

Comments regarding each of these applications must be received at the Reserve Bank indicated or the offices of the Board of Governors, Ann E. Misback, Secretary of the Board, 20th Street and Constitution Avenue NW, Washington, DC 20551-0001, not later than December 4, 2023.

A. Federal Reserve Bank of Boston (Prabal Chakrabarti, Senior Vice President) 600 Atlantic Avenue, Boston, Massachusetts 02210-2204. Comments can also be sent electronically to BOS.SRC.Applications.Comments@bos.frb.org:

1. *Mutual Bancorp MHC, Hyannis, Massachusetts*; to merge with Fidelity Mutual Holding Company Leominster, Massachusetts, and thereby indirectly acquire Life Design Holding Company, Hyannis, Massachusetts, and Fidelity Co-Operative Bank, Leominster, Massachusetts.

Board of Governors of the Federal Reserve System.

Michele Taylor Fennell,

Deputy Associate Secretary of the Board.

[FR Doc. 2023-24243 Filed 11-1-23; 8:45 am]

BILLING CODE P

DEPARTMENT OF HEALTH AND HUMAN SERVICES

Agency for Healthcare Research and Quality

Supplemental Evidence and Data Request on Trauma Informed Care

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

ACTION: Request for supplemental evidence and data submission.

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 *Trauma Informed Care*, 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 Access to published and unpublished pertinent scientific information will improve the quality of this review.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 December 4, 2023.

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: Kelly Carper, Telephone: 301-427-1656 or Email: epc@ahrq.hhs.gov.

SUPPLEMENTARY INFORMATION: The Agency for Healthcare Research and Quality has commissioned the Evidence-based Practice Centers (EPC) Program to complete a review of the evidence for *Trauma Informed Care*. AHRQ is conducting this review pursuant to Section 902 of the Public Health Service Act, 42 U.S.C. 299a.

The EPC Program is dedicated to identifying as many studies as possible that are relevant to the questions for each of its 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 *Trauma Informed Care*. The entire research protocol is available online at: <https://effectivehealthcare.ahrq.gov/products/trauma-informed-care/protocol>.

This is to notify the public that the EPC Program would find the following information on *Trauma Informed Care* helpful:

- A list of completed studies that your organization has sponsored for this topic. 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, if relevant: 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 topic.* In the list, please provide the *ClinicalTrials.gov* trial number or, if the trial is not registered, the protocol for the study including, if relevant, 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 topic 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 topics 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 4 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 review will answer the following questions. This information is provided as background. AHRQ is not requesting that the public provide answers to these questions.

Key Questions (KQ)

TIC for Adult Patients/Clients

- KQ 1. What is the evidence of benefits and/or harms of TIC on outcomes for patients/clients?
 - KQ 1a. Which components (e.g., education and training of providers about trauma, screening patients, delivering point-of-care interventions [note this is not meant to include established evidence-based treatments for trauma-related disorders], referring patients/clients for various forms of additional assessment and treatment for indicated needs) of TIC models, and organizational and practice characteristics, are associated with benefits and/or harms?
 - KQ 1b. Do outcomes vary by patient/client or clinical or organizational characteristics, including the nature, extent and timing of exposure (e.g., recent or ongoing vs. prior exposure in childhood)?

TIC for Child and Adolescent Patients/Clients

- KQ 2. What is the evidence of benefits and/or harms of TIC on outcomes for patients/clients?
 - KQ 2a. Which components (e.g., education and training of providers about trauma, screening patients, delivering point-of-care interventions [note this is not meant to include indicated evidence-based treatments for trauma-related disorders], referring clients for various forms of additional assessment and treatment for indicated needs) of TIC models, organizational and practice characteristics, are associated with benefits and/or harms?

○ KQ 2b. Do outcomes vary by patient/client (as well as parent) or clinical or organizational characteristics including the nature, extent, and timing of exposure (e.g., recent or ongoing vs. prior exposure)?

PICOTS (POPULATIONS, INTERVENTIONS, COMPARATORS, OUTCOMES, TIMING, AND SETTING)

PICOTS	KQ1	KQ2
Population	Adults 18 years and older, regardless of trauma exposure. 1b. Patient/client and clinical characteristics including type, time since, and duration of trauma exposure; gender; race/ethnicity; age; clinical condition; or disorder (e.g., anxiety, depression, substance use).	Youth <18 years, regardless of trauma exposure. 2b. Patient/client and clinical characteristics including type, time since, and duration of trauma exposure; gender; race/ethnicity; age; clinical condition; or disorder, (e.g., anxiety, depression, ADHD, conduct disorder, substance use).
Intervention	TIC models/components of care (e.g., education and training of providers about trauma, screening patients/clients for trauma exposure using ACEs or other tools, screening for symptoms, delivering point-of-care interventions, referring patients/clients for various forms of additional assessment and treatment for indicated needs). 1a. single or multi-component, individual or group, targeting organizations, providers, patients/clients, caregivers, or a combination, training, screening.	TIC models/components of care (e.g., education and training of providers about trauma, screening patients/clients for trauma exposure using ACEs or other tools, screening for symptoms, delivering point of care interventions, referring patients/clients for various forms of additional assessment and treatment for indicated needs). 2a. single or multi-component, individual or group, targeting organizations, providers, patients/clients, caregivers, or a combination, training, screening.
Comparator	No TIC model of care/usual or routine care (CAU) Other TIC model or component(s) of care, evidence-based therapies for trauma-related conditions (e.g., prolonged exposure, cognitive processing therapy) or approaches (e.g., Collaborative Care).	No TIC model of care/usual or routine care (CAU). Other TIC model or component(s) of care, evidence-based therapies for trauma-related conditions (e.g., trauma-focused CBT) or approaches (e.g., Collaborative Care).
Outcome	<i>Trauma-Specific:</i> Additional or repeat trauma exposure from the point-of-care in the course of care/service delivery (e.g., retraumatization). <i>Process outcomes:</i> Health care outcomes/utilization/referral, provider burnout/mental health. <i>Organizational/practice/systems outcomes:</i> Intake and referral processes (e.g., wait times), disseminated policies, trainings, staffing (e.g., scribes), administrative requirements, access to treatment, workforce diversity. <i>Patient/client-centered outcomes:</i> Physical and mental health outcomes, functioning, clinical improvement, patient/client engagement, trust, comfort or satisfaction, and strengths-based outcomes (e.g., quality of life). <i>Harms:</i> Includes displacement of evidence based care (e.g., screening for anxiety, depression, substance use, suicide risk), increase in patient/client aggression or other behavioral misconduct.	<i>Trauma-Specific:</i> Additional or repeat trauma exposure from the point-of-care in the course of care/service delivery (e.g., retraumatization). <i>Process outcomes:</i> Healthcare outcomes/utilization/referral, provider outcomes burnout/mental health. <i>Organizational/practice/systems outcomes:</i> Intake and referral processes (e.g., wait times), disseminated policies, trainings, staffing (e.g., scribes), administrative requirements, access to treatment, workforce diversity, anti-racism principles. <i>Patient/client-centered outcomes:</i> Physical and mental health outcomes, functioning, clinical improvement, patient/client engagement, trust, comfort or satisfaction, and strengths-based outcomes (e.g., quality of life). <i>Harms:</i> Includes displacement of evidence based care (e.g., screening for developmental milestones, ADHD, depression, anxiety, suicide risk, substance use), increase in patient/client aggression or other behavioral misconduct.
Timing	Any	Any.
Setting	Routine or emergency healthcare in any setting that provides human or social services, including in non-traditional settings (e.g., HIV clinics providing behavioral health care).	Routine or emergency healthcare in any setting that provides human or social services, including in non-traditional settings (e.g., school-based clinics providing behavioral health care).

Dated: October 27, 2023.

Marquita Cullom,

Associate Director.

[FR Doc. 2023-24214 Filed 11-1-23; 8:45 am]

BILLING CODE 4160-90-P

DEPARTMENT OF HEALTH AND HUMAN SERVICES

Centers for Disease Control and Prevention

Notice of Closed Meeting

Pursuant to 5 U.S.C. 1009(d), notice is hereby given of the following meeting.

The meeting will be closed to the public in accordance with the provisions set forth in sections 552b(c)(4) and 552b(c)(6), title 5 U.S.C., as amended, and the Determination of the Director, Office of Strategic Business Initiatives, Office of the Chief Operating Officer, CDC, pursuant to Public Law 92-463. The grant applications and the discussions could disclose confidential trade secrets or commercial property such as patentable material, and personal information concerning individuals associated with the grant applications, the disclosure of which

would constitute a clearly unwarranted invasion of personal privacy.

Name of Committee: Disease, Disability, and Injury Prevention and Control Special Emphasis Panel (SEP)-TS24-010, Identify and Evaluate Potential Risk Factors for Amyotrophic Lateral Sclerosis (ALS).

Date: April 9, 2024.

Time: 8:30 a.m.-5 p.m., EDT.

Place: Videoconference.

Agenda: To review and evaluate grant applications.

For Further Information Contact: Mikel L. Walters, Ph.D., Scientific Review Officer, National Center for