Spring 2014 Funding Cycle

PCORI Funding Announcement:
Communication and Dissemination Research

Published February 5, 2014
Latest Revision February 20, 2014

This PCORI Funding Announcement applies to the funding cycle that closes May 6, 2014, at 5:00 p.m. (ET). Application guidelines, templates, and other resources are available at pcori.org/PFA/spring-2014/communication.
About PCORI

PCORI is committed to transparency and a rigorous stakeholder-driven process that emphasizes patient engagement. PCORI uses a variety of forums and public comment periods to obtain public input to enhance its work. PCORI helps people make informed healthcare decisions and improves healthcare delivery and outcomes by producing and promoting high-integrity, evidence-based information that comes from research guided by patients, caregivers, and the broader healthcare community.

PCORI was authorized by the Patient Protection and Affordable Care Act of 2010 as a non-profit, nongovernmental organization. PCORI’s purpose, as defined by the law, is to help patients, clinicians, purchasers, and policy makers make better informed health decisions by “advancing the quality and relevance of evidence about how to prevent, diagnose, treat, monitor, and manage diseases, disorders, and other health conditions.”

Patient-Centered Outcomes Research Institute
1828 L St., NW, Suite 900
Washington, DC 20036
Phone: (202) 827-7700
Fax: (202) 355-9558
Email: info@pcori.org

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## Overview

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<td>Letter of Intent Due</td>
<td>March 7, 2014, by 5:00 p.m. (ET)</td>
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<tr>
<td>Summary</td>
<td>PCORI seeks to fund projects that address critical knowledge gaps in the communication and dissemination process—both the communication and dissemination of research results to patients, their caregivers, and clinicians, as well as the communication between patients, caregivers, and clinicians in the service of enabling patients and caregivers to make the best possible decisions in choosing among available options for care and treatment.</td>
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### Applicant Resources

See [pcori.org/pfa/spring-2014/communication](http://pcori.org/pfa/spring-2014/communication)

### Key Dates

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<td>Letter of Intent (LOI) Deadline</td>
<td>March 7, 2014, by 5:00 p.m. (ET)</td>
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<td>Applicant Town Hall Session</td>
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<td>Merit Review</td>
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<td>Awards Announced</td>
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<td>Earliest Project Start Date</td>
<td>December 2014</td>
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### Maximum Project Budget (Direct Costs)

$1.5 million

### Maximum Project Period

Three years

### Funds Available Up To (Direct Costs)

$8 million

### Eligibility

Applications may be submitted by any private sector research organization, including any nonprofit or for-profit organization, and any public sector research organization, including any university or college hospital or healthcare system, laboratory or manufacturer, or unit of local, state, or federal government. All US applicant organizations must be recognized by the Internal Revenue Service. Non-domestic components of organizations based in the United States and foreign organizations may apply, as long as there is demonstrable benefit to the US healthcare system and US efforts in the area of patient-centered research can be clearly shown. Organizations may submit multiple applications for funding. Individuals are not permitted to apply.
### Review Criteria

1. Impact of the condition on the health of individuals and populations  
2. Potential for the study to improve healthcare and outcomes  
3. Technical merit  
4. Patient-centeredness  
5. Patient and stakeholder engagement

### Budget/Time Limits

Applicants must submit a Greater Than Time/Budget Request with their Letter of Intent if the proposed project’s budget or duration exceed limits specified in this funding announcement.

### Contact Us

**Programmatic Inquiries:** Please contact the PCORI Helpdesk via email ([pfa@pcori.org](mailto:pfa@pcori.org)), phone (202-627-1884), or online ([http://www.pcori.org/PFA/inquiry](http://www.pcori.org/PFA/inquiry)). PCORI will provide a response within three business days. However, we cannot guarantee that all questions will be addressed three business days prior to a Letter of Intent or application deadline.

**Administrative, Financial, or Technical Inquiries:** Please contact the PCORI Helpdesk at [pfa@pcori.org](mailto:pfa@pcori.org). PCORI will provide a response within two business days. Please note that during the week of the application deadline, response times may exceed two business days. One week prior to an application deadline, applicants may also call the PCORI Helpdesk (202-627-1885). Applicants are asked to plan accordingly. It is the applicant’s responsibility to submit the application or before the application deadline.
I. Introduction

Summary of Program

The Patient-Centered Outcomes Research Institute (PCORI) invites applications to study the comparative effectiveness of strategies to increase patient and clinician awareness of the uncertainty associated with specific healthcare interventions, with the goal of increasing knowledge about—and the use of—comparative clinical effectiveness research (CER) results. This announcement is designed to include the context of the type of healthcare decision the patient faces as an important variable affecting the information needed and how it is provided. PCORI seeks studies that will provide information of value to patients, their caregivers and clinicians, as well as to healthcare leaders regarding which features of systems lead to better patient-centered outcomes.

Background

Every day, patients and their caregivers are faced with crucial healthcare decisions while lacking key information that they need. This information would not necessarily deliver verdicts or tell people what to do, but it would inform them of the trade-offs associated with the options they have—and enable them to make better decisions for themselves in collaboration with their clinicians—based on the facts, perhaps even personalized for them, and their own values, preferences, and goals.

PCORI is entrusted by the public to fund research that will matter to patients, their caregivers, and other stakeholders [defined as clinicians and clinician societies, hospitals, and health systems, payers (insurance), purchasers (business), industry, researchers, policymakers, and training institutions]. PCORI seeks to change how research is conducted by emphasizing the role of diverse research teams that include varying perspectives. PCORI distinguishes itself by supporting research in which patients, caregivers, practicing clinicians, and the broader stakeholder community are actively engaged in generating the research questions, reviewing the proposals, conducting the research, disseminating the findings, promoting the implementation of the findings, and using the results to understand and address patient and other stakeholder needs.

Because many patients and caregivers are not aware that they may have more than one viable option for prevention, diagnosis, or treatment decisions, the value of CER may not be immediately recognized. However, strategies can be developed to increase patient and clinician awareness of the uncertainty associated with specific healthcare interventions, with the goal of increasing knowledge about—and the use of—CER results. It should be noted that the type of healthcare decision the patient faces is an important variable affecting the information needed and how it is provided. (For example, the information needs of a patient weighing options for treating high blood pressure will be different from those of a patient facing a terminal cancer diagnosis with complicated treatment options.) Additionally, although a majority of patients prefers an active role in clinical decision making, the reasons some choose not to participate are unclear. Knowledge gaps in this area include the role of cultural norms and values in shaping preferences for participation in clinical decision making. Communication skills of both patients and healthcare providers are an important issue for the effective use of CER results. Research on doctor-patient communication has focused primarily on the doctor-
patient dyad, but still little is known about the potential role of the patient’s family members or significant others in shaping the decision-making process.

**Clinician Engagement with CER**

Changes in practice on the part of providers in response to CER has been limited. It is unclear which methods for translating CER results into clinical care will prove to be most effective in terms of reaching the greatest proportion of patients and improving patient outcomes. Further research is needed to understand clinicians’ attitudes toward CER and shared medical decision making. Strategies can then be developed to increase clinicians’ utilization of CER and to increase clinicians’ willingness to engage their patients in the decision-making process. Little is known about how clinical decision making could be structured to reduce the potential time burden in individual clinical encounters. Additional information is also needed on how community-based healthcare resources are engaging, if at all, with CER findings.

**Translating Research, Decision Support Interventions, and Risk Communication**

Another important area of research in both clinical and community-based settings is translating existing scientific research into accessible and usable formats that clearly outline the risks and benefits of preventive, diagnostic, and treatment options for patients, caregivers, and healthcare providers. In clinical care, decision support intervention is one of the primary ways in which medical evidence is translated into a format that is usable by patients, families, and caregivers. The integration of patient decision support, electronic medical records, and associated patient systems holds considerable promise, but little, if any, evidence is available to guide best practices. More research is needed about how decision support interventions perform using different media, what level of information and detail they require, and how they perform in patient populations with lower levels of literacy and numeracy. A further significant gap is the limited research on risk communication, in general, and with underserved individuals and those with limited health literacy and numeracy, in particular. To date, research on effective methods for communicating risk information to healthcare providers and enabling them to use the information effectively is lacking.

As part of the current funded portfolio, PCORI has funded many decision making projects, including those studies that test a decision aid. PCORI wants to balance its decision-making research portfolio. To that end, PCORI is developing a framework for supporting a diverse set of research topics in decision making. For the current funding announcement, PCORI welcomes proposals about decision making but does not encourage new proposals that develop and test decision aids. PCORI welcomes resubmissions that test decision aids, but the number of awards will be limited within this funding cycle.

**Distribution of CER**

The distribution of CER information to patients, caregivers, and providers (in both clinical and community-based settings) is an area that has not received sufficient research attention. Little is known about which methods and approaches are most effective or the various impacts of different approaches. More research is needed to identify effective approaches to distribute CER results to healthcare providers, with the goals of sustained changes in clinical practice and effective distribution of results to patients in order to enable changes in behavior (for example, adherence and self-care). Research is also needed to identify trusted intermediaries and trusted channels of communication most often turned to by patients, caregivers, and clinicians.
Additionally, further investigation is needed to explore how strategies used in public health communication and social marketing can be adapted to distributing the results of CER, and to identify creative ways of combining multiple channels of communication and dissemination to increase exposure to CER. Further exploration is also needed to understand the disparities that may remain regarding access to social media resources to ensure that the “e-health revolution” does not widen existing health-related knowledge gaps among low-income and racial and ethnic minority populations. Finally, further research is needed to examine the reliability of any CER data currently available through social media sites and to understand how individuals evaluate and use this information in their prevention, screening, diagnosis, and treatment decision-making processes. More specifically, there is a lack of information on how these media may influence patient self-care and adherence to treatment recommendations.

**Research of Interest**

The Communication and Dissemination Research program is interested in the following broad topical areas:

- Research that compares alternative communication, dissemination, health literacy and/or implementation strategies that aim to improve patients’ health outcomes, by increasing patient, caregiver, and/or provider awareness of healthcare options in clinical or community-based settings.
- Research that compares the effectiveness of alternative approaches across a range of patient-centered outcomes to increase or encourage effective patient, caregiver, or clinician participation in care decisions and in shared decision making.
- Studies to develop and compare alternative methods and tools to elicit and include patient-desired outcomes in the healthcare decision-making process.
- Studies comparing alternative approaches, including use of public health strategies or social media, for providing new information to patients, caregivers, or clinicians, with attention to differences in effectiveness in different populations.
- Research that compares innovative approaches in the use of existing electronic clinical data and other electronic modalities from the healthcare system or from a network of systems to enhance clinical decision making by patients and providers.

Research studies may focus on patient populations with a single condition or involve patients with a range of conditions. Studies addressing care for patients with rare conditions are of interest. Rare diseases are defined as life-threatening or chronically debilitating diseases that are of such low prevalence in populations that special efforts, such as combining data across large populations, may be needed to address them. By “low prevalence” we mean conditions that affect fewer than 200,000 individuals in the United States or have a prevalence of less than 1 in 1,500 persons.

**Sample Research Questions**
The following research questions are meant as examples of the types of questions that your research may help answer. This list is by no means exhaustive. **All questions must have a comparative component.**

- How do designs for decision support interventions compare in their ability to assist patients and/or caregivers with lower levels of literacy/numeracy, and how do strategies for communicating risk information to vulnerable populations compare?
- How do methods for distributing CER findings to patients, caregivers, or healthcare providers compare in their ability to improve patients’ health outcomes?
- To whom are clinicians most likely to turn for trustworthy information about the effectiveness, relative effectiveness, benefits, and harms of different treatment options for a given condition, and how do they access that information?
- How do strategies learned from public health communication and social marketing compare in their ability to promote the distribution of CER to patients and/or their caregivers and to their clinicians?
- How do strategies in community-based settings compare with strategies in clinical-based settings in their ability to promote the distribution of CER to patients and/or their caregivers?
- How—and how effectively—can strategies using social media be deployed to distribute CER to patients and/or their caregivers and to their clinicians?
- How do patient outcomes compare when patient preferences around screening, diagnosis, treatment, and management strategies have been elicited and accounted for in the decision-making process?
- How do strategies compare in their ability to effectively engage patients with lower levels of literacy and/or numeracy in clinical decision making?
- How do strategies for training healthcare providers in imparting information about risk to patients and their caregivers compare in their ability to improve patient outcomes?
- How do interventions to promote shared decision making compare in their ability to influence patients’ health behaviors and self-care (e.g., adherence to medication) or patients’ behavior in the clinical encounter?

II. **Guidance for Proposing Research**

**Research Priorities**

PCORI funds patient-centered outcomes research, a type of comparative clinical effectiveness research. The studies PCORI supports must include the perspectives of patients and other healthcare stakeholders. To be considered responsive, applications must describe research that:

- Studies the benefits and harms of interventions and strategies delivered in actual settings. By “delivered in actual settings,” we mean delivered and received in
typical “real-life” clinical settings, not just in restrictive trials of experimental care or at selected academic centers. PCORI is interested in innovative studies that provide practical information that can help patients and other stakeholders make informed decisions about their health care and health outcomes.

- Compares at least two alternative approaches. The types of interventions tested can include specific drugs, devices, and procedures, as well as other types of alternatives, such as medical and assistive devices and technologies, diagnostic testing, behavioral change, and a wide variety of strategies for improving delivery systems, but the studies must be comparative. “Usual care” (or no specific intervention) may be an appropriate comparator if this is a realistic choice faced by patients and other stakeholders, but the clinical characteristics must be specified.

- Is based on health outcomes that are meaningful to the patient population under study. While most patient-centered outcomes directly impact the patient’s quality of life, certain physiological measurements, such as blood pressure and serum cholesterol, are strongly linked to complications or other outcomes that patients care about and have become outcomes of interest to patients because of increased awareness. Therefore, an application to PCORI that proposes to conduct a study comparing two approaches to helping people control their blood pressure would be well-aligned with PCORI’s focus on patient-centeredness, assuming that the study would also compare the two approaches’ effects on any other relevant outcomes that are important to patients, such as treatment-related symptoms (side effects).

Non-responsiveness

Applications will be considered non-responsive if the proposed research:

- Conducts a formal cost-effectiveness analysis in the form of dollar-cost per quality-adjusted life-year to compare two or more alternatives.

- Measures the relative costs of care of two or more alternative approaches as the primary criteria for choosing the preferred alternative.

PCORI does have an interest, however, in studies that address questions about conditions that lead to high costs to the individual or to society. This is included in our review criterion on impact of the condition on the health of individuals and populations. PCORI is also interested in studies that examine differentials in healthcare resources or costs as a determinant of, or barrier to, good outcomes. Examples include ways in which out-of-pocket costs may constitute a barrier to the receipt of care.

Further, PCORI considers it important for applicants to discuss cost-related issues such as the resources needed to implement, replicate or disseminate a successful intervention. PCORI also is interested in evaluation of interventions intended to reduce health system waste or increase health system efficiency. Proposals that include studies of these issues without utilizing a formal cost-effectiveness analysis or comparing the costs of alternatives are considered responsive.
PCORI discourages proposals that include studies of the natural history of disease, instrument development, pharmacodynamics, and fundamental science or study of biological mechanisms. It also discourages studies that have the primary purpose of developing and evaluating new decision aids or clinical prognostication tools.

Features of Patient-Centered Outcomes Research

PCOR helps people and their caregivers communicate and make informed healthcare decisions, allowing their voices to be heard in assessing the value of healthcare options. This research:

- Assesses the benefits and harms of preventive, diagnostic, therapeutic, palliative, or health-delivery-system features to inform decision making, highlighting the choices that matter to people.
- Is inclusive of an individual’s preferences, autonomy, and needs, focusing on outcomes that people notice and care about, such as survival, functioning, symptoms, and health-related quality of life.
- Incorporates a wide variety of settings and diversity of participants to address individual differences and barriers to implementation and dissemination.
- Directly compares clinical interventions that are generally available in the clinical settings that people use to access health care.
- Obtains the perspectives of stakeholders to address the burdens to individuals, availability of services, and requirements for technology and personnel.

Comparative Clinical Effectiveness Research

Applications submitted in response to this funding announcement should evaluate the comparison of two or more clinical interventions. The efficacy of each intervention must be known prior to the initiation of the proposed project. The application must provide information from systematic reviews or other credible literature reviews on the nature of the research gap being addressed and the data about efficacy of the clinical interventions that will be compared. Projects that aim to develop new or novel interventions will be considered out of scope.

Leveraging Existing Resources

Investigators are encouraged to propose studies that leverage existing resources, such as adding PCOR to an existing large clinical trial or analyzing existing large databases that contain valuable, relevant information that may be used to answer important comparative clinical effectiveness research questions. PCORI is also interested in seeking proposals for meta-analyses that use individual participant data.

Preliminary Data and Use of Accepted Measures

PCORI encourages investigators to design their research using valid patient-centered outcomes measures. Include preliminary data that supports the proposed measures. Investigators are encouraged to consider those
measures described in the Patient Reported Outcomes Measurement Information System (PROMIS). ¹

Documentation of Assumptions

PCORI specifically seeks studies that are sufficiently powered to detect clinically meaningful effects. To that end, please justify the proposed sample sizes by explaining the assumptions used in all study power calculations. The application should clearly state all the necessary assumptions (i.e., the primary outcome measure, the estimated difference in the mean value of this measure between study arms, standard deviation of the measure, type I error rate, and any other assumptions). All such estimates must be justified by referring to prior published research or preliminary data.

Studies in Rare Diseases

PCORI is interested in the investigation of strategies that address care for patients with rare conditions. Rare diseases are defined as life-threatening or chronically debilitating diseases that are of such low prevalence in populations that special efforts, such as combining data across large populations, may be needed to address them. The term low prevalence is defined as conditions that affect fewer than 200,000 individuals in the United States or have a prevalence of less than 1 in 1,500 persons.

Review Criteria

PCORI’s review panels rate all submitted applications on the following five criteria:

Criterion 1. Impact of the condition on the health of individuals and populations
The proposal addresses the following questions:
• Is the condition or disease associated with a significant burden in the US population, in terms of prevalence, mortality, morbidity, individual suffering, or loss of productivity?
• Alternatively, does the condition or disease impose a significant burden on a smaller number of people who have a rare disease?
• Does the proposal include a particular emphasis on patients with one or more chronic condition?

Criterion 2. Potential for the study to improve health care and outcomes
The proposal has the potential to lead to meaningful improvement in the quality and efficiency of care and to improvements in outcomes that are important to patients. It addresses the following questions:
• Does the research question address a critical gap in current knowledge as noted in systematic reviews, guideline development efforts, or previous research prioritizations?
• Has it been identified as important by patient, caregiver, or clinician groups?
• Do wide variations in practice patterns suggest current clinical uncertainty?
• Is the research novel or innovative in its methods or approach, in the population?

¹ Available at nihpromis.org
being studied, or in the intervention being evaluated in ways that make it likely to improve care?

- Do preliminary studies indicate potential for a sizeable benefit of the intervention relative to current practice? How likely is it that positive findings could be disseminated and implemented quickly, resulting in improvements in practice and patient outcomes?

**Criterion 3. Technical merit**

The proposal has sufficient technical merit to ensure that the study goals will be met. It includes:

- A clear research plan with rigorous methods that demonstrates adherence to PCORI’s Methodology Standards
- A realistic timeline that includes specific scientific and engagement milestones
- A research team with the necessary expertise and an appropriate organizational structure
- A research environment sufficient to support the conduct of the work with appropriate resources
- A diverse study population with respect to age, gender, race, ethnicity, and clinical status, as appropriate for the proposed research

**Criterion 4. Patient-centeredness**

The proposal demonstrates patient-centeredness at every stage of the research. It addresses the following questions:

- Is the research focused on questions that affect outcomes of interest to patients and their caregivers?
- Does the research address one or more of the key questions mentioned in PCORI’s definition of patient-centered outcomes research?

**Criterion 5. Patient and stakeholder engagement**

The proposal demonstrates that people representing the population of interest and other relevant stakeholders are engaged in ways that are appropriate and necessary in a given research context.

- Are patients and other stakeholders engaged in:
  - Formulating research questions
  - Defining essential characteristics of study participants, comparators, and outcomes
  - Identifying and selecting outcomes that the population of interest notices and cares about (e.g., survival, function, symptoms, health-related quality of life) and that inform decision making relevant to the research topic
  - Monitoring study conduct and progress
  - Designing/suggesting plans for dissemination and implementation activities

- Are the roles and the decision making authority of all research partners clearly stated?
- Does the proposal demonstrate the principles of reciprocal relationships, co-
learning, partnership, trust, transparency, and honesty?

Patient and Stakeholder Engagement

PCORI encourages all applicants to clearly describe the patient and stakeholder engagement in their research proposals. PCORI understands that patient and stakeholder engagement in research can take many forms; it is not seeking one particular type or method of engagement. Rather, applicants should communicate how patients (those with lived experience), family members, caregivers, and the organizations that represent them, as well as any other relevant stakeholders, will be involved in study activities. Because this type of engagement in research is a relatively new concept, PCORI has developed a Patient and Family Engagement Rubric (see the appendix to the Engagement Template) to guide both applicants and merit reviewers. Additionally, studies are expected to adhere to PCORI’s Methodology Standards Associated with Patient-Centeredness and to the PCOR Engagement Principles found within the rubric. These and additional resources are available in PCORI’s Funding Center.²

Methodological Considerations

Regardless of study design, proposals must adhere to all relevant PCORI Methodology Standards. PCORI Methodology Standards include 47 individual standards that fall into 11 categories. The first five categories are cross-cutting and are relevant to most PCOR studies. Researchers should refer to all of these standards when planning and conducting their research projects. These categories are:

- Standards for Formulating Research Questions
- Standards Associated with Patient-Centeredness
- Standards on Data Integrity and Rigorous Analyses
- Standards for Preventing and Handling Missing Data
- Standards for Heterogeneity of Treatment Effect (HTE)

Six other categories of standards will be applicable to particular study designs and methods. The standards in each of these categories should be used for guidance when they are relevant to a particular study:

- Standards for Data Registries
- Standards for Data Networks as Research-facilitating Infrastructures
- Standards for Causal Inference Methods
- Standards for Adaptive and Bayesian Trial Designs
- Standards for Studies of Diagnostic Tests
- Standards for Systematic Reviews

² Available at pcori.org/apply
Most of these standards should be considered “minimal” standards. Additional best practices, including relevant guidelines for the conduct of clinical trials developed by other organizations, should be addressed in the application for PCORI funding.

All applicants should specifically discuss their capacity to measure factors such as differential adherence to chosen treatments (or participation in intervention programs) that could create or explain apparent differences in the effectiveness of the alternative interventions being compared in clinical populations.

Populations Studied

PCORI seeks to fund research that includes diverse populations with respect to age, gender, race, ethnicity, geography, or clinical status, so that possible differences in comparative clinical effectiveness research may be examined. PCORI recognizes that some proposed studies may represent important PCOR opportunities even in the absence of a broadly diverse study population. However, the burden is on the applicant in such cases to justify the importance of the study in the absence of diversity. Alternatively, PCORI is interested in the inclusion of previously understudied populations for whom effectiveness information is particularly needed, such as hard-to-reach populations or patients with multiple conditions. Thus, comparisons should examine the impact of the strategies in various subpopulations with attention to the possibilities that the effects of the strategy might differ across various populations. Populations of interest include those that are less frequently studied. PCORI has developed the following list of priority populations to guide our efforts in research and engagement, which includes:

- Racial and ethnic minority groups
- Low-income groups
- Women
- Children (age 0–17 years)
- Older adults (age 65 years and older)
- Residents of rural areas
- Individuals with special healthcare needs, including individuals with disabilities
- Individuals with multiple chronic diseases
- Individuals with rare diseases
- Individuals whose genetic make-up affects their medical outcomes
- Patients with low health literacy/numeracy and/or limited English proficiency
- Lesbian, gay, bisexual, and transsexual (LGBT) persons

Protection of Human Subjects

PCORI adopts, by reference, the Human Subjects requirements of 45 CFR Part 46. If the proposed research will involve human subjects, refer to the Supplemental Instructions for Preparing the Protection of Human Subjects Section of the Research Plan in Part II of the Instructions for the PHS 398 Form provided by the National
Institute of Health. Note: PCORI requires engagement in the research by patients and/or other stakeholders, as research partners. Research subjects protection requirements do not apply to co-investigators, members of the research team, or research partners.

Budget and Project Duration

If your proposed project exceeds $1.5 million in total direct costs and/or exceeds three years, please submit, before the LOI deadline, a Greater Than request form with a justification for the increased budget and/or extended length of your study. The form will be reviewed by the program staff, and you will receive a notification for approval or denial of this request within two weeks of the LOI deadline.

III. How to Submit a Proposal

Submission Dates

Letters of Intent and applications must be submitted in accordance with the published dates and times listed in the Overview in this PFA and in the PCORI Funding Center.

PCORI Online System

To submit a proposal, you must register with the PCORI Online System and submit both a Letter of Intent (LOI) and an application for each cycle in which you are applying.

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3 Available at grants.nih.gov/grants/funding/phs398/phs398.html
4 Available at pcori.org/funding-center
5 Available at https://pcori.fluxx.io
Applicant Resources

PCORI Funding Center          pcori.org/pfa/spring-2014/communication
PCORI Online System          https://pcori.fluxx.io
PCORI Funding Awards          pcori.org/pfaawards

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