COMPARATIVE EFFECTIVENESS RESEARCH

Activities Funded by the Patient-Centered Outcomes Research Trust Fund
**Why GAO Did This Study**

In 2010, the Patient Protection and Affordable Care Act (PPACA) authorized the establishment of the Patient-Centered Outcomes Research Institute (PCORI) to carry out comparative effectiveness research (CER) and improve its quality and relevance. PPACA also established new requirements for the Department of Health and Human Services (HHS) to, among other things, disseminate findings from federally funded CER, including findings published by PCORI; and coordinate with relevant federal health programs to build data capacity for this research. To fund CER activities, PPACA established the Trust Fund from which PCORI and HHS are expected to receive an estimated $4.0 billion from fiscal years 2010 through 2019.

PPACA included a provision for GAO to review PCORI’s and HHS’s use of the Trust Fund. This report examines (1) PCORI’s use of the Trust Fund for CER activities, including the dissemination and use of research findings; and (2) HHS’s use of the Trust Fund for these activities.

GAO examined PCORI and HHS documents and data related to use of the Trust Fund, such as commitment, obligation, and expenditure data; PCORI’s audited financial statements; and descriptions of CER activities. GAO also interviewed PCORI and HHS officials responsible for planning and carrying out CER activities and interviewees from stakeholder organizations representing potential users of CER, including public and private payer organizations, provider organizations, and patient organizations. PCORI and HHS provided technical comments, which GAO incorporated as appropriate.

**What GAO Found**

The Patient-Centered Outcomes Research Institute (PCORI) made about $2 billion in commitments for awards in fiscal years 2010 through 2017. PCORI is a federally funded, nonprofit corporation established to carry out and improve comparative clinical effectiveness research (CER), which evaluates and compares the health outcomes and the clinical effectiveness, risks, and benefits of two or more medical treatments, services, or items. PCORI provides funding through award commitments from the Patient Centered Outcomes Research Trust Fund (Trust Fund) and may pay these awards over multiple years. Of the $2 billion PCORI committed as of the end of fiscal year 2017, about $1.6 billion (or 79 percent of its commitments) is for research awards, and $325 million (or 16 percent) is for building the capacity to use existing health data for research. Through fiscal year 2017, commitments for dissemination and implementation awards—intended to share CER findings with potential users of this research—were limited because most PCORI-funded research was still underway. PCORI projects to commit an additional $721 million for awards in fiscal years 2018 through 2021. In addition to awards, PCORI spent $310 million on program and administrative support services in fiscal years 2010 through 2017 and projects to spend an additional $206 million for these services through fiscal year 2024.

**PCORI’s Actual and Projected Commitments and Expenditures, Fiscal Years 2010 through 2024 (Dollars in Millions)**

<table>
<thead>
<tr>
<th>Year</th>
<th>Research award commitments</th>
<th>Building data capacity award commitments</th>
<th>Engagement and workforce award commitments</th>
<th>Dissemination and implementation award commitments</th>
<th>Program and administrative support expenditures</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Actual, fiscal years 2010 through 2017</strong></td>
<td>514</td>
<td>70</td>
<td>46</td>
<td>1,591</td>
<td>325</td>
</tr>
<tr>
<td><strong>Projected, fiscal years 2018 through 2024</strong></td>
<td>91</td>
<td>91</td>
<td>206</td>
<td>310</td>
<td>12</td>
</tr>
</tbody>
</table>

Source: GAO analysis of Patient-Centered Outcomes Research Institute (PCORI) information | GAO-18-311

Note: Amounts listed are current as of Jan. 5, 2018.

From fiscal years 2011 through 2017, the Department of Health and Human Services (HHS) obligated about $448 million from the Trust Fund. Of this amount, HHS obligated about $260 million (or 58 percent of all obligations) to the dissemination and implementation of CER findings. As most PCORI-funded CER had not yet been completed due to the time needed to conduct this research, HHS efforts focused instead on the dissemination and implementation of CER funded by other federal entities. Additionally, HHS obligated funds for efforts to train researchers on conducting CER, build data capacity, and on administrative activities. HHS projects to obligate an additional $120 million for these activities in fiscal years 2018 through 2020.
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Abbreviations

AHRQ  Agency for Healthcare Research and Quality
ASPE  Office of the Assistant Secretary for Planning and Evaluation
CDC  Centers for Disease Control and Prevention
CDS  clinical decision support
CER  comparative clinical effectiveness research
HHS  Department of Health and Human Services
NIH  National Institutes of Health
PCORI  Patient-Centered Outcomes Research Institute
PCRF  People-Centered Research Foundation
PPACA  Patient Protection and Affordable Care Act
Treasury  Department of the Treasury
Trust Fund  Patient-Centered Outcomes Research Trust Fund

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Congressional Committees

Clinical research is often conducted to evaluate the effectiveness of a specific treatment for a disease or condition; however, research is less often conducted to compare the relative effectiveness of two or more available treatment options. Comparative clinical effectiveness research (CER) evaluates and compares the health outcomes and the clinical effectiveness, risks, and benefits of two or more medical treatments, services, or items. For example, a recent CER study found that the life expectancy of women over age 50 with breast cancer in one breast, but who do not carry the breast cancer gene mutation, is similar whether they have one or both breasts removed.\(^1\) Another recent study compared the effectiveness and safety of treatments for childhood anxiety disorders and found that a combination of medications and therapy is likely more effective than either treatment alone.\(^2\) According to the National Academies of Medicine, CER can help clinicians, patients, payers, and others make informed decisions to improve health care.\(^3\)

In 2010, the Patient Protection and Affordable Care Act (PPACA) authorized the establishment of the Patient-Centered Outcomes Research Institute (PCORI) as a federally funded, nonprofit corporation to improve the quality and relevance of CER.\(^4\) In addition, PPACA


\(^3\)National Academy of Medicine (formally the Institute of Medicine), *Initial National Priorities for Comparative Effectiveness Research* (Washington, D.C.: June 2009).

\(^4\)Pub. L. No. 111-148, §§ 6301(a), 10602, 124 Stat. 119, 727, 1005 (codified at 42 U.S.C. § 1320e). PCORI’s patient-centered outcomes research builds upon the definition for CER previously established by the Federal Coordination Council for Comparative Effectiveness Research. See Department of Health and Human Services, *Federal Coordinating Council for Comparative Effectiveness Research: Report to the President and the Congress* (Washington, D.C.: June 30, 2009). For the purposes of this report, we will refer to the work PCORI and HHS conduct as CER.
established new requirements for the Department of Health and Human Services (HHS) to, among other things, broadly disseminate findings from federally funded CER, including findings published by PCORI; train researchers on CER methodological approaches; and coordinate relevant federal health programs to build data capacity for this research. To fund PCORI’s and HHS’s CER activities, PPACA established the Patient-Centered Outcomes Research Trust Fund (Trust Fund) through which PCORI and HHS expect to receive an estimated total of about $4 billion from fiscal years 2010 through 2019.

PPACA included a provision for us to report on the use of the Trust Fund for CER activities. In this report, we examine:

1. PCORI’s use of the Trust Fund for CER activities, including the dissemination and use of research findings, and
2. HHS’s use of the Trust Fund for these activities.

To examine PCORI’s and HHS’s use of the Trust Fund for CER activities, including the dissemination and use of research findings, we reviewed PCORI award commitment and expenditure data and HHS data on obligated funds. We also reviewed descriptions of PCORI’s and HHS’s CER activities, relevant legislation, and previous GAO reports. Furthermore, we conducted interviews with PCORI and HHS officials responsible for planning and carrying out CER activities. Within HHS, we interviewed officials from the Agency for Healthcare Research and Quality (AHRQ), the Office of the Assistant Secretary for Planning and Evaluation

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5PPACA included provisions that we review PCORI’s research activities by 2015 and its use of federal funding by 2018. We completed the first of these reviews in 2015. See GAO, Comparative Effectiveness: Initial Assessment of the Patient-Centered Outcomes Research Institute, GAO-15-301 (Washington, D.C.: March 9, 2015). In addition, PPACA requires the Comptroller General to appoint members to serve on PCORI’s Board of Governors and Methodology Committee.

6According to PCORI, “commitments” represent the amount of funding PCORI intends to award or has awarded to contractors. HHS obligations in a given year represent a definite commitment of funds that create a legal liability for the payment of goods and services ordered or received, although funds may be expended in subsequent years.

We also interviewed officials coordinating PCORI’s PCORnet initiative, the National Patient-Centered Clinical Research Network, to build data capacity for research. We reviewed funding from fiscal year 2010, the first year funding was made available, through 2017, the most recent data available at the time of our analysis. To the extent available, we also reviewed the amount of funding projected beyond fiscal year 2017. Finally, to gather the views of potential users of CER findings, including public and private payers, providers, and patients, we interviewed officials from several large organizations that broadly represent each group of potential users.

To assess the reliability of PCORI commitment and expenditure data and HHS obligation data, we collected information from PCORI and HHS officials regarding the reliability of the data, including the accuracy of data entry and the systems that contain the data. We also reviewed PCORI’s annual audited financial statements. On the basis of these steps, we determined the data were sufficiently reliable for the purposes of our reporting objectives.

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8HHS conducts CER activities through AHRQ and ASPE. AHRQ’s mission is to produce evidence to make health care safer, higher quality, more accessible, equitable, and affordable, and to work within HHS and with other partners to make sure that the evidence is understood and used. AHRQ also coordinates with NIH on certain aspects of its work, such as disseminating research findings. ASPE advises the Secretary of HHS on policy development in health, disability, human services, data, and science, and provides advice and analysis on economic policy. ASPE leads special initiatives, coordinates the department’s evaluation, research and demonstration activities, and manages cross-department planning activities such as strategic planning, legislative planning, and review of regulations.

9PCORnet is a PCORI-funded initiative that supports the translation of existing clinical, patient, and health plan networks’ data into a common data model resulting in the availability of a nationally representative sample of over 128 million individuals that can be used in randomized clinical trials, large observational studies, and other research to make it faster, easier, and less costly to conduct clinical research. Its Coordinating Center leads data and engagement activities, connects with research partners, and supports the PCORnet infrastructure.

10Specifically, to gather the views of public and private payers, we interviewed officials from the Medicaid Medical Directors Network and America’s Health Insurance Plans (a trade group representing health insurers), as well as officials from two health plans. Likewise, to gather the views of providers, we interviewed officials from the American Medical Association and the American College of Physicians. Finally, to gather the views of patients, we interviewed officials from the National Health Council and the Partnership to Improve Patient Care.
We conducted this performance audit from May 2017 to March 2018 in accordance with generally accepted government auditing standards. Those standards require that we plan and perform our work to obtain sufficient, appropriate evidence to provide a reasonable basis for our findings and conclusions based on our audit objectives. We believe that the evidence obtained provides a reasonable basis for our findings and conclusions based on our audit objectives.

In 2010 PPACA authorized the establishment of PCORI to improve CER quality and relevance. PPACA also established requirements for HHS to, among other things, disseminate findings from federally funded CER, including findings published by PCORI, and coordinate with relevant federal health programs to build data capacity for research. PPACA established a Trust Fund to fund these CER activities by PCORI and HHS through fiscal year 2019.

PPACA authorized the establishment of PCORI as a federally funded, nonprofit corporation aimed at advancing the quality and relevance of evidence through research to help patients, clinicians, purchasers, and policy-makers to make informed health care decisions. PCORI is required to identify research priorities, establish a research project agenda, fund research consistent with its research agenda, and disseminate research findings, among other responsibilities. In 2015 we reported that PCORI had conducted activities consistent with its legislative requirements. For example, we reported that since its inception in 2010, PCORI established and implemented priorities for funding CER and related activities, developed plans to disseminate funded research and track its utilization, and took steps to make its research more centered on outcomes prioritized by patients. Further, PCORI developed PCORnet as a distributed research network initiative.

11While clinical research is often conducted to evaluate the effectiveness of a specific treatment for a disease or condition, research is less often conducted to compare among available treatment options. For example, pharmaceutical companies conduct clinical trials to evaluate the safety and effectiveness of a new drug to treat a specific disease or condition. However, prior to federal funding for CER, there was little funding available for research to determine whether one drug is safer or more effective than another.

12See GAO-15-301.
that enables electronic health-related data from multiple sources to be available for research.¹³

PPACA requires HHS to perform several requirements related to CER, which it has implemented through AHRQ and ASPE.¹⁴ Specifically, AHRQ is required to disseminate and support the incorporation of CER funded by PCORI and other federal entities, as well as to foster capacity for conducting CER by supporting training in the methods used to conduct such research. ASPE, in turn, is required to build data capacity for conducting CER. In 2015, we reported that AHRQ had taken some steps to disseminate research findings, but had not taken other actions to help it fully address its dissemination requirements. Furthermore, we reported that ASPE coordinated among various agencies to fund projects intended to build data capacity for research, but that its approach lacked key elements—such as defined objectives, milestones, and time frames—that are necessary to ensure effectiveness. In our 2015 report, we made five recommendations to HHS to direct AHRQ and ASPE to address these issues, as appropriate. HHS concurred with these recommendations and

¹³According to PCORI officials, PCORnet was developed to take existing electronic health-related data from multiple sources and translate certain fields of that disparate data into a common data model, so that the translated data can be used to conduct research.

¹⁴PPACA enacted the new CER requirements by adding section 937 to Title IX of the Public Health Service Act. Pub. L. No. 111-148, § 6301(b), 124 Stat. 119, 738-740 (codified at 42 U.S.C. § 299b-37). Section 937 requires HHS’s AHRQ to carry out specified functions with respect to the dissemination of CER, incorporation of research findings, collection of feedback, and the establishment of a grant program to train researchers, which AHRQ has carried out in consultation with NIH. Section 937(a)-(c), (e). In contrast, section 937 does not specify a particular HHS entity to carry out the functions specified under subsection (f) related to the coordination of federal programs to build data capacity for CER. The Secretary of HHS has delegated responsibility for carrying out section 937(f) to HHS’s ASPE.
PPACA Funding for CER

PPACA established the Trust Fund through which PCORI and HHS receive funds for CER activities. The law provides that for fiscal years 2010 through 2019, the Trust Fund will receive appropriations from the general fund of the Treasury, transfers from the Medicare trust funds, and fees collected by the Department of the Treasury (Treasury) from private insurance and self-insured health plans. Eighty percent of the amounts in the Trust Fund must be made available to PCORI in fiscal years 2011 through 2019, and Treasury must transfer the remaining 20 percent to the Secretary of HHS in each of those years. Under current law, appropriations and transfers to the Trust Fund will end in fiscal year 2019. The law also provides that no amounts shall be available for expenditure from the Trust Fund after September 30, 2019, and specifies that any amounts remaining in the Trust Fund after that time will be transferred to the general fund of the Treasury. (See fig. 1 for an overview of transfers to the Trust Fund and distribution of funds to PCORI and HHS).

15See GAO-15-280. To address our recommendations, AHRQ developed time frames for the implementation and distribution of marketing plans and informational tools, increased dissemination efforts to health plans and vendors of health information technology, and developed plans for collaborating with NIH on dissemination efforts. ASPE implemented a monitoring system to track progress toward milestones and deliverables on its projects to build data capacity. As of January 2018, AHRQ was in the process of implementing the remaining recommendation, which directs AHRQ to document and complete plans to develop a publicly available CER database, including plans to meet the needs of various potential users in the general public. While AHRQ has developed a website that includes instructions and links for searching multiple different publicly available databases that have information on CER, AHRQ has not yet provided information on how it plans to meet the needs of various potential users in the general public, as specified in our recommendation.

16The law provides that amounts transferred to HHS will remain available until expended. For fiscal year 2010, all amounts appropriated to the Trust Fund were made available to PCORI.

17PCORI plans to draw down all remaining funds in the Trust Fund prior to this date to have funds available for anticipated expenditures after fiscal year 2019.
PPACA Limitations on Use of CER

PPACA limits the use of CER in certain ways; for example, the law prohibits PCORI from developing or using a dollars-per-quality adjusted life-year to establish what type of health care is cost effective or recommended, and prohibits the Secretary of HHS from using such measures as a threshold to determine coverage, reimbursement, or incentive programs under Medicare. HHS may use CER findings to help inform Medicare coverage decisions, but PPACA does not allow Medicare coverage to be denied solely on the basis of CER findings.

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18. 42 U.S.C. § 1320e-1(e). A quality adjusted life-year is an economic evaluation that assesses the monetary value of both the quality and the quantity of life and the cost of medical interventions to determine the value of health outcomes.

PCORI Committed Funds Primarily to Research and Data Capacity Efforts; Awards for Dissemination and Implementation of Findings Were Limited as Most Research Was Still Underway

In fiscal years 2010 through 2017, PCORI committed about $1.6 billion (or 79 percent of its total award commitments of $2.0 billion) to awards for conducting CER and $325 million (or 16 percent) to awards for building data capacity for research. In addition, PCORI committed $93 million for engagement and workforce awards to involve stakeholders in the research process and expand the research workforce, and committed $12 million for awards to disseminate and implement its research findings. Awards for the dissemination and implementation of its research findings were limited as of the end of fiscal year 2017, as most of this research was still underway.\(^{20}\) (See table 1 for PCORI’s award commitments for fiscal years 2010 through 2017.)

<table>
<thead>
<tr>
<th>Activity</th>
<th>2010-2013</th>
<th>2014</th>
<th>2015</th>
<th>2016</th>
<th>2017</th>
<th>Total(^a)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Research awards</td>
<td>274</td>
<td>304</td>
<td>372</td>
<td>327</td>
<td>314</td>
<td>1,591</td>
</tr>
<tr>
<td>Building data capacity awards</td>
<td>9</td>
<td>95</td>
<td>142</td>
<td>44</td>
<td>35</td>
<td>325</td>
</tr>
<tr>
<td>Engagement and workforce awards</td>
<td>0</td>
<td>3</td>
<td>15</td>
<td>23</td>
<td>52</td>
<td>93</td>
</tr>
<tr>
<td>Dissemination and implementation awards</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>10</td>
<td>2</td>
<td>12</td>
</tr>
<tr>
<td>Total(^a)</td>
<td>283</td>
<td>402</td>
<td>529</td>
<td>404</td>
<td>403</td>
<td>2,021</td>
</tr>
</tbody>
</table>

Source: GAO analysis of Patient-Centered Outcomes Research Institute (PCORI) award commitment data. \(^{a}\) Totals may not add up due to rounding.

By the end of fiscal year 2024, PCORI projects to spend a total of almost $3.3 billion, which reflects its projected Trust Fund revenue through fiscal year 2019 plus interest income. This total amount encompasses the commitments PCORI has made for awards through fiscal year 2017, as well as $514 million in projected additional research award commitments\(^{20}\). These award commitments include expenditures through fiscal year 2017 as well as remaining payments for these commitments to be made by 2024. Of the $2.0 billion in award commitments PCORI made through fiscal year 2017, PCORI spent $948 million through fiscal year 2017 and expects to spend the remaining $1.1 billion in fiscal years 2018 through 2024.
to be made by the end of fiscal year 2019 and $207 million for other award commitments to be made by the end of fiscal year 2021. In addition to awards, the total includes PCORI’s expenditures for program and administrative support services in fiscal years 2010 through 2017, as well as projected expenditures for these services through fiscal year 2024.21 (See fig. 2 for PCORI’s actual and projected commitments and expenditures and see app. I for an overview of PCORI’s awards.)

The following information provides details on PCORI’s awards related to research, building data capacity, engagement and workforce activities, and the dissemination and implementation of its research findings.

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21PCORI spent $193 million in fiscal years 2010 through 2017 for program support services, such as staff salaries and benefits, as well as $117 million for administrative support services. PCORI officials anticipate additional expenditures of $206 million for these services through 2024 in order to see these award commitments through to completion. Descriptions of the activities under each PCORI award category may include some program support services. PCORI projects that its administrative expenses will average 6 percent of its total expenditures for fiscal years 2010 through 2024.
PCORI committed $1.6 billion, or 79 percent of its total award commitments, for research in fiscal years 2010 through 2017. In fiscal years 2018 and 2019, PCORI projects to commit an additional $514 million for research awards. PCORI research awards have increasingly focused on conditions that impose a substantial health or financial burden on patients and the healthcare system. (See table 2 for information on the health conditions that received the highest research award funding.)

Table 2: Heath Conditions That Received the Highest PCORI Research Award Commitments, Fiscal Years 2010 through 2017

<table>
<thead>
<tr>
<th>Condition</th>
<th>Total commitments (dollars in millions)</th>
<th>Description and examples</th>
</tr>
</thead>
<tbody>
<tr>
<td>Mental and behavioral health</td>
<td>327</td>
<td>PCORI awarded funding for 93 studies on depression, substance abuse, bipolar disorder, schizophrenia, anxiety disorders, autism, and related projects. For example, in one study researchers are determining whether it is more effective to treat patients with post-traumatic stress and bipolar disorders through integrated care or by using telemedicine technology to facilitate referrals to local mental health specialists.</td>
</tr>
<tr>
<td>Cancer</td>
<td>286</td>
<td>PCORI awarded funding for 78 studies on prevention or treatments for breast, colorectal, lung, prostate, cervical, blood, and other cancer projects. For example, one study compared surgery, radiation or active surveillance outcomes for men with localized prostate cancer.</td>
</tr>
<tr>
<td>Cardiovascular disease</td>
<td>271</td>
<td>PCORI awarded funding for 62 studies on congestive heart failure, hypertension, strokes, and other cardiovascular conditions. For example, one study demonstrated that stroke survivors who used the blood-thinning drug warfarin after leaving the hospital were able to stay in their homes an average of 46 more days than those who did not take the drug.</td>
</tr>
<tr>
<td>Neurological disorders</td>
<td>260</td>
<td>PCORI awarded funding for 65 studies related to Multiple Sclerosis, Alzheimer’s disease, dementia, and other cognitive impairment issues. For example, one study is comparing the effects of each of the five components of an intensive, multipronged, behavioral intervention that is designed to delay or prevent onset of dementia in people with mild cognitive impairment.</td>
</tr>
<tr>
<td>Multiple chronic conditions</td>
<td>225</td>
<td>PCORI awarded funding for 55 studies on patients with two or more chronic conditions such as diabetes, hypertension, obesity, or depression, and other projects focused on multiple chronic conditions. For example, one study focused on comparing case management with standard care for older patients with multiple chronic conditions.</td>
</tr>
</tbody>
</table>

Source: Patient-Centered Outcomes Research Institute (PCORI) information. | GAO-18-311

Note: Studies may be counted in more than one category.

Similar to certain types of CER that may take many years, the entire research award process for PCORI-funded CER may span multiple years from the funding announcement to the dissemination of completed results.

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22This amount includes $785 million PCORI has paid on research awards through fiscal year 2017 as well as $806 million in remaining payments for these commitments to be made by 2024.
Specifically, the process PCORI established can take as many as 6 years, which includes requesting and reviewing proposals, awarding contracts, recruiting participants or obtaining data, conducting and reviewing research, and disseminating findings and typically involves awards that span multiple years. For example, PCORI estimates that the typical timeframe for announcing funding and selecting applications to receive research awards can take 8 to 11 months as PCORI brings scientists, patients, payers, and other stakeholders together to prioritize proposals based on the impact of the condition, potential to improve health, technical merit, patient-centeredness, and engagement. (See fig. 3.)

PCORI officials told us that PCORI-funded research is considered complete only after its findings are posted on the PCORI website.

For example, in 2016 PCORI awarded $15 million for a multi-year study *Comparing Three Medicines to Prevent Blood Clots in Patients Who Need Long Term Blood-Thinning Treatment*. This Duke University randomized control trial will examine 3,000 patients in 60 locations to determine whether those taking two newer medications have fewer bleeding complications than patients taking the older medicine and will compare the two newer blood thinners to each other.
Most of PCORI’s research projects, awarded through fiscal year 2017, were still underway. Only 53 of its 543 research projects had been completed as of the end of fiscal year 2017—in part because PCORI’s research award process typically takes 2 to 6.5 years to complete, and because almost two-thirds of the funds committed for research projects were awarded in fiscal years 2015 through 2017. While most PCORI-funded research is underway, a larger number of research studies are projected to be completed by the end of each year from 2018 to 2022, with all of the remaining studies to be completed by 2024. (See fig. 4.) PCORI officials told us that the institute attempts to manage its funds to ensure that its research awards are funded and managed through completion, including peer review and the distribution of research.

25PCORI officials also told us that, after PPACA authorized the establishment of PCORI in 2010, it took over a year to form its board of governors, hire staff, and develop policies and procedures for funding awards before it could begin making award commitments.
findings, in recognition of the time needed to conduct this research as well as the uncertainty regarding the total amount of funding available.  

Officials from all but one of the stakeholder organizations we interviewed—public and private payers, health care providers, and patient advocacy organizations that represented potential users of CER—generally supported PCORI’s research award priorities. Most of the stakeholders we interviewed stressed the importance of research conducted by unbiased organizations, such as the federally funded research funded by PCORI and HHS. In addition, most stakeholders also told us that PCORI’s efforts to engage patients in the research process

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The fees collected by Treasury from private insurance and self-insured health plans depend upon the average number of covered lives in a policy in a given year and therefore can vary annually.
have changed the way research is conducted for the better, such as prioritizing research outcomes that are most meaningful to patients. However, officials from an organization representing payers (and from an individual health plan) told us that PCORI’s priorities did not fully align with their needs, such as their needs for CER on certain high-cost conditions, medications or treatments.

Building Data Capacity Awards

PCORI committed the second largest portion of award funding—$325 million through fiscal year 2017—for awards to build data capacity for research through the development of PCORnet.27 PCORI officials told us that the institute supported the development of the PCORnet initiative in order to use existing medical records and claims data and to transform much of that data into a common data model to be used for clinical research, until such time when such data will have been standardized in electronic health records so that they can easily be used for research. As of December 2017, PCORnet included 36 partner networks agreeing to link their electronic claims and health data.28 PCORI officials told us that this distributed data network already comprises a nationally representative sample of approximately 128 million individuals whose data can be used in randomized clinical trials, large observational studies, and other research. In fiscal years 2018 and 2019, PCORI projects to commit an additional $70 million for these awards to continue building this data capacity.

PCORnet research is managed through its Coordinating Center, which oversees the translation of certain categories of the partner networks’ data into the common data model and forges agreements with each of the partners to share results of queries using their data with researchers.29

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27This amount includes $115 million PCORI has paid for building data capacity awards through fiscal year 2017 as well as $210 million in remaining payments for these commitments to be made by 2024.

28These partner networks include 21 patient-powered research networks operated and governed by groups of patients and their partners; 13 clinical data research networks based in healthcare systems such as hospitals, practice-based networks, and health centers; and 2 health plan research networks operated by insurers.

29The Coordinating Center oversees the research infrastructure, ensures the reliability of the partner networks’ data, provides for data security, and develops software for the system. PCORnet continues to expand the categories of data that can be translated into the common data model, thereby expanding the data available for research. It has also developed Institutional Review Board systems to facilitate and speed the review of PCORnet research for clinical trials and is developing other resources to facilitate research using partners’ data.
This research process generally starts when a researcher requests to query data on a specific population, after which PCORnet may approve the request and invite network partners to participate. Participating network partners then run queries on their data following established parameters and submit the results to a secure portal that the researcher can access in order to analyze the results for research. (See fig. 5.) PCORI officials told us that there were 32 research projects using PCORnet that received funding through PCORI’s research award process as of the end of December 2017, as well as 45 research projects funded by other parties, including federal agencies and private industry.

Figure 5: Process for Researcher to Receive Data Query Results from PCORnet Partners

Further, as part of its building data capacity awards, in fiscal year 2017 PCORI committed $25 million to the People-Centered Research Foundation, a nonprofit foundation formed in March 2017 to support the network partners and other entities conducting research using PCORnet. This funding was provided to support this foundation’s development of a business plan, as well as its governance structure, to ensure the continuity of the PCORnet network partnership efforts after PCORI.
funding for PCORnet ends. PCORI has indicated it may provide additional funding to the foundation, provided that the foundation and the networks make progress toward self-sustainability.

Officials from most stakeholder organizations we interviewed generally agreed that PCORnet offers value by improving the data available to conduct CER. Officials from two organizations told us that PCORnet has made it possible to use network partners’ aggregated data to make conducting research more efficient than in the past.

Through fiscal year 2017, PCORI also committed $93 million for engagement and workforce awards.\(^{30}\) For example, PCORI committed a total of $63 million for engagement awards, intended to involve a variety of stakeholders in the research process and to improve the methodology for carrying out CER. Engagement awards include “Eugene Washington Engagement Awards” that are intended to bring patients, caregivers, clinicians, and other healthcare stakeholders into the research process and to disseminate study results.\(^{31}\) In addition, “Pipeline to Proposal Awards” are intended to bring together stakeholders with strong interests in a specific health issue to develop research proposals to address their needs. Officials from the two patient advocacy organizations we interviewed told us that PCORI’s engagement awards have helped to support patient involvement in the research process. For example, one official noted that, while it has not been easy to find patients willing to participate, these awards have been important to train and support patients in the research process.

PCORI also committed $30 million to workforce training awards for clinicians and researchers. For example, one of PCORI’s career development programs, conducted in partnership with AHRQ, is designed to train clinician and research scientists to conduct patient-centered

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\(^{30}\)This amount includes $44 million PCORI has paid for engagement and workforce awards through fiscal year 2017 as well as $49 million in remaining payments for these commitments to be made by 2024.

\(^{31}\)For example, the National Kidney Foundation received an award to foster collaboration among patients, clinicians, caregivers, researchers, health payers, and healthcare industry representatives to facilitate the development of research designs to measure home dialysis quality as well as interventions to address the barriers to maintaining dialysis treatment at home.
outcomes research and to actively engage stakeholders in efforts to improve the quality and safety of care.\(^{32}\)

Dissemination and implementation awards for PCORI-funded research findings thus far have been limited as most of the research was still underway, but, according to PCORI officials, awards for this work will substantially increase as research is completed. Specifically, through fiscal year 2017, PCORI committed a total of $12 million for awards to disseminate and implement PCORI-funded research by helping researchers and other stakeholders to publicize findings and by supporting patients and providers to utilize findings.\(^{33}\) PCORI projects to commit an additional $91 million for these awards in fiscal years 2018 through 2021.

Dissemination and implementation awards are intended to encourage PCORI awardees that have completed research and their patient and stakeholder partners to pursue strategic activities to disseminate and implement their findings. For example, PCORI awarded about $0.4 million to increase awareness and promote the use of research findings on using technology to deliver virtual care home visits for those with Parkinson’s disease. According to PCORI, these funds will be used to train neurologists and other health professionals to provide virtual care for patients in their homes. In addition, as part of its efforts to summarize research findings, PCORI also awarded funds to the American Institutes for Research to establish a Translation Center that develops two summaries of each of PCORI’s research findings: a public abstract for general audiences that is also translated into Spanish and a professional abstract for clinicians.

In addition to awards, PCORI has fostered the dissemination and implementation of its research findings in other ways, including through its website, publications, and roundtable briefings. For example, according to PCORI, it posts research findings on its website within 90 days of receiving final peer-reviewed research results so that patients and providers have access to the information to make healthcare decisions. In addition, according to PCORI, it pays journals’ open access fees to allow

\(^{32}\)We describe AHRQ’s workforce training efforts later in this report.

\(^{33}\)This amount includes $4 million PCORI has paid for dissemination and implementation awards through fiscal year 2017 as well as $8 million in remaining payments for these commitments to be made by 2024.
free public access to selected research and plans to support research awardees to place accepted journal manuscripts in the PubMed Central database. PCORI also facilitates roundtable briefings that bring together clinicians, patients, and others with interests in recent findings in order to build support for immediate use of the findings. PCORI also coordinates its dissemination efforts with AHRQ.34

PCORI considers the implementation of its research methods and findings to be an integral part of its dissemination efforts and a culmination of its work and so has begun efforts to track implementation, such as the number of its findings published in peer-reviewed journals, and the use of its findings in clinical care. For example, PCORI officials told us that there were 891 publications in peer-reviewed journals that resulted from studies fully or partially funded by PCORI through October 2017. According to PCORI, two PCORI-funded studies on prostate cancer, one study on oral versus intravenous antibiotics for certain children, and one study on self-monitoring of blood glucose were included in medical resource software that is used by nearly 90 percent of academic medical centers in the United States.

Most of the stakeholder officials we interviewed noted the importance of disseminating research findings quickly and in ways that are readily available and understandable to both experts and the general public to raise awareness about the findings. While officials representing two payers noted limitations to the usefulness of PCORI’s research findings because they do not take treatment costs into account, most stakeholder officials noted the importance of the PCORI-funded research underway and looked forward to utilizing the research findings once they become available.35 In particular, officials representing provider and patient advocacy organizations told us that they were interested in ensuring that the most important research findings would be quickly implemented by patients and clinicians.

34We describe AHRQ’s dissemination efforts later in this report.

35Officials representing two payers told us that cost is a critical component of the information they need to make decisions about coverage policies. However, according to PCORI, it does not fund research that analyzes cost effectiveness because of restrictions in PPACA prohibiting it from developing or using a dollars-per-quality adjusted life-year to establish what type of health care is cost effective or recommended. See 42 U.S.C. § 1320e-1(e).
Between fiscal years 2011 and 2017, HHS’s AHRQ obligated about $260 million (or 58 percent of HHS’s $448 million in total obligations) for the dissemination and implementation of CER findings. According to AHRQ officials, because most PCORI-funded research had not been completed by the end of fiscal year 2017, these efforts were primarily focused on the dissemination and implementation of research funded by other entities, including NIH and the Centers for Disease Control and Prevention (CDC). Additionally, AHRQ obligated $94 million for efforts to train researchers on conducting CER, and ASPE obligated $85 million for efforts to build data capacity. AHRQ and ASPE have obligated a total of $9 million for administrative activities during those years. Table 3 provides an overview of HHS’s obligations in each fiscal year.

### Table 3: HHS Obligations for Comparative Clinical Effectiveness Research Activities, Fiscal Years 2011 through 2017 (Dollars in Millions)

<table>
<thead>
<tr>
<th></th>
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<th></th>
<th></th>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>Dissemination and implementation</td>
<td>4.9</td>
<td>31.1</td>
<td>28.3</td>
<td>82.6</td>
<td>73.0</td>
<td>40.2</td>
<td>260.1</td>
<td></td>
</tr>
<tr>
<td>Training</td>
<td>2.4</td>
<td>11.4</td>
<td>19.9</td>
<td>22.2</td>
<td>22.9</td>
<td>14.9</td>
<td>93.8</td>
<td></td>
</tr>
<tr>
<td>Building data capacity</td>
<td>3.0</td>
<td>7.1</td>
<td>11.3</td>
<td>20.7</td>
<td>17.3</td>
<td>25.8</td>
<td>85.2</td>
<td></td>
</tr>
<tr>
<td>Administrative</td>
<td>0.0</td>
<td>0.4</td>
<td>0.9</td>
<td>1.8</td>
<td>2.1</td>
<td>2.1</td>
<td>2.0</td>
<td>9.3</td>
</tr>
<tr>
<td>Total</td>
<td>0.0</td>
<td>10.7</td>
<td>50.5</td>
<td>61.3</td>
<td>127.6</td>
<td>115.3</td>
<td>82.9</td>
<td>448.4</td>
</tr>
</tbody>
</table>

Source: GAO analysis of Department of Health and Human Services (HHS) obligation data. (GAO-18-311)

### Notes
- HHS’s dissemination and implementation as well as its training activities are managed by the Agency for Healthcare Research and Quality, while HHS’s data capacity initiatives are managed by the Office of the Assistant Secretary for Planning and Evaluation.
- Totals may not add up due to rounding.

AHRQ and ASPE plan to obligate an additional $120 million for dissemination and implementation, training, building data capacity, and administrative activities during fiscal years 2018 through 2020. They expect to have $245 million available to fund ongoing and future CER activities, based on expected transfers from the Trust Fund in fiscal years 2018 and 2019.36 (See fig. 6.)

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36As stated previously, for each of fiscal years 2011 through 2019, Treasury must transfer 20 percent of Trust Fund revenue to HHS. Amounts transferred to HHS remain available until expended. The fees collected by Treasury from private insurance and self-insured health plans depend upon the average number of covered lives in a policy in a given year and therefore can vary annually.
The following information provides details on HHS-funded projects related to dissemination and implementation, training on conducting CER, and building data capacity.

Dissemination and Implementation

During fiscal years 2011 through 2017, AHRQ obligated a total of $260 million for CER dissemination and implementation initiatives and plans to obligate an additional $93 million for these initiatives in fiscal years 2018 through 2020. According to officials, AHRQ plans to fund additional dissemination and implementation initiatives in fiscal years 2018 and 2019 but had not finalized those plans as of January 2018. (See app. II for an overview of all of AHRQ’s dissemination and implementation initiatives.) AHRQ’s dissemination and implementation initiatives comprise efforts to synthesize CER findings, translate and communicate research findings to potential users, and implement them:

- **Synthesis of CER findings:** According to AHRQ officials, AHRQ’s Evidence-Based Practice Centers developed 48 systematic reviews of CER findings based on completed research. As of the end of fiscal year 2017, 40 of these reviews had been published, while 8 were still in progress. Officials told us that these systematic reviews have likely
not included PCORI-funded research, as most of that research had not been completed by the end of fiscal year 2017.

- **Translation and communication of CER findings:** AHRQ funded initiatives, which—according to the agency—are aimed at making CER findings accessible and understandable to health care professionals, patients, and others. For example, AHRQ developed a “Library of Patient-Centered Outcomes Research Resources” website with links to CER databases maintained by other entities including NIH and PCORI. Another example is AHRQ’s “John M. Eisenberg Center for Clinical Decisions and Communications Science,” which translates research findings into information that can be used by consumers, health care providers, and policymakers.

- **Implementation of CER findings:** AHRQ funded four key initiatives to implement CER findings. According to AHRQ officials, one of the four initiatives includes PCORI-funded research, while the other three have thus far focused on implementing existing CER funded by other entities:
  - The “Dissemination and Implementation Initiative” was designed to disseminate and implement government-funded CER findings—including PCORI-funded findings—relevant to physicians, healthcare providers, patients, and others. This initiative consists of a multi-step approach for identifying several areas of CER each year that—according to AHRQ officials—have the greatest potential for impact and are feasible to implement. (See figure 7 for an overview of this process.) According to AHRQ officials, as of December 2017, 37 findings have been nominated for consideration under AHRQ’s Dissemination and Implementation Initiative, including 5 findings nominated by PCORI. According to these officials, 1 of the findings PCORI has nominated is under consideration for implementation. Two were rejected—1 because

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37The website is intended to assist researchers who may be conducting new studies, as well as clinicians, policymakers, consumers, and others who are seeking access to evidence-based health information. AHRQ did not use any resources from the Trust Fund to support this effort during the time of our review but plans to in the future.

38According to AHRQ, the Eisenberg Center develops products to support informed, shared, and evidence-based health care decisions regarding tests or treatments for different types of health conditions. The products, including research summaries and patient decision aids, are presented in short, easy-to-use formats. In addition, the Eisenberg Center works with local and national patient and consumer organizations to identify opportunities to disseminate research findings.
of insufficient impact and the other because of challenges in implementation feasibility. (Two are still under review.)

Figure 7: Overview of the AHRQ Dissemination and Implementation Initiative Process

AHRQ receives nominations to fund the dissemination and implementation of Comparative Clinical Effectiveness Research (CER) findings.

1. **Reviews against criteria**
   - Reviews nominations to ensure findings fit the definition of CER and are supported by peer-reviewed research.
   - Projects proceed to the next step
   - Projects are dropped from proceeding

2. **Assesses evidence and impact**
   - Assesses strength of the evidence, consistency with prior evidence, and potential impact on health outcomes, delivery, and disparities.
   - Projects proceed to the next step
   - Projects are dropped from proceeding

3. **Assesses feasibility**
   - Assesses for implementation feasibility, including fit with organizational capabilities and alignment with AHRQ's mission.
   - Projects proceed to the next step
   - Projects are dropped from proceeding

4. **Explores implementation approaches**
   - Convenes stakeholders to assess current implementation issues, relevant approaches, and strategies, settings, and populations.
   - Projects proceed to the next step
   - Projects are dropped from proceeding

5. **Supports funding**
   - Selects CER for dissemination and implementation funding with considerations for available funds, agency priorities and stakeholder support.
   - Projects proceed to the next step
   - Projects are dropped from proceeding

6. **Assesses evidence and impact**
   - Evaluates efforts for changes in practice or health outcomes to inform subsequent efforts.
   - Projects proceed to the next step
   - Projects are dropped from proceeding

Source: GAO analysis of Agency for Healthcare Research and Quality (AHRQ) information. | GAO-18-311

- The “Evidence Now” initiative disseminates CER evidence directly to primary care practices and supports them in implementing clinical and organizational evidence in practice through regional cooperatives.
- The “Comparative Health System Performance Initiative” established three centers of excellence and a coordinating center to identify, classify, track, and compare health systems. AHRQ’s goal is to understand the factors that affect health systems’ use of CER and to identify best practices in disseminating and using CER.
• The “Clinical Decision Support (CDS) Initiative” is designed to use CDS to promote the timely incorporation of CER findings into clinical practice.39

Some of AHRQ’s dissemination and implementation initiatives—such as “Evidence Now” and “CDS Initiative”—include an evaluation component, as described in app. II. According to AHRQ officials, as of January 2018 results from these evaluations were not yet available.

Training on Conducting CER

Between fiscal years 2011 and 2017, AHRQ obligated a total of $94 million for awards supporting training in the methods used to conduct CER. AHRQ plans to obligate an additional $14 million for training on conducting CER by fiscal year 2020. AHRQ has funded eight categories of awards for individual researchers or research institutions. For example, AHRQ’s “Infrastructure Development Program in Patient-Centered Outcomes Research” award supports institutions in the development of their capacity to conduct and implement CER. Its “Institutional Mentored Career Development Award Program in Patient-Centered Outcomes Research” award supports the development of researchers in academic and applied settings. (See app. III for an overview of these awards.) Starting in fiscal year 2018, AHRQ plans to fund an additional training award category in conjunction with PCORI. AHRQ developed a plan to evaluate its training activities and, according to AHRQ officials, the evaluation is expected to be funded in fiscal year 2018.

Building Data Capacity

Between fiscal years 2012 and 2017, ASPE obligated a total of $85 million for 30 projects designed to build data capacity for conducting CER and plans to obligate an additional $6 million to existing projects and 1 new project through fiscal year 2019. Officials told us that ASPE plans to fund additional projects to build data capacity in fiscal years 2018 and 2019, based on HHS leaders’ priorities, but had not finalized those plans as of January 2018. ASPE manages these projects, which are largely carried out by other HHS agencies through interagency agreements and are intended to develop and maintain a comprehensive, interoperable data network to collect, link, and analyze data on outcomes and

39CDS provides health care providers and other individuals with knowledge and person-specific information, intelligently filtered or presented at appropriate times, to enhance health and health care. CDS encompasses a variety of tools, such as computerized alerts and clinical guidelines, to enhance decision-making in the clinical workflow.
effectiveness from multiple sources for CER.40 (See app. IV for an overview of these activities.)41

In response to a recommendation in our 2015 report on HHS’s CER activities, ASPE implemented a monitoring system to track progress toward its milestones and deliverables for these projects.42 ASPE also contracted to evaluate its projects to build data capacity for CER. The evaluation, completed in December 2017, found that ASPE made progress managing these projects towards the core functionalities outlined in its strategic framework. However, among other things, the evaluation found that additional efforts are needed to explore how to enhance data privacy and security, ensure data quality, and operationalize related standards. According to ASPE officials, the evaluation will inform the development and implementation of future ASPE projects to build data capacity for conducting CER.

We provided a draft of this report to PCORI and HHS for review and comment. PCORI and HHS provided technical comments, which we incorporated as appropriate.

We are sending copies of this report to the appropriate congressional committees, the Executive Director of PCORI, the Secretary of Health and Human Services, the Director of AHRQ, the Assistant Secretary for

40The activities are carried out by the following HHS agencies: AHRQ, ASPE, CDC, Centers for Medicare and Medicaid Services, Food and Drug Administration, Health Resources and Services Administration, NIH, National Libraries on Medicine, and Office of the National Coordinator for Health Information Technology. In fiscal year 2012, ASPE obligated about $0.2 million for the establishment of intra-departmental partnerships between ASPE and other agencies within HHS.

41ASPE’s projects are designed to align with the five core functionalities identified in HHS’s Patient-Centered Outcomes Research Trust Fund strategic plan: 1) use of clinical data for research; 2) standardized collection of standardized clinical data; 3) linking clinical and other data; 4) collection of participant-provided information via new data collection technologies that provide a means for collecting patient generated information critical to patient-centered research; and 5) use of enhanced publicly-funded data systems for research.

42See GAO-15-280. In response to GAO’s recommendation, in each quarter the agencies responsible for implementing each project are required to report on performance towards goals and objectives, among other requirements. ASPE developed “individual dashboard” reports for the projects that include objectives, milestones, and timeframes, as specified in GAO’s recommendation.
ASPE, and other interested parties. In addition, the report will be available at no charge on GAO's website at http://www.gao.gov.

If you or your staff have any questions about this report, please contact me at (202) 512-7114 or dickenj@gao.gov. Contact points for our Offices of Congressional Relations and Public Affairs may be found on the last page of this report. GAO staff who made major contributions to this report are listed in app. V.

John E. Dicken
Director, Health Care
List of Committees

The Honorable Orrin G. Hatch
Chairman
The Honorable Ron Wyden
Ranking Member
Committee on Finance
United States Senate

The Honorable Lamar Alexander
Chairman
The Honorable Patty Murray
Ranking Member
Committee on Health, Education, Labor, and Pensions
United States Senate

The Honorable Greg Walden
Chairman
The Honorable Frank Pallone Jr.
Ranking Member
Committee on Energy and Commerce
House of Representatives

The Honorable Kevin Brady
Chairman
The Honorable Richard Neal
Ranking Member
Committee on Ways and Means
House of Representatives
## Appendix I: Patient-Centered Outcomes Research Institute (PCORI) Award Commitments Made During Fiscal Years 2010 through 2017

<table>
<thead>
<tr>
<th>Award category</th>
<th>Award category description</th>
<th>Total commitments (dollars in millions)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Research awards</td>
<td>These awards generally fund research studies in priority areas for conditions that impose a substantial burden on patients and the healthcare system. Information about individual research awards can be found at: <a href="https://www.pcori.org/research-results?f%5B0%5D=field_project_type%3A298&amp;f%5B0%5D=field_project_type%3A298#search-results">https://www.pcori.org/research-results?f%5B0%5D=field_project_type%3A298&amp;f%5B0%5D=field_project_type%3A298#search-results</a></td>
<td>1,591</td>
</tr>
<tr>
<td>Building data capacity awards</td>
<td>These awards fund infrastructure projects to build data capacity through the development of PCORnet and support for clinical and patient-powered data research networks. Information about individual building data capacity awards can be found at: <a href="https://www.pcori.org/research-results?f%5B0%5D=field_project_type%3A441#search-results">https://www.pcori.org/research-results?f%5B0%5D=field_project_type%3A441#search-results</a></td>
<td>325</td>
</tr>
<tr>
<td>Engagement and workforce awards</td>
<td>Engagement awards fund projects to improve the methodology for carrying out research by involving patients, caregivers, clinicians, and other healthcare stakeholders into the research process. Workforce training awards provide accredited continuing education opportunities, in coordination with the Agency for Healthcare Research and Quality, for researchers and clinicians. Information about individual engagement awards and workforce awards can be found at: <a href="https://www.pcori.org/research-results?f%5B0%5D=field_project_type%3A299#search-results">https://www.pcori.org/research-results?f%5B0%5D=field_project_type%3A299#search-results</a> and <a href="https://www.pcori.org/research-results/2017/k12-institutional-mentored-career-development-program">https://www.pcori.org/research-results/2017/k12-institutional-mentored-career-development-program</a></td>
<td>93</td>
</tr>
<tr>
<td>Dissemination and implementation awards</td>
<td>Dissemination and implementation awards are intended to help researchers and other stakeholders to publicize findings, and support the utilization of findings for patients and providers. Information about individual dissemination and implementation awards can be found at: <a href="https://www.pcori.org/research-results?f%5B0%5D=field_project_type%3A308#search-results">https://www.pcori.org/research-results?f%5B0%5D=field_project_type%3A308#search-results</a></td>
<td>12</td>
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</table>

Source: GAO analysis of PCORI documentation. | GAO-18-311

*All websites accessed on January 23, 2018.*
## Appendix II: Agency for Healthcare Research and Quality’s (AHRQ) Dissemination and Implementation Initiatives, as of September 30, 2017

<table>
<thead>
<tr>
<th>Initiative title</th>
<th>Description</th>
<th>Total obligations (dollars in millions)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Focusing on the ABCs: Accelerating the Dissemination and Implementation of Patient-Centered Outcomes Research (PCOR) Findings into Primary Care Practice (Evidence Now)*</td>
<td>Disseminates evidence to primary care practices and supports them in implementing clinical and organizational evidence in practice through regional cooperatives. AHRQ awarded a separate grant to establish an independent, external evaluation to study improvements in the delivery of the ABCs.</td>
<td>96.1b</td>
</tr>
<tr>
<td>Comparative Health System Performance in Accelerating PCOR</td>
<td>Develops three Centers of Excellence on “Comparative Health System Performance in Accelerating PCOR Dissemination.” According to AHRQ, the Centers of Excellence will identify and classify characteristics of health care systems over 5 years. They will also identify ways to assess the quality and cost of such systems, including their use of PCOR, understand the characteristics of high performing systems, and identify what system characteristics are associated with more rapid adoption and diffusion of PCOR-recommended practices throughout a system.</td>
<td>34.2c</td>
</tr>
<tr>
<td>The Evidence-Based Practice Centers – Evidence Synthesis and Translation</td>
<td>Evaluates and synthesizes research findings to aid decision-making for patients, providers, and payers, among others.</td>
<td>27.9d</td>
</tr>
<tr>
<td>John M. Eisenberg Clinical Decisions and Communications Science Center</td>
<td>Translates PCOR findings into tools, such as research summaries and decision aids, designed to help patients and consumers, clinicians, and policymakers make informed and evidence-based health care decisions.</td>
<td>15.2</td>
</tr>
<tr>
<td>Disseminating PCOR to Improve Healthcare Delivery Systems</td>
<td>Expands an existing initiative or creates a new initiative that supports multi-site, multi-region, multi-stakeholder dissemination and implementation of evidence.</td>
<td>13.2</td>
</tr>
<tr>
<td>Methods for Dissemination and Translation</td>
<td>Develops and tests methods for translating and disseminating PCOR findings to hard-to-reach audiences, including patients with low health literacy, disadvantaged populations, isolated clinicians and policy makers, and other decision makers who may not have had the benefit of more traditional translation and dissemination efforts.</td>
<td>11.4</td>
</tr>
<tr>
<td>Dissemination and Implementation of PCOR Findings to Clinicians</td>
<td>Integrates PCOR into clinical practice using various methods shown to improve the uptake of scientific evidence in clinical decision making. Grantees were asked to consider both educational theory and the relevance of “new media” as they designed their programs.</td>
<td>6.9</td>
</tr>
<tr>
<td>Horizon Scanning System: Dissemination of Information on Emerging Interventions</td>
<td>Searches for emerging interventions, prioritizes those most likely to have a large impact in the near future, and disseminates the information to the public. According to AHRQ, the Horizon Scanning System screened more than 22,000 potential intervention leads and tracked over 2,300 intervention topics.</td>
<td>5.5</td>
</tr>
<tr>
<td>National Guideline Clearinghouse</td>
<td>Provides targeted audiences—such as providers and payers—with an accessible tool for obtaining objective, detailed information on evidence-based clinical practice guidelines to further their dissemination, implementation, and use.</td>
<td>5.3</td>
</tr>
<tr>
<td>Registry of Patient Registries</td>
<td>Promotes collaboration, reduces redundancy, and improves transparency in patient registries.</td>
<td>5.1</td>
</tr>
</tbody>
</table>
### Comparative Clinical Effectiveness Research Funding

<table>
<thead>
<tr>
<th>Initiative title</th>
<th>Description</th>
<th>Total obligations (dollars in millions)</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Electronic Data Methods Forum</strong></td>
<td>Created informational tools to support the dissemination and implementation of PCOR findings, including best practices and new knowledge about the use of electronic health record data for research and quality improvement.</td>
<td>5.1</td>
</tr>
<tr>
<td><strong>Supporting Decisions With PCOR: PCOR Clinical Decision Support Initiative</strong></td>
<td>Promotes the timely incorporation of PCOR findings into clinical practice—which encompasses a variety of tools to enhance clinical decision-making.</td>
<td>5.0</td>
</tr>
<tr>
<td><strong>National Initiative (Publicity Center)</strong></td>
<td>Collaborated with 176 national organizations to disseminate materials for the Effective Health Care Program.</td>
<td>4.2</td>
</tr>
<tr>
<td><strong>“Value of Using Medical Evidence” – A Paid Multi-Media Campaign</strong></td>
<td>Multi-media campaign to educate health care consumers about the value of reviewing medical evidence when weighing treatment options.</td>
<td>3.4</td>
</tr>
<tr>
<td><strong>Educating the Educators</strong></td>
<td>Conducted and disseminated research to develop a process for shared decision making that includes exploring and comparing the benefits, harms, and risks of each option through meaningful dialogue about what matters most to the patient.</td>
<td>3.1</td>
</tr>
<tr>
<td><strong>Patient-Generated Health Outcomes Data and Clinical Decision Support Using Smart Device Technology</strong></td>
<td>Collects patient-generated health data, integrates patient-generated health data with PCOR evidence, and disseminates PCOR findings using mobile health technology.</td>
<td>3.0</td>
</tr>
<tr>
<td><strong>Closing the Gap in Disparities with PCOR</strong></td>
<td>Identifies ways to reduce health care differences across diverse populations with a particular focus on minority populations in under-resourced healthcare settings.</td>
<td>2.9</td>
</tr>
<tr>
<td><strong>Regional Partnership Development Offices</strong></td>
<td>Established five regional offices, responsible for developing and cultivating dissemination partnerships within each region.</td>
<td>2.5</td>
</tr>
<tr>
<td><strong>Implementation of a Systematic Review Data Repository Collaborative</strong></td>
<td>This repository houses study data extracted from primary research publications during the course of conducting systematic reviews. It is designed to increase the transparency of comparative effectiveness reviews, improve the ability to update systematic reviews, improve the quality of abstracted data, and enhance the efficiency and reduce the costs of conducting reviews.</td>
<td>1.9</td>
</tr>
<tr>
<td><strong>Deliberative Approaches for Patient Involvement in Implementing Evidence-Based Health Care</strong></td>
<td>Gathers input from patients on a complex topic related to the implementation of evidence-based health-care decision making.</td>
<td>1.9</td>
</tr>
<tr>
<td><strong>Methods Guidance to Improve the Dissemination and Implementation of Systematic Reviews</strong></td>
<td>Increases the relevance of AHRQ systematic reviews for patients, clinicians, and policymakers by examining and addressing challenging topic areas that may affect the credibility and utility of the review for end users and that are areas of inconsistency or variation among AHRQ systematic reviews.</td>
<td>1.3</td>
</tr>
<tr>
<td><strong>Developing and Evaluating Methods for Record Linkage and Reducing Bias in Clinical Patient Registries</strong></td>
<td>Conducted three projects to improve the development of registries, a major activity of AHRQ’s Effective Health Care Program.</td>
<td>1.1</td>
</tr>
<tr>
<td>Initiative title</td>
<td>Description</td>
<td>Total obligations (dollars in millions)</td>
</tr>
<tr>
<td>---------------------------------------------------------------------------------</td>
<td>---------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------</td>
<td>----------------------------------------</td>
</tr>
<tr>
<td>Continuing Education of Comparative Clinical Effectiveness Research (CER)</td>
<td>Provides online continuing education materials that inform physicians and other health care providers about PCOR from the Effective Health Care Program.</td>
<td>0.9</td>
</tr>
<tr>
<td>Evaluating the Knowledge and Education Needs to Students of Health Professions on PCOR</td>
<td>Worked with health professional student associations to evaluate students’ understanding of the importance and clinical applicability of PCOR and shared decision-making to their practice and evaluated students’ educational needs and preferences related to integrating PCOR findings into their training curricula.</td>
<td>0.7</td>
</tr>
<tr>
<td>Methods Center in Decision and Simulation Modeling</td>
<td>Created a decision-modeling methods center that reviewed the existing research and guidance published on modeling methods with input from a multidisciplinary group of experts.</td>
<td>0.7</td>
</tr>
<tr>
<td>Enhancing CER Data Resources</td>
<td>Provides for maintenance and updating of existing data resources to conduct future CER through a grant competition. The grants fund three to four 1-year pilot projects aimed at enabling a future, larger competition to enhance the data infrastructure and move the resources to self-sustaining models.</td>
<td>0.7</td>
</tr>
<tr>
<td>Dissemination &amp; Implementation of PCOR</td>
<td>Disseminates CER findings published by the Patient-Centered Outcomes Research Institute (PCORI) and other government entities to providers, patients, payers, and others. This initiative consists of a seven-step approach for identifying research findings that have the greatest potential for implementation.</td>
<td>0.4[^4]</td>
</tr>
<tr>
<td>Integrating PCOR into Clinical Decision Support Systems: An Environmental Scan</td>
<td>Provided an understanding of how AHRQ could effectively disseminate and promote PCOR findings and tools in the development and maintenance of clinical decision support systems. The project included a market analysis and an assessment of potential stakeholders and audiences, including vendors of health information technology focused on clinical decision support. Information gathered from this project directly informed the concept for the PCOR clinical decision support initiative that was launched in 2016.</td>
<td>0.3</td>
</tr>
<tr>
<td>PCOR Dissemination: In-Store Promotion</td>
<td>Promoted PCOR through public service announcements nationwide.</td>
<td>0.2</td>
</tr>
<tr>
<td>Increasing Access to PCOR/CER Findings</td>
<td>Created a new page on AHRQ’s website that highlights the agency’s own resources, as well as directs researchers, health professionals, patients, caregivers, and families to additional databases that collect information on CER. These databases provide summaries of findings from a wide range of CER findings and research that is in progress.</td>
<td>0.0[^9]</td>
</tr>
</tbody>
</table>

[^4] PCOR is a form of CER.
[^6] AHRQ plans to obligate an additional $24.7 million for this initiative in fiscal year 2018.
[^7] AHRQ plans to obligate an additional $16.0 million for this initiative in fiscal years 2018 through 2020.
[^8] AHRQ plans to obligate an additional $30.0 million for this initiative in fiscal years 2018 through 2020.
[^9] AHRQ plans to obligate $0.4 million for this initiative in fiscal year 2018.
[^10] AHRQ plans to obligate $0.2 million for this initiative in fiscal year 2018.
### Appendix III: Agency for Healthcare Research and Quality’s (AHRQ) Training Awards, as of September 30, 2017

<table>
<thead>
<tr>
<th>Award title</th>
<th>Description</th>
<th>Total obligations (dollars in millions)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Infrastructure Development Program in Patient-Centered Outcomes Research (PCOR)*</td>
<td>Funds a 5-year, renewable effort to support the development of PCOR capacity among institutions that have basic health services research capacity but need to develop capacity to conduct and implement PCOR. The program would potentially include institutions located in geographic areas that lack capacity, and institutions that serve predominantly minority populations.</td>
<td>33.3</td>
</tr>
<tr>
<td>Institutional Mentored Career Development Award Program in PCOR</td>
<td>Supports the development of researchers in academic and applied settings. The program combines didactic and experiential opportunities, focusing on the generation, adoption, and spread of new scientific evidence. The goal is to improve population-specific health outcomes by developing and disseminating evidence-based information to patients, clinicians, and other decision-makers, responding to their expressed needs, about which interventions are most effective for which patients under specific circumstances.</td>
<td>26.3&lt;sup&gt;b&lt;/sup&gt;</td>
</tr>
<tr>
<td>Researcher Training and Workforce Development in Methods and Standards for Conducting PCOR Studies</td>
<td>Provides basic, advanced, and experiential training on the methods to conduct PCOR, particularly prospective observational research, registries, and clinical trials. The program was open to researchers employed in both the public and private sectors, particularly those who serve minorities, economically or medically disadvantaged populations.</td>
<td>8.9&lt;sup&gt;c&lt;/sup&gt;</td>
</tr>
<tr>
<td>Pathway to Independence in PCOR</td>
<td>Facilitates the transition of postdoctoral candidates from mentored to independent research positions, accelerating research independence for PCOR researchers.</td>
<td>7.9</td>
</tr>
<tr>
<td>Individual Mentored Career Development Award Program in PCOR</td>
<td>Provides support for intensive, research career development for individual investigators in academic or applied settings, leading to research independence in the field of PCOR and the generation and translation of new scientific evidence and analytic tools.</td>
<td>7.6&lt;sup&gt;d&lt;/sup&gt;</td>
</tr>
<tr>
<td>Research Career Enhancement Awards for Established Investigators in PCOR</td>
<td>Provides career development awards for established investigators to further develop their research expertise in PCOR methodologies. This concept seeks to accelerate the development of the research workforce capable of conducting PCOR.</td>
<td>5.5</td>
</tr>
<tr>
<td>PCOR Training</td>
<td>Provides 2-year fellowships for training in PCOR. A focus for these fellowships is recruitment of trainees from diverse disciplines, including social and behavioral sciences, business, and engineering. The expected output of these fellowships is trained PCOR researchers.</td>
<td>4.2</td>
</tr>
<tr>
<td>Training the Next Generation of PCOR Investigators to Lead Research on the Learning Healthcare System</td>
<td>Establishes an expert panel, comprised of 7 to 10 leaders in the fields of learning healthcare system, health services research, and PCOR, to assess the current state of health services research and PCOR training and recommend ways to improve core competencies/curriculum to meet the needs of the health system. Develops a report summarizing the panel’s recommendations concerning current deficiencies and recommendations regarding skills and competencies needed to meet the challenges.</td>
<td>0.2</td>
</tr>
</tbody>
</table>

*PCOR is a form of CER.

<sup>a</sup>AHRQ plans to obligate an additional $7.3 million for this award in fiscal year 2018.

<sup>b</sup>AHRQ plans to obligate an additional $2.4 million for this award in fiscal year 2018.

<sup>c</sup>AHRQ plans to obligate an additional $4.1 million for this award in fiscal years 2018-2020.

Source: GAO analysis of AHRQ documentation. | GAO-18-311
### Appendix IV: Office of the Assistant Secretary for Planning and Evaluation’s (ASPE) Projects to Build Data Capacity, as of September 30, 2017

<table>
<thead>
<tr>
<th>Project title</th>
<th>Description</th>
<th>Total obligations (dollars in millions)</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Complete</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Creating the Foundational Blocks for the Learning Health Care System: Data Access Standards for Electronic Health Records</td>
<td>Developed technical standards for how health care providers, researchers, and the public health community access and extract data from electronic health records to conduct Patient-Centered Outcomes Research (PCOR). ³</td>
<td>5.2</td>
</tr>
<tr>
<td>Creating the Foundational Blocks for the Learning Health Care System: Structured Data Capture</td>
<td>Identified and developed the functional and technical specifications necessary to enable electronic health record systems to retrieve, display, and fill a structured form or template and store and submit the completed form to an external repository.</td>
<td>4.2</td>
</tr>
<tr>
<td>Maintenance and Support of Chronic Conditions Warehouse for Comparative Clinical Effectiveness Research (CER)</td>
<td>Provided researchers with access to the Centers for Medicare &amp; Medicaid Services’ Chronic Conditions Warehouse, which contains Medicare and Medicaid beneficiary, claims, and assessment data, and supported infrastructure enhancements to conduct CER.</td>
<td>3.0</td>
</tr>
<tr>
<td>Expanding Data Collection for the National Program of Cancer Registries for CER</td>
<td>Longitudinal follow-up of certain cancer patients to assess vital statistics, disease recurrence, disease progression, and additional treatment types. Treatment data submitted each year to the Centers for Disease Control and Prevention and provided to researchers through the National Center for Health Statistics Research Data Center.</td>
<td>2.6</td>
</tr>
</tbody>
</table>
| Strengthening and Expanding Community Health Applied Research Network Registry to PCOR | • Included clinical encounters for all patients and all conditions seen at the community health centers from 2006 to 2013 in the Community Health Applied Research Network Registry data warehouse, a research network comprising 18 community health centers. A de-identified analytic file and associated data codebook were developed to support the use of analytic files by researchers outside of the network.  
• Established a process for investigators to access the data warehouse through the development of a data access plan.  
• Maintained the infrastructure for PCOR and for quality improvement in the safety net. | 2.0                                     |
| Development of Data Infrastructure for Use of Electronic Health Records in CER | Developed common data elements and standards for CER. The results were the initial entries into the National Institutes of Health’s National Library of Medicine common data element repository. | 2.0                                     |
| Strategic Opportunities for Building Data Infrastructure for CER              | Developed a conceptual framework and environmental scan; produced policy documents ranging from patient-initiated data, through research data on care processes, transitions and coordination, to researcher access to claims data; and developed the ‘HHS Strategic Roadmap for Building Data Capacity for Clinical Comparative Effectiveness Research.’ | 1.5                                     |
| CER Inventory                                                                | The overall CER Inventory project was to design and implement a system for the categorization and cataloguing of CER activities through a web-based tool. Due to the rapidly evolving technologies supporting web-based search engines, and the improved methods for identification of more recent CER, the development of the CER Inventory (as a web-based search engine using a retrospective algorithm) was determined to have been superseded by existing search engine tools available. | 0.8                                     |
### Appendix IV: Office of the Assistant Secretary for Planning and Evaluation’s (ASPE) Projects to Build Data Capacity, as of September 30, 2017

<table>
<thead>
<tr>
<th>Project title</th>
<th>Description</th>
<th>Total obligations (dollars in millions)*</th>
</tr>
</thead>
<tbody>
<tr>
<td>Office of the Secretary Patient-Centered Outcomes Research Trust Fund evaluation</td>
<td>Designed and conducted an independent evaluation of the ASPE portfolio to systematically assess progress related to the strategic framework functionalities.</td>
<td>0.7</td>
</tr>
<tr>
<td>Multi-Payer Claims Database Beta Test</td>
<td>Conducted CER analyses on the beta release of the Multi-Payer Claims Database and evaluated beta testers’ experiences requesting and using data from the MPCD for research. Results of the beta test found that the project was successful in achieving the key objectives of building a pilot database.</td>
<td>0.2</td>
</tr>
<tr>
<td>Improving Beneficiary Access to Health Information “Blue Button” to Enable a ‘Data-as Service’ Platform</td>
<td>Planned for development and implementation of the Centers for Medicare &amp; Medicaid Services’ Blue Button—a service that allows patients to access their own health information in electronic form.</td>
<td>0.2</td>
</tr>
<tr>
<td><strong>In Progress</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Enhancing Data Resources for Studying Patterns and Correlates of Mortality in PCOR</td>
<td>Linkage of data on fact, cause, and manner of death from the National Death Index to several federal population-based health data platforms in order to demonstrate the feasibility of such linkage, enable PCOR on patterns and correlates of mortality via the resulting linked data; and to facilitate collaboration between federal partners regarding strengthening the infrastructure and methods for linking healthcare data to mortality outcomes and using such linked data for PCOR.</td>
<td>6.6</td>
</tr>
<tr>
<td>Improving the Mortality Data Infrastructure for Patient-Centered Outcomes</td>
<td>Improve the infrastructure to support timely and complete mortality data collection through more timely delivery of state death records to the National Death Index database and by linking National Death Index database records with nationally collected hospital datasets to obtain a more complete picture of patient care.</td>
<td>6.5</td>
</tr>
<tr>
<td>Security and Privacy Standards for Patient Matching, Linking and Aggregation</td>
<td>Identify the best patient attributes to address the challenge of linking patients’ data across research, clinical, and claims data sets in order to support the PCOR data infrastructure that enables standardization and sharing of patient data across organizations.</td>
<td>5.5</td>
</tr>
<tr>
<td>Developing a Strategically Coordinated Registry Network for Women’s Health Technologies</td>
<td>Create a coordinated registry network for women’s health technologies that will collect patient reported outcomes and employ structured data capture from electronic health records for data collection and exchange.</td>
<td>5.1</td>
</tr>
<tr>
<td>Harmonization of Various Common Data Models and Open Standards for Evidence Generation</td>
<td>Build data infrastructure for conducting PCOR using data from routine clinical settings. The sources of these data may include, but are not limited to, insurance billing claims, electronic health records, and patient registries. This project intends to harmonize several existing common data models, potentially including PCORnet and other networks.</td>
<td>4.1</td>
</tr>
<tr>
<td>Advancing the Collection and Use of Patient-Reported Outcomes through Health Information Technology</td>
<td>Develop technical tools for collecting and integrating patient-reported outcome assessments into electronic health records or other health information technology products.</td>
<td>3.9</td>
</tr>
<tr>
<td>Improving Beneficiary Access to Their Health Information through an Enhanced Blue Button Service</td>
<td>Create an interface that enables CMS beneficiaries to connect their MyMedicare.gov data to applications and services they trust, including research platforms related to research studies in which the beneficiary may be interested in participating.</td>
<td>3.8</td>
</tr>
<tr>
<td>Resource Center</td>
<td>Provide technical assistance to the Trust Fund awardees in informatics and assist ASPE in setting up additional oversight processes and procedures to monitor progress.</td>
<td>3.5</td>
</tr>
<tr>
<td>Project title</td>
<td>Description</td>
<td>Total obligations (dollars in millions)</td>
</tr>
<tr>
<td>-----------------------------------------------------------------------------</td>
<td>---------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------</td>
<td>----------------------------------------</td>
</tr>
<tr>
<td>PCOR: Privacy and Security Blueprint, Legal Analysis and Ethics Framework for Data Use, and Use of Technology for Privacy</td>
<td>Develop a privacy and security data infrastructure blueprint, legal analysis, and ethical framework to address legal and privacy and security related policy issues that affect the use of data for various types of PCOR.</td>
<td>3.4</td>
</tr>
<tr>
<td>Harmonization of Clinical Data Element Definitions for Outcome Measures in Registries</td>
<td>Convene clinical topic-specific working groups to discuss the data definitions currently in use and how these definitions can be harmonized to promote common definitions for outcome measures across systems. These common definitions are to be made publicly available to PCOR researchers and analysts.</td>
<td>3.1</td>
</tr>
<tr>
<td>Development of a Natural Language Processing Web Service for Public Health Use</td>
<td>Develop a natural language processing service that will be accessible and publicly available to researchers on the Public Health Community Platform – a cooperative platform for sharing interoperable technologies to address public health priority areas aimed at improving population health outcomes and health equity (e.g., tobacco use).</td>
<td>3.0</td>
</tr>
<tr>
<td>Technologies for Donating Medicare Beneficiary Claims Data to Research Studies</td>
<td>Leverage the Sync for Science and Blue Button application programming interface programs to enable Medicare beneficiaries to donate their medical claims data for scientific research studies.</td>
<td>2.5</td>
</tr>
<tr>
<td>Utilizing Data from Various Data Partners in a Distributed Manner</td>
<td>Develop and test the capability to conduct timely and secure distributed regression analysis in distributed data networks. Additionally, explore the feasibility of creating virtual linkage capabilities to utilize data from multiple data sources and data for one specific patient with information at different institutions.</td>
<td>2.5</td>
</tr>
<tr>
<td>Collection of Patient-Provided Information through a Mobile Device Application for Use in Comparative Effectiveness and Drug Safety Research</td>
<td>Create the infrastructure for collecting data from patients through a mobile device application, allowing patient-generated data to be linked with a single data partner that participates in the Food and Drug Administration’s Sentinel distributed network. The project will develop and pilot a mobile application to capture data from pregnant women who volunteer to participate.</td>
<td>2.0</td>
</tr>
<tr>
<td>Conceptualizing a Data Infrastructure for the Capture and Use of Patient-Generated Health Data</td>
<td>Develop a policy framework for the use of patient-generated data in research and care delivery that addresses data collection tools, data donation policies, regulatory gaps, combining data with medical record data, and interoperability of data across health information systems and devices.</td>
<td>2.1</td>
</tr>
<tr>
<td>Standardization and Querying of Data Quality Metrics and Characteristics for Electronic Health Data</td>
<td>Create and implement a metadata standard data capture and querying system for data quality and characteristics, data source and institutional characteristics, and “fitness for use.”</td>
<td>1.8</td>
</tr>
<tr>
<td>Cross-Network Directory Service</td>
<td>Create an interoperable service that allows data partners to participate in multiple data research networks, query across the networks, and share analytic capabilities and knowledge across networks. The project will be piloted across two existing networks: Food and Drug Administration’s Sentinel and PCORnet.</td>
<td>1.5</td>
</tr>
<tr>
<td>Use of ADAPTABLE Trial to Strengthen Methods to Collect and Integrate Patient-Reported Information with Other Data Sets and Assess Its Validity</td>
<td>Generate tools and data standards that could be deployed in other CER studies by leveraging the infrastructure of an existing research study called the ADAPTABLE trial (Aspirin Dosing: A Patient-Centric Trial Assessing Benefits and Long Term Effectiveness). This trial is the first major randomized comparative effectiveness trial to be conducted by PCORnet.</td>
<td>1.1</td>
</tr>
</tbody>
</table>
### Appendix IV: Office of the Assistant Secretary for Planning and Evaluation's (ASPE) Projects to Build Data Capacity, as of September 30, 2017

<table>
<thead>
<tr>
<th>Project title</th>
<th>Description</th>
<th>Total obligations (dollars in millions)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Creation of LOINC Equivalence Classes</td>
<td>Create a flexible, extensible, and computable mechanism for rolling data into clinically relevant equivalence groups that enable more efficient processing aggregation of laboratory data and other data from diverse health information technology systems. The primary focus of this work will be on laboratory tests.</td>
<td>1.0</td>
</tr>
<tr>
<td>Source Data Capture from Electronic Health Records: Using Standardized Clinical Research Data</td>
<td>Create a single point data capture approach from the electronic health record to electronic data capture systems using the Retrieve Form for Data Capture standard. Stakeholders will be provided with a tool to seamlessly integrate electronic health record and electronic data capture systems.</td>
<td>0.7</td>
</tr>
</tbody>
</table>

*Source: GAO analysis of ASPE documentation. [GAO-18-311](#)*

---

*In addition to the projects listed, ASPE plans to obligate $2.0 million for one new project starting in fiscal year 2018.

*PCOR is a form of CER.

*ASPE plans to obligate an additional $0.4 million for this project in fiscal year 2018.

*ASPE plans to obligate an additional $0.4 million for this project in fiscal year 2018.

*ASPE plans to obligate an additional $0.1 million for this project in fiscal year 2018.

*ASPE plans to obligate an additional $2.0 million for this project in fiscal year 2019.

*ASPE plans to obligate an additional $1.4 million for this project in fiscal year 2018.
## Appendix V: GAO Contact and Staff

### Acknowledgments

In addition to the contact named above, Karin Wallestad, Assistant Director; Michael Zose, Analyst-in-Charge; Kye Briesath; Laurie Pachter; Vikki Porter, and Jennifer Whitworth made key contributions to this report.

<table>
<thead>
<tr>
<th>GAO Contact</th>
<th>John E. Dicken, (202) 512-7114 or <a href="mailto:dickenj@gao.gov">dickenj@gao.gov</a></th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Staff</strong></td>
<td>In addition to the contact named above, Karin Wallestad, Assistant Director; Michael Zose, Analyst-in-Charge; Kye Briesath; Laurie Pachter; Vikki Porter, and Jennifer Whitworth made key contributions to this report.</td>
</tr>
<tr>
<td><strong>Acknowledgments</strong></td>
<td></td>
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</tbody>
</table>
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