Implementing Malaria in Pregnancy Programs in the Context of World Health Organization Recommendations on Antenatal Care for a Positive Pregnancy Experience

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Background

MiP is a major public health problem with substantial risks for mothers and their babies. Each year, MiP is responsible for 20% of stillbirths in sub-Saharan Africa, 11% of all newborn deaths in sub-Saharan Africa, and 10,000 maternal deaths globally. WHO recommends a package of interventions for controlling malaria and its effects during pregnancy. In areas where malaria is a risk, WHO recommends delivery and use of insecticide-treated nets (ITNs) and effective management of cases by providing prompt quality diagnosis and effective treatment of malaria infections. In areas with moderate to high transmission of Plasmodium falciparum, WHO additionally recommends the administration of intermittent preventive treatment during pregnancy using sulfadoxine-pyrimethamine (IPTp-SP) that is quality assured. SP is the only drug currently recommended for administration in the context of IPTp, and it is

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important to note that SP continues to show benefit for both the mother and her baby, even in areas of SP resistance. Further, a recent study by Chico et al. found women who received two or more doses of IPTp-SP were protected not only from adverse outcomes related to malaria, but also from some sexually transmitted infections/reproductive tract infections.

The delivery of high-quality antenatal care (ANC) is essential for successful MiP programming. WHO’s Recommendations on Antenatal Care for a Positive Pregnancy Experience now promote a minimum of eight contacts between pregnant women and the health system versus the previously recommended four ANC visits. This new WHO ANC model highlights that a woman’s contact with her provider should be more than a simple visit. It should be an opportunity for comprehensive, high-quality care, including medical care, support, and the provision of timely and relevant information throughout pregnancy. Depending on the context of the country, the definition of contact may include scheduled ANC visits and information sessions for pregnant women with relevant caretakers at the household, community, and health facility levels. These increased opportunities to support women during their pregnancies are an incentive for countries to deliver comprehensive care, including MiP interventions, to pregnant women.

Considerations for the Implementation of MiP Programming

Timing of IPTp-SP

The new ANC recommendations need to be adapted to each country’s context. Complementing the use of an ITN, and prompt and effective case management, the ANC contact schedule for MiP should be applied flexibly so that pregnant women always receive IPTp-SP when eligible, starting as early as possible during the second trimester of pregnancy. Table 1 highlights the WHO ANC recommended schedule and corresponding MiP interventions.

It is important to keep in mind that:

- Determining gestational age by clinical examination, especially early in pregnancy, can be challenging. WHO recommends that countries continue to use what is currently practiced for dating—either abdominal palpation or symphysis-fundal height. Doing one ultrasound scan, ideally during the first trimester, where available, is another opportunity to determine early gestational age, among other potential benefits for the pregnancy.

- The period between 13 and 20 weeks is critical for irreversible negative consequences of MiP, when parasite densities are highest, and major benefit can be achieved from malaria prevention. For effective MiP programming, contact with a health provider early in the second trimester (between 13 and 16 weeks) is critical to ensuring timely access to the first dose of IPTp-SP for maximal impact.

- WHO’s Recommendations on Antenatal Care for a Positive Pregnancy Experience and Optimizing Health Worker Roles for Maternal and Newborn Health promote task shifting of components of ANC, including the provision of IPTp, from staff in health facilities to a broad range of cadres, including auxiliary nurses, nurses, midwives, and doctors. As countries consider the application of the new WHO ANC recommendations and acceleration of MiP programming, delivery approaches at the community level that complement ANC offer promise to achieve increased coverage during the antenatal period.

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including malaria prevention. Countries could consider piloting and scaling up community approaches that help to increase IPTp uptake and ANC coverage.

### Table 1: 2016 ANC contact schedule with timelines for implementation of malaria in pregnancy interventions

<table>
<thead>
<tr>
<th>ANC Contact Schedule and Proposed Time of IPTp-SP Administration (To be adapted to country context, also considering disease burden and health needs)</th>
<th>MiP-related Interventions and Considerations during ANC Contacts</th>
</tr>
</thead>
<tbody>
<tr>
<td>Contact 1: Up to 12 weeks</td>
<td>• Register pregnant women, provide ITNs, and counsel on their use. Screen for HIV.</td>
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<tr>
<td></td>
<td>• Administer 30 to 60 mg of elemental iron and 400 μg (0.4 mg) of folic acid.</td>
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<tr>
<td></td>
<td>• Counsel to return for a visit at 13 to 16 weeks (see contact 1a below) to receive the first dose of IPTp-SP (as directed by national guidelines).*</td>
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<tr>
<td></td>
<td>• Counsel on prompt diagnosis and effective treatment/malaria case management during pregnancy.</td>
</tr>
<tr>
<td>Additional contact (1a): In moderate to high malaria transmission areas in Africa where IPTp-SP is policy, a contact should be made early in the second trimester (13 to 16 weeks) to administer SP as early as possible.</td>
<td>IPTp-SP dose 1</td>
</tr>
<tr>
<td>Contact 1a: Up to 12 weeks</td>
<td>Remember:</td>
</tr>
<tr>
<td></td>
<td>• Do not administer IPTp-SP before week 13 of pregnancy.</td>
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<tr>
<td></td>
<td>• Administer the first IPTp-SP dose as early as possible in the second trimester to fully benefit from the protective capacity in this critical period of pregnancy.†</td>
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<tr>
<td></td>
<td>• Administer the second dose of IPTp-SP one month later.</td>
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<tr>
<td></td>
<td>• Administer the following doses of IPTp-SP starting from the scheduled contact at 20 weeks, observing at least one-month intervals between SP doses.</td>
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<td></td>
<td>• SP can be safely administered from the beginning of the second trimester until the time of delivery.</td>
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<td></td>
<td>• One full dose of IPTp-SP consists of 1,500 mg/75 mg SP (i.e., three tablets of 500 mg/25 mg SP).</td>
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<td>• Provide IPTp-SP by directly observed treatment.</td>
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<td></td>
<td>• Pregnant women on co-trimoxazole should not receive IPTp-SP due to an increased risk of adverse events when both drugs are given in parallel.</td>
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<tr>
<td></td>
<td>• Continue to administer 30 to 60 mg of elemental iron and 400 mcg (0.4 mg) of folic acid.</td>
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<tr>
<td></td>
<td>• Continue counseling as above.</td>
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<tr>
<td>Contact 2: 20 weeks</td>
<td>IPTp-SP dose 2</td>
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<tr>
<td>Contact 3: 26 weeks</td>
<td>IPTp-SP dose 3</td>
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<tr>
<td>Contact 4: 30 weeks</td>
<td>IPTp-SP dose 4</td>
</tr>
<tr>
<td>Contact 5: 34 weeks</td>
<td>IPTp-SP dose 5</td>
</tr>
<tr>
<td>Contact 6: 36 weeks</td>
<td>No SP administration if last dose was received at contact 5 in week 34</td>
</tr>
<tr>
<td>Contact 7: 38 weeks</td>
<td>IPTp-SP dose 6 (if no dose was received at contact 6 in week 36)</td>
</tr>
<tr>
<td>Contact 8: 40 weeks</td>
<td>No SP administration if last dose was received at contact 5 in week 34</td>
</tr>
</tbody>
</table>

Pregnant women should receive MiP interventions as appropriate, even when they come at weeks not designated in the contact schedule.

Despite the known side effects associated with sulfonamides, SP for intermittent preventive treatment in pregnancy is generally very well tolerated. Mild and transient side effects including nausea, vomiting, weakness and dizziness have been reported by some women, particularly with the first dose of SP. Studies have demonstrated that side effects tend to decrease with the administration of further doses (§,‡). Side effects should be discussed openly and managed in the ANC.

* The first dose of IPTp-SP is recommended to be given as early as possible in the second trimester of pregnancy to ensure optimal protection from malaria for the mother and her baby. However, pregnant women who come later in pregnancy can and should receive their first dose anytime (as long as it is not in the first trimester), with following doses being given at least one month apart. When malaria-endemic countries are planning their ANC programming, they may wish to add another contact to allow for monthly dosing of IPTp-SP.

† Pregnant women should receive their first dose of IPTp-SP as early as possible at the beginning of the second trimester, defined as 13 weeks gestation (i.e., 12 completed weeks or 13 weeks and zero days).


Frequency of IPTp-SP
Following administration of the first dose of IPTp as early as possible in the second trimester (i.e., 13 to 16 weeks), pregnant women should receive an additional dose of IPTp-SP at each contact with a health care worker trained to deliver IPTp-SP until the time of delivery, ensuring that doses of IPTp-SP are administered at least one month apart. WHO does not recommend a maximum number of doses of IPTp-SP. SP can be safely administered from the beginning of the second trimester until delivery.

Sourcing of quality-assured SP
The availability of quality-assured SP for IPTp is critical to ensure pregnant women have optimal protection from malaria, in addition to using an ITN and accessing effective case management. Countries should procure the drug from manufacturers who produce quality-assured SP (see checklist below) and ensure supporting partners are doing the same.

ITN use
All pregnant women should sleep under an ITN as early as possible in pregnancy, though ideally before becoming pregnant. Providing an ITN at the first contact will help to keep the pregnant woman and her fetus safe from malaria. Additionally, all efforts should be made to ensure women of reproductive age have access to and sleep under an ITN so that they are protected against malaria if they become pregnant.

Key points regarding ITN use include:

- Free delivery of an ITN at the first ANC visit is an incentive to attend antenatal care and provides the pregnant woman with a lifesaving tool for herself and her baby. Sleeping under the ITN will also protect her baby during the first year of life.

- Countries need to plan and budget for continuous ITN distribution to pregnant women at the first ANC contact, in addition to forecasting, procuring, and distributing ITNs for campaigns targeting the whole population.

Effective case management
Pregnant women with signs and symptoms of malaria need immediate access to quality diagnosis and effective treatment. Health care providers must be able to consistently assess all women of reproductive age for pregnancy, and test and treat these women for malaria in accordance with national and WHO guidelines.13

As malaria prevalence in a country declines, the clinical manifestations of malaria infection in pregnant women become more severe due to reduced immunity. Having strong public and private health systems in place to rapidly detect and treat MiP becomes increasingly important as malaria transmission levels fall.

Women living with HIV
Women living with HIV are at increased risk of all adverse consequences of malaria infections due to their compromised immune responses. All pregnant women should be screened for HIV at first ANC contact. Pregnant women living with HIV and taking co-trimoxazole prophylaxis should not receive SP, as concomitant administration of SP and co-trimoxazole could increase adverse drug reactions. When taken daily, co-trimoxazole provides protection against MiP. Despite this, it is especially important that pregnant women living with HIV sleep under an ITN, and seek prompt diagnosis and receive effective treatment if they experience symptoms of malaria.

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Iron and folic acid supplementation

Since iron and folic acid requirements increase in pregnancy, WHO recommends supplementation with 30 to 60 mg of elemental iron and 400 mcg (0.4 mg) of folic acid for pregnant women to prevent maternal anemia, puerperal sepsis, low birthweight, and preterm birth.14

To improve maternal and newborn outcomes, intermittent oral iron and folic acid supplementation with 120 mg of elemental iron and 2,800 mcg (2.8 mg) of folic acid once weekly is recommended for pregnant women who cannot take daily iron supplements due to side effects and for populations in which less than 20% of pregnant women have anemia.

Every effort should be made to ensure that low-dose folic acid (i.e., 0.4 mg, equivalent to 400 mcg) is available and provided as part of routine antenatal care. High doses of 5 mg of folic acid and greater counteract the antimalarial efficacy of SP and should not be given along with SP. In areas where only high-dose folic acid is available, there is presently no scientific consensus on how long high doses of folic acid should be withheld following the dose of SP. Many countries suggest withholding high doses of folic acid (5 mg or more) for two weeks after administration of SP, but this may shorten the duration of efficacy of SP. Countries should advocate for procurement of low-dose folic acid, which does not interfere with the efficacy of SP. In cases where high-dose folic acid is resumed two weeks following SP dosing, the health care provider should strongly advise the pregnant woman to use her ITN, and seek care immediately for proper diagnosis and treatment if signs and symptoms of malaria are present.15

KEY MESSAGES

WHO ANC Contacts

1. Each contact between a pregnant woman and the corresponding provider/caretaker should be an opportunity for high-quality care, including medical care, support, and the provision of timely and relevant information throughout pregnancy.

2. Depending on the context of the country, the definition of contact may include scheduled ANC visits and information sessions for pregnant women with relevant caretakers at the household, community, and health facility levels.

Malaria in Pregnancy

1. All pregnant women living in areas at risk for malaria transmission should:
   - Sleep under an ITN.
   - Seek prompt quality diagnosis when signs and symptoms of malaria are present, and receive effective malaria case management with an appropriate drug at the correct dose.

2. Pregnant women living in moderate to high malaria transmission areas in Africa should also receive:
   - IPTp-SP under directly observed therapy (DOT), starting as early as possible in the second trimester, with doses given at least one month apart until the time of delivery.
   - To enable pregnant women in endemic areas to start IPTp-SP at the beginning of the second trimester, policymakers should put in place supportive policies to ensure that women have an ANC contact at 13 weeks gestation. See Table 1.
     - IPTp-SP should be given to a pregnant woman at every ANC contact starting from 13 to 16 weeks, with each dose being given at least one month (four weeks) apart.
     - Pregnant women who have an ANC contact twice between 13 and 20 weeks, at least one month apart, should receive IPTp-SP by DOT at both contacts.
     - If a woman comes for her first second-trimester contact anytime between 13 and 20 weeks, she should receive IPTp-SP, and at every following contact, with doses one month apart.
     - Pregnant women can receive IPTp-SP safely starting as early as possible in their second trimester up until the end of pregnancy.
   - SP should not be administered to women living with HIV who are receiving co-trimoxazole.

3. Countries should only provide quality-assured SP for IPTp to ensure effective care for pregnant women.
   - Current procurement sources of quality-assured SP can be found on The Global Fund List of Pharmaceutical Products compliant with the quality assurance policy, accessible via: https://www.theglobalfund.org/media/4756/psm_productsmalaria_list_en.pdf.

4. Iron and folic acid requirements increase during pregnancy:
   - Administer 30 to 60 mg of elemental iron and 400 mcg (0.4 mg) of folic acid.

WHO Department of Maternal, Newborn, Child and Adolescent Health
http://www.who.int/maternal_child_adolescent

WHO Department of Reproductive Health and Research
http://www.who.int/reproductivehealth

WHO Global Malaria Programme
http://www.who.int/malaria

WHO Department of Nutrition for Health and Development
http://www.who.int/nutrition/en/