BACKGROUND

The United States (U.S.) has the highest maternal mortality rate among high-income countries;\(^1,2\) moreover, significant racial and ethnic disparities exist in maternal mortality.\(^3\) MMRC data has shown that 60% of these deaths are preventable.\(^1\) Maternal mortality review committees (MMRCs) are in a unique position to collect and analyze information on maternal deaths to identify opportunities for prevention.\(^4,5\) MMRCs are intended to be multidisciplinary committees that conduct comprehensive reviews of pregnancy-related deaths to prevent future deaths.\(^4,6\) However, MMRCs have struggled to identify, engage, and meaningfully include diverse community members who can speak to the essential context, including the strengths and needs of the communities they represent.

In 2021, the Black Mamas Matter Alliance (BMMA) conducted an environmental scan of nine state MMRCs in the U.S. to gather information about the opportunities and challenges MMRCs face as they work to broaden committee membership and community engagement. The purpose of this scan was to capture the challenges and opportunities as described by community members who have engaged, or attempted to engage, with MMRCs in an effort to help strengthen the capacity of public health leaders, including those administering and serving on MMRCs, and to better integrate strategies toward equitable practices across MMRC processes.

MMRCs are in a unique position to collect and analyze information on maternal deaths to identify opportunities for prevention.

PARTICIPANTS OF ENVIRONMENTAL SCAN DISCUSSIONS AND CONVERSATIONS

**BMMA KINDRED PARTNERS AND COLLABORATORS** \(N=11\)

Black women-led organizations and individuals who are guided by birth justice, reproductive justice, and human rights

**COMMUNITY REPRESENTATIVES** \(N=11\)

Members of state MMRCs who represent local community constituents

**OTHER MMRC MEMBERS** \(N=8\)

Chairs, coordinators, and abstractors of select state MMRCs
More information on the project and findings have been reported elsewhere, but this issue brief highlights one of three major themes that arose from these conversations: the challenges experienced due to lack of transparency, inconsistent data provision, and legislative barriers as inhibitors to community involvement. The brief centers the experiences of KP/Cs and community representatives (henceforth referred to as “participants”) in reporting the findings. We also share potential opportunities for improvement that CDC and MMRCs can take to better integrate and value the expertise of community members in the MMRC process.

KEY FINDINGS: LACK OF TRANSPARENCY, INCONSISTENT DATA PROVISION, AND LEGISLATIVE BARRIERS AS INHIBITORS TO COMMUNITY INVOLVEMENT

Participants discussed the barriers they faced in attempting to join MMRCs, which included lack of transparency in the recruitment process and legislative barriers around compensation, background checks, and the number of seats available for community members on MMRCs. In addition, participants described the challenges of helping their communities due to inconsistent and/or delayed reporting of MMRC findings and their lack of involvement in the creation of MMRC recommendations.

CHALLENGES AND STRATEGIES ENGAGING WITH MMRCs AS IDENTIFIED BY COMMUNITY MEMBERS

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<th>CHALLENGES</th>
<th>STRATEGIES</th>
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<td>Participation in MMRCs can be detrimental to communities:</td>
<td>→ Ensure proportional representation and BIPOC leadership</td>
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<td>→ Culture and environment are unwelcoming and harmful</td>
<td>→ Eliminate burden of proof mentality for evaluating racism and discrimination</td>
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<td>→ Harmful and disrespectful conversations about patients</td>
<td>→ Integrate family interviews</td>
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<td>→ Community representatives carry burden of teaching about racism and discrimination</td>
<td>→ Improve orientation of MMRCs</td>
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<td>→ Centering of provider narrative, not the deceased birthing person</td>
<td>→ Institute anti-racism trainings</td>
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<td>→ Provide training and guidance on trigger tools</td>
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The following **CHALLENGES** were identified by participants:

1. **Lack of knowledge, transparency, and diversity in the MMRC recruitment process**

   Participants described a lack of knowledge and transparency on how decisions were made around who was invited or allowed to join MMRCs. Participants said that information on MMRC applications (if they even exist), nominations, and selection processes was not readily available, accessible, or openly communicated to the public. One participant exemplified this sentiment by stating that MMRCs were “so secretive that it’s prohibitive,” which ultimately prevented new membership or diversity on the committee. Knowledge about and nominations for MMRCs were thought to only be reserved for a select few, which were usually colleagues of clinical members on a committee. Participants expressed that MMRCs were mostly white and clinical and that more Black, Indigenous, People of Color (BIPOC), community representatives, and birth workers who use holistic birthing practices were needed to ensure a diverse committee that is more representative of the communities it aims to serve.

2. **Lack of timely and consistent access to data and communication of recommendations**

   Participants expressed that communities need access to and awareness of timely MMRC data for their own advocacy and organizing efforts. They stated that having significant multi-year data, reporting, and publication delays hindered community members from optimally serving their communities. Moreover, the lack of community representation on MMRCs also inhibited the development and inclusion of recommendations for prevention that were responsive to community needs.

3. **Legislative hurdles and lack of structural support like compensation impede community participation**

   Participants discussed state laws that prevented or limited MMRCs from adding community members to the committee. Legislation often only allowed one or two community members to join the MMRC and even then, community members faced barriers to joining such as lengthy background checks that required credit reports, work history, and other vetting strategies. Participants stated the process was “overwrought and created a lot of bureaucratic barriers,” thus further prohibiting community inclusion. Lack of compensation and other financial support to participate in all day MMRC meetings was another challenge, which most other members who were salaried, clinical or health department MMRC members did not face.

The following **OPPORTUNITIES FOR IMPROVEMENT** were identified:

1. **Institute open calls for recruitment**

   Participants recommended that all MMRCs should implement open calls for applications to provide equal access to membership and increase transparency of the recruitment process. In addition, community groups should be consulted for input on the application process.
Participants presented a variety of challenges and potential opportunities for MMRCs to integrate equitable practices throughout their processes. They maintained that all conversations regarding maternal deaths, policy solutions, and improvements needed to address the systems contributing to maternal mortality should start and end with communities. Community members have the expertise and ability to bring new solutions to MMRCs. However, this requires incorporating BIPOC leadership and proportional representation* onto MMRCs to shift power to community members, and to move beyond token efforts to diversify membership.

**Compensate community members**
Participants advocated for MMRCs to work with their local legislative bodies to secure compensation for community members. This was essential for community members who often were not compensated by their jobs while serving on the committee like some of their health department, physician, and clinical staff counterparts. Compensation can be an important strategy to increase equity and bolster community engagement.

**Increase accountability, frequency, and transparency of communications**
Participants stated that MMRCs should prioritize the provision of timely, consistent, and clear data to lay audiences. This could be done by working with local communities and anti-racism experts to develop and disseminate a communications plan that includes press releases, public campaigns, and infographics that multiple groups can understand and use. These efforts could be supported through the provision of additional resources and funding for MMRCs to have increased capacity to review cases and dedicated anti-racist experts who help MMRCs disseminate information in a timely manner. In addition, participants discussed the need for oversight and accountability around data releases.

**Involve community in the creation of recommendations**
Communities need to be involved in the creation of MMRC recommendations. While MMRCs are charged with developing recommendations to prevent maternal mortality, community members were often expected to implement the recommendations despite not being included in the discussion or development of those recommendations. As one participant said, “when community members’ voices weren’t truly heard, the recommendations still tended to come back to the status quo.” Overall, participants identified the need for collaboration between communities and MMRCs to develop and create a plan of accountability to determine who is responsible for implementing the recommendations.

**CONCLUSION**
Participants presented a variety of challenges and potential opportunities for MMRCs to integrate equitable practices throughout their processes. They maintained that all conversations regarding maternal deaths, policy solutions, and improvements needed to address the systems contributing to maternal mortality should start and end with communities. Community members have the expertise and ability to bring new solutions to MMRCs. However, this requires incorporating BIPOC leadership and proportional representation* onto MMRCs to shift power to community members, and to move beyond token efforts to diversify membership.

Integrating equity into MMRCs requires systemic change and that can only be achieved when the communities who are most impacted are meaningfully engaged throughout the data collection, review, recommendations, and implementation phases. State and local MMRCs need funding, training, and support to accomplish this, and CDC has the opportunity to provide these supports to MMRCs in their pursuit to create a more equitable review process. Below are brief recommendations that participants identified as useful starting points for both CDC and MMRCs to consider in addressing the identified challenges including lack of transparency, inconsistent data provision, and legislative barriers as inhibitors to community involvement on MMRCs.

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* The intent of this recommendation is to broadly increase community representation on an MMRC by encouraging MMRCs to shift its representation and leadership towards the Black and Brown people who disproportionately represent maternal deaths nationally. This recommendation is not intended to be applied as a formula to fill a quota on an MMRC.
RECOMMENDATIONS

Listen to and center the experiences of community members

Diversify membership and meaningfully engage communities

Provide training, guidance, and resources to strengthen the capacity of MMRCs

Provide additional funding to MMRCs

Increase transparency of MMRC processes and data

REFERENCES


SUGGESTED CITATION:


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