clarity, we have modified this language to indicate that the textual warning statement that is inserted into the electronic images must comply with the requirements of section 4(b)(2) of FCLAA. As explained in the NPRM (75 FR 69524 at 69538), proposed § 1141.10(b)(4) would have required regulated entities to obtain color graphics for foreign language required warnings, other than Spanish language warnings, from the electronic files contained in “Cigarette Required Warnings—Other Foreign Language Advertisements,” and regulated entities would have to insert a true and accurate foreign language translation of the textual warning required by FCLAA into the electronic file to generate the required warning (as explained previously, these electronic files are now contained in the document entitled “Cigarette Required Warnings”). While the electronic file obtained from “Cigarette Required Warnings” contains some of the elements required by FCLAA (e.g., a rectangular border to enclose the required warnings and the color graphic to accompany the label statement), the textual warning statement that regulated entities insert into the electronic file in accordance with § 1141.10(b)(4) must comply with the requirements of section 4(b)(2) of FCLAA. This section provides, among other things, format specifications related to the textual warning statements in cigarette advertising, including required type sizes and color specifications (i.e., the text of the label statement shall be black if the background is white and white if the background is black), and requires that the statements appear in conspicuous and legible type.

In addition, we wish to clarify our intent regarding whether the same warning statement must appear on both the front and rear panels of an individual cigarette package. We believe that section 4(a)(1) of FCLAA is ambiguous as to whether it mandates the use of the same required warning on both the front and rear panels of an individual cigarette package or allows two different required warnings to be used, one on the front panel and the other on the rear panel. We believe that the latter interpretation is reasonable. It is consistent with Congress’ intent that all of the required warnings, each of which conveys somewhat different health information, are required to be displayed in the marketplace at the same time (see section 4(c)(1) and (c)(3) of FCLAA). While it is possible that two copies of the same statement on a single package might increase the likelihood of the warning being noticed and remembered, we also note that different statements on a single package could lead to greater consumer exposure as well as delay the wear out of the required warnings. Proposed
§ 1141.10(a)(1), along with the description of this provision in the preamble to the proposed rule (75 FR 69524 at 69536), however, implied that the same required warning must appear on both the front and the back of the package. Therefore, we are revising §1141.10(a)(1) to state, “It shall be unlawful for any person to manufacture, package, sell, offer to sell, distribute, or import any cigarettes the package of which fails to bear* * * one of the required warnings on the front and the rear panels.”

We received comments regarding the format of required warnings on packages and advertisements, the applicability of the requirements to cigarette cartons, and the need for the warnings to remain clearly visible and permanently affixed to packages. A summary of these comments and our responses is provided in the following paragraphs.

(Comment 140) Many comments, including those from health institutions, nonprofit organizations, academics, and consumers, expressed the belief that there was no adequate justification for the amount of space mandated for the new required warnings. The comments asserted that the current size and placement of the warnings on cigarette packages and advertising have contributed to “complete awareness levels of the dangers of cigarettes.”

(Response) We disagree. As we stated in the preamble to the proposed rule, our assessment of the literature and our experience as a public health agency supports the requirement that the new warnings comprise the top 50 percent of the area of each of the front and rear panels of cigarette packages and the top 20 percent of the area of cigarette advertisements in the United States (75 FR 69524 at 69533). For example, researchers have found that larger graphic warnings are likely to have the greatest impact and that “larger (label) size means higher visibility and better ability to compete with other package elements” (Ref. 40 at p. 30). Smokers are more likely to recall larger warnings, and have been found to correlate the size of the warning with the seriousness of the risk (Ref. 61). One Canadian study found that smokers judged warnings that covered 80 percent of the package to be most effective (Ref. 11). In a New Zealand study gauging responses to different sized graphic health warnings (one sized 50 percent of the front of the pack, and another sized 30 percent of the front of the pack), participants strongly preferred the larger sized warning (Ref. 40 at p. 31). Participants felt that the larger sized warning was more prominent, more likely to stand out from product branding, and that some of the messages on the front of the pack remained visible when the pack was open (Id. at p. 30). The 50 percent requirement also is consistent with the FCTC (i.e., the required warnings should occupy 50 percent or more of the principal display areas of packages), which was among the substantial evidence considered by Congress when enacting the Tobacco Control Act (FCTC art. 11.1(b)). “Congress and its supporters knew the warning requirement by looking at the use of a nearly identical warning requirement in Canada.”

(Comment 141) A few comments expressed the belief that there was no adequate justification for the amount of space mandated for the new required warnings (i.e., 50 percent of the front and back panels of packages and the top 20 percent of the area of advertisements). One comment noted that Congress enacted the 50 percent requirement without committee testimony or other fact-finding as to whether a smaller-sized warning would be effective. The comments asserted that the current size and placement of the warnings on cigarette packages and advertising have contributed to “complete awareness levels of the dangers of cigarettes.”

(Response) We disagree. As we stated in the preamble to the proposed rule, our assessment of the literature and our experience as a public health agency supports the requirement that the new warnings comprise the top 50 percent of the area of each of the front and rear panels of cigarette packages and the top 20 percent of the area of cigarette advertisements in the United States (75 FR 69524 at 69533). For example, researchers have found that larger graphic warnings are likely to have the greatest impact and that “larger (label) size means higher visibility and better ability to compete with other package elements” (Ref. 40 at p. 30). Smokers are more likely to recall larger warnings, and have been found to correlate the size of the warning with the seriousness of the risk (Ref. 61). One Canadian study found that smokers judged warnings that covered 80 percent of the package to be most effective (Ref. 11). In a New Zealand study gauging responses to different sized graphic health warnings (one sized 50 percent of the front of the pack, and another sized 30 percent of the front of the pack), participants strongly preferred the larger sized warning (Ref. 40 at p. 31). Participants felt that the larger sized warning was more prominent, more likely to stand out from product branding, and that some of the messages on the front of the pack remained visible when the pack was open (Id. at p. 30). The 50 percent requirement also is consistent with the FCTC (i.e., the required warnings should occupy 50 percent or more of the principal display areas of packages), which was among the substantial evidence considered by Congress when enacting the Tobacco Control Act (FCTC art. 11.1(b)). “Congress and its supporters knew the warning requirement by looking at the use of a nearly identical warning requirement in Canada.”


In addition, as described more fully in section II.C of this document, the existing warnings have not been effective in communicating the health risks of smoking, resulting in significant portions of the population that misunderstand or underestimate the health risks of smoking. The new size and placement requirements are needed to increase the salience of cigarette health warnings, which are now considered “invisible,” in order to educate the public about the health risks of smoking, which in turn, can positively impact smoking intentions and behaviors (Ref. 3 at p. 291).

(Comment 142) Some comments suggested that the regulation include a font size requirement.

(Response) We note that the proposal included a requirement related to font size and this is retained in the final rule. The final rule mandates that the required warnings be accurately reproduced from the document incorporated by reference entitled “Cigarette Required Warnings.” The required font style and font size already will be included in the options within the downloadable files that allow the user to select English and Spanish language warning statements.

For advertisement in foreign languages other than Spanish, companies must comply with the font size requirements in section 4(b)(2) of FCLAA and any format requirements included in the document incorporated by reference (see section V.B.4 of this document). In all situations, the textual statements must be conspicuous and legible as required by section 4 of FCLAA.

(Comment 143) One comment from an industry group took issue with FDA’s authority to require the new graphic warnings on cigarette cartons, claiming that Congress’ intent was to require the new graphic warnings on individual cigarette packs only, not cartons. The submitter recommended that FDA expressly exempt cartons from this requirement.

(Response) We disagree with this comment. FCLAA defines the term “package” to mean a “pack, box, carton, or container of any kind in which cigarettes are offered for sale, sold, or otherwise distributed to consumers” (section 3(4) of FCLAA (15 U.S.C. 1332(4)) (emphasis added)). Similarly, section 900(13) of the FD&C Act defines the term “package” to mean a “pack, box, carton, or container of any kind or if no other container, any wrapping (including cellophane), in which a tobacco product is offered for sale, sold, or otherwise distributed to consumers.” (21 U.S.C. 387(13) (emphasis added)). Given these definitions, it is clear that when Congress decided to require graphic warnings that occupy 50 percent of the front and back panels of cigarette “packages,” it intended for this requirement to apply to both individual
packs and cartons. Therefore, § 1141.10(a)(4) continues to mandate that the required warnings must constitute 50 percent of the left side of the front and rear panels of cigarette cartons.

(Comment 144) One comment recommended that FDA require the nine new textual warning statements, included in section 4(a) of FCLAA, to be displayed in the same manner as the display of the existing warnings, because that format has contributed to the public being fully informed about the health risks of smoking. (Response) We disagree. First, as explained in section II.C. of this document, the public is not adequately informed about the health risks of smoking and frequently underestimates those risks. Second, Congress mandated that the format of the new health warnings change from the small warning on the side panel of the pack, covering only 4 percent of the pack, to health warnings that “comprise the top 50 percent of the front and rear panels of the package” and “at least 20 percent of the area of the advertisement.” (15 U.S.C. 1333(a)(2) and (b)(2)). This is consistent with the FCTC (FCTC art. 11.1(b)). Therefore, we decline to change the format of the required warnings from that included in the proposed rule.

(Comment 145) One comment suggested that the required warnings on cigarette advertisements cover at least 50 percent of the advertisement’s principal surface and match the advertisement’s primary language. (Response) As stated in the preamble to the proposed rule and as required by section 4 of FCLAA, § 1141.10(b)(5) mandates that the required warnings comprise at least the top 20 percent of the area of the advertisement. Section 4 of FCLAA also requires that the warning statement appear in conspicuous and legible type. At this time, we conclude these requirements are sufficient to ensure that the required warnings are appropriately clear, conspicuous, and legible by consumers. Moreover, as stated in the preamble to the proposed rule and as indicated in section IV.C of this document, while the textual portion of the required warning in a cigarette advertisement must generally be in English, if an advertisement is presented in a language other than English, the textual portion of the required warning must be presented in the language principally used in the advertisement (see § 1141.10(b)(2)(ii)). Therefore, we have determined that modifications to the codified text are not necessary.

(Comment 146) Proposed § 1141.10(a)(5) provided that the “required warning shall be positioned such that the text of the required warning and the other information on that panel of the package have the same orientation.” One comment expressed concern that this provision could be problematic if a manufacturer places the brand name and other information vertically on the front and/or back of the cigarette package. The comment believed that this provision would require the warning, or the text of the warning, to appear sideways on the cigarette package. (Response) The intent of this provision is to ensure that the textual statement in the required warning and other information on the front and rear panels of the package have the same orientation. As explained in the NPRM, this will in turn ensure that the warnings are noticed and read by consumers that are reading the other information found on the package (75 FR 69524 at 69537). Therefore, in the unusual circumstance where a manufacturer chooses to place its brand name or other information such that viewers do not read along the horizontal axis (i.e., from left to right) to read this information, the manufacturer must place the required warning in the same orientation.

(Comment 147) Two comments suggested that the FDA require health warnings on 100 percent of only the front or the rear panel of the cigarette package. (Response) We disagree. First, section 4(a)(2) of FCLAA specifically requires that the cigarette health warnings “comprise the top 50 percent of the front and rear panels of the package.” Second, Article 11 of the FCTC states that the health warnings “should be 50% or more of the principal display areas but shall be no less than 30% of the principal display areas” (Ref. 60). FDA’s new warnings implement Congress’ directive and are consistent with the FCTC.

(Comment 148) A few comments suggested that FDA require health warning statements on cigarette papers and/or filters. (Response) We decline to require warnings on cigarette papers and/or filters. In section 4(d) of FCLAA, Congress directed FDA to issue regulations to require color graphic images to accompany the warnings statements required by section 4(a)(1) of FCLAA. FCLAA requires that the statements be included on advertisements and cigarette packages, not individual cigarette papers or filters. While we may be able to require warnings on papers or filters under other authority, that is outside the scope of this rulemaking.

(Comment 149) One comment suggested that FDA amend the regulation to prohibit distributors from obscuring any portion of the warning label with revenue stamps. (Response) As written, the proposed rule would prohibit distributors from obscuring any portion of the required warning with revenue stamps. Cigarette packages must comply with the requirement in § 1141.10(a)(3) that the new required warnings be clearly visible. Moreover, in order for the required warnings to appear conspicuously and legibly as mandated by section 4 of FCLAA, the warnings must not be obscured. Thus, if the placement of revenue stamps by a distributor causes the required warnings to not be clearly visible or legible, the distributor would be in violation of these regulations. Therefore, we do not agree that any revisions to § 1141.10 are necessary.

(Comment 150) One comment suggested that FDA require the use of onsets affixed to cigarette packages in addition to the new required warnings, stating that they would enhance the effectiveness of the new health warnings. Similarly, another comment stated that, in addition to the new required warnings, FDA should require that cigarette packages contain inserts with animated warnings containing supplementary or distinct warning messages to enhance the overall warning impression and further engage individuals.

(Comment 151) One comment stated that there is no empirical basis for concluding that the nine warning statements required under section 4 of FCLAA should be written in large text on the front and back panels of packages in order to convey the health risk information. (Response) We disagree with this comment and conclude that there is a sufficient empirical basis for concluding that the warning statements should be in large text that is conspicuous and legible. Research has shown that increasing the salience of warnings increases the likelihood of consumers reading warnings and that the salience of a visual warning can be enhanced by using large, bold print (Ref. 62). In addition, after Australia changed their health warnings to include large, bold warnings on cigarette packages with a cessation resource and additional explanatory text in 1995,
researchers found that the increased text size was the most salient feature (Ref. 63). Furthermore, the IOM Report, which provides a summary of the available research on the efficacy of graphic warnings, found that larger, graphic health warnings (including large text and a large graphic) would promote greater public knowledge of the health risks and would help reduce consumption of tobacco products (Ref. 3). The placement of the large text and graphic image on the front and back panels of cigarette packages is consistent with the FCTC, i.e., that health warnings should occupy 50 percent or more of the principal display areas of packages (FCTC art. 11.1(b)).

(Comment 152) One comment claimed that the format of the new required warnings is inconsistent with FDA’s drug warning label regime. For example, the comment stated that even for very severe risks, the drug regulations do not require warning information to appear in large text or to occupy a large portion of the packaging. The comment also noted that, in drug advertising, the FDA requires important risk information to be included in a section of the advertisement entitled “Brief Summary.”

(Response) We have acknowledged that the warning requirements for cigarettes are, and should be, different than the warnings for other FDA-regulated products. As we explained in the preamble to the proposed rule, “(1) The warning information for cigarettes is different in its applicability than the warning information for other products, and (2) the disclosure requirements for other products have a different purpose than the cigarette warnings, and (3) the mechanisms for exposure to warning information are different for tobacco products than for other products FDA regulates.” (75 FR 69524 at 69539). In contrast to medical products regulated by FDA, there is no population that cigarettes are medically appropriate for, and there is no safe method of using cigarettes; the required warnings for these products thus have an inherently different purpose than medical product warning information. The different warning schemes that apply to tobacco products versus medical products are necessary to most effectively communicate the health risks for tobacco products and for other FDA-regulated products.

(Comment 153) One comment claimed that FDA did not provide an adequate justification for requiring the same health warning messages in multiple media, including print advertisements, point-of-sale displays, cartons, and the front and back of individual cigarette packs. This comment claimed that the publication of health warning messages in multiple media will not foster awareness of the information (because it is already known) or belief in it (because it is already believed).

(Response) We disagree. As explained in section II.D of this document, despite existing warning requirements on packages and in advertisements, consumers lack knowledge of the health risks and underestimate the health risks of smoking. It is critical that the negative health consequences of cigarette smoking, which is the leading cause of preventable death and disease in the United States, be clearly, accurately, and effectively conveyed in all advertisements and on all cigarette packages sold or distributed in the United States.

This is consistent with the requirements of FCLAA. As explained more fully in response to Comment 143, FCLAA’s requirements apply to cigarette packages (including cartons), and to advertisements generally.

Further, with its passage of the Tobacco Control Act, Congress noted the pervasiveness of tobacco advertising and how it impacts use, especially promotions directed to attract youths to tobacco products, and found that comprehensive advertising restrictions will have a positive effect on the smoking rates of young people (section 2(15) and 2(25) of the Tobacco Control Act). Therefore, the requirement that the warnings appear in all advertisements, regardless of the medium used for the advertisement, is also consistent with Congress’ intent.

(Comment 154) One comment noted that the Federal government warnings on alcoholic beverages are mandated on packages only, presented in small font, and not required on the prominent faces of containers or packaging. According to the comment, this suggests that Congress believes a configuration like the one for alcoholic beverages also would be sufficient for cigarette warnings, particularly given the more widespread use of alcoholic beverages in this country.

(Response) We disagree. Congress clearly intended for the warnings for cigarettes and alcoholic beverages to be different, as evidenced by the different statutory schemes that govern the warning requirements for cigarettes and alcohol products. For cigarettes, Congress clearly set out the location of the health warnings for cigarette packages and advertisements, the area of the product that must be covered by the warnings and the requirements for text and background color of the warnings. In addition, Congress provided specific font size requirements for the cigarette warnings (while also affording FDA the authority to initiate a rulemaking proceeding to adjust the format, type sizes, and certain other aspects of the health warnings under sections 4(b)(4) and (d) of FCLAA and section 202(b) of the Tobacco Control Act. In contrast, Congress’ health warning requirements for alcoholic beverages, published at 27 U.S.C. 215, do not set forth area, location, and color requirements with as much specificity.

(Comment 155) One comment from an individual consumer expressed concerns that manufacturers may alter their packaging to subvert § 1141.10(c), which mandates that the required warnings on packages and advertisements must be irremovable or permanent.

(Response) The regulation, as drafted, should address the comment’s concern. Section 1141.10(c) of the final rule, which is unchanged from what appeared in the proposed rule, states that the “required warnings shall be indelibly printed on or permanently affixed to the package or advertisement.” Therefore, regardless of the type of packaging used by manufacturers, all cigarette packages must contain required warnings that are irremovable or permanently affixed to the cigarette packages.

4. Section 1141.12—Incorporation by Reference of Required Warnings

Proposed § 1141.12 proposed that two documents, “Cigarette Required Warnings—English and Spanish” and “Cigarette Required Warnings—Other Foreign Language Advertisements,” be incorporated by reference in accordance with 5 U.S.C. 552(a) and 1 CFR part 51. Draft versions of both documents were made available in the docket with the NPRM.

We did not receive comments regarding the use of the incorporated by reference mechanism provided in 5 U.S.C. 552(a) and 1 CFR part 51 and the proposed codified language, or regarding the two draft documents proposed for incorporation by reference. However, as explained in section V.B.3 of this document, the material of which was proposed to be contained in the two documents entitled “Cigarette Required Warnings—English and Spanish” and “Cigarette Required Warnings—Other Foreign Language Advertisements” is now contained in a single document entitled “Cigarette Required Warnings.” As a result, we have made nonsubstantive changes to the language used in § 1141.12 to indicate that we are
incorporating “Cigarette Required Warnings” by reference (rather than “Cigarette Required Warnings—English and Spanish” and “Cigarette Required Warnings—Other Foreign Language Advertisements”). In addition, we also have updated the incorporation by reference document to include the final electronic files for the required warnings and to add additional formats and instructions for regulated entities to use to place the required warnings on various sizes of cigarette packages (including cartons) and in different sizes and shapes of advertisements, as is discussed in more detail in section VI of this document.

“Cigarette Required Warnings,” including the electronic files for all of the required warnings and the instructions for their use, is available from a variety of sources. For example, this material is available on a Web site at http://www.fda.gov/cigarettewarningfiles. In addition, regulated entities can request a copy of “Cigarette Required Warnings” by submitting a request to FDA at the following e-mail address—cigarettewarningfiles@fda.hhs.gov—or by contacting the Center for Tobacco Products, Food and Drug Administration, Office of Health Communication and Education, ATTN: Cigarette Warning File Requests, 9200 Corporate Blvd., Rockville, MD 20850, 1–877–CTP–1373.

5. Section 1141.14—Misbranding of Cigarettes

Proposed § 1141.14(a) provided that a cigarette shall be deemed to be misbranded unless its labeling and advertising bear one of the required warnings. Under section 903(a)(1) and (a)(7)(A) of the FD&C Act (21 U.S.C. 387c(a)(1) and (a)(7)(A)), a tobacco product, including a cigarette, is deemed misbranded if its labeling or advertising is false or misleading in any particular. Under section 201(n) of the FD&C Act (21 U.S.C. 321(n)), in determining whether something is misleading, it: “shall be taken into account * * * not only representations made or suggested * * * but also the extent to which the labeling or advertising fails to reveal facts * * * material with respect to consequences which may result from the use of the article to which the labeling or advertising relates * * * under such conditions of use as are customary or usual.” As explained in the NPRM (75 FR 69524 at 69539), the required warnings are clearly material with respect to consequences that may result from the use of cigarettes.

Proposed § 1141.14(b) provided that a cigarette advertisement or package will be deemed to include a brief statement of relevant warnings for the purposes of section 903(a)(8) of the FD&C Act if it bears one of the required warnings. It also proposed that a cigarette advertisement or package offered for sale in any State shall be deemed to be misbranded under section 903(a)(8) of the FD&C Act unless the manufacturer, packer, or distributor includes in all advertisements and packages issued or caused to be issued by the manufacturer, packer, or distributor with respect to the cigarette one of the required warnings. We received two comments on the issue, which we have summarized and responded to in the following paragraphs.

(Comment 156) One comment from a tobacco product manufacturer stated that FDA should replace the word “labeling” with the word “packages” in § 1141.14(a). The comment indicated that FDA should avoid using the word “labeling” because that term has a broader meaning under the FD&C Act than it does under FCLAA, and therefore its use in the regulation could create unnecessary ambiguity. The comment also stated that FCLAA only requires warnings on cigarette packages and advertisements.

(Response) We agree that cigarettes can be deemed misbranded under the FD&C Act for a number of reasons. We also agree that, although compliance with the requirements of part 1141 is necessary to comply with certain provisions of section 903 of the FD&C Act, this does not guarantee that a cigarette product satisfies all the requirements of FCLAA. However, as explained in the NPRM, the goal is to provide a place where smokers and other members of the public can obtain smoking cessation information from staff trained specifically to help smokers quit by delivering current, unbiased, and evidence-based information, advice, and support. The NPRM identified a number of possible alternatives for a cessation resource, including use of an existing or new quitline or Web site.

6. Section 1141.16—Disclosures Regarding Cessation

Section 1141.16 of the NPRM proposed that one or more of the required warnings include specified information about an appropriate smoking cessation resource. As explained in the NPRM, the goal is to provide a place where smokers and other members of the public can obtain smoking cessation information from staff trained specifically to help smokers quit by delivering current, unbiased, and evidence-based information, advice, and support. The NPRM identified a number of possible alternatives for a cessation resource, including use of an existing or new quitline or Web site.
Although we did not include a specific cessation resource on the proposed images published with the NPRM, we proposed that the final rule would include one or more required warnings containing a cessation resource. We proposed that the resource must meet specific criteria designed to ensure that the cessation information, advice, and support provided are unbiased and evidence-based.

As explained more fully in the following paragraphs, we have decided, based on our authority in section 906(d) of the FD&C Act, to require that all nine required warnings refer to a cessation resource, and we have included this resource in the nine graphic warnings in “Cigarette Required Warnings,” which is incorporated by reference (IBR) document as described in section V.B.4 of this document. This final rule specifies the criteria that will be required of any responsible entity providing services through the chosen cessation resource. The resource we have selected is the existing National Network of Tobacco Cessation Quitlines (Network), which uses the telephone portal 1–800–QUIT–NOW. This telephone portal, provided by the National Cancer Institute (NCI), routes calls to the appropriate State quitline, based on the area code of the caller. The Network includes a designated quitline run by or on behalf of each of the 50 states as well as the District of Columbia, Puerto Rico, and Guam (hereinafter referred to as “State quitlines” or “State-run quitlines”).

We conclude that this resource will provide the broadest access for smokers throughout the United States to unbiased, evidence-based cessation information, advice, and support. The Centers for Disease Control and Prevention (CDC) already provides significant support and oversight to these State-run quitlines. Beginning with the effective date of this rule, CDC’s cooperative agreements with State health departments will specify that the State quitlines must meet the criteria described in §1141.16(b) to qualify for cessation funding under the cooperative agreement. HHS will monitor the quitlines for compliance with the criteria, and if it determines that a State quitline does not meet the criteria, it will take appropriate steps to bring the State quitline into compliance. What is appropriate will depend on the circumstances of the particular situation. For example, it might involve CDC working with the State quitline to ensure staff are adequately trained. If warranted, it could also include more serious measures such as CDC working with NCI to re-route calls to another resource. Because the record indicates that quitlines that are members of the Network generally comply with the criteria already, we anticipate that any measures to bring quitlines into compliance will be rare.

a. Rationale and authority for requiring inclusion of a cessation resource. The NPRM explained that reducing the number of Americans who smoke by increasing the likelihood that smokers will quit smoking would provide substantial public health benefits by reducing the life-threatening consequences associated with continued cigarette use. The NPRM also cited studies finding that health warnings are more effective if they are combined with cessation-related information. Consequently, FDA proposed requiring information about an appropriate smoking cessation resource under section 906(d) of the FD&C Act as appropriate for the protection of the public health (75 FR 69524 at 69540 through 69541). We received a number of comments regarding our rationale and authority to require a cessation resource on the graphic health warnings, which we summarized and responded to in the following paragraphs.

(Comment 158) A large majority of comments that addressed the issue strongly supported inclusion of a cessation resource on all the required warnings. These include comments from public health advocacy groups, medical organizations, academics, State and local public health agencies, and representatives of quitlines. The comments provided a variety of reasons supporting inclusion of a cessation resource on the required warnings. Many comments asserted that a majority of smokers want to quit, and referring smokers to a smoking cessation resource will help them to quit. Some comments cited statistics regarding the number of smokers who actually attempt to quit smoking—about 40 percent of smokers try to quit in a calendar year—and the very low percentage of smokers who are successful—95 percent of those who try to quit on their own relapse (citing, e.g., Ref. 65 and Ref. 66). One comment from a State public health agency asserted that media campaigns and educational efforts, while effective, still do not reach all smokers. According to this comment, after extensive outreach, about 25 percent of smokers in that city had never heard of the quitline being promoted and 25 percent of smokers reported that it is not easy for a person interested in quitting smoking to obtain information about ways to quit.

Several comments noted that the purpose of graphic warnings is to inform smokers about the risks of smoking and motivate smokers to want to quit, but this message will be more effective if there is information in the graphic warnings on how smokers can obtain help quitting. Some comments argued that health warnings should not just inform smokers about the dangers of tobacco use, but also provide assurance that quitting is possible and assistance is available. One comment cited research that shocking, fear-arousing images can be more effective when combined with encouragement or...
empowering messages (citing, e.g., Ref. 74). Another comment from an academic institution claimed that when people perceive that there is a strategy for them to take positive action to reduce the threat in a fear message, fear appeals successfully changed health-related attitudes and behaviors (citing, e.g., Refs. 75, 76, 77, and 78). However, if people do not believe they have an effective means of avoiding a threat, they may suppress thoughts about the risk, and, as a result, not process the threat information (citing, e.g., Refs. 79, 80, and 81). As one comment from an academic institution explained, under fear appraisal theory, a fear communication message will cause aversive anxiety, which individuals will try to ameliorate through behaviors that reduce the perceived threat. This comment asserted that the positive effects of a fear message depend upon the existence of an available coping option that is perceived to be potentially effective at reducing the threat. In addition, comments cited research that smokers may be more likely to attempt to quit when they know a quitline is available (Ref. 82).

One comment from a submitter representing a State quitline claimed that health care providers are more likely to address tobacco use in their patients when they know of an effective program to which they can refer their patients, and that adding a cessation resource to the required warnings will dramatically increase awareness of this resource. Several comments from submitters representing State quitlines noted that they receive referrals from clinicians via fax referral services.

One comment from an academic researcher submitted results from a study that tested one of the proposed required warnings included in the proposed rule with and without a cessation resource. This study found that when youth and adult participants were asked to rank order six images tested for use with one of the warning statements, based on which image would be most effective for discouraging smoking, the image with the cessation resource was ranked as the most effective by more study participants than any other image.

(Response) We agree with comments that there is strong support for including a smoking cessation resource on the required warnings. As required by section 906(d) of the FD&C Act, we find that addition of a cessation resource is appropriate for the protection of the public health because of the benefits, and lack of risks, to the population as a whole. This is due, in part, to the increased likelihood that existing smokers will become aware of the cessation resource and, consequently, the increased likelihood that existing smokers who want to quit will be successful. It is also due to the likelihood that the reference to a smoking cessation resource will enhance the effectiveness of the warnings required under FCLAA at conveying information about the risks to health from smoking.

As stated in the comments, the majority of smokers want to quit and about 40 percent of smokers attempt to quit each year. In addition, the warnings required under FCLAA and this regulation convey information and promote greater understanding about the significant health risks associated with smoking, which will likely lead additional smokers to decide that they want to quit smoking to address these risks. Also, as discussed in the comments, the vast majority of those attempts are unsuccessful. By including a cessation resource on required warnings, the many smokers who want to quit will receive information about a resource that has been demonstrated to be effective in helping smokers to quit (see section V.B.6.c of this document). Media campaigns are helpful in reaching some smokers who want to quit, and can be used in conjunction with the inclusion of a cessation resource on the required warnings. It is important to ensure that this information reaches a broad number of smokers. Inclusion of a cessation resource on the required warnings is likely to have a broader reach than media campaigns alone. The evidence from one comment is that, even after an extensive media campaign, approximately one quarter of smokers surveyed were not aware of the existence of the quitline or that help was available to obtain information about ways to quit. The cessation information will be there each time a consumer looks at a package of cigarettes or a cigarette advertisement; a pack-a-day smoker potentially would be exposed to the cessation information more than 7,000 times per year. This evidence highlights that cigarette packages are useful communication tools for ensuring that smokers are aware of cessation resources.

Based on experience in other countries, we anticipate that including a reference to a cessation resource as part of the required warnings will increase the utilization of that resource. Many foreign countries have included cessation resources on cigarette package warnings. As described in the comments, these countries have generally experienced a large increase in the number of calls to the quitlines following their appearance on cigarette packages. For example, in the Netherlands, the number of callers to the quitline increased more than threefold after a smoking cessation message (“Ask for help with smoking cessation”) and the national quitline number were included on cigarette packages (Ref. 72). Similarly, in Australia, the number of calls to the quitline nearly doubled, compared with the previous 2 years, following the introduction of new color graphic warnings with a prominent quitline number. The increase in call volume persisted in the following year, although it was about 40 percent lower than in the year in which the graphic warnings were first introduced. Although there was a series of mass media campaign activities that accompanied the new graphic warnings, one study concluded it was very unlikely that the mass media campaign alone explained the observed increase in calls because the introduction of the graphic warnings had an independent effect (Ref. 67). In New Zealand, after the introduction of pictorial warnings with a supportive cessation message and quitline information, the average number of new monthly calls increased and the percentage of first-time callers who reported obtaining the quitline number from tobacco product packaging doubled (Ref. 83). In Brazil, there was a progressive increase in calls to a quitline in the 6 months following the requirement for graphic warnings and the inclusion of a quitline number on cigarette packages. People who called the quitline showed that over 92 percent knew about the quitline number because it appeared on cigarette packs (Ref. 73). We also note that Canada has recently proposed including a quitline number on the graphic warnings that will appear on its packages.

Although we are not aware of any studies regarding the inclusion of cessation information on graphic warnings in cigarette advertisements, it seems likely that a reference to a cessation resource to cigarette advertisements would have a similar effect as including the reference on cigarette packages.

Inclusion of a cessation resource on the required warnings is also consistent with the advice of the FCTC. Although the United States has not yet ratified the FCTC and therefore is not bound by the treaty, the United States is a signatory and the Guidelines for implementation of the Treaty provide further support for the inclusion of a cessation resource. The Guidelines for implementation of
Article 11 of the FCTC (Packaging and labeling of tobacco products) explain that the provision of advice on cessation and specific sources for cessation help on tobacco packaging, such as a Web site address or a toll-free telephone number, can be important in helping tobacco users to change their behavior, and is expected to increase demand for cessation-related services.

In addition to providing information to increase the likelihood that smokers will become aware of the cessation resource and use it to successfully quit, including a cessation resource will also help to make the required warnings more effective at conveying information about the health risks of smoking. As noted in the NPRM, studies have found that health warnings are more effective when they are combined with cessation-related information (75 FR 69524 at 69541). Risk communication research indicates that messages that arouse fear about the health risks of smoking should be combined with information on concrete steps that can be taken to reduce those risks (Ref. 81 (Messages that arouse fear "appear to be effective when they depict a significant and relevant threat * * * and when they outline effective responses that appear easy to accomplish * * *"); see also Ref. 55 (explaining the importance of giving smokers who are motivated to quit smoking upon seeing a graphic health warning an immediate way to act on this impulse and access cessation assistance)). In addition, the results from one study conducted by an academic researcher and submitted to the docket also suggest that adding a cessation resource to the required warnings is beneficial. When youth and adult participants were asked to rank order six images (including one image with and without a cessation resource) tested for use with one of the warning statements, based on which image would be most effective for discouraging smoking, the image with the cessation resource was ranked as the most effective by more study participants than any other image.

(Comment 159) Several tobacco industry comments claimed that it was difficult to comment on the issue of a cessation resource, because the proposed rule did not identify the resource FDA proposed to reference or suggest alternative resources from which FDA would choose. Tobacco industry comments also claimed that the NPRM did not indicate how FDA proposed to reference the resource or integrate it into the proposed warning images. For these reasons, some tobacco industry comments contended that the NPRM did not provide adequate notice for requiring inclusion of a cessation resource, and that FDA should not require a cessation resource without providing an additional opportunity to comment on specific proposed cessation resources.

(Response) We disagree. The Administrative Procedure Act requires that a notice of proposed rulemaking include "either the terms or substance of the proposed rule or a description of the subjects and issues involved" (5 U.S.C. 553(b)(3)). Consistent with this requirement, the NPRM provided adequate notice that FDA was considering the inclusion of a cessation resource in the required warnings and the factors it would consider in choosing a specific smoking cessation resource. Proposed § 1141.16 specifically stated that one or more of the required warnings "shall include a reference to a smoking cessation assistance resource" (75 FR 69524 at 69564). The preamble to the proposed rule explained the goal "would be to provide a place where smokers and other members of the public can obtain smoking cessation information from staff trained specifically to help smokers quit by delivering unbiased and evidence-based information, advice, and support" (75 FR 69524 at 69540). The preamble also explained the range of alternatives available, including use of an existing or new quitline or Web site (75 FR 69524 at 69540; see Small Refiner Lead Phase-Down Task Force v. EPA, 705 F.2d 506, 549 (DC Cir. 1983) ("Agency notice must describe the range of alternatives being considered with reasonable specificity."). In addition, proposed § 1141.16(b) identified specific criteria that any referenced cessation resource would need to meet as well as two additional criteria that the resource would need to meet if the resource was a toll-free telephone number (proposed § 1141.16(d) and two additional, but different, criteria that the resource would need to meet if it was a Web site (proposed § 1141.16(c)). The NPRM further explained that the reference to a smoking cessation resource was proposed to "be included as part of one or more of the required warnings and therefore would not appear outside of the areas specified for the required warning" (75 FR 69524 at 69541). Thus, the "notice was sufficiently descriptive of the subjects and issues involved so that interested parties [could] offer informed criticism and comments" (Air Transport Ass'n of America v. CAB, 732 F.2d 219, 224 (DC Cir. 1980) (quoting National Small Shipments Traffic Conference, Inc. v. CAB, 618 F.2d 819, 834 (DC Cir. 1980)) (internal quotations omitted)).

Our choice of a specific smoking cessation resource, 1–800–QUIT–NOW and the State quinelines to which it links, is a logical outgrowth of the proposed rule. We received many comments that discussed whether FDA should use a toll-free telephone number and/or a Web site. We also received a comment advocating that the Agency include information about contacting a physician for help quitting (see Comment 170). Numerous comments identified an existing resource (primarily 1–800–QUIT–NOW) as the preferred cessation resource for the required warnings. As discussed in section V.B.6.b of this document, many comments addressed the specific criteria proposed for the cessation resource and several comments provided reasons why 1–800–QUIT–NOW meets the criteria identified in the NPRM. In addition to comments received about whether to include a resource and, if so, what resource, as discussed in section V.B.6.d of this document, the proposed rule was sufficiently detailed for comments to raise issues regarding implementation details, such as the words surrounding the cessation resource.

We are generally adopting the criteria identified in the NPRM, including the criteria specific to a toll-free number. Our changes to the criteria are minor clarifications that were informed by comments. Thus, the requirement that the graphic warnings include a reference to a cessation resource is a logical outgrowth of the proposed rule and further notice and opportunity for comment is not necessary (Air Transport Ass’n of America, 732 F.2d at 224 ("An Agency adopting final rules that differ from its proposed rules is required to renotice when the changes are so major that the original notice did not adequately frame the subjects for discussion. * * * The agency need not renotice changes that follow logically from or that reasonably develop the rules it proposed originally") (quoting Connecticut Light and Power Co. v. NRC, 673 F.2d 525, 533 (DC Cir. 1982))). An agency is permitted to add specific details to a rule in response to comments even if the proposed rule described the requirement in a more general manner (Chemical Manufacturers Ass’n v. EPA, 870 F.2d 177, 202 (5th Cir. 1989) (finding that EPA provided adequate notice for final rule appendices, one of which established limits for charge of certain metals, even though the appendices were not included in the
proposed rule, because there was adequate notice that the agency was considering establishing limitations “and this was all the APA demands”); Trans-Pacific Freight Conference of Japan/Korea v. Federal Maritime Comm’n, 650 F.2d 1235, 1248–49 (DC Cir. 1980) (finding that the final rule merely enumerates more specifically the type of information which the Commission sought, but parties were on notice that a requirement of more detailed reports was under consideration)).

b. Criteria for cessation resource. The NPRM included three paragraphs in proposed § 1141.16 detailing criteria that would apply, on an ongoing basis, to any cessation resource chosen in the final rule. The purpose of these proposed criteria was to ensure that the cessation information, advice, and support provided by the cessation resource are unbiased and evidence based (75 FR 69524 at 69540). Proposed § 1141.16(b) described 10 criteria that would be applied to any cessation resource chosen. Proposed § 1141.16(c) described two additional criteria that would apply if the cessation resource chosen were a Web site, and proposed § 1141.16(d) described two additional criteria that would apply if the cessation resource chosen were a toll-free telephone number. In addition, the preamble to the proposed rule provided examples and additional explanation to help clarify the proposed criteria (75 FR 69524 at 69540).

As discussed more fully in section V.B.6.c of this document, we have decided that the appropriate cessation resource is a toll-free telephone number (1–800–QUIT–NOW). Therefore, our final rule does not include the criteria proposed for a cessation resource that is a Web site. We have incorporated the two criteria proposed for a cessation resource that is a toll-free telephone number into § 1141.16(b) as paragraphs 11 and 12, deleted the proposed criteria for a Web site, and added a paragraph clarifying an issue raised in the comments.

In the following paragraphs, we summarize and respond to comments regarding our general criteria for a cessation resource, as well as criteria relating to a cessation resource that is a telephone quitline. However, because we are not choosing a Web site as the cessation resource, we do not respond to specific suggestions regarding the criteria in proposed § 1141.16(c) and other comments about criteria for a cessation resource that is a Web site.

(Comment 160) One comment suggested that the rule does not need to specify criteria for the cessation resource. Instead, this comment proposed that FDA rely on the most recent version of the Public Health Service Guideline on Treating Tobacco Use and Dependence (2008 PHS Guideline) (Ref. 66). The rationale for this suggestion was that this guideline is regularly updated to reflect new effective treatments for tobacco dependence and, therefore, the criteria would not become out-of-date. In addition, the comment asserted that the 2008 PHS Guideline is the gold standard for tobacco cessation in the United States, because it is produced by leading cessation experts, updated on a regular basis, and published by HHS.

(Response) We agree with the comment that the 2008 PHS Guideline is a valuable resource for evidence-based smoking cessation treatments. However, the purpose of FDA’s criteria is not to reference particular treatment strategies. Rather, these criteria are designed to ensure that the resource’s information, advice, and support are unbiased and evidence-based. By setting forth, requirement that the cessation resource provide evidence-based treatment strategies, the resource will be able to employ newer strategies as more research is done on the most effective approaches to smoking cessation treatments.

(Comment 161) Comments representing tobacco product manufacturers claimed that the criteria set forth in proposed § 1141.16 are unspecific or that this section uses vague terminology. One comment argued that the terminology is subject to conflicting interpretations.

(Response) We disagree. The criteria in the proposed rule, and generally adopted in this final rule, are extensive and detailed. In addition, the notice and comment process gave the public an opportunity to raise questions about our use and interpretation of specific terms. The proposed rule provided adequate detail for a number of comments to request revisions and clarifications. We have responded to the significant issues raised in the comments. As explained more fully in response to Comments 163 and 164, in the final rule, we revised the criteria to clarify that quitlines may tailor their services to meet the needs of individual callers and added more explanation and examples to the preamble to further clarify issues raised by comments. The criteria we are adopting will ensure that smokers using the referenced cessation resource receive unbiased and evidence-based services suited to their individual needs.

(Comment 162) Several comments that supported the choice of 1–800–QUIT–NOW as the cessation resource expressed concern that State quitlines would be subject to two sets of potentially inconsistent requirements because the CDC already maintains standards for these quitlines. These comments proposed that FDA specify that quitlines authorized by CDC for connection to the 1–800–QUIT–NOW network are qualified to be the cessation resource included on the required warnings.

(Response) We believe that it is important to establish criteria for the cessation resource as part of this rule to ensure that the standards reflected in these criteria will be followed for as long as the rule is in effect. We do not believe there will be any conflict between these criteria and CDC’s requirements for State quitlines that are associated with our chosen resource (1–800–QUIT–NOW). We have worked closely with CDC regarding the choice of the cessation resource and the criteria that will be required. Moreover, CDC will include the criteria in this rule in its State grantee funding requirements, and will work with leading quitline experts to review, and where necessary, update existing scripting such as to accurately reflect current FDA-approved cessation medications.

(Comment 163) Many comments from public health advocacy groups and representatives of quitlines expressed concern about the criterion in proposed § 1141.16(b)(7) regarding providing accurate and evidence-based unbiased (including with respect to products, services, persons, and other entities) and relevant to tobacco cessation.” The focus of the cessation resource should be about changing a
smoker’s behavior by providing factual information and evidence-based advice and support about tobacco cessation. Our purpose in adding to the preamble the example about derogatory statements was to emphasize that our chosen cessation resource must not provide biased information about, for example, tobacco companies. The preamble to the proposed rule contrasted derogatory statements as well as statements advocating public policy changes with factual information relevant to tobacco cessation. We conclude that this distinction should be retained in the final rule. Nonetheless, as discussed in the response to Comment 164, the final rule clarifies the distinction between providing factual information, advice, and support and providing biased opinions or advice.

(Comment 164) One comment representing quitlines expressed concern that many of the cessation resource criteria described in proposed § 1141.16(b) and the preamble to the proposed rule may interfere with the ability of counselors at a telephone quitline to tailor information to a specific caller. Specifically, this comment requested that FDA delete many of the criteria or clarify that they refer to the capacity of the quitline overall, and not to each interaction with a caller. Also, this comment requested that FDA either delete the term “unbiased” in proposed § 1141.16(b)(7), or define that term to include the concept of tailoring a call to the needs of an individual caller. In addition, this comment asked the FDA to remove the word “unbiased” from proposed § 1141.16(d)(1) regarding staff training for a telephone quitline.

(Response) We agree that this issue needs to be clarified. It was not our intent that the criteria described in proposed § 1141.16 would limit the ability of the cessation resource to tailor an interaction to the needs of the individual smoker seeking help. In fact, as discussed below, we believe that one of the many benefits of choosing a telephone quitline as the cessation resource is the ability of the resource to tailor counseling sessions to individual callers. Although we do not agree that it is appropriate to delete any of the general criteria or the word “unbiased” from § 1141.16(b)(7), we have revised the rule to reorganize the criteria described in proposed § 1141.16(b) and (d). The final rule includes a paragraph (b) describing the types of services that a cessation resource must provide generally. The criteria in § 1141.16(b)(1) through (b)(7) were previously described in proposed § 1141.16(b)(1) through (b)(7), however, we revised the introductory language to clarify that a quitline may tailor individual calls as appropriate to meet the smoking cessation needs of individual callers. Thus, for example, if a caller says that he or she has attempted to quit many times and knows what to expect, the quitline does not need to provide factual information about what smokers can expect when trying to quit. Instead, the quitline might focus the counseling on practical advice about how to deal with common issues faced by users trying to quit or evidence-based information about effective relapse prevention strategies. In addition, we changed “users” to “smokers” in § 1141.16(b)(3) for consistent terminology with the rest of the paragraph.

The final rule also contains a paragraph (c) in § 1141.16 that addresses general requirements for the cessation resource, rather than the types of information to be provided to consumers seeking information or assistance. Section 1141.16(c) is primarily composed of the criteria in proposed § 1141.16(b)(8) through (b)(10) and (d). Except for the requirements regarding staff training and the maintenance of appropriate controls, this paragraph lists prohibitions for the cessation resource. For example, the cessation resource must not provide or otherwise encourage the use of any drug or other medical product that FDA has not approved for tobacco cessation. As described more fully in the response to Comment 166, we have clarified that the cessation resource may tailor information about cessation products to meet the particularized needs of an individual caller and may provide particular FDA-approved cessation products to callers, based on availability of those products to the resource. With respect to the comment expressing concern about the use of the term “unbiased” in the staff training criterion precluding the ability to tailor information, the revisions to paragraph (b) address concerns about the ability of cessation resource staff to tailor information to the needs of an individual caller. The criterion in paragraph (c) about staff training, when read in conjunction with paragraph (b), does not preclude tailoring of information during individual calls. Therefore, it is unnecessary to delete the term “unbiased” from § 1141.16(c)(8) to address this concern. We conclude that the revised criteria in paragraphs (b) and (c) of § 1141.16 will ensure that the cessation resource has the flexibility to provide counseling about smoking cessation that is appropriate to the needs of an individual caller while still ensuring that the resource does not provide opinions, advice, or support that are biased or not supported by appropriate evidence.

(Comment 165) One comment representing quitlines suggested that FDA either delete the criterion described in proposed § 1141.16(b)(10) that prohibits the cessation resource from encouraging “the use of any non-evidence-based smoking cessation practices,” or replace the word “practices” with “treatment.” This comment explained that practices such as coping strategies for dealing with cravings have not been as rigorously tested as medications and may not be considered evidence-based. This comment asserted that the criterion in proposed § 1141.16(b)(3), requiring a cessation resource to provide practical advice about how to deal with common issues faced by users trying to quit, adequately addresses this issue.

(Response) We understand the concerns expressed by this comment and agree that a cessation resource should be permitted to discuss coping strategies for dealing with cravings (e.g., chewing gum) that may not have been rigorously tested in a scientific manner. However, because the distinction between treatment and practices is unclear, we conclude that a broad term such as “practices” is appropriate in order to ensure that evidence-based research is being used to provide callers with effective services. Using the broader term “practices” also avoids the possibility that definitional questions about whether something is a treatment will interfere with the ability of the cessation resource to provide effective cessation services to smokers. Deleting proposed § 1141.16(b)(10) completely, or replacing the word “practices” with “treatment,” may result in cessation resources encouraging non-evidence-based practices even though evidence-based practices are available. Section 1141.16(b)(3) permits the cessation resource to provide practical advice, and the practices described in the comment would be considered “practical advice” rather than “non-evidence-based practices.” In addition, as discussed in the comment, a cessation resource is permitted to tailor each counseling session to the needs of the individual caller.

(Comment 166) FDA received several comments relating to the cessation resource providing or discussing particular smoking cessation drug products. One comment representing a manufacturer of smoking cessation drug products suggested that the Agency permit the resource to provide one or more FDA-approved over-the-counter...
cessation products, but not include language in the rule that prohibits the cessation resource from “advertising or promoting a particular product.” This comment claimed that there is evidence that recognizable brands of smoking cessation products can be important tools to promote cessation (Ref. 84). Comments representing telephone quitlines and a public health advocacy group requested that FDA clarify that simply mentioning a particular cessation product does not constitute advertising or promoting a particular product, so long as the resource makes clear it does not recommend the use of one cessation product or brand over another.

(Response) The final rule has been revised to clarify that a cessation resource may tailor a discussion of cessation medications for an individual caller. As noted in the preamble to the proposed rule, under the criteria the cessation resource may provide one or more FDA-approved over-the-counter cessation products, provided that it does so in a manner that does not advertise or promote a particular product (75 FR 69524 at 69540). We agree that, in the context of individual counseling, one medication may be suggested over another, based on an individual smoker’s health needs and prior experience with cessation medications. For example, a quitline counselor may take into account warnings, precautions, and contraindications identified in the labeling of a specific drug product in relation to an individual caller. Also, a quitline counselor may suggest a particular medication based on the caller’s prior experience with cessation medications (e.g., not recommend a medication that previously caused significant side effects or did not work; recommend a medication that worked well in the past). In addition, a cessation resource may provide one or more FDA-approved over-the-counter cessation products, based on availability of the product(s) to the resource. A cessation resource may also mention the availability of free medication, provided it does so in a manner that does not advertise or promote a particular product. However, the resource must not advocate or promote a cessation product, such as by recommending the use of particular cessation products or brands over others to callers generally. All products that have been approved with smoking cessation claims have been found by FDA to be safe and effective for the approved indication. Even if there might be benefits associated with brand recognition for a smoking cessation drug product, we do not believe that it is appropriate for the cessation resource that we include in a required warning to promote any particular product.

(Comment 167) Several comments proposed that additional criteria be added to the criteria proposed in the NPRM. One comment suggested adding an additional criterion that the cessation resource must provide evidence-based advice regarding the protection of children and other nonsmokers from secondhand smoke. This comment reasoned that two of the warning statements address the dangers of secondhand smoke and the cessation resource should be prepared to counsel smokers who seek assistance after seeing these messages. Another comment recommended adding a criterion to prohibit the cessation resource from promoting a tobacco industry cessation program. This comment claimed that research has demonstrated that tobacco industry sponsored cessation resources either have no effect on smoking prevalence or actually cause increased smoking (Refs. 85 and 86). One comment from a submitter representing quitlines recommended the addition of a new criterion that would require the cessation resource to provide proactive, multi-call counseling services. The comment claimed that there is evidence these types of services are effective.

(Response) We recognize that there could be additional criteria for a cessation resource that would require the resource to provide broader services. However, we have designed the criteria in this final rule to focus on the minimum services that must be provided by an effective cessation resource and the minimum standards the resource must meet. We are mindful that existing cessation resources have varied budgets and do not want to require additional standards that, while possibly beneficial, would disqualify some effective treatment programs that do not have the resources to provide these services. We note, however, that the criteria described in § 1141.16 (b) and (c) do not preclude any cessation resource from providing additional unbiased, evidence-based cessation information, advice, and support. With respect to prohibiting the promotion of a tobacco industry cessation program on the basis that they are not effective, we conclude that the addition of a separate criterion is unnecessary. The cessation resource that will appear in the required warnings—1–800–QUIT–NOW—is run by government entities, and the criteria are designed to ensure that the resource provides cessation information, advice, and support that are unbiased and evidence-based.

(Comment 168) One comment recommended that an additional role of a cessation resource should be to direct smokers (who request it) to local specialist face-to-face treatment services and to provide accessible information on Medicaid, Medicare, and other large insurers’ coverage for tobacco dependence treatment.

(Response) Our primary objective in requiring that referenced cessation resources comply with the criteria is to ensure that the cessation resource chosen provides evidence-based counseling to help smokers quit. Our criteria are designed to ensure that the cessation resource will continue to meet certain minimum standards. While not required by the criteria in this regulation, a referenced cessation resource is not precluded from providing additional relevant factual information, such as information about reimbursement for tobacco dependence treatments.

c. Choice of cessation resource. The NPRM did not specify a particular cessation resource. Rather, it noted that there are a number of possible alternatives, including use of an existing or new quitline or Web site, where smokers and other members of the public can obtain current unbiased, factual smoking cessation information (75 FR 69524 at 69540). Based on the information before the Agency, including the information provided in the comments, we have chosen the Network, which uses the toll-free telephone number 1–800–QUIT–NOW (1–800–784–8669), as the cessation resource to include on all nine required warnings. The Network is the single point of access to reach State-based quitlines in all 50 states, the District of Columbia, Puerto Rico, and Guam. Since 2005, CDC and NCI have partnered with States to create the Network. NCI manages the 1–800–QUIT–NOW telephone number, along with appropriate telecommunications and routing infrastructure, to ensure that calls are transferred to the appropriate State or territory quitline based on the area code of the caller. Calls from U.S. territories that do not have a quitline are routed to an NCI-run quitline. CDC and individual States or territories provide the funding for the quitlines. CDC provides funding through cooperative agreements as part of the National Tobacco Control Program.

It is discussed more fully in the context of comments and responses in the following paragraphs, we find that this cessation resource, which was strongly
favored in many comments, will provide people in the United States with access to unbiased, evidence-based smoking cessation information, advice, and support. We have determined that including this cessation resource as part of the required warnings will increase the likelihood that smokers will quit smoking and thereby provide substantial public health benefits by reducing the life-threatening consequences associated with continued cigarette use. Therefore, we conclude that including a reference to 1–800–QUIT–NOW as part of all the required warnings is appropriate for the protection of the public health.

(Comment 169) Comments favoring inclusion of a cessation resource generally preferred the use of a telephone quitline. In particular, most of these comments advocated the use of 1–800–QUIT–NOW. The comments pointed to a robust body of evidence showing that proactive telephone counseling is effective in helping smokers to quit successfully. Several comments cited statistics from individual State quitlines about the types of services provided and success rates. In addition, several comments asserted that quitlines associated with 1–800–QUIT–NOW generally meet the criteria for a cessation resource specified in the NPRM.

Many comments discussed the advantages of choosing 1–800–QUIT–NOW. In support of the choice of a telephone quitline over a Web-based cessation resource, several comments noted the broad penetration of telephone access, including among low income and minority populations. These comments noted that Internet access has much lower penetration among the American public, particularly in many groups with high rates of smoking (e.g., low income, low level of education). Many comments that advocated the use of 1–800–QUIT–NOW noted that it has an existing infrastructure that is available in all 50 states, the District of Columbia, Puerto Rico, and Guam. One comment stated that all quitlines associated with 1–800–QUIT–NOW are at least several years old.

Several comments argued that inclusion of 1–800–QUIT–NOW on cigarette packages could address issues relating to poorer smoking cessation outcomes among racial and ethnic minorities, as well as populations with low income and/or low education. One academic noted that smokers in these groups try to quit as often as other smokers but are less likely to use effective treatments (citing Ref. 87). The comment claimed that adding 1–800–QUIT–NOW to the required warnings holds unprecedented potential to close the gaps and disparities in treatment awareness and use. One comment representing a State quitline argued that quitlines can help address racial or ethnic disparities in access to effective tobacco treatment. For example, African-Americans have been significantly overrepresented among quitline callers in California, relative to the proportion of African-American tobacco users in that State. Several comments stated that quitlines provide services in languages other than English, particularly Spanish, and provide materials to important population groups (e.g., youth, pregnant women, racial/ethnic populations). One comment representing a State quitline asserted that quitlines can help address disparities related to socioeconomic status. In California, utilization of quitline service is highest among low socioeconomic status tobacco users. This comment also claimed that the attractiveness of quitlines to tobacco users with low socioeconomic status is related to the fact that services are provided without a charge and are accessible by telephone, eliminating the need to arrange for transportation or child care. According to this comment, these factors can be significant barriers for individuals with modest resources. Another quitline provider stated that quitlines are disproportionately used by the chronically ill and those who are socially and economically stressed. This comment claimed that, arguably, these groups have the greatest need for support because they have a higher prevalence of smoking and are disproportionately affected by tobacco-related health concerns.

One comment representing a public health advocacy group pointed out that designation of a single quitline number would avoid the difficulty of manufacturers having to print different dialing information depending on where the cigarette package will be sold. (Response) We agree with comments that a telephone quitline is the most effective means of ensuring that all Americans have access to unbiased, evidence-based smoking cessation information, advice, and support. We have decided to use the Network as the cessation resource and its portal number, 1–800–QUIT–NOW, will be included as part of electronic files for the required warnings that are available in the IBR document described in section V.B.4 of this document.

A key factor in our decision is that the evidence underscoring the effectiveness of telephone quitlines is well documented. The 2008 PHS Guideline found that quitlines significantly increase abstinence rates compared to minimal or no counseling interventions. The 2008 PHS Guideline also found that use of quitline counseling in conjunction with cessation medication significantly improves abstinence rates compared to the use of medication with minimal or no counseling (Ref. 66 at pp. 91–92; see also Ref. 88). Consequently, quitlines are an important part of the HHS Tobacco Control Strategic Action Plan (Ref. 89).

In addition, there is evidence that knowing about the availability of a quitline increases quit attempts and successful cessation even among smokers who do not call the quitline (Ref. 88 (finding “[t]elephone quitlines provide an important route of access to support for smokers, and call-back counselling enhances their usefulness”). For example, one study of the effect of a smokers’ hotline as an adjunct to self-help manuals found “it is unlikely that higher abstinence rates among users of the hotline) accounted for the total differences in outcome between hotline and manual only counties. It is possible that simply knowing that telephone help was there if needed enhanced abstinence even among nonusers” (Ref. 82). A CDC report hypothesized that a possible explanation is that “knowledge of cessation services, engendered through promotion, increases tobacco users’ belief in the normalcy of quitting, which may lead to increased quit attempts among people who have access to the services, even those who do not use them” (Ref. 90).

Another factor that we considered in choosing a telephone quitline is that telephone access within the United States is nearly universal. According to a 2010 Federal Communications Commission statistical report, household telephone subscribership in the United States was 96 percent in March 2010. This report shows that, even among households with annual incomes as low as $25,000, telephone penetration was over 90 percent in 2000, including among African-Americans and Hispanics (Ref. 91). Currently, Internet use and broadband penetration is much lower than telephone penetration in the United States, particularly among low income groups, certain racial and ethnic minorities, and households with low education levels (Ref. 92).

Beyond their wide accessibility, quitlines are also successful in helping certain high risk populations and other important demographic groups. One comment asserted that low income and uninsured smokers, those with the
lowest levels of formal education, and those in racial/ethnic populations with the highest smoking rates try to quit as often as other smokers, but are far less likely to use effective treatments. For example, smokers in several racial and ethnic groups attempt to quit as often as or more often than nonminority smokers but use effective treatments less often and have lower success rates (Ref. 66 at p. 156). Similarly, low socioeconomic status smokers or those with limited education express significant interest in quitting and appear to benefit from treatment. However, these smokers are less likely to receive cessation assistance (Id. at p. 151). One study concluded that non-Hispanic black and Hispanic smokers who attempted to quit smoking were significantly less likely to use cessation aids, and that this has implications for successful quitting among minority smokers (Ref. 87). Several comments, however, explained that at least some quitlines receive a disproportionate numbers of calls from certain minority or disadvantaged populations (see, e.g., Ref. 93). In light of the overall low rates of calls to quitlines (approximately 1 percent of smokers call quitlines, although this percentage varies by State and how much the State promotes its quitline), even a disproportionately high volume of calls from important demographic groups is not enough to alter the overall quit rates for these groups. However, as discussed in section V.B.6.a of this document, there is strong evidence that there will be an increase in call volume to quitlines after the required warnings appear on cigarette packages and in cigarette advertising. This increase in use of quitlines could have an important impact on high risk and other important demographic groups if they continue to constitute a significant percentage of calls to quitlines.

In addition, a telephone quitline provides an excellent opportunity to tailor counseling sessions and provide additional materials for specific populations. The 2008 PHS Guideline also found that individually tailoring materials to address smoker-specific variables (e.g., support sources available, time lapse since quitting, concerns about quitting) has been shown to be effective and have broad reach (Ref. 66 at p. 92). Several comments noted that virtually all State quitlines associated with 1–800–QUIT–NOW provide specialized materials to special populations, including pregnant women, racial and ethnic populations, and youth. Quitlines can also provide information (e.g., about the negative health consequences of smoking or the health benefits of quitting) to smokers who are not ready to quit but who want additional information.

With respect to our choice of the Network and its telephone number, 1–800–QUIT–NOW, for the quitline cessation resource, we have determined that this resource will fulfill the goal to provide a place where smokers and other members of the public can obtain smoking cessation information from staffed trained specifically to help smokers quit by delivering current, unbiased, and evidence-based information, advice, and support. The quitlines that compose the Network, the telecommunications infrastructure supporting the Network, and the telephone number, 1–800–QUIT–NOW, are already well established and provide smoking cessation services to people throughout the United States. Comments that advocated the use of a specific quitline referred to 1–800–QUIT–NOW as the preferred cessation resource. By using an existing resource, resource, and telephone number, we can leverage the Network’s established structure and experience providing cessation services. This choice also avoids the costs associated with establishing a new quitline.

In addition, we agree with comments that the individual State and territory quitlines that are associated with 1–800–QUIT–NOW generally meet the criteria specified in § 1141.16(b). We understand, however, that these quitlines have some differences in funding resources and consequently provide differing levels of service. For example, some State quitlines provide longer hours of service than others. Based on the statistics provided in some comments, it is possible that not all of the individual State and territory quitlines associated with 1–800–QUIT–NOW meet all of the criteria we are adopting in § 1141.16(b). To assure that these criteria are met, CDC will include these criteria beginning with its 2013 National Tobacco Control Program funding opportunity announcement and HHS will monitor the quitlines for compliance with the criteria on an ongoing basis and will take appropriate steps to address any noncompliance.

(Comment 170) One medical organization suggested that the reference to the smoking cessation resource in the required warnings should also include a message encouraging smokers to contact their physician or health care provider. This comment cited studies to support the proposition that physician advice is effective in encouraging smoking cessation (citing, e.g., Ref. 94). This comment also noted that both Australian and European Union graphic warnings recognize the role that physicians play in assisting patients’ cessation efforts.

(Response) We agree that physicians, particularly primary care physicians, and other health care providers are a very helpful resource for encouraging smokers to quit (Ref. 66 at p. 35). However, we decline to include language on the required warnings encouraging smokers to see their doctor. Many Americans do not have an ongoing relationship with a physician. Recent evidence indicates that the United States may be suffering from a shortage of primary care physicians, making it less likely that they would be available to provide cessation information to smokers (see Ref. 95 for statistics on decreasing numbers of U.S. medical school graduates selecting a family medicine career). In addition, unlike the selected quitline, we would not have a practical means to monitor health care provider compliance with the criteria the Agency is establishing in § 1141.16(b). Studies indicate that rates of physician adherence to similar practice guidelines for smoking cessation advice vary widely (see Ref. 96). For these reasons, it is preferable to include a reference to 1–800–QUIT–NOW on the required warnings.

With respect to our choice of the quitlines that are associated with 1–800–QUIT–NOW, we would be a useful cessation resource in addition to a telephone quitline. For example, some State quitlines frequently refer people to their primary care physicians (e.g., if a caller has further questions about the use of medications).

In addition, there is limited space available for including information about a cessation resource. The size of the required warnings is relatively small and the textual warning statement and color graphic image included in each warning must be clear, conspicuous, and legible as required by section 4 of FCLAA. In light of the limited space available, we have determined that including an additional message encouraging smokers to contact their physician or health care provider is not appropriate at this time.

(Comment 171) Some comments urged FDA to include a Web site as a cessation resource. Generally these comments suggested that a Web site would be a useful cessation resource in addition to a telephone quitline. For example, one public health advocacy group noted that there are advantages to utilizing both quitlines and Internet resources. According to this comment, while quitlines provide individualized telephone counseling, a Web site provides support 24 hours per day. One comment from a public health advocacy group claimed that smokers search online for smoking cessation assistance every year, and it is
particularly important for the required warnings to include Web-based resources because there are a large number of Internet sites that ostensibly offer quitting assistance but do not offer evidence-based cessation help. Several comments acknowledged that the 2008 PHS Guideline did not find enough evidence to recommend computer-based interventions, but noted that the 2008 PHS Guideline also concluded that these interventions remain promising. Some comments also noted that Internet use is low in many groups with high rates of smoking (e.g., low-income, racial and ethnic minority groups).

However, several comments advocating inclusion of a Web site resource noted that many cessation services, including many quitlines and health plans, are utilizing the Internet to provide combined telephone counseling and Web-based cessation treatment. One comment suggested that as American culture adopts different forms of communication, it will be important to assess the effectiveness of using new technologies and approaches. This comment encouraged FDA to fund research to learn which approaches will encourage the most people to quit smoking.

One comment from the tobacco industry claimed that reference to a smoking cessation Web site may raise additional implementation issues and requested an opportunity to comment in advance of such a requirement. This comment did not identify any specific issues associated with reference to a smoking cessation Web site.

(Response) We recognize that Web sites are another important source of smoking cessation information and interventions. Although the 2008 PHS Guideline did not recommend the use of Web-based interventions, it concluded that "[g]iven the potential reach and low costs of such interventions * * * they remain a highly promising delivery system for [treatment] tobacco dependence" (Ref. 66 at p. 94). We also recognize that Internet use is highest among younger populations, and thus might be a useful tool to intervene with young smokers, given that maximum cessation benefits are gained by quitting at a younger age. Furthermore, Web sites can provide information to smokers who are not ready to quit but who are seeking additional information about cessation.

However, we have decided not to include a Web site as the cessation resource incorporated in the required warnings. For the reasons explained more fully above, we find that a telephone quitline is a better overall cessation resource than a Web site.

There is stronger scientific support that telephone quitlines are effective, they are more widely available to a broader cross section of Americans, particularly groups with higher rates of smoking and lower access to cessation services, and there is a strong national quitline infrastructure in place. In light of the limited space available on the required warnings and the need to ensure that the graphic images and textual warning statements are clear, conspicuous, and legible, we do not think it is appropriate at this time to include both a telephone quitline and a Web site address on all required warnings. We intend to evaluate this possibility in the future when we are designing and testing revised versions of the required warnings.

d. Implementation issues. Proposed § 1141.16(a) stated that a required warning must include a reference to a smoking cessation assistance resource as specified in the IBR document. The preamble to the proposed rule explained that the smoking cessation information would be included as part of the required warning and would not appear outside of the areas specified for the required warning. In other words, the cessation resource would be within the top 50 percent of the front and rear panels of cigarette packages and within the 20 percent of the area of advertisements occupied by the required warning (75 FR 69524 at 69541). We received several comments regarding how a cessation resource should appear in the required warning and other implementation issues relating to inclusion of a cessation resource in the required warning. These comments and our responses are summarized in the following paragraphs.

(Comment 172) A comment representing small tobacco product manufacturers expressed confusion about whether FDA would add the reference to a cessation resource to the required warnings or whether a manufacturer would have to select the cessation resource and incorporate it into the required warning. The comment noted a preference that FDA provide the specific language for the cessation resource. However, one small tobacco product manufacturer asked that FDA provide a variety of options for cessation resources and include those options in the electronic files for the required warnings provided by the Agency.

(Response) We have selected 1–800–QUIT–NOW as the cessation resource that appears on the required warnings. The required warnings in the IBR document include the reference to the cessation resource, 1–800–QUIT–NOW. We disagree with the request that we provide a variety of options for cessation resources and include those options in the electronic files for the required warnings. Such an approach could be confusing to consumers, because the required warnings would appear with a different cessation resource on different packages of cigarettes and in different advertisements. Also, it would be difficult to monitor many cessation resources to ensure that each one meets the criteria established in § 1141.16(b) and (c). By choosing one, existing toll-free telephone number that is under the control of NCI, provides access to consumers throughout the country, and includes State quitlines that have cooperative agreements with CDC, we have assurances that our cessation resource criteria will be followed.

(Comment 173) Several comments mentioned that an increase in the volume of calls to State quitlines may increase funding needs. These comments suggested that additional resources should be provided to State quitlines.

(Response) We expect that inclusion of 1–800–QUIT–NOW on the required warnings will increase the volume of calls to State quitlines. While some quitlines may currently have some additional capacity, there will likely be need for additional resources. In the fiscal year 2012 President’s Budget, there is $25 million from the Prevention and Public Health Fund allocated for CDC to spend on the National Network of Tobacco Cessation Quitlines. Additionally, the Centers for Medicare and Medicaid Services is working with the State Medicaid Directors to permit tobacco quitlines as an allowable Medicaid administrative activity.

(Comment 174) One comment encouraged FDA to require that the cessation resource be displayed as a telephone number (1–800–784–8669 in addition to 1–800–QUIT–NOW because some wireless phones do not have letters on the keypad. However, another comment representing a quitline expressed the view that it is important to use the letters in 1–800–QUIT–NOW rather than the telephone number because it is itself a cogent cessation message.

(Response) We agree there would be benefits to identifying the cessation resource using 1–800–QUIT–NOW as well as the telephone number 1–800–784–8669. However, as explained previously, there is very limited space for identifying the cessation resource. The use of 1–800–QUIT–NOW is a way to provide the number for people to call.
while in the same space providing information about what the number is for. Using less space for the cessation resource helps ensure the required warning remains clear, conspicuous, and legible and appears within the specified area. Moreover, the use of letters is likely to be easier for people to remember. The Agency also believes most telephones in use still include letters on keypads and that toll-free telephone numbers are frequently identified using these letters. As stated previously, we will also conduct research and keep abreast of scientific developments regarding the efficacy of various required warnings and the types and elements of various warnings that improve efficacy, including elements related to identifying cessation resources.

(Comment 175) Several comments addressed the words that would signal the appearance of a cessation resource. These comments described experience from New Zealand that showed increases in both quitline number recognition and the number of callers reporting cigarette packages as the source for learning the quitline number after the introduction of new graphic warnings with a redesigned reference to a cessation resource (i.e., “You CAN quit smoking. Call Quitline 0800 778 778, or talk to a quit smoking provider”). The prior warning said “For more information call” next to a telephone number. According to one study, there was a 24 percent increase in reported recognition of the quitline number after this change (Ref. 69). Also, in the first full year after the introduction of the new graphic warnings, the volume of calls to the quitline increased significantly and 26 percent of callers reported cigarette packages as the source of the number (compared to 7.5 percent the prior year) (Id., Wilson 10/10). One academic researcher suggested a short, direct “call to action” phrase to motivate cessation behavior. Similarly, another comment from an academic institution suggested that the warnings provide the smoker with avenues to take in order to quit and simultaneously instill confidence in the user that he or she can take action.

(Comment 176) One comment from an academic institution encouraged FDA to require, in addition to a quitline number, clear encouragement of action steps for quitting. This comment recognized that space on the required warnings is limited and suggested that package inserts and onserts are one way of accomplishing this without compromising the visual impact of the graphic warnings.

(Response) A requirement to add onserts or inserts is beyond the scope of this rulemaking and, therefore, we decline to require them here.

VI. Comments Regarding Implementation Issues

A. Technical Issues Regarding Compliance

Section 1141.12 refers to “Cigarette Required Warnings,” which is incorporated by reference (IBR) in accordance with 5 U.S.C. 552(a) and 1 CFR part 51. The IBR document includes electronic files of images that must be included on all cigarette packages, and in all cigarette advertisements.

In response to the proposed rule, some comments, including comments from cigarette manufacturers and tobacco industry trade associations, raised issues relating to the electronic files and the implementation of the graphic warnings on cigarette packages and in cigarette advertisements. Those comments, and FDA’s responses, are discussed in the following paragraphs.

(Comment 177) Comments from two tobacco product manufacturers stated that they would need to make certain technical adjustments to the single sized graphic warnings published with the proposed rule in order to ensure that the warning fits packaging of varying sizes and shapes. According to the comments, if FDA provided only the single warning format published with the proposed rule, the company would need to adjust the height-to-width ratio (i.e., aspect ratio) of that warning in order to cover 50 percent of the front and rear panels of various package configurations. However, adjusting the aspect ratio, such as by elongating or compressing the warning, could distort the graphic image and/or textual warning statement. These comments recommended that FDA ensure that manufacturers are able to adapt the graphic warnings to fit cigarette packages of varying sizes and shapes and provide guidance about how to adapt the warnings.

(Response) We agree that the size and shape of certain packages might require companies to adapt the electronic files provided in the IBR document. To help prevent distortion of the image and text and to minimize the need for adaptation, we are providing electronic files in different formats designed to fit packaging of various sizes and shapes. We are adding language to the IBR document that provides instructions as to when each of the formats must be used. The instructions are based on the aspect ratio of the display area where the required warning must appear. This language also describes the requirements companies must follow when adapting the electronic files provided in the IBR document. For example, the requirements state that each of the different elements of the warning (i.e., the image, the textual warning statement and reference to the cessation resource) must, to the extent possible, maintain the relative scale and proportions of the elements as displayed in the relevant electronic file, and the positions of each of these elements must be maintained relative to each other.

(Comment 178) Two comments from cigarette manufacturer requested clarification concerning how companies should incorporate the required warnings on packages with hinged lids. These comments stated that the content of warnings printed on the hinged lids can shift up or down by about 1 mm at the point where the lid meets the front of the pack due to normal variations in production of the packaging. These comments recommended that FDA design the warnings with all text located either above or below the hinge, or allow for minor variations in how the graphic warnings appear on cigarette packs due to this manufacturing variability.

(Response) We agree that the integrity of the warning must be maintained on packages to ensure that the warning is clear and legible. To clarify the requirements that companies must follow when they adapt the electronic files for hinged lid packages, we have added language to the IBR document that permits companies to separate the segments of text or add or subtract one line of text within the textual warning statement so that the line at the location where the lid is to open cuts across the background space between two lines rather than through a line of text. This provision will allow companies to adapt the electronic files provided in the IBR document to ensure that the textual warning statement is not severed when the package is opened and is clear, conspicuous, and legible in accordance with section 4 of FCLAA. According to this language in the IBR document, companies are specifically prohibited from severing any word in the textual
to move the upper boundary of the display area of the warning so that it runs along a line that is parallel to and not more than 0.375 inches from the top edge of the package. The companies compress the vertical size of the image and then shift it down (so that it stays within the top 50 percent of the package). This language also requires companies who do this to ensure that, to the extent the file must be adapted to fit the dimensions of the warning area below the closure, the proportions of the required warning must be maintained. In addition, the instructions in the IDR document specify that the closure and the portion of the packaging that appears between the top edge of the package and the upper boundary of the display area of the required warning must be either solid black or solid white. This will allow companies to continue to produce “soft pack” style packaging with closures at the top center of the pack without obstructing the required warning. However, if we determine that it would be technologically feasible to incorporate the required warnings on “soft pack” style packaging without the need to adapt the warning as set out in the electronic files, we will permit the required warnings to appear on the bottom 50 percent of the packaging. We have determined that requiring the warnings appear in the upper portion of the package, as specified by the Tobacco Control Act, will result in warnings that are more prominent, more salient, and more effective than warnings appearing at the bottom of the package.

(Response) We recognize the technological difficulty of incorporating the required warnings on “soft pack” style packaging. Given the paramount need to incorporate the warning without obstructing any of the discrete elements of the warning (i.e., the image and the textual warning statement) or the reference to a cessation resource, the final rule permits companies to adapt the warnings on “soft pack” style packaging by moving the warning below the closure in accordance with the requirements included in the IDR document. The IDR document states that this is only permitted when it is not technologically feasible to incorporate the required warnings on “soft pack” style packaging without the need to adapt the warning as set out in the electronic files provided in the IDR document. The requirements included in the IDR document allow companies using “soft pack” style packaging only to move the upper boundary of the display area of the warning so that it runs along a line that is parallel to and not more than 0.375 inches from the top edge of the package. The companies compress the vertical size of the image and then shift it down (so that it stays within the top 50 percent of the package). This language also requires companies who do this to ensure that, to the extent the file must be adapted to fit the dimensions of the warning area below the closure, the proportions of the required warning must be maintained. In addition, the instructions in the IDR document specify that the closure and the portion of the packaging that appears between the top edge of the package and the upper boundary of the display area of the required warning must be either solid black or solid white. This will allow companies to continue to produce “soft pack” style packaging with closures at the top center of the pack without obstructing the required warning. However, if we determine that it would be technologically feasible to incorporate the required warnings on “soft pack” style packaging without the need to adapt the warning as set out in the electronic files, we will permit the required warnings to appear on the bottom 50 percent of the packaging. We have determined that requiring the warnings appear in the upper portion of the package, as specified by the Tobacco Control Act, will result in warnings that are more prominent, more salient, and more effective than warnings appearing at the bottom of the package.

(Response) We have determined that companies can use cellophane tear tapes, and the final regulation does not prohibit such use on cigarette packaging. We further have determined that it is technologically feasible to use clear tear tapes in a manner that does not obstruct the required warning before the cigarette package is opened for the first time, and note that clear tear tape is widely used on product packaging in other countries that require graphic warnings. We are not aware that this has created any substantial technical difficulty in the production of cigarette packages, nor are we aware that clear tear tape has led to any significant obstruction of the graphic warnings. If a company has a unique problem with regard to packaging, it should raise this issue with us, and the difficulty can be addressed on an individual basis. We decline to change the final regulation to allow the required warnings to appear on the bottom 50 percent of the packaging. We have determined that requiring the warnings appear in the upper portion of the package, as specified by the Tobacco Control Act, will result in warnings that are more prominent, more salient, and more effective than warnings appearing at the bottom of the package.

(Response) We have revised the proposed IDR document and the...
Within the specified areas, especially given the variety of font styles included in the nine selected warnings.

(Comment 184) Several comments requested that FDA provide fonts for the textual warning statements in each of the required warnings.

(Response) For English and Spanish language warnings, the font size and font style is built into the electronic files provided in the IBR document. For advertisements in foreign languages other than Spanish, companies must comply with the font size requirements in section 4(b)(2) of FCLAA and any format requirements included in the IBR document. In all situations, it is the advertiser’s responsibility to ensure that the textual statements appear in conspicuous and legible type and that the required warning complies with the format specifications set forth in section 4 of FCLAA.

(Comment 185) One comment requested that FDA provide instructions on how companies should combine and display the images developed for use in small advertisements less than 12 square inches with the required textual warning statements.

(Response) We recognize that the small size of these advertisements presents additional challenges. We are providing an electronic file of the graphic that must be used for warnings appearing in advertisements that are less than 12 square inches. Companies may combine the graphic and the textual warning statement or otherwise adjust the layout of the warning so long as each warning includes the specified graphic and an appropriate textual warning statement. It is the advertiser’s responsibility to ensure that the textual warning statement appears in conspicuous and legible type and that the combined warning complies with the format specifications set out in section 4 of FCLAA.

(Comment 186) Several comments recommended that FDA require that companies reproduce the color graphics in the industry standard four-color (CMYK) printing process.

(Response) The electronic files provided in the IBR document were built with CMYK printing standards. The directions in the IBR document specify the use of CMYK printing standards.

(Comment 187) One comment requested that FDA make available “printers proofs” for each of the required warnings in order to ensure optimal clarity.

(Response) We have determined that the electronic files provided in the IBR document will be adequate to ensure necessary clarity. Thus, we do not believe it is necessary to provide “printers proofs” for the warnings.

(Comment 188) One comment requested that FDA adopt required warnings with consistent dimensions to allow for accurate incorporation into manufacturers’ packages and advertisements.

(Response) We decline to adopt this recommendation. As discussed previously, our selection of the nine final required warnings was based in part on our desire for a diverse set of warnings in a variety of different styles (e.g., photographic and illustrative, different fonts and font sizes) and diversity of human images (e.g., race, gender, age) in order to reach the broadest range of target audiences. We have determined that this variety will enhance the effectiveness of the warnings and help to delay potential wear out of the warnings. Because of the diversity of styles and images, some warnings have slightly different dimensions than others.

(Comment 189) One comment recommended that FDA provide layered high resolution .tif or .eps files, with text supplied as a separate layer. Another comment recommended that FDA provide images as .jpeg files.

(Response) The electronic files included in the IBR document are built as .eps files, with separate layers for text and images. Companies will be able to convert the files into .jpeg files if needed.

B. Textual Statement Color Formats

In the document entitled “Proposed Required Warning Images” included in the docket for the NPRM, FDA provided two formats for each proposed required warning: one with the warning statement in white text on a black background and one with the warning statement in black text on a white background, under section 4(a)(2) and (b)(2) of FCLAA. Several comments offered suggestions regarding the use of the color combinations, which we have summarized and responded to in the following paragraphs.

(Comment 190) A few comments suggested that FDA specify that the required warnings on cigarette packages and advertisements contain required warnings in either the white text on black background or the black text on white background format, whichever the Agency chooses to most effectively communicate the warnings.

(Response) We disagree. Section 4(a)(2) of FCLAA states that for cigarette packages, the “text shall be black on a white background, or white on a black background.” Similarly, for advertisements, section 4(b)(2) of
FCLAA states that the text of the statement in the required warning “shall be black if the background is white and white if the background is black.” We interpret these statutory requirements to mean that companies can use either of these two text/background color combinations on the package or in the advertisement.

(Comment 191) One comment recommended that the word “CANCER” always appear in red as part of the health warnings on cigarette packages and advertisements.

(Response) We disagree. As stated previously, section 4(a)(2) and (b)(2) of FCLAA prescribe the colors for the textual statements on packages and advertisements (e.g., white text on black background or black text on white background). FDA has the authority to change the format of the textual statements if such a change would promote greater understanding of the health risks associated with cigarette smoking. If we determine at a later date, that requiring the word “CANCER” to appear in red font will promote a greater understanding of smoking’s risks, we may propose new iterations of the required warnings in future rulemakings.

C. Random Display and Rotation of Warnings

The proposed rule did not specifically address the statutory requirements for the warnings on cigarette packages to be randomly displayed in each 12-month period and for quarterly rotation of the required warnings in advertisements, under section 4 of FCLAA. However, FDA received several comments on this issue. These comments, and FDA’s responses, are included in the following paragraphs.

(Comment 192) One comment expressed concern that cigarette manufacturers may only use some of the nine new required warnings on their cigarette packages and requested that FDA require companies to use all the required warnings in equal numbers.

(Response) We agree that all cigarette manufacturers must use all of the nine required warnings on their cigarette packages and requested that FDA require companies to use all the required warnings in equal numbers.

(Comment 193) One comment recommended that retailers be exempted from any requirement to rotate the required warnings for each brand they sell in stores.

(Response) We decline to address this issue here, as it is beyond the scope of the current rulemaking.

(Comment 194) Several comments recommended that FDA rotate the graphic warnings to prevent overexposure. The comments also noted that different warnings will have different impacts on the various segments of the population, further emphasizing the need to rotate the warnings.

(Response) It is unclear whether these comments were referring to the quarterly rotation of the required warnings in advertisements or the need to refresh the warnings on a regular basis. We agree that rotation of the warnings is important to delay wear out and to ensure that all population segments are exposed to the different warnings in as equal a number of times as is possible. In accordance with section 4(c)(2) of FCLAA, the required warnings must be rotated quarterly in cigarette advertisements. See section ILE of this document for additional information regarding FDA’s efforts to delay or prevent wear out.

(Comment 195) One comment recommended that FDA monitor the rotation of required warnings in cigarette advertisements to ensure compliance by all manufacturers, distributors, importers, and retailers.

(Response) We agree with this comment. We will monitor rotation and ensure compliance, which will include the review and approval of warning plans submitted to the Agency in accordance with section 4(c) of FCLAA.

(Comment 196) One comment suggested that manufacturers be given broad discretion in complying with the requirements that they include the required warnings on all cigarette packages such that in each 12-month period all of the different warnings appear in as equal a number of times as is possible on each brand of the product (see 15 U.S.C. 1333(c)). The comment stated that its printing machines, and in particular the print cylinders, used to produce “soft pack” style packaging only allows the company to print five images per roll and does not allow for warnings to be die cut and collated. Because “soft pack” style packaging only accounts for about 10 percent of all packages distributed and sold, this style of packaging frequently is printed in small batches and for some, is printed only once per year. The comment stated that in light of these production constraints, it would be impossible to apply and distribute “soft pack” style packages displaying the nine required warnings randomly and in approximately equal numbers. The comment recommended that, for “soft pack” style packages, FDA apply a policy of enforcement discretion that relieves companies of the obligation to display the nine required warnings randomly and equally as long as companies have taken reasonable steps to distribute the warnings as randomly and equally as possible. Another comment expressed general concerns about a manufacturer’s ability to comply with the requirement that the warnings be randomly displayed in as equal a number of times as possible.

Several comments requested additional guidance on the filing of warning plans, including how to hold parties responsible for meeting FCLAA and the Tobacco Control Act’s rotation and random display requirements.

In addition, one comment asked that FDA adopt a formal process for approval of required warnings on packages and warning plans. Some comments from manufacturers suggested that, to add predictability for companies on the transition to the new warnings, FDA should consider adopting a procedure to allow pre-approval or pre-submission review of cigarette packaging and advise manufacturers of any deficiencies so the manufacturer can remedy them before production. One comment requested that FDA use Federal Trade Commission (FTC) procedures for pre-approval review of packaging.

(Response) We have opted not to address these issues as part of this rulemaking proceeding. Under section 4(c) of FCLAA, warning plans must be submitted to FDA for approval. As noted in the NPRM, we intend to separately address the requirements of section 4(c) of FCLAA related to the submission of plans regarding the random display of warnings on packages and rotation in advertisements (75 FR 69524 at 69538). This is still our plan, and we believe the issues raised in these comments would be better addressed in that context.

(Comment 197) One comment suggested that FDA provide sample pre-approved layouts for required warnings on cigarette packages.

(Response) By providing the electronic files of the required warnings, we are providing formats that the companies must use for their packages. The final rule includes a document incorporated by reference, entitled “Cigarette Required Warnings,” which contains the final images to be required on cigarette packages. Cigarette manufacturers also should refer to § 1141.10(a), which mandates that the required warnings be on the top 50
percent of both the front and back of the cigarette packages.

(Comment 198) One comment requested that FDA issue a tobacco product advertising guide for industry. This comment noted that while product labeling and advertising present some similar issues, there are specific issues that relate solely to advertising communications with consumers. Another comment suggested that FDA should issue separate advertising guidance for industry that includes recommendations for display of required warnings in each common advertising form.

One comment stated that FDA should require that cigarettes displayed at the point of sale should be required to be displayed in a manner so that the graphic warnings are visible.

One comment submitted on behalf of several nonprofit organizations suggested that FDA modify proposed §1141.10 to include two paragraphs regarding the use of images of cigarette packs in advertisements and in other communications. They requested that FDA add one paragraph to state that any image of a cigarette pack in an advertisement must include a required warning on the cigarette pack image. In addition, they requested that FDA add a paragraph to state that no manufacturer, importer, distributor, or retailer may alter any image used to depict cigarette packs as legally distributed or sold to consumers in any public communication (including, but not limited to, movies, Web sites, and television programs) so that the required warning on the cigarette pack image is removed or obscured in any way.

(Response) We recognize that the range of advertising materials covered by the new graphic warning rules may create additional complexities. As stated previously, we intend to issue separate regulatory documents to provide information on compliance with the random display and rotation requirements. We will consider whether any other actions that are within the scope of our authority under the Tobacco Control Act may be warranted, such as addressing requests for additional guidance regarding advertising or suggested regulatory changes.

VII. Legal Authority and Responses to Comments

A. FDA’s Legal Authority

As set forth in the preamble to the proposed rule (75 FR 69524 at 69524 through 69525), the Tobacco Control Act provided FDA with the authority to regulate tobacco products, and section 201 of the Tobacco Control Act modifies section 4 of FCLAA to require that nine new health warning statements appear on cigarette packages and in cigarette advertisements and to require that “the Secretary [of Health and Human Services] shall issue regulations that require color graphics depicting the negative health consequences of smoking” to accompany the nine new health warning statements.

Under section 4(d) of FCLAA (as amended by section 201(a) of the Tobacco Control Act), FDA may adjust the type size, text, and format of the required warnings as FDA determines appropriate so that both the textual warning statements and the accompanying graphics are clear, conspicuous, and legible and appear within the specified area. Furthermore, section 202(b) of the Tobacco Control Act amends section 4 of FCLAA to permit FDA to, after notice and an opportunity for the public to comment, adjust the format, type size, color graphics, and text of any health warning statement if such a change would promote greater public understanding of the risks associated with the use of tobacco products.

In addition, provisions of the FD&C Act provide authority to require disclosures. For example, section 906(d) of the FD&C Act (21 U.S.C. 387(d)) authorizes FDA to issue regulations restricting the sale or distribution of cigarettes and other tobacco products, including restrictions on the advertising and promotion of such products, if FDA determines the restriction is appropriate for protecting the public health.

These requirements are supplemented by the FD&C Act’s misbranding provisions, which require that product advertising and labeling include proper warnings (see 21 U.S.C. 321(a); 387(c)(a)(1), (a)(7)(A), (a)(7)(B), and (a)(8)(B)). In addition, under section 701(a) of the FD&C Act (21 U.S.C. 371(a)), FDA has authority to issue regulations for the efficient enforcement of the FD&C Act.

While we did not receive comments regarding our authority to issue these regulations under the provisions referenced in the previous paragraphs, we did receive comments regarding the constitutionality of the warning requirements, which are summarized and responded to in sections VII.B and VII.C of this document.

B. First Amendment Commercial Speech Issues

FDA received several comments related to First Amendment commercial speech issues. These comments are summarized and responded to in the following paragraphs.

(Comment 199) Several comments from the tobacco industry, advertising industry associations, and private citizens expressed concern that the graphic warning requirements proposed by FDA violate the First Amendment of the United States Constitution. Specifically, comments alleged that the proposed required warnings are unconstitutional because, rather than conveying factual information to consumers, they contain “disturbing,” “lurid” images that are designed to elicit emotions, such as “loathing, disgust, and repulsion.” Thus, the comments state, they force tobacco companies to “stigmatize their own products” and compel them to convey the government’s “ideological message” that “the risks associated with smoking cigarettes outweigh the pleasure that smokers derive from them” and that no one should use these lawful products. The comments also asserted that the warning requirements are unjustified because the health risks of smoking are already well known, and that they are unduly burdensome because the size and positioning requirements for the warnings on packages and advertisements would effectively rule out the companies own attempts to convey information about their products. For these reasons, the comments asserted that the graphic warning requirements constitute compelled speech regulation that is content-based and presumptively invalid and that the requirements can only be upheld if they satisfy strict scrutiny, i.e., if they further a compelling government interest by the least restrictive means available. The comments stated that the graphic warning requirements cannot satisfy this standard because they will have no material impact on consumers’ beliefs about the health risks of smoking or on smoking behavior and because the government bypassed less speech-restrictive alternatives in favor of the requirements.

The comments from the tobacco industry also stated that the warning requirements violate the First Amendment because they restrict tobacco companies’ speech. They stated that requiring the warnings to occupy the top 50 percent of the front and back display panels of cigarette packages and the top 20 percent of cigarette advertisements impairs the communication value of the tobacco product manufacturers’ trademarks and trade dress and narrows their avenues of communications with adult smokers, which are already limited because of the
Master Settlement Agreement and the other requirements of the Tobacco Control Act. Indeed, one of the comments argued that delegating tobacco companies’ message to the bottom half of cigarette packages would render their speech on packaging “wholly ineffective” and that the collective requirements with respect to packaging and advertisements would “effectively rule out” the companies’ attempts to convey information about their products to consumers. The comments asserted that the warning requirements do not satisfy the test governing restrictions on commercial speech articulated by the Supreme Court in Central Hudson Gas & Electric Corp. v. Public Service Commission, 447 U.S. 557 (1980), which requires that government restrictions on commercial speech directly advance a substantial government interest and be no more extensive than necessary to serve that interest. Similar to their assertions with respect to compelled speech, the comments asserted that, to the extent that the warning requirements restrict speech, they do not pass muster under the First Amendment because they will have no material impact on consumers’ beliefs about, or understanding of, the health risks of smoking or on smoking behavior, and because the government bypassed less speech-restrictive alternatives in favor of the requirements.

Other comments, including comments from a law firm, a public health advocacy group, and a private citizen, disagreed that the warning requirements violate the First Amendment. Specifically, two comments noted that the warning requirements have been upheld by a Federal court in Commonwealth Brands v. United States, 678 F. Supp. 2d 512, 529–32 (W.D. Ky. 2010), appeal pending sub nom., Discount Tobacco City & Lottery, Inc. v. United States, Nos. 10–5234 & 10–5235 (6th Cir.). One comment noted that the court rejected an argument that the new warnings required under the Tobacco Control Act are too large and too prominent and stated that Congress has made findings with respect to the required size of the warnings, their placement on packages and advertisements, and the text of the warnings based on a substantial record. The comment also stated that Congress’ findings are supported by the voluminous authority cited in FDA’s NPRM. Another comment stated that, although tobacco companies will have to redesign their packages as a result of the warning requirements, they will still be able to communicate with their customers through packaging, advertising, and other channels. In addition, the comment stated that the warning requirements do not offend manufacturers’ First Amendment rights because the required warnings are factual disclosures that accurately depict the real consequences of smoking cigarettes and the benefits and importance of quitting. The comment asserted that the warning requirements support the public interest by providing consumers with truthful information that is helpful in making informed purchasing decisions. The comment also stated that the government constitutionally regulates the advertising and labeling for a wide variety of industries in the interest of providing consumers with accurate information about products that affect their health and that no product affects consumers’ health more than cigarettes. Finally, one comment stated that requiring warnings for cigarettes is well established legally and that the addition of graphic images to the warnings represents a difference in form that will not change the fundamental message content of the warnings. As a result, the comment concluded that there is no constitutional basis to delay the implementation of the warning requirements.

(Response) We have carefully considered these comments and we disagree that the warning requirements violate the First Amendment under either of the theories set forth in the comments. To the extent that the warning requirements compel commercial speech, they are permissible under Zauderer v. Office of Disciplinary Counsel of Supreme Court of Ohio, 471 U.S. 626 (1985), and to the extent that they restrict commercial speech, they satisfy the Central Hudson requirements.

The Warning Requirements Permissibly Compel Disclosure of Factual Information. The comments do not dispute that required warnings and other disclosure requirements “trench much more narrowly on an advertiser’s interests than do flat prohibitions on speech” and may appropriately be required “in order to dissipate the possibility of consumer confusion or deception” (Zauderer, 471 U.S. at 651 (citation omitted)). Accordingly, regulations that compel “purely factual and uncontroversial” commercial speech are subject to more lenient review than regulations that restrict accurate commercial speech and will be sustained if they are “reasonably related” to the government’s asserted interest (see also Milavetz, Gallop & Milavetz, P.A. v. United States, 130 S. Ct. 1324, 1339 (2010) (disclosure requirements are subject to “less exacting scrutiny” than affirmative limitations on speech)). “Commercial disclosure requirements are treated differently from restrictions on commercial speech because mandated disclosure of accurate, factual, commercial information does not offend the core First Amendment values of promoting efficient exchange of information or protecting individual liberty interests” (Nat’l Electric Manufacturers Ass’n v. Sorrell, 272 F.3d 104, 113–14 (2d Cir. 2001), cert. denied, 536 U.S. 905 (2002)). Instead, such disclosure advances “the First Amendment goal of the discovery of truth and contributes to the efficiency of the ‘marketplace of ideas’” (Id. at 114).

“Protection of the robust and free flow of accurate information is the principal First Amendment justification for protecting commercial speech” (Id.). The nine new health warning statements and the accompanying graphic images selected by FDA convey information that is factual and uncontroversial. Therefore, the warning requirements are subject to the “reasonable relationship” test in Zauderer, rather than strict scrutiny as suggested by some of the comments. The comments do not dispute that the warning statements are true. Indeed, as detailed in the NPRM and in section II.A.2 of this final rule, there is substantial scientific evidence to support the information conveyed in the new required warnings. The NPRM summarizes a large body of scientific evidence showing that cigarettes cause a wide range of negative health consequences, including various types of cancer; all the major cardiovascular diseases, including heart disease and stroke; COPD and other respiratory diseases; and a variety of negative health effects in infants born to women who smoke and in nonsmokers exposed to secondhand smoke (75 FR 69524 at 69527 through 69529). The NPRM also sets forth scientific evidence describing the negative effects of nicotine addiction and the major and immediate health benefits of smoking cessation (75 FR 69524 at 69528 through 69529). As the court in Commonwealth Brands correctly observed, the content of the warnings “is objective and has not been controversial for many decades” (Commonwealth Brands, 678 F. Supp. 2d at 531).

The images we have selected to accompany the nine warning statements also convey information that is factual and uncontroversial regarding the negative health consequences of smoking. These images are consistent with the information conveyed in the
accompanying textual warning statements; each image depicts themes and subjects that provide visual context for the textual warning statements. The images also play a crucial role in the communication of the textual warning information; as discussed extensively in the NPRM, the addition of graphic images to health warning messages causes consumers to notice and attend to the warning information in the first instance, and increases recall of the warning message and the depth of cognitive processing of the message (75 FR 69524 at 69531).

The comments did not dispute that the images proposed to accompany the warning statements accurately depict the negative health consequences of smoking. Rather, they faulted our proposed images for being “disturbing” or eliciting emotions. For example, one of the comments cited as disturbing several of the images selected by FDA in this rule, including the images entitled “hole in throat,” depicting a man smoking through a tracheostomy opening; “healthy/diseased lungs...” depicting healthy lungs juxtaposed with lungs damaged by smoking; “cancerous lesion on lip,” depicting a lesion consistent with that caused by oral cancer; and “man with chest staples,” depicting a man with an autopsy scar. The comment did not assert, however, that the effects shown in the images are false, i.e., that they are not manifestations of negative health consequences of smoking, such as throat, lung, and oral cancer, and death. The fact that the images are disturbing or evoke emotion does not mean that they are not factual representations of the effects of smoking. In fact, the severe, life-threatening and sometimes disfiguring health effects of smoking conveyed in the required warnings are disturbing and the images we have selected appropriately reflect this fact. As such, it is not surprising that the warnings regarding the negative health consequences of smoking would evoke emotions such as fear of being stricken with life-threatening cancer or disgust at what it has to have that happen. If the required warnings failed to elicit emotional reactions, they would also fail to communicate the described negative health consequences of smoking in a truthful, forthright manner.

Some comments also stated that “non-factual cartoon images” proposed by FDA removed any doubt that the proposed warnings convey an ideological message. For this final rule, one of the images we have selected is, indeed, a graphic illustration. That image shows a “baby in incubator” and accompanies the warning statement, “Smoking during pregnancy can harm your baby.” As set forth in the NPRM, there is ample evidence to show that smoking during pregnancy has negative effects, including increasing rates of preterm delivery and shortened gestation and increasing the likelihood of low birth weight infants, among other things (75 FR 69524 at 69528). Thus, the image “baby in incubator” accurately depicts the health consequences smoking during pregnancy can have for infants born to mothers who smoke. The style of the depiction—here, a graphic illustration—does not make it less factual. The style is just a means to convey the information.

The remaining images we have selected also factually depict the negative health consequences of smoking when viewed in context with their accompanying warning statements. As explained in section III of this document, the image “smoke approaching baby” accompanying the statement “WARNING: Tobacco smoke can harm your children” effectively conveys the factual message that exposure to tobacco smoke is harmful for children by realistically showing a baby being exposed to secondhand smoke. The image “oxygen mask on man’s face,” which accompanies the statement “WARNING: Cigarettes cause strokes and heart disease,” accurately depicts a typical intervention for a patient suffering acute cardiac distress or stroke. The image “woman crying,” which is paired with the statement “WARNING: Tobacco smoke causes fatal lung disease in nonsmokers,” is a realistic portrayal of the emotional suffering experienced as a result of disease caused by secondhand smoke exposure. Finally, the image “man I Quit t-shirt,” which is paired with the statement “WARNING: Quitting smoking now greatly reduces serious risks to your health,” realistically portrays an image of a man that is consistent with and supportive of this factual warning statement, although, unlike the other required warnings, this warning is presented in a less literal manner (i.e., it conveys factual information about the negative health consequences of smoking by educating consumers about the positive health consequences of refraining from smoking).

The comments also asserted that some of the proposed images, including some now selected by FDA in this final rule, appear to use technologically-enhanced photographs to emphasize the effects of sickness and disease. While we acknowledge that some of the photographs were technologically modified to depict the negative health consequences of smoking, the effects shown in the photographs are, in fact, accurate depictions of the effects of sickness and disease caused by smoking, and the comments did not dispute this fact.

As one of the comments noted, the addition of graphics to warnings for cigarettes is a difference in form only and does not change the fundamental content of the messages, which convey factual information about the health consequences of smoking. The court in Commonwealth Brands was correct when it stated that it “does not believe that the addition of a graphic image will alter the substance of such [warning] messages, at least as a general rule” (Commonwealth Brands, 678 F. Supp. 2d at 532). Rather, these images alter the effectiveness of the warnings by enhancing their ability to communicate factual information to consumers.

Despite the factual nature of the messages conveyed by the required warnings as described previously, some comments asserted that the government’s goal is to force cigarette companies to stigmatize their products by including the government’s ideological, antismoking message on their packages and advertisements. These comments claimed that the size of the warnings and the FDA study endpoints assessing consumers’ emotional and cognitive reactions to the required warnings and whether the warnings were “difficult to look at,” belie any suggestion that they are purely factual.

We disagree with these comments. The size of the warnings and their ability to evoke cognitive and emotional responses are consistent with the government’s interest in ensuring that the required warnings effectively communicate factual information about the negative health consequences of smoking to consumers. The NPRM (75 FR 69524 at 69531 through 69534) and section II.D of this final rule summarize the significant research literature supporting FDA’s conclusion that larger, graphic warnings more effectively communicate health risks to consumers than the existing smaller, text-only warnings on cigarette packages and in advertisements.

Likewise, our decision to use images that elicit strong cognitive and emotional responses is consistent with established models of risk communication. Our research study included three measures to assess the salience (i.e., noticeability and readability) of the proposed required warnings. Emotional reactions were measured in cognitive reactions, and whether the warning was difficult to look at. Use of
these measures is well-established in the scientific literature. As discussed in the study report (Ref. 49, study report) and in comments discussed in section III of this document, risk information is most readily conveyed by warnings that elicit strong responses on these measures—eliciting strong emotional and cognitive reactions to graphic warnings enhances recall and information processing, which helps to ensure that the warnings are better understood and remembered. These responses in turn influence short-term outcomes, such as later recall of the message and changes in knowledge, attitudes, and beliefs related to the dangers of tobacco use and exposure to secondhand smoke, and eventually lead to long-term changes in behavior. Thus, contrary to the comments discussed previously, our use of these reaction measurements does not demonstrate the Agency’s intent to stigmatize tobacco products. Rather, these measures are appropriate indicators of how effectively health warning messages are communicated, and were used in FDA’s research study to provide valuable information regarding the relative ability of the 36 proposed required warnings to effectively convey the very real adverse health consequences of smoking to the public.

Indeed, the court in Commonwealth Brands rejected an argument that the purpose of the new, larger warnings with their graphic image component is to “browbeat potential tobacco consumers” with the government’s antismoking message. The court stated that “the government’s goal is not to stigmatize the use of tobacco products on the industry’s dime; it is to ensure that the health risk message is actually seen by consumers in the first instance” (Commonwealth Brands, 678 F. Supp. 2d at 530 (emphasis in original)). We agree with these findings of the district court.

Because the warning requirements compel the disclosure of information that is purely factual and noncontroversial, they are permissible under Zauderer if they are reasonably related to the government’s asserted interest. As stated repeatedly in the NPRM and this rule (see, e.g., section I.D of this document), the Agency’s primary interest is to effectively convey the negative health consequences of smoking on cigarette packages and in advertisements, a necessary part of which, as the court in Commonwealth Brands recognized, is “to ensure that the health risk message is actually seen by consumers in the first instance.” The warning requirements are clearly reasonably related to this interest.

Both the research literature and FDA’s study of the proposed required warnings indicate that the required warnings are effective at communicating the health consequences of smoking to consumers. We have cited extensive literature in the NPRM and in section I.D of this final rule discussing the greater effectiveness of larger, graphic warnings over the current warnings at getting consumers’ attention (see 75 FR 69524 at 69531 through 69532). For example, in one study in which students were shown images of the Canadian graphic warnings and the current warnings in use in the United States, the Canadian graphic warnings significantly increased aided recall of the warnings, increased depth of message processing, and increased the perceived strength of the message (75 FR 69524 at 69531, citing Ref. 97). In addition, as discussed in section III of this document, FDA’s study report (Ref. 49) demonstrates that eight of the nine required warnings selected for the final rule showed highly significant effects relative to the text-only control on all the salience measures (emotional reaction scale, cognitive reaction scale, and difficult to look at measure) across all of the target audiences (youth, young adults, and adults). The ninth warning, which communicates the message that “Quitting smoking now greatly reduces serious risks to your health,” also showed strong effects relative to the text-only control, with significant effects in at least some audiences on the emotional and cognitive reaction scales. Again, these results with respect to the salience measures are important because they have been shown to enhance recall and information processing, which helps to ensure that warnings are better understood and remembered.

As set forth previously, to the extent that the warning requirements compel speech, they are permissible under Zauderer because they require disclosure of factual information and are reasonably related to FDA’s goal of effectively communicating the health consequences of smoking to consumers. Accordingly, it is necessary to address the strict scrutiny analyses set forth in the comments.

We are not persuaded to the contrary by the comments’ assertions that the warning requirements are unjustified and unduly burdensome. The industry comments discussed previously contended that the warnings are unjustified because the health risks of smoking are already universally known and overestimated and the FDA study results show that the required warnings will have no impact on smoking beliefs or behavior. To support their argument, they cite Ibanez v. Florida Department of Business and a major study into tobacco policy in the United States by the IOM in 2007 concluded that U.S. package warnings are both “unnoticed and stale” and found that they fail to communicate relevant information in an effective way (Ref. 3 at 281). The Chair of the IOM’s Committee on Reducing Tobacco Use described the warnings on cigarette packs as “invisible” in
testimony in 2007 on a precursor to what was enacted as the Tobacco Control Act (75 FR 69524 at 69530). Research regarding warning statements in cigarette advertisements has shown similar results (Id., and studies cited therein). As discussed in the NPRM, the IOM expressed concern about the ability of consumers with less education to recall the information included in text-based messages. The IOM further explained that smokers are more likely to recall larger warnings as well as warnings that appear on the front of packages than those on the side, as is the case for the current warnings (75 FR 69524 at 69531). As the court in Commonwealth Brands likewise concluded, the evidence before Congress clearly demonstrates that the new warning requirements are justified (678 F. Supp. 2d at 530–31).

Substantial evidence showing consumer ignorance regarding the health risks of smoking and the ineffectiveness of the current warnings at communicating such risks clearly supports the need for the required warnings. The results of our research study showing significant effects on salience measures for all of the required warnings, along with the substantial international evidence showing that larger, graphic warnings effectively communicate health risks, demonstrate that, unlike the disclosures in the cases cited in the comments, the required warnings will have more than a speculative effect on consumer confusion about the risks of smoking.\footnote{In Zauderer, the asserted government interest was preventing consumers from being misled by a legal advertisement, and thus, the Court noted that warnings or disclosures could be appropriately required “in order to dissipate the possibility of consumer confusion or deception” (Zauderer, 471 U.S. at 651 (citations omitted)). In articulating the applicability of the Lemon Amendment scrutiny for disclosure requirements, the Court stated that such requirements must be “reasonably related to the State’s interest in preventing deception of consumers” (Id.). However, appellate courts have held that Zauderer’s holding was not limited to disclosure requirements that addressed potentially deceptive advertising, but rather applied to disclosures aimed at better informing consumers about the products that they purchase (see Sorrell, 272 F.3d at 115 (applying the Zauderer standard and upholding a disclosure statute aimed at increasing consumer awareness of the presence of mercury in various products because the statute’s goal was consistent with the policies underlying First Amendment protection of commercial speech and the distinction between compelled and restricted commercial speech); see also New York State Restaurant Assoc. v. New York City Board of Health, 556 F.3d (2d Cir. 2009) (upholding under Zauderer a requirement that restaurants disclose calorie content on menus because it was reasonably related to the city’s goal of education). Care Mgmt. Ass’n v. Rowe, 429 F. 3d 294, 310 n. 8 (1st Cir. 2005) (stating that the court did not find any cases limiting Zauderer to “potentially deceptive advertising directed at consumers”).}

Equally unavailing is the assertion that the warning requirements are unduly burdensome because the required size and positioning of warnings on packages and in advertisements effectively rule out tobacco companies’ own attempts to convey information. Because this part of the compelled speech argument overlaps with the assertion that the warning requirements restrict speech in violation of the First Amendment, it is addressed in the following paragraphs.

The Warning Requirements Are Permissible Under Central Hudson. To the extent that the challenged provisions restrict commercial speech, the restrictions are analyzed under the framework established in Central Hudson Gas & Electric Corp. v. Public Service Commission, 447 U.S. 557 (1980). “The First Amendment’s concern for commercial speech is based on the informational function of advertising” (Id. at 563). Consequently, there is no protection for “commercial messages that do not accurately inform the public about lawful activity” or that are “related to illegal activity” (Id. at 563–64). If the communication is neither misleading nor related to unlawful activity, the government may impose restrictions that directly advance a substantial government interest and are not more extensive than is necessary to serve that interest (Id. at 566). That standard does not require the legislature to employ “the least restrictive means” of regulation or to achieve a perfect fit between means and ends (Board of Trustees v. Fox, 492 U.S. 469, 480 (1989)). It is sufficient that the legislature achieve a “reasonable” fit by adopting regulations “in proportion to the interest served” (Id., quoting In re R.M.J., 455 U.S. 191, 203 (1982); accord Puglisi v. Trucchin, 402 F.3d 786, 771 (6th Cir. 2007) (en banc)). The Supreme Court has emphasized that “[t]he Constitution gives to Congress the role of weighing conflicting evidence in the legislative process” (Turner Broadcasting System, Inc. v. FCC, 520 U.S. 180, 199 (1997)).

“Even in the realm of First Amendment questions where Congress must base its conclusions upon substantial evidence, deference must be accorded to its findings as to the harm to be avoided and to the remedial measures adopted for that end, lest [a court] infringe on traditional legislative authority to make predictive judgments when enacting nationwide regulatory policy” (Id. at 196). Thus, “the question is not whether Congress, as an objective matter, was correct” in its determinations (Id. at 211). “Rather, the question is whether the legislative conclusion was reasonable and supported by substantial evidence in the record before Congress” (Id.).

Comments from tobacco product manufacturers argued that the warning requirements restrict tobacco companies’ speech because the warnings must occupy the top 50 percent of the front and back display panels of cigarette packages and 20 percent of the area of cigarette advertisements. They stated that these size and positioning requirements are unduly burdensome and will significantly impair their ability to convey information about their products to adult consumers. In essence, their argument is that the new warnings are too large and too prominent, which, as recognized by some of the comments discussed previously, has already been rejected by the court in Commonwealth Brands (see Commonwealth Brands, 678 F. Supp. 2d at 531).

It is important to note that the comments did not identify any specific statements that will be restricted by the warning requirements. Nonetheless, we will assume for the purpose of argument that any speech that possibly could be restricted as a result of this rule would be nonmisleading and relate to lawful activity and, thus, would be commercial speech protected by the First Amendment.

The comments did not dispute that the government has a substantial interest in effectively communicating the health risks of smoking to the public or, as the court in Commonwealth Brands characterized it, in “ensur[ing] that the health risk message is actually seen by consumers in the first instance” (Id. at 530). This substantial interest satisfies the first step of the Central Hudson analysis.

With respect to the second step, we have repeatedly discussed in the NPRM and this final rule evidence demonstrating that the required warnings will directly advance that interest. Such evidence includes the FDA study results showing significant effects on salience measures for all of the nine required warnings (see section III of this document) and the international experience demonstrating the enhanced communication value of larger, graphic warnings (see 75 FR 69524 at 69531 through 69533). It also
includes studies showing the improved effectiveness of Canada’s larger, graphic warnings at communicating health risks. For example, national surveys conducted on behalf of Health Canada indicate that approximately 95 percent of youth smokers and 75 percent of adult smokers report that the Canadian pictorial warnings have been effective in providing them with important health information (see Ref. 3 at p. 294). In another study of adult smokers, more than half of the study participants reported that the pictorial warnings made them think about the health risks of smoking (Ref. 44). A study comparing Canadian and United States warnings found that while “83 percent of Canadian students mentioned health warnings in a recall test of cigarette packs,” only “7 percent of U.S. students” did the same (see Ref. 3 at C–3 to C–4).

The comments that argued that the warning requirements are unconstitutionally restrictive ignored this evidence. Instead, they suggested that, to satisfy this step, FDA’s research study would have to show a material impact on consumers’ beliefs about, or understanding of, the health risks of smoking or smoking behavior. We disagree. The evidence showing that the required warnings will directly advance the government’s primary goal of effectively communicating the negative health consequences of smoking by first ensuring that the warnings will be seen and processed by consumers is sufficient to satisfy the second step of Central Hudson. A showing with respect to other goals, such as impacts on consumer beliefs or smoking behavior, is not necessary for purpose of this analysis. However, we note that there is significant evidence that these goals will also be advanced by the warning requirements.

The comments repeatedly cited to FDA’s study report to support the proposition that the required warnings will have no effect on consumer beliefs or behavior. However, such an assertion fails to take into account the study design and the extensive evidence in the literature indicating that the required warnings will positively impact beliefs and behavior. As we note in section III of this document, it is not surprising that the proposed required warnings, as a whole, did not elicit strong responses on the beliefs and intentions measures because study participants had only a single exposure to one warning; the study was not designed to show long-term effects on behavior. However, the study cannot be ignored in isolation from the overall body of scientific evidence regarding the positive effects of larger, graphic health warnings on smoking beliefs and behavior, which we summarized in the NPRM (75 FR 69524 at 69531 through 69534).

Finally, the comments stated that the warning requirements do not satisfy the third step of the Central Hudson test because the mandated size and positioning of the warnings on packages and advertisements will effectively rule out tobacco companies’ ability to convey information about their products. They stated that the requirements are more extensive than necessary to achieve the government’s interests and suggested that less-speech restrictive alternatives, including alternatives to the warning size and positioning requirements included by Congress in the Tobacco Control Act, would be equally as effective.

The comments provided no basis for setting aside Congress’ judgment as to the appropriate specifications. As the court in Commonwealth Brands explained, Congress considered extensive evidence starting with the 1994 Surgeon General’s Report and ending with the 2007 IOM Report, which is discussed in the NPRM (75 FR 69524 at 69530), demonstrating that the existing warnings are “unnoticed” and “stale” and decided that the content and format of the warnings needed to be revised (Commonwealth Brands, 678 F. Supp. 2d at 530–31). In so doing, Congress chose specifications for the warnings that accord with FCTC, which calls for warnings that “shall be large, clear, visible and legible,” “should be 50% or more of the principal display areas but shall be no less than 30% of the principal display areas,” and “may be in the form of or include pictures or pictograms” (FCTC art. 11.1(b)). The FCTC has been signed by the United States and ratified by 167 countries. As the Commonwealth Brands court correctly found, “Congress also informed its warning requirements by looking at the use of a nearly identical warning requirement in Canada” (Commonwealth Brands, 678 F. Supp. 2d at 531). Like the required warnings, the Canadian warnings occupy the top half of the two main panels of cigarette packages.

Thus, Congress based its legislative decision to revise the warnings in the first instance and to mandate certain size and placement specifications for the warnings on substantial evidence in the record. At this time, we do not intend to change those specifications. Although comments from tobacco companies asserted that the larger size leaves inadequate room for their own commercial messages, they identified no information that is suppressed by virtue of the larger warnings, even though they have complied with similar requirements in other countries for years. The tobacco companies retain more than half of their cigarette packaging and 80 percent of their advertisements for their own commercial speech.

Moreover, extensive disclosure requirements are by no means unique to cigarettes. For example, for products such as pain relievers, certain allergy medications, and products to treat a variety of cold symptoms, the required warnings together with other FDA-required information typically encompass more than 50 percent of the product packaging.

For these reasons, “the warning requirement is sufficiently tailored to advance the government’s substantial interest under Central Hudson” (Id. at 532).

The reliance by two comments on the Seventh Circuit’s decision in Entertainment Software Association v. Blagojevich, 469 F.3d 641 (7th Cir. 2006), does not persuade us to the contrary. In that case, the court invalidated a State law requiring video-game retailers to place a four-square-inch label with the numerals “18” on any “sexually explicit” video game. Unlike here, the court concluded that the sticker “communicates a subjective and highly controversial message—that the game’s content is sexually explicit,” a term capable of multiple definitions, and expressly rejected the comparison to the “surgeon general’s warning of the carcinogenic properties of cigarettes, the analogy the State attempts to draw” (Id. at 652). “Applying strict scrutiny,” the court noted that “[t]he State has failed to even explain why a smaller sticker would not suffice” (Id.). Here, by contrast, Congress has required accurate and objective warnings in a format that accords with the provisions of the FCTC, to which the United States is a signatory, and whose effectiveness has been demonstrated by international experience, after concluding existing, yet smaller, warnings were ineffective at conveying important health information.

We also disagree with the assertion in the comments that the warning requirements fail to meet the third step of Central Hudson because the government failed to consider numerous less speech-restrictive alternatives. One of the comments suggested that the government disseminate information

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8 See 21 CFR 201.66; see also http://www.accessdata.fda.gov/druginfodocs/docs/label/2009/022032s003lbl.pdf (example of packaging for OTC heartburn medication).
about health risks as one alternative for communicating health risks to consumers. However, government dissemination of the message already occurs—for example, HHS currently has several hundred tobacco-related Web sites, which provide informative messages regarding, for example, the harmful effects of tobacco use (Ref. 89), and CDC’s Office on Smoking and Health funds health departments in all 50 states, the District of Columbia, and seven U.S. territories for comprehensive tobacco prevention and control and provides access to tobacco control advertising material for use in this comprehensive regulation of the tobacco industry. That decision seems eminently reasonable, too, since every other tool in the government’s arsenal is made less effective and more costly by Plaintiffs’ use of advertising “to stimulate underage demand.” (Government’s Response, p. 40). Accordingly, the Court rejects Plaintiffs’ contention that the existence of “numerous obvious non-speech-restrictive alternatives” renders the Act’s speech restrictions unconstitutional for lack of tailoring. (670 F. Supp. 2d at 530). For all of the reasons set forth in the previous paragraphs, we conclude that the warning requirements do not violate the First Amendment.

(Comment 200) One tobacco industry comment also claimed that requiring a reference to a cessation resource in the required warnings would violate the First Amendment because it is compelled speech that does not convey factual information about the product that is being sold. This comment claimed that requiring a cessation resource communicates a subjective policy message that consumers should not buy or use the product. (Response) We disagree. As explained previously, the requirement in this rule for graphic warnings on cigarette packages and advertisements is consistent with the First Amendment. Contrary to the comment, the reference to a cessation resource, when considered in context with the rest of the required warnings, conveys factual information to consumers and is permissible under the Zauderer standard for compelled disclosures because it is reasonably related to our interest in increasing the likelihood that existing smokers will become aware of the cessation resource and, consequently, increasing the likelihood that those who want to quit will be successful. It is also reasonably related to our interest in effectively communicating the health risks of smoking to consumers.

As discussed in detail in section V.B.6 of this document, the rule requires each required warning to include a reference to the existing National Network of Tobacco Cessation Quitlines (Network), which uses the telephone portal 1–800–QUIT–NOW. This rule will require that the cessation resource be displayed on the required warning images: “1–800–QUIT–NOW.” The NPRM cited evidence that more than 70 percent of smokers in the United States report that they want to quit, and approximately 44 percent report that they try to quit each year (75 FR 69524 at 69529; Ref. 66 at p. 13). However, as a result of nicotine addiction and, consequently, the resets of these smokers achieve success (75 FR 69524 at 69528 through 69529).

Instead of advocating a subjective policy message as suggested by the comment, including a cessation resource on required warnings will provide factual information for the many smokers who have already developed a desire to quit, either prior to or after viewing the health risk information in the required warnings. The reference is designed to inform such smokers and others that a resource exists that can help smokers quit and to inform them how they can access that resource. The factual nature of this information is underscored by our explanation in the NPRM that the Agency’s goal is “to provide a place where smokers and other members of the public can obtain smoking cessation information from staff trained specifically to help smokers quit by delivering unbiased and evidence-based information, advice, and support” (75 FR 69524 at 69540 (emphasis added)). In addition, our adoption of detailed criteria designed to ensure that the resource’s information, advice, and support are unbiased and evidence-based further emphasizes that the required reference to a cessation resource is factual in nature.

We disagree that a reference to a cessation resource does not convey information about the product being sold. The reference must be considered in context with the rest of the required warnings, which consist of textual statements and accompanying graphic images conveying to consumers factual information regarding the negative health consequences of smoking and the benefits of quitting. The reference to a smoking cessation resource naturally complements this information; instead of leaving consumers who are motivated to quit by the health risk information unassisted, it provides them with a concrete step to take action on this information.

Because the reference to a smoking cessation resource conveys factual information, it is permissible under Zauderer if it is reasonably related to the government’s asserted interest. Here, the reference is reasonably related to FDA’s interest in increasing the likelihood that existing smokers will become aware of the cessation resource and, consequently, increasing the likelihood that they will successfully quit smoking. As set forth in the discussion of the comments in section V.B.6 of this document, foreign countries that have included cessation resources on cigarette package warnings have generally experienced large increases in volume of calls to quitlines following their appearance on cigarette packages. In addition, as also discussed
in section V.B.6 of this document, the effectiveness of telephone quitlines is well documented; there is evidence that significant numbers of smokers are unaware of such assistance, even after extensive media campaigns; and there is evidence that knowing about the availability of a quitline increases quit attempts and successful cessation even among smokers who do not call the quitline.

Moreover, requiring a smoking cessation resource is also reasonably related to FDA’s interest in effectively communicating the health risks of smoking to consumers. As noted in the NPRM (75 FR 69524 at 69541) and in section V.B.6 of this final rule, there is evidence to show that including a reference to a smoking cessation resource in graphic warnings can enhance the effectiveness of graphic warnings at conveying health risk information to the public. We have determined that it is also important to inform smokers about a specific tool they can use to help them to quit smoking at the time they are looking at the warnings and thinking about the health consequences of smoking and the positive health benefits of quitting. Risk communication research indicates that messages that arouse fear about the health risks of smoking should be combined with information on concrete steps that can be taken to reduce those risks (Ref. 81 [Messages that arouse fear “appear to be effective when they depict a significant and relevant threat * * * and when they outline effective responses that appear easy to accomplish * * *”]). As one comment stated, providing information about how to reduce a risk that arouses fear helps to prevent consumers from suppressing thoughts about such risks, and thereby, failing to process the risk information. For this reason, too, we do not agree that the requirement to refer to a smoking cessation resource on cigarette packages and advertisements violates the First Amendment.

C. Takings Under the Fifth Amendment

We received a comment related to the Takings Clause of the Fifth Amendment. That comment is summarized and responded to in the following paragraphs.

(Comment 201) One comment submitted by several tobacco companies argued that the new health warning requirements unconstitutionally deprive them of their property rights in violation of the Takings Clause of the Fifth Amendment. The tobacco companies asserted that the new required warnings constitute a per se physical taking of their packaging and advertising space, as well as a regulatory taking of their property interests in their trademarks. (Response) We disagree that the rule effects a taking under either theory. The Takings Clause provides that “private property [shall not] be taken for public use, without just compensation.” A takings analysis begins with a threshold determination of what interest a person has in the thing that is allegedly taken (see Ruckelshaus v. Monsanto Co., 467 U.S. 986, 1001 (1984)). In order to assert a taking, a person must first identify a specific, concrete property interest that has been invaded or destroyed by the government (Penn Central Transp. Co. v. New York City, 438 U.S. 104, 124-25 (1978)). Once a concrete property interest is identified, it is necessary to determine whether the government’s action constitutes a taking of that interest.

The graphic warning requirements do not effect a per se taking. To conclude that a categorical, or per se, taking has occurred when the government directly appropriates or physically invades property is another way of saying that the government action so onerously burdens an important property right that the inquiry ends there. As the Supreme Court has explained: “A permanent physical invasion, however minimal the economic cost it entails, eviscerates the owner’s right to exclude others from entering and using her property—perhaps the most fundamental of all property interests” (Lingle v. Chevron U.S.A. Inc., 544 U.S. 528, 539 (2005); see also Loretto v. Teleprompter Manhattan CATV Corp., 458 U.S. 419, 433 (1982) [citation omitted] (“[T]he landowner’s right to exclude is ‘one of the most essential sticks in the bundle of rights that are commonly characterized as property.’”)). Viewed in this light, a requirement that tobacco companies display graphic health warnings as part of the package label on their products cannot be equivalent to the “physical invasion” of real property in the cases that the comments cites to support its per se takings argument (see Loretto, 458 U.S. at 441 (“Our holding today is very narrow.”)). The warnings involve personal property of a type that is already subject to extensive government regulation. Indeed, given the ubiquitous nature of government-mandated warnings on all kinds of consumer products, manufacturers of inherently dangerous products such as cigarettes cannot be said to have a categorical right to exclude health warnings from their products’ labels.9 Therefore, the tobacco companies have failed to identify the sort of property right the destruction of which would result in a per se taking.

Furthermore, as the Supreme Court has explained, the Takings Clause exists “to bar Government from forcing some people alone to bear public burdens which, in all fairness and justice, should be borne by the public as a whole” (Armstrong v. United States, 364 U.S. 40, 49 (1960); see Monongahela Nav. Co. v. United States, 148 U.S. 312, 325 (1893)). The tobacco companies’ argument amounts to an assertion that they must be compensated because they have been required to allow health warnings on their property. The point of the warnings is to protect the public health by informing consumers about the many harmful effects of the companies’ products, which kill an estimated 443,000 Americans every year. Therefore, the proposition that the public must pay for the cost of the warnings on tobacco products is simply not compatible with how “the burden of common citizenship” is proportioned in our system of modern government (see Keystone Bituminous Coal Ass’n v. DeBenedictis, 480 U.S. 470, 488–91 (1987); Pennsylvania Coal Co. v. Mahon, 260 U.S. 393, 413 (1922) (“Government hardly could go on if to some extent values incident to property could not be diminished without paying for every such change in the general law.”)).

In addition, the graphic warning requirements do not effect a regulatory taking. The tobacco companies also argue that the warnings constitute a regulatory taking because they have a reasonable expectation that their property rights will be protected based on statutory and common law protections provided to trademarks and trade dress. The tobacco companies do not identify the specific statutory or common law protections that led to their expectation that their property would be protected. Also lacking is an explanation of how the rule would interfere with such expectations. In any event, we do not agree that the rule effects a regulatory taking of the tobacco companies’ property.

The Supreme Court has declined to prescribe a “set formula” for identifying takings and instead has characterized a takings analysis as an “essentially ad hoc, factual” inquiry (Penn Central, 438 U.S. at 124). Nonetheless, the Court has identified three factors for consideration in assessing whether a regulatory taking has occurred: (1) The character of the

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9For example, for products such as pain relievers, certain allergy medications, and products to treat a variety of cold symptoms, the required warnings together with other FDA-required information typically encompass more than 50 percent of the product packaging (see 21 CFR 201.66).
V. Rose Acre Farms, Inc.

The second factor of the analysis also supports the conclusion that no taking will occur as a result of the rule. The vague suggestion that the rule interferes with tobacco companies' "reasonable investment-backed expectations" is similarly unpersuasive. To be reasonable, expectations must take into account the power of the State to regulate in the public interest (Ruckelshaus v. Shriversby Township, 808 F.2d 1023, 1033 (3d Cir.), cert. denied, 482 U.S. 906 (1987)). The nature of the property, and whether it has historically been, or potentially could be, subject to regulation also aids in determining whether any expectation in remaining free from regulation is reasonable. "[I]n the case of personal property, by reason of the State's traditionally high degree of control over commercial dealings, [the property owner] ought to be aware of the possibility that new regulation might even render his property economically worthless." (Lucas v. South Carolina Coastal Council, 505 U.S. 1003, 1027–28 (1992)). This is particularly true with respect to cigarettes, which are lethal and addictive—features the industry masked decades while stimulating underage demand (see United States v. Philip Morris USA, Inc., 566 F.3d 1095, 1124 (D.C Cir. 2009); United States v. Philip Morris USA, Inc., 449 F. Supp. 2d 1, 580 (Finding 2717) (D.D.C. 2006); Ref. 54 at p. 211). Commerce in tobacco products has been regulated for decades, subject to increasingly more restrictive Federal, State, and local measures over time. Indeed, Congress has mandated warnings on cigarette packs since 1965 (see Federal Cigarette Labeling and Advertising Act of 1965 (FCLAA), Pub. L. 89–92, 79 Stat. 282). Congress later amended FCLAA to update the text of the cigarette warnings and mandate them in cigarette advertisements as well (see Comprehensive Smoking Education Act of 1984, Pub. L. 98–474, 98 Stat. 2200). In light of this long history of regulation, companies that package and advertise cigarettes lack a reasonable investment-backed expectation that they will be able to continue to use their property without modification of the regulatory requirements that protect the public health. Any expectation that the industry would escape comprehensive regulation, such as the Tobacco Control Act, was eminently unreasonable. For these reasons, the third factor of the takings analysis, like the first two factors, compels the conclusion that the rule does not amount to a regulatory taking of property that requires compensation under the Fifth Amendment.

VIII. Implementation Date

In the preamble to the proposed rule, FDA stated that the final rule would become effective 15 months after the date the final rule publishes in the Federal Register. This time period is consistent with section 201(b) of the Tobacco Control Act, which specifies that the requirements for health warnings on cigarette packages and in advertisements are effective 15 months after the issuance of the regulations that FDA issues in this rulemaking.

In particular, we proposed that as of the effective date, no manufacturer, importer, distributor, or retailer of cigarettes may advertise or cause to be advertised within the United States any cigarette product unless the advertising complies with the final rule. With respect to cigarette packages, we explained that cigarettes must not be manufactured after the effective date unless their packages comply with the regulation. If any packaged cigarette product was manufactured prior to the effective date and does not comply with the regulation, a manufacturer may continue to introduce that package into commerce in the United States for an additional 30 days after the effective date of the final rule. After 30 days following the effective date, a manufacturer may not introduce into domestic commerce any cigarette the package of which does not meet the requirements of the final rule (75 FR 69524 at 69541). We noted that this limitation applied only to manufacturers and requested comments regarding mechanisms for enforcing this rule and its effective date, including ways to differentiate cigarette packages sold from inventory manufactured prior to the effective date rather than from inventory manufactured after the effective date.

We received several comments about the effective date, particularly requesting clarification regarding its application to manufacturers, distributors, and retailers after the 30-day period in which manufacturers may continue to sell noncompliant packages. Based on the comments and our review of the language in section 201(b) of the Tobacco Control Act, we find:

- The effective date should be 15 months after the date of publication in the Federal Register of this final rule;
- No manufacturer, importer, distributor, or retailer may advertise any cigarette product after the effective date if the advertisement does not comply with this rule;
- After the effective date, no person may manufacture for sale or distribution within the United States any cigarette...
the package of which does not comply with this rule;
• Beginning 30 days after the effective date of this rule, a manufacturer may not introduce into domestic commerce any cigarette, irrespective of the date of manufacture, if its package does not comply with the requirements of this rule;
• After the effective date, an importer, distributor, or retailer may not sell, offer to sell, distribute, or import for sale or distribution within the United States any cigarette the package of which does not comply with this regulation, unless the cigarette was manufactured prior to the effective date; and
• After the effective date, however, a retailer may sell cigarettes the packages of which do not have a required warning if the retailer demonstrates it falls outside the scope of this rule as described in §1141.1(c).

In the following paragraphs, we describe the individual comments concerning the effective date and respond to these comments.

(Comment 202) Several comments expressed the view that 15 months is an excessive amount of time to allow the tobacco industry before it must comply with the new requirements of this rulemaking. For example, some comments contended that tobacco companies have employed marketing and advertising experts and are continuously changing cigarette packaging and advertisements. These comments also noted that the tobacco industry has known that they will need to update packaging and advertising to comply with this regulation since the passage of the Tobacco Control Act. Some comments estimated the number of Americans that will become new smokers or die due to smoking during the 15 months prior to the effective date. Other comments recognized that the statute specifies a 15-month effective date, but requested that FDA make clear that cigarette packages manufactured after the effective date must comply with the requirements of the regulation.

(Response) The Tobacco Control Act specifies a 15-month implementation period for cigarette manufacturers to include required warnings on their packages and for all cigarette advertisements to comply with this rule. We agree this is an appropriate amount of time for implementation of the rule.

(Comment 203) One tobacco product manufacturer indicated in its comment that all manufacturers should be required to implement the same warning requirements within the same time period, and that there should not be a separate implementation period for small manufacturers.

(Response) As in the proposed rule, the implementation date in the final rule is the same for all manufacturers, regardless of size.

(Comment 204) One comment requested that FDA delay implementation of the rule until Constitutional issues raised in the comment are resolved either administratively or through litigation.

(Response) We disagree that the effective date of this rule should be delayed beyond the 15 months proposed in the NPRM. As explained in section VII of this document, we disagree that there are any Constitutional deficiencies associated with this rule and, therefore, there is no need to revise the rule or issue a new proposed rule to address these alleged deficiencies. Furthermore, section 201(b) of the Tobacco Control Act specifies that the requirements for health warnings on cigarette packages and in advertisements for cigarettes are effective 15 months after the issuance of this final rule.

(Comment 205) Several comments addressed the 30-day period for manufacturers to sell noncompliant packages that were manufactured prior to the effective date. One comment asserted that it is unnecessary to permit this 30-day sell-off period if there is adequate time for manufacturers to make necessary changes to cigarette packages prior to the effective date. The comment cited the United Kingdom as an example of a jurisdiction where tobacco product manufacturers had adequate lead time (1 year to implement changes to cigarette packages and 2 years to introduce picture warnings on other tobacco products) to meet implementation deadlines so that only compliant packages were sold after the compliance deadline. Other comments recognized that the statute grants manufacturers 30 days to sell noncompliant cigarette packages; however, these comments emphasized that FDA does not have the discretion to lengthen the 30-day period. Comments also stressed that any additional delay of implementation would needlessly delay the important public health benefits of the rule.

(Response) As explained previously, section 201(b) of the Tobacco Control Act specifies that manufacturers have an additional 30 days to sell cigarette packages that do not meet the requirements of the regulation if those packages were manufactured prior to the effective date.

(Comment 206) A small tobacco product manufacturer requested that FDA discontinue the use of the term “introduce into domestic commerce.” The comment asked whether the term means out of the manufacturer’s possession. The comment raised this question in the context of expressing concern that distributors and retailers might want to return product to a manufacturer if there is doubt about a distributor or retailer being permitted to sell cigarette packages that do not have a required warning, but were introduced into domestic commerce by the manufacturer during the 30-day sell-through period for manufacturers.

(Response) We agree with this comment that when a cigarette package has been sold by the manufacturer and is in the possession of a distributor or retailer, the product would be considered introduced into domestic commerce. However, we do not agree that a definition of “introduce into domestic commerce” is needed at this time. The comment recognized that there was similar language in the context of a statutory prohibition on the use of “light,” “low,” and “mild” descriptors and related FDA guidance for industry, however, that guidance did not define the phrase “introduce into domestic commerce.” We are not aware of confusion regarding this phrase in the context of “light,” “low,” and “mild” descriptors and decline to define that phrase here.

(Comment 207) Public health advocacy groups expressed concern that manufacturers will seek to sell a disproportionate number of noncompliant cigarette packages immediately prior to the expiration of the 30-day sell-off period and, therefore, FDA should take steps to ensure that all these sales are fully documented. The comment recommended that FDA impose certain requirements for selling noncompliant cigarette packages, such as a requirement to mark these packages with a statement that the product was manufactured prior to September 22, 2012, or with a readily identifiable symbol. This comment also recommended that each manufacturer be required to certify that all cigarettes so marked were manufactured before that date and submit an accounting of the number of packages on hand as of the effective date, the number of cigarette packages introduced into commerce during the 30-day period, and the number of packages on hand as of the expiration of the 30-day period. This comment also suggested that FDA not permit manufacturers to introduce into commerce in any calendar month a number of noncomplying cigarette packages that exceeds 10 percent of the average total number of cigarette packages introduced per month during the preceding year.
We disagree that such specific requirements are necessary to address a one-time sell-off period of 30 days. We recognize that some manufacturers may try to increase their sales of cigarette packages prior to the effective date and prior to the expiration of the sell-off period. However, there will be some limit to the demand for these cigarette packages. Manufacturers may increase manufacturing prior to the effective date at their own risk. After the 30-day sell-off period, a manufacturer may not sell noncompliant cigarette packages and would need to repack or destroy any noncompliant cigarettes manufactured prior to the effective date and sold after the effective date.

This comment did not provide a statutory interpretation that would justify this approach. Section 201(b) of the Tobacco Control Act states the effective date “shall be with respect to the date of manufacture” and that 30 days after the effective date, a manufacturer is precluded from introducing into domestic commerce any product that is not in conformance with section 4 of FCLAA. No similar statutory provision applies to importers or distributors.

Public health advocacy groups requested that FDA clarify that manufacturers are not prohibited from introducing into commerce cigarette packages that comply with the regulation prior to the effective date.

We agree that manufacturers are not precluded from introducing into commerce cigarette packages that contain required warnings in accordance with the regulation prior to the effective date. We also note that a cigarette manufacturer, importer, or retailer may include a required warning in an advertisement prior to the effective date. However, because the health warning requirements in FCLAA do not change until the effective date of this rule, any manufacturer, importer, or retailer that, prior to the effective date, includes a new required warning on a cigarette advertisement or advertisement must also comply with the warning requirements under the current version of FCLAA and any warning plan approved by the FTC.

Many comments requested clarification regarding whether there is any limitation on the period during which distributors and retailers may sell cigarettes that were manufactured prior to the effective date that are not compliant with the rule. Several comments submitted by organizations representing manufacturers and retailers asked that FDA clarify that distributors and retailers have an unlimited period to sell cigarette packages that do not comply with the regulation as long as the cigarettes were manufactured prior to the effective date. Several comments noted that this approach would be consistent with FDA’s treatment of cigarettes with the descriptors “light,” “low,” and “mild.” One manufacturer commented that any restraint on the ability of distributors or retailers to sell through their lawfully acquired product would unfairly deprive them of the benefit of their investment. Small tobacco product manufacturers noted that small manufacturers cannot afford to have distributors and retailers returning product based on a potential labeling concern. Retailer comments contended that limiting a sell-off period may cause a severe financial burden on small retailers because manufacturers generally do not allow cigarettes to be returned. Retailers also claimed that cigarettes do not have an indefinite shelf life and both distributors and retailers generally turn over their cigarette inventory in a timely manner. One comment suggested that retailers should be allowed to sell noncompliant cigarette packages at least through their “sell by” date, as indicated on the cigarette package by the manufacturer. On the other hand, one comment claimed it is essential that there be a fixed implementation deadline at the retail level or old stock can be expected to remain on retail store shelves for 6 months and more after the effective date.

As explained in the NPRM, section 201(b) of the Tobacco Control Act describes no limitation on the period during which distributors and retailers may sell cigarette packages that were manufactured prior to the effective date of this rule. In addition, there is no requirement that manufacturers include a “sell by” date on all cigarette packages. We note, however, that distributors, importers, and retailers are responsible for complying with this rule. After the rule’s effective date, they may not sell, offer to sell, distribute, or import for sale or distribution within the United States any cigarette the package of which does not comply with this regulation, unless the cigarette was manufactured prior to the effective date. After the effective date, however, retailers may sell cigarettes the packages of which do not have a required warning if they demonstrate they meet the provisions of § 1141.1(c) and are exempt from the requirements of 21 CFR part 1141 that apply to the display of health warnings on cigarette packages.

FDA has analyzed this final rule in accordance with the principles set forth in Executive Order 13132. Section 4(a) of the Executive order requires agencies to “construe * * * a Federal statute to preempt State law only where the statute contains an express preemption provision or there is some other clear evidence that the Congress intended preemption of State law, or where the exercise of State authority conflicts with the exercise of Federal authority under the Federal statute.” This rule is being issued under section 4 of FCLAA, as amended by the Tobacco Control Act, and sections 701(a), 903, and 906 of the FD&C Act (21 U.S.C. 371(a), 387c, and 387f), as amended by the Tobacco Control Act. Federal law includes an express preemption provision that preempts any requirement, except under the Tobacco Control Act, for a “statement relating to smoking and health, other than the statement required by section 4 of [FCLAA], * * * on any cigarette package.” (section 5(a) of FCLAA (15 U.S.C. 1334(a))). It also includes an express preemption provision that preempts any “requirement or prohibition based on smoking and health * * * imposed under State law with respect to the advertising or promotion of any cigarettes the packages of which are labeled in conformity with the provisions of [FCLAA],” which includes section 4 of FCLAA (section 5(b) of FCLAA). However, section 5(b) of FCLAA does not preempt any State or local statutes and regulations “based on smoking and health, that take effect after [June 22, 2009], imposing specific bans or restrictions on the time, place, and manner, but not content, of the advertising or promotion of any cigarettes” (section 5(c) of FCLAA).

In addition, section 916(a)(2) of the FD&C Act (21 U.S.C. 387p) expressly preempts any State or local requirement “which is different from, or in addition to, any requirement under [Chapter IX of the FD&C Act] relating to,” among other things, advertising and labeling. This express preemption provision, however, “does not apply
requirements relating to” among other things “the sale, distribution, * * * access to, [or] the advertising and promotion of * * * tobacco products.”

X. Environmental Impact

FDA has determined under §25.30(k) (21 CFR 25.30(k)) that this action is of a type that does not individually or cumulatively have an impact on the human environment. Therefore, neither an environmental assessment (EA) nor an environmental impact statement (EIS) is required. We received one comment on this issue, which we have summarized and responded to in the following paragraphs.

(Comment 211) One comment expressed concern regarding FDA’s statement in the proposed rule that this action does not individually or cumulatively have an impact on the human environment. The comment stated that there is an impact on the environment due to the fact that a reduction in the number of cigarettes consumed will result in a reduction of cigarette-related waste. The comment explained that cigarette butts pose a greater health hazard than most other litter, because they contain toxins that can be leached into water systems. The comment requested that this be included in FDA’s analysis to understand the large positive impact the required warnings have on the human environment.

(Response) We have considered this comment, but have concluded that neither an EA nor an EIS is required under §25.30(k). We have determined that a categorical exclusion applies in this instance, because (1) the action meets the criteria of the exclusion, i.e., there are no increases in existing levels of use or changes in intended use, and (2) there are no extraordinary circumstances.

According to the National Environmental Policy Act of 1969 (NEPA) and the Agency’s corresponding regulations, FDA must prepare an EIS for major Federal actions “significantly affecting the quality of the human environment” (see 40 CFR 1501.4; 21 CFR 25.22). If the action “may” have such a significant environmental effect, an agency must prepare an EA to provide sufficient evidence and analysis for the agency to determine whether to prepare an EIS or a finding of no significant impact (FONSI) (see 40 CFR 1501.3; 21 CFR 25.20). Agencies can establish categorical exclusions for categories of actions that do not individually or cumulatively have a significant effect on the human environment and for which, therefore, neither an EA nor an EIS is required (see 40 CFR 1508.4). However, FDA will require at least an EA for any specific action that ordinarily would be excluded if extraordinary circumstances indicate that “the specific proposed action may significantly affect the quality of the human environment” (see 21 CFR 25.21; 40 CFR 1508.4).

A regulation to modify labeling regulations constitutes a major Federal action under NEPA (see 40 CFR 1508.18), and typically requires at least an EA under 21 CFR 25.20(f). However, regulations establishing labeling requirements for marketed articles are categorically excluded, if the action will not result in (1) increases in the existing levels of use of the article or (2) changes in the intended use of the article (§25.30(k)). Therefore, FDA would not be required to file an EA if it meets these requirements.

We have determined that this regulation meets the requirements for a categorical exclusion. First, this regulation is clearly not expected to increase cigarette usage. In fact, this regulation is expected to cause a reduction in overall smoking rates and initiation, and we estimate that this rule will reduce the number of smokers by 213,000 in 2015, with smaller additional reductions through 2031. Second, the rule will not affect the way in which cigarettes are used among smokers and it does not change the intended use of cigarettes.

In addition, we have determined that there is no potential for serious harm to the environment resulting from the final rule that would otherwise constitute an extraordinary circumstance (see 21 CFR 25.21). Our action to regulate cigarette labeling does not lead to an increase in the level of use of these articles or a change in the intended use of these articles or their substitutes. The primary effect of this regulation will be to reduce smoking initiation and increase cessation efforts. Accordingly, there is no extraordinary circumstance that requires the filing of an EA.

XI. Analysis of Impacts

A. Introduction and Summary

FDA has examined the impacts of the final rule under Executive Order 12866, Executive Order 13563, the Regulatory Flexibility Act (5 U.S.C. 601–612) and the Unfunded Mandates Reform Act of 1995 (Pub. L. 104–4). Executive Orders 12866 and 13563 direct agencies to assess all costs and benefits of available regulatory alternatives and, when regulation is necessary, to select regulatory approaches that maximize net benefits (including potential economic, environmental, public health and safety, and other advantages; distributive impacts; and equity). This rule is an economically significant regulatory action under Executive Order 12866.

The Regulatory Flexibility Act requires agencies to analyze regulatory options that would minimize any significant impact of a rule on small entities. This rule will have a significant economic impact on a substantial number of small entities.

Section 202(a)(1) of the Unfunded Mandates Reform Act of 1995 requires that agencies prepare a written statement, which includes an assessment of anticipated costs and benefits, before proposing “any rule that includes any Federal mandate that may result in the expenditure by State, local, and Tribal governments, in the aggregate, or by the private sector, of $100,000,000 or more (adjusted annually for inflation) in any one year.” The current threshold after adjustment for inflation is $136 million, using the most current (2010) implicit Gross Domestic Product Deflator for the Gross Domestic Product. This rule will result in a 1-year expenditure that meets or exceeds this amount.

Conducting an impact analysis under Executive Order 12866, Executive Order 13563, the Regulatory Flexibility Act, and the Unfunded Mandates Reform Act involves assembling any available information that is relevant to the assessment of a regulation’s benefits and costs. It is not uncommon in scientific pursuits for there to be a lack of definitive information on some aspects of the question under investigation, and the impact analysis of this final rule is no exception. In light of this situation, we identify and present a range of possible benefits and costs.

The benefits, costs, and distributional effects of the final rule are summarized in table 1a of this document. As the table shows, the midpoints of the estimates for benefits annualized over 20 years is approximately $630.5 million at a 3-percent discount rate and $2.3 billion at a 7-percent discount rate. The midpoint for costs annualized over 20 years is approximately $29.1 million at a 3-percent discount and $37 million at a 7-percent discount rate.

The total benefits and costs of the final rule can also be expressed as present values. The midpoints of the estimates for the present value of benefits over 20 years is approximately $9.4 billion at a 3-percent discount rate and $2.3 billion at a 7-percent discount rate. The midpoint of the estimates for the present value of costs over 20 years is approximately $434 million at a 3-percent discount rate and $392 million.
at a 7-percent discount rate. With both discount rates, our midpoint estimates indicate that the benefits of the rule greatly exceed the costs. Executive Order 13563, section 1(b), requires that, to the extent permitted by law, agencies proceed with a regulation “only upon a reasoned determination that its benefits justify its costs.” The regulation is consistent with this requirement.

### Table 1a: Summary of Benefits, Costs and Distributional Effects

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<th>Economic Data: Costs and Benefits Statement</th>
<th>Notes</th>
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<td>$0</td>
</tr>
<tr>
<td>High Estimate</td>
<td>Units Year</td>
</tr>
<tr>
<td>2009</td>
<td>7%</td>
</tr>
<tr>
<td>2009</td>
<td>3%</td>
</tr>
<tr>
<td>Many of the health benefits included in the totals are realized after 2031 (as far out as 2113), but the smoking preventions that generate these benefits are estimated only for the period from 2012-2031.</td>
<td></td>
</tr>
</tbody>
</table>

| Annulized Quantified                       | 7%    |
| Qualitative                                | 3%    |
| All quantified benefits are also monetized. |
| Reduction in morbidity for dissuaded smokers who do not reach ages 18-24 between 2012 and 2031, reduction in passive smoking, reduction in infant and child health effects due to mothers smoking during pregnancy. |

| **Costs**                                  |       |
| Annualized Monetized $ millions/year       |       |
| $37.0  | $34.7  | $52.7  | Units Year | Discount | Period | Covered |
| 2009   | 7%  | 2012-31  |
| 2009   | 3%  | 2012-31  |
| One-time costs to change cigarette package labels and remove point-of-sale promotions that do not comply with the new restrictions, smaller ongoing costs for equal random display and for government activities. |

| Annulized Quantified                       | 7%    |
| Qualitative                                | 3%    |

| **Transfers**                              |       |
| Federal                                    |       |
| Annualized Monetized $ millions/year       |       |
| $36.6  | $0  | $237.8  | Units Year | Discount | Period | Covered |
| 2009   | 7%  | 2012-31  |
| 2009   | 3%  | 2012-31  |
| Some of the transfers included in the totals occur after 2031 (as far out as 2113), but the smoking preventions that generate these transfers are estimated only for the period from 2012-2031. Numbers reflect the assumption that the Federal cigarette excise tax will rise, on average, at the rate of inflation from 2012-2113. Numbers also include effects on Medicare, Social Security, Medicaid, other government insurance programs and income taxes. |

| From/To                                    |       |
| Federal Government (more specifically, general taxpayers and recipients of government services) | To: Individuals who would have been smokers in the absence of the rule but will not be smokers in the presence of the rule |
| $12.6  | $0  | $81.7  | Units Year | Discount | Period | Covered |
| 2009   | 7%  | 2012-31  |

| Other                                      |       |

| $12.6  | $0  | $81.7  | 2009  | 7%  | 2012-31  |
| Some of the transfers included in the
The term “actuarially fair” refers to insurance premiums that are exactly equal to expected losses.

Our primary estimate of annualized net benefits equals $601.4 million, with a 3-percent discount rate, or $184.5 million, with a 7-percent discount rate.

As shown in table 1b of this document, these net benefits are associated with 16,544 smoking preventions and 5,802 quality-adjusted life-years (QALYs) saved, annualized at a 3-percent discount rate, or 19,687 smoking preventions and 1,749 QALYs saved, annualized at a 7-percent discount rate.

As shown in table 1a of this document, the primary estimate of annualized monetized $ millions/year totals occur after 2031 (as far out as 2113), but the smoking preventions that generate these transfers are estimated only for the period from 2012-2031. Numbers reflect the assumption that State cigarette excise tax rise, on average, at the rate of inflation from 2012-2113. Numbers also include effects on Medicaid, other government insurance programs, income taxes, private insurance, pensions and life insurance programs.

**Table 1a.---Summary of Benefits, Costs and Distributional Effects**

<table>
<thead>
<tr>
<th>Economic Data: Costs and Benefits Statement</th>
<th>Notes</th>
</tr>
</thead>
<tbody>
<tr>
<td>Category</td>
<td>Primary Estimate</td>
</tr>
<tr>
<td>Annualized</td>
<td>$23.0</td>
</tr>
<tr>
<td>Monetized $ millions/year</td>
<td></td>
</tr>
</tbody>
</table>

| From/To | From: Individuals who would have been smokers in the absence of the rule but will not be smokers in the presence of the rule | To: General public (in some cases, via State government) |

**Effects**

State, Local or Tribal Government: Each year, State governments will lose approximately $25.1 million in excise tax revenue. There will be additional changes in Medicaid and other government health insurance receipts and outlays.

Small Business: The proposed rule would affect small entities in several industries, from tobacco farming to the retail industry. In particular, at least 20 of the 24 domestic cigarette manufacturers are small, and the one-time labeling change cost could be a significant proportion of average annual sales receipts of these firms.

Wages: No Estimated Effect

Growth: No Estimated Effect

**Table 1b.---Annualized Net Benefits, Smoking Preventions and Quality-Adjusted Life-Years Saved**

<table>
<thead>
<tr>
<th>Discount Rate</th>
<th>Net Benefits ($) mil</th>
<th>Smoking Preventions</th>
<th>Quality-Adjusted Life-Years Saved</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Primary Estimate</td>
<td>Low Estimate</td>
<td>High Estimate</td>
</tr>
<tr>
<td>7%</td>
<td>184.5</td>
<td>-52.7</td>
<td>3,326.0</td>
</tr>
<tr>
<td>3%</td>
<td>601.4</td>
<td>-40.8</td>
<td>10,889.2</td>
</tr>
</tbody>
</table>

FDA’s estimate of the benefits of the rule is determined by the predicted reduction in the number of U.S. smokers and the consequent reduction in the number of people who will ultimately become ill or die from diseases caused by smoking. In the first step of our analysis, we conclude that graphic warnings on cigarette packages will reduce smoking rates (both by encouraging smokers to quit and by deterring nonsmokers from starting). This conclusion is based on an analysis of the experience of Canada, which introduced graphic warnings on cigarette packages in December 2000. By comparing smoking rates in the United States with those in Canada and accounting for other relevant differences between the two countries, we are able to isolate the effect of graphic warnings on smoking rates from the effects of other interventions to reduce smoking in Canada and the United States. This comparison yields an estimate of how the graphic warnings required by this rule will reduce smoking rates in the United States. FDA estimates that this rule will reduce the number of smokers by 213,000 in 2013, with smaller additional reductions through 2031.

This estimated drop in the smoking rate in turn allows us to estimate benefits that will accrue to dissuaded smokers and to other members of society. Some individuals whose smoking status is not affected by the required graphic warning labels will receive benefits from the rule-induced reductions in smoking-related fires and certain financial outlays, such as life insurance premiums that are not actuarially fair, that implicitly subsidize smoking. Individuals who are dissuaded from smoking by the rule receive benefits equal to the value of cessation or avoided initiation. We use two methods of estimating this value, one that extrapolates from the price of actual cessation programs and one that measures the excess value of health improvements, over and above what smokers give up by not engaging in the activity of smoking. Our estimates of health improvements include the monetized value of life extensions, the monetized benefits from improved

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10 The term “actuarially fair” refers to insurance premiums that are exactly equal to expected losses.
health status (avoided nonfatal health consequences or morbidity from smoking), and reductions in medical costs. We do not have direct estimates for the value smokers attach to the activity of smoking, which adds some uncertainty to the second benefits estimation method. We therefore present several benefits estimates for which there is some justification in the literature or in comments on the proposed rule. For each discount rate and value of a statistical life-year (VSLY), our primary benefits result is the midpoint between the lower and upper bound values generated by the multiple estimation methods. Table 2 of this document shows the benefits broken down into the value of gained life-years, improved health status, medical cost reductions, other financial effects, and reduced fire-related losses. Most of the public health benefits from the rule will be realized in the future, perhaps several decades after the rule takes effect.

The estimated totals may understate the full public health benefits of the rule because they fail to quantify reductions in external effects attributable to passive smoking and the reduction in infant and child morbidity and mortality caused by mothers smoking during pregnancy. These benefits are likely to be significant, but FDA has been unable to obtain reliable data with which to quantify them with greater precision than an order-of-magnitude approximation which will be discussed in the “Benefits” section of this Analysis of Impacts. In particular, we were not able to project future levels of exposure to secondhand smoke (passive smoking) from historical trends. We were also unable to quantify reductions in the cost of excess cleaning and maintenance costs caused by smoking.

Table 2.--Benefits of Regulation

<table>
<thead>
<tr>
<th>Impacts of the Rule</th>
<th>3 percent</th>
<th></th>
<th>7 percent</th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Low</td>
<td>Medium</td>
<td>High</td>
<td>Low</td>
</tr>
<tr>
<td>Smokers’ Life-Years Saved</td>
<td>237.6</td>
<td>465.1</td>
<td>692.7</td>
<td>66.1</td>
</tr>
<tr>
<td>Health Status Improvements</td>
<td>49.9</td>
<td>97.8</td>
<td>145.6</td>
<td>22.8</td>
</tr>
<tr>
<td>Medical Expenditure Reduction</td>
<td>28.0</td>
<td>27.7</td>
<td>27.6</td>
<td>22.8</td>
</tr>
<tr>
<td>Other Financial Effects</td>
<td>27.4</td>
<td>27.5</td>
<td>27.6</td>
<td>15.4</td>
</tr>
<tr>
<td>Fire Loss Averted</td>
<td>7.1</td>
<td>12.4</td>
<td>17.6</td>
<td>3.2</td>
</tr>
<tr>
<td>TOTAL</td>
<td>349.9</td>
<td>630.5</td>
<td>911.1</td>
<td>130.3</td>
</tr>
</tbody>
</table>

Note: Table entries are annualized over 20 years, but many of the benefits represented will not be realized until well beyond the 20th year of the rule’s implementation. (Details of timing appear in Technical Appendix X3.) The ranges in the table are generated by three values of a statistical life-year: $106,308 (low), $212,615 (medium), and $318,923 (high).

The total estimated costs of implementing cigarette graphic warning labels include $319.5 million to $518.4 million in one-time costs and $6.6 to $7.1 million in annual recurring costs. Annualized over 20 years, the total costs range from $27.4 million to $40.8 million with a 3-percent discount rate and from $34.7 million to $52.7 million with a 7-percent discount rate, as shown in table 3 of this document. These totals include the costs to manufacturers of changing cigarette labels, the administrative and recordkeeping costs to manufacturers of ensuring equal and random display of the nine different warning labels over time, the costs to large manufacturers of market-testing new cigarette package labels, and the costs to manufacturers and retailers of removing point-of-sale advertising that does not comply with the rule. There are also costs to the Government of administering and enforcing the rule. FDA could not quantify every regulatory cost. Some commercial sectors will experience costs for short-term dislocations of current business activities, but the costs will be mitigated for those businesses that anticipate the industry’s adjustments to the final rule. In addition to the costs described previously, the rule will lead to private costs in the form of reduced revenues for many firms in the affected sectors. These sector-specific revenue reductions are for the most part distributional effects and cannot be counted as social costs.

Table 3.--Costs of Regulation

<table>
<thead>
<tr>
<th>Requirements of the Rule</th>
<th>Annualized Costs ($ million)</th>
<th></th>
<th>7 percent</th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>3 percent</td>
<td>Low</td>
<td>Med</td>
<td>High</td>
</tr>
<tr>
<td>Private Sector</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Label Change</td>
<td>17.8</td>
<td>19.3</td>
<td>30.3</td>
<td>24.0</td>
</tr>
<tr>
<td>Market Testing</td>
<td>0.1</td>
<td>0.1</td>
<td>0.5</td>
<td>0.1</td>
</tr>
<tr>
<td>Point-of-Sale Advertising</td>
<td>3.0</td>
<td>3.0</td>
<td>3.0</td>
<td>4.0</td>
</tr>
<tr>
<td>Continuing Admin and Recordkeeping</td>
<td>0.4</td>
<td>0.6</td>
<td>0.8</td>
<td>0.3</td>
</tr>
<tr>
<td>Subtotal</td>
<td>21.2</td>
<td>23.0</td>
<td>34.7</td>
<td>28.5</td>
</tr>
<tr>
<td>Government</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>FDA</td>
<td>6.2</td>
<td>6.2</td>
<td>6.2</td>
<td>6.2</td>
</tr>
<tr>
<td>Other (Cessation Resource)</td>
<td>6.2</td>
<td>6.2</td>
<td>6.2</td>
<td>6.2</td>
</tr>
<tr>
<td>Subtotal</td>
<td>27.4</td>
<td>29.1</td>
<td>40.8</td>
<td>34.7</td>
</tr>
</tbody>
</table>
As tobacco industry revenues decline, State and Federal tobacco tax revenues will also fall. If excise tax rates on tobacco products remain at current levels, annual State tax revenues will fall by approximately $25.1 million and annual Federal tax revenues by $19.3 million.

In the following section, FDA responds to comments on the economic analysis of the proposed rule. The full economic analysis of the final rule begins in section XLC of this document.

B. Comments on the Preliminary Regulatory Impact Analysis

1. General

In the Preliminary Regulatory Impact Analysis (PRIA), FDA estimated various benefits, costs and transfers brought about by the graphic warning label rule. We received comments on the PRIA from approximately seven tobacco manufacturers or industry groups, one advertising industry group, four nonprofit organizations, a group of researchers and an individual researcher affiliated with a medical school, two economists submitting on behalf of the tobacco industry, one additional economist, and several private citizens. Two comments related to the scope of the effects that should have been estimated in the PRIA and to a parameter choice that affected several portions of the analysis.

(Comment 212) One comment stated that FDA’s use of a 7-percent discount rate is not appropriate.

(Response) The use of both 3-percent and 7-percent discount rates is standard practice in regulatory impact analysis and is required by OMB Circular A-4 (Ref. 103).

(Comment 213) One comment stated that FDA should measure the scope of the following potentially rule-induced phenomena: Increases in the purchase of illicit cigarettes (counterfeits, contraband, cheap whites, etc.), increases in the presence of nondomestic products (duty-free, etc.), and decreases in the presence of legal domestic products.

(Response) FDA has performed a quantitative analysis of the regulation’s effect on domestic cigarette consumption (sections XID.1 and Technical Appendix X6) and a qualitative analysis of the international effects of the regulation (section X1.H of this document). FDA agrees that it would be useful to include the effect of the rule on illicit cigarette trading in the regulatory impact analysis. However, due to data limitations, FDA has been unable to quantify this effect.

2. Need for the Rule

In the preliminary impact analysis of the graphic warning label rule, FDA cited our statutory mandate as the primary need for the regulation. We received a comment stating that we had failed to discuss the economic rationale for the rule.

(Comment 214) One comment stated that FDA, in the preliminary Analysis of Impacts, failed to identify the market failure that the regulation is addressing. The comment went on to state that warning labels are a means of disseminating information, and if consumers are already fully informed about a particular product, there can be no increase in consumer welfare due to the addition or revision of a warning label.

(Response) An absence of adequate information is a well-established market failure, one which provides a rationale for disclosure requirements. There is evidence that smokers may not be fully informed of the risks associated with cigarette smoking and that large graphic warning labels can be more effective at providing information than small, text-only warnings. There is also evidence that those who have an accurate understanding of the statistical risks may underestimate their personal risks; and even where consumers have an accurate understanding, the risk might not be considered at the time of purchase (Ref. 183).

Evidence on some of these points is provided by O’Hegarty et al. (Ref. 111), who find that young American consumers are aware of some health consequences of smoking, such as the increased probability of lung cancer, but not of others, such as the increased probability of stroke. Other evidence on this question comes from Khwaja et al. (Ref. 112), who find that smokers aged 50 to 65, unlike their nonsmoking counterparts, underestimate their personal probability of dying within the next 10 years. Borland and Hill (Ref. 63, Borland 1997) find that Australia’s requirement of larger warning labels increased tobacco consumers’ knowledge that smoking causes cancer, heart and circulatory illnesses, and pregnancy-related problems. O’Hegarty et al. (Ref. 111) report that American focus group members anticipate that Canadian-style large, graphic warning labels would be more effective at communicating health information than the labels currently required in the United States. Evidence from the International Tobacco Four-Country Survey (Ref. 26, Hammond 2006) supports this conclusion, with Canadian smokers more likely than smokers from the United States, United Kingdom, or Australia—countries that required only text warnings at the time of the survey—to know that smoking causes heart disease, stroke, and impotence and that cigarettes contain such chemicals as carbon monoxide and cyanide.

The U.S. Census indicates that nearly 11 million respondents in the year 2000 did not speak English well or very well (Ref. 102); the non-English-speaking population has likely increased in the intervening years. Moreover, the Department of Education reports that, in 2003, 30 million American adults, aged 16 and over, possessed “below basic” prose literacy skills (Ref. 113). Images of smoking’s consequences and translation of warnings into Spanish and other languages can provide health information to consumers who lack English literacy.

FDA also notes that the economics and psychology literatures suggest several rationales, other than incomplete or imperfect information, for policy intervention that is aimed at a market failure. The growing literature on myopia, self-control, and time-inconsistency examines situations in which consumers may overvalue (relatively modest) short-term benefits and undervalue (relatively large) mid-term or long-term harms. The theoretical and empirical evidence suggests the possibility that through their decisions at early stages, smokers may impose significant costs on their future selves, producing net losses in terms of welfare; if so, these costs might legitimately be taken into account for purposes of policy. Helping to inaugurate the modern literature, Thomas Schelling suggests in a series of papers that smoking and similar behaviors characterized by attempts to quit and relapses can be interpreted as a contest between two selves: One self trying to stop smoking for health reasons and the other self wanting to continue to smoke. These alternating preferences violate the assumption of stable preferences and can provide a rationale for policy interventions (Refs. 106, 107, and 108).

Discussing another potential rationale for policy intervention, Gruber and Köszegi (2001) (Ref. 104) state: “While the rational addiction model implies that the optimal tax on addictive bads should depend only on the externalities that their use imposes on society, the time inconsistent alternative suggests a much higher tax that depends also on the ‘internalities’ that use imposes on consumers.” With the graphic warning label rule, FDA is undertaking a policy option that, like a tax, limits cigarette consumption, and we reach a conclusion similar to that of Gruber and
Közegi; we find that individuals who are dissuaded from smoking are made better off (i.e., they receive a net benefit) as a result of government policy intervention. (We note that Gruber and Mullainathan (Ref. 182), using subjective well-being data, find that one regulatory tool—excise taxation—has a positive effect on the happiness of those with a propensity to smoke, a result consistent with the results we present in this analysis.)

Bernheim and Rangel (Ref. 105) find that the benefits of smoking (realized by smokers themselves) are less than the realized health costs, but chemical reactions in the brain cause the consumer to mistakenly forecast more benefits when making consumption choices than he or she actually realizes from consuming the addictive product. These authors suggest that this overestimation occurs through a flawed hedonic forecasting mechanism in which particular environmental cues lead a smoker to move into a “hot” state in which he or she overestimates the pleasure from smoking. This analysis suggests that graphic warning labels may be able to serve as counter-cues that prevent movement into the hot state and allow the addict to continue to exercise self-control.

Laux (Ref. 109) identifies other reasons that smokers may not fully internalize the costs of their addictive behavior, including teen addiction as an intrapersonal (two selves) externality, partially myopic adult behavior, and peer effects.

According to the model developed by Gul and Pesendorfer (Ref. 110), if graphic warning labels reduce the temptation associated with the addictive product, they will reduce smoking and increase social welfare.

3. Benefits

In the preliminary impact analysis, FDA estimated a variety of welfare-enhancing effects of the graphic warning label rule; these included reductions in smoking-related mortality, morbidity, medical expenditures, and fire damage. We received many comments on the methods, assumptions, choice of sources, and results that were reported in the benefits analysis.

(Comment 215) One comment stated that FDA’s preliminary estimate of the rule-induced smoking rate reduction was too low, in that it ignored the rule’s effect on initiation, in favor of a cessation-only analysis.

(Response) For both the proposed rule and the final rule, FDA has analyzed the national adult smoking rate (i.e., the nation’s smoking population divided by the nation’s total population). The smoking rate at any particular moment is a function of all past initiation, cessation, birth, death, and migration of smokers and nonsmokers across national borders. Therefore, our approach includes the effect of the rule on initiation.

(Comment 216) One comment stated that FDA’s preliminary estimate, that only 82,000 individuals would be dissuaded from smoking between 2014 and 2031, was too low.

(Response) FDA’s estimate that the rule-induced reduction in the United States smoking population will occur mostly during the first year after implementation of graphic warning labels is a product of the simplicity of our empirical model. We agree that a time trend of the effect of the rule is to be preferred over a single average effect. However, our attempts to estimate linear or quadratic time trends have produced highly implausible results, especially for projections furthest into the future. We are then left with a best estimate of how the rule would decrease the U.S. smoking rate in which number of dissuaded smokers is smaller for any year from 2014 to 2031 than for 2013. This estimated change is not a decrease from year to year (e.g., 2013 to 2014), but a net decrease for a given year in the presence of the rule compared with the same year in the absence of the rule.

(Comment 217) Two comments stated that FDA’s preliminary estimate of smoking rate reduction was too low, in that it ignored the fact that someone who is dissuaded from smoking in 1 year will likely remain a nonsmoker in future years.

(Response) FDA notes that the likelihood that an individual dissuaded from smoking in a particular year will likely continue to be a nonsmoker in subsequent years was accounted for by our preliminary estimate, which had the U.S. smoking rate continuing to be lower than it otherwise would have been in years 2014 through 2031, not just in 2013. The same characterization holds for the estimate in FDA’s Final Regulatory Impact Analysis.

(Comment 218) One comment stated that “Canada has used graphic warnings for years, and in the last decade their smokers dropped from 23% to 22% of the population.”

(Response) Canada’s smoking rate has decreased by around seven percentage points, not one, since the implementation of graphic warning labels in late 2000. Even if the one percentage point statistic was correct, a one percentage point decrease in the smoking rate is not enough to completely offset the small change when applied to the large population of the United States; in fact, it would imply that there would be more than 3 million dissuaded American smokers.

(Comment 219) One comment stated that the required label change would have very little impact on smoking rates because minors, who form the bulk of new smokers, obtain their cigarettes from parents rather than from retail establishments.

(Response) Due to lack of data, FDA’s estimates of the amount of smoking cessation or avoided initiation brought about by the rule include only adults aged 18 and above, or young persons who reach age 18 by the year 2031. The number of minors dissuaded from smoking by the rule may be substantial. Whether they obtain cigarettes from friends, through theft, or by purchasing them from retail establishments operating in violation of youth access laws, young people will be exposed to new graphic warning labels because the labels are printed directly on cigarette packages.

(Comment 220) One comment stated that FDA’s preliminary estimate of the rule-induced smoking rate reduction was too high, in that it did not address potential competitive responses of the cigarette companies to the proposed rule. The comment went on to state that, under the proposed rule, graphic warning labels would take up a substantial portion of the area in packaging and advertising where firms establish brand recognition, thus reducing consumers’ ability to distinguish premium from discount brands. This would cause premiums for branded cigarettes to decrease and price competition to intensify, which in turn would likely lead to an increase in cigarette usage.

(Response) FDA believes that, even for well-known brands only have half a package with which to advertise themselves, they still have name recognition. We expect that consumers will continue to be able to find their preferred brands; as a result, any change in prices due to competitive pressures is likely to be small.

The cigarette producers’ strategic responses suggested by the comment should have occurred in Canada when that country implemented graphic warning labels. Because FDA’s estimate of the effect of graphic warning labels is based on the Canadian experience, we implicitly account for any decrease in the price of cigarettes caused by competition between premium and discount brands. Our point estimate indicates that the net effect of graphic warning labels is an increase in the national smoking rate in spite of this possible offsetting effect.
(Comment 221) One comment stated that FDA’s preliminary estimate of the rule-induced smoking rate reduction was too high, in that it failed to recognize or control for other regulatory changes (such as smoking bans) affecting cigarette consumption at the State, provincial, or municipal levels. (Response) FDA acknowledges that our model does not explicitly allow for many potential confounding factors, but we note that our estimates of the effect of graphic warning labels could as easily be underestimates as overestimates. More specifically, our model will produce an overestimate if: Smoking-reducing phenomena (other than graphic warning labels) were growing in prevalence or effectiveness at a faster rate in Canada after 2000 than before 2001, smoking-reducing phenomena (other than graphic warning labels) were more prevalent or effective in Canada than in the United States after 2000, or smoking-reducing phenomena (other than graphic warning labels) were less prevalent or effective in Canada than in the United States before 2001. In the opposite cases, our model will produce an underestimate. In the absence of extensive high-quality data, we assume that trends in smoking-reducing phenomena (other than graphic warning labels) were about the same before and after the year 2000 and about the same in Canada and the United States.

(Comment 222) One comment stated that FDA’s preliminary estimate of the rule-induced smoking rate reduction was too high, in that it did not account for potential differences in responder bias between United States and Canadian surveys created by different levels of stigma associated with smoking in the two countries. (Response) FDA generates its estimate not only by comparing Canada with the United States but also by comparing each country with itself. Specifically, we find the difference between each country’s actual 1994 through 2009 smoking rates with rates predicted by a pre-2000 trend (which accounts for changes in cigarette taxes), and then calculate how the average difference for 2001 through 2009 compares with the average difference for 1994 through 2000. The trend at least partially controls for any steady change over time in responder bias within a given survey, and the within-country comparison of pre-2001 and post-2000 rates controls for any difference in responder bias between the two countries.

(Comment 223) One comment stated that FDA’s preliminary estimate of the rule-induced smoking rate reduction was too high, in that it did not account for differences in cigarette prices over time in the United States and Canada. (Response) For the analysis of the final rule, FDA has incorporated changes in Canadian and United States tax rates into its estimates.

This comment suggests elsewhere that graphic warning labels will cause prices to decrease. FDA agrees that this is a possibility. Thus, for the non-tax portion of cigarette prices, we are faced with what economists call an endogeneity problem; it is difficult to determine, in an empirical analysis in which price is used directly as a control variable, the direction and magnitude of causality. However, if the changes in the non-tax portion of prices in the United States and Canada follow the same pattern post-2000 as they did pre-2001, and if the relationship between smoking status and cigarette prices was also relatively constant between the two time periods, then our smoking rate trends successfully control for the effect of non-tax price changes on smoking rates.

(Comment 224) One comment stated that FDA’s preliminary estimate of the rule-induced smoking rate reduction was too high, in that it did not account for the fact that Canada’s Tobacco Act’s prohibitions on advertising and promotion came into full effect after the introduction of the graphic cigarette labels. The comment went on to state that other local regulations (such as restrictions on the retail display of tobacco products and advertisements) that came into effect in Canada after the year 2000 also may have had an effect on smoking rates in Canada, and thereby would have inflated FDA’s estimate of the expected rule-induced reduction in smoking rates.

(Response) From 2001 to 2008, at least 41 states, plus the District of Columbia, enacted or substantially updated legislation regarding tobacco advertising and promotion, youth access or sampling and distribution (Ref. 114). FDA concludes, therefore, that the U.S. experience provides a reasonably good control for the effect of local and regional policy changes on national smoking rates.

(Comment 225) One comment stated that FDA’s preliminary estimate of the rule-induced smoking rate reduction was too high, in that it failed to account for the fact that, in April 2001, the Government of Canada launched a Federal public education, outreach, and mass media campaign that had a goal of reducing tobacco-related death and disease among Canadians.

(Response) The U.S. experience provides a reasonably good control for the effect of media campaigns on smoking rates because antismoking initiatives have been active in the United States in the past decade. For example, the “Truth” Campaign, a nationwide advertising effort aimed at discouraging youth smoking, launched in the United States in 2000 and continued into the 2000s.

(Comment 226) One comment stated that FDA’s preliminary estimate of the rule-induced smoking rate reduction was too high, in that it failed to account for the fact that individuals over age 65 are less likely to be smokers than younger individuals and Canada’s population is aging more rapidly than that of the United States. Specifically, during the period 2001 through 2009, Canada’s over-65 population grew by 21 percent while the U.S. over-65 population grew by only 12 percent. Canada’s over-65 population represented 13.9 percent of its total population in 2009, up from 12.9 percent in 2001. This compares to the U.S. over-65 population which increased to 12.9 percent in 2009, up from 12.4 percent in 2001.

(Response) FDA notes that the comment’s finding (that individuals over age 65 have a lower probability of being smokers than individuals aged 65 and below) does not necessarily imply that aging causes individuals to cease smoking. Smoking rates are much lower in the over-65 age category than in the 65-and-under category because smokers are less likely than nonsmokers to survive to and live past the age of 65.

Possible reasons for the aging of a nation’s population include: A decrease in the birth rate, net emigration of relatively young people, net immigration of relatively old people, a decrease in the death rate of relatively old people, or an increase in the death rate of relatively young people. If the changes in these population phenomena in the United States and Canada follow the same pattern post-2000 as they did pre-2001, and if the relationship between smoking status and the population phenomena was also relatively constant between the two time periods, then our smoking rate trends successfully control for the effect of population changes on smoking rates. (Of course, there is a correlation between smoking rates and death rates, but it operates with sufficient lag so as not to confound our results to a meaningful degree.)

(Comment 227) Several comments suggested that the lack of statistical significance of FDA’s estimate of the effect of graphic warning labels on Canada’s smoking rate implies that there is no sound basis for the conclusion that the proposed (and now final) rule’s benefits exceed costs and that this creates a
violation of Executive Order 12866, which requires government agencies to show the quantitative benefits exceed the quantitative cost from a regulation. One comment further noted that FDA did not, in the preliminary analysis, report whether its secondary methodology (in the Uncertainty Analysis) produced an estimate that was statistically significant.

(Response) Executive Order 12866 states that: “Each agency shall assess both the costs and the benefits of the intended regulation and, recognizing that some costs and benefits are difficult to quantify, propose or adopt a regulation only upon a reasoned determination that the benefits of the intended regulation justify its costs.” The point estimates indicate that the benefits of the rule justify the costs. Although our analysis concludes, on this basis, that graphic warning labels will be effective at reducing smoking, we recognize there is large uncertainty about the size of the effect. The lack of statistical significance in FDA’s smoking rate estimate reflects this uncertainty, as well as the noisiness of data derived from surveys and the small number of observations.

The use of a point estimate (which indicates that graphic warning labels have decreased the smoking rate in Canada) is appropriate for the main portion of our analysis as long as we state clearly the lack of statistical significance. Moreover, in the final analysis, we report the results of Monte Carlo simulations to better show the uncertainty. To do so, we follow the advice of Vining and Weimer (Ref. 115): “In view of the large number of uncertain effects and shadow prices involved in applying BCA [benefit-cost analysis] to social policies, analysts must take special care in dealing with uncertainty. Rather than setting estimates of effects equal to zero when their estimates are statistically insignificant, a more appropriate approach is to take account of the uncertainty of these effects in Monte Carlo simulations.”

In addition to reporting Monte Carlo results, FDA has added additional discussion which will allow the interested reader to examine our empirical approaches in greater detail.

(Response) The study research commissioned by FDA and included in the docket analyzes the reactions of consumers to each image. We cannot yet know the effectiveness of each image on improving health outcomes (such as avoidance of cancer) because the images have not yet appeared on cigarette packages or advertisements. Our best estimate of the images’ collective effect comes from Canada’s experience with a collection of graphic warning labels.

(Comment 229) One comment stated that FDA should use worldwide data if its model of smoking reduction cannot achieve statistical significance using only Canadian data.

(Response) FDA disagrees because, culturally and geographically, Canada provides a closer comparison for the United States than any other country. Moreover, in many countries, graphic warning labels have been implemented for only a few years, so any international additions to our data set would likely contribute only a small number of data points while simultaneously necessitating the addition of extra variables (for example, geographic and time fixed effects) into the model, thus producing only a small overall increase in degrees of freedom and introducing potential errors due to more omitted variables.

(Comment 230) One comment stated that FDA should use data from New York City’s experience with a graphic image media campaign, which reduced smoking prevalence in that State by 1.4 percentage points between 2005 and 2006.

(Response) FDA prefers the Canada-United States empirical model over a potential New York model both because Canada’s graphic warning policy is much more similar to the present rule than is New York’s television-based campaign and because Canada’s policy has been in place for a longer period of time than New York’s, thus providing more data points. Furthermore, we note that the New York experience would likely yield a much lower (than 1.4 percentage points) estimate of the effect of graphic images if only the excess smoking rate changes, beyond New York’s own trend and the changes experienced simultaneously in comparable cities or States, were included.

(Comment 231) Several comments stated that Sloan and coauthors’ estimates of the number of life-years lost by smokers are too low and recommended that FDA use other, higher estimates that appear in the scholarly literature.

(Response) The comments making this point have confused the life-years lost for a lifetime smoker (compared with a nonsmoker or quitter) with the measure that FDA needs for its analysis: the adjusted life expectancy changes that make up the incremental effects of reduced smoking rates induced by the final rule.

Regarding life-years lost for a lifetime smoker (compared with a nonsmoker or quitter), Sloan and coauthors’ estimates (Ref. 116) do not differ much from those reported in other studies. Specifically, Sloan et al. use results from the Taylor et al. (Ref. 117) study, which reports that men who quit smoking at age 35 gain 8.5 years of life expectancy and male never-smokers gain 10.5 years. In comparison, Doll et al. (Ref. 118) find that if an individual avoids smoking entirely or quits at age 30, he increases his life expectancy by 10 years. Strandberg et al. (Ref. 119) find that smoking shortens life expectancy for males by 7 to 10 years.

Sloan et al. adjust the Taylor et al. results to account for the probability that an individual who smokes at a given age will quit sometime later in his or her life and for confounding factors, such as differences in demographic characteristics and behaviors between average smokers and nonsmokers. Unlike Sloan et al., the studies cited in comments estimate the longevity gains to an individual from not smoking or from quitting at a given age but do not incorporate the probabilities of quitting at each age or isolate the effect of cigarette consumption from other risk factors that tend to be correlated with smoking. These studies are therefore inappropriate for a regulatory impact analysis estimating the incremental effects of warning labels on lifetime mortality consequences related to smoking at a particular age.

(Comment 232) Two comments expressed concern that Sloan and his coauthors’ analysis is outdated. One of the comments went on to state that Sloan et al.’s literature review contains some studies that have been funded by the tobacco industry and their “defense of rational addiction” may be undermining FDA’s effort to “ensure that its economic analysis is based on empirical evidence, not theoretical predictions from the rational addiction model.”

(Response) The Sloan et al. results that FDA uses are empirical, not theoretical. In producing these empirical results, Sloan and coauthors use data from the 1990s; while this is somewhat out-of-date, no analysis as
detailed as that of Sloan et al. has been released more recently. The comment critiques some of the literature reviewed by Sloan and coauthors but not the methods Sloan et al. use to produce their life tables and other results. FDA has thus continued to use these results in its Final Regulatory Impact Analysis.

(Comment 233) One comment stated that the FDA provided in its preliminary Analysis of Impacts virtually no details on its calculation of the benefit of expected life-years saved.

(Response) FDA has added a more detailed explanation to the final Analysis of Impacts.

(Comment 234) One comment stated that, in its estimate of rule-induced emphysema reductions, FDA did not provide any documentation supporting its calculations.

(Response) FDA has replaced its analysis of rule-induced emphysema reductions with an analysis of general health effects. Simultaneous with this change has been an expansion of our explanation of methodology.

(Comment 235) Several comments stated that morbidity effects other than emphysema were inappropriately excluded from FDA’s preliminary analysis.

(Response) FDA has expanded its morbidity estimates for the final Analysis of Impacts. Instead of analyzing individual diseases, we have calculated rule-induced changes in general health status (categorized as poor, fair, good, very good, or excellent).

(Comment 236) Several comments stated that benefits due to reductions in secondhand smoke exposure and mothers smoking during pregnancy were inappropriately excluded from FDA’s preliminary analysis.

(Response) FDA did not exclude discussion of these effects from the preliminary Analysis of Impacts, but we were not able to quantify them due to the difficulty of projecting future secondhand smoke exposure levels from historical trends. Similarly, we were not able to project future reductions in maternal smoking during pregnancy. In the Final Regulatory Impact Analysis, FDA has again been unable to quantify these benefits.

(Comment 237) One comment stated that FDA’s analysis includes only health benefits that accrue in the distant future, not immediate benefits of cessation or avoided initiation.

(Response) FDA’s preliminary and final estimates of morbidity and mortality effects include discounted totals of all future effects, both short-term and long-term. For example, we obtained our life expectancy estimates from Sloan et al.’s life tables. Calculated for 24-year-olds, these tables include survival probability differences for smokers and nonsmokers as early as the 25th birthday.

(Comment 238) One comment stated that FDA’s assumptions regarding the distribution of benefits over dissuaded smokers’ lifetimes were incorrect.

(Response) In many cases, FDA’s sources reported smoking-related effects only as present values calculated with a single discount rate and for a particular age group. In order to expand our results to other age groups or discount rates, it was necessary that we make assumptions about the timing of benefits. The absence of data prevents FDA from confirming the degree of inaccuracy of our assumptions. For the final analysis, we have expanded our discussion of the likely direction of estimation error that may be caused by our assumptions and, in one case, have accounted for uncertainty related to assumption-making in our Monte Carlo analysis.

(Comment 239) One comment stated that Sloan et al.’s estimates of smoking-attributable medical cost ($3,757 per female and $2,617 per male) are too low. The comment went on to recommend the use of Thomas Hodgson’s estimate (Ref. 120) that this cost, in 2009 dollars and discounted at a 3 percent rate, is $18,967.

(Response) FDA believes that Sloan et al.’s estimates are to be preferred over Hodgson’s because Hodgson does not adjust for confounding effects (by analyzing “nonsmoking smokers,” a theoretical comparison group Sloan et al. used to account for the effects of other risky behaviors) and Sloan et al.’s data sets are more recent (from the 1990s, rather than 1978 through 1988). The comment calculates the present-dollar value of Hodgson’s medical cost estimates using the medical component of the consumer price index (CPI). For the Final Regulatory Impact Analysis, FDA will do the same because medical costs have risen at a very different rate than overall price levels and thus the measure of inflation used in the PRIA—the gross domestic product (GDP) deflator—is not the best available option for updating medical costs.

(Comment 240) One comment stated that FDA’s medical cost results were not adjusted for inflation in the preliminary Analysis of Impacts.

(Response) FDA’s medical cost estimates were adjusted for inflation in the analysis of the proposed rule; however, our language on this issue was unclear and has been revised for the analysis of the final rule.

(Comment 241) One comment stated that, in the preliminary analysis, FDA provided only a very high-level and cursory description of how it arrived at its estimate of reduced fire costs.

(Response) For the final analysis, FDA has expanded the discussion of how fire loss reductions were calculated.

(Comment 242) One comment stated that FDA’s assumption that the introduction of self-extinguishing cigarettes would reduce the incidence of smoking-related fires, with or without the proposed rule, by 50 percent was arbitrary.

(Response) FDA agrees that the 50 percent assumption lacked empirical support. For the final analysis, we use a data-driven estimate of the effectiveness of self-extinguishing cigarettes at preventing accidental fires.

(Comment 243) Two comments stated that FDA’s preliminary benefits analysis inappropriately excluded effects of the rule on employee productivity.

(Response) FDA estimates morbidity and mortality effects using a willingness-to-pay approach, estimated using the QALY metric as the base. Willingness-to-pay to avoid morbidity, as we use it in this analysis, includes the subjective value of avoiding an illness that affects mobility, self-care, usual activities (including work), pain or discomfort, and anxiety or depression. These elements encompass the value of market and nonmarket productivity, and much else. Therefore, in general, the value to smoking employees of productivity effects is implicitly included in both morbidity and mortality benefits; adding productivity effects separately would almost certainly lead to double counting of some of the benefits that accrue to dissuaded smokers. Economic theory predicts that, for employers, rule-induced productivity effects generate no long-term net benefit or cost because greater firm output will be offset by the greater wages commanded by the more productive employees.

(Comment 244) One comment stated that “FDA’s analysis could benefit from a more fulsome explanation of the concept of QALY.”

(Response) FDA has added the final analysis accordingly.

(Comment 245) FDA received several comments in regard to its downward adjustment of benefits estimates to account for consumer surplus loss. One comment stated that such an adjustment should not be performed at all because doing so requires an inaccurate assumption that smokers enjoy smoking. Three comments suggested that, if an adjustment is performed, it should not be 50 percent of gross health benefits, as suggested in FDA’s cited reference, because that analysis assumes perfect
rationality on the part of smokers. Another comment objected to the model in the cited reference because it is very simplified and stylized, with a linear demand curve for smoking. One of the comments suggested FDA should instead consider modern economic analyses of addiction that account for time inconsistencies in preferences, including the work of Fritz Laux (Ref. 109) or Jonathan Gruber and Botond Köszegi (Ref. 104). Another of the comments suggested past regulatory changes and their effect on smoking be used to measure demand and the lost surplus associated with those changes to get a more empirically relevant measure of the effect of the proposed rule.

(Response) The concept of consumer surplus is a basic tool of welfare economics. If consumers respond to price, information, or other market changes, there will be a change in consumer surplus. Although some economists describe consumer surplus as a measure of the pleasure, satisfaction, or usefulness that a product provides to consumers, others simply say that whatever generates a demand for the product generates consumer surplus. Moreover, how we qualitatively describe consumer surplus does not affect how it is measured—the measurement is independent of the description. In an analysis of benefits based on willingness-to-pay, we cannot reject this tool and still fulfill our obligation to conduct a full and an objective economic analysis under Executive Orders 12866 and 13563. Although it does not affect our use of consumer surplus, we note that virtually all studies of the economics of smoking and addiction assume that smoking is pleasurable to smokers. In their 2001 paper in The Quarterly Journal of Economics, Gruber and Köszegi state that “smoking is a short-term pleasure” (emphasis added) (Ref. 104). Economists Warner and Mendez state: “Many members of the tobacco control community dismiss the notion that smoking can be pleasurable. But those people were never smokers or, if they were, have selective memory. For some smokers, the relief of withdrawal symptoms might suffice as a ‘pleasure.’ But smokers derive much more from their cigarettes, including everything from ‘mouth feel’ to the nicotine drug rush, from relaxation to self-image (think Marlboro Man), and from enhanced ability to concentrate to companionship” (Ref. 121).

FDA’s approach to the economics of smoking treats it as an addiction and draws on many economic theories of addiction, including the studies cited in the comments, as already detailed in our response to comments on market failure. FDA agrees that the model we used in the PRIA to explain changes in consumer surplus is not detailed enough to fully explain the assumptions about consumer behavior underlying our estimates. In the revised analysis, we have made some important changes in the presentation and the model used to adjust our estimates and account for uncertainty. The key assumption made explicit in the new model is that, on average, smokers are informed of, and able to internalize, some but not all health and life expectancy effects of their smoking. Full graphical and algebraic analyses have been added to the final analysis, as has a discussion of the implications of Gruber and Köszegi’s work in the context of the new model. Moreover, we have supplemented our benefits analysis with another approach, in which we replace the steps of summing all health effects and then subtracting lost consumer surplus with a direct estimation of the value to smokers and potential smokers of cessation and avoided initiation, as shown by their willingness-to-pay for cessation programs.

(Comment 246) One comment stated that FDA’s preliminary benefits analysis inappropriately excluded the effects of the rule on employer and government cleaning and maintenance costs. (Response) Reductions in the cost of cleaning and maintenance were not included in the analysis because we did not find reliable data.

(Comment 248) Two comments stated that FDA should conduct its uncertainty analysis by performing a Monte Carlo simulation. (Response) FDA agrees and has conducted a Monte Carlo simulation for the Final Regulatory Impact Analysis.

(Comment 249) One comment stated that FDA’s preliminary analysis inappropriately excluded the effects of the rule on government-funded health care and Social Security expenditures. (Response) In our analysis of the proposed rule, FDA did not exclude government health care costs. In section VIII.C.6 of the PRIA, FDA reported estimates of reductions in smoking-related medical expenditures, paid for both by smokers themselves and by nonsmokers via insurance premiums or, notably, taxes used to fund government health care. For the Distributional Effects portion of the Final Regulatory Impact Analysis, we have expanded the discussion of this effect of the rule to include greater detail.

(Comment 250) One comment took issue with FDA’s characterization of the up-front costs associated with a major label change as “large” by pointing out: “In the context of tobacco marketing, with the companies spending $12.5 billion on marketing and promotion in 2006, the amounts of money being described are not ‘large.’” (Response) FDA has added stream-of-benefits and -costs tables as appendices to the final analysis.

4. Costs

In the analysis of the proposed rule, FDA focused on three main costs to industry: The cost of changing cigarette package labels, the cost of conducting market testing for redesigned packages, and the cost of removing noncompliant point-of-sale advertising. FDA received several comments about costs, which are summarized and responded to in the following paragraphs.

(Comment 251) One comment asserted that the cost section was systematically biased, and that all costs were upper bound estimates as opposed to “best” point estimates. (Response) FDA did not rely on upper bound estimates of any costs. The label change costs (the largest single cost component FDA estimated) and the market testing costs have low, medium, and high estimates. For the other cost components, we use our best estimates.

(Comment 252) One comment argued that because tobacco manufacturers spend large amounts of money on marketing activities, changing labels is just an ordinary cost of business to them, and one that they can “write off.” Furthermore, the comment argued that manufacturers can, to some extent, pass the costs on to consumers. The comment ends by stating: “It is not appropriate for the FDA to fear that its regulatory efforts on this industry might impose costs on them, and to use these costs as a reason not to proceed with its
regulations. The agency is supposed to act in the public interest, not the interest of a particular industry to protect it from protecting the public in the first place.”

(Response) The baseline expenditures of the tobacco industry are irrelevant. There is a cost to society when its scarce resources are expended to comply with this rule. The costs the comment refers to are economic or opportunity costs. Cost estimation is concerned with the value of the resources used to carry out some activity, not their incidence (i.e., who ultimately pays), which is a separate question. As acknowledged in the proposed rule (section VII.D, Costs), although cigarette manufacturers are legally responsible for complying with this rule, the costs may be borne at least in part by tobacco consumers. The potential for “passing costs on” to consumers is a matter of economic incidence but does not negate the fact that there are costs, nor does it change those costs.

In the cost-benefit analysis we estimate costs and benefits that accrue to citizens and residents of the United States (Ref. 103) regardless of who we think may bear them. The “interest of a particular industry” is a subject we rightly leave to the “Distributional Effects” section of our analysis.

(Comment 253) A comment stated that FDA should estimate “the marginal cost of changing the warning labels that the cigarette companies would incur accounting for ongoing expenses associated with producing cigarette packages and assuming that the companies implemented the new labels using economical strategies.”

(Response) The labeling cost model’s baseline already accounts for ongoing expenses associated with producing cigarette packages. Manufacturers change product labels at regular intervals without regulatory changes in labeling requirements. Based on both product type and compliance period, the model provides an estimate of the percent of UPCs that can be coordinated with a previously scheduled labeling change. For those UPCs, the only costs assumed by the model are a small fraction of the administrative labor cost and recordkeeping costs.

If anything, this approach taken by the model quite possibly understates the labeling costs for so-called coordinated UPCs. For example, even though a graphic designer can redesign a label to satisfy both regulatory and nonregulatory goals at once, such a redesign would plausibly take longer than if the designer only satisfied nonregulatory requirements, and time devoted to regulatory compliance must be taken away from other activities. However, because this rule requires a set of 9 plates for the 9 different graphic labels, we manually adjust the model to add back the 8 extra plates.

(Comment 254) A comment asserted that although there are 3,324 different UPCs, each UPC would not have to be redesigned because product varieties within a brand family share essential trade dress and package design features. The comment asserted that using a number equal to 10 percent of the number of UPCs, 332, would still result in an overestimate of costs.

(Response) Although products within a brand family share certain package design features, the packages for different UPCs still contain unique features. Thus, every individual UPC represents a separate design job. Furthermore, the labeling cost model presents an average cost per UPC of similar types within a product category, not the cost of changing one UPC. The model therefore accounts for the existence of brand families with similar label designs.

(Comment 255) A comment asserted that FDA overestimates production and printing costs by “not accounting for the realities of how such work is actually done.” The comment provided the following quote from an unknown large job printer: “In looking at the costs associated with each label, this might be fairly accurate for 1 label, but they don’t take into account the economies of scale. After the first one, the second and subsequent package costs will go down exponentially. The only costs that might remain static would be the costs of printing plates, which depending on how they print them, could be reduced if they gang run several different packages of similar production runs together on the same sheet. All the nonproduction costs would be amortized over the whole.”

(Response) The labeling cost model does not measure the cost of changing one label, but the average cost when a large number of labels are changed at once. Due to resource constraints, the economic cost could be higher when a large number of labels are changed at once. The comment did not provide either alternate cost estimates for FDA to consider, or potential sources for such data.

(Comment 256) A comment asserted that design costs should not be inflated due to the requirement to use nine different warnings because all warnings would occupy the same portion of each package, so the design would only have to be done once regardless of which warning would be used.

(Response) The comment appears to misunderstand which cost elements are affected by the need for nine labels. The term “Design costs,” as used in the labeling cost model, could refer to all per-UPC costs associated with a labeling change or specifically to graphic design labor costs. FDA inflated some, but not all, per-UPC labeling change costs by a factor of nine.

For graphic design labor costs, FDA agrees that the part of the package design that is under the control of the manufacturer will probably be the same regardless of which of the nine warning labels is used. Therefore, the work of designing the new package label only has to be done once for each UPC; in the cost estimates, graphic design labor costs were not inflated by a factor of nine.

Likewise, FDA assumed that the need to incorporate nine different warnings on every package would have a negligible impact on administrative labor costs, prepress labor costs, and recordkeeping labor costs. These costs therefore were not inflated by a factor of nine.

It was only for materials costs, which specifically includes prepress materials and printing plate costs, that FDA assumed costs increased by a factor of nine due to the need to incorporate nine separate warning labels. We employed this assumption because nine times as many printing plates will be needed upfront.

(Comment 257) A comment argued that some of the costs attributed to the label change would be incurred on an ongoing basis. The example provided is that printing plates wear out after a few million impressions and have to be replaced at regular intervals. The comment argued our cost estimates need to be adjusted to account for this. An analysis follows which claims to demonstrate that the average cigarette label printing plate has to be replaced every 3 weeks.

(Response) The calculation provided in the comment contains errors. Once those errors are fixed, the calculation no longer supports the assertion that printing cylinders are being constantly replaced, as discussed in the following paragraphs. Furthermore, the model accounts for possible coordination with previously scheduled labeling changes, which provides the most likely opportunity for cigarette manufacturers to avoid some of the incremental cost from new printing plates (cylinders). New cylinders must be engraved when a nonregulatory labeling change takes place. Given the expense of the printing cylinders, manufacturers would avoid engraving new cylinders right before a
nonregulatory labeling change. In other words, we would expect some coordination between cylinder wear out and nonregulatory changes.

Rotogravure plates are the longest lasting, good for making millions of labels. The comment assumed a life of only 3 million labels and did not justify this point estimate. For rotogravure, this estimate is too low.

In attempting to determine weekly sales per UPC, the comment divided weekly cigarette sales (in packs) by their estimate of the number of brands, not by the number of UPCs. Dividing by the number of UPCs, even under the assumption that plates wear out after 3 million labels, yields a life of 29 weeks for the average brand. Updating this analysis for the revised number of cigarette UPCs yields a life of 38 weeks for the average brand.

Additional calculations can be performed for the “average” brand, but it is important to keep in mind that most brands are not average. A few products will be a high volume. A large number of lesser-known products will have low volume.

Because manufacturers will have to buy nine plates up front for each UPC, those nine plates would have a life of 346 weeks, or 6.6 years, based on the comment’s assumptions about the life of a rotogravure plate and the updated UPC count. Manufacturers of the average product would not wear out all these plates before they changed labels again for nonregulatory reasons.

(Comment 258) Multiple comments argued that FDA should not include 10 percent rush charges in calculating the cost of changing labels in 15 months. In particular, the argument was made that cigarette manufacturers have known this was coming before publication of the final rule.

(Response) Although it is true that manufacturers have known this rule was coming, in some form, since the passage of the Tobacco Control Act, it is only with the publication of the final rule that they will know its exact form, i.e., what the images will be. Tobacco companies will need to see the final images and the exact provisions of the final rule before the bulk of the work for a labeling change can be undertaken.

In evaluating the need for rush charges, it is important to keep in mind that the labeling model is designed to measure the cost of changing a large number of labels at once. Resources are scarce and a large number of labeling changes cannot be simultaneously rushed without increasing costs.

The previous labeling cost model assumed 10 percent rush charges for compliance periods shorter than 2 years. The new labeling cost model assumes constant rush charges equal to 40 percent for compliance periods of 3 to 15 months. In reality, rush charges are likely to decline continuously as the compliance period increases. The rush charges under a 3-month compliance period could exceed 40 percent, and the rush charges for a 15-month compliance period are likely to be far less. FDA has therefore retained the original assumption of 10 percent rush charges for a 15-month compliance period.

(Comment 259) One comment stated that FDA has underestimated costs because of technical implementation difficulties associated with providing for equal, random, simultaneous display of nine different images.

(Response) FDA does not agree that there is a technical infeasibility. Similar requirements have been successfully implemented in other countries. The cost analysis for the label change includes administrative labor and recordkeeping costs, part of which would be the devising and implementing a method for ensuring equal random display. However, FDA is now persuaded that there will be some ongoing cost associated with equal, random display. In other words, once a system for compliance is designed and implemented, it will require some work to ensure continuing compliance with equal, random display. Therefore, in the Final Regulatory Impact Analysis FDA has added recordkeeping costs and administrative costs as ongoing costs in years 2 through 20 after the final rule takes effect.

(Comment 260) Comments argued that market testing costs undertaken by the tobacco industry should not be counted. Various arguments were presented: Such costs would be beyond the minimal cost required to implement the law “effectively and in good faith.” Such costs would be incurred in order to “undermine the effect of Congressionally-mandated warning labels.” Such costs would not be societal costs at all, but distributional effects because the cost to the tobacco companies would be a benefit to employees or contractors paid to do the work. If FDA includes market testing costs, it should also include legal fees for potential challenges to this rule and lobbying fees to get the statute repealed. (Response) We do not simply estimate the cost of minimal compliance. In benefit-cost analyses of regulations, we assume agents react to a new regulation by changing behavior in many ways. The analysis itself then compares the expected outcomes with and without the rule. Regardless of whether the rule requires it, if manufacturers conduct market testing as a direct result of this rule, the costs are attributable to this rule. Resources devoted to this market testing have an opportunity cost, so there is a social cost. We have been unable to obtain reliable data with which to quantify potential costs incurred to challenge the rule in litigation. Lobbying costs associated with the repeal of the statute do not represent incremental costs of this rule and therefore are appropriately excluded from the analysis.

(Comment 261) A comment stated that cigarette manufacturers and retailers change advertisements and labels frequently and only the incremental cost of replacements that would not have otherwise been made should be attributed to this rule. The comment asserted that this incremental cost is negligible.

(Response) FDA only looked at the cost of removing point-of-sale advertisements. Other forms of cigarette advertisements are now relatively rare. The comment assumes that some or all manufacturers and retailers could perform the removal of noncompliant point-of-sale advertising at zero cost by coordinating it with the usual replacement schedule for point-of-sale advertising. Manufacturers and retailers would only remove noncompliant advertising early if the benefit from keeping them longer did not justify the modest cost (between $12 and $198 per establishment) of removing the advertising at the deadline. FDA expects that the most likely response will be for most establishments to continue displaying noncompliant advertisements up until the enforcement deadline and resources will therefore be expended to achieve compliance at the deadline.

(Comment 262) One comment stated that the cost analysis needs to include reduced government revenue from lost taxes due to lowered cigarette sales. (Response) FDA notes that, leaving aside potential deadweight loss, there are two principal effects of tax reductions: Gains to former payers and losses to former recipients. Because these effects exactly offset each other, there is no net social cost or benefit associated with the reduction in excise tax collections induced by the rule. As such, we discuss rule-induced changes in tax collections in the Distributional Effects section of our analysis (section XI.G.5 of this document).

(Comment 263) One comment stated that the disturbing nature of the graphic warning labels will cause adverse mental reactions in those who view them, especially cashiers at cigarette-
selling retail establishments because they must handle these products daily. (Response) FDA is not aware of any scientific evidence that mental or emotional costs would be incurred by the general public as a result of this regulation, and the comment did not provide any.

5. Distributional Effects

In the analysis of the proposed rule, FDA estimated a variety of effects that are equal transfers away from some segments of society and as roughly equal transfers to other segments of society. FDA received several comments about these distributional effects.

(Comment 264) One comment stated that FDA’s preliminary analysis of the rule’s effect on tax collections ignored offsetting effects due to increased sales of other taxable goods and services even though the Joint Committee on Taxation estimates this offset at 25 percent of a policy’s direct effect.

(Response) FDA agrees with the comment and has adjusted its analysis of rule-induced changes in tax collections accordingly.

(Comment 265) One comment stated that, in its preliminary analysis of the rule’s impact on tax collection, FDA suggested that inelastic demand for cigarettes means that some or all lost tax revenue could be offset through higher tax rates. The comment went on to note that FDA undertook no analysis of whether State and local governments could or would increase excise taxes on cigarettes in response to the graphic warning label rule and that the political environment, as demonstrated by recent elections, may not be amenable to tax increases.

(Response) FDA did not claim any increases in State or Federal cigarette taxes are likely to occur. Instead, we merely pointed out that cigarette demand has been shown to be inelastic; therefore, an increase in tax levels will increase revenue. For the final analysis, we have removed some of our more confusing language on this issue. We continue to assume that tax rates will rise at the rate of inflation because, without such an assumption, we need a reliable forecast of inflation in order to express the stream of future tax revenue changes in current dollars. However, we have added discussion of alternative approaches, including the possible forecasting of inflation using the difference between interest rates for Treasury Inflation-Protected Securities (TIPS) and standard Treasury bills.

(Comment 266) One comment stated that, to the extent that State and local excise taxes are based on the price of cigarettes, increased price competition that could result from the proposed rule would reduce tax revenues beyond what FDA reports in its analysis.

(Response) At present, all State and Federal cigarette taxes are applied per unit, not ad valorem; therefore, changes in the pre-tax price of cigarettes will not change the total excise tax collection separately from changes caused by decreases in the quantity sold. Sales taxes, on the other hand, are applied to cigarettes on the basis of price. FDA has not quantified the effect of the rule on sales tax collections, but we expect it to be small, both because sales taxes make up a very small portion of total cigarette-related tax collections and because any rule-induced change in cigarette prices is also likely to be small.

(Comment 267) One comment stated that, in its preliminary analysis, FDA failed to note that research indicates that U.S. employment will increase if smoking decreases.

(Response) In the PRIA (section VIII.F.2), FDA stated that decreases in smoking may cause increases in national employment, citing (Ref. 122) the same paper to which the comment refers.

(Comment 268) One comment stated that, in its preliminary analysis, estimated that the proposed rule would result in 500 to 600 displaced jobs among manufacturers, warehouses and wholesalers but failed to note that these lost jobs probably would occur during a period of high unemployment, when the displaced individuals would likely have difficulty obtaining new jobs with similar remuneration. The comment went on to state that the average unemployment duration in November 2010 was 34.5 weeks and that one could, by multiplying the average wage by the average duration of unemployment, obtain a rough estimate of lost wages.

(Response) The wages lost are not the appropriate cost to attribute to the rule; instead, we must include the difference between wages lost from tobacco-related jobs and the value of next-best options. FDA is unable to quantify this difference. For instance, average unemployment tenure from late 2010 would likely give a skewed estimate of length of rule-induced unemployment because compliance with the rule is not required until 2012. Unemployment may change substantially between now and then, especially because the United States is currently in the early stages of recovery from a recession.

(Comment 269) One comment stated that manufacturing, warehouse, and wholesaler jobs displaced by the rule would be permanent losses to the economy. In addition to failing to note this permanence, FDA did not account for any job losses in the retail sector. The comment went on to state that convenience stores are highly dependent on tobacco sales, both in terms of cigarette sales’ portion of profit margins and as a generator of customer traffic to spur the sale of ancillary products. Even the small reductions in revenue caused by the graphic warning label rule could cause retailers to reduce employment, with some stores possibly going out of business entirely.

(Response) The portion of dissuaded smokers’ budgets that would, in the absence of the rule, have been spent on cigarettes will, in the presence of the rule, be spent on other goods and services, thus creating jobs in other segments of the economy. Only the difference between losses borne by individuals losing cigarette-related jobs and gains realized by individuals obtaining employment in other sectors represents a net social cost. FDA believes this difference to be small and possibly negative (that is, the losses are less than the gains), as found by Warner et al. (Ref. 122).

(Comment 270) One comment stated that, in its preliminary analysis, FDA incorrectly concluded that there would be no rule-induced losses experienced by tobacco growers. The comment went on to state that FDA’s assumption that acreage taken out of tobacco production could be easily shifted to other crops, with no net loss, is not consistent with economic theory because economic theory indicates that land currently planted in tobacco is being used in its highest-valued use. Another comment suggested that FDA work with the Department of Agriculture on estimating the impact of the rule on tobacco farmers.

(Response) FDA agrees that a transition from tobacco cultivation to the next-best option entails some loss for farmers, but only the difference between first- and second-best uses of land represents a net social cost in terms of reduced efficiency.

(Comment 271) One comment stated that the requirement that cigarette manufacturers print half of their packaging with images supplied by the government would be a burden to all cigarette companies, the costs of which would ultimately be paid by consumers.

(Response) FDA has estimated the cost to cigarette producers of adding graphic warning labels; however, we have not assessed whether cigarette consumers or shareholders of cigarette-producing firms will bear the burden of the cost. We expect that the costs will be shared by consumers and producers but we are unable to estimate the...
portions borne by each group. In the cigarette market, increases in variable costs are borne almost entirely by consumers. In the case of the addition of graphic warning labels, however, most of the cost does not vary with the quantity of cigarettes produced. We therefore expect that producers will be unable to pass all of the cost on to consumers through increased prices. Consumer prices could, however, be affected in the long run. For example, one possibility is that some cigarette product lines will be discontinued and this decrease in supply would lead to increased prices paid by consumers. FDA lacks the detailed market data that would be necessary for predicting which of these or other possible outcomes would likely be realized.

(Comment 272) One comment argued that retailers must lose profit when reallocating space away from cigarettes to other products because it was suboptimal to make such an allocation in the absence of the rule.

(Response) This comment ignores the fact that the final rule will reduce demand for cigarettes and increase demand for other products. While it is clear by observation that allocating shelf space away from cigarettes to other products in the absence of this rule would be suboptimal, this need not imply that retailers’ profits will be lower after they optimally respond to changes in the demand for cigarettes and the demand for other products.

(Comment 273) Some comments argued that retailers (including small retailers such as convenience stores) may not be able to simply shift shelf space to other goods.

(Response) FDA argued in the distributional effects section of the proposed rule, section VIII.F.3, that the retail sector (as a whole) will shift shelf space to other products to take advantage of the increase in demand for noncigarette products. FDA acknowledges that this substitution may not take place wholly within each retail establishment. If cigarette-reliant retailers have some (but less than complete) success shifting shelf space to take advantage of the increase in demand for noncigarette products, they will suffer an overall loss in revenue that is less than their loss of cigarette sales revenue. Other parts of the retail sector would gain sales. This would be a purely distributional effect within the retail sector. Such an effect would be small because this rule is only projected to reduce cigarette consumption by less than one quarter of a percent.

6. Impact on Small Entities

In the initial regulatory flexibility analysis, FDA considered the potential effects on small cigarette manufacturers of having to change all cigarette labels in accordance with this rule. FDA also considered the potential impact on small retailers of having to remove noncompliant point-of-sale advertising. FDA received comments from industry pertaining to these matters, which are summarized in the following paragraphs.

(Comment 274) A comment stated that FDA “grossly underestimates” costs, referring specifically to the estimates of the label change costs and their impact on small manufacturers. The comment argued that the necessary changes will cost at least $500,000 to $1 million, including such factors as package redesign, dye cuts, and the number of colors needed for the artwork. Further, “these changes represent global changes for the manufacturers’ products, and that change will have a far greater effect on the small manufacturer as opposed to larger entities.” Many aspects of compliance will require the work of outside contractors.

(Response) It is not clear whether the comment intends to argue that the cost is on average $500,000 to $1 million per UPC, when many UPC labels are being changed at once, or that the total cost would be at least this much per firm, among some subset of small manufacturers. FDA does not agree that the average cost per UPC could be nearly this high. Although FDA estimates much higher total costs for the average small manufacturer, $500,000 to $1 million could describe the total costs for a subset of especially small manufacturers.

The cost estimate with which the comment takes issue was based on a combination of the old FDA labeling cost model and early estimates of some values from the new FDA labeling cost model. Costs have been updated in the analysis for the final rule to more fully reflect the estimates of the new model. Interviews with manufacturers and trade associations were conducted in the process of building the new model. FDA believes the model provides the best estimate of the average cost of changing a product label. FDA inflates materials costs by a factor of nine to account for the requirement to use nine separate warnings.

The comment also argued that FDA has underestimated the costs to small businesses but is not specific enough about whether there are additional factors, beyond the results of the labeling cost model, with which the comment disagrees.

FDA agrees that small tobacco product manufacturers are more likely to hire outside contractors for tasks required to comply with this rule. However, from a societal point of view, it makes no difference to costs whether a manufacturer conducts the functions required for compliance in-house or contracts them out.

(Comment 275) A comment argued that small manufacturers do not carry a small inventory of supplies, but must buy materials in bulk to be cost effective (often as much as 6-months worth). The comment stated therefore that it is untrue that all label inventories will be exhausted during the 15-month compliance period. Small manufacturers will have to discard large amounts of advertising and labeling material. Another similar comment argued that small manufacturers purchase long-term quantities of “advertising pieces such as pole signs and shelf talkers,” in order to get better prices. FDA should take this into account and give small manufacturers time to use up existing inventories of printed materials. The comment suggested that manufacturers could provide FDA with inventory counts and usage rates.

(Response) FDA believes the first comment combines two separate issues: Label inventory assumptions (the matter at hand in the quote from the preliminary analysis) and advertising inventory assumptions.

FDA stands by its conclusion that the costs of discarded label inventory will be small under a 15-month compliance period. With modern just-in-time inventory control methods, firms keep far less inventory on hand than in decades past. However, rather than assume that there is zero cost for discarded inventory, FDA will accept the new labeling cost model’s default assumptions regarding discarded inventory. This assumption results in a low inventory cost being attributed to this final rule, as very little inventory is expected to remain after a 15-month compliance period. While it may be the case that some small manufacturers keep large amounts of inventory on hand, the evidence used to construct the labeling cost model implies that most manufacturers would not have much (if any) label inventory remaining after 15 months and the output of the labeling model accurately represents the average inventory cost.

While it is possible that some manufacturers will have some point-of-sale advertising materials in inventory that will be discarded as a result of this
rule, FDA doubts that this inventory cost is substantial. Manufacturers will have 15 months to use up existing inventory. Cigarette manufacturers are known to be sophisticated advertisers, and effective advertising changes to reflect the times. Therefore, the value of existing advertisements would decline over time as they become more dated and less effective. Additionally, the comments themselves do not provide data with which to estimate any effect that may exist.

(Comment 276) One comment estimated that the label change cost would be between $2.1 million and $5.5 million per average small tobacco product manufacturer, based on an average number of UPCs per firm of 44. The comment asserted that small manufacturers cannot absorb the cost of changing all their cigarette labels and many will leave the cigarette manufacturing business. Two relief options were suggested: Phasing in the rotational warnings over a longer period of time or running the warnings sequentially rather than simultaneously.

(Response) According to this comment, small tobacco product manufacturers have fewer UPCs each than FDA originally estimated. If the UPC estimate from the comment holds, the compliance costs for small firms would be lower than FDA originally estimated. FDA has retained the original method for estimating the number of UPCs for small firms so as to take care not to understate the burden on them. FDA acknowledges that this rule may put some small manufacturers at risk of going out of business. However, we do not have the information necessary to estimate this risk. In the initial regulatory flexibility analysis, FDA considered the relief that would be provided by allowing small (or all) tobacco product manufacturers additional time to comply with the rule, even though this not in keeping with the statutory mandate. Running nine warnings sequentially rather than in parallel is a complicated alternative for which it is difficult to estimate the amount of relief provided. A very large reduction in costs would only materialize if the warnings were only changed as often as the usual frequency of nonregulatory label changes (every couple of years). However, FDA has now included an analysis of the potential impact of a related relief option, that of letting small manufacturers randomly assign one label to each distinct UPC.

(Comment 277) Some comments argued that some small retailers, such as convenience stores, may go out of business as a result of reduced cigarette sales and loss of revenue from ancillary products, and that this effect of the rule on small entities needs to be reflected in the analysis. Beyond the effect on the retailers themselves, closure of convenience stores would result in loss of convenience to nearby customers and could also adversely affect suppliers.

(Response) Although in the small entity analysis we are only able to quantify the cost of removing noncompliant advertising, we acknowledge that small retailers selling cigarettes could also lose some net sales revenue (to other retailers), to the extent that shifting shelf space to other goods less than fully offsets the reduction in revenue from cigarettes. We expect any such loss of revenue to be modest because the expected reduction in cigarette consumption is modest to begin with. Convenience store closures as a result of this final rule are therefore unlikely.

(Comment 278) One comment recommended that FDA reconsider exempting small cigarette producers. (Response) The initial regulatory flexibility analysis considered exempting small manufacturers from the label change requirements as a relief option. Exempting small manufacturers from any part of this regulation would cause a significant proportion of consumers to be exposed to cigarette packages or advertising lacking the new graphic warnings. In 2008, the combined market share of all but the four largest firms was 10.3 percent (Ref. 123). This situation would be inconsistent with the public health objective of the rule as well as FDA’s statutory mandate.

C. Need for the Rule

Written with the goal of ameliorating the large toll on public health that is directly attributable to the consumption of tobacco, the Tobacco Control Act mandates the publication of this rule. Section 201 of the Tobacco Control Act modifies section 4 of FCLAA to require that nine new health warning statements, along with color graphics depicting the health consequences of smoking, appear on cigarette packages and in cigarette advertisements. As discussed in detail in FDA’s response to comments in section X.B.2 of this document, the economics literature suggests several sources of market failure 11 that the new graphic warning labels will address; these include myopia, lack of salience, time inconsistency, and incomplete information. In the following analysis, we do not attempt to choose among the many models of smoking and addiction that potentially cause market failure, but the models have similar policy implications.

D. Benefits

We estimate the benefits of the final rule by comparing expected life-cycle events of smokers with those of nonsmokers. Nonsmokers tend to live longer and develop fewer cancers, cardiovascular, pulmonary, and other diseases, so the benefits in our analysis include the discounted value of life-years gained, health status improvements and medical services freed for other uses. We also include an estimate of the monetary value of the property and lives saved as a result of the rule-induced reduction in the number of accidental fires caused by smoking. There are other benefits, such as reductions in nonsmokers’ morbidity and mortality associated with both passive smoking and mothers smoking during pregnancy, that are likely generated by the final rule, but FDA has been unable to obtain reliable data with which to quantify them. In particular, we were not able to project future levels of exposure to secondhand smoke from historical trends, nor predict future decreases in maternal smoking during pregnancy.

1. Reduced Cigarette Smoking Rates

The changes outlined in this rule are projected to decrease smoking initiation and increase smoking cessation. For each of the first 20 years of the rule’s implementation (2012 through 2031), FDA calculates the predicted decrease in the number of U.S. smokers by multiplying together the following:

(a) The estimated effect (percentage point change) of cigarette warning labels on the national cigarette smoking rate and

(b) The population in a particular year in the absence of the regulation (as projected by the U.S. Census Bureau).

To obtain estimates of the effect of cigarette warning labels on smoking rates (item (a) in the list above), we look to the experience of Canada, which has required the use of graphic warning labels since December 2000 (Ref. 124). The advantage of this approach lies in our ability to observe actual consumer behavior—in the form of smoking rates—before and after a graphic warning label requirement went into

11 A situation in which a market left to itself does not allocate resources efficiently.
effect. The warning labels to be required in the final rule are generally similar to those developed by Health Canada and authorities in other foreign countries. As in Canada, the labels required by the rule will occupy at least half the front and rear display panels of a cigarette package. Moreover, under the rule, there will be a mix of warning statements and images that depict the negative consequences of smoking. Although the rule will follow much the same approach as the Canadian warning label requirements, it will differ in some ways: Canada has 16 labels in rotation, rather than 9; warning statements appear in English on one side of a Canadian package and in French on the other; and health and cessation information is included on leaflets within Canadian cigarette packages (Ref. 125). These details, combined with general differences in legal and social trends, indicate that Canada's experience with warning labels can give only a general idea of the changes in smoking rates to be expected as a result of the rule. Many factors, such as tobacco advertising restrictions, youth access restrictions, educational campaigns regarding the health effects of smoking, restrictions on smoking in indoor public places, and taxes on tobacco products have influenced smoking rates in the two countries. In order to estimate the incremental effect of the present rule, we need to isolate the impact of graphic warning labels on the Canadian smoking rate.

In order to accomplish this, as discussed in detail in Technical Appendix X1, we begin by using data from Health Canada (Refs. 126 and 127), the National Center for Health Statistics (Ref. 129), and the National Health Interview Survey (Ref. 128) to estimate pre-2001 smoking rate trends for both the United States and Canada. Because tax-induced changes in the price of cigarettes have been shown to substantially reduce smoking, in each trend estimation we include the effects of Federal and State or provincial cigarette tax changes on national smoking rates. (After decreasing substantially in the early 1990s, Canada’s real average cigarette excise tax level grew by 9 percent between 1995 and 2000 and by 123 percent between 2001 and 2009. Real average cigarette tax levels in the United States grew by 29 percent between 1995 and 2000 and by 117 percent between 2001 and 2009.) Using the estimated trends, we predict smoking rates for the United States and Canada, and the difference between them, for years up to and including 2009. We then subtract the predicted United States-Canada smoking rate differences from the actual differences observed in the data. Implicit in this method is the assumption that these otherwise-unexplained differences may be attributed solely to the presence in Canada of graphic warning labels. We do not account for potential confounding variables or for possible substitution by consumers from cigarettes to other products (such as little cigars) that may produce similar health effects; our method is therefore a rudimentary approach to estimating the smoking reduction that will be effected by the new graphic warning labels and may be producing results that are off by one or more orders of magnitude.
Using this rudimentary approach, FDA estimates that the average unexplained difference between United States and Canadian national smoking rates is 0.088 percentage points higher for the 2001 through 2009 period than for 1994 through 2000. Applying this estimate to population projections (Ref. 130 provides annual projections only through 2030, so we assume cohort populations will remain the same from 2030 to 2031); summing over all age groups yields an estimate that the rule will reduce (either through cessation or avoided initiation) the United States' smoking population by approximately 213,000 in 2013, with the total decrease rising to approximately 246,000 in 2031 due to the predicted smoking rate decrease being applied to a growing population. FDA has not quantified rule-induced decreases in cigarette consumption among smokers who do not quit entirely, although such decreases have the potential to improve health outcomes for affected individuals.

2. Quantifying Benefits That Accrue to Dissuaded Smokers

a. Smokers' willingness-to-pay for cessation programs. One method for estimating dissuaded smokers' net internal benefits involves using the amount smokers are willing to pay to participate in cessation programs. This willingness-to-pay will reduce the value of cessation (i.e., the value of health and other benefits of cessation minus any value that smokers attribute to the activity of smoking) multiplied by the participation-related probability of success. Warner et al. (Ref. 131) report that the choke price, or the price at which no smokers would participate in cessation programs, may be around $350 (in 2000 dollars), while a maximum of 10 percent of the smoking population would participate in cessation programs even if those programs had a money price of zero. With a linear demand curve, these parameters produce an average willingness-to-pay among potential cessation program participants of $175. Warner and coauthors report that approximately 15 percent of smoking cessation program participants successfully quit without eventual relapse. These parameters indicate that the average value of cessation is $175/0.15 = $1,167, or $1,444 when updated for inflation (using Ref. 132).

We estimate in section XI.D.1 of this document that the final graphic warning label rule would reduce the U.S. adult smoking population by 213,000 in 2013. In the absence of the rule, the baseline 2013 smoking population would be approximately 213,000 in 2013. In the absence of the rule, the baseline 2013 smoking population would be approximately 49.5 million, so a decrease of 213,000 represents a 0.43 percent effectiveness of graphic warning labels. The value to an individual smoker of graphic warning labels equals their effectiveness multiplied by the value of cessation, or 0.0043*51,444 = $6.22. Multiplying by the predicted 2013 smoking population yields a value of $6.22*94.5 million = $307.9 million. For each year from 2014 to 2031, we perform an analogous calculation, but we replace the entire smoking population with only the particular year's newly exposed cohort (consisting of 18-year-olds and new immigrants). This results in a present value of net intrapersonal benefits of $370.3 million, calculated with a 3-percent discount rate, or $322.4 million, calculated with a 7-percent discount rate.

While these values can provide rough estimates of the benefits of the final rule, there are several reasons to believe they are only approximations and probably reflect lower bounds. First, we are implicitly assuming that the value of avoided smoking initiation is equal to the value of cessation and that the value of cessation is equal across the entire smoking population. In fact, we have willingness-to-pay data only from those smokers who are potential participants in cessation programs. The value of avoided initiation is likely much higher than the value of cessation, which would tend to make the present estimates of rule-induced benefits too low. A second reason willingness-to-pay represents a lower bound on the rule's benefits is because it captures only the misinformation and time-inconsistent preferences that smokers themselves recognize and act upon via participation in cessation programs.

b. Gross and net health benefits. We now turn to the literature on time inconsistency, which is one of the principal forms of market failure relevant to tobacco, to develop an alternative approach to estimating rule-induced benefits that accrue to dissuaded smokers. The papers we will...
The applicability of any of the suggested net-to-gross internal benefits ratios requires an estimate of the gross benefits realized by individuals who are dissuaded from smoking. Gruber and Köszegi admit that their $30.45 per pack estimate is not exhaustive, so we now turn to quantifying morbidity, mortality, and other effects of smoking cessation and avoided initiation.

1. Expected life-years saved. The largest health consequence of smoking is the increased rate of mortality from pulmonary and cardiovascular disease, cancer, and certain other illnesses. As a result, the largest benefits of this rule stem from the increased life expectancies for those individuals who, in the absence of the rule, would be smokers and thus susceptible to premature mortality from one of these often-fatal diseases. We calculate the number of life-years saved using differences in the probabilities of survival for smokers and nonsmokers. Sloan et al. (Ref. 116) construct life tables for various categories of individuals, including “nonsmoking smokers” and typical 24-year-old smokers. A nonsmoking smoker is someone who does not use cigarettes but otherwise exhibits the lifestyle and personal characteristics of the average smoker. A typical 24-year-old smoker does not necessarily smoke for his or her entire life, but instead faces cessation probabilities that are in line with values observed for all ages in the National Health Interview Survey; the life expectancy effects of cessation at older ages are netted out of life expectancy effects of avoiding smoking at age 24 (results reported below). Sloan et al.’s life tables allow us to calculate how many additional deaths, per 100,000 population, may be expected among typical smokers than among nonsmoking smokers between the 24th and 25th birthdays, the 25th and 26th, and so on until the 100th birthday. (FSA assumes that differences in yearly survival probabilities for smokers and nonsmokers are negligible below age 24 and above age 100.)

Overall, Sloan et al. find that an average (or what Sloan et al. call “typical”) 24-year-old female smoker can expect to live another 55.5 years, while a comparable nonsmoker can expect another 57.8 years of life, producing an overall regulation-induced gain of 2.4 undiscounted life-years per individual who is prevented from starting to smoke. Comparing male 24-year-old typical and nonsmoking smokers, life expectancy increases from 49.8 to 54.2 years, producing a gain of 4.4 undiscounted years. The gap between male and female life expectancy results may be due to different physiological responses to equal amounts of smoking, different lifetime cessation patterns, or different smoking intensities. Taylor et al. (Ref. 117), for instance, find that male smokers are more likely than female smokers to consume more than a pack a day. Sloan et al. do not report how much of the male-female difference in their estimated life expectancy effects may be attributed to each possible mechanism.

We assume that each person who reaches ages 18 to 24 during the 20 years (2012 to 2031) of our analysis and is dissuaded from smoking extends his or her life by the gender-specific amount Sloan and coauthors report. For older individuals, whose post-smoking cessation survival probabilities cannot be plausibly assumed to equal those of individuals who were nonsmokers at age 24, we predict life extensions using former smoker life tables that we construct using Sloan et al.’s results and cessation probabilities from the 1996 National Health Interview Survey (Ref. 128). The details of these adjustments appear in Technical Appendix X2.

ii. Benefits of reduced premature mortality. OMB Circular A–4 (Ref. 103) advises that the best means of valuing benefits of reduced fatalities is to measure the affected group’s willingness-to-pay to avoid fatal risks. Three life-year values (also known as values of a statistical life-year, or VSLY) used frequently in the literature and in previous analyses are $100,000, $200,000, and $300,000 (Refs. 134 and 135; FR 33030, July 9, 2009), which we update to $106,308, $212,615, and $318,923 in 2009 prices. These values constitute our estimates of willingness-to-pay for a year of life preserved in the present. The economic assessment of a future life-year requires discounting its value to make it commensurate with the value of present events. As required by OMB Circular A–4, we use 3-percent and 7-percent discount rates to calculate the present value of the life-years we predict will be saved.

For each dissuaded smoker, we multiply a VSLY by the relevant age and gender-specific life extension and...
then discount appropriately to arrive at a per-person value of reduced mortality. For 24-year-olds, this value ranges from $9,280 (for a female applying a VSLY of $106,308 and a 7-percent discount rate to her 2.4 life-years gained due to smoking avoidance) to $363,333 (for a male applying a VSLY of $318,923 and a 3-percent discount rate to his 4.4 life-years gained due to smoking avoidance). Multiplying the per-person values by the predicted number of dissuaded smokers and discounting the results back to year 2011 yields estimates of rule-induced mortality benefits that range from $1.45 to $22.56 billion.

### Table 5.--Gross Present Value of Lifetime Reduced Smoker Mortality ($ mil)

<table>
<thead>
<tr>
<th>Value of a Statistical Life-Year</th>
<th>3% Discount Rate</th>
<th>7% Discount Rate</th>
</tr>
</thead>
<tbody>
<tr>
<td>$106,308</td>
<td>7,520.9</td>
<td>1,450.6</td>
</tr>
<tr>
<td>$212,615</td>
<td>15,041.7</td>
<td>2,901.1</td>
</tr>
<tr>
<td>$318,923</td>
<td>22,562.6</td>
<td>4,351.7</td>
</tr>
</tbody>
</table>

These totals may underestimate the full value of rule-induced reductions in mortality because they do not account for increasing trends in life expectancy. Sloan et al.’s results, from which our mortality estimates are derived, are based on data from the late 1990s. Sloan (Ref. 136) reports that between 1999 to 2001 and 2006 (the most recent year for which life tables have been developed), life expectancy at age 25 increased from 50.54 to 51.5 years, or 1.90 percent, for males and from 55.41 to 56.1 years, or 1.25 percent, for females. If these percentage changes are approximately correct for the typical smoker and nonsmoking smoker populations, then our estimates of smoking-related life expectancy effects would need to be adjusted upward accordingly (or perhaps by different percentages because life expectancy has continued to change since 2006).

A further reason to believe the values in table 5 of this document may be underestimates is their lack of quantification of any reduction in either the external effects attributable to passive smoking or the infant and child fatalities caused by mothers smoking during pregnancy. Sloan et al. (Ref. 116) indicate that, historically, the inclusion of spouse and infant deaths from exposure to secondhand smoke or mothers smoking while pregnant increased estimates of smoking’s mortality effects by approximately 26.3 percent. We do not incorporate this adjustment into our analysis, however, because recent restrictions on indoor public smoking and educational campaigns have significantly reduced, though not eliminated, nonsmokers’ exposure to secondhand smoke. In other words, an analysis of the rule’s impact on health benefits that accrue to individuals other than smokers themselves requires three pieces of estimation: (1) The rule-induced change in the number of U.S. smokers, (2) the relationship between the number of smokers and exposure of nonsmoking individuals to the harmful effects of cigarettes, and (3) the effect of cigarette exposure on nonsmokers’ mortality. The ever-changing level of nonsmoker cigarette exposure means that a simple extrapolation from the recent past provides a much less reliable prediction of the near future for element (2) than for other pieces of this analysis. Any estimation of (2) would therefore be highly data-intensive and subject to an unacceptable level of potential error. In general, FDA has been unable to obtain data with which to solve this problem; it is for this reason that we do not quantify health benefits that will accrue to individuals other than smokers themselves.

We do, however, note that the Robert Wood Johnson Foundation (Ref. 137) reports that the percentage of the U.S. population living in homes where smoking was permitted decreased from 56.9 percent in 1992 to 1993 to 20.9 percent in 2006 to 2007. This may indicate that the ratio of spouse and infant mortality effects (related to passive smoking) to smoker mortality effects is now approximately 36.7 (20.9/56.9) percent as large as the 26.3 percent ratio derived from Sloan et al.’s results (which were calculated using data from the 1990s). Using this very rough approximation yields a present value of spouse and infant mortality benefits ranging from $140.3 million ($0.263*$0.367*$1.45 billion) to $2.18 billion ($0.263*0.367*$22.56 billion). Although there are serious weaknesses with this estimation approach that make it inappropriate to include in our overall benefits analysis, the results may give a sense of the magnitude of mortality benefits generated by the rule via reductions in spousal and fetal smoking exposure.

**iii. Improved health status (or reduced morbidity).** In the previous section, we estimated the benefits that will accrue as a result of the rule-induced reduction in premature deaths from cancer, pulmonary and cardiovascular disease, and other smoking-caused illnesses. Cigarette smoking also imposes costs on smokers in the form of pain, distress, and impaired function even before these illnesses cause fatalities. As with premature death, individuals are assumed to be willing to give up valuable resources in order to avoid reductions in quality of life associated with smoking-related illnesses.

Sloan et al. (Ref. 116) examine survey respondents’ self-reported health status (which can be categorized as poor, fair, good, very good, or excellent) and estimate that a 24-year-old smoker can expect, on average, an extra 1.086 discounted years (using a discount rate of 3 percent and averaging over Sloan’s estimates for males and females) or 0.521 discounted years (using a discount rate of 7 percent and again averaging over males and females) of fair or poor health over his or her lifetime, as compared with a nonsmoking smoker.

In order to quantify the value of rule-induced reductions in years spent in fair or poor health, we scale our estimates of the VSLY ($106,308, $212,615, and $318,923, as discussed in the previous section of this document) by a ratio representing the trade-off individuals are willing to make between time spent in best-possible and lesser levels of health. Nyman et al. (Ref. 138) estimate this trade-off by matching survey respondents’ self-reported subjective health statuses with their EuroQol–5D (EQ–5D) health index scores. The EQ–5D survey responses—to questions about five areas of health, including mobility, self-care, pain, anxiety, and ability to perform usual activities—are mapped so that a score of one represents best measurable health, a score of zero represents death, and fractional values represent intermediate levels of health. Nyman et al.’s analysis indicates that, relative to the health index score of an individual with excellent health, a very good health score will be lower by 0.03,
a good health score will be lower by 0.078, a fair health score will be lower by 0.194 and a poor health score will be lower by 0.392. Weighting by Nyman et al.’s reported percentages of respondents in each health category, FDA finds that the health index score for the average individual in good, very good, or excellent health is lower than the index for excellent health by 0.036 and the health index score for the average individual in fair or poor health is lower than the index for excellent health by 0.244; the difference between these averages is 0.208. This result may be interpreted as follows: The harm experienced by an individual whose health changes, for 1 year, from good, very good, or excellent to fair or poor is equal, on average, to the harm experienced by an individual in the best possible health whose death is hastened by 0.208 years. Thus, the welfare effect of smoking-related health status changes may be found by multiplying a plausible life-year value (such as $106,308, $212,615, or $318,923) by 0.208; this multiplication yields estimates of $21,800, $43,600, and $65,400 for the amounts individuals are willing to pay to avoid a year of reduced health status. The U.S. Census Bureau (Ref. 130) predicts that the nation’s 24-year-old cohort will be 2.17 million females and 2.25 million males in 2013 and rise steadily to approximately 2.25 million females and 2.33 million males in 2031. FDA’s estimate of a 0.088 percentage point reduction in the U.S. smoking rate thus translates to a decrease of 3,906 24-year-old smokers in 2013, with the decrease rising to approximately 4,154 in 2037. Multiplying these estimates of the rule-induced reduction in the number of smokers by Sloan et al.’s predictions of discounted reduced health-years per smoker and the quality-of-life loss per year of fair or poor health derived from Nyman et al., and discounting appropriately, yields a rule-induced welfare gain of $0.5 to $4.7 billion. Detailed results appear in table 6 of this document.

<table>
<thead>
<tr>
<th>Value of a Statistical Life-Year = $106,308</th>
<th>Value of a Statistical Life-Year = $212,615</th>
<th>Value of a Statistical Life-Year = $318,923</th>
</tr>
</thead>
<tbody>
<tr>
<td>3% Discount Rate 7% Discount Rate</td>
<td>3% Discount Rate 7% Discount Rate</td>
<td>3% Discount Rate 7% Discount Rate</td>
</tr>
<tr>
<td>1,580.7 500.5</td>
<td>3,161.4 1,001.0</td>
<td>4,742.2 1,501.5</td>
</tr>
</tbody>
</table>

Sloan and coauthors do not report the effect of smoking on fair or poor health years for dissuaded smokers of ages other than 24; in the absence of a reliable estimate of the morbidity effect of smoking cessation for individuals aged 25 and above, FDA takes the conservative approach of estimating benefits only for adults who are at or below that age sometime during the first 20 years of the rule’s implementation. Smoking cessation brought about by this rule will improve health status, in some cases substantially, for many individuals who are over age 24 at the time of the rule’s implementation. Our omission of these benefits to older individuals produces an underestimate of the rule’s morbidity benefits (which is why we describe our estimate as conservative) but there are several reasons to believe the magnitude of the underestimate may not be overwhelmingly large. First, although individuals aged 24 and below make up a fairly small portion of the smokers we estimate will be dissuaded from smoking in 2013, they make up the vast majority of smokers newly dissuaded in years 2014 to 2031 because it is young people and a few immigrants who will be exposed to graphic warning labels for people and a few immigrants who will be exposed to graphic warning labels for individuals aged 25 and above who decrease smoking as a result of the rule requires some assumptions. For this analysis, we assume that smoking-related annual excess medical costs are the same whether smokers are compared with never-smokers or former smokers and that the payments, reported by Sloan et al. as present values for 24-year-olds, are distributed equally from ages 24 to 100 (in other words, we annualize Sloan et al.’s estimated present value over the 77 years between ages 24 and 100). With these assumptions, given FDA’s projected 20-year reductions in smoking prevalence, we anticipate that the regulation will cause smoking-related medical expenditures to fall by $859.9 million, of which $458.2 million will be realized as savings by smokers themselves and $401.7 million by nonsmokers. With a 7-percent discount rate, the total decrease in expenditure becomes $491.3 million, with $261.2 million of those savings accruing to smokers and $230.1 million to nonsmokers. Further details about the nonsmoker portion of expenditures appear in the Distributional Effects portion of this analysis.

In the absence of the rule, some portion of smoking-related medical expenditures accrues to health service providers as economic rent (also known
as producer surplus 14). Any reduction of this portion will not contribute to the social benefit of the rule but will instead be a transfer of resources from health service providers to consumers, public and private insurers, and others. A further complication in the analysis of the market for health is generated because nonsmokers’ payments take the form of a subsidy for smoking-related medical services and thus some portion of their expenditure in the absence of the rule is greater than smokers’ own willingness-to-pay for those medical services. Both for this reason and due to the existence of economic rent, the avoidance of at least some portion of nonsmokers’ smoking-related spending will transfer value from one portion of society to another but not contribute to an overall social benefit of the rule. We do not know the size of this portion relative to nonsmokers’ overall rule-induced expenditure change, so we assume that 50 percent of nonsmokers’ smoking-related spending accrues as a net social benefit of the rule. This produces an overall estimate of rule-induced reductions in medical expenditures of $659.0 million, calculated with a 3-percent discount rate, or $376.3 million, calculated with a 7-percent discount rate.

v. Other financial effects of smoking cessation. In section XI.F.6 of this document, we will discuss in detail the effects of the rule on Social Security, income taxes, private pensions, and life insurance. Summaries of these effects will appear in table 23 of this document. For the most part, we will characterize the values appearing in table 23 as transfers, having equal and offsetting effects on various members of society. There are, however, some additional consequences of these transfers that must be considered in light of the optimal internality tax estimation approach and the related need to estimate gross internal benefits and costs of dissolved smoking. The mixture of positive and negative values in table 23 shows that societal transfers can take the form of both subsidies and additional costs of smoking; when summed together, the positive and negative effects in table 23 show a net smoking subsidy, which individuals relinquish when they avoid initiating or quit smoking.

There is a difficulty in quantifying the effect of the types of transfers appearing in table 23 of this document on internal benefits. Smokers’ experience of these transfers may already be included in the section XI.D.2.b.ii and XI.D.2.b.iii of this document estimates of gross health benefits because the willingness-to-pay measure on which we base our morbidity and mortality calculations includes all the effects a person will likely experience as a result of improving his or her health and extending his or her life. These effects include increased opportunities to collect Social Security and defined benefit pension payments, a decreased chance of leaving survivors enough life insurance to make up for the amount paid in premiums, and increases in pension and income tax payments (due to working longer and receiving higher wages in compensation for higher productivity). If the results in section XI.D.2.b.ii and XI.D.2.b.iii of this document already reflect these phenomena, what is missing from our analysis is not the intrapersonal effect associated with smokers’ experience of table 23 transfers but the direct benefit to the general public of no longer providing a net smoking subsidy; in this case, the total value of the subsidy, or 100 percent of the values in table 23, would need to be added to our net benefits estimate. Because morbidity and mortality are the primary but not the only ways in which smoking affects Social Security, income tax, pension, and life insurance payments and receipts, we do not know the extent to which our morbidity and mortality willingness-to-pay measures capture smokers’ experience of these transfers. We will assume that 50 percent of the midpoint values in table 23 are included in our morbidity and mortality estimates; with this assumption, our estimated net benefits will change in two opposing directions: They will increase by 100 percent of the midpoint values in table 23 (representing the reduced subsidy payment from the general public), but will decrease by an amount equal to 50 percent of the table 23 midpoint values times the net-to-gross benefits ratio (representing the effects on dissuaded smokers that are not included in the morbidity and mortality estimates).

Summing our estimates of rule-induced life-year extensions, health status improvements, medical cost reductions, and financial effects, we find that the present value of health-related and financial benefits accruing to dissuaded smokers totals $9.29 to $27.50 billion (with a 3-percent discount rate) or $2.10 to $6.01 billion (with a 7-percent discount rate). As shown in table 7 of this document, the present value of financial benefits accruing to the general public totals $733.1 million (with a 3-percent discount rate) or $330.3 million (with a 7-percent discount rate).

Table 7.--Financial Benefits Accruing to General Public ($ million)

<table>
<thead>
<tr>
<th>Description</th>
<th>Discount Rate = 3%</th>
<th>Discount Rate = 7%</th>
</tr>
</thead>
<tbody>
<tr>
<td>Smoking-Related Medical Cost Subsidies, Net of Reduced Producer Surplus for Health Care Providers</td>
<td>200.9</td>
<td>115.0</td>
</tr>
<tr>
<td>Social Security Outlays</td>
<td>-649.2</td>
<td>-263.2</td>
</tr>
<tr>
<td>Income Taxes on Social Security-Taxable Earnings</td>
<td>746.5</td>
<td>301.5</td>
</tr>
<tr>
<td>Defined Benefit Private Pension Outlays</td>
<td>-906.8</td>
<td>-366.1</td>
</tr>
<tr>
<td>Life Insurance Outlays</td>
<td>1,341.7</td>
<td>543.1</td>
</tr>
<tr>
<td>Total</td>
<td>733.1</td>
<td>330.3</td>
</tr>
</tbody>
</table>

Note: Positive entries in the table represent transfers of value from individuals dissuaded from smoking to the general public. Negative entries represent transfers in the opposite direction.

vi. Summary of benefits accruing to dissuaded smokers. Table 8 of this document presents benefits estimates that reflect a variety of net-to-gross ratios, ranging, as discussed in Technical Appendix X5, from the 7
percent derived from the work of Gruber and Köszegi to the 90 percent suggested in a public comment. Also presented are the net internal benefits results derived from Warner et al.'s work on the value to smokers of cessation programs. For each discount rate and VSLY, we also report the midpoint between the lower and upper bound benefits estimates, where the upper bound is yielded by the 90 percent net-to-gross benefits ratio and the lower bound by the 7-percent ratio in some cases and by the cessation value approach in others. Given the great variation in estimates of net benefits to dissuaded smokers, we follow the recommendation of OMB Circular A–4 and use the midpoints for our primary calculations in the remainder of this analysis. The resulting midpoints range from $4.37 to $12.56 billion (with a 3-percent discount rate) or $1.02 to $2.86 billion (with a 7-percent discount rate). We emphasize that all the net benefits appearing in table 8 are intrapersonal and thus could not be positive if all tobacco consumers were time-consistent, fully rational, self-controlled, able to resist temptation, and in possession of perfect and complete information; instead, our results are qualitatively consistent with policy implications of economic models in which consumers are characterized by hyperbolic discounting, incorrect forecasting, temptation utility or self-control problems (in addition to Gruber and Köszegi (Ref. 104), see Bernheim and Rangel (Ref. 105) and Gul and Pesendorfer (Ref. 110)) and with Gruber and Mullainathan's (Ref. 182) examination of the effect of cigarette excise taxes on the happiness of individuals with a high propensity to smoke.

3. Reduced Fire Costs

Each year, fires started by lighted tobacco products kill and injure people and destroy structures and other property. In the United States in 2007, civilian deaths caused by smoking-related fires totaled 720, with direct property damage of $530 million (Ref. 141). A reduction in the number of smokers, and the coinciding number of cigarettes smoked, will reduce the number of future fires.

FDA estimates the rule-induced decrease in cigarettes smoked by multiplying together the percentage change in smoking whose calculation was described in section XI.D.1 of this document, the projected population in a given year (Ref. 130) and age-appropriate discounted lifetime cigarette consumption (in packs) per smoker. FDA calculates average consumption for 18- to 23-year-olds using the May 2006, August 2006, and January 2007 Tobacco Use Supplements to the Current Population Survey (Ref. 142). Sloan et al. (Ref. 116) report lifetime discounted consumption for typical 24-year-old smokers. Comparing against total consumption in 2006 (the most recent year for which the FTC (Ref. 143) reports cigarette sales), we find that discounted lifetime cigarette consumption will decrease by an amount equivalent to 3.9 percent (using a 3-percent discount rate) or 2.1 percent (using a 7-percent discount rate) of a present-day annual total as a result of the final rule.

The rule-induced percentage reduction in fires may not equal the percentage reduction in cigarette consumption, however, because all 50 States have passed legislation that requires cigarettes to be self-extinguishing or fire-safe (Ref. 144). FDA acknowledges some uncertainty in the effectiveness rate of fire-safe cigarettes; for this analysis, we present the rule-induced percentage reduction in fires at 90 percent of the projected reduction in cigarettes smoked.

Table 8.—Present Value of Net Internal (i.e., Intrapersonal) Benefits ($ millions)

<table>
<thead>
<tr>
<th></th>
<th>VSLY=$106,308</th>
<th>VSLY=$212,615</th>
<th>VSLY=$318,923</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>3% Discount Rate</td>
<td>7% Discount Rate</td>
<td>3% Discount Rate</td>
</tr>
<tr>
<td>Totals Calculated with Alternative Methods or Net-to-Gross Benefits Ratios:</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>90 Percent Ratio Derived from Public Comment</td>
<td>8,364.3</td>
<td>1,894.2</td>
<td>16,555.7</td>
</tr>
<tr>
<td>24 Percent Ratio Derived from Gruber (Ref. 133)</td>
<td>2,254.7</td>
<td>506.9</td>
<td>4,478.0</td>
</tr>
<tr>
<td>7 Percent Ratio Derived from Gruber and Köszegi (Ref. 104)</td>
<td>624.2</td>
<td>137.7</td>
<td>1,250.7</td>
</tr>
<tr>
<td>Value of Cessation Derived from Warner et al. (Ref. 131)</td>
<td>370.3</td>
<td>322.4</td>
<td>370.3</td>
</tr>
<tr>
<td>Midpoint Between Lower and Upper Bounds</td>
<td>4,367.3</td>
<td>1,016.0</td>
<td>8,463.0</td>
</tr>
<tr>
<td>Allocation of Midpoint Total:</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Life-Years</td>
<td>3,534.2</td>
<td>700.2</td>
<td>6,920.2</td>
</tr>
<tr>
<td>Health Status</td>
<td>742.8</td>
<td>241.6</td>
<td>1,454.5</td>
</tr>
<tr>
<td>Medical Expenditure Reduction</td>
<td>215.3</td>
<td>126.1</td>
<td>210.8</td>
</tr>
<tr>
<td>Other Financial Effects</td>
<td>-125.0</td>
<td>-51.9</td>
<td>-122.4</td>
</tr>
</tbody>
</table>

15 One of the first States to enact these laws, New York, requires cigarettes to self-extinguish 75 percent of the time (Ref. 145). Data from New York show a reduction in smoking-caused fires of about 10.6 percent from the average of the 4 years (2000 to 2003) prior to passage of the fire-safe cigarette law to the first 2 years (2006 to 2007) after implementation was complete (Ref. 146).
estimate that 10.6 percent of apparently rule-induced future fire reductions would have been avoided even without this final rule due to fire-safe cigarette design.

The National Fire Protection Association (Ref. 147) reports the percentages of fire fatalities by age category; along with the CDC’s estimate of average American life expectancy (Ref. 136), these data allow FDA to calculate that the average number of life-years lost by fire victims is approximately 37.3; we project that total discounted life-years saved as a result of the rule will be 317.4 (at a 7-percent discount rate) or 1,198.5 (at a 3-percent discount rate). Using—as in sections XLD.2.b.ii and XLD.2.b.iii of this document—VSLY ranging from $106,308 to $318,923, FDA estimates total rule-induced fire-cost savings of $106.0 to $262.5 million (at a 3-percent discount rate) or $34.1 to $76.5 million (at a 7-percent discount rate); of these totals, $12.9 (7-percent discount rate) or $27.7 million (3-percent discount rate) consisted of averted property damage, with the remainder being the value of life-years saved. These estimated savings may significantly underestimate the final rule’s fire-related benefits because they exclude noncivilian mortality and the value of reduction in fire-caused nonfatal injuries. There will, however, be some double counting between the estimated fire-related mortality benefits and the mortality benefits estimated in section XLD.2.b.ii of this document to the extent that it is smokers themselves who are killed in cigarette-caused fires.

4. Summary of Benefits

The discussion above demonstrates the considerable magnitude of the economic benefits available from smoking reduction efforts. As shown in table 9a of this document, our midpoint benefits estimates range from $5.21 to $13.55 billion (with a 3-percent discount rate) or $1.38 to $3.27 billion (with a 7-percent discount rate). Estimates are presented as annualized values in table 9b of this document, reported over time in Appendix X3, and subjected to Uncertainty Analysis in Technical Appendix X6. Nonquantified benefits include reductions in nonsmoker morbidity and mortality associated with passive smoking and mothers smoking during pregnancy.

<table>
<thead>
<tr>
<th>Table 9a.--Present Value of Benefits ($ mil)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
</tr>
<tr>
<td>VSLY=$106,308</td>
</tr>
<tr>
<td>VSLY=$212,615</td>
</tr>
<tr>
<td>VSLY=$318,923</td>
</tr>
<tr>
<td>3% Discount Rate</td>
</tr>
<tr>
<td>------------------------------------------------</td>
</tr>
<tr>
<td>Totals Calculated with Alternative Methods or Net-to-Gross Benefits Ratios:</td>
</tr>
<tr>
<td>90 Percent Ratio Derived from Public Comment</td>
</tr>
<tr>
<td>24 Percent Ratio Derived from Gruber (Ref. 133)</td>
</tr>
<tr>
<td>7 Percent Ratio Derived from Gruber and Köszegi (Ref. 104)</td>
</tr>
<tr>
<td>Value of Cessation Derived from Warner et al. (Ref. 131)</td>
</tr>
<tr>
<td>Midpoint Between Lower and Upper Bounds</td>
</tr>
<tr>
<td>Allocation of Midpoint Total:</td>
</tr>
<tr>
<td>Life-Years</td>
</tr>
<tr>
<td>Health Status</td>
</tr>
<tr>
<td>Medical Expenditure Reduction</td>
</tr>
<tr>
<td>Other Financial Effects</td>
</tr>
<tr>
<td>Fire Loss</td>
</tr>
</tbody>
</table>
All of the upfront costs of this rule are assumed to occur in the first period of the time horizon of this rule (2012). The cost tables present raw undiscounted calculations of these one-time costs. For summary tables requiring a present value, these costs are discounted 1 year back to the present (2011).

### Table 9b.--Annualized Value of Benefits ($ mil)

<table>
<thead>
<tr>
<th></th>
<th>VSLY=$106,308</th>
<th>VSLY=$212,615</th>
<th>VSLY=$318,923</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>3% Discount Rate</td>
<td>7% Discount Rate</td>
<td>3% Discount Rate</td>
</tr>
<tr>
<td>Totals Calculated with Alternative Methods or Net-to-Gross Benefits Ratios:</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>90-Percent Ratio</td>
<td>569.3</td>
<td>182.0</td>
<td>1,125.2</td>
</tr>
<tr>
<td>Derived from Public</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Comment</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>24-Percent Ratio</td>
<td>158.7</td>
<td>51.1</td>
<td>313.4</td>
</tr>
<tr>
<td>Derived from Gruber</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>(Ref. 133)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>7-Percent Ratio Derived</td>
<td>49.1</td>
<td>16.2</td>
<td>96.5</td>
</tr>
<tr>
<td>from Gruber and Köszegi</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>(Ref. 104)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Value of Cessation</td>
<td>32.0</td>
<td>33.6</td>
<td>37.3</td>
</tr>
<tr>
<td>Derived from Warner et</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>al. (Ref. 131)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Midpoint Between Lower</td>
<td>349.9</td>
<td>130.3</td>
<td>630.5</td>
</tr>
<tr>
<td>and Upper Bounds</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Allocation of Midpoint</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Total</td>
<td>237.6</td>
<td>66.1</td>
<td>465.1</td>
</tr>
<tr>
<td>Life-Years</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Health Status</td>
<td>49.9</td>
<td>22.8</td>
<td>97.8</td>
</tr>
<tr>
<td>Medical Expenditure</td>
<td>28.0</td>
<td>22.8</td>
<td>27.7</td>
</tr>
<tr>
<td>Reduction</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Other Financial Effects</td>
<td>27.4</td>
<td>15.4</td>
<td>27.5</td>
</tr>
<tr>
<td>Fire Loss</td>
<td>7.1</td>
<td>3.2</td>
<td>12.4</td>
</tr>
</tbody>
</table>

### E. Costs

Implementation of this final rule, and the statutory requirements directly linked to it, will create new burdens for cigarette manufacturers. In particular, manufacturers will incur the upfront costs associated with a major labeling change. There will be additional ongoing costs associated with equal and random display of the warnings required in this rule, as mandated by the Tobacco Control Act. Cigarette manufacturers and retailers will be responsible for the removal of noncompliant point-of-sale advertising. Consumers are likely to ultimately bear a share of these costs in the form of increased prices. In addition, the tobacco industry and possibly other sectors will experience lost sales and employment, but these revenue transfers will be offset by gains to other sectors, as discussed in the "Distributional Effects" section of this document.

1. Number of Affected Entities

Labeling and advertising requirements will affect domestic cigarette manufacturers and importers of foreign-made cigarettes. Statistics of U.S. Businesses data show that there were 24 cigarette manufacturing firms in the United States in 2007 (Ref. 148). An undetermined number of importers will also be affected.

Noncompliant point-of-sale advertising will be removed by manufacturers (or importers) and retailers. We use detailed data from the 2002 Economic Census report on product line sales for establishments with payroll to estimate the percentage of various types of retail establishments that sell tobacco products. Searching by the Economic Census product line 20150 (cigars, cigarettes, tobacco, & smokers’ accessories), we find accommodation and food service establishments (NAICS 72) and retail trade establishments (NAICS 44–45) that report tobacco sales (Refs. 149 and 150). Although some establishments in other industries may have unreported sales of tobacco products, the product line sales data provide a reasonable basis to determine which establishments will be affected by the rule.
Because the 2007 Census data on product line sales for retail establishments with employees are not yet available, we update the number of various types of retail establishments using 2007 Statistics of U.S. Businesses data but assume the share of establishments selling tobacco products is unchanged (since 2002) within each category. Likewise, we lack 2007 Census data on product line sales for nonemployer establishments. Without additional information, we assume that, within a NAICS category, the share of establishments selling tobacco products will be the same for nonemployer establishments in 2007 as for establishments with payroll in the 2002 Census. As shown in table 11 of this document, we estimate that about 249,000 retail establishments with payroll and 126,000 nonemployer establishments sell tobacco products.
Rotogravure, the most expensive printing method, is used for cigarette package labels.

2. Costs of Changing Cigarette Labels

We have updated our analysis of the cost of changing cigarette labels based on the availability of improved estimates generated by the new FDA labeling cost model. Unless stated otherwise, our estimates in this analysis come from the new model.

The front and back of every cigarette package must be redesigned to incorporate graphic warnings that will occupy the entire top half, and the current warning will be eliminated. This is classified by the labeling model as a major change. (Any change that affects more than one color or changes the layout enough to require a redesign is major.) In addition, the requirement to incorporate nine different warnings will increase costs beyond what the labeling model estimates. FDA accounted for the additional warnings by first calculating the standard cost of a major change for cigarette labels and then inflating specific cost components expected to increase as a direct result of the requirement for nine warnings.

The FDA labeling cost model incorporates three potential cost components of a labeling change: Label design costs (incurred on a per-UPC basis), inventory costs (incurred on a per-unit basis), and testing costs (incurred on a per-formulation basis). Because the model has a greater focus on analytic testing (e.g., measuring fat grams in a candy bar) than on market testing (which is the aspect of testing applicable to cigarettes), we perform several modifications to the model's testing cost estimation. First, we calculate costs on a per-brand, rather than per-formulation, basis and, second, we restrict the calculation of market testing costs to the largest firms. The large cigarette manufacturers can plausibly be expected to conduct quantitative studies and focus group testing for each of their brands to gauge the effect of the new graphic warnings and to study how they might best be able to mitigate their effects. By contrast, small manufacturers with lower sales revenues are highly unlikely to conduct expensive market testing in response to the new requirements. Further details of our estimation approach will be discussed in section XI.E.4 of this document.

The labeling model estimates that a total of 4,312 cigarette UPCs (3,789 branded and 523 private label) will be affected by this rule. However, it is estimated that label changes for 335 UPCs (8 percent of branded and 6 percent of private label) can be coordinated with previously scheduled, nonregulatory labeling changes.

Coordination of a regulatory change with a nonregulatory change reduces the incremental burden of the regulatory change.

As discussed in the responses to comments, FDA follows its previous labeling cost model (Ref. 152) in assuming 10-percent rush charges under a 15-month compliance period. Using the labeling model cost estimates for uncoordinated changes and incorporating 10-percent rush charges, we estimate that labor costs for label design, including administrative labor costs as well as graphic design and prepress labor costs, are $4,147 to $10,890. Materials costs are estimated to be $6,644 to $10,934; included in this total are both prepress materials and printing plate costs. Recordkeeping costs are estimated to be $55 to $99.

Summing labor, materials, and recordkeeping costs yields a per-UPC label design cost of $10,846 to $21,923. The model estimates that for coordinated labeling changes, there is a per-UPC cost of $340 to $840. This cost is nonzero because there will still be

---

Table 11.--Establishments That Sell Tobacco Products

<table>
<thead>
<tr>
<th>Kind of Business</th>
<th>NAICS</th>
<th>Percentage Selling Tobacco Products</th>
<th>Establishments With Payroll</th>
<th>Nonemployer Establishments</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td></td>
<td>Numberb</td>
<td>Estimated Number Selling Tobacco Products</td>
</tr>
<tr>
<td>General merchandise stores</td>
<td>452</td>
<td>17%</td>
<td>47,456</td>
<td>8,147</td>
</tr>
<tr>
<td>Food &amp; beverage stores</td>
<td>445</td>
<td>55%</td>
<td>122,858</td>
<td>67,037</td>
</tr>
<tr>
<td></td>
<td>excluding 44512</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Convenience stores</td>
<td>44512</td>
<td>85%</td>
<td>28,173</td>
<td>23,986</td>
</tr>
<tr>
<td>Gasoline stations with convenience stores</td>
<td>44711</td>
<td>92%</td>
<td>95,389</td>
<td>87,713</td>
</tr>
<tr>
<td>Health &amp; personal care stores</td>
<td>446</td>
<td>22%</td>
<td>89,406</td>
<td>19,413</td>
</tr>
<tr>
<td>Other retail stores</td>
<td>D</td>
<td>1%</td>
<td>600,537</td>
<td>3,499</td>
</tr>
<tr>
<td>Accommodation and food services</td>
<td>72</td>
<td>2%</td>
<td>358,541</td>
<td>13,991</td>
</tr>
<tr>
<td></td>
<td>excluding 7224</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Drinking places</td>
<td>7224</td>
<td>24%</td>
<td>46,948</td>
<td>11,041</td>
</tr>
<tr>
<td>Tobacco stores</td>
<td>455991</td>
<td>100%</td>
<td>4,558</td>
<td>4,458</td>
</tr>
<tr>
<td>Nonstore retailers</td>
<td>454</td>
<td>2%</td>
<td>42,565</td>
<td>737</td>
</tr>
<tr>
<td></td>
<td>excluding 4542</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Vending machine operators</td>
<td>4542</td>
<td>15%</td>
<td>5,158</td>
<td>777</td>
</tr>
<tr>
<td>Total</td>
<td></td>
<td>15%</td>
<td>1,690,633</td>
<td>249,147</td>
</tr>
</tbody>
</table>

---

a Percentage of establishments with payroll from table 10 of this document.
b Ref. 148
c Ref. 151
d Includes NAICS 441, 443, 444, 448, 451, 453 excluding 453991
e Data on nonemployer establishments unavailable for this NAICS category

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17 Rotogravure, the most expensive printing method, is used for cigarette package labels.
some administrative labor and recordkeeping associated with coordinating a regulatory change with a previously scheduled, nonregulatory change. Total label design costs of this change are thus estimated to be $43 to $87 million.

Manufacturers incur costs if they discard unused label inventory at the end of the compliance period and thus have to print new labels instead of using that inventory. (There is also a small cost associated with disposal.) The labeling model estimates that 767,016 labels will be discarded at the end of the 15-month compliance period, each having a cost of $0.028 to $0.039. The inventory-replacement cost of this labeling change would then be $21,000 to $30,000. Table 12 of this document summarizes the total cost of a standard major labeling change (one warning per UPC), which is estimated to be $43 to $88 million.

Table 12.--Cost of a Standard Major Label Change for Cigarettes

<table>
<thead>
<tr>
<th></th>
<th>Low Cost</th>
<th>Medium Cost</th>
<th>High Cost</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Label Design Costs</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Number of uncoordinated UPCs</td>
<td>3,978</td>
<td>3,978</td>
<td>3,978</td>
</tr>
<tr>
<td>Labor cost ($)</td>
<td>4,147</td>
<td>6,380</td>
<td>10,890</td>
</tr>
<tr>
<td>Materials cost ($)</td>
<td>6,644</td>
<td>6,996</td>
<td>10,934</td>
</tr>
<tr>
<td>Recordkeeping cost ($)</td>
<td>55</td>
<td>88</td>
<td>99</td>
</tr>
<tr>
<td>Per-UPC cost ($)</td>
<td>10,846</td>
<td>13,464</td>
<td>21,923</td>
</tr>
<tr>
<td><strong>Label Design Costs for Uncoordinated UPCs ($)</strong></td>
<td>43,145,388</td>
<td>53,559,792</td>
<td>87,209,694</td>
</tr>
<tr>
<td>Number of coordinated UPCs</td>
<td>335</td>
<td>335</td>
<td>335</td>
</tr>
<tr>
<td>Labor cost ($)</td>
<td>310</td>
<td>550</td>
<td>790</td>
</tr>
<tr>
<td>Materials cost ($)</td>
<td>0</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>Recordkeeping cost ($)</td>
<td>30</td>
<td>40</td>
<td>50</td>
</tr>
<tr>
<td>Per-UPC cost ($)</td>
<td>340</td>
<td>590</td>
<td>840</td>
</tr>
<tr>
<td><strong>Label Design costs for Uncoordinated UPCs ($)</strong></td>
<td>113,900</td>
<td>197,650</td>
<td>281,400</td>
</tr>
<tr>
<td><strong>Total Label Design Costs ($)</strong></td>
<td>43,259,288</td>
<td>53,757,442</td>
<td>87,491,094</td>
</tr>
<tr>
<td><strong>Inventory Costs</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Number of discarded Labels</td>
<td>767,016</td>
<td>767,016</td>
<td>767,016</td>
</tr>
<tr>
<td>Unit cost per discarded label ($)</td>
<td>0.028</td>
<td>0.033</td>
<td>0.039</td>
</tr>
<tr>
<td><strong>Total Inventory Costs ($)</strong></td>
<td>21,093</td>
<td>25,312</td>
<td>29,530</td>
</tr>
<tr>
<td><strong>Total Cost ($)</strong></td>
<td>43,280,381</td>
<td>53,782,754</td>
<td>87,520,624</td>
</tr>
</tbody>
</table>

We expect materials costs for printing plates and prepress activities to be approximately nine times as large as previously calculated for uncoordinated UPCs because of the requirement for nine separate warnings. Each UPC will require nine printing plates, one for each warning label. Additionally, the incremental materials cost of a coordinated label change will be eight times the uncoordinated materials costs, because eight extra printing plates will be needed. We assume that this adjustment accounts for all the one-time costs that arise from the requirement to use nine warnings. Table 13 of this document shows the total costs of the cigarette labeling change, making the adjustment for the nine-warning requirement. The labeling cost range increases to $273 million to $465 million.

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18 Some of the subcomponents of other cost categories might increase due to the nine-warning requirement, but there is far less reason to believe there will be a direct, proportional relationship between those cost categories and the number of warnings. For example, the part of the label that is under the manufacturer’s control only has to be designed once because the same design will be paired with all nine labels. Likewise, the amount of unused inventory discarded is unaffected by the number of warnings used under the new requirements.
3. Ongoing Costs of Equal and Random Display

The Tobacco Control Act calls for equal and random display of the graphic warning images required by this rule. Although the initial design and implementation of a system for equal and random display will be part of the upfront label change, continued operation of such a system in subsequent years will have incremental ongoing administrative and recordkeeping costs. Such a system will be more burdensome than the current system of quarterly rotation of four warnings. FDA assumes that the ongoing yearly administrative labor cost per UPC will be equal to 10 percent of the (non-rush) administrative labor cost of an uncoordinated labeling change, and the yearly recordkeeping cost will be equal to 50 percent of the (non-rush) recordkeeping cost of an uncoordinated labeling change. As shown in table 14 of this document, FDA estimates that, under these assumptions, ongoing annual administrative and recordkeeping costs equal $375,000 to $876,000.


As stated previously, FDA expects that only the large manufacturers will conduct market tests for their brands. Using several State directories of certified tobacco products, FDA estimates that 75 brands are marketed by the 4 largest domestic manufacturers (Refs. 153 through 158). If we assume (as in the labeling model) that 8 percent of changes for these brands are coordinated, then changes for the remaining 69 brands are not coordinated. Including rush charges, the cost of focus group testing is estimated to range from $8,000 to $14,000 per brand, and the cost of a quantitative study is estimated to range from $14,000 to $105,000 per brand. Assuming both types of testing are conducted for 69 brands yields a total cost estimate ranging from $1.5 to $8.2 million with a medium estimate of $2.1 million, as shown in table 15 of this document. We assume that the requirement to use nine
different color graphic-text pairs does not affect these costs.

Table 15.--Cost of Market Testing

<table>
<thead>
<tr>
<th></th>
<th>Low Cost</th>
<th>Medium Cost</th>
<th>High Cost</th>
</tr>
</thead>
<tbody>
<tr>
<td>Number of brands to be tested</td>
<td>69</td>
<td>69</td>
<td>69</td>
</tr>
<tr>
<td>Cost of focus group testing ($)</td>
<td>8,030</td>
<td>11,000</td>
<td>13,970</td>
</tr>
<tr>
<td>Cost of quantitative studies ($)</td>
<td>13,750</td>
<td>19,800</td>
<td>105,160</td>
</tr>
<tr>
<td>Market testing cost per brand ($)</td>
<td>21,780</td>
<td>30,800</td>
<td>119,130</td>
</tr>
<tr>
<td>Total Market Testing Cost ($)</td>
<td>1,502,820</td>
<td>2,125,200</td>
<td>8,219,970</td>
</tr>
</tbody>
</table>

a Undiscounted amount assumed to be incurred in the first period of the time horizon of this rule.

5. Advertising Restrictions: Removal of Noncompliant Point-of-Sale Advertising

The principal effect of the restrictions on advertising in the rule stem from the requirement that retailers and manufacturers of cigarettes remove any point-of-sale advertising for cigarettes that fails to conform to the requirements. In this analysis, we estimate the social resource costs for the removal. In the analysis of FDA’s 1996 final tobacco rule, we based much of our estimate of the cost of removing noncompliant point-of-sale advertising on a report from the Barents Group that used average removal costs for seven types of retail establishments, calculated using in-store surveys conducted by A.T. Kearney, Inc. (61 FR 44396 at 44580). We retain our assumptions from 1996 about the level of effort required to remove point-of-sale advertising. We acknowledge, however, that this approach may overstate or understate the costs for a particular action or type of business.

Table 16 of this document regroups the information from table 11 of this document according to the categories studied by A.T. Kearney. Because our analysis considers only the removal of point-of-sale advertising from physical retail locations, we do not include nonstore establishments. Table 17 of this document shows that, in current dollars, one-time per-establishment costs range from about $12 for “other establishments” to about $198 for convenience stores. To estimate the total costs to comply with the restriction on point-of-sale advertising, we apply the updated per-establishment costs from table 17 to affected establishments. As shown in table 16 of this document, the one-time costs to remove point-of-sale materials will total $45.4 million.

Table 16.--Estimated Number of Establishments Selling Cigarettes Products Affected by the Rule

<table>
<thead>
<tr>
<th>Kind of Business</th>
<th>Establishments With Payroll</th>
<th>Nonemployer Establishments</th>
<th>Total</th>
</tr>
</thead>
<tbody>
<tr>
<td>AT Kearney Category</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>General Merchandise</td>
<td>8,147</td>
<td>5,661</td>
<td>13,808</td>
</tr>
<tr>
<td>Supermarket &amp; Grocery</td>
<td>67,037</td>
<td>56,761</td>
<td>123,799</td>
</tr>
<tr>
<td>Convenience Stores</td>
<td>23,986</td>
<td>23,986</td>
<td>47,972</td>
</tr>
<tr>
<td>Convenience Stores with Gas</td>
<td>87,713</td>
<td>87,713</td>
<td>175,426</td>
</tr>
<tr>
<td>Service Stations</td>
<td>6,347</td>
<td>2,979</td>
<td>9,326</td>
</tr>
<tr>
<td>Drug Stores</td>
<td>19,413</td>
<td>30,138</td>
<td>49,552</td>
</tr>
<tr>
<td>Specialty Tobacco Stores</td>
<td>6,458</td>
<td></td>
<td>6,458</td>
</tr>
<tr>
<td>Other establishments b</td>
<td>28,531</td>
<td>17,391</td>
<td>45,922</td>
</tr>
<tr>
<td>Total</td>
<td>247,633</td>
<td>112,931</td>
<td>360,564</td>
</tr>
</tbody>
</table>

b Source: Table 11 of this document
b Includes miscellaneous retail establishments and accommodations and food services establishments (including drinking places), but excludes nonstore retailers.
6. Government Administration and Enforcement Costs

FDA’s estimated internal costs for administering and enforcing this regulation are uncertain. As a best estimate, however, FDA projects that 25 full-time equivalent employees (FTEs) will be needed to implement the rule. Fully loaded employee costs vary with the type of employee (e.g., field inspectors versus administrative), but an average of $247,049 per FTE places the dollar cost at approximately $6.2 million per year.

An additional cost of the final rule, borne by government but not necessarily FDA, arises due to the required reference to the cessation resource. The rule requires the final graphic warning labels to refer to an already-existing cessation resource. Therefore, only costs associated with additional traffic to that resource are attributable to this final rule. FDA has not quantified these costs.

7. Summary of Costs

Table 19 of this document summarizes the cost estimates from the preceding sections and table 20 of this document displays the present value and annualized value of costs. The tables in Technical Appendix X4 show the undiscounted stream of costs. The range of total costs presented in table 20 of this document is an approximate 90 percent confidence interval and, as such, corresponds to the uncertainty range of benefits presented in table 51 of this document. The distributions of costs and benefits, however, are not correlated; in other words, it may be the case that the actual effects of the rule fall in the high end of the cost range and the low end of the benefits range, or vice versa.
F. Cost-Effectiveness Analysis

We measure the effectiveness of the final rule as the sum of saved life-years and QALYs. In order to assess the cost-effectiveness of the rule, we must adjust the costs to account for effects that are not captured by life-years or QALYs. As shown in detail in the previous section, we calculated the first 20 years’ costs attributable to the rule and found present values of $367.6 to $558.4 million (using a 7-percent discount rate) or $407.3 to $607.4 million (using a 3-percent discount rate). We add to each total the estimated monetary value of lost consumer surplus (as discussed in detail in Technical Appendix X5, this was implicitly netted out of life-years and health improvement benefits estimated calculated in section XI.D.2.b of this document); this yields overall costs of $4.146 to $3.70 billion (using a 7-percent discount rate) or $5.33 to $15.55 billion (using a 3-percent discount rate). In order to focus on the costs associated with extensions of quality-adjusted life (see Ref. 103 at pp. 11–12), we then subtract both medical cost reductions and the value of property savings due to reductions in accidental fires and arrive at a net cost of $0.94 to $3.19 billion (using a 7-percent discount rate) or $4.38 to $14.59 billion (using a 3-percent discount rate). Discounting over the same 20-year time period, we calculate that this rule will lead to 208,535 to 246,137 discounted smoking preventions or cessations. Similarly, we find that 18,534 to 86,326 discounted QALYs will be saved (this includes both fractional life-years associated with reduced morbidity and full life-years associated with reduced premature mortality—both for smokers themselves and for others caught in the path of cigarette-related fires). This yields a cost per smoking prevention of $4,530 to $59,287, and a cost per QALY saved of $50,746 to $172,082.

Braithwaite et al. (Ref. 159) find that preferences in the United States are such that the threshold for cost-effective interventions is somewhere in the range of $109,000 to $297,000 per QALY saved.

Table 19.--Summary of Costs

<table>
<thead>
<tr>
<th>Requirements of the Rule</th>
<th>Annual ($ million)</th>
<th>One-Time ($ million)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Low</td>
<td>Med</td>
</tr>
<tr>
<td>Private Sector</td>
<td>272.5</td>
<td>295.2</td>
</tr>
<tr>
<td>Label Change</td>
<td>1.5</td>
<td>2.1</td>
</tr>
<tr>
<td>Market Testing</td>
<td>45.4</td>
<td>45.4</td>
</tr>
<tr>
<td>Point-of-Sale Advertising</td>
<td>0.4</td>
<td>0.6</td>
</tr>
<tr>
<td>Continuing Admin and Recordkeeping</td>
<td>0.4</td>
<td>0.6</td>
</tr>
<tr>
<td>Subtotal</td>
<td>0.4</td>
<td>0.6</td>
</tr>
<tr>
<td>Government</td>
<td></td>
<td></td>
</tr>
<tr>
<td>FDA</td>
<td>6.2</td>
<td>6.2</td>
</tr>
<tr>
<td>Other (Cessation Resource)</td>
<td>6.2</td>
<td>6.2</td>
</tr>
<tr>
<td>Subtotal</td>
<td>6.2</td>
<td>6.2</td>
</tr>
<tr>
<td>TOTAL</td>
<td>6.6</td>
<td>6.8</td>
</tr>
</tbody>
</table>

a Undiscounted value of one-time costs assumed to be incurred in the first period of the time horizon of this rule.

b Ongoing cost assumed to be incurred in years 2 through 20.
c Annual costs assumed to be incurred in each period for a total of 20 years.

Table 20.--Present Value and Annualized Value of Costs

<table>
<thead>
<tr>
<th>Requirements of the Rule</th>
<th>Present Value ($ million)</th>
<th>Annualized Costs ($ million)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>3 percent</td>
<td>7 percent</td>
</tr>
<tr>
<td></td>
<td>Low</td>
<td>Med</td>
</tr>
<tr>
<td>Private Sector</td>
<td>264.6</td>
<td>286.6</td>
</tr>
<tr>
<td>Label Change</td>
<td>1.5</td>
<td>2.1</td>
</tr>
<tr>
<td>Market Testing</td>
<td>44.1</td>
<td>44.1</td>
</tr>
<tr>
<td>Point-of-Sale Advertising</td>
<td>5.2</td>
<td>9.0</td>
</tr>
<tr>
<td>Continuing Admin and RK</td>
<td>315.4</td>
<td>341.8</td>
</tr>
<tr>
<td>Subtotal</td>
<td>91.9</td>
<td>91.9</td>
</tr>
<tr>
<td>Government</td>
<td>91.9</td>
<td>91.9</td>
</tr>
<tr>
<td>FDA</td>
<td>91.9</td>
<td>91.9</td>
</tr>
<tr>
<td>Other (Cessation Resource)</td>
<td>407.3</td>
<td>433.6</td>
</tr>
<tr>
<td>Subtotal</td>
<td>407.3</td>
<td>433.6</td>
</tr>
</tbody>
</table>

Table 21.--Cost-Effectiveness

<table>
<thead>
<tr>
<th>Cost ($)</th>
<th>3 percent</th>
<th>7 percent</th>
</tr>
</thead>
<tbody>
<tr>
<td>Per Smoking Prevention</td>
<td>179.79</td>
<td>38.243</td>
</tr>
<tr>
<td>Per QALY Saved</td>
<td>50.746</td>
<td>109,040</td>
</tr>
</tbody>
</table>
G. Distributional Effects

This final rule will lead to losses to some segments of U.S. society that will most likely be offset by equal gains to some other segments of society; as such, these effects do not constitute net social costs or benefits and have not yet been discussed in detail in this Analysis of Impacts. In general, sectors affiliated with tobacco and tobacco products will lose sales revenues as a result of this final rule. Simultaneously, nontobacco-related industries will gain sales, because dollars not spent for tobacco products will be spent on other commodities.

1. Tobacco Manufacturers, Distributors, and Growers

FDA estimates that implementation of the regulation may reduce the annual cigarette consumption of U.S. smokers by 30.8 million packs (in 2013) to 40.5 million packs (in 2031). Meanwhile, the FTC (Ref. 143) reports that, in 2006, 17.5 billion cigarette packs were manufactured and distributed to consumers. These numbers imply that tobacco manufacturer revenues will be 0.176 percent lower in the rule’s first year, and 0.231 percent lower in 2031, than they were in 2006. The U.S. Census Bureau (Ref. 160) reports that tobacco manufacturers’ revenues totaled $41.6 billion in 2006; hence, the rule-induced decrease in annual tobacco sales will range from approximately $73.1 to $96.2 million. These estimates would rise somewhat higher if we were accounting for the decrease in price that accompanies the decrease in demand for a good (in this case, cigarettes). Experimental evidence from Mexico (Ref. 101) indicates that graphic warning labels may decrease smokers’ willingness-to-pay for cigarettes by 17 percent; however, without supply elasticity data, we cannot determine how much this decline in willingness-to-pay will change cigarettes’ market price.

We estimate that the tobacco manufacturing, warehousing, and wholesale trade sectors employ about 74,000 full-time workers (Ref. 148). Under the assumption of constant production-to-employment ratio, we project that a 0.176 to 0.231 percent reduction in sales will result in the displacement of 130 to 171 jobs among manufacturers, warehouse workers, and wholesalers.

Effects of the rule will also be observed in the agricultural sector. According to USDA’s 2007 Census of Agriculture (Ref. 161), there are 16,234 tobacco farms. Upon implementation of the rule, these farms may shift some of their acreage from growing tobacco to producing other agricultural products.

2. National and Regional Employment Patterns

Several studies estimate the contribution of tobacco to the U.S. economy or, alternatively, the losses to the U.S. economy that will follow a decline in tobacco-related consumption. Economists have shown both theoretically and empirically that, for the nation as a whole, employment gains from spending on other products will offset any employment losses from reduced spending on tobacco products (Ref. 162). The major tobacco-growing states, however, will experience some adverse economic effects. An economic simulation of the regional impacts of spending on tobacco products carried out in 1994 found that after 8 years, a 2-percent per year fall in tobacco consumption (which substantially exceeds the FDA forecast for the effects of this final rule) would cause the loss of 36,600 jobs for the Southeast Tobacco region of the United States (0.2 percent of regional employment), whereas the nontobacco regions of the United States would gain 56,300 jobs (Ref. 122). That study, if carried out today, would find a much smaller net effect because total employment in tobacco-related industries has fallen. Overall, FDA finds that the income and employment effects associated with the estimated reduction in tobacco consumption will be small.

3. Retail Sector

As will tobacco growers, distributors, and manufacturers, tobacco retailers will be affected by any decrease in cigarette sales. Retailers will, however, be in a position to shift shelf space and promotional activities to nontobacco products, in order to take advantage of the increase in demand for other products that will be expected to accompany the decrease in spending on cigarettes. It is possible that some retailers who rely heavily on cigarette sales may not be able to fully offset their reduction in cigarette sales with sales of other products. Other retailers would then experience some of the gain in sales associated with an increase in demand for other products. This would be a distributional effect within the retail sector.

4. Advertising Industry

The overall impact of the rule on the advertising industry is uncertain. Advertiser revenue may decrease because advertisements with graphic warning labels are less desirable from a cigarette seller’s standpoint and thus tobacco manufacturers will choose to conduct less advertising. On the other hand, advertising industry revenue may increase due to cigarette sellers’ need to redesign advertisements to accommodate new warning labels and to devise new promotional strategies. In either case, few net social costs or benefits will be generated. Moreover, the effect on advertising revenue will likely be relatively small because spending on cigarette advertising has declined substantially in recent years and is now quite small compared with the 1980s and 1990s (Ref. 143). By 2006, expenditures on magazine advertising had fallen to about $50 million and outdoor advertising to under $1 million. Most of the remaining affected advertising expenditures were point-of-sale promotions, which totaled $240 million (Ref. 143).

5. Excise Tax Revenues

In 2009, Federal tobacco tax revenues totaled $16.3 billion, while State and local tobacco tax revenues totaled $16.5 billion (Ref. 143). This rule will decrease government tobacco tax revenues as fewer Americans consume cigarettes. Sales tax revenues generated through tobacco sales will also fall as a result of the rule, but those changes will be much smaller than the changes in excise tax collections and have not been quantified by FDA.

FDA estimates this change in excise tax revenues by multiplying together the percentage change in smoking rate, whose calculation was described in section XI.D.1 of this document; the projected population in a given year (Ref. 130); age-appropriate discounted lifetime cigarette consumption (in packs) per smoker; and current Federal and average State tax rates (Refs. 164 and 165). FDA calculates average consumption for 18- to 23-year-olds using the May 2006, August 2006, and January 2007 Tobacco Use Supplements to the Current Population Survey (Ref. 142). Sloan et al. (Ref. 116) report lifetime discounted consumption for typical 24-year-old smokers.

FDA estimates that average direct annual rule-induced decreases in excise tax collections will be approximately $33.4 million for State governments and $25.7 million for the Federal government. Approximately 25 percent of this reduction may be offset by increased sales of other taxable goods and services (Ref. 166); thus, the annual reductions in tax collections will be $25.1 million for State governments and $19.3 million for the Federal government. Assuming that excise taxes rise in line with average, the rate of inflation allows us to sum these values over the time horizon of our analysis, yielding an...
overall revenue loss to State
governments of $454.9 million (present
value with a 7-percent discount rate) to
$977.5 million (present value with a 3-
percent discount rate) and to the Federal
government of $348.1 million (present
value with a 7-percent discount rate) to
$749.8 million (present value with a 3-
percent discount rate).

Because we cannot know if nominal
cigarette excise taxes actually will
increase at the rate of inflation, we also
calculate these discounted present
values for the case in which tax rates
remain at their current nominal levels.
In this case, the real tax rate will fall at
the rate of inflation, which we forecast
using the difference between interest
rates for standard and inflation-
protected long-term Treasury bills. The
U.S. Department of the Treasury (Ref.
167) reports that, as of February 11,
2011, the composite rate for long-term
standard bills was 4.33 percent, while
the composite rate for long-term
inflation-protected bills was 2.00
percent; the difference yields an
inflation forecast of 2.33 percent per
year. At this rate of inflation, the overall
rule-induced tax revenue loss to State
governments will be $327.8 to $590.0
million and to the Federal government
will be $250.6 to $451.9 million. FDA
emphasizes that these estimates would
be altered, possibly a great deal, either
by future changes in tax rates or
inaccuracy in the inflation forecast.

We note that, leaving aside potential
deadweight loss, there are two principal
effects of tax reductions: Gains to former
payers and losses to former recipients.
Because these transfers exactly offset
each other, there is no net social cost or
benefit associated with the reduction in
cigarette tax collections induced by the
rule.
6. Government-Funded Medical
Services, Insurance Premiums, and
Social Security

Sloan et al. (Ref. 116) estimate that
smokers use more medical services over
their life cycles than do comparable
nonsmokers: in 2000 dollars and
discounted at a 3-percent rate, specific
net costs are $3,757 per female 24-year-
old smoker and $2,617 per male 24-
year-old smoker. Smokers bear a portion
of these net costs themselves, but a
portion equaling $1,726 per female
smoker or $1,245 per male smoker is
borne by nonsmokers through increased
private insurance premiums or taxes
used to fund government health care
programs; hence, a reduction in the U.S.
smoking population will transfer value
from smokers (who receive medical
services paid partially by the general
public) to nonsmokers. If nonsmokers’
payment portions are adjusted for
inflation and distributed over ages 24 to
100 as described in section XI.D.2.b.iv
of this document (“Medical Services”),
given FDA’s projected 20-year
reductions in smoking prevalence, this
transfer totals $401.7 million. With a 7-
percent discount rate, the total becomes
$230.1 million. Sloan et al. indicate that
this reduction will be distributed
unequally across Medicare, Medicaid,
and other insurance types. Details
appear in table 22 of this document.

Sloan et al. (Ref. 116, at p. 255)
estimate the effect of smoking, per male
and female smoker, on net Social
Security, private pension, and life
insurance outlays, as well as on income
tax payments. In the cases of Social
Security and private pension outlays,
smoking-related premature mortality
causes smokers to collect less from the
programs than they contribute during
their lifetimes. Therefore, any rule-
induced reduction in the U.S. smoking
population will shift value from
members of the general public who pay
Social Security taxes and who
contribute to private pension plans to
the individuals who are dissuaded from
smoking by the regulation. A transfer in
the opposite direction—from
individuals dissuaded from smoking by
the regulation to the general public—
will occur in the realms of life insurance
programs and income taxes.

Because Sloan et al. only report
effects for 24-year-olds, we can only
directly calculate these transfer effects
for cohorts who are no older than 24
during the period from 2012 to 2031.
The sum of these effects appears in the
lower bound columns of table 23 of this
document. For the upper bounds, we
assume that effects are the same for
smokers aged 25 and above as they are
for 24-year-olds. In converting Sloan et
al.’s present values, calculated with a 3-
percent discount rate, to present values
calculated with a 7-percent discount
rate, further assumptions are necessary.
We calculate the ratios of 7-percent
present values to 3-percent present
values for all gross benefits categories
(life-years, health status, medical cost
reductions, and fire loss reductions) and
use the lowest and highest ratios for the
lower and upper bounds in table 23.
Finally, we note that we update Sloan
et al.’s estimates using the most recent
annual GDP deflator (Ref. 132).

<table>
<thead>
<tr>
<th>Discount Rate</th>
<th>Medicaid</th>
<th>Medicare Part A</th>
<th>Medicare Part B</th>
<th>Other Government</th>
<th>Private</th>
<th>Uninsured</th>
<th>Total</th>
</tr>
</thead>
<tbody>
<tr>
<td>3%</td>
<td>104.2</td>
<td>-13.1</td>
<td>-174.1</td>
<td>50.4</td>
<td>359.1</td>
<td>75.2</td>
<td>401.7</td>
</tr>
<tr>
<td>7%</td>
<td>50.3</td>
<td>-14.5</td>
<td>-109.1</td>
<td>28.5</td>
<td>231.0</td>
<td>43.9</td>
<td>230.1</td>
</tr>
</tbody>
</table>

Note: Positive entries in the table represent transfers of value from individuals dissuaded from smoking to the general public. Negative entries represent transfers in the opposite direction.
The increase in the proportion of UPCs that can be coordinated is also expected to affect the number of brands that are market tested.

H. International Effects

Of the $87.9 billion worth of tobacco products consumed in the United States in 2009 (Ref. 168), only $156 million consisted of imported cigarettes, with another $897 million imported as tobacco in a less-processed state (Refs. 169 and 170). As in the United States, foreign manufacturers, distributors, and growers of tobacco and tobacco products will lose revenue as a result of the rule, though their loss will be a small fraction of the overall revenue loss. As consumers who would have been smokers purchase other products, there could be a shift in patterns of international trade, depending on where the preferred substitute products are made.

The rule does not apply to cigarettes manufactured for export, whose value totaled $417 million in 2009 (Ref. 169).

I. Regulatory Alternatives

We compare the rule to two hypothetical alternatives: An otherwise identical rule with a 24-month compliance period and an otherwise identical rule with a 6-month compliance period. Even though we estimate costs and benefits for these alternatives, they do not provide viable regulatory options, as they are inconsistent with FDA’s statutory mandate. We also describe alternatives associated with different graphical warnings.

1. 24-Month Compliance Period

Extension of the compliance period to 24 months reduces the one-time costs of this rule through three avenues: The number of UPCs that can be coordinated with a previously scheduled labeling change is increased, rush charges for the label design and market testing costs are eliminated, and discarded inventory costs are eliminated.

Table 24 of this document shows that extending the compliance period to 24 months would reduce the upfront label change cost by $30 to $53 million, to a total of $242 to $411 million. Table 25 of this document shows that market testing costs would be reduced by $0.3 to $1.8 million to a total of $1.2 to $6.4 million.19 Extending the compliance period to 24 months would also delay all costs by about 9 months. We account for this by discounting the present value of costs an extra 9 months in the summary of alternatives table at the end of this section.

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19 The increase in the proportion of UPCs that can be coordinated is also expected to affect the number of brands that are market tested.
Extending the compliance period to 24 months would delay the accrual of health and fire reduction benefits by 9 months. An approximation of the effect of this delay may be found by discounting, at 3- and 7-percent discount rates, the previously calculated total benefits. As shown in table 26 of this document, FDA finds that a 24-month compliance period would decrease the present value of benefits by between $65.4 and $294.6 million.

2. 6-Month Compliance Period

With a 6-month compliance period, the labeling cost model assumes that there is not enough time for any of the labeling changes to be coordinated with previously scheduled changes. Also, FDA accepts the labeling model’s assumption of 40 percent rush charges, rather than assuming 10-percent rush charges as we did with a 15-month compliance period. The labeling model further assumes that 12 months is the shortest compliance period that can be met without resorting to covering up the old labels with stickers as a temporary solution. Therefore, with a 6-month compliance period, the cost of discarded inventory is the same as under a 12-month compliance period, but there is an additional cost for applying appropriate stickers to cover the old package label design.
The model, based on current sales data, estimates the number of units sold annually to be about 8 billion. Therefore, 4 billion units would be relabeled with stickers. The per-unit cost for the sticker and application is between $0.045 and $0.323. Reducing the compliance period to 6 months would then increase label change costs by $258 to $1,430 million to a total of $531 to $1,895 million. It would also increase the market testing costs by $0.6 to $3 million to a total of $2 to $11 million. Finally, shortening the compliance period to 6 months would move all costs up by about 9 months.

We account for this by compounding the present value of costs 9 months in the summary of alternatives table at the end of this section.

Reducing the compliance period to 6 months would hasten the accrual of health and fire reduction benefits by 9 months. An approximation of the effect of this change in timing may be found by compounding, at 3- and 7-percent discount rates, the previously calculated total benefits. As shown in table 29 of this document, FDA finds that a 6-month compliance period would increase benefits by between $68.8 and $301.2 million.

### Table 27.--Cost of a Major Cigarette Label Change With Nine Warning Labels (6-Month)

<table>
<thead>
<tr>
<th>Per-UPC Costs(^a)</th>
<th>Low Cost</th>
<th>Medium Cost</th>
<th>High Cost</th>
</tr>
</thead>
<tbody>
<tr>
<td>Number of uncoordinated UPCs</td>
<td>4,312</td>
<td>4,312</td>
<td>4,312</td>
</tr>
<tr>
<td>Labor cost ($)</td>
<td>5,278</td>
<td>8,120</td>
<td>13,860</td>
</tr>
<tr>
<td>Materials cost ($)</td>
<td>76,104</td>
<td>80,136</td>
<td>125,244</td>
</tr>
<tr>
<td>Recordkeeping cost ($)</td>
<td>70</td>
<td>112</td>
<td>126</td>
</tr>
<tr>
<td>Per-UPC cost ($)</td>
<td>81,452</td>
<td>88,368</td>
<td>139,230</td>
</tr>
<tr>
<td>Per-UPC costs for Uncoordinated UPCs ($)</td>
<td>351,221,024</td>
<td>381,042,816</td>
<td>600,359,760</td>
</tr>
<tr>
<td>Total Per-UPC Costs ($)</td>
<td>351,221,024</td>
<td>381,042,816</td>
<td>600,359,760</td>
</tr>
</tbody>
</table>

### Per-Unit Costs

| Number of discarded labels | 1,087,966 | 1,087,966 | 1,087,966 |
| Unit cost per discarded label ($) | 0.035 | 0.042 | 0.049 |
| Discarded Inventory Cost | 38,079 | 45,695 | 53,310 |
| Sticker and application costs per unit ($) | 0.0448 | 0.115 | 0.3234 |
| Number of units sold in 6 months | 4,002,097,332 | 4,002,097,332 | 4,002,097,332 |
| Sticker cost ($) | 179,293,960 | 459,440,774 | 1,294,278,277 |
| Total Per-Unit Costs | 179,332,039 | 459,486,468 | 1,294,331,588 |
| Total Cost ($) | 530,553,063 | 840,529,284 | 1,894,691,348 |
| Change from 15-month Compliance Period | 258,028,106 | 545,356,547 | 1,429,903,987 |

\(^a\) Undiscounted value of costs assumed to be incurred in the first period of the time horizon of this rule.

### Table 28.--Market Testing Cost With a 6-Month Compliance Period

<table>
<thead>
<tr>
<th>Market Testing Cost(^b)</th>
<th>Low Cost</th>
<th>Medium Cost</th>
<th>High Cost</th>
</tr>
</thead>
<tbody>
<tr>
<td>Number of brands to tested</td>
<td>75</td>
<td>75</td>
<td>75</td>
</tr>
<tr>
<td>Cost of focus group testing ($)</td>
<td>10,220</td>
<td>14,000</td>
<td>17,780</td>
</tr>
<tr>
<td>Cost of quantitative studies ($)</td>
<td>17,500</td>
<td>25,200</td>
<td>133,840</td>
</tr>
<tr>
<td>Market testing cost per brand ($)</td>
<td>27,720</td>
<td>39,200</td>
<td>151,620</td>
</tr>
<tr>
<td>Total Market Testing Cost ($)</td>
<td>2,079,000</td>
<td>2,940,000</td>
<td>11,371,500</td>
</tr>
<tr>
<td>Change from 15-month Compliance Period</td>
<td>576,180</td>
<td>814,800</td>
<td>3,151,530</td>
</tr>
</tbody>
</table>

\(^b\) Undiscounted value of costs assumed to be incurred in the first period of the time horizon of this rule.
3. Alternative Graphic Images

A legally available alternative to this rule would be to select a different set of graphic images. Although we are unable to quantify the effects of different graphic images, we note that some images may have a larger impact on smoking rates than other images. Another alternative suggested would be to use more than nine graphic images to accompany the nine statutory warnings. We cannot assess the effect of additional images on the benefits of the rule but more images would increase costs. Although not all costs rise in proportion to the number of graphic images, the materials cost, which is the largest cost component, would rise in proportion to the number of images.

4. Summary of Regulatory Alternatives

Table 30 of this document summarizes the regulatory alternatives related to the compliance period by displaying ranges for the present values of the total benefits and total costs. Estimated ranges for the cost ratios (per smoking prevention and per life-year saved) of the rule and its regulatory alternatives appear in table 31 of this document.

<table>
<thead>
<tr>
<th>Compliance Period</th>
<th>Present Value of Total Benefits ($ million)</th>
<th>Present Value of Total Costs ($ million)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>3%</td>
<td>7%</td>
</tr>
<tr>
<td>24-Month Total</td>
<td>5,092.2 to 13,257.1</td>
<td>1,312.0 to 3,109.2</td>
</tr>
<tr>
<td>(Final Rule) 15-Month Total</td>
<td>5,206.4 to 13,554.3</td>
<td>1,380.3 to 3,271.0</td>
</tr>
<tr>
<td>6-Month Total</td>
<td>5,323.1 to 13,858.1</td>
<td>1,452.2 to 3,441.3</td>
</tr>
</tbody>
</table>

- Range in benefits is based on a VSLY of $106,308 to $318,923.
- Range in costs is based on low cost and high cost values.

<table>
<thead>
<tr>
<th>Discount Rate = 3 percent</th>
<th>Discount Rate = 7 percent</th>
</tr>
</thead>
<tbody>
<tr>
<td>Low</td>
<td>Incremental CE*</td>
</tr>
<tr>
<td>24-Month Compliance:</td>
<td></td>
</tr>
<tr>
<td>Per Smoking Prevention</td>
<td>$17,677</td>
</tr>
<tr>
<td>Per QALY Saved</td>
<td>$50,401</td>
</tr>
<tr>
<td>15-Month Compliance:</td>
<td></td>
</tr>
<tr>
<td>Per Smoking Prevention</td>
<td>$17,798</td>
</tr>
<tr>
<td>Per QALY Saved</td>
<td>$50,746</td>
</tr>
<tr>
<td>6-Month Compliance:</td>
<td></td>
</tr>
<tr>
<td>Per Smoking Prevention</td>
<td>$18,818</td>
</tr>
<tr>
<td>Per QALY Saved</td>
<td>$53,655</td>
</tr>
</tbody>
</table>

* As the compliance period decreases, the number of rule-induced smoking preventions and life-years saved increases. Hence, the incremental costs of 15-Month Compliance are calculated relative to 24-Month Compliance, and the incremental costs of 6-Month Compliance are calculated relative to 15-Month Compliance.
J. Impact on Small Entities

The Regulatory Flexibility Act requires agencies to prepare a final regulatory flexibility analysis if a final rule will have a significant effect on a substantial number of small entities. We expect this rule to have a significant effect on a substantial number of small entities. Consequently, this analysis, together with other relevant sections of this document, serves as the Final Regulatory Flexibility Analysis, as required under the Regulatory Flexibility Act.

1. Description and Number of Affected Small Entities

The final rule will affect small entities in several industries, from tobacco farming to the retail industry. Most of the Nation’s 16,234 tobacco farms are small; between 90.7 and 95.8 percent (between 14,732 and 15,555) of the farms growing tobacco in 2007 had total farm sales under the U.S. Small Business Administration (SBA) small business size standard of $750,000 (Refs. 161 and 171).

Statistics of U.S. Businesses data show that 1,067 of 1,159 tobacco wholesale trade firms (92 percent) employ fewer than the 100-employee threshold that constitutes a small business according to the SBA (Refs. 148 and 171). If the size distribution of cigarette importers is similar to that of all tobacco wholesale trade firms, then 92 percent of them will be affected small businesses.

Also likely to be affected by the regulation are small retail and service entities that sell cigarettes. Retail establishments bear shared responsibility with manufacturers for the cost of removing noncompliant advertising. SBA size standards for the retail trade and the accommodations and food services industries differ from size categories used by the U.S. Census. Table 33 of this document shows the 2002 Census size categories that most closely match the SBA size standards. In all cases, the closest Census size category is smaller than the SBA size standard. As a consequence, any estimate based on the Census size categories may underestimate the number of affected small entities.

### Table 32.--Cigarette Manufacturers by Number of Employees

<table>
<thead>
<tr>
<th>Size by Number of Employees</th>
<th>Number of Firms</th>
</tr>
</thead>
<tbody>
<tr>
<td>Less than 20</td>
<td>9</td>
</tr>
<tr>
<td>20 to 99</td>
<td>7</td>
</tr>
<tr>
<td>100 to 499</td>
<td>4</td>
</tr>
<tr>
<td>Source: Ref. 171</td>
<td></td>
</tr>
<tr>
<td>SBA size standard: 1,000 employees</td>
<td></td>
</tr>
</tbody>
</table>

### Table 33.--SBA Size Standards and Census Size Categories for Retail and Service Firms in NAICS Categories With Tobacco Product Line Sales

<table>
<thead>
<tr>
<th>NAICS with Tobacco Product Line Sales</th>
<th>Description of NAICS Category</th>
<th>SBA Size Standard ($ million)</th>
<th>Census Size Category ($ million)</th>
</tr>
</thead>
<tbody>
<tr>
<td>General Merchandise</td>
<td>Other General Merchandise</td>
<td>11</td>
<td>10</td>
</tr>
<tr>
<td>452990</td>
<td>Department, Discount Department, Warehouse Clubs and Superstores</td>
<td>27</td>
<td>25</td>
</tr>
<tr>
<td>452102</td>
<td>Other Food and Beverage Stores</td>
<td>7</td>
<td>5</td>
</tr>
<tr>
<td>445110</td>
<td>Supermarkets and Grocery</td>
<td>27</td>
<td>25</td>
</tr>
<tr>
<td>44502</td>
<td>Convenience Stores</td>
<td>27</td>
<td>25</td>
</tr>
<tr>
<td>445110</td>
<td>Convenience Stores with Gas</td>
<td>27</td>
<td>25</td>
</tr>
<tr>
<td>445190</td>
<td>Service Stations</td>
<td>9</td>
<td>5</td>
</tr>
<tr>
<td>446</td>
<td>Health and Personal Care Stores</td>
<td>7</td>
<td>5</td>
</tr>
<tr>
<td>453991</td>
<td>Specialty Tobacco Stores</td>
<td>7</td>
<td>5</td>
</tr>
<tr>
<td>Other Kinds of Business</td>
<td>Varies</td>
<td>Varies</td>
<td>Varies</td>
</tr>
</tbody>
</table>

Source: Refs. 171 through 173.

1. Includes only firms with payroll.

2. Includes NAICS 4413, 443112, 444, 448, 451, 4532, 453998, 72 (excluding 72231), 722310.
The Census reports establishment numbers for business by product line, and establishment and firm size by type of business, but provides no size data by type of business and product line. To estimate the number of affected entities that SBA classifies as small, we begin by counting the number of firms that fall below the Census size standard shown in table 33 of this document, including only firms in NAICS categories with tobacco product line sales. Next, we calculate the percentage of small firms in each NAICS category. Depending on the category of business, the percentage of small firms ranges from 41 percent for Discount Department, Warehouse Clubs and Superstores to almost 100 percent for Convenience Stores.

Table 34.--Estimated Percentage of Small Retail and Service Firms in NAICS Categories With Tobacco Product Line Sales

<table>
<thead>
<tr>
<th>NAICS</th>
<th>Description of NAICS Category</th>
<th>Number of Firms</th>
<th>Number of Firms Below Census Size Standard</th>
<th>Percentage of Small Firms (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>452110</td>
<td>Discount Department, Warehouse Clubs and Superstores</td>
<td>88</td>
<td>36</td>
<td>40.9</td>
</tr>
<tr>
<td>452910</td>
<td>Other General Merchandise</td>
<td>7,451</td>
<td>7,320</td>
<td>98.2</td>
</tr>
<tr>
<td></td>
<td>General Merchandise Subtotal</td>
<td>7,539</td>
<td>7,356</td>
<td>97.6</td>
</tr>
<tr>
<td>445110</td>
<td>Supermarkets &amp; Grocery</td>
<td>34,017</td>
<td>33,328</td>
<td>98.0</td>
</tr>
<tr>
<td>4452 and 4453</td>
<td>Other Food and Beverage Stores</td>
<td>34,807</td>
<td>34,082</td>
<td>97.9</td>
</tr>
<tr>
<td></td>
<td>Supermarket &amp; Grocery Subtotal</td>
<td>68,824</td>
<td>67,410</td>
<td>97.9</td>
</tr>
<tr>
<td>445120</td>
<td>Convenience Stores</td>
<td>18,705</td>
<td>18,676</td>
<td>99.8</td>
</tr>
<tr>
<td>447110</td>
<td>Convenience Stores with Gas</td>
<td>37,437</td>
<td>36,848</td>
<td>98.4</td>
</tr>
<tr>
<td>447190</td>
<td>Service Stations</td>
<td>19,822</td>
<td>18,103</td>
<td>91.3</td>
</tr>
<tr>
<td>4461</td>
<td>Drug Stores</td>
<td>36,198</td>
<td>33,894</td>
<td>93.6</td>
</tr>
<tr>
<td>453991</td>
<td>Tobacco Stores</td>
<td>3,238</td>
<td>3,017</td>
<td>93.2</td>
</tr>
<tr>
<td></td>
<td>Other Kinds of Business</td>
<td>589,400</td>
<td>572,619</td>
<td>97.2</td>
</tr>
</tbody>
</table>

Source: Refs. 172, 173, 149, and 150.

* Includes only firms with payroll.

Finally, we apply the percentages in table 34 of this document to our current estimate of the number of affected establishments with payroll (table 16 of this document). This approach implicitly assumes that small establishments are similar whether or not they sell tobacco products. In addition, we classify all nonemployer establishments as small. In total, we estimate that about 355,000 small retail and service establishments will be affected by the rule. This number represents about 98 percent of the estimated 361,000 establishments selling tobacco products.

Table 35.--Estimated Number of Small Establishments With Tobacco Product Line Sales By Kind of Business

<table>
<thead>
<tr>
<th>Kind of Business</th>
<th>Percentage of Small (%)</th>
<th>Number with Payroll</th>
<th>Small with Payroll</th>
<th>Non-employers</th>
<th>Estimated Total Number of Small Establishments</th>
</tr>
</thead>
<tbody>
<tr>
<td>General Merchandise</td>
<td>97.6</td>
<td>8,147</td>
<td>7,949</td>
<td>5,661</td>
<td>13,611</td>
</tr>
<tr>
<td>Supermarket &amp; Grocery</td>
<td>98.0</td>
<td>67,037</td>
<td>65,679</td>
<td>56,761</td>
<td>122,441</td>
</tr>
<tr>
<td>Convenience Stores</td>
<td>99.8</td>
<td>23,986</td>
<td>23,949</td>
<td>0</td>
<td>23,949</td>
</tr>
<tr>
<td>Convenience Stores with Gas</td>
<td>98.4</td>
<td>87,713</td>
<td>86,333</td>
<td>0</td>
<td>86,333</td>
</tr>
<tr>
<td>Service Stations</td>
<td>91.3</td>
<td>6,347</td>
<td>5,797</td>
<td>2,979</td>
<td>8,775</td>
</tr>
<tr>
<td>Drug Stores</td>
<td>93.6</td>
<td>19,413</td>
<td>18,178</td>
<td>30,138</td>
<td>48,316</td>
</tr>
<tr>
<td>Specialty Tobacco Stores</td>
<td>93.2</td>
<td>6,458</td>
<td>6,017</td>
<td>0</td>
<td>6,017</td>
</tr>
<tr>
<td>Other Establishments</td>
<td>97.2</td>
<td>28,531</td>
<td>27,719</td>
<td>17,391</td>
<td>45,110</td>
</tr>
<tr>
<td>Total</td>
<td></td>
<td>247,633</td>
<td>241,621</td>
<td>112,931</td>
<td>354,552</td>
</tr>
</tbody>
</table>

* From table 34 of this document.

* From table 16 of this document.
2. Description of the Potential Impacts of the Final Rule on Small Entities

   a. Effect on manufacturers. In order to estimate how much of the label change and rotation costs will be incurred by small domestic cigarette manufacturers, FDA subtracts from the total costs those costs estimated to be incurred by large domestic manufacturers and foreign manufacturers. Scanner data from AC Nielsen indicate that approximately 49 percent of UPCs can be readily identified as belonging to a brand marketed by one of the four largest cigarette firms by volume (Refs. 153 through 158). Because the costs of label changes are roughly proportional to the number of UPCs, FDA then attributes 49 percent of the total label design and inventory costs to the four firms employing at least 500 people. FDA attributes an additional 3 percent of the label change costs to foreign manufacturers. 20 These adjustments leave 48 percent of costs, or $131 to $223 million in upfront costs and $180,000 to $420,000 in ongoing costs, to be incurred by the 20 small manufacturers. Assuming costs are distributed equally among these firms implies one-time costs of $6.5 to $11.2 million and ongoing costs of $9,000 to $21,000 per firm. Table 36 of this document compares these estimated compliance costs to average annual receipts in order to gauge the potential impact of labeling change requirements on small cigarette manufacturing firms. Because the number of UPCs is probably larger for larger firms, costs are likely greater for larger firms than for smaller firms; if so, this method overstates the impact on the smallest firms and understates the impact on the largest firms (within the category of firms employing fewer than 500 people).

<table>
<thead>
<tr>
<th>Size by Number of Employees</th>
<th>Number of Firms</th>
<th>Average Annual Receipts ($)</th>
<th>Average Compliance Costs ($)</th>
<th>Average Compliance Costs as a % of Average Annual Receipts</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Size</td>
<td>Lower Bound</td>
<td>Upper Bound</td>
<td>Lower Bound</td>
</tr>
<tr>
<td>Less than 20</td>
<td>9</td>
<td>11,195,000</td>
<td>6,541,000</td>
<td>11,155,000</td>
</tr>
<tr>
<td>20 to 99</td>
<td>7</td>
<td>21,265,000</td>
<td>6,541,000</td>
<td>11,155,000</td>
</tr>
<tr>
<td>100 to 499</td>
<td>4</td>
<td>147,896,000</td>
<td>6,541,000</td>
<td>11,155,000</td>
</tr>
</tbody>
</table>

   b. Effect on retailers. As shown in table 37 of this document, retail trade businesses account for almost all sales of tobacco products (Refs. 149 and 150). About 90 percent of tobacco product line sales occur at gasoline stations, food and beverage stores, general merchandise stores, or tobacco stores. Convenience stores (with gasoline stations and stand-alone convenience stores) account for about half of all tobacco product line sales.

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20 In 2006, 9.9 billion out of 345.3 billion individual cigarettes sold were imported (Ref. 123).

FDA assumes the same proportion holds for UPCs. These UPCs should not overlap with those produced by the four largest domestic producers.
To illustrate the effects of the rule on a typical small retail store, we look at one-time costs for a convenience store and a convenience store with gasoline. We select these businesses because, as illustrated in table 37 of this document, sales of tobacco products in these stores account for about 50 percent of all tobacco sales. In addition, tobacco products are an important part of overall revenue for these stores, composing over 12 percent of total sales (as shown in table 38 of this document).
For both types of convenience stores, table 39 of this document shows that for the smallest firms with less than $250,000 in annual sales, the one-time costs of the rule will equal less than 2 percent of annual average sales of tobacco products. Furthermore, one-time costs total less than 0.1 percent of annual average sales of tobacco products for stores with $1 million or more in average annual sales. Although the impact on other small retail and service entities is uncertain, this example suggests that the rule will be unlikely to create a significant direct burden on small retail stores or service establishments.

If individual small retailers are unable to fully offset reduced cigarette sales with increased sales of other items, their sales revenue may fall. Although this decline would not be a social cost (as discussed in the distributional effects section) it would be a cost to the retailers who experience it. FDA has not quantified this additional potential effect, but believes that it is minor because the overall reduction in cigarette consumption is predicted to be less than one half of a percent, the demand for other goods is expected to increase, and retailers can be expected to shift shelf space to the other goods for which demand increases.

Table 38.--The Importance of Tobacco Sales by Kind of Business: Ranked by the Percentage of Total Sales From Tobacco Product Line

<table>
<thead>
<tr>
<th>Kind of Business</th>
<th>Sales From Tobacco Product Line&lt;sup&gt;a&lt;/sup&gt; ($ billion)</th>
<th>Total Sales From All Product Lines&lt;sup&gt;b&lt;/sup&gt; ($ billion)</th>
<th>Percentage of Total Sales From Tobacco Product Line (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Tobacco Stores</td>
<td>5.7</td>
<td>6.5</td>
<td>86.9</td>
</tr>
<tr>
<td>Convenience Stores</td>
<td>4.5</td>
<td>18.1</td>
<td>25.0</td>
</tr>
<tr>
<td>Nonstore Retailers</td>
<td>0.5</td>
<td>2.4</td>
<td>20.3</td>
</tr>
<tr>
<td>Convenience Stores with Gas</td>
<td>21.2</td>
<td>173.4</td>
<td>12.2</td>
</tr>
<tr>
<td>Vending Machine Operators</td>
<td>0.2</td>
<td>1.7</td>
<td>11.2</td>
</tr>
<tr>
<td>Miscellaneous store retailers</td>
<td>0.1</td>
<td>1.2</td>
<td>11.2</td>
</tr>
<tr>
<td>Liquor Stores</td>
<td>1.2</td>
<td>12.8</td>
<td>9.7</td>
</tr>
<tr>
<td>Other Kinds of Business</td>
<td>0.1</td>
<td>1.4</td>
<td>6.5</td>
</tr>
<tr>
<td>Drinking places</td>
<td>0.1</td>
<td>3.9</td>
<td>3.5</td>
</tr>
<tr>
<td>Gasoline Stations</td>
<td>1.0</td>
<td>29.4</td>
<td>3.5</td>
</tr>
<tr>
<td>General Merchandise</td>
<td>7.1</td>
<td>246.1</td>
<td>2.9</td>
</tr>
<tr>
<td>Supermarket &amp; Grocery</td>
<td>7.7</td>
<td>383.5</td>
<td>2.0</td>
</tr>
<tr>
<td>Drug Stores</td>
<td>1.5</td>
<td>80.0</td>
<td>1.9</td>
</tr>
<tr>
<td>Other Accommodation &amp; Foodservice</td>
<td>0.3</td>
<td>33.3</td>
<td>0.8</td>
</tr>
<tr>
<td>Total</td>
<td>51.2</td>
<td>993.9</td>
<td>5.2</td>
</tr>
</tbody>
</table>

<sup>a</sup> Tobacco sales from table 37 of this document.

<sup>b</sup> Includes total sales for firms with tobacco product line sales (Refs. 149 and 150).
3. Alternatives To Minimize the Burden on Small Entities

   a. Increase the compliance period to 24 months for small manufacturers or all manufacturers. Allowing all manufacturers, or only small manufacturers, 24 months to comply with the label changes would eliminate overtime and rush charges, eliminate costs for replacing discarded inventory, and increase the number of UPCs for which the addition of graphic warning labels could be coordinated with previously scheduled label changes. Under a 24-month compliance period, the one-time label change costs would fall by an average of $0.7 to $1.3 million per small firm. Table 40 of this document compares the reduced estimated compliance costs to average annual receipts in order to gauge the potential impact of this regulatory alternative on cigarette manufacturing firms employing fewer than 500 people. As a comparison with table 36 of this document shows, this option would provide some relief, but the burden would remain significant. It would also delay the public health benefits of the rule and be inconsistent with FDA’s statutory mandate.

<table>
<thead>
<tr>
<th>Sales Size of Firm</th>
<th>Number of Establishments</th>
<th>Sales</th>
<th>Sales of Tobacco Products</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td></td>
<td>($ million)</td>
</tr>
<tr>
<td>Convenience Store-NAICS 445120 ¹</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Less than $250,000</td>
<td>4,231</td>
<td>653</td>
<td>0.0</td>
</tr>
<tr>
<td>$250,000 to $499,999</td>
<td>5,296</td>
<td>1,920</td>
<td>0.1</td>
</tr>
<tr>
<td>$500,000 to $999,999</td>
<td>5,150</td>
<td>3,646</td>
<td>0.2</td>
</tr>
<tr>
<td>$1,000,000 to $2,499,999</td>
<td>3,586</td>
<td>4,915</td>
<td>0.3</td>
</tr>
<tr>
<td>$2,500,000 to $4,999,999</td>
<td>659</td>
<td>1,601</td>
<td>0.6</td>
</tr>
<tr>
<td>5,000,000 to 9,999,999</td>
<td>324</td>
<td>712</td>
<td>0.5</td>
</tr>
<tr>
<td>10,000,000 to 24,999,999</td>
<td>215</td>
<td>440</td>
<td>0.5</td>
</tr>
<tr>
<td>Convenience Stores with Gasoline-</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>NAICS 447110 ²</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Less than $250,000</td>
<td>2,246</td>
<td>343</td>
<td>0.0</td>
</tr>
<tr>
<td>$250,000 to $499,999</td>
<td>3,801</td>
<td>1,425</td>
<td>0.0</td>
</tr>
<tr>
<td>$500,000 to $999,999</td>
<td>7,667</td>
<td>5,624</td>
<td>0.1</td>
</tr>
<tr>
<td>$1,000,000 to $2,499,999</td>
<td>14,309</td>
<td>22,303</td>
<td>0.2</td>
</tr>
<tr>
<td>$2,500,000 to $4,999,999</td>
<td>7,977</td>
<td>22,786</td>
<td>0.3</td>
</tr>
</tbody>
</table>

Source: Ref. 167.

¹ Tobacco product line sales account for 25.0 percent of sales for all firms in NAICS 445120 (see Table 38 of this document); one-time costs equal $198.16 (see Table 17 of this document).

² Tobacco product line sales account for 12.2 percent of sales for all firms in NAICS 447110 (see Table 38); one-time costs equal $193.42 (see Table 17).

b. Allow small manufacturers to use one warning per UPC. Allowing small cigarette manufacturers to use only one randomly selected warning and graphic image per UPC would reduce their upfront label change cost substantially. The costs to small businesses of implementing this option can be approximated by assuming that the 20
The average cost per small manufacturer would be reduced by $5.5 to $9 million per firm. Additionally, there would be some small cost at the beginning to ensure random selection of the warnings, but the ongoing annual rotation cost of $9,000 to $21,000 per firm would be eliminated. Table 41 of this document compares the reduced estimated compliance costs to average annual receipts in order to gauge the potential impact of this regulatory alternative on cigarette manufacturing firms employing fewer than 500 people. As a comparison with table 36 of this document shows, this alternative would provide significant relief. However, it is inconsistent with FDA’s statutory mandate. Smokers who use only one specific product would not be exposed to all the warnings, which would likely hinder the effectiveness of this rule.

Table 41.—Potential Impact of Compliance Costs on the 20 Small Cigarette Manufacturers With One Label per UPC

<table>
<thead>
<tr>
<th>Size by Number of Employees</th>
<th>Number of Firms</th>
<th>Average Annual Receipts ($)</th>
<th>Average Compliance Costs ($)</th>
<th>Average Compliance Costs as a Percentage of Average Annual Receipts</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Lower Bound</td>
<td>Upper Bound</td>
<td>Lower Bound</td>
<td>Upper Bound</td>
</tr>
<tr>
<td>Panel 1: Upfront Label Change Costs</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Less than 20</td>
<td>9</td>
<td>11,195,000</td>
<td>1,039,000</td>
<td>2,100,000</td>
</tr>
<tr>
<td>20 to 99</td>
<td>7</td>
<td>21,265,000</td>
<td>1,039,000</td>
<td>2,100,000</td>
</tr>
<tr>
<td>100 to 499</td>
<td>4</td>
<td>147,896,000</td>
<td>1,039,000</td>
<td>2,100,000</td>
</tr>
</tbody>
</table>

Source: Statistics of U.S. Businesses, 2007 (Ref. 148)  
SBA size standard: 1,000 employees

c. Exempt small manufacturers from the labeling change requirements. Exempting small manufacturers from the label change requirements would eliminate their label change costs and ongoing rotation costs (an average reduction of $6.5 to $11.2 million in upfront costs and $9,000 to $21,000 in ongoing costs), thus providing maximum relief. The combined market share of the four largest manufacturers was 89.7 percent in 2008 (Ref. 123). The immediate impact of exempting small manufacturers would therefore be to allow 10.3 percent of cigarettes to be marketed without graphic warning labels. This proportion would grow over time, however, as some consumers would be expected to switch to brands marketed without graphic warnings. This approach would be inconsistent with both FDA’s statutory mandate and the public health objectives of this rule.

d. Exempt small cigarette retailers from the point-of-sale advertising requirements. Exempting small cigarette retailers from the point-of-sale advertising requirements would eliminate their need to remove noncompliant advertising, reducing their direct costs to zero. However, table 35 of this document shows that the overwhelming majority of retail establishments selling cigarettes are small. Although the few establishments operated by large firms might be expected to have higher volume, a significant proportion of consumers would continue to be exposed to advertising lacking the new graphic warnings. This situation would be inconsistent with the public health objective of the rule as well as FDA’s statutory mandate.

XII. Paperwork Reduction Act of 1995

The required warning disclosures are the “public disclosure of information originally supplied by the Federal government to the recipient for that purpose,” and are, therefore, not within the scope of the Paperwork Reduction Act (see 5 CFR 1320.3(c)(2)).

XIII. References

10. Breslau, N., and Peterson, E., “Smoking Cessation in Young Adults: Age at Initiation of Cigarette Smoking and Other


81. Li, J., and Grigg, M., “New Zealand: New Graphic Warnings Encourage Registrations with the Quitline,” Tobacco Control, 18(1); 72, 2009.


§ 1141.1 Scope.

(a) This part sets forth the requirements for the display of health warnings on cigarette packages and in advertisements for cigarettes. FDA may require additional statements to be displayed on packages and in advertisements under the Federal Food, Drug, and Cosmetic Act or other authorities.

(b) The requirements of this part do not apply to manufacturers or distributors of cigarettes that do not manufacture, package, or import cigarettes for sale or distribution within the United States.

(c) A cigarette retailer shall not be considered in violation of this part as it applies to the display of health warnings on a cigarette package if the package:

(1) Contains a health warning;

(2) Is supplied to the retailer by a license- or permit-holding tobacco product manufacturer, importer, or distributor; and

(3) Is not altered by the retailer in a way that is material to the requirements of section 4(a) of the Federal Cigarette Labeling and Advertising Act (15 U.S.C. 1333(a)) or this part, including by obscuring the warning, by reducing its size, by severing it in whole or in part, or by otherwise changing it in a material way.

(d) A cigarette retailer shall not be considered in violation of this part as it applies to the display of health warnings in an advertisement for cigarettes if the advertisement is not created by or on behalf of the retailer and the retailer is not otherwise responsible for the inclusion of the required warnings. This paragraph shall not relieve a retailer of liability if the retailer displays, in a location open to the public, an advertisement that does not contain a health warning or that contains a warning that has been altered by the retailer in a way that is material to the requirements of section 4(b) of the Federal Cigarette Labeling and Advertising Act (15 U.S.C. 1333(b)), this part, or section 4(c) of the Federal Cigarette Labeling and Advertising Act (15 U.S.C. 1333(c)), including by obscuring the warning, by reducing its size, by severing it in whole or in part, or by otherwise changing it in a material way.

§ 1141.10 Required warnings.

(a) The requirements of this part do not apply to manufacturers or distributors of cigarettes that do not

(b) Manufacture, package, or import cigarettes for sale or distribution within the United States.

(d) Manufacturer means any person who imports any cigarette that is intended for sale or distribution to consumers in the United States.

(e) Importer means any person who imports any cigarette that is intended for sale or distribution to consumers in the United States.

(f) Person means an individual, partnership, corporation, or any other business or legal entity.

(g) Required warning means the combination of one of the textual warning statements and its accompanying color graphic, which are set forth in “Cigarette Required Warnings,” which is incorporated by reference at § 1141.12.

(h) Retailer means any person who sells cigarettes to individuals for personal consumption, or who operates a facility where vending machines or self-service displays of cigarettes are permitted.

(i) United States, when used in a geographical sense, includes the several States, the District of Columbia, the Commonwealth of Puerto Rico, Guam, the U.S. Virgin Islands, American Samoa, Wake Island, Midway Islands, Kingman Reef, or Johnston Island and any place outside thereof.

(j) Commerce between points in any State, the District of Columbia, the Commonwealth of Puerto Rico, Guam, the U.S. Virgin Islands, American Samoa, Wake Island, Midway Islands, Kingman Reef, or Johnston Island, but through any place outside thereof; or

(k) Commerce wholly within the District of Columbia, Guam, the U.S. Virgin Islands, American Samoa, Wake Island, Midway Islands, Kingman Reef, or Johnston Island.

§ 1141.12 Incorporation by reference at § 1141.12.

§ 1141.16 Disclosures regarding cessation.


Subpart A—General Provisions

§§ 1141.1 Scope.

Subpart B—Cigarette Package and Advertising Warnings

§ 1141.3 Definitions.

(a) Person means an individual, partnership, corporation, or any other business or legal entity.

(b) Importer means any person, including any repacker or relabeler, who manufactures, fabricates, assembles, processes, or labels a finished cigarette product.

(c) Package means a pack, box, carton, or container of any kind in which cigarettes are offered for sale, sold, or otherwise distributed to consumers.

(d) Person means an individual, partnership, corporation, or any other business or legal entity.

(e) Required warning means the combination of one of the textual warning statements and its accompanying color graphic, which are set forth in “Cigarette Required Warnings,” which is incorporated by reference at § 1141.12.

(f) Retailer means any person who sells cigarettes to individuals for personal consumption, or who operates a facility where vending machines or self-service displays of cigarettes are permitted.

(g) United States, when used in a geographical sense, includes the several States, the District of Columbia, the Commonwealth of Puerto Rico, Guam, the U.S. Virgin Islands, American Samoa, Wake Island, Midway Islands, Kingman Reef, and Johnston Island. The term “State” includes any political division of any State.

Subpart B—Cigarette Package and Advertising Warnings

§ 1141.10 Required warnings.

(a) Packages—(1) It shall be unlawful for any person to manufacture, package,
sell, offer to sell, distribute, or import for sale or distribution within the United States any cigarettes the package of which fails to bear, in accordance with section 4 of the Federal Cigarette Labeling and Advertising Act (15 U.S.C. 1333) and this part, one of the required warnings on the front and the rear panels.

(2) The required warning shall be obtained from the electronic images contained in “Cigarette Required Warnings,” which is incorporated by reference at §1141.12, and accurately reproduced as specified in “Cigarette Required Warnings.”

(3) The required warning shall appear directly on the package and shall be clearly visible underneath the cellophane or other clear wrapping.

(4) The required warning shall be located in the upper portion of the front and rear panels of the package and shall comprise at least the left 50 percent of these panels; Provided, however, that on cigarette cartons, the required warning shall be located on the left side of the front and rear panels of the carton and shall comprise at least the left 50 percent of these panels.

(5) The required warning shall be positioned such that the text of the required warning and the other information on that panel of the package have the same orientation.

(b) Advertisements—(1) It shall be unlawful for any manufacturer, importer, distributor, or retailer of cigarettes to advertise or cause to be advertised within the United States any cigarettes unless its advertising bears, in accordance with section 4 of the Federal Cigarette Labeling and Advertising Act (15 U.S.C. 1333) and this part, one of the required warnings.

(2) The text in each required warning shall be in the English language, except that:

(i) In the case of an advertisement that appears in a non-English publication, the text in the required warning shall appear in the predominant language of the publication whether or not the advertisement is in English; and

(ii) In the case of an advertisement that appears in an English language publication but that is not in English, the text in the required warning shall appear in the same language as that principally used in the advertisement.

(3) For English-language and Spanish-language warnings, each required warning shall be obtained from the electronic images contained in “Cigarette Required Warnings,” which is incorporated by reference at §1141.12, and accurately reproduced as specified in “Cigarette Required Warnings.”

(4) For foreign-language warnings, except for Spanish-language warnings, each required warning shall be obtained from the electronic images contained in “Cigarette Required Warnings,” which is incorporated by reference at §1141.12, and accurately reproduced as specified in “Cigarette Required Warnings,” including the insertion of a true and accurate translation of the textual warning. The inserted textual warning must comply with the requirements of section 4(b)(2) of the Federal Cigarette Labeling and Advertising Act (15 U.S.C. 1333(b)(2)).

(5) The required warning shall occupy at least 20 percent of the area of each advertisement, and shall be placed in accordance with the requirements in the Federal Cigarette Labeling and Advertising Act.

(c) Irremovable or permanent warnings. The required warnings shall be indelibly printed on or permanently affixed to the package or advertisement. Such warnings, for example, must not be printed or placed on a label affixed to a clear outer wrapper that is likely to be removed to access the product within the package.

§1141.12 Incorporation by reference of required warnings.

“Cigarette Required Warnings” Edition 1.0 (June 2011), consisting of electronic files, U.S. Food and Drug Administration, referred to at §1141.3, §1141.10(a) and (b), and §1141.16(a), is incorporated by reference into this section with the approval of the Director of the Federal Register under 5 U.S.C. 552(a) and 1 CFR part 51. To enforce any edition other than that specified in this section, FDA must publish notice of change in the Federal Register and the material must be available to the public. All approved material is available for inspection at the National Archives and Records Administration (NARA). For information on the availability of this material at NARA, call 202–741–6030 or go to http://www.archives.gov/federal_register/code_of_federal_regulations/ibr_locations.html. Also, you may obtain a copy of the material by contacting the Center for Tobacco Products, Food and Drug Administration, Office of Health Communication and Education, ATTN: Cigarette Warning File Requests, 9200 Corporate Blvd., Rockville, MD 20850, 1–877–CTP–1373, or cigarettewarningfiles@fda.hhs.gov. You may also obtain the material at http://www.fda.gov/cigarettewarningfiles.

§1141.14 Misbranding of cigarettes.

(a) A cigarette shall be deemed to be misbranded under section 903(a)(1) of the Federal Food, Drug, and Cosmetic Act if its package does not bear one of the required warnings in accordance with section 4 of the Federal Cigarette Labeling and Advertising Act (15 U.S.C. 1333) and this part. A cigarette shall be deemed to be misbranded under section 903(a)(7)(A) of the Federal Food, Drug, and Cosmetic Act if its advertising does not bear one of the required warnings in accordance with section 4 of the Federal Cigarette Labeling and Advertising Act (15 U.S.C. 1333) and this part.

(b) A cigarette advertisement or package will be deemed to include a brief statement of relevant warnings for the purposes of section 903(a)(8) of the Federal Food, Drug, and Cosmetic Act if it bears one of the required warnings in accordance with section 4 of the Federal Cigarette Labeling and Advertising Act (15 U.S.C. 1333) and this part. A cigarette distributed or offered for sale in any State shall be deemed to be misbranded under section 903(a)(8) of the Federal Food, Drug, and Cosmetic Act unless the manufacturer, packer, or distributor includes in all advertisements and packages issued or caused to be issued by the manufacturer, packer, or distributor with respect to the cigarette one of the required warnings in accordance with section 4 of the Federal Cigarette Labeling and Advertising Act (15 U.S.C. 1333) and this part.

Subpart C—Additional Disclosure Requirements for Cigarette Packages and Advertising

§1141.16 Disclosures regarding cessation.

(a) The required warning shall include a reference to a smoking cessation assistance resource in accordance with, and as specified in, “Cigarette Required Warnings” (incorporated by reference at §1141.12).

(b) In meeting the smoking cessation needs of an individual caller, the smoking cessation assistance resource required to be referenced by paragraph (a) of this section must, as appropriate:

(1) Provide factual information about the harms to health associated with cigarette smoking and the health benefits of quitting smoking;

(2) Provide factual information about what smokers can expect when trying to quit;

(3) Provide practical advice (problem solving/skills training) about how to deal with common issues faced by smokers trying to quit;
(4) Provide evidence-based advice about how to formulate a plan to quit smoking;
(5) Provide evidence-based information about effective relapse prevention strategies;
(6) Provide factual information on smoking cessation treatments, including FDA-approved cessation medications; and
(7) Provide information, advice, and support that is evidence-based, unbiased (including with respect to products, services, persons, and other entities), and relevant to tobacco cessation.
(c) The smoking cessation resource must:
(1) Other than as described in this section, not advertise or promote any particular product or service;
(2) Except to meet the particularized needs of an individual caller as determined in the context of individual counseling, not selectively present information about a subset of FDA-approved cessation products or product categories while failing to mention other FDA-approved cessation products or product categories;
(3) Not provide or otherwise encourage the use of any drug or other medical product that FDA has not approved for tobacco cessation;
(4) Not encourage the use of any non-evidence-based smoking cessation practices;
(5) Ensure that staff providing smoking cessation information, advice, and support are trained specifically to help smokers quit by delivering unbiased and evidence-based information, advice, and support; and
(6) Maintain appropriate controls to ensure the criteria described in paragraphs (b) and (c) of this section are met.
(d) If the Secretary of the Department of Health and Human Services (Secretary) determines that a part of the smoking cessation assistance resource referenced by paragraph (a) of this section does not meet the criteria described in paragraphs (b) and (c) of this section, the Secretary shall take appropriate steps to address the noncompliance.
Dated: June 9, 2011.
Margaret A. Hamburg, Commissioner of Food and Drugs.
Dated: June 9, 2011.
Kathleen Sebelius, Secretary of Health and Human Services.

Appendices
I. Technical Appendix X1: Smoking Rates
II. Technical Appendix X2: Life-Years
III. Technical Appendix X3: Timing of Benefits
IV. Technical Appendix X4: Timing of Costs
V. Technical Appendix X5: Additional Diagrams on Benefits
VI. Technical Appendix X6: Uncertainty Analysis
A. Alternative Estimation of Smoking Rate Reduction
B. Monte Carlo Simulation

I. Technical Appendix X1: Smoking Rates
FDA’s primary and secondary methods for estimating the reduction in smoking rates realized in Canada due to that country’s introduction, in December 2000, of graphic warning labels both involve several steps. In both methods, the first step is to estimate the smoking rate trend for Canada in the years from 1991 up to and including 2000. (We perform a similar analysis for the United States, but this will be used only in the primary method.)

In response to comments on the Preliminary Regulatory Impact Analysis of the proposed rule, we refine our estimate of the Canadian smoking rate trend by accounting for tax changes at the Federal and provincial levels. The Ontario Flue-Cured Tobacco Growers’ Marketing Board (Ref. 174) reports time series of cigarette taxes for Canadian provinces and territories. (Because these time series only extend back to 1991, we have had to estimate a shorter time trend than the one used in the analysis of the proposed rule.) We find average tax levels for all of Canada by weighting by provincial and territorial populations (using Ref. 175). We then adjust nominal cigarette taxes for general inflation using the broad Canadian CPI (Ref. 176). (Canada has estimated a GDP deflator only since 2002, so we use the Canadian CPI, even though consumer price indices tend to be characterized by slight upward biases in their estimates of inflation.) Our results, along with results from an analogous estimation for the United States, are reported in Table 42.

<table>
<thead>
<tr>
<th>Regression Results, Canadaa</th>
<th>Regression Results, United Statesb</th>
</tr>
</thead>
<tbody>
<tr>
<td>Intercept</td>
<td>Intercept = 4.455</td>
</tr>
<tr>
<td>Standard Error</td>
<td>0.215</td>
</tr>
<tr>
<td>t-statistic</td>
<td>20.715</td>
</tr>
<tr>
<td>Coefficient</td>
<td>-0.377</td>
</tr>
<tr>
<td>Time Trend = ln(Year - 1985)</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Standard Error = 0.063</td>
</tr>
<tr>
<td></td>
<td>t-statistic = -6.012</td>
</tr>
<tr>
<td></td>
<td>Coefficient = -0.215</td>
</tr>
<tr>
<td></td>
<td>Standard Error = 0.080</td>
</tr>
<tr>
<td></td>
<td>t-statistic = -2.688</td>
</tr>
<tr>
<td>N</td>
<td>7</td>
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<tr>
<td></td>
<td>7</td>
</tr>
<tr>
<td></td>
<td>Standard Error = 0.074</td>
</tr>
<tr>
<td></td>
<td>t-statistic = -1.551</td>
</tr>
<tr>
<td></td>
<td>Coefficient = -0.101</td>
</tr>
<tr>
<td></td>
<td>Standard Error = 0.106</td>
</tr>
<tr>
<td></td>
<td>t-statistic = -0.950</td>
</tr>
</tbody>
</table>

a Underlying smoking rate data appear in table 4 of this document.
b Regression equation: ln(SmokingRate) = Intercept + Coefficient*ln(Year-1985) + Coefficient*ln(ExciseTax) + error.
c Standard errors and t-statistics reported here are not adjusted for uncertainty introduced by the use of survey data.
d Regression equation: ln(SmokingRate) = Intercept + Coefficient*ln(Year-1985) + Coefficient*ln(ExciseTax) + error.
Using the estimated time trend, we forecast the Canadian smoking rate that would have been realized post-2000 had graphic warning labels not been introduced in that country. The difference between the smoking rate forecast and the actual Canadian smoking rate yields the portion of the smoking rate that is unexplained apart from the introduction of graphic warning labels. Calculating the difference in the average unexplained smoking rate between 1994–2000 and 2001–09 yields the estimate of the effect of graphic warning labels, 0.574 percentage points, that appears in part (a) of Technical Appendix X6.

In our preferred estimation method (see section XI.D.1, above), we use the U.S. experience as an additional control. We find the unexplained smoking rate in the United States using calculations analogous to those used for Canada and tax data from the Centers for Disease Control and Prevention (Ref. 177) and Jamison et al. (Ref. 178), population data from the U.S. Census Bureau (Refs. 179 and 180), and inflation data from the U.S. Bureau of Economic Analysis (Ref. 132). We then calculate the difference in unexplained smoking rates between the United States and Canada. Finally, we again subtract the average for 1994–2000 from the average for 2001–09; this produces the estimate that graphic warning labels decrease the national smoking rate by 0.088 percentage points. Details appear in Table 44.

### Table 43.--Impact of Graphic Warning Labels on Canadian Smoking Rate

<table>
<thead>
<tr>
<th></th>
<th>Smiling Rate, Canada</th>
<th>Time Trend Forecast Smoking Rate, Canada</th>
<th>Unexplained Smoking Rate, Canada</th>
</tr>
</thead>
<tbody>
<tr>
<td>1994-95</td>
<td>30.5</td>
<td>30.391</td>
<td>0.109</td>
</tr>
<tr>
<td>1996-97</td>
<td>28.6</td>
<td>28.172</td>
<td>0.428</td>
</tr>
<tr>
<td>1998-99</td>
<td>27.7</td>
<td>26.237</td>
<td>1.463</td>
</tr>
<tr>
<td>1999</td>
<td>25.2</td>
<td>25.855</td>
<td>-0.655</td>
</tr>
<tr>
<td>2000</td>
<td>24.4</td>
<td>25.099</td>
<td>-0.699</td>
</tr>
<tr>
<td>2001</td>
<td>21.7</td>
<td>24.088</td>
<td>-2.388</td>
</tr>
<tr>
<td>2002</td>
<td>21.4</td>
<td>22.247</td>
<td>-0.847</td>
</tr>
<tr>
<td>2003</td>
<td>20.9</td>
<td>20.274</td>
<td>0.626</td>
</tr>
<tr>
<td>2004</td>
<td>19.6</td>
<td>19.596</td>
<td>0.004</td>
</tr>
<tr>
<td>2005</td>
<td>18.7</td>
<td>19.242</td>
<td>-0.542</td>
</tr>
<tr>
<td>2006</td>
<td>18.6</td>
<td>18.950</td>
<td>-0.350</td>
</tr>
<tr>
<td>2007</td>
<td>19.2</td>
<td>18.607</td>
<td>0.593</td>
</tr>
<tr>
<td>2008</td>
<td>17.9</td>
<td>18.291</td>
<td>-0.391</td>
</tr>
<tr>
<td>2009</td>
<td>17.25</td>
<td>17.957</td>
<td>-0.707</td>
</tr>
</tbody>
</table>

a Source: Health Canada (Refs. 126 and 127).

b Mean for 1994-2000 is 0.129; mean for 2001-09 is -0.445; difference in means is 0.574.

### Table 44.--Impact of Graphic Warning Labels on Difference Between Unexplained United States and Canadian Smoking Rates

<table>
<thead>
<tr>
<th></th>
<th>Smoking Rate, United States(^a)</th>
<th>Standard Error, Smoking Rate, United States(^b)</th>
<th>Time Trend Forecast Smoking Rate, United States</th>
<th>Unexplained Smoking Rate, United States</th>
<th>Difference in Unexplained Smoking Rates (United States-Canada)(^c)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1994-95</td>
<td>24.6</td>
<td>b</td>
<td>24.742</td>
<td>-0.142</td>
<td>-0.251</td>
</tr>
<tr>
<td>1996-97</td>
<td>24.558</td>
<td>0.29</td>
<td>24.213</td>
<td>0.344</td>
<td>-0.083</td>
</tr>
<tr>
<td>1998</td>
<td>23.918</td>
<td>0.30</td>
<td>23.971</td>
<td>-0.053</td>
<td>-1.516</td>
</tr>
<tr>
<td>1999</td>
<td>23.302</td>
<td>0.32</td>
<td>23.564</td>
<td>-0.261</td>
<td>0.393</td>
</tr>
<tr>
<td>2000</td>
<td>23.065</td>
<td>0.32</td>
<td>23.005</td>
<td>0.060</td>
<td>0.759</td>
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<tr>
<td>2001</td>
<td>22.644</td>
<td>0.30</td>
<td>22.869</td>
<td>-0.226</td>
<td>2.162</td>
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<tr>
<td>2002</td>
<td>22.262</td>
<td>0.32</td>
<td>22.141</td>
<td>0.121</td>
<td>0.967</td>
</tr>
<tr>
<td>2003</td>
<td>21.310</td>
<td>0.30</td>
<td>21.945</td>
<td>-0.635</td>
<td>-1.261</td>
</tr>
<tr>
<td>2004</td>
<td>20.724</td>
<td>0.31</td>
<td>21.538</td>
<td>-0.814</td>
<td>-0.272</td>
</tr>
<tr>
<td>2005</td>
<td>20.564</td>
<td>0.35</td>
<td>21.447</td>
<td>-0.882</td>
<td>-0.533</td>
</tr>
<tr>
<td>2006</td>
<td>19.449</td>
<td>0.40</td>
<td>21.211</td>
<td>-1.762</td>
<td>2.356</td>
</tr>
<tr>
<td>2007</td>
<td>20.409</td>
<td>0.38</td>
<td>20.948</td>
<td>-0.539</td>
<td>-0.148</td>
</tr>
<tr>
<td>2008</td>
<td>20.513</td>
<td>0.37</td>
<td>20.190</td>
<td>0.323</td>
<td>1.030</td>
</tr>
</tbody>
</table>

\(^a\) Sources: National Center for Health Statistics (Ref. 129) and FDA analysis of National Health Interview Survey (Ref. 128).

\(^b\) Not reported for 1994, but likely to be near the standard error of 0.3 found for years 2000-03.

\(^c\) Mean for 1994-2000 is -0.140; mean for 2001-09 is -0.051; difference in means is 0.088.
II. Technical Appendix X2: Life-Years

In calculating expected life-years saved per dissuaded smoker, FDA relies heavily on the life tables developed by Sloan et al. (Ref. 116). The life tables are calculated from the perspective of 24-year-olds, so the calculation of rule-induced effects on males and females who turn 24 sometime after the rule takes effect is relatively straightforward. In the following example, we will show the calculation of expected rule-induced effects for 24-year-old females, under the assumption of a 3 percent discount rate; the calculations for males or for a 7 percent discount rate would be analogous.

The life tables show that, of one hundred thousand females who smoke at their 24th birthdays, 99,939 will survive to their 25th birthday and 99,876 to their 26th birthdays. Of one hundred thousand 24-year-old female nonsmoking smokers, 99,946 will survive to their 25th birthdays and 99,889 to their 26th birthdays. These numbers imply that, for every one hundred thousand females who smoke at their 24th birthdays, smoking will cause seven deaths between birthdays 24 and 25 and six deaths between birthdays 25 and 26. The tables continue to show number of survivors in each category (and thus the smoking-related excess probability of dying) for every birthday up to age 100; the discontinuation of the tables at this point requires us to assume no survival in either category to the one-hundred-and-first birthday.

Someone who dies at the age of 24 lose all the life-years up to and including age 100. Without discounting, this would be a total of 77 years; with a 3 percent discount rate, however, the total is 29.9 years. Similarly, someone who dies at age 25 loses 76 undiscounted or 29.8 discounted life-years. By multiplying together the age-specific discounted life-year loss and the age-specific smoking-related excess probability of dying, then summing over all ages, we arrive at the overall expected number of life-years saved per dissuaded female smoker. Using a discount rate of 3 percent, this result is (7/100,000)*29.9 + (6/100,000)*29.8 + ... = 0.524.

For individuals who are older than 24 at the time of the rule’s implementation, we want to perform a similar calculation; however, direct application of the nonsmoking smoker life tables is inappropriate because the life expectancy effect of smoking cessation at a particular age is almost certainly different than the effect of having refrained from smoking since at least the age of 24. Thus, it is necessary to develop age-specific survival probabilities for former smokers.

There are four possible events that a 24-year-old smoker can experience between any two birthdays: staying alive and remaining a smoker, staying alive and becoming a former smoker, dying in the state of being a smoking, or dying in the state of being a former smoker. The percentage of former smokers who do not experience the last of these events is the former smoker survival probability that we seek to calculate. We will illustrate this calculation for 25-year-old females, under the assumption of a 3 percent discount rate; the calculation for males or other discount rates or age categories would be analogous.

We again consider one hundred thousand female smokers at their 24th birthdays. According to the National Health Interview Survey (Ref. 128), 3.4 percent of them will become former smokers by their 25th birthdays. Following Sloan et al., we use the 1998 NHIS and define former smokers as individuals who quit at least 5 years in the past. Sloan et al.’s life tables indicate that another 61 of the original one hundred thousand will die before their 25th birthdays; all 61 die in the state of being smokers (because no time has elapsed since they were smokers at the definitional age of 24). This leaves 96,540 who are alive and still smoking and 3,399 who are living former smokers at the 25th birthday.

Sloan et al.’s typical smoker life table indicates that 63 of these 25-year-old survivors will die before their 26th birthdays; we must calculate how many of them die in the state of being smokers and how many in the state of being former smokers. To find death probabilities for those individuals who are still smoking at age 25, we look to Sloan et al.’s life table for lifetime smokers. Whereas the typical smoker life table shows survival patterns for individuals who smoke at age 24 and may quit sometime later in life, the lifetime smoker life table isolates survival patterns for individuals who smoke at age 24 and continue to a specific age. The lifetime smoker life table will begin to diverge from the typical life table at later ages, but for birthdays 25 and 26, the results are once again 99,939 and 99,876 survivors; therefore, the percentage of 25-year-old female smokers who survive to birthday 26 is 99,876/99,939. Multiplying this percentage by the 96,540 smokers alive at birthday 25 yields 61 deaths.

The results for the life expectancy benefits being too great for those who quit at an advanced age and too low for those who quit at an early age.)

To find the expected number of life-years gained for a female who quits smoking at age 25, we subtract from 0.99937 the survival probability for a smoker of the same age (calculated from Sloan et al.’s typical smoker life table), then multiply by the discounted number of life-years lost if death occurs at age 25 (previously found to be 29.8), and finally add the expected value of life-years saved by quitting at age 26, discounted 1 year. Because there is no extension of life brought about by quitting at age 100, this addition is feasible for age 99, and then for age 98, and so on back to age 25. The final result for females who quit smoking at age 25 is 0.081 discounted life-years saved.

For the year 2013, we multiply our estimated age-specific expected discounted life-years saved by the cohort sizes (for ages 18 and above) projected by the U.S. Census Bureau (Ref. 130). For years 2014–31, we multiply our estimated age-specific expected discounted life-years saved by the cohorts that would not have been included in our 2013 calculation, specifically new 24-year-olds and older individuals whose cohorts grow from one year to another (for example, if the projected number of 35-year-olds in 2014 is greater than the projected number of 34-year-olds in 2013, the difference is included in the 2014 calculation). Finally, we estimate effects for individuals who are 18–23 in the year 2031 by discounting the present value of benefits accruing to 24-year-olds by the number of years until each cohort reaches that age threshold.

Results are further multiplied by FDA’s estimate of the rule-induced reduction in the U.S. smoking rate to yield our final estimate of the number of life-years saved by the regulation.

III. Technical Appendix X3: Timing of Benefits

FDA’s estimated benefits appear as undiscounted streams in Table 45. Parts 1 through 12. Benefits are realized as late as 2113 because we calculate effects over lifetimes extending to age 100 for
cohorts aged 18 and above during the first 20 years (2012 to 2031) of the final rule’s implementation. Because many of our sources report only present values of smoking-related effects, estimating the timing of those effects requires us to make various assumptions. Changing those assumptions would change the results appearing in Table 45. Similarly, because many of our sources report present values calculated only with a discount rate of 3 percent, changing our assumptions about the timing of effects would change the present values we have reported at the 7 percent discount rate (an important exception being the present value of reduced mortality for 24-year-olds because Sloan et al.’s life tables allow us to know the timing of those benefits).

<p>| Table 45.—Undiscounted Stream of Benefits and Consumer Surplus Costs ($ mil), Part 1 |
|------------------|------------------|------------------|------------------|------------------|------------------|------------------|------------------|------------------|------------------|
| Mortality, Age = 18-24 during 2013-31, with VSLY = $212,615 ( ^a ) | 0.0              | 0.0              | 0.0              | 0.0              | -0.1             | -0.1             | -0.1             | -0.1             | -0.2             |
| Mortality, Age &gt; 24 in 2013, with VSLY = $212,615 ( ^{b, h} ) | 0.0              | 10.4             | 20.9             | 31.3             | 41.7             | 52.0             | 62.2             | 72.2             | 82.1             |
| Health Status, with VSLY = $212,615 ( ^{c} ) | 0.0              | 6.1              | 12.3             | 18.5             | 24.6             | 30.6             | 36.5             | 42.4             | 48.2             |
| Medical Costs Reductions, Age = 18-24 during 2013-31 ( ^d ) | 0.0              | 1.2              | 2.5              | 3.8              | 5.0              | 6.3              | 7.5              | 8.7              | 9.9              |
| Medical Costs Reductions, Age &gt; 24 in 2013 ( ^e ) | 0.0              | 21.7             | 21.7             | 21.7             | 21.7             | 21.7             | 21.7             | 21.6             | 21.6             |
| Financial Effects ( ^f ) | 0.0              | 1.1              | 1.6              | 2.1              | 2.6              | 3.1              | 3.6              | 4.1              | 4.6              |
| Fire-Related Mortality with 3% Disc. Rate and VSLY = $212,615 ( ^{g, h} ) | 0.0              | 4.9              | 4.9              | 5.0              | 5.1              | 5.2              | 5.2              | 5.3              | 5.4              |
| Fire-Related Mortality with 7% Disc. Rate and VSLY = $212,615 ( ^{h, h} ) | 0.0              | 2.8              | 2.9              | 2.9              | 3.0              | 3.0              | 3.1              | 3.1              | 3.2              |
| Fire-Related Property Damage ( ^i ) | 0.0              | 0.9              | 0.9              | 0.9              | 0.9              | 0.9              | 0.9              | 0.9              | 1.0              |
| Consumer Surplus with 3% Disc. Rate and VSLY = $212,615 ( ^{j} ) | 0.0              | 308.1             | 313.0             | 317.8             | 322.6             | 327.4             | 332.4             | 337.7             | 343.0             |
| Consumer Surplus with 7% Disc. Rate and VSLY = $212,615 ( ^{k, j} ) | 0.0              | 139.9             | 142.2             | 144.3             | 146.5             | 148.7             | 151.0             | 153.4             | 155.8             |</p>
<table>
<thead>
<tr>
<th>Description</th>
<th>2021</th>
<th>2022</th>
<th>2023</th>
<th>2024</th>
<th>2025</th>
<th>2026</th>
<th>2027</th>
<th>2028</th>
<th>2029</th>
</tr>
</thead>
<tbody>
<tr>
<td>Mortality, Age = 18-24 during 2013-31, with VSLY = $212,615^d</td>
<td>-0.2</td>
<td>-0.2</td>
<td>-0.1</td>
<td>-0.1</td>
<td>0.7</td>
<td>2.2</td>
<td>4.5</td>
<td>7.7</td>
<td>11.6</td>
</tr>
<tr>
<td>Mortality, Age &gt; 24 in 2013, with VSLY = $212,615^a,b</td>
<td>91.8</td>
<td>101.3</td>
<td>110.6</td>
<td>119.7</td>
<td>128.3</td>
<td>136.4</td>
<td>144.0</td>
<td>151.1</td>
<td>157.8</td>
</tr>
<tr>
<td>Health Status, with VSLY = $212,615^c</td>
<td>53.9</td>
<td>59.6</td>
<td>65.3</td>
<td>71.1</td>
<td>77.1</td>
<td>83.2</td>
<td>89.3</td>
<td>95.4</td>
<td>101.6</td>
</tr>
<tr>
<td>Medical Costs Reductions, Age = 18-24 during 2013-31^d</td>
<td>11.0</td>
<td>12.2</td>
<td>13.4</td>
<td>14.6</td>
<td>15.8</td>
<td>17.0</td>
<td>18.3</td>
<td>19.5</td>
<td>20.8</td>
</tr>
<tr>
<td>Financial Effects^f</td>
<td>5.0</td>
<td>5.5</td>
<td>5.9</td>
<td>6.4</td>
<td>6.9</td>
<td>7.3</td>
<td>7.8</td>
<td>8.3</td>
<td>8.8</td>
</tr>
<tr>
<td>Fire-Related Mortality with 3% Disc. Rate and VSLY = $212,615^e</td>
<td>5.5</td>
<td>5.6</td>
<td>5.7</td>
<td>5.8</td>
<td>5.8</td>
<td>5.9</td>
<td>6.0</td>
<td>6.1</td>
<td>6.2</td>
</tr>
<tr>
<td>Fire-Related Mortality with 7% Disc. Rate and VSLY = $212,615^f,b</td>
<td>3.2</td>
<td>3.3</td>
<td>3.3</td>
<td>3.4</td>
<td>3.4</td>
<td>3.5</td>
<td>3.5</td>
<td>3.6</td>
<td>3.6</td>
</tr>
<tr>
<td>Fire-Related Property Damage^g</td>
<td>1.0</td>
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<td>Consumer Surplus with 3% Disc. Rate and VSLY = $212,615^h</td>
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<td>359.4</td>
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<tr>
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<td>Mortality, Age &gt; 24 in 2013, with VSLY = $212,615&lt;sup&gt;b&lt;/sup&gt;</td>
<td>163.9 169.5 174.6 179.2 183.3 186.8 189.6 191.8 193.4</td>
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<td>181.8 184.1 182.4 180.7 178.9 177.1 175.1 173.2 172.3</td>
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<td>203.2</td>
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<td>374.9</td>
<td>372.6</td>
<td>370.1</td>
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<td>170.3</td>
<td>169.2</td>
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<td>Mortality, Age = 18-24 during 2013-31, with VSLY = $212,615^{d}</td>
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<td>Mortality, Age &gt; 24 in 2013, with VSLY = $212,615^{ah}</td>
<td>177.9</td>
<td>173.4</td>
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Table 45.—Undiscounted Stream of Benefits and Consumer Surplus Costs ($ mil), Part 6

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<td>VSLY = $212,615^{ij}</td>
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### Table 45.--Undiscounted Stream of Benefits and Consumer Surplus Costs ($ mil), Part 7

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<td>4.0</td>
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<td>2.0</td>
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<td>Fire-Related Property Damage&lt;sup&gt;i&lt;/sup&gt;</td>
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<td>255.9</td>
<td>250.9</td>
<td>245.9</td>
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<td>118.5</td>
<td>116.2</td>
<td>113.9</td>
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<td>109.3</td>
<td>106.9</td>
<td>104.5</td>
<td>102.2</td>
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Table 45.—Undiscounted Stream of Benefits and Consumer Surplus Costs ($ mil), Part 8

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<th>2075</th>
<th>2076</th>
<th>2077</th>
<th>2078</th>
<th>2079</th>
<th>2080</th>
<th>2081</th>
<th>2082</th>
<th>2083</th>
</tr>
</thead>
</table>
| Mortality, Age = 18-24 during 2013-31, with VSLY = $212,615
during 2013-31 | 19.8  | 16.8  | 14.1  | 11.8  | 9.7   | 8.0   | 6.5   | 5.2   | 4.1   |
| Mortality, Age > 24 in 2013, with VSLY = $212,615 | 152.4 | 152.4 | 152.4 | 152.4 | 152.4 | 152.4 | 152.4 | 152.4 | 152.4 |
| Health Status, with VSLY = $212,615 | -34.1 | -36.1 | -38.1 | -40.1 | -40.1 | -40.1 | -40.1 | -40.1 | -40.1 |
| Medical Costs Reductions, Age = 18-24 during 2013-31 | 6.7   | 6.2   | 5.8   | 5.4   | 5.0   | 4.6   | 4.1   | 3.7   | 3.2   |
| 2013 | Financial Effects | 62.1  | 62.1  | 61.7  | 60.9  | 59.9  | 58.6  | 57.0  | 55.3  | 53.3  |
| Fire-Related Mortality with 3% Disc. Rate and VSLY = $212,615 | 3.4   | 3.3   | 3.2   | 3.2   | 3.1   | 3.0   | 2.9   | 2.8   | 2.7   |
| Fire-Related Mortality with 7% Disc. Rate and VSLY = $212,615 | 2.0   | 1.9   | 1.9   | 1.8   | 1.8   | 1.7   | 1.7   | 1.6   | 1.6   |
| Fire-Related Property Damage | 0.6   | 0.6   | 0.6   | 0.6   | 0.5   | 0.5   | 0.5   | 0.5   | 0.5   |
| Consumer Surplus with 3% Disc. Rate and VSLY = $212,615 | 215.4 | 210.6 | 205.7 | 200.9 | 195.8 | 190.7 | 185.3 | 179.8 | 174.3 |
| Consumer Surplus with 7% Disc. Rate and VSLY = $212,615 | 97.9  | 95.6  | 93.5  | 91.2  | 89.0  | 86.6  | 84.1  | 81.7  | 79.2  |
### Table 45.--Undiscounted Stream of Benefits and Consumer Surplus Costs ($ mil), Part 9

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<th>2086</th>
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<th>2091</th>
<th>2092</th>
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<td>Mortality, Age = 18-24 during 2013-31, with VSLY = $212,615(a)</td>
<td>1,692.7</td>
<td>1,611.9</td>
<td>1,526.4</td>
<td>1,436.4</td>
<td>1,342.4</td>
<td>1,245.4</td>
<td>1,147.1</td>
<td>1,047.9</td>
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<td>Mortality, Age &gt; 24 in 2013, with VSLY = $212,615(b)</td>
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<td>1.5</td>
<td>1.1</td>
<td>0.9</td>
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<tr>
<td>Health Status, with VSLY = $212,615(c)</td>
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<td>152.4</td>
<td>152.4</td>
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<td>146.4</td>
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<td>-40.1</td>
<td>-40.1</td>
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<td>-40.1</td>
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<td>Fire-Related Mortality with 3% Disc. Rate and VSLY = $212,615(g)</td>
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<td>2.6</td>
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<td>2.2</td>
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<td>Fire-Related Mortality with 7% Disc. Rate and VSLY = $212,615(h)</td>
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<td>1.5</td>
<td>1.5</td>
<td>1.4</td>
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</tr>
<tr>
<td>Mortality, Age = 18-24 during 2013-31, with VSLY = $212,615&lt;sup&gt;a&lt;/sup&gt;</td>
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<td>96.4</td>
<td>67.2</td>
<td>45.0</td>
<td>28.7</td>
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<td>9.5</td>
<td>4.7</td>
<td>1.9</td>
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<td>0.0</td>
<td>0.0</td>
<td>0.0</td>
<td>0.0</td>
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<tr>
<td>Health Status, with VSLY = $212,615&lt;sup&gt;c&lt;/sup&gt;</td>
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<td>69.3</td>
<td>63.2</td>
<td>57.0</td>
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<td>44.5</td>
<td>38.3</td>
<td>32.0</td>
<td>25.6</td>
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<tr>
<td>Medical Costs Reductions, Age = 18-24 during 2013-31&lt;sup&gt;d&lt;/sup&gt;</td>
<td>-19.8</td>
<td>-18.2</td>
<td>-16.6</td>
<td>-15.0</td>
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<td>Financial Effects&lt;sup&gt;g&lt;/sup&gt;</td>
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<td>Fire-Related Mortality with 3% Disc. Rate and VSLY = $212,615&lt;sup&gt;h&lt;/sup&gt;</td>
<td>1.9</td>
<td>1.8</td>
<td>1.7</td>
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<td>1.0</td>
<td>1.0</td>
<td>1.0</td>
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<td>Fire-Related Property Damage&lt;sup&gt;l&lt;/sup&gt;</td>
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<td>0.3</td>
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<td>Consumer Surplus with 3% Disc. Rate and VSLY = $212,615&lt;sup&gt;j&lt;/sup&gt;</td>
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<tr>
<td>Mortality, Age &gt; 24 in 2013, with VSLY = $212,615^{ab}</td>
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<td>Health Status, with VSLY = $212,615^{ac}</td>
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<td>12.9</td>
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<tr>
<td>Medical Costs Reductions, Age = 18-24 during 2013-31^g</td>
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<td>-1.7</td>
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<td>Medical Costs Reductions, Age &gt; 24 in 2013^g</td>
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<td>Financial Effects^f</td>
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<td>Fire-Related Mortality with 3% Disc. Rate and VSLY = $212,615^{kg}</td>
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<td>Fire-Related Mortality with 7% Disc. Rate and VSLY = $212,615^{kh}</td>
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<td>0.3</td>
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<tr>
<td>Fire-Related Property Damage^i</td>
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<tr>
<td>Consumer Surplus with 3% Disc. Rate and VSLY = $212,615^{ij}</td>
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<td>33.4</td>
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<tr>
<td>Consumer Surplus with 7% Disc. Rate and VSLY = $212,615^{jk}</td>
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<td>15.2</td>
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</tbody>
</table>
IV. Technical Appendix X4: Timing of Costs

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\( ^a \) Numbers in this row may be multiplied by 0.5 to produce results for VSLY=$106,308 or by 1.5 to produce results for VSLY=$318,923.

\( ^b \) Also includes individuals who turn 24 between 2013 and 2031 but are first exposed to graphic warning labels at later ages due to immigration. Underlying assumptions discussed in detail in Technical Appendix X2.

\( ^c \) Underlying assumption: Sloan et al.’s present value of years with fair/poor health status distributed equally over ages 24 to 100. Result: this row shows benefits being accrued in a pattern somewhat less concentrated in the middle years of life than the likely reality. Because Sloan et al. report undiscounted effects of 2.69 years for females and 1.41 year for males, and discounting reduces the effects to 1.27 and 0.90 years, this concentration, on average, centers on females’ forty-ninth birthdays and males’ thirty-ninth birthdays.

\( ^d \) Underlying assumption: Sloan et al.’s medical cost present value distributed equally within age bins (24-50, 51-64 and 65+).

\( ^e \) Also includes individuals who turn 24 between 2013 and 2031 but are first exposed to graphic warning labels at later ages due to immigration. Underlying assumption: Sloan et al.’s medical costs present value distributed equally over ages 24 to 100. Result: this row shows benefits being accrued somewhat later and in lesser amounts than the likely reality for relatively young quitters and somewhat earlier and in greater amounts than the likely reality for relatively old quitters.

\( ^f \) Includes Social Security outlays, income taxes on Social Security-taxable earnings, defined benefit private pension outlays and life insurance outlays. Underlying assumption: net financial effect distributed over time in the same pattern as the sum of mortality, morbidity and medical cost effects.

Underlying assumption for quitters aged 25 and above: Sloan et al.’s cigarette consumption present value distributed equally over ages 24 to 100. Result: this row shows benefits being accrued somewhat later and possibly in slightly greater amounts than the likely reality. Fire-related death loss is a present value, calculated at the time of death with a discount rate of 3 percent, of future VSLY.

Underlying assumption for quitters aged 25 and above: Sloan et al.’s cigarette consumption present value distributed equally over ages 24 to 100. Result: this row shows benefits being accrued somewhat later and possibly in slightly greater amounts than the likely reality. Fire-related death loss is a present value, calculated at the time of death with a discount rate of 7 percent, of future VSLY.

Underlying assumption for quitters aged 25 and above: Sloan et al.’s cigarette consumption present value distributed equally over ages 24 to 100. Result: this row shows benefits being accrued somewhat later and possibly in slightly greater amounts than the likely reality.

\( ^g \) Numbers in this row may be multiplied by approximately 0.496 to produce results for VSLY=$106,308 or by approximately 1.504 to produce results for VSLY=$318,923.

\( ^h \) Numbers in this row may be multiplied by approximately 0.520 to produce results for VSLY=$106,308 or by approximately 1.500 to produce results for VSLY=$318,923.
### Table 46.--Undiscounted Stream of Costs, Low Estimate ($ mil), Part 1

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<td>Point-of-Sale Advertising</td>
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<td>FDA</td>
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### Table 46.--Undiscounted Stream of Costs, Low Estimate ($ mil), Part 2

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<tr>
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### Table 47.--Undiscounted Stream of Costs, Medium Estimate ($ mil), Part 1

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V. Technical Appendix X5: Additional Diagrams on Benefits

Consumer Surplus Model. The benefits estimated in sections XLD.2.b.ii, XLD.2.b.iii, XLD.2.b.iv and XLD.2.b.v overstate, all else held equal, the net internal (i.e., intrapersonal) benefits (or costs, in the case of section XLD.2.b.v) of reduced smoking because they include only the increased welfare from improved health and expected longevity (and decreased welfare due to subsidy loss) and do not account for any lost consumer surplus associated with the activity of smoking. In the Preliminary Regulatory Impact Analysis (see page 75 FR 69524 at 69544), FDA adjusted benefits estimates with a 50 percent consumer surplus reduction, based on a model created by Cutler (Ref. 134). Several comments on the proposed rule expressed concern about the appropriateness of Cutler’s assumptions, so FDA has revised the model to make it more applicable to the present analysis. Our revised model is illustrated in Figure E1.

\[^{21}\text{The difference between what a consumer would be willing to pay for a good or service and what that consumer actually has to pay.}\]
We begin with a downward-sloping demand for typical lifetime smoking. A negative relationship between price and consumption of cigarettes has been demonstrated empirically many times over (Chaloupka and Warner (Ref. 162) review this literature).

The height of line DCS_{full} marks the full cost, including the cost of adverse health and life expectancy effects, of typical lifetime smoking (thus, the “Discounted Cost of Smoking” or DCS), while the height of line DCS_{money} marks only the after-tax price of cigarettes. The height difference between these two lines is the sum of the per-person effects we calculated in sections XI.D.2.b.ii, XI.D.2.b.iii and XI.D.2.b.iv. Also belonging in DCS_{full} are the effects calculated in section XI.D.2.b.v because the concept of the full cost of smoking, as used in the model, is defined from the private perspective of the smoker (and thus it is irrelevant whether or not there is someone else in society who experiences an effect that offsets the cost or benefit experienced by the smoker—which is what distinguishes the entries in Tables 22 and 23 from the effects in sections XI.D.2.b.ii, XI.D.2.b.iii and XI.D.2.b.iv). While the elements in Tables 22 and 23 do contribute to DCS_{full}, we posit that they should not be thought of as included in DCS_{money} because they are intricately related to the mortality and morbidity effects of smoking that, unlike the after-tax price of cigarettes, are likely characterized by time inconsistency, incomplete information or other sources of market failure.

Society will be at the intersection of Demand and DCS_{money} if the health costs associated with smoking are not known or, if known, cannot be “internalized” and incorporated into consumption decisions. The current widespread awareness that smoking poses health risks and the significant decline in smoking rates over the past 50 years make it highly implausible that actual consumption is near that hypothetical level. The intersection of the Demand line and DCS_{full} represents the other extreme. At that hypothetical level, consumers are fully aware of all known risks and have internalized all health costs and incorporated them into consumption decisions. The economic models and empirical studies of addiction, self-control, and time inconsistency (which we discuss in detail in our response to comments on the preliminary analysis) strongly suggest that health costs are not fully internalized; the behaviors that lead to less-than-full internalization appear to be common. In surveys, many smokers express a desire to quit and report that they have tried to stop smoking. The demand for various aids to smoking cessation provides further evidence of less-than-full internalization. Moreover, the immature judgments, short time horizons and lack of self-control of most children and adolescents—who make up the vast majority of new smokers—suggest that policy interventions that prevent initiation and encourage cessation can increase social welfare.

For these reasons, we find it implausible that actual consumption is at the intersection of Demand and DCS_{full}. The number of current smokers is therefore found at the intersection of Demand with a line falling somewhere between DCS_{full} and DCS_{money}. We have drawn this as line DCS_{money}. Our finding that the graphic warning label regulation will reduce smoking rates is represented by an upward shift of this line to DCS_{rule}. (This may seem less intuitive to some readers than shifting the demand curve—which is the approach taken by Weimer et al. (Ref. 161)—but the two analytic methods will produce equivalent results, as we illustrate below.) The intersections of DCS_{absence} and DCS_{rule} with the demand curve show the number of smokers, Q_{absence} and Q_{rule}, in the absence and in the presence of the final rule.

In the absence of the final rule, total cost, including health costs, for smokers is shown by the sum of areas B through K. We reiterate that, even though consumers do not internalize all costs upfront, they do ultimately incur them. The gross value smokers place on cigarette consumption (known as willingness-to-pay) is the area under the demand curve as far right as Q_{absence}, or $A + B + E + F + H + I + J + K$. The net value to smokers of cigarette consumption is thus $(A + B + E + F + H + I + J + K) - (B + C + D + E + F + G + H + I + J + K) = A - (C + D + G)$.

In the presence of the final rule, total expenditure, including health costs, by smokers is $B + C + E + H + J$. Smokers' willingness-to-pay is the area under the demand curve as far right as $Q_{rule}$, or $A + B + E + H + J$. The net value to smokers of cigarette consumption is thus $(A + B + E + H + J) - (B + C + E + H + J) = A - C$. As a result, the effect of the rule is to increase net value by $(A - C) - [A - (C + D + G)] = D + G$. 

![Figure E1. The Market for Smoking, Before and After Rule Implementation](image-url)
The calculations appearing in sections XI.D.2.b.ii, XI.D.2.b.iii, XI.D.2.b.iv and XI.D.2.b.v each consist of multiplying \((Q_{\text{absence}} - Q_{\text{rule}})\) by some portion of \((DCS_{\text{full}} - DCS_{\text{money}})\); therefore, summing the results of D2b.ii, D2b.iii, D2b.iv and D2b.v produces an estimate of \((D+F+G+I)\). Because we have already established that the benefit of the rule is \((D+G)\), reporting the unadjusted sum of results from sections XI.D.2.b.ii, XI.D.2.b.iii, XI.D.2.b.iv and XI.D.2.b.v would cause us to overestimate the benefits of the final rule by an amount equal to \((D+F+G+I) - (D+G) = (F+I)\). As drawn in Figure E1, \((F+I)\) is approximately 50 percent of the unadjusted estimate, \((D+F+G+I)\). FDA does not claim that 50 percent is the correct ratio; the correct ratio of \((D+F+G+I)\) to \((D+F+G+I)\) is determined by the shape of the demand curve as it divides areas \(F\) and \(G\) and, more pertinently, by the relative height differences between DCS_{\text{full}} and DCS_{\text{rule}} and between DCS_{\text{absence}} and DCS_{\text{money}}. \((DCS_{\text{full}} - DCS_{\text{rule}})\) may be much greater than \((DCS_{\text{absence}} - DCS_{\text{money}})\) or it may be much less, yielding a ratio that may be near zero or may be near 100 percent, depending on the starting height of DCS_{\text{absence}} and the size of the policy-induced reduction in smoking.

We now parameterize this model using the literature on the economics of habits and addiction. (We note, however, that rigorous quantitative welfare analyses of tobacco control interventions are rare in published, peer-reviewed literature, so the estimates generated below should not be viewed as definitive.) First, the Robert Wood Johnson Foundation (Ref. 137) reports that, as of 2009, State and Federal taxes made up 40.4 percent of the total retail price of cigarettes. With the Federal cigarette excise tax being $1.01 per pack (Ref. 164) and the population-weighted average State tax being $1.33 per pack (Ref. 165, with population weights from Ref. 130), we estimate the average after-tax price of a pack of cigarettes, or the height of DCS_{\text{money}}, to be $5.78. FDA’s analysis in section XI.D.2.b of the benefits of smoking reduction has produced an estimate of discounted internal health and financial effects (reduced mortality, morbidity, medical costs and implicit smoking subsidy) that ranges from $2.10 billion to $27.80 billion in total, or from $4.56 to $27.69 per pack; this range indicates the range of potential height differences between DCS_{\text{full}} and DCS_{\text{money}}. We can derive the heights of the remaining DCS curves from a simulation conducted by Gruber and Köszegi (Ref. 104), in which they estimate the tax rate that would allow time-inconsistent smokers to consume the quantity that would be optimal under perfect rationality. Because this quantity is found at the intersection of the demand curve and DCS_{\text{full}}, Gruber and Köszegi’s tax result provides an estimate of DCS_{\text{full}} - DCS_{\text{absence}}. Gruber and Köszegi first estimate an internal health cost of $30.45 per pack. From this, they calculate an internality tax that ranges from $0.98 to $2.89 (depending on technical parameters of their model), with an average of $2.17. FDA’s internal health and financial cost estimates differ from Gruber and Köszegi’s in a number of respects, including discount rate and use of a VSLY rather than value of a statistical life approach. We therefore scale the $2.17 internality tax estimate according to the ratio between our internal health and financial cost estimates and the $30.45 result found by Gruber and Köszegi; this produces internality tax estimates ranging from $0.33 to $1.98. Subtracting these values from our estimates of DCS_{\text{full}} yields estimates of DCS_{\text{absence}} ranging from $10.01 to $31.49. Knowing DCS_{\text{absence}} and Q_{\text{absence}}, we can use a Gruber and Köszegi elasticity estimate, – 0.803, to find the height of DCS_{\text{rule}}. This calculation yields estimates of the difference between DCS_{\text{rule}} and DCS_{\text{full}} that range from $0.27 to $1.81. If we assume a linear demand curve (in which case \(F\) will be 50 percent of the sum of \(F\) and \(G\)), this indicates that consumer surplus loss offsets roughly 93 percent of rule-induced internal health benefits. An analogous calculation using the $7.50 per pack tax suggested by Gruber (Ref. 133) indicates that consumer surplus loss offsets roughly 76 percent of rule-induced internal health benefits.

Figures E2 and E3 illustrate the underlying model for the benefits analysis and the uncertainty associated with the changes in consumer surplus resulting from the final rule and other tobacco control policies. The diagrams are elaborations on Figure E1, and lines and areas should be interpreted as discussed in the explanation of that figure. (Full internalization in Figure E2 corresponds to DCS_{\text{full}} in Figure E1; no internalization in Figure E2 corresponds to DCS_{\text{money}} in Figure E1.) Both of the diagrams below show the effects on lifetime smoking of differing degrees of average internalization of the full costs of smoking. Figure E2 shows a rise in the full price (equal to the money price plus the internalized cost), while Figure E3 shows a downward shift in demand equal to the level where all costs are internalized; both diagrams illustrate how the market evolves as it moves leftward from the no-internalization equilibrium to the full-internalization equilibrium. We note that the net internal benefits to smokers of smoking reductions, shown as shaded triangles or trapezoids above the full-internalization demand curve, are the same size in each diagram. Moreover, the area representing benefits decreases in size as the size of the smoking population decreases. We assume that the market is currently at some intermediate point given by the intersection of one of the dashed (partial internalization) price lines with the solid demand curve or the intersection of one of the dashed (partial internalization) demand curves with the solid money price line, but we are not able to definitively estimate where that point is today or where it will be after this final rule takes effect.
VI. Technical Appendix X6: Uncertainty Analysis

Estimation of the effectiveness of the rule (on reducing the future U.S. smoking rate) is subject to a large uncertainty that is not fully reflected in the benefits estimates appearing in the preceding sections, which only reflect different estimates of the VSLY and different discount rates. In this section, we show the uncertainty associated with our estimate of the effectiveness of the rule.

A. Alternative Estimation of Smoking Rate Reduction

Our primary estimate, that the U.S. smoking rate will decrease by 0.088 percentage points, was calculated in the following steps. First, we found the decrease in Canadian smoking rates...
that this proxy is inappropriate. To account for this possibility, we calculate the unexplained difference in Canadian smoking rates before and after graphic warning labels were introduced, this time omitting any U.S. adjustments. We assume that antismoking policies and programs other than the graphic warning labels are incorporated in the pre-2001 trend, with no additional effects of these variables occurring after the introduction of graphic warning labels. This approach indicates that graphic warning labels may have been responsible for a 0.574 percentage point decrease in the Canadian smoking rate. If the rule were to achieve this effectiveness level in the United States, benefits would be approximately six times larger than those reported earlier in this analysis. For example, our benefits estimates calculated with a VSLY of $318,923 and a net-to-gross benefits ratio of 90 percent rise from $1,681.0 million with a 3 percent discount rate and $517.5 million with a 7 percent discount rate (see Table 9b) to $10,916.6 and $3,360.7 million. We use these last two numbers as global upper bounds in Table 1.

Although both of the estimation methods discussed thus far lead to the conclusion that graphic warning labels will reduce smoking rates, FDA has had access to very small data sets, so our effectiveness estimates are in general not statistically distinguishable from zero; we therefore cannot reject, in a statistical sense, the possibility that the rule will not change the U.S. smoking rate. Therefore, the appropriate lower bound on benefits is zero. Ranges of benefits, representing the zero-effect case and the Canada-only modeling approach, appear in Table 49. The wide ranges shown in the table heighten the uncertainty inherent in our approach.

### Table 49.--Ranges of Benefits ($ billion)

<table>
<thead>
<tr>
<th>VSLY=$106,308</th>
<th>VSLY=$212,615</th>
<th>VSLY=$318,923</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>3% Discount Rate</td>
<td>7% Discount Rate</td>
</tr>
<tr>
<td>Present Value</td>
<td>[0, 33.8]</td>
<td>[0, 9.0]</td>
</tr>
<tr>
<td>Annualized Value (Over Twenty Years)</td>
<td>[0, 2.3]</td>
<td>[0, 0.8]</td>
</tr>
</tbody>
</table>

### B. Monte Carlo Simulation

In addition to the uncertainty surrounding the effectiveness of graphic warning labels at reducing smoking rates, the other principal uncertainty in our benefits analysis is the value to smokers of cessation or avoided initiation. As discussed in section XI.D.2, we use two methods and several net-to-gross benefits ratios to produce a range of value estimates. For every percentage point reduction in the national smoking rate, these estimates become $4.2 to $281.6 billion (with a 3 percent discount rate) or $1.3 to $61.1 billion (with a 7 percent discount rate). Similarly, for every percentage point reduction in the national smoking rate, estimates of benefits accruing to the general public (including fire loss and financial effects) range from $6.1 to $14.7 billion (with a 3 percent discount rate) or $4.3 to $11.6 billion (with a 7 percent discount rate). Details appear in Table 50.

### Table 50.--Benefits Ranges, Per Percentage Point Reduction in Smoking Rate ($ mil)

<table>
<thead>
<tr>
<th>Accruing to Dissuaded Smokers:</th>
<th>3% Discount Rate</th>
<th>7% Discount Rate</th>
</tr>
</thead>
<tbody>
<tr>
<td>VSLY=$106,308</td>
<td>VSLY=$212,615</td>
<td>VSLY=$318,923</td>
</tr>
<tr>
<td>Lower Bound</td>
<td>4,191.0</td>
<td>4,191.0</td>
</tr>
<tr>
<td>Upper Bound</td>
<td>96,196.9</td>
<td>188,907.5</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Accruing to General Public:</th>
<th>3% Discount Rate</th>
<th>7% Discount Rate</th>
</tr>
</thead>
<tbody>
<tr>
<td>VSLY=$106,308</td>
<td>VSLY=$212,615</td>
<td>VSLY=$318,923</td>
</tr>
<tr>
<td>Lower Bound</td>
<td>6,097.1</td>
<td>6,982.8</td>
</tr>
<tr>
<td>Upper Bound</td>
<td>12,895.9</td>
<td>13,781.7</td>
</tr>
</tbody>
</table>

We estimate the 90th percentile range for the present and annualized values of total benefits with a Monte Carlo simulation. We model the distribution of the decline in smoking rates with a non-parametric bootstrap, in which we draw from discrete uniform distributions an individual year's United States-Canada adjusted smoking rate difference from the graphic warning label period (in Canada) and an individual year's difference from the pre-graphic warning label period. To account for uncertainty in the value to dissuaded smokers of cessation or avoided initiation, we use for each discount rate and VSLY a uniform distribution running from the lower...
bound estimate to the upper bound estimate, as shown in Table 50. Benefits accruing to the general public are modeled analogously, with a uniform distribution bounded below and above by the values appearing in the table. We run 100,000 iterations for each simulation and report our results in Table 51. Both positive and negative results appear in the table because some paired-year United States-Canada differences show graphic warning labels decreasing the Canadian smoking rate and some paired-year differences show them increasing the smoking rate. (The second finding is almost certainly due to survey noise. More specifically, ordinary sampling variation will cause the percentage of smokers contained in a survey sample to change from one year or country to the next; this is separate from any underlying change in the true smoking rate. Depending on the sizes and directions of the relative changes, a comparison of country-year pairs can show the smoking rate increasing even when it has actually decreased, or vice versa. Because we expect this survey noise to overestimate the smoking rate change in some years and underestimate it in others, in our primary estimate, we take an average over all the years for which we have data in order to estimate as reliably as possible the true underlying change.) The wide differences in benefits shown in the table highlight the uncertainty inherent in our analysis.

<table>
<thead>
<tr>
<th>Present Value</th>
<th>VSLY=$106,308</th>
<th>VSLY=$212,615</th>
<th>VSLY=$318,923</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>3% Discount Rate</td>
<td>7% Discount Rate</td>
<td>3% Discount Rate</td>
</tr>
<tr>
<td>[ -54.0 , 69.4 ]</td>
<td>[ -14.3 , 18.1 ]</td>
<td>[ -100.3 , 127.1 ]</td>
<td>[ -24.2 , 30.9 ]</td>
</tr>
<tr>
<td>Annualized Value (Over 20 Years)</td>
<td>[ -3.6 , 4.7 ]</td>
<td>[ -1.3 , 1.7 ]</td>
<td>[ -6.7 , 8.5 ]</td>
</tr>
</tbody>
</table>

Table 51.—Monte Carlo Simulation Ranges of Benefits ($ billion)