

From	To	MEA	
*9500—MRA *7600—MCA FRIDA, AK FIX, WBND #MEA IS ESTABLISHED WITH A GAP IN NAVIGATION SIGNAL COVERAGE			
<b>§ 95.6440 Alaska VOR Federal Airway V440 Is Amended to Read in Part</b>			
WINOR, AK FIX ..... *9500—MRA *7600—MCA FRIDA, AK FIX, WBND #MEA IS ESTABLISHED WITH A GAP IN NAVIGATION SIGNAL COVERAGE	*FRIDA, AK FIX .....	#10000	
From	To	MEA	MAA
<b>§ 95.7001 Jet Routes</b> <b>§ 95.7537 Jet Route J537 Is Amended to Read in Part</b>			
ROME, OR VOR/DME .....	MULLAN PASS, ID VOR/DME .....	22000	45000

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**SUSQUEHANNA RIVER BASIN COMMISSION**

**18 CFR Part 806**  
**Review and Approval of Projects**

*CFR Correction*

In Title 18 of the Code of Federal Regulations, Part 400 to End, revised as of April 1, 2011, on page 118, in § 806.6, (b)(1)(i) and (ii) are removed.

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**DEPARTMENT OF HEALTH AND HUMAN SERVICES**

**Food and Drug Administration**

**21 CFR Part 558**

**New Animal Drugs for Use in Animal Feeds**

*CFR Correction*

In Title 21 of the Code of Federal Regulations, Parts 500 to 599, revised as of April 1, 2011, on page 490, in § 558.500, (e)(1)(i) is reinstated to read as follows;

**§ 558.500 Ractopamine.**

\* \* \* \* \*

(e) \* \* \*

(1) \* \* \*

Ractopamine in grams/ton	Combination in grams/ton	Indications for use	Limitations	Sponsor
(i) 4.5 to 9 .....	.....	For increased rate of weight gain, improved feed efficiency, and increased carcass leanness in finishing swine, weighing not less than 150 lbs, fed a complete ration containing at least 16% crude protein for the last 45 to 90 lbs of gain prior to slaughter.	Feed continuously as sole ration.	000986

\* \* \* \* \*  
[FR Doc. 2012-5838 Filed 3-8-12; 8:45 am]  
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**DEPARTMENT OF HEALTH AND HUMAN SERVICES**

**Food and Drug Administration**

**21 CFR Part 866**

[Docket No. FDA-2012-N-0165]

**Medical Devices; Immunology and Microbiology Devices; Classification of Norovirus Serological Reagents**

**AGENCY:** Food and Drug Administration, HHS.

**ACTION:** Final rule.

**SUMMARY:** The Food and Drug Administration (FDA) is classifying norovirus serological reagents into class II (special controls). The special control that will apply to these devices is the guidance document entitled “Class II Special Controls Guidance Document: Norovirus Serological Reagents.” The Agency is classifying these devices into class II (special controls) because special controls, in addition to general controls, will provide a reasonable assurance of safety and effectiveness of these devices and there is sufficient information to establish special controls.

**DATES:** *Effective Date:* April 9, 2012. The classification was effective February 23, 2011.

**FOR FURTHER INFORMATION CONTACT:** Steven Gitterman, Center for Devices and Radiological Health, Food and Drug Administration, 10903 New Hampshire Ave., Bldg. 66, Rm. 5518, Silver Spring, MD 20993-0002, 301-796-6694.

**SUPPLEMENTARY INFORMATION:**

**I. Legal Authority**

In accordance with section 513(f)(1) of the Federal Food, Drug, and Cosmetic Act (FD&C Act) (21 U.S.C. 360c(f)(1)), devices that were not in commercial distribution before May 28, 1976, the date of enactment of the Medical Device Amendments of 1976 (the 1976