- To meet the requirements at § 482.42(b)(1), the Joint Commission added a new EP to require that an accredited hospital delineate the responsibilities of the chief medical officer, medical staff, and director of nursing, to ensure that problems identified by the infection control officer are addressed and that corrective action plans are successfully implemented.
- To meet the requirements at § 482.45(b)(2), the Joint Commission added the definition of "organ" to its glossary
- To meet the requirements at § 482.51(a)(4), the Joint Commission added a new EP to address the hospital's responsibility to maintain a roster of practitioners specifying the surgical privileges of each practitioner.
- To meet the requirements at § 482.51(b)(2), the Joint Commission revised its EPs to require an accredited hospital to place a properly executed informed consent form in each patient's chart before surgery, except in emergencies.
- To meet the requirements at § 482.51(b)(3), the Joint Commission added a note to its standards to clarify that the hospital must have the necessary resuscitation equipment available in the operating room.
- To meet the requirements at § 482.52(a), the Joint Commission added a new EP to include the requirements for individuals qualified to administer anesthesia in an accredited hospital.
- To meet the requirements at § 482.52(c), the Joint Commission added a new EP to incorporate the permissive exemption from physician supervision of certified registered nurse anesthetists.
- To meet the requirements at § 482.53(a)(2), the Joint Commission added a new EP to require that an accredited hospital's service director and medical staff approve the qualifications, training, functions, and responsibilities of nuclear medicine personnel.
- To meet the requirements at § 482.53(c)(2), the Joint Commission revised its EPs to require an accredited hospital to inspect, test, and calibrate nuclear medicine equipment annually.
- To meet the requirements at § 482.53(d)(3), the Joint Commission added the definition
- "radiopharmaceuticals" to its glossary.
- To meet the requirements at § 482.54(b)(1), the Joint Commission added a new EP to require that an accredited hospital assign responsibility for outpatient services to one individual.
- To meet the requirements at § 488.55(a)(1) and § 482.55(b)(1), the

- Joint Commission added a new EP to require an accredited hospital's emergency services to be directed and supervised by a qualified member of the medical staff.
- To meet the requirements at § 482.56(a)(2), the Joint Commission revised its EPs to include qualifications for physical therapy, occupational therapy, speech-language pathology, and audiology services when these services are provided by accredited hospitals
- To render a decision regarding the deemed status of an accredited hospital, The Joint Commission revised its accreditation decision letters to ensure that they are accurate and contain all the required elements for the CMS Regional Office.
- To meet the requirements at § 488.28(a), the Joint Commission updated its guidelines for submission of Evidence of Standards Compliance (ESC) to emphasize that the person responsible for implementation of corrective action and assessment of ongoing compliance must be documented in the ESC.
- To clearly identify whether an identified deficient practice represented condition-level or standard-level noncompliance, the Joint Commission modified its survey report.
- To meet the requirements of section 2728 of the SOM, the Joint Commission modified its policies regarding timeframes for sending an ESC.
- To meet the requirements at section 5075.9 of the SOM, the Joint Commission revised its policies to ensure complaint surveys triaged as non-immediate jeopardy (IJ) high and non-IJ medium are conducted within 45 calendar days.
- To meet the survey process requirements in Appendix A of the SOM, the Joint Commission developed a policy outlining the minimum number of inpatient records required for review during a certification survey.
- To meet the requirements at § 488.3(a), section 2026A of the SOM and Appendix A, the Joint Commission developed a new policy to ensure all areas and locations receiving payment under the Medicare's provider agreement are surveyed for compliance with the conditions of participation independently.
- To meet the requirements at section 2700A of the SOM, the Joint Commission revised its survey activity guide to ensure all deemed status surveys are unannounced.
- To meet the requirements at § 489.18 and section 3210 of the SOM, the Joint Commission revised its policies to state that if an organization

acquires a new service, program, or site which requires an extension survey, the survey will be conducted within 6 months, and the results of the survey will immediately impact the accreditation status of the acquiring organization.

To verify the Joint Commission's continued compliance with the provisions of this final notice, we will conduct a follow-up corporate onsite visit and survey observation within 1 year of the effective date of this notice.

### B. Term of Approval

Based on the review and observations described in section III of this final notice, we have determined that the Joint Commission's requirements for hospitals meet or exceed our requirements. Therefore, we approve the Joint Commission as a national accreditation organization for hospitals that request participation in the Medicare program, effective July 15, 2010 through July 15, 2014.

# V. Collection of Information Requirements

This document does not impose information collection and recordkeeping requirements.
Consequently, it need not be reviewed by the Office of Management and Budget under the authority of the Paperwork Reduction Act of 1995 (44 U.S.C. 35).

(Catalog of Federal Domestic Assistance Program No. 93.778, Medical Assistance Program; No. 93.773, Medicare—Hospital Insurance Program; and No. 93.774, Medicare—Supplementary Medical Insurance Program)

Dated: October 15, 2009.

### Charlene Frizzera,

Acting Administrator, Centers for Medicare & Medicaid Services.

[FR Doc. E9–27973 Filed 11–25–09; 8:45 am] **BILLING CODE 4120–01–P** 

# DEPARTMENT OF HEALTH AND HUMAN SERVICES

# Centers for Medicare & Medicaid Services

[CMS-2305-FN]

Medicare and Medicaid Programs; Approval of the Accreditation Commission for Health Care for Deeming Authority for Hospices

**AGENCY:** Centers for Medicare & Medicaid Services, HHS.

**ACTION:** Final notice.

**SUMMARY:** This final notice announces our decision to approve the

Accreditation Commission for Health Care (ACHC) for recognition as a national accreditation program for hospices seeking to participate in the Medicare or Medicaid programs.

**DATES:** *Effective Date:* This final notice is effective November 27, 2009 through November 27, 2013.

FOR FURTHER INFORMATION CONTACT: Cindy Melanson, (410) 786–0310. Patricia Chmielewski, (410) 786–6899.

#### SUPPLEMENTARY INFORMATION

### I. Background

Under the Medicare program, eligible beneficiaries may receive covered services in a hospice provided certain requirements are met. Section 1861 (dd)(1) of the Social Security Act (the Act) establishes distinct criteria for facilities seeking designation as a hospice program. Under this authority, the regulations at 42 CFR part 418 specify the conditions that a hospice must meet in order to participate in the Medicare program, the scope of covered services, and the conditions for Medicare payment for hospice care. Provider agreement regulations are located in 42 CFR part 489 and regulations pertaining to the survey and certification of facilities are located in 42 CFR part 488.

Generally, in order to enter into an agreement, a hospice facility must first be certified by a State survey agency as complying with the conditions or requirements set forth in part 418 of our regulations. Then, the hospice is subject to regular surveys by a State survey agency to determine whether it continues to meet these requirements. There is an alternative, however, to surveys by State agencies.

Section 1865(a)(1) of the Act provides that, if a provider entity demonstrates through accreditation by an approved national accreditation organization that all applicable Medicare conditions are met or exceeded, we would "deem" those provider entities as having met the requirements. Accreditation by an accreditation organization is voluntary and is not required for Medicare participation.

If an accreditation organization is recognized by the Secretary as having standards for accreditation that meet or exceed Medicare requirements, any provider entity accredited by the national accrediting body's approved program would be deemed to meet the Medicare conditions. A national accreditation organization applying for approval of deeming authority under part 488, subpart A must provide us with reasonable assurance that the accreditation organization requires the

accredited provider entities to meet requirements that are at least as stringent as the Medicare conditions.

# II. Deeming Applications Approval Process

Section 1865(a)(3)(A) of the Act provides a statutory timetable to ensure that our review of deeming applications is conducted in a timely manner. The Act provides us with 210 calendar days after the date of receipt of an application to complete our survey activities and application review process. Within 60 days of receiving a completed application, we must publish a notice in the Federal Register that identifies the national accreditation body making the request, describes the request, and provides no less than a 30 day public comment period. At the end of the 210day period we must publish a notice in the **Federal Register** of our approval or denial of the application.

### III. Provisions of the Proposed Notice

On July 24, 2009 we published a proposed notice (74 FR 36720) announcing ACHC's request for initial approval as a deeming organization for hospices. In this notice, we specified in detail our evaluation criteria. Under section 1865(a)(2) of the Act and in our regulations at § 488.4 (Application and reapplication procedures for accreditation organizations), we conducted a review of ACHC's application in accordance with the criteria specified in our regulation, which include, but are not limited to the following:

- An onsite administrative review of ACHC's (1) corporate policies; (2) financial and human resources available to accomplish the proposed surveys; (3) procedures for training, monitoring, and evaluation of its surveyors, (4) ability to investigate and respond appropriately to complaints against accredited facilities; and (5) survey review and decisionmaking process for accreditation.
- A comparison of ACHC's accreditation standards to our current Medicare conditions for participation (CoPs).
- A documentation review of ACHC's survey processes to:
- + Determine the composition of the survey team, surveyor qualifications, and the ability of ACHC to provide continuing surveyor training.
- + Compare ACHC's processes to that of State survey agencies, including survey frequency, and the ability to investigate and respond appropriately to complaints against accredited facilities.
- + Evaluate the ACHC's procedures for monitoring providers or suppliers found to be out of compliance with ACHC

program requirements. The monitoring procedures are used only when ACHC identifies noncompliance. If noncompliance is identified through validation reviews, the survey agency monitors corrections as specified at § 488.7(d).

- + Assess ACHC's ability to report deficiencies to the surveyed facilities and respond to the facility's plan of correction in a timely manner.
- + Establish ACHC's ability to provide us with electronic data and reports necessary for effective validation and assessment of ACHC's survey process.
- + Determine the adequacy of staff and other resources.
- + Review ACHC's ability to provide adequate funding for performing required surveys.
- + Confirm ACHC's policies with respect to whether surveys are announced or unannounced.
- + Obtain ACHC's agreement to provide us with a copy of the most current accreditation survey together with any other information related to the survey as we may require, including corrective action plans.

In accordance with section 1865(a)(3)(A) of the Act, the July 24, 2009 proposed notice (74 FR 36720) also solicited public comments regarding whether ACHC's requirements met or exceeded the Medicare CoPs for hospices. We received no public comments in response to our proposed notice.

## IV. Provisions of the Final Notice

A. Differences Between ACHC's Standards and Requirements and Medicare's Conditions and Survey Requirements

We compared ACHC's accreditation requirements and survey process with the Medicare CoPs and survey process as outlined in the State Operations Manual (SOM). Our review and evaluation of ACHC deeming application, which were conducted as described in section III of this notice yielded the following:

- ACHC modified its survey report to clearly identify whether an identified deficient practice represented condition level or standard level noncompliance.
- ACHC revised it accreditation decision letters to ensure that they are accurate and contain all of the required elements for the CMS Regional Office to render a decision regarding the deemed status of an accredited hospice.
- ACHC modified its policies regarding timeframes for sending and receiving a plan of correction (PoC) in accordance with section 2728 of the SOM.

- To meet the CMS requirements related to a PoC, ACHC amended its policies to ensure approved PoCs contain all elements specified in section 2728 of the SOM.
- To meet the requirements at § 488.28(a) and section 2726 of the SOM, ACHC developed and implemented new policies that require a written PoC for all deficiencies cited.
- ACHC revised its policies to ensure complaints triaged as immediate jeopardy are investigated within 2 business days of receipt in accordance with the requirements at section 5075.9 of the SOM.
- To meet the requirements at § 418.3, ACHC revised its standards to include the definitions used in the Medicare hospice CoPs.
- To meet the requirements at § 418.52(a)(3), ACHC revised its standards to require that the hospice obtain the patient's or patient's representative signature confirming that he or she received a copy of the notice of rights and responsibilities.
- To meet the requirements at § 418.54(c)(8), ACHC revised its standards to require that the comprehensive assessment consider the patient's need for referrals and further evaluation by appropriate health professionals.
- To meet the requirements at § 418.58(d)(1), ACHC revised its standards to include the requirement that the hospice governing body determine the number and scope of performance improvement projects conducted annually.
- To meet the requirements at § 418.110(c), ACHC revised its standards to ensure the hospice maintains a safe physical environment free of hazards for patients, staff and visitors.
- To meet the requirements at § 418.110(m)(15), ACHC revised its standards to require that hospices document in the patient clinical record: the one hour face to face medical and behavioral evaluation if restraint or seclusion is used to manage violent or self-destructive behavior; a description of the patient behavior and intervention used; alternatives or other less restrictive interventions attempted; the patient condition or symptom(s) that warranted the use of restraint and seclusion; and the patient response to the intervention(s) used, including the rationale for continued use of the intervention.

## B. Term of Approval

Based on the review and observations described in section III of this final notice, we have determined that ACHC's requirements for hospices meet or exceed our requirements. Therefore, we recognize ACHC as a national accreditation organization for hospices that request participation in the Medicare program, effective November 27, 2009 through November 27, 2013.

# V. Collection of Information Requirements

This final notice does not impose any information collection and record keeping requirements. Consequently, it does not need to be reviewed by the Office of Management and Budget (OMB) under the authority of the Paperwork Reduction Act of 1995 (44 U.S.C. Chapter 35).

## VII. Regulatory Impact Statement

In accordance with the provisions of Executive Order 12866, the Office of Management and Budget did not review this final notice.

In accordance with Executive Order 13132, we have determined that this final notice will not have a significant effect on the rights of States, local or tribal governments.

**Authority:** Section 1865 of the Social Security Act (42 U.S.C. 1395bb).

Catalog of Federal Domestic Assistance Program No. 93.778, Medical Assistance Program; No. 93.773 Medicare—Hospital Insurance Program; and No. 93.774, Medicare—Supplemental Medical Insurance Program)

Dated: November 5, 2009.

### Charlene Frizzera,

Acting Administrator, Centers for Medicare & Medicaid Services.

[FR Doc. E9–28010 Filed 11–25–09; 8:45 am]  $\tt BILLING\ CODE\ 4120-01-P$ 

## DEPARTMENT OF HEALTH AND HUMAN SERVICES

## National Institutes of Health

# National Institute of General Medical Sciences; Notice of Closed Meeting

Pursuant to section 10(d) of the Federal Advisory Committee Act, as amended (5 U.S.C. App.), notice is hereby given of the following meeting.

The meeting will be closed to the public in accordance with the provisions set forth in sections 552b(c)(4) and 552b(c)(6), Title 5 U.S.C., as amended. The grant applications and the discussions could disclose confidential trade secrets or commercial property such as patentable material, and personal information concerning individuals associated with the grant applications, the disclosure of which would constitute a clearly unwarranted invasion of personal privacy.

Name of Committee: National Institute of General Medical Sciences Special Emphasis Panel; Review of Applications for 'R13' Scientific Conference Grants.

Date: December 10, 2009.

Time: 1 p.m. to 5 p.m.

*Agenda:* To review and evaluate grant applications.

Place: National Institutes of Health, Natcher Building, Room 3AN18, 45 Center Drive, Bethesda, MD 20892. (Virtual Meeting)

Contact Person: Margaret J. Weidman, Ph.D., Scientific Review Officer, Office of Scientific Review, National Institute of General Medical Sciences, National Institutes of Health, 45 Center Drive, Room 3AN18B, Bethesda, MD 20892, 301–594–3663, weidmanma@nigms.nih.gov.

This notice is being published less than 15 days prior to the meeting due to the timing limitations imposed by the review and funding cycle.

(Catalogue of Federal Domestic Assistance Program Nos. 93.375, Minority Biomedical Research Support; 93.821, Cell Biology and Biophysics Research; 93.859, Pharmacology, Physiology, and Biological Chemistry Research; 93.862, Genetics and Developmental Biology Research; 93.88, Minority Access to Research Careers; 93.96, Special Minority Initiatives, National Institutes of Health, HHS)

Dated: November 20, 2009.

### Jennifer Spaeth,

Director, Office of Federal Advisory Committee Policy.

[FR Doc. E9-28413 Filed 11-25-09; 8:45 am]

BILLING CODE 4140-01-P

# DEPARTMENT OF HEALTH AND HUMAN SERVICES

### **National Institutes of Health**

### Center for Scientific Review; Cancellation of Meeting

Notice is hereby given of the cancellation of the Center for Scientific Review Special Emphasis Panel, December 10, 2009, 1 p.m. to December 10, 2009, 4 p.m., National Institutes of Health, 6701 Rockledge Drive, Bethesda, MD 20892 which was published in the **Federal Register** on November 18, 2009, 74 FR 59567.

The meeting is cancelled due to the reassignment of the applications.

Dated: November 20, 2009.

#### Jennifer Spaeth,

Director, Office of Federal Advisory Committee Policy.

[FR Doc. E9–28404 Filed 11–25–09; 8:45 am] BILLING CODE 4140–01–P