Summary

The Project BioShield Act of 2004 (P.L. 108-276) established a 10-year program to acquire civilian medical countermeasures to chemical, biological, radiological, and nuclear (CBRN) agents for the Strategic National Stockpile. Provisions of this act were designed to encourage private companies to develop these countermeasures by guaranteeing a government market for successfully developed countermeasures.

Congress has expressed concern about the implementation of Project BioShield. It has held multiple oversight hearings and considered several pieces of legislation to improve the execution of this program, including the Pandemic and All-Hazards Preparedness Act (P.L. 109-417), H.R. 1089, and H.R. 1684. Stakeholders and policymakers have criticized specific contract award decisions and the rate at which they are made. Additionally, contract awards reported by the Department of Health and Human Services (HHS) do not directly correspond with figures provided in the President’s annual budget documents, which may suggest problems with interagency coordination and communication.

Both the Department of Homeland Security (DHS) and HHS have responsibilities in this program. Funds for this program are appropriated to DHS, while contracts are executed through HHS. The interagency process responsible for deciding which countermeasures to procure has changed multiple times since this program’s inception.

The Homeland Security Appropriations Act, 2004 (P.L. 108-90) provided an advance appropriation of $5.6 billion to acquire CBRN countermeasures over a 10-year period (FY2004-FY2013). This act also limited the amount that could be obligated during specified time periods. The Project BioShield Act of 2004 (P.L. 108-276) assigned the $5.6 billion advance appropriation to Project BioShield countermeasure acquisitions. Two separate rescissions reduced the total amount available for Project BioShield by a total of $25 million. Congress retains the power to make additional appropriations and rescissions to this account.

HHS has awarded Project BioShield contracts for a countermeasures against anthrax, smallpox, botulinum toxin, and radiological or nuclear agents. These awards total approximately $2.331 billion. However, the largest contract, $878 million for an anthrax vaccine, was cancelled in December 2006 for failure to meet a contract milestone. Taking this into account, approximately $1.889 billion remains available for obligation through FY2008 and $4.064 billion available for obligation through the end of the program in FY2013.

This report discusses actions taken by Congress and the Administration that have affected this program, describes the decision-making process for choosing countermeasures, describes the countermeasures for which the Department of Health and Human Services (HHS) has contracted, and discusses accounting discrepancies in Project BioShield budget documents. This report will be updated periodically.
Contents

Overview of Project BioShield ........................................... 1
  Project BioShield Procurement Process .............................. 2
  DHS Roles ................................................................... 2
  HHS Roles .................................................................. 4
  Presidential Roles .......................................................... 6
  Interagency Roles ............................................................ 6

Appropriations, Rescissions, and Future Funding Options ................. 9
  Appropriations ............................................................... 9
  Rescissions ................................................................ 10
  Future Funding Options ................................................... 10

Acquisitions ........................................................................ 12
  Anthrax ...................................................................... 16
    rPA Vaccine .............................................................. 17
    AVA Vaccine .............................................................. 18
    ABthrax ................................................................ 19
    Anthrax Immune Globulin ........................................... 19
  Smallpox .................................................................... 20
  Botulinum Toxin ............................................................ 20
  Radiological and Nuclear Agents ........................................ 21
    Potassium Iodide ......................................................... 21
    Chelators ................................................................. 21

Differences in HHS Contract Awards and Annual Budget Document
  Accounting ................................................................... 22

Remaining Available Funds ................................................... 25

Concluding Observations ...................................................... 26

List of Figures

  Figure 1. Project BioShield Acquisition Process .......................... 3
  Figure 2. Project BioShield Acquisition Activity ........................... 14

List of Tables

  Table 1. Project BioShield Appropriation and Rescissions
          by Years Funding Is Available ........................................ 11
  Table 2. Project BioShield Appropriation and Rescissions
          by Year Money First Becomes Available ........................... 11
  Table 3. HHS Reported Project BioShield Contract Awards ............. 13
  Table 4. Disposition of Project BioShield Special Reserve Fund
          According to the President’s Budget ............................... 23
  Table 5. Comparison of HHS Award Reporting
          and DHS Budget Accounting .......................................... 24
Project BioShield: Appropriations, Acquisitions, and Policy Implementation
Issues for Congress

Following the terrorist attacks of 2001, the federal government determined that it would need additional medical countermeasures (e.g., diagnostic tests, drugs, vaccines, and other treatments) to respond to an attack using chemical, biological, radiological, or nuclear (CBRN) agents. The enactment of the Project BioShield Act of 2004 (P.L. 108-276) was designed to be an important part of federal efforts to obtain new civilian medical countermeasures. It provides countermeasure developers with a guaranteed government market for their products. As Congress continues oversight of federal efforts to protect the United States, the effectiveness and efficiency of the Project BioShield implementation may draw legislative attention.

This report discusses actions taken by Congress and the Administration that have affected this program, describes the decision-making process for choosing countermeasures, describes the countermeasures for which the Department of Health and Human Services (HHS) has contracted, and discusses accounting discrepancies between the President’s Budget and HHS reporting of Project BioShield awards.

Overview of Project BioShield

The Project BioShield Act of 2004 (P.L. 108-276) contains three major provisions. One relaxes some procedures for bioterrorism-related procurement, hiring, and research grant awarding. Another permits the emergency use of countermeasures not approved by the Food and Drug Administration (FDA). The third authorizes a 10-year program to encourage the development and production of new countermeasures for chemical, biological, radiological, and nuclear (CBRN) agents. This last provision is usually referred to as Project BioShield and is the focus of this report.

In contrast to federal programs that directly fund research and development of biomedical countermeasures, Project BioShield is a procurement program. It acts as a guarantee that the federal government will buy successfully developed countermeasures for the Strategic National Stockpile (SNS). It allows the government to enter into contracts to procure countermeasures while they still are in development, up to eight years before product delivery is expected. The government

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1 For brief overview of this program and information on other aspects of this act, see CRS Report RS21507, Project BioShield: Purposes and Authorities, by Frank Gottron.

2 The SNS contains pharmaceuticals, vaccines, medical supplies, and medical equipment to respond to terrorist attacks and other emergencies.
guarantees that it will buy a certain quantity at a specified price, once the countermeasure meets specific requirements. The government pays the agreed-upon amount only after these requirements are met and the product is delivered to the Strategic National Stockpile. If the product does not meet the requirements within the specified time frame, the contract can be cancelled without any payment to the contractor. Thus, Project BioShield is intended to reduce the developer’s market risk; that is, the possibility that no customer will buy the successfully developed product. However, it does not reduce the development risk; that is, the possibility that the countermeasure will fail during development. The Pandemic and All-Hazards Preparedness Act (P.L. 109-417) modified the Project BioShield Act to allow for milestone-based payments for up to half of the total award before countermeasure delivery.

**Project BioShield Procurement Process**

The Project BioShield procurement process requires actions by the Department of Homeland Security (DHS), HHS, and the President, and relies on interagency working groups. Figure 1 illustrates the Project BioShield decision-making and acquisition process.

**DHS Roles.** The first step in the BioShield acquisition process is to determine whether a particular CBRN agent poses a material threat to national security. This analysis, generally referred to as a Material Threat Assessment (MTA), is performed by DHS. Between 30 and 40 subject matter experts are consulted during an MTA.\(^3\) On the basis of this assessment, the DHS Secretary determines whether that agent poses a material threat to national security. The Project BioShield Act of 2004 requires such a written Material Threat Determination (MTD) for procurement using BioShield funds and authorities. This declaration neither addresses the relative risk posed by an agent nor determines the priority of its acquisition. Furthermore, the issuance of an MTD does not guarantee that the government will pursue countermeasures against that agent.

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\(^3\) Dr. John Vitko, Director, DHS Chemical and Biological Division, Public comments at the Project BioShield Stakeholders Meeting, Washington DC, September 25, 2006.
DHS has issued MTDs for 13 agents. These included the biological agents that cause anthrax, multi-drug resistant anthrax, botulism, glanders, melioidosis, tularemia, typhus, smallpox, plague, and the hemorrhagic fevers Ebola, Marburg, and Junin. Additionally, DHS issued a single MTD covering radiological and nuclear agents. According to HHS, the first four MTDs (anthrax, radiological/nuclear agents, botulinum toxin, and smallpox) were completed before or shortly after the enactment of the Project BioShield Act. No other MTDs were issued until

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September 2006, when nine were issued. HHS predicted no additional MTDs would be issued unless “technology advances or if our understanding of the potential threats changes.”

Homeland Security Presidential Directive (HSPD)-10 and HSPD-18 direct DHS to perform additional risk assessments. HSPD-10 directs DHS to develop, and periodically update, risk assessments that include a ranking of relative risks for biological agents. HSPD-10 states that this overall biological agent risk assessment is to be used to prioritize federal government-wide planning and response to the threat of biological agent attacks. The first iteration of this assessment was delivered in 2006. Following its completion, this overall biological agent risk ranking helped determine which agents should have MTAs and MTDs. HSPD-18 requires DHS to develop a comprehensive risk assessment that integrates all CBRN agents into a single ranking of relative risk. This risk assessment is required to be completed by June 1, 2008. HSPD-18 directs that this assessment be used to prioritize CBRN countermeasure research, development, and acquisition.

In addition to making MTDs and performing risk assessments, DHS contributes to the interagency process by developing credible attack scenarios to help establish countermeasure requirements and response planning.

**HHS Roles.** For agents that have received an MTD, HHS assesses the public health consequences of an attack using that agent. This analysis relies on interagency working groups (see below) and is now coordinated by the HHS Office of Public Health Emergency Medical Countermeasures (OPHEMC). OPHEMC is within the Office of the Assistant Secretary for Preparedness and Response (ASPR).

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6 Dr. Gerald Parker, Principal Deputy Assistant Secretary for Preparedness and Response HHS, Testimony before the House Committee on Appropriations, Subcommittee on Labor, Health and Human Services, Education and Related Agencies, March 8, 2007.


9 Dr. John Vitko, Director, DHS Chemical and Biological Division, Public comments at the Project BioShield Stakeholders Meeting, Washington DC, September 25, 2006.

10 These offices have undergone several name changes. The Office the Assistant Secretary for Preparedness and Response (ASPR) was formerly the Office of Public Health Emergency Preparedness (OPHEP), and was renamed pursuant to P.L. 109-417, The Pandemic and All-Hazards Preparedness Act, in December 2006. The name OPHEP was created administratively in August 2004 (69 Fed. Reg. 51679-51680). Prior to that change, the office was called the Office of the Assistant Secretary for Public Health Emergency Preparedness (ASPHEP), pursuant to P.L. 107-188, the Public Health Security and Bioterrorism Preparedness and Response Act of 2002 (67 Fed. Reg. 48903-48905). Briefly, before that change, it had been called the Office of Public Health Preparedness, which was created administratively in January 2002 (67 Fed. Reg. 1980). In July 2006, Office of...
Following this assessment, HHS determines whether this material threat lacks an existing, effective countermeasure and whether a countermeasure should be procured using Project BioShield authorities and funds. If so, the HHS and DHS Secretaries may jointly submit a recommendation for presidential approval to use BioShield funds to acquire such a countermeasure.

The HHS Secretary is also responsible for establishing countermeasure requirements, such as dosage, patient administration method (e.g., injection or pill), minimum effectiveness, and quantity. This process is coordinated by OPHEMC and relies on input from interagency working groups. HHS is responsible for the entire Project BioShield contracting process, including issuing Requests for Information, Requests for Proposals, awarding contracts, managing awarded contracts, and determining whether contractors have met the minimum requirements for payment. OPHEMC maintains a website detailing all Project BioShield solicitations and awards.11

HHS implementation of Project BioShield and its management of the procurement process have been widely criticized.12 These issues provided some of the impetus for creating the Biodefense Advanced Research and Development Authority (BARDA) through the Pandemic and All-Hazards Preparedness Act (P.L. 109-417).13 Despite concerns that OPHEMC was not optimally executing its BioShield responsibilities, HHS has chosen to implement P.L. 109-417 by adding the new BARDA responsibilities and authorities to this office.14 To reflect this increase in responsibilities, HHS also plans to rename OPHEMC as the Biodefense Advanced Research and Development Authority. These new duties include directly

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10 (...continued)
Public Health Emergency Medical Countermeasures, an office within ASPR, was renamed, replacing the name Office of Research and Development Coordination (ORDC, 71 Fed. Reg. 38403-38405). ORDC was created administratively within ASPHEP in December 2002 (67 Fed. Reg. 71568).

11 See online at [http://www.hhs.gov/ophep/ophemc/bioshield/PBSPrctPrjct/]. OPHEMC is also responsible for procurement of countermeasures (e.g., vaccines, antiviral drugs, and tests) for a possible influenza pandemic. Pandemic procurements are not financed through Project BioShield. See CRS Report RS22576 Pandemic Influenza: Appropriations for Public Health Preparedness and Response, by Sarah A. Lister. See also OPHEMC online at [http://www.hhs.gov/aspr/ophemc/PanFlu/procurement_activities/index.html].

12 For examples, see Elizabeth MacDonald and Robert Langworth, “Spore Wars,” Forbes, Vol. 175, No. 12, p. 162, June 6, 2005; and Eric Lipton, “Bid to Stockpile Bioterror Drugs Stymied by Setbacks,” New York Times, September 18, 2006. Additionally, both chambers of Congress have held hearings examining these issues.


funding the advanced development of countermeasures which are not yet deemed eligible for Project BioShield contract awards.15

**Presidential Roles.** Presidential approval is required before HHS enters into any Project BioShield countermeasure procurement contract or issues a call for countermeasure development. The President may only make such approval subsequent to a joint recommendation from the Secretaries of HHS and DHS. The President delegated this approval responsibility to the Director of the Office of Management and Budget.16

The Executive Office of the President also had coordinated the interagency process, largely through the Homeland Security Council (HSC), the National Security Council (NSC), and the National Science and Technology Council (NSTC). This was changed by HSPD-18, which directed the HHS Secretary to lead the interagency process (see below).

**Interagency Roles.** Much of the priority-setting and requirement-determining activities have input from multiple agencies, such as HHS, DHS, Department of Defense, and some of the intelligence agencies. The interagency process has been changed multiple times in the past, most recently by the issuance of HSPD-18 and the enactment of the Pandemic and All-Hazards Preparedness Act (P.L. 109-417).

**Weapons of Mass Destruction Medical Countermeasures Subcommittee.** In the past, the interagency process relied on expertise resident in the Weapons of Mass Destruction Medical Countermeasures (WMD MCM) Subcommittee. As part of the National Science and Technology Council (NSTC), this interagency group predated Project BioShield. The NSTC, a cabinet-level council, acts to coordinate science and technology policy across the federal research and development enterprise.17 The WMD MCM Subcommittee is a part of the NSTC Committee on Homeland and National Security. According to HHS, the charter of the WMD MCM Subcommittee was changed in 2005, and it began reporting to the joint HSC/NSC Biodefense Policy Coordinating Committee.18 According to NSTC, the Subcommittee also continues to remain within NSTC.19 The WMD MCM Subcommittee contains representatives from Centers for Disease Control and Prevention, Food and Drug Administration, National Institutes of Health, DHS, Department of Defense, Department of Agriculture, Nuclear Regulatory Commission, Department of Energy, Department of Veterans Affairs, Environmental

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17 See online at [http://www.ostp.gov/nstc/index.html].


19 See online at [http://www.ostp.gov/nstc/html/_committees.html].
Protection Agency, Homeland Security Council, National Security Council, Office of the Vice President, Office of Science and Technology Policy, Office of Management and Budget, and various intelligence agencies.\textsuperscript{20}

The WMD MCM Subcommittee’s role in the Project BioShield process appears to have been assumed by the Public Health and Emergency Countermeasure Enterprise Governance Board (see below).

\textbf{Public Health and Emergency Medical Countermeasures Enterprise.} The Public Health and Emergency Medical Countermeasures Enterprise (PHEMCE) is an interagency working group that was established in July 2006 during a HHS Office of Public Health Emergency Preparedness reorganization. It is to:

1. define and prioritize requirements for public health medical emergency countermeasures,
2. coordinate research, early and late stage product development and procurement activities addressing the requirements [including BioShield procurement], and
3. set deployment and use strategies for medical countermeasures held in the Strategic National Stockpile.\textsuperscript{21}

PHEMCE is distinct from the HHS Office of Public Health Emergency Medical Countermeasures (OPHEMC). PHEMCE is an interagency working group while OPHEMC resides solely within HHS. However, the Director of OPHEMC is also responsible for coordinating PHEMCE. Neither its establishing regulation nor the PHEMCE strategy\textsuperscript{22} states to whom this interagency group reports nor details its membership.

According to HHS, the WMD MCM Subcommittee’s duties were transferred to the PHEMCE Governance Board.\textsuperscript{23} However, the apparent continuance of the WMD MCM Subcommittee in the NSTC suggests that not all of its duties have transferred to PHEMCE.\textsuperscript{24} It is unclear what effect this transfer of duties from a subcommittee of a Cabinet-level Council to an interagency working group associated with an office under the Assistant Secretary for Preparedness and Response will have.


\textsuperscript{21} HHS, “Office of Public Health Emergency Preparedness; Statement of Organization, Functions, and Delegations of Authority,” 71 Fed. Reg. 38404, July 6, 2006. This regulation establishes the name of this group as the Public Health Medical Countermeasures Enterprise (PHMCE). This name was apparently changed to include “Emergency” in the title, making it the Public Health Emergency Medical Countermeasures Enterprise (PHEMCE). See 71 Fed. Reg. 53097, September 8, 2006.


\textsuperscript{24} See online at [http://www.ostp.gov/nstc/index.html].
on the interagency process and the efficiency of the Project BioShield acquisition process.

**HSPD-18.** Homeland Security Presidential Directive 18 (HSPD-18) was issued on January 31, 2007. When fully implemented, HSPD-18 may change the interagency process described above. HSPD-18 establishes a government-wide strategy for developing and acquiring civilian WMD countermeasures. One of its provisions requires the HHS Secretary to establish an interagency committee to provide advice in setting medical countermeasure requirements and coordinate HHS research, development, and procurement activities.\(^{25}\)

HSPD-18 also requires the HHS Secretary to establish a dedicated strategic planning activity to integrate risk-based requirements across the threat spectrum and of the full range of research, early-, mid- and late-stage development acquisition and life-cycle management of medical countermeasures.\(^{26}\)

The Secretary is to align all relevant HHS programs to support this plan.

These roles are similar to those of PHEMCE whose draft strategy was published prior to the issuance of HSPD-18.\(^{27}\) The final PHEMCE strategy appears to support the interpretation that HHS intends PHEMCE to fulfill the interagency committee and dedicated strategic planning activity requirements of HSPD-18.\(^{28}\) HSPD-18 requires the interagency committee to “apprise” the joint HSC/NSC Biodefense Policy Coordination Committee of countermeasure development and acquisition progress.

**The Pandemic and All-Hazards Preparedness Act.** The Pandemic and All-Hazards Preparedness Act (P.L. 109-417), enacted December 19, 2006, may also affect the Project BioShield interagency decision-making process.\(^{29}\) It gives the HHS Secretary until June 19, 2007 to develop and make public a strategic plan to integrate biodefense and emerging infectious disease requirements with the advanced research and development, strategic initiatives for innovation, and the procurement of... countermeasures.\(^{30}\)

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\(^{30}\) The Pandemic and All-Hazards Preparedness Act (P.L. 109-417), 120 Stat. 2866. 42
This role is similar to that directed by HSPD-18. The finalized *PHEMCE Strategy* and *PHEMCE Implementation Plan* appear to only partially fulfill this requirement in that they address the biodefense plan but do not address emerging infectious diseases. HHS is preparing a separate strategic plan to fulfill the requirements of P.L. 109-417.\(^3^1\)

### Appropriations, Rescissions, and Future Funding Options

#### Appropriations

The Department of Homeland Security Appropriations Act, 2004 (P.L. 108-90) provided an advance appropriation of $5.593 billion to procure civilian medical countermeasures for a 10-year period (FY2004-FY2013).\(^3^2\) This appropriation was enacted October 1, 2003, almost a year before the July 21, 2004 enactment of the Project BioShield Act of 2004 (P.L. 108-276). The appropriations act established the “Biodefense Countermeasures” account for “necessary expenses for securing medical countermeasures against biological terror attacks.”\(^3^3\)

Although all the funds for this account were provided in the 2004 appropriations act, only a portion became available for obligation upon enactment. The Department of Homeland Security Appropriations Act, 2004 specified that no more than $890 million could be obligated in FY2004, and no more than $3.418 billion could be obligated from FY2004 through FY2008 (Table 1). Any money not obligated within these defined periods would remain available through FY2013. Thus, before rescissions were enacted, DHS had $890 million available as budget authority for this account in FY2004. In FY2005, an additional $2.528 billion would have become available. The remaining $2.175 billion would become available in FY2009 (Table 2).

The Project BioShield Act of 2004 (P.L. 108-276) designated the “Biodefense Countermeasures” account established by the Department of Homeland Security Appropriations Act, 2004 (P.L. 108-90) as the special reserve fund for Project

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\(^3^0\) (...continued)

U.S.C. 247d-7e.

\(^3^1\) Dr. Carol Linden, Acting Director for Public Health Emergency Medical Countermeasures Office of the HHS Assistant Secretary for Preparedness and Response, Public Comments at “Implementation of Project BioShield and BARDA: A Dialogue with HHS,” Washington, DC, May 18, 2007.

\(^3^2\) This section was written with the assistance of Bill Heniff Jr., CRS Analyst in American National Government, Government and Finance Division.

BioShield acquisitions. P.L. 108-276 placed additional restrictions on the use of these funds, including requiring a determination that an agent constitutes a material threat to national security, requiring Presidential approval before a countermeasure can be purchased, and restricting these funds to procurements only (i.e., not for administrative costs). It also broadened the types of countermeasures that may be acquired from this account to include those against biological, chemical, radiological, and nuclear agents.

**Rescissions**

Although Congress provided the entire appropriation for the 10-year program, Congress retains the power to increase or decrease the amount available for Project BioShield. Two separate rescissions have removed a total of $25 million from the Project BioShield special reserve fund.

The Consolidated Appropriations Act, 2004 (P.L. 108-199) contained an across-the-board rescission of 0.59%. This rescission applied to the amount of the Project BioShield advance appropriation that became available for obligation in FY2004 (Table 2). This rescission removed $5 million from the amounts available for obligation in FY2004, as well as reducing the total special reserve fund by an equal amount. Thus, the amount available for obligation in FY2004 was reduced from $890 million to $885 million, and the total amount available for FY2004-FY2013 was reduced from $5.593 billion to $5.588 billion (Table 1 and Table 2).

The Consolidated Appropriations Act, 2005 (P.L. 108-447) contained an across-the-board rescission of 0.8%. This rescission applied to the $2.528 billion that became available for obligation in FY2005 (Table 2). This removed $20 million from the amount available for obligation for FY2005-FY2008 as well as reducing the total special reserve fund by an equal amount. Thus, the amount that became available for obligation in FY2005 was reduced from $2.528 billion to $2.508 billion, and the total amount available until FY2013 was reduced from $5.588 billion to $5.568 billion (Table 1 and Table 2).

**Future Funding Options**

Across-the-board rescissions generally only affect those amounts that become available in that fiscal year. Therefore, the special reserve fund is unlikely to be affected by future across-the-board rescissions, except in FY2009, when the remaining $2.175 billion becomes available (Table 2). However, Congress retains the power to make both specific appropriations and rescissions to this account and could thus directly increase or decrease the amount available for Project BioShield obligations.

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34 The Project BioShield Act of 2004 (P.L. 108-276), 118 STAT. 852, (6 U.S.C. 320 (b)).
Table 1. Project BioShield Appropriation and Rescissions by Years Funding Is Available

<table>
<thead>
<tr>
<th></th>
<th></th>
<th></th>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>P.L. 108-90</td>
<td>Appropriation</td>
<td>890</td>
<td>3,418</td>
<td>5,593</td>
</tr>
<tr>
<td>P.L. 108-199</td>
<td>0.59% Rescission</td>
<td>(-5)</td>
<td>(-5)</td>
<td>(-5)</td>
</tr>
<tr>
<td>P.L. 108-447</td>
<td>0.8% Rescission</td>
<td>n.a.</td>
<td>(-20)</td>
<td>(-20)</td>
</tr>
<tr>
<td><strong>Total</strong></td>
<td></td>
<td>885</td>
<td>3,393</td>
<td>5,568</td>
</tr>
</tbody>
</table>

Note: Amounts rounded to nearest million.

Table 2. Project BioShield Appropriation and Rescissions by Year Money First Becomes Available

<table>
<thead>
<tr>
<th></th>
<th></th>
<th></th>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>P.L. 108-90</td>
<td>Appropriation</td>
<td>890</td>
<td>2,528</td>
<td>2,175</td>
</tr>
<tr>
<td>P.L. 108-199</td>
<td>0.59% Rescission</td>
<td>(-5)</td>
<td>n.a.</td>
<td>n.a.</td>
</tr>
<tr>
<td>P.L. 108-447</td>
<td>0.8% Rescission</td>
<td>n.a.</td>
<td>(-20)</td>
<td>n.a.</td>
</tr>
<tr>
<td><strong>Total</strong></td>
<td></td>
<td>885</td>
<td>2,508</td>
<td>2,175</td>
</tr>
</tbody>
</table>

Note: Amounts rounded to nearest million.
Acquisitions

The HHS has reported awarding $2.331 billion worth of Project BioShield contracts (Table 3).  These contracts address four material threats: *Bacillus anthracis* (the bacteria which cause anthrax), smallpox, botulinum toxin, and radiological and nuclear agents. The distribution of contract awards has been uneven between these threats, with $1.429 million against *Bacillus anthracis* (61%), $500 million against smallpox (21%), $364 million against botulinum toxin (16%) and $38 million against radiological and nuclear weapons (2%). While HHS has made additional requests for information from companies developing CBRN countermeasures, none have resulted in contract offers.

On December 17, 2006, HHS terminated an anthrax countermeasure contract for failure to meet a contract milestone. This contract was the first, and largest to date, awarded using Project BioShield funds. This cancellation took place after the preparation of both the HHS’ *Project BioShield Annual Report to Congress* and the President’s FY2008 Budget. Thus, neither of these documents reflect the recovery of these funds. Taking this cancellation into account, the HHS has obligated $1.454 billion to date (Table 3).

Government acquisitions often follow a pattern of gathering information about available products, contract solicitation, award of the contract, and finally product delivery. Figure 2 displays a time line of Project BioShield acquisition activity.

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36 These requests addressed acute radiation syndrome, chemical agents, and a general request for information from all companies developing CBRN countermeasures. For details, see [http://www.hhs.gov/ophep/ophemc/bioshield/PBSPrcrtPrjct/].

# Table 3. HHS Reported Project BioShield Contract Awards

<table>
<thead>
<tr>
<th>Material Threat</th>
<th>Product</th>
<th>Doses (thousands)</th>
<th>Cost ($ millions)</th>
<th>Company</th>
<th>Award Date</th>
</tr>
</thead>
<tbody>
<tr>
<td>Anthrax</td>
<td>rPA vaccine</td>
<td>75,000</td>
<td>877.5</td>
<td>VaxGen, Inc.</td>
<td>11/4/04; Cancelled 12/19/06</td>
</tr>
<tr>
<td></td>
<td>AVA vaccine</td>
<td>10,000</td>
<td>242.7</td>
<td>Emergent BioSolutions (formerly BioPort Corp.)</td>
<td>5/6/05 and 5/5/06</td>
</tr>
<tr>
<td></td>
<td>ABthrax</td>
<td>20</td>
<td>165.2</td>
<td>Human Genome Sciences</td>
<td>6/19/06</td>
</tr>
<tr>
<td></td>
<td>Anthrax Immune Globulin</td>
<td>10</td>
<td>143.8</td>
<td>Cangene Corp.</td>
<td>7/28/06</td>
</tr>
<tr>
<td>Smallpox</td>
<td>MVA vaccine</td>
<td>20,000</td>
<td>500.0</td>
<td>Bavarian Nordic A/S</td>
<td>6/4/07</td>
</tr>
<tr>
<td>Botulinum Toxin</td>
<td>Botulinum Antitoxin</td>
<td>200</td>
<td>362.6</td>
<td>Cangene Corp.</td>
<td>6/1/06</td>
</tr>
<tr>
<td>Radiological/</td>
<td>Potassium Iodide (Liquid)</td>
<td>4,800</td>
<td>17.5</td>
<td>Fleming &amp; Company</td>
<td>3/18/05 and 2/8/06</td>
</tr>
<tr>
<td>Nuclear</td>
<td>Ca-DTPA</td>
<td>395</td>
<td>21.9</td>
<td>Akorn, Inc.</td>
<td>2/13/06</td>
</tr>
<tr>
<td></td>
<td>Zn-DTPA</td>
<td>80</td>
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Total Announced Obligations to Date: $2,331.2 million  
Total Current Obligations: a $1,453.7 million


a. Accounts for cancellation of the rPA vaccine.
Figure 2. Project BioShield Acquisition Activity

A Request for Information (RFI) is a mechanism for the government to determine what products are available or that are under development that might fulfill a specified government need. It can cover a broad area or be narrowly focused. For example, in September 2006, HHS issued an relatively broad RFI to help in “identifying and characterizing the current and projected status of the research and development programs related to CBRN medical countermeasures” (CBRN General in Figure 2). 38 In contrast, an RFI issued in December 2003 focused on a specific type of treatment for a specific disease, anthrax therapeutics, based on antibodies (Figure 2). 39

Agencies can use the information in RFI responses to help shape policy and to help develop requirements for a contract solicitation. However, RFIs do not necessarily lead to contract solicitations. Four of the eight Project BioShield RFIs have not lead to contract solicitations. These RFIs were seeking countermeasures against CBRN in general, nerve agents, one of the two anthrax therapeutic RFIs, and one of the two acute radiation syndrome RFIs (Figure 2). RFIs are also not required before issuing a contract solicitation. Four of the eight contract solicitations did not have an RFI. These contracts were for AVA based anthrax vaccine, botulinum antitoxin, and the radiation treatments Zn- and Ca-DTPA and potassium iodide (KI) (Figure 2). These contract solicitations were for specific products from specific companies and not subject to open competition. 41

Contract solicitations are invitations for companies to submit proposals to provide goods or services to fulfill government needs. Project BioShield solicitations fall into two basic categories, sole source and Requests for Proposals (RFP). The sole source solicitations were for specific products from specific companies and not subject to open competition. Four of the eight contract solicitations were sole source. These are the same four contracts which did not go through the RFI process discussed above, AVA-based anthrax vaccine, botulinum antitoxin, and the radiation treatments Zn- and Ca-DTPA, and KI (Figure 2). Four of the eight contract solicitations were RFPs. Each RFP specified certain characteristics required by the government and multiple companies could submit proposals. The government could then choose the proposal or proposals that best fit its requirements needs or decide that none of the proposals met the minimum requirements. The contract solicitations which went through the RFP process were those seeking an rPA-based anthrax vaccine, an MVA-based smallpox vaccine, and treatments for acute radiation syndrome (Figure 2).

38 HHS, “Medical Countermeasures to Address Chemical, Biological, Radiological and Nuclear Threats,” RFI-OPHEMC-60-03, September 14, 2006.
40 Zinc diethylenetriaminepentaacetate and calcium diethylenetriaminepentaacetate (Zn- and Ca-DTPA) are also known as referred to as chelators due to how they remove radiation from the body, see the section Chelators below. Potassium iodide is often abbreviated KI, its chemical formula.
41 Details of these and all other Project BioShield-related solicitations can be found on the HHS Project BioShield Procurement Activities website, [http://www.hhs.gov/aspr/ophemc/bioshield/procurement_activities/PBSPrcrtPrjct/index.html].
Three of the four RFPs have resulted in contract awards to date, rPA-based anthrax vaccine, anthrax therapeutics, and MVA-based smallpox vaccine. The anthrax therapeutics RFP resulted in contracts with two companies for two different products. The government may decide that none of the companies responding to an RFP have products that meet the government’s minimum requirements. This appears to be the case with the acute radiation syndrome RFP, which was terminated without an award on March 7, 2007.

HHS has awarded ten Project BioShield contracts to six different companies. Of these contracts, four have been completed (two for AVA-based anthrax vaccine, one for the radiation treatments Zn-DTPA and Ca-DTPA, and one of the two for the radiation treatment KI), four remain open (one of the two for the radiation treatment KI, two for anthrax therapeutics, and one for smallpox vaccine), and one was terminated (rPA-based anthrax vaccine). All of the completed contracts resulted from sole source contracting rather than an open bidding RFP process. These completed contracts were for products which required no further development time. It is not clear why HHS chose to acquire these products through the Project BioShield process rather than using the standard process for acquiring similar off-the-shelf products for the Strategic National Stockpile.

Of the ten contracts awarded by HHS, five were for products that required further development: rPA-based anthrax vaccine, smallpox vaccine, botulinum antitoxin, and the two anthrax therapeutics. None of these contracts have yet resulted in deliveries to the Strategic National Stockpile. The rPA anthrax vaccine contract was cancelled and development continues on the remaining four products with open contracts.

**Anthrax**

The Project BioShield countermeasures against anthrax fall into two categories, vaccines and treatments. The vaccines would likely be used after an attack to prevent those people who were exposed to *Bacillus anthracis* from developing the disease anthrax, a procedure called postexposure prophylaxis. This contrasts with the manner in which most vaccines (e.g., childhood vaccines) are administered before exposure.

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rPA Vaccine. The vaccine based on recombinant Protective Antigen (rPA) is often referred to as the “second generation anthrax vaccine,” to differentiate it from the anthrax vaccine adsorbed (AVA) vaccine, which is currently used by the Department of Defense (DOD).\textsuperscript{44} In 2002, the Institute of Medicine (IOM) stated, “Although AVA appears to be sufficiently safe and effective for use, it is far from optimal.”\textsuperscript{45} The IOM supported the development of a new anthrax vaccine. Officials at HHS believe that, when fully developed, the rPA vaccine will address many of the shortcomings of the AVA vaccine as identified in the IOM report.\textsuperscript{46}

In November 2004, HHS awarded VaxGen, Inc. an $877.5 million contract for the delivery of 75 million doses of rPA vaccine to the Strategic National Stockpile ($11.70 per dose). On December 17, 2006, HHS terminated this contract for VaxGen’s failure to meet a contract milestone.\textsuperscript{47}

HHS had planned that each person would require a three dose regimen of this vaccine for protection.\textsuperscript{48} Thus, 75 million doses would be sufficient for 25 million people. The Food and Drug Administration (FDA) has not licensed this vaccine. Although FDA licensing is not required for delivery to the stockpile, this vaccine required additional clinical testing before it could be accepted by the government. Under the contract with VaxGen, delivery was to begin by the end of 2006 and be completed by the end of 2007. Technical difficulties repeatedly delayed delivery.\textsuperscript{49}

This first, largest Project BioShield contract has drawn intense scrutiny. Critics of this contract award point to VaxGen’s previous unsuccessful attempts to develop products, financial difficulties, and problems meeting the contract deadlines as indicative of problems in HHS’ implementation of Project BioShield authorities.\textsuperscript{50} HHS responded to such criticisms by stating VaxGen won the contract through open competition after all the proposals were subjected to “a robust technical and business
evaluation process.” HHS portrayed the delays as part of the normal drug development process. VaxGen reportedly denied responsibility for the delays, stating that they arose from the government changing its requirements.

Following the cancellation of the contract, HHS restated its commitment to obtain an rPA-based anthrax vaccine for the Strategic National Stockpile.

AVA Vaccine. The AVA anthrax vaccine was originally licensed in 1970. It is currently approved by the FDA for use in 18- to 65-year-olds prior to exposure to *Bacillus anthracis* (pre-exposure prophylaxis). Neither this vaccine nor the rPA vaccine is approved by the FDA for post-exposure prophylaxis. The FDA-approved regimen for pre-exposure prophylaxis requires a series of six doses administered over the course of 18 months.

The DOD currently uses this vaccine for troops and other personnel deployed in certain areas, including South Korea, Afghanistan, and Iraq. Complaints of adverse reactions and questions about the vaccine’s efficacy prompted judicial review of its use. In October 2004, a federal judge ordered the DOD to stop mandatory vaccinations pending FDA review. After this order, DOD continued to use this vaccine on a voluntary, rather than mandatory, basis. The FDA completed its review in December 2005. In October 2006, DOD announced plans to resume mandatory vaccinations. Reportedly, several DOD employees plan to sue to block implementation of mandatory vaccinations.

In May 2005 and May 2006, HHS awarded contracts to Emergent BioSolutions (formerly BioPort Corp.) for the delivery of AVA vaccine to the Strategic National Stockpile. Combined, the contracts are for 10 million doses of AVA vaccine for $242.7 million ($24.27 per dose). According to the company, 9 million doses have been delivered to the government, and the remainder is to be delivered in 2007.

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51 Stewart Simonson, Assistant Secretary, Department of Health and Human Services, Office of Public Health and Emergency Preparedness, Testimony before the Senate Committee on Appropriations, Subcommittee on Homeland Security, April 28, 2005.


54 Dr. Gerald Parker, Principal Deputy Assistant Secretary for Preparedness and Response HHS, Testimony before the House Committee on Appropriations, Subcommittee on Labor, Health and Human Services, Education and Related Agencies, March 8, 2007.


This contract award has also drawn criticism on the basis of cost and questions of policy. Despite the manufacturer carrying no developmental risk, the AVA vaccine cost per dose is twice the cost per dose of rPA. Additionally, critics observe that DOD studies indicate that up to 35% of people have adverse reactions to this vaccine and that 6% of vaccine recipients have reported serious complications to the FDA’s Vaccine Adverse Event Reporting System. Critics point to this and observations in the IOM report to support their conclusion that AVA is an inferior product. Lastly, since AVA is the only currently licensed vaccine, critics question whether its acquisition has resulted from its unique status rather than filling a Project BioShield need. Emergent BioSolutions defended its product stating that both the IOM report and the FDA found its product safe and that, as the only FDA-approved anthrax vaccine available, it is filling an urgent need.

**ABthrax.** ABthrax is an antibody-based treatment that works in a manner similar to anti-venom treatments for snake bites. It is currently under development and it is not yet licensed by the FDA. In June 2006, HHS awarded a $165.2 million contract to Human Genome Sciences for the delivery of 20 thousand doses of ABthrax ($8,260 per dose). Human Genome Sciences expects to complete the delivery of ABthrax to the government in 2008. This high cost per dose, the mechanisms of action, and method of patient administration suggest that ABthrax would be used as a treatment for people who have already developed the symptoms of anthrax, rather than as a post-exposure prophylactic.

**Anthrax Immune Globulin.** Anthrax Immune Globulin is also an antibody-based therapeutic. It is derived from the blood of people who have received the anthrax vaccine. It is currently under development and is not yet licensed by the FDA. In July 2006, HHS awarded a $143.8 million contract to Cangene Corp. for the delivery of 10 thousand doses of Anthrax Immune Globulin ($14,380 per dose). This high cost per dose, the mechanism of action, and likely method of patient administration suggest that Anthrax Immune Globulin would be used as a treatment for people who have already developed the symptoms of anthrax, rather than as a post-exposure prophylactic.

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58 (...)continued


Smallpox

Although the World Health Organization eradicated naturally occurring smallpox, it remains a terrorist threat. Following the terrorist attacks of 2001, the United States acquired for the Strategic National Stockpile enough of the currently FDA-licensed vaccine (Dryvax® made by Wyeth Laboratories) to vaccinate 300 million people. However, this vaccine has a high rate of complications, which could be especially serious in people with certain conditions including pregnancy, compromised immune systems, and eczema. The HHS determined that a different smallpox vaccine is required to protect such vulnerable populations.

In June 2007, HHS awarded a $500 million contract to Bavarian Nordic A/S for 20 million doses of smallpox vaccine ($25 per dose), enough for 10 million people. This vaccine is based on the Modified Vaccinia Ankara (MVA) viral strain, which is a different viral strain than the currently licensed vaccine. Experts at HHS believe that this will make it safer for use in vulnerable populations. HHS plans to use this vaccine as a pre-exposure prophylactic in those populations following a known or suspected smallpox release. Additional research is required before this vaccine can be accepted into the stockpile and licensed by the FDA. According to the company, this contract contains options worth up to $1.1 billion for 60 million additional doses and clinical research to extend the license to include children, the elderly, and people infected with HIV.

Botulinum Toxin

Botulinum antitoxin is an antibody-based treatment for botulism, a life threatening illness caused by a toxin produced by Clostridium botulinum bacteria. In June 2006, HHS awarded a $362.6 million contract to Cangene Corp. for 200 thousand doses of a botulinum antitoxin ($1,813 per dose). The company expects to begin delivery by the end of 2007. Botulinum toxin has several different types; an antitoxin against one type will not be effective against other types. This contract calls for a combination of antitoxins that will work against seven types of botulinum toxins. This combination is known as heptavalent antitoxin. Following an intentional release of botulinum toxin, this antitoxin would probably be administered to people who have developed symptoms of toxin exposure, consistent with the way that similar trivalent products are currently used to treat naturally occurring exposures.

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63 President Bush, Remarks by the President on Smallpox Vaccination, December 13, 2002.
64 Dryvax® FDA-approved package insert, June 2003.
Botulinum antitoxin is produced in a manner similar to anthrax immune globulin, except in this case, it is extracted from horse blood instead of human blood. In 2004, after the Department of Homeland Security Appropriations Act, 2004 provided the advance appropriation, but before the Project BioShield Act was enacted, HHS obligated $50 million from this account to support the botulinum antitoxin program. These funds were used to process existing horse blood that had been collected by the DOD and to establish horse farms needed to provide new horse blood. This expenditure would probably not have been eligible for funding from this account after enactment of the Project BioShield Act, as it limited the use of these funds to procuring products. Because these funds were not obligated as part of Project BioShield, they are not included in Table 3, but they are included in Table 4 (see below).

**Radiological and Nuclear Agents**

In addition to direct blast effects, attacks using radiological or nuclear agents can produce injuries resulting from ionizing radiation, which can damage or kill living cells. HHS determined that the threat posed by both acute radiation sickness and internal contamination with radioactive particles require countermeasures. HHS has contracted for two types of countermeasures designed to reduce internal contamination. An RFP for countermeasures to address acute radiation sickness did not lead to a contract award. The RFP was cancelled, apparently because none of the proposals met the minimum requirements determined by HHS.

**Potassium Iodide.** The HHS awarded contracts in March 2005 and February 2006 to Fleming & Company Pharmaceuticals for the delivery of a total of 4.8 million doses of liquid potassium iodide (KI) for a total cost of $15.9 million ($3.31 per dose). This product is FDA-approved and available without a prescription to treat people exposed to radioactive iodine.

Potassium iodide might be distributed following a release of radioactive iodine into the air, possibly following an attack on a nuclear power plant. Because the thyroid gland extracts and stores iodine present in the blood, it is vulnerable to injury from radioactive iodine. If administered in time, potassium iodide would block extraction and storage of radioactive iodine by the thyroid. Potassium iodide does not protect against the effects of any other type of radioactive material. Even before these acquisitions, potassium iodide tablets were included in the Strategic National Stockpile, but the tablet formulation was considered poorly suited for children. This liquid preparation, in contrast, is designed for pediatric use.

**Chelators.** In February and April 2006, HHS awarded a $21.9 million contract to Akorn, Inc. for 395 thousand doses of calcium diethylenetriaminepentaacetate (Ca-DTPA) and 80 thousand doses of zinc diethylenetriaminepentaacetate (Zn-DTPA).

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69 HHS, Personal communication, November 11, 2006.


(a nominal average of $46 per dose). These chelators might be used to treat those exposed to radioactive material through the detonation of a radiological dispersal device (“dirty bomb”), improvised nuclear device, or terrorist attack against stored radioactive material. These products are FDA-approved for this type of internal decontamination.

Radioactive materials that may be inhaled or ingested following a dirty bomb or nuclear attack are treated as minerals in the body. Thus, they enter into biological processes like other minerals and become incorporated into internal organs. Once incorporated, they are very difficult to remove and continue to emit radiation, potentially sickening those exposed. Chelators help remove these radioactive particles from the body by binding to them and facilitating their excretion through normal physiological processes.

**Differences in HHS Contract Awards and Annual Budget Document Accounting**

The Project BioShield special reserve fund, established by the Department of Homeland Security Appropriations Act, 2004, is managed by DHS. In FY2006, the DHS management of this appropriations account passed internally from the Federal Emergency Management Agency to the Preparedness Directorate. However, the contracts obligating the appropriated funds are executed through the HHS OPHEMC.

Table 4 shows the accounting from the President’s annual budget documents. In FY2004, $885 million from the advance appropriation became available for obligation. According to the DHS section of the budget, all available budget authority was obligated in FY2004; no budget authority was carried into the following fiscal year. In FY2005, another $2.508 billion became available for obligation. The budget documents state that $189 million of this was obligated in FY2005, leaving $2.324 billion to be carried over into FY2006. For FY2006, the budget states that $856 million was obligated, leaving $1.468 billion to be carried over into FY2007. DHS anticipates obligations of $1.045 billion in FY2007, leaving only $423 million available for obligation in FY2008. The next part of the advance appropriation does not become available for obligation until FY2009 (see Table 2).

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76 Budget of the United States Government, Fiscal Year 2008 — Appendix, p. 479.
### Table 4. Disposition of Project BioShield Special Reserve Fund According to the President’s Budget

($ in Millions)

<table>
<thead>
<tr>
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<th></th>
<th></th>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>Unobligated Balance from Previous Years</td>
<td>0</td>
<td>0</td>
<td>2,324</td>
<td>1,468</td>
<td>423(^b)</td>
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<tr>
<td>Recovered Obligations from Previous Year</td>
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<td>5</td>
<td>0</td>
<td>0(^b)</td>
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<tr>
<td>New Budget Authority</td>
<td>885(^a)</td>
<td>2,508(^a)</td>
<td>0</td>
<td>0</td>
<td>0</td>
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<tr>
<td>Total Amount Available for Obligation</td>
<td>885</td>
<td>2,513</td>
<td>2,324</td>
<td>1,468(^b)</td>
<td>423(^b)</td>
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<tr>
<td>New Obligations by Fiscal Year</td>
<td>885</td>
<td>189</td>
<td>856</td>
<td>1,045</td>
<td>423</td>
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<tr>
<td>Unobligated Balance Carried Forward</td>
<td>0</td>
<td>2,324</td>
<td>1,468</td>
<td>423(^b)</td>
<td>0(^b)</td>
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<tr>
<td>Cumulative Total of Obligations at End of Fiscal Year</td>
<td>885</td>
<td>1,074</td>
<td>1,930</td>
<td>2,975(^b)</td>
<td>3,398(^b)</td>
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</table>

**Source:** *Budget of the United States Government, Fiscal Year 2006 — Appendix, pp. 513-514; Budget of the United States Government, Fiscal Year 2007 — Appendix, p. 512; Budget of the United States Government, Fiscal Year 2008 — Appendix, p. 479; and CRS calculations.*

\(^a\) Includes rescissions made by P.L. 108-199 and P.L. 108-447. See Table 2.

\(^b\) This figure was estimated by the Administration before the cancellation of the $878 million rPA anthrax vaccine contract.

These figures conflict with totals calculated from the countermeasure awards reported by HHS (Table 3). Table 5 lists all of the contracts that HHS has announced for this account along with their dates of award and fiscal year subtotals.\(^77\)

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\(^77\) HHS, *Project BioShield Annual Report to Congress: July 2004-July 2006*, January 26, 2007. HHS also maintains an updated list of awards on their *Project BioShield: Procurement Activities* website available online at [http://www.hhs.gov/ophep/ophenc/bioshield/PBSPrctPrjct/]. HHS asserts that all of the obligations from this account are detailed on that website. Personal communication with HHS staff, November 20, 2006.
### Table 5. Comparison of HHS Award Reporting and DHS Budget Accounting

<table>
<thead>
<tr>
<th></th>
<th>HHS announcements ($ millions)</th>
<th>DHS' Obligations in President’s Budget ($ millions)</th>
<th>Difference ($ millions)</th>
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</thead>
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<tr>
<td>FY04</td>
<td>Botulinum Antitoxin Program&lt;sup&gt;a&lt;/sup&gt; 50</td>
<td>885</td>
<td>-835</td>
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<tr>
<td><strong>FY04 Total</strong></td>
<td>50</td>
<td>885</td>
<td>-835</td>
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<td>FY05</td>
<td>rPA, 11/04 878</td>
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<td>123</td>
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<td>KI, 3/05 123</td>
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<td></td>
<td>AVA, 5/05 123</td>
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<tr>
<td><strong>FY05 Total</strong></td>
<td>1,008</td>
<td>189</td>
<td>819</td>
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<td>FY06</td>
<td>KI, 2/06 10</td>
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<td>Chelators (Zn- and Ca-DTPA), 2/06 22</td>
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<td>AVA, 5/06 120</td>
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<td></td>
<td>Botulinum Antitoxin, 6/06 363</td>
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<td>ABthrax, 6/06 165</td>
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<td>AIG, 7/06 144</td>
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<td><strong>FY06 Total</strong></td>
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<td><strong>Grand Total</strong></td>
<td>1,882</td>
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</table>


**Note:** Amounts rounded to nearest million.

<sup>a</sup> HHS states that this obligation occurred in FY2004 and does not provide a precise date.

According to HHS, the only obligation from this account in FY2004 was $50 million to support the botulinum antitoxin program. In contrast, the President’s FY2006 Budget documents state that $885 million was obligated in FY2004. Additionally, it describes this obligation as falling under two object classifications; with $190 million for “other services” (object classification 25.2) and $695 million for “other purchases of goods and services from Government accounts” (object
classification 25.3). It is not clear what these amounts represent. The $50 million HHS obligated for the botulinum antitoxin program support could fall under the “other services” category, since it was not an acquisition per se, but the amount of this contract does not correlate to the amount categorized as “other services.”

Another possibility is that President’s Budget accounted for the rPA vaccine contract (awarded in November 2004) in FY2004 rather than FY2005. This interpretation is supported by the FY2007 Budget reporting that only $189 million was obligated in FY2005. However, the total of the $878 million rPA obligation and the $50 million botulinum antitoxin program obligation is greater than the budget authority made available in FY2004 ($885 million). This interpretation also would not account for the division of the funds into the two object classifications. Furthermore, the FY2007 DHS Preparedness Directorate BioDefense Countermeasures Congressional Justification materials list acquiring the rPA vaccine as one of its FY2005 accomplishments. The source of the FY2004 account discrepancy of $835 million is not apparent.

In FY2005, HHS reported awarding three contracts for a total of $1.008 billion. The FY2007 Budget states that the actual amount obligated in FY2005 was $189 million. The DHS FY2007 Congressional Justification documents state that its FY2005 accomplishments include the rPA, KI, and AVA contracts. These would equal the $1.008 billion calculated from the HHS figures. It is not apparent to what the $189 million stated in the Budget correlates.

Like the preceding two years, the stated obligations for FY2006 are different according to HHS and the President’s Budget. For FY2006, HHS reported awarding six contracts, with obligations totaling $824 million. This is $32 million less than the $856 million stated as “actual obligations” in FY2006 in the President’s FY2008 Budget.

Combining all of the differences in reporting through FY2006, the President’s Budgets state that $48 million more have been obligated than the HHS documents report.

### Remaining Available Funds

Effective management and Congressional oversight of Project BioShield require specific and clear knowledge of the funds remaining available. For the Administration to most effectively plan and prioritize future acquisitions, it must know the amount of funds remaining available. For Congress, knowing the amount of funds remaining can be important in assessing program management, the implementation pace, and general program effectiveness. Due to conflicting

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80 Budget of the United States Government, Fiscal Year 2008 — Appendix, p. 479.
statements from executive branch agencies, the amount of funds remaining available for obligation for this program is not clear.

According to HHS, as of June 2007, it has obligated $2.331 billion from this account. This figure does not include the $878 million that should be recovered in FY2007 from the cancellation of the rPA anthrax vaccine contract. Taking this recovery into account, $1.889 billion would be available for obligation in FY2007-FY2008 and $4.064 billion would be available until the end of the program in FY2013. As stated above, using the President’s Budget figures to calculate obligations would reduce these numbers by $48 million.

Concluding Observations

Project BioShield plays a key role in the federal government’s response to the threat of chemical, biological, radiological, and nuclear terrorism. It created a process for the government to agree to purchase countermeasures while they still are in development. In addition to increasing the holdings of the Strategic National Stockpile, it was hoped that this government market guarantee would encourage companies to continue to develop promising countermeasures that they might have otherwise abandoned, and induce other companies to begin countermeasure development. It remains unclear how well Project BioShield is meeting these goals.

Many stakeholders, industry leaders, and policymakers have criticized the rate at which DHS completes Material Threat Determinations. To address these concerns, legislation has been introduced in the previous and current Congresses. In the 110th Congress, the Project BioShield Material Threats Act of 2007 (H.R. 1089, Langevin) and the Department of Homeland Security Authorization Act for Fiscal Year 2008 (H.R. 1684, Thompson) would require an assessment, and an MTD if appropriate, for all currently known CBRN agents likely to pose a significant national security threat. These assessments would be required to be completed by December 31, 2007. By assessing all known threats and issuing those MTDs necessary, the full spectrum of material threats may be considered when developing a countermeasure acquisition strategy. Such a comprehensive acquisition strategy may allow for more efficient prioritization and balance of countermeasures, providing optimized protection from CBRN attacks using finite funds in the shortest time. Since HHS has not issued contracts for the all of the agents that already have MTDs, an increase in this number may not increase the rate of countermeasure awards. However, HHS has predicted that no additional MTDs would be issued unless “technology advances or if our understanding of the potential threats changes.”


Appropriators set limits on how much could be obligated during specified periods of time. The pace by which HHS awards countermeasure contracts roughly corresponds to these limits. By this criterion, this program is on track to fulfill its goals; HHS cannot obligate the money faster than it becomes available.

Stakeholders, industry leaders and policymakers have criticized HHS for some of the countermeasures it has chosen. In decisions as complicated and weighty as these, any choice is likely to be criticized. Given the failure of the largest contract to date, some critics may conclude that Project BioShield has fallen short of its goals, since the majority of the money that has been obligated, though not the majority of contracts, has not yet resulted in products in the stockpile. However, one of the unique features of Project BioShield contracts is that the government may contract for products that require up to eight years more of development. It was designed to allow the government to promise to buy something, but only pay for it on delivery. Thus the company, rather than the government, bears the majority of the development risk, i.e. that the product will never be deliverable. One industry group estimates that more than half of all pharmaceuticals will fail during the last eight years of development. Thus, it may be expected that at least some Project BioShield contracts will be cancelled. The government bears some development risk in the form of opportunity costs since the money available for obligation is finite, i.e., money obligated to a countermeasure that will ultimately fail in development cannot be simultaneously obligated to another needed countermeasure.

It is possible that the establishment of the Biodefense Advance Research and Development Authority (BARDA) in HHS will reduce the likelihood that future Project BioShield contracts will fail during the advanced development phase. Established by the Pandemic and All-Hazards Preparedness Act (P.L. 109-417), one of BARDA’s roles is to support the advanced research and development of promising countermeasures. In theory, funding this part of the development process through such a dedicated mechanism could allow countermeasures to further mature through the development process longer before competing for a Project BioShield contract. This could reduce the risk that a countermeasure will fail while under a Project BioShield contract. P.L. 109-417 included authorization for approximately $1 billion to support this type of activity for FY2007 through FY2008. Although Congress did not appropriate money for BARDA in FY2007, the U.S. Troop Readiness, Veterans’ Care, Katrina Recovery, and Iraq Accountability Appropriations Act, 2007 (P.L. 110-28) transferred $99 million from National Institutes of Health accounts to fund BARDA. Even if BARDA becomes operational in FY2007, it will take some time to determine what projects to fund, provide funding, and receive returns on this investment. It remains to be seen how HHS’ decision to combine BARDA with the HHS office responsible for executing Project BioShield (Office of Public Health Emergency Medical Countermeasures) will affect the execution of both programs.

Additional criticism of the Project BioShield procurement process may stem from the perceived opacity of its decision-making process. HHS is moving to address some of these issues by publishing its PHEMCE Strategy for Chemical,
Biological, Radiological and Nuclear Threats, inviting public comment, and reaching out to the public and companies that might develop needed countermeasures through stakeholder meetings.

Some critics also suggest that the Project BioShield process has been poorly managed overall. Such suggestions are reinforced by the annual accounting discrepancies between HHS and DHS. It remains to be seen whether these concerns will be allayed through the management changes being implemented subsequent to: the establishment of the Public Health and Emergency Medical Countermeasures Enterprise (PHEMCE) and publication of its strategy; the enactment of the Pandemic and All-Hazards Preparedness Act (P.L. 109-417); and the issuance of HSPD-18.

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