Supplemental Evidence and Data Request on Maternal and Fetal Effects of Mental Health Treatments in Pregnant and Breastfeeding Women: A Systematic Review of Pharmacological Interventions

AGENCY: Agency for Healthcare Research and Quality (AHRQ), HHS.

ACTION: Request for supplemental evidence and data submissions.

SUMMARY: The Agency for Healthcare Research and Quality (AHRQ) is seeking scientific information submissions from the public. Scientific information is being solicited to inform our review on Maternal and Fetal Effects of Mental Health Treatments in Pregnant and Breastfeeding Women: A Systematic Review of Pharmacological Interventions, which is currently being conducted by the AHRQ’s Evidence-based Practice Centers (EPC) Program. Access to published and unpublished pertinent scientific information will improve the quality of this review.

DATES: Submission Deadline on or before 30 days after date of publication.

ADDRESSES: Email Submissions: epc@ahrq.hhs.gov.

Print Submissions:
Mailing Address: Center for Evidence and Practice Improvement, Agency for Healthcare Research and Quality, ATTN: EPC SEADs Coordinator, 5600 Fishers Lane, Mail Stop 06E53A, Rockville, MD 20857.
Shipping Address (FedEx, UPS, etc.): Center for Evidence and Practice Improvement, Agency for Healthcare Research and Quality, ATTN: EPC SEADs Coordinator, 5600 Fishers Lane, Mail Stop 06E77D, Rockville, MD 20857.

FOR FURTHER INFORMATION CONTACT: Jenae Benns, Telephone: 301–427–1496 or Email: epc@ahrq.hhs.gov.

The EPC Program is dedicated to identifying as many studies as possible that are relevant to the questions for each of the reviews. In order to do so, we are supplementing the usual manual and electronic database searches of the literature by requesting information from the public (e.g., details of studies conducted). We are looking for studies that report on Maternal and Fetal Effects of Mental Health Treatments in Pregnant and Breastfeeding Women: A Systematic Review of Pharmacological Interventions, including those that describe adverse events. The entire research protocol, including the key questions, is also available online at: https://effectivehealthcare.ahrq.gov/topics/mental-health-pregnancy/protocol.

This is to notify the public that the EPC Program would find the following information on Maternal and Fetal Effects of Mental Health Treatments in Pregnant and Breastfeeding Women: A Systematic Review of Pharmacological Interventions helpful:

- A list of completed studies that your organization has sponsored for this indication. In the list, please indicate whether results are available on ClinicalTrials.gov along with the ClinicalTrials.gov trial number.
- For completed studies that do not have results on ClinicalTrials.gov, a summary, including the following elements: Study number, study period, design, methodology, indication and diagnosis, proper use instructions, inclusion and exclusion criteria, primary and secondary outcomes, baseline characteristics, number of patients screened/eligible/enrolled/lost to follow-up/withdrawn/analyzed, effectiveness/efficacy, and safety results.
- A list of ongoing studies that your organization has sponsored for this indication. In the list, please provide the ClinicalTrials.gov trial number or, if the trial is not registered, the protocol for the study including a study number, the study period, design, methodology, indication and diagnosis, proper use instructions, inclusion and exclusion criteria, and primary and secondary outcomes.
- Description of whether the above studies constitute ALL Phase II and above clinical trials sponsored by your organization for this indication and an index outlining the relevant information in each submitted file.

Your contribution is very beneficial to the Program. Materials submitted must be publicly available or able to be made public. Materials that are considered confidential; marketing materials; study types not included in the review; or information on indications not included
in the review cannot be used by the EPC Program. This is a voluntary request for information, and all costs for complying with this request must be borne by the submitter.

The draft of this review will be posted on AHRQ’s EPC Program website and available for public comment for a period of four weeks. If you would like to be notified when the draft is posted, please sign up for the email list at: https://www.effectivehealthcare.ahrq.gov/email-updates.

The systematic review will answer the following questions. This information is provided as background. AHRQ is not requesting that the public provide answers to these questions.

The Key Questions

**Key Question 1:** Among pregnant and postpartum women, what is the effectiveness of pharmacologic interventions on maternal outcomes?

a. Among those with a new or preexisting anxiety disorder?

b. Among those with a new or preexisting depressive disorder?

c. Among those with a new or preexisting bipolar disorder?

d. Among those with new or preexisting schizophrenia?

**Key Question 2:** Among pregnant and postpartum women, what is the comparative effectiveness of pharmacologic interventions on maternal outcomes?

a. Among those with a new or preexisting anxiety disorder?

b. Among those with a new or preexisting depressive disorder?

c. Among those with a new or preexisting bipolar disorder?

d. Among those with new or preexisting schizophrenia?

**Key Question 3:** Among reproductive-aged women with any mental health disorder, what are the maternal and fetal harms associated with pharmacologic interventions for a mental health disorder during preconception, pregnancy, and postpartum?

**Key Question 4:** Among reproductive-aged women with any mental health disorder, what are the comparative maternal and fetal harms of pharmacologic interventions for a mental health disorder during preconception, pregnancy, and postpartum?

**Contextual Question 1:** Among women who are preconceptional, pregnant, or postpartum, within a given disorder, what are the harms of NOT treating or stopping a pharmacological treatment, or of switching medications?

<table>
<thead>
<tr>
<th>TABLE 1—PICOTS (POPULATIONS, INTERVENTIONS, COMPARATORS, OUTCOMES, TIMING, SETTINGS) AND INCLUSION/EXCLUSION CRITERIA</th>
</tr>
</thead>
<tbody>
<tr>
<td>PICOTS</td>
</tr>
<tr>
<td>Population ...</td>
</tr>
<tr>
<td></td>
</tr>
<tr>
<td></td>
</tr>
<tr>
<td></td>
</tr>
<tr>
<td>Intervention †</td>
</tr>
<tr>
<td></td>
</tr>
<tr>
<td></td>
</tr>
<tr>
<td></td>
</tr>
<tr>
<td></td>
</tr>
<tr>
<td></td>
</tr>
<tr>
<td></td>
</tr>
<tr>
<td></td>
</tr>
<tr>
<td></td>
</tr>
<tr>
<td></td>
</tr>
<tr>
<td>Comparator ...</td>
</tr>
<tr>
<td></td>
</tr>
<tr>
<td>Outcomes ‡ ...</td>
</tr>
<tr>
<td></td>
</tr>
<tr>
<td></td>
</tr>
<tr>
<td></td>
</tr>
<tr>
<td></td>
</tr>
<tr>
<td></td>
</tr>
<tr>
<td></td>
</tr>
<tr>
<td></td>
</tr>
<tr>
<td></td>
</tr>
<tr>
<td></td>
</tr>
</tbody>
</table>
Disease Registry.

Agency for Toxic Substances and Disease Registry.

September 8, 2017) is amended to

recently at 82 FR 42555, dated

June 17, 1985, as amended most

most

Services (50 FR 25129–25130, dated

Section J–C, Order of Succession:

Dated: December 4, 2019.

Virginia Mackay-Smith,
Associate Director.

FR Doc. 2019–26510 Filed 12–9–19; 8:45 am
BILLING CODE 4160–90–P

DEPARTMENT OF HEALTH AND HUMAN SERVICES

Agency for Toxic Substances and Disease Registry

Statement of Organization, Functions, and Delegations of Authority

Part J (Agency for Toxic Substances and Disease Registry) of the Statement of Organization, Functions, and Delegations of Authority of the Department of Health and Human Services (50 FR 25129–25130, dated June 17, 1985, as amended most recently at 82 FR 42555, dated September 8, 2017) is amended to reflect the Order of Succession for the Agency for Toxic Substances and Disease Registry.

Section J–C, Order of Succession:

Delete in its entirety the Section J–C, Order of Succession, and insert the following:

During the absence or disability of the Administrator, Agency for Toxic Substances and Disease Registry (ATSDR), or in the event of a vacancy in that office, the first official listed below who is available shall act as Administrator, except during a planned period of absence, the Administrator may specify a different order of succession:

1. Assistant Administrator, ATSDR
2. Deputy Director for Non-Infectious Diseases
3. Principal Deputy Director
4. Chief Medical Officer
5. Director, Center for Preparedness and Response

Sherri A. Berger,
Chief Operating Officer, Centers for Disease Control and Prevention.

FR Doc. 2019–26494 Filed 12–9–19; 8:45 am
BILLING CODE 4163–70–P

DEPARTMENT OF HEALTH AND HUMAN SERVICES

Centers for Medicare & Medicaid Services


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

AGENCY: Centers for Medicare & Medicaid Services, HHS.

ACTION: Notice.

SUMMARY: The Centers for Medicare & Medicaid Services (CMS) is announcing an opportunity for the public to comment on CMS’ intention to collect information from the public. Under the Paperwork Reduction Act of 1995 (PRA), federal agencies are required to publish notice in the Federal Register concerning each proposed collection of information, including each proposed extension or reinstatement of an existing collection of information, and to allow a second opportunity for public comment on the notice. Interested

TABLE 1—PICOTS (POPULATIONS, INTERVENTIONS, COMPARATORS, OUTCOMES, TIMING, SETTINGS) AND INCLUSION/EXCLUSION CRITERIA—Continued

<table>
<thead>
<tr>
<th>PICOTS</th>
<th>Inclusion</th>
<th>Exclusion</th>
</tr>
</thead>
</table>

Time frame ... Followup .................................................................
KQ 1, KQ 2: From conception up to 1 year postpartum for maternal outcomes.
KQ 3, KQ 4: All .................................................................

Followup
• KQ 1, KQ 2: More than 12 weeks preconception for maternal preconception outcomes, more than 1 year for maternal postpartum outcomes
• KQ 3, KQ 4: None.

Settings § ..... Clinical setting .................................................................
All settings .................................................................

Clinical setting
None.

Study design
• RCTs, CCTs, case-control studies, cohort studies with comparison arms.
• Reference lists of relevant systematic reviews published in 2013 or later will be used to ensure our search strategies captured all relevant studies.

All other designs and studies using included designs that do not meet the sample size criterion.

Language ..... Studies published in English .................................................................
Studies published in languages other than English.

* We will limit included outcomes to those using validated measures. Another potential exclusion, depending on volume of yield, includes studies that fail to control for confounding.
† Drugs such as brexanolone that are awaiting FDA approval will be included in the review once they are approved
‡ We will focus strength of evidence (SOE) grades on outcomes prioritized by the Technical Expert Panel (TEP).
§ Depending on volume, we may limit the primary analysis to studies from geographic settings with resources comparable or applicable to the United States.