Indicator: # mothers with infants under 12 months who received intermittent preventive treatment (IPTp) according to national policy during their last pregnancy. (HE_MNCH_134)

Health Sector: Maternal, Newborn, and Child Health

Project Area: Maternal, Newborn, and Child Health

Type: Outcome

Unit of Measure: Individual

Disaggregation: None

Related Objective: Increase knowledge and skills of women to adopt practices that contribute to a healthy pregnancy, safe delivery, good postpartum, and a healthy newborn (Objective 1)

Precise definitions

Mothers with infants under 12 months: The mothers included in the outcome indicator should have attended a multisession training on maternal and newborn health, ideally during their pregnancy, co-facilitated by the PCV and their counterpart or local health staff/volunteer. The individual should have attended at least 75 percent of sessions to be counted in the denominator for HE_MNCH_079: Number of target population reached with individual or small group education on maternal and newborn care services, and therefore this indicator.

IPTp: is defined as intermittent preventive treatment in pregnancy (IPTp). It is also known as intermittent presumptive therapy or intermittent protective treatment in pregnancy. It involves the administration of a single curative dose of an efficacious antimalarial drug (e.g., Sulphadoxine-pyremethemine, or Fansidar) as prescribed by national policy at specific intervals at least twice during pregnancy—regardless of whether or not the woman is infected with malaria. WHO recommends that pregnant women in malarious zones receive IPTp at each antenatal clinic visit starting at quickening provided those visits are at least a month apart. This translates to as many as five doses per pregnancy, but national policies vary.

Data collection

Tool: Maternal Health Outcomes Survey

This survey is intended to be given to the pregnant women who participated in a multisession training focused on maternal and newborn health facilitated by the PCV and their counterpart. The pregnant woman should have attended at least 75 percent of the defined information sessions to be included in the survey. Data for the outcome indicators ideally would be collected three to six months after the birth of their baby. If the timing of a PCV’s close of service (COS) or other factors will not allow for this, a survey could be conducted earlier, but indicators that reference specific timeframes may need to be removed such as the one focused on exclusive breastfeeding for six months. While it is the pregnant woman/mother who is taking the survey, some of the questions are focused on the infant. The survey should be conducted by the PCV and their counterpart or co-facilitator of the group following the instructions provided in the outcomes survey tool closely. The survey should include an indicator for each area of prevention of maternal and newborn health that was covered in the training. See the Maternal Health Outcomes Survey for a complete list.
**Reporting**

To be counted for this indicator the following criteria must be met:

- The woman gave birth in the 12 months prior to being surveyed.
- During her pregnancy, the woman received training on the essential maternal care services and/or participated in a group that was facilitated by a PCV or their partner and was encouraged to attend antenatal care as a result of being in the group.
- The training was provided by the PCV or their partner in an individual or small group setting of 25 or less.
- Attendance at the educational sessions was documented by the Volunteer or their partner.