How to use heat-stable carbetocin and tranexamic acid for the prevention and treatment of postpartum haemorrhage in low-resource settings

Nguyen Toan Tran, Catrin Schulte-Hillen, Sarah Bar-Zeev, Agnes Chidanyika, Willibald Zeck

BACKGROUND

The least developed countries, which include those affected by fragility and humanitarian crises, account for 44% of all maternal deaths globally. Postpartum haemorrhage (PPH) is a leading cause of maternal mortality in these low-resource settings. Because uterine atony accounts for approximately two-thirds of PPH cases, WHO recommends that every woman receives a prophylactic uterotonic immediately after birth to prevent PPH as part of the active management of the third stage of labour.

Some PPH prevention and treatment medicines are well evidenced with a long implementation history, including oxytocin, misoprostol and ergometrine. Heat-stable carbetocin (HSC), a uterotonic recommended for PPH prevention and tranexamic acid (TXA), an antifibrinolytic recommended for PPH treatment, were recently added to the core list of reproductive health medicines in the 2019 Model List of Essential Medicines by the WHO. Unlike heat-sensitive oxytocin or ergometrine, HSC and TXA have the operational advantage of overcoming the logistic costs and challenges inherent to ensuring a cold chain system. Therefore, they could play a critical role in resource-challenged and warm climate settings, where cold chain transport and storage is often not available, which compromises the quality of oxytocin.

The growing number of medications now available in the PPH prevention and treatment toolkit can make it difficult for policymakers, programme managers and clinicians operating in resource-constrained settings to decide where and how to invest limited resources to achieve the best possible maternal health outcomes.

WHAT, WHERE, AND HOW

First, the critical features of HSC and TXA should be understood in light of other established PPH medications. Established medications include oxytocin (a first-line and highly effective injectable uterotonic to prevent and treat PPH in all births), misoprostol (a non-injectable uterotonic for PPH prevention and treatment when there are no trained

Summary box

⇒ Heat-stable carbetocin, a uterotonic used for postpartum haemorrhage (PPH) prevention and tranexamic acid, an antifibrinolytic indicated for PPH treatment, are recently recommended medications.
⇒ The growing number of medications in the PPH prevention and treatment toolkit can challenge policymakers, programme managers and clinicians operating in resource-constrained settings in deciding where and how to invest limited resources to achieve the best possible maternal health outcomes.
⇒ This paper argues that there is no one-size-fits-all approach to implementing international PPH prevention and treatment guidance.
⇒ A programmatic strategy tailored to the different levels of maternity care and the availability of skilled providers and cold chain systems is proposed.
providers to give injectable uterotonic or oxytocin is not available or of questionable quality) and ergometrine (an injectable uterotonic for PPH prevention and treatment). Table 1 synthesises the main features of these medications. Notably, some uterotonic, for example, oxytocin and misoprostol, have multiple obstetric and gynaecological applications, such as labour induction and augmentation as well as abortion and postabortion care. In contrast, HSC and TXA have currently a single obstetric application.

HSC is an injectable uterotonic recommended only for PPH prevention. WHO recommends HSC in situations when (1) oxytocin is unavailable or of dubious quality, (2) there is no cold transportation and storage capability, (3) its cost is comparable to that of other effective uterotonic and (4) there is skilled health personnel to inject it. TXA is an antifibrinolytic administered intravenously. It is not a uterotonic—therefore, not a uterotonic substitute. TXA is recommended only for PPH treatment in complement with uterotonic as part of the standard PPH treatment package. TXA decreases mortality from bleeding in women with PPH, irrespective of the aetiology, be it uterine atony, trauma to the genital tract, retained tissue or clotting disorder.

Second, prioritising the different medications should align with WHO guidance. For PPH prevention, the following uterotonic hierarchy is recommended: (1) in settings where multiple uterotonic are available, oxytocin (10 IU, intramuscular/intravenous) is the recommended uterotonic in all births, (2) in settings where oxytocin is unavailable (or its quality cannot be guaranteed), the use of other injectable uterotonic (HSC, or, if appropriate, ergometrine/methylergometrine or oxytocin-ergometrine fixed-dose combination) or oral misoprostol is recommended and (3) in settings where skilled health personnel are not present to administer injectable uterotonic, the administration of misoprostol (400 μg or 600 μg orally) by community healthcare workers and lay health workers

### Table 1: Summary of clinical indications and health system requirements of uterotonic and tranexamic acid

<table>
<thead>
<tr>
<th>Use and health system requirements</th>
<th>Uterotonics</th>
<th>Non-uterotonic</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>PPH prevention</strong></td>
<td>Recommended</td>
<td>Not recommended</td>
</tr>
<tr>
<td><strong>PPH treatment</strong></td>
<td>Recommended</td>
<td>Not recommended</td>
</tr>
<tr>
<td><strong>Labour induction</strong></td>
<td>Recommended</td>
<td>Not recommended</td>
</tr>
<tr>
<td><strong>Labour augmentation</strong></td>
<td>Recommended</td>
<td>Not recommended</td>
</tr>
<tr>
<td><strong>Abortion care</strong></td>
<td>Not recommended</td>
<td>Not recommended</td>
</tr>
<tr>
<td><strong>Postabortion care</strong></td>
<td>Not recommended</td>
<td>Not recommended</td>
</tr>
</tbody>
</table>

| Administration route | Intravenous, IM | Oral, sublingual, vaginal

| Skilled health provider requirement | Yes | No |
| Cold chain transport and storage requirement | Yes (2°C–8°C) | No (but ≤25°C) |
| Presentation** | 10 IU ampoule | 200 mcg tablet** |
| Price per unit** | US$0.334 per ampoule | US$0.054 per oral tablet of 200 mcg |

The colour shadings reflecting a traffic light approach (red, orange and green) were used to give the reader an instant recognition of the level of caution that is needed in regards to the use of the uterotonic and non-uterotonic in different clinical situations. The use of lighter orange and grey in rows 10 and below of table 1 were to help the reader easily identify which uterotonic and non uterotonic required a skilled health provider for their administration and which required a cold chain transport and storage requirement.

**Heat-stable carbetocin: only in contexts where its cost is comparable to that of other effective uterotonic.
†Ergometrine: only in contexts where hypertensive disorders can be safely ruled out before use. Ergometrine refers to ergometrine/methylergometrine.
‡First line.
§Alone or preceded by mifepristone.
**Per UNFPA Product Catalogue (2021).
††The vaginal route is not recommended for PPH prevention and treatment.
¶The vaginal route is not recommended for PPH prevention and treatment.
IM, intramuscular; IU, international units; PPH, postpartum haemorrhage; UNFPA, Product Catalogue of the United Nations Population Fund.
is recommended. As for PPH treatment, (1) intravenous oxytocin is the recommended uterotonic; (2) the early use of intravenous TXA within 3 hours of birth in addition to standard care is recommended in all cases of PPH, regardless of whether the bleeding is due to genital tract trauma or other causes and (3) if intravenous oxytocin is unavailable, or if the bleeding does not respond to oxytocin, the use of intravenous ergometrine, oxytocin-ergometrine fixed-dose, or a prostaglandin drug (including misoprostol) is recommended (ergometrine and oxytocin-ergometrine are not recommended in case of hypertensive disorder).

Third, these recommendations must, in practice, account for various health system requirements, as summarised in table 1. To help programme managers operationalise the information in table 1, the different health system requirements can be streamlined by focusing on the availability of skilled providers and cold chain transport and storage of 2°C–8°C, two critical constraints often encountered in resource-challenged settings (figure 1).

Additionally, figure 1 offers examples of implementation settings, such as comprehensive obstetric care facilities with skilled providers and reliable cold storage (yellow box). Basic emergency obstetric care facilities with skilled providers are subdivided according to the availability (yellow box) or non-availability (orange box) of a consistent electric power source to ensure cold storage. Community-based programmes with the distribution of misoprostol for home births are captured in the grey box.

Fourth, the reflection on whether HSC and TXA should be integrated into PPH prevention and treatment strategies offers an opportunity to examine the health system and health service gaps that prevent optimal maternal health outcomes. There is, however, no single solution that matches the resources and needs of all the different settings. Therefore, policymakers and programme managers should consider a stepwise programmatic approach to averting maternal deaths. First, existing emergency obstetric care services should undertake a continuous quality improvement process, notably in terms of staff competencies, facility materials and supplies, functional referral mechanisms and performance and accountability. Health facilities should have a working supply chain to avail uterotonics for PPH prevention round-the-clock in addition to essential equipment, medications and up-to-date protocols and job aids for emergency obstetric treatment if PPH or other complications occur. As a complement to uterotonics for PPH treatment, the introduction of TXA for all PPH cases should be considered at this step. This aligns with the WHO guidance and the recommendations of the International Federation of Gynecologists and Obstetricians and the International Confederation of Midwives. Second, the integration of HSC—and TXA if not done in the previous step—into the

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**Figure 1** Practical considerations for procuring and using medicines for PPH prevention and treatment with examples of implementation settings. In blue: 2°C–8°C cold chain transport and storage required (oxytocin and ergometrine). *Heat-stable carbetocin: only in contexts where its cost is comparable to that of other effective uterotonics. §Ergometrine: only in contexts where hypertensive disorders can be safely ruled out before use. PPH, postpartum haemorrhage. ¶For PPH treatment in contexts where there are competent health providers, it is imperative to procure other essential supplies according to the level of care, including crystalloid for fluid replacement and oxygen.
If the decision is made to include heat-stable carbetocin and/or tranexamic acid into the national PPH prevention and treatment package, the following actions should be considered:

**National policy and planning readiness**
- Update existing national PPH prevention and treatment guidelines, including PPH algorithms integrating heat-stable carbetocin and/or tranexamic acid into the standard PPH prevention and treatment package.
- Define where (levels of care) heat-stable carbetocin and/or tranexamic acid may be positioned and who (health cadres) can administer it.
- Establish a plan for addressing the financial and program needs in accordance with revised national guidelines.
- Assess the status of procurement, distribution, and storage of existing PPH supplies, heat-stable carbetocin, and/or tranexamic acid.
- Review and update national pre-service education curricula for PPH prevention and treatment in accordance with revised national guidelines, focusing on competency-based training.
- Develop updated provider job aids and decision support tools to help providers acquire necessary skills to integrate heat-stable carbetocin and/or tranexamic acid into routine PPH prevention and treatment (e.g., pre- and in-service education/training, onsite job aids, and clinical algorithms).
- Incorporate heat-stable carbetocin and/or tranexamic acid into the national health management information system indicators relevant for PPH surveillance, prevention, and treatment, prioritizing indicators pertinent for quality improvement action (e.g., stock-outs).
- Incorporate heat-stable carbetocin and/or tranexamic acid into patient records for childbirth and postpartum care (e.g., time and dose of PPH prevention measures, diagnostic time and cause of PPH, time and dose of PPH treatments, PPH outcome).

**Health facility readiness**
- Ensure provider competencies and positive behavior change by enlisting respected clinical champions and leaders to:
  1. Offer competency-based in-service refresher training and follow-up supportive supervision on PPH prevention and treatment, inclusive of heat-stable carbetocin and/or tranexamic acid.
  2. Address barriers and enablers to provider behavior change, including potential concerns and areas of resistance.
- Make copies of the updated PPH prevention and treatment protocols, job aids, decision support tools, and standardized patient charts immediately accessible in labor, delivery, and postpartum areas.
- Support a functioning supply chain to ensure 24/7 availability of heat-stable carbetocin and/or tranexamic acid and other essential emergency obstetric care supplies in labor, delivery, and postpartum areas.

**Figure 2** Policy and programme considerations for establishing an enabling environment to introduce heat-stable carbetocin or tranexamic acid, or both. PPH, postpartum haemorrhage.

Health system should be carefully considered at the appropriate level of care. **Figure 2** outlines key actions to establishing an enabling policy and programme environment for the introduction of HSC or TXA, or both. Finally, it is important to recognise that contraception remains the most cost-effective intervention to avert maternal deaths globally by preventing unintended pregnancies and allowing healthy timing and spacing of pregnancies. Therefore, in conjunction with the previous two steps, health systems should invest in generating demand for family planning and optimising the quality of contraceptive programmes and services (e.g., commodities, providers’ clinical skills, provider-client decision-making tools). This includes the first year postpartum when the unmet need for family planning is particularly high.
CONCLUSIONS

HSC and TXA are recently recommended medications for inclusion in the PPH prevention and treatment toolbox and could potentially play a critical role in decreasing maternal deaths in low-resource settings. Securing their availability in-country should be embedded in a thorough understanding of their clinical indications and the enabling health system environment. Considering whether HSC and TXA should be included in maternal health services provides a new opportunity to engage key national and local stakeholders, including health professionals, to look at the overall health system challenges and opportunities that hinder or support the reduction of maternal mortality and morbidity in general, and that due to PPH specifically.

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