care costs, higher quality of care, and improved service offerings. Left unremedied, the proposed acquisition likely would cause anticompetitive harm by enabling UHS to profit by unilaterally raising the reimbursement rates negotiated with commercial health plans. These costs are ultimately borne by consumers in the form of higher premiums, co-pays, and other out-of-pocket costs. The loss of competition also reduces UHS's incentive to improve quality and provide better service. New entry or expansion is unlikely to deter or counteract the anticompetitive effects of the proposed acquisition. While regulatory barriers to opening a new psychiatric facility or unit are lower in Texas and New Mexico than in other states (e.g., there are no Certificate of Need regulations in either state), local zoning regulations, Medicaid and Medicare certifications, and the need to develop strong relationships with local patient referral sources hinder the ability of firms to enter the market. Cuts to Medicaid funding may also affect the financial incentive of a provider to offer inpatient psychiatric services. Thus, it is unlikely that new entry or expansion sufficient to achieve a significant market impact will occur in a timely manner.

IV. The Proposed Consent Agreement

The proposed Consent Agreement wholly remedies the anticompetitive effects in the El Paso/Santa Teresa market by requiring UHS to divest Peak, located in Santa Teresa, New Mexico, and its associated operations and businesses within six months after issuance of the Decision and Order. The potential acquirer of Peak is subject to prior approval of the Commission. The Consent Agreement also provides that, if Peak is not sold to an approved acquirer within six months, a Divestiture Trustee will be appointed and empowered to divest both Peak and Mesilla Valley. The purpose of this provision is to address the uncertainty of whether Peak alone is sufficient to attract an acquirer that would compete as effectively as UHS competed prior to the merger.

Until completion of the requisite divestiture(s), UHS is required to abide by the Order to Hold Separate and Maintain Assets, which includes a requirement that UHS hold Peak separate from its other businesses and facilities, and a requirement to take all actions necessary to maintain the economic viability, marketability, and competitiveness of both the Peak and Mesilla Valley assets. The Consent Agreement also requires UHS to provide transitional services to the approved acquirer for one year, as needed to assist the acquirer with operating the divested assets as a viable and ongoing business. In addition, the proposed order allows the Commission to appoint a Hold Separate Trustee to oversee UHS’s compliance with the Order to Hold Separate and Maintain Assets. Finally, the proposed order contains a ten-year prior notice requirement for acquisitions of acute inpatient psychiatric service providers in the local area, as well as compliance reporting requirements.

The sole purpose of this analysis is to facilitate public comment on the Consent Agreement. This analysis does not constitute an official interpretation of the Consent Agreement or modify its terms in any way.

By direction of the Commission.

Donald S. Clark,
Secretary.

Dated: October 5, 2012.

John Kastenbauer,
Acting Director, Management Analysis and Services Office, Centers for Disease Control and Prevention.

DEPARTMENT OF HEALTH AND HUMAN SERVICES

Centers for Disease Control and Prevention

Advisory Board on Radiation and Worker Health (ABRW/Advisory Board), National Institute for Occupational Safety and Health (NIOSH)

In accordance with section 10(a)(2) of the Federal Advisory Committee Act (Pub. L. 92–463), the Centers for Disease Control and Prevention announces the following meeting of the aforementioned committee:

Time and Date: 11:00 a.m. – 3:00 p.m., November 5, 2012.

Place: Audio Conference Call via FTS Conferencing. The USA toll-free, dial-in number is 1–866–659–0537 and the pass code is 9933701.

Status: Open to the public, but without a verbal public comment period. Written comment should be provided to the contact person below in advance of the meeting.

Background: The Advisory Board was established under the Energy Employees Occupational Illness Compensation Program Act of 2000 to advise the President on a variety of policy and technical functions required to implement and effectively manage the new compensation program. Key functions of the Advisory Board include providing advice on the development of probability of causation guidelines, which have been promulgated by the Department of Health and Human Services (HHS) as a final rule; advice on methods of dose reconstruction, which have also been promulgated by HHS as a final rule; advice on the scientific validity and quality of dose estimation and reconstruction efforts being performed for purposes of the compensation program; and advice on petitions to add classes of workers to the Special Exposure Cohort (SEC).

In December 2000, the President delegated responsibility for funding, staffing, and operating the Advisory Board to HHS, which subsequently delegated this authority to the CDC. NIOSH implements this responsibility for CDC. The charter was issued on August 3, 2001, renewed at appropriate
intervals, most recently, August 3, 2011, and will expire on August 3, 2013.

Purpose: This Advisory Board is charged with (a) providing advice to the Secretary, HHS, on the development of guidelines under Executive Order 13179; (b) providing advice to the Secretary, HHS, on the scientific validity and quality of dose reconstruction efforts performed for this program; and (c) upon request by the Secretary, HHS, advising the Secretary on whether there is a class of employees at any Department of Energy facility who were exposed to radiation but for whom it is not feasible to estimate their radiation dose, and on whether there is reasonable likelihood that such radiation doses may have endangered the health of members of this class.

Matters to be Discussed: The agenda for the conference call includes: SEC Petition for Battelle Laboratories—King Avenue (Columbus, Ohio); Subcommittee and Work Group Updates; SEC Petition Evaluations Update for the December 2012 Advisory Board Meeting; Plans for December 2012 Advisory Board Meeting; and Advisory Board Correspondence.

The agenda is subject to change as priorities dictate. Because there is not an oral public comment period, written comments may be submitted. Any written comments received will be included in the official record of the meeting and should be submitted to the contact person below in advance of the meeting.

Contact Person for More Information: Theodore M. Katz, M.P.A., Executive Secretary, NIOSH, CDC, 1600 Clifton Road NE., Mailstop: E–20, Atlanta, Georgia 30333, Telephone (513)533–6800, Toll Free 1–800–CDC–INFO, Email ocas@cdc.gov.

The Director, Management Analysis and Services Office, has been delegated the authority to sign Federal Register notices pertaining to announcements of meetings and other committee management activities, for both the Centers for Disease Control and Prevention and the Agency for Toxic Substances and Disease Registry.

Dated: October 5, 2012.

John Kastenbauer,
Acting Director, Management Analysis and Services Office, Centers for Disease Control and Prevention.

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BILLING CODE 4163–18–P

DEPARTMENT OF HEALTH AND HUMAN SERVICES

Centers for Medicare & Medicaid Services

[Document Identifier: CMS–10237 and CMS–10137]

Agency Information Collection Activities: Submission for OMB Review; Comment Request

AGENCY: Centers for Medicare & Medicaid Services, HHS.

In compliance with the requirements of section 3506(c)(2)(A) of the Paperwork Reduction Act of 1995, the Centers for Medicare & Medicaid Services (CMS), Department of Health and Human Services, is publishing the following summary of proposed collections for public comment. Interested persons are invited to send comments regarding this burden estimate or any other aspect of this collection of information, including any of the following subjects: (1) The necessity and utility of the proposed information collection for the proper performance of the Agency’s function; (2) the accuracy of the estimated burden; (3) ways to enhance the quality, utility, and clarity of the information to be collected; and (4) the use of automated collection techniques or other forms of information technology to minimize the information collection burden.


In general, coverage for the prescription drug benefit is provided through prescription drug plans (PDPs) that offer drug-only coverage or through Medicare Advantage (MA) organizations that offer integrated prescription drug and health care products (MA–PD plans). PDPs must offer a basic drug benefit. Medicare Advantage Coordinated Care Plans (MA–CCPs) either must offer a basic benefit or may offer broader coverage for no additional cost. Medicare Advantage Private Fee for Service Plans (MA–PFFS) may choose to offer enrollees a Part D benefit. Employer Group Plans may also provide Part D benefits. If any of the contracting organizations meet basic requirements, they may also offer supplemental benefits through enhanced alternative coverage for an additional premium.

Organizations wishing to provide healthcare services under MA and/or MA–PD plans must complete an application, file a bid, and receive final approval from CMS. Existing MA plans may request to expand their contracted service area by completing the Service Area Expansion (SAE) application. Applicants may offer a local MA plan in a county, a portion of a county (i.e., a partial county) or multiple counties. Applicants may offer a MA regional plan in one or more of the 26 MA regions.

Since the publication of the 60-day notice, the information collection request has been revised to provide clarification to applicants, to ensure consistency throughout the entire application, and to reduce confusion among applicants. As a result of those changes, the overall burden associated with the collection has decreased from 22,995 to 21,581 hours. Form Number: CMS–10237 (OCN 0938–0093).

Frequency: Yearly. Affected Public: Private Sector (Business or other for-profits, Not-for-profit institutions).

Number of Respondents: 566. Total Annual Responses: 566. Total Annual Hours: 21,581. (For policy questions regarding this collection contact Barbara Gullick at 410–786–0563. For all other issues call 410–786–1326.)

2. Type of Information Collection Request: Revision of a currently approved collection; Title: Application for New and Expanding Medicare Prescription Drug Plans and Medicare Advantage Prescription Drug (MA–PD), including Cost Plans and Employer Group Waiver Plans; Use: The Medicare Prescription Drug Benefit program was established by section 101 of the Medicare Prescription Drug, Improvement, and Modernization Act of 2003 (MMA) and is codified in section 1860D of the Social Security Act (the Act). Section 101 of the MMA amended Title XVIII of the Social Security Act by redesignating Part D as Part E and inserting a new Part D, which establishes the voluntary Prescription Drug Benefit Program (“Part D”). The MMA was amended on July 15, 2008 by the enactment of the Medicare Improvements for Patients and Providers Act of 2008 (MIPPA), on March 23, 2010 by the enactment of the Patient Protection and Affordable Care Act and on March 30, 2010 by the