17th Invitation to Manufacturers of Antimalarial Medicines to Submit an Expression of Interest (EOI) for Product Evaluation to the WHO Prequalification Team: medicines

To support national and global efforts to increase access to and the affordability of care and treatment of malaria, WHO, together with UNICEF, UNAIDS and UNITAID, invite manufacturers of selected pharmaceutical products to submit Expressions of Interest (EOIs) for product evaluation. The first Invitation to EOI for antimalarial medicines was published in 2002.

ARTICLE 1. PROCEDURE FOR THIS INVITATION TO EOI

The current Invitation is published in accordance with the Procedure for Prequalification of Pharmaceutical Products, adopted in 2001 by the 37th WHO Expert Committee on Specifications for Pharmaceutical Preparations, and amended subsequently as part of the 45th report of the Committee, published as No. 961 of the WHO Technical Report Series in 2011.

Assessment of product(s) submitted under this Invitation will include evaluation of:

• product dossiers, which must include product data and information as specified in the Procedures & Fees section of the website of the WHO Prequalification Team: medicines (PQTm)
• manufacturing sites, which must adhere to good manufacturing practices (GMP)
• clinical sites (if applicable), which must adhere to good clinical practices (GCP).

If evaluation demonstrates that a product and its corresponding manufacturing (and clinical) site(s) meet WHO recommended standards, it will be included in the list of medicinal products that are considered to be acceptable for procurement by UN organizations and others.

ARTICLE 2. MEDICINAL PRODUCTS INCLUDED ON THE 17TH INVITATION

The ultimate aim of this 17th Invitation is to increase the range of selected products and sources available in relation to treatment for malaria. The medicines listed in the 17th Invitation have been identified by WHO Global Malaria Programme as vital to effective treatment for people living with malaria. These products are included either in the WHO Model List of Essential Medicines and/or in the WHO Guidelines for the Treatment of Malaria.

Products included in the WHO Model List of Essential Medicines are those which satisfy the priority health care needs of a population. They are selected on the basis of disease prevalence, evidence on efficacy and safety, and comparative cost-effectiveness.

Products included in WHO treatment guidelines are selected on the basis of an assessment of the quality of evidence for benefits, harms, costs, and appropriateness for use in a variety of situations, taking into account needs of special populations, and the values and preferences of the groups (professional and patient) using them.

Interested manufacturers are encouraged to submit documentation for recommended dosage forms and strengths, as specified below, of medicinal products in the following categories. The appropriate solid dosage formulations, which are scored for flexible dosing purposes, should be supported by relevant evidence on equal distribution of active ingredients in the scored products, especially in case of fixed-dose combination products.
1. Artemisinin-based fixed dose oral combination formulations
   - Artemether/Lumefantrine
     tablet 20 mg/120 mg
     tablet 40 mg/240 mg
     tablet 60 mg/360 mg
     tablet 80 mg/480 mg
   - Artesunate/Amodiaquine
     tablet 50 mg/135 mg
     tablet 100 mg/270 mg
   - Artesunate/Mefloquine
     tablet 100 mg/200 mg
   - Artesunate/Pyronaridine
     tablet 60 mg/180 mg
   - Dihydroartemisinin/Piperaquine Phosphate
     tablet 60 mg/480 mg
     tablet 80 mg/640 mg

2. Artemisinin-based fixed dose combination oral paediatric formulations, preferably dispersible
   - Artemether/Lumefantrine,
     tablet 20 mg/120 mg
   - Artesunate/Amodiaquine,
     tablet 25 mg/67.5 mg
   - Artesunate/Mefloquine,
     tablet 25 mg/50 mg
   - Artesunate/Pyronaridine
     tablet 20 mg/60 mg
   - Dihydroartemisinin/Piperaquine, phosphate,
     tablet 20 mg/160 mg
     tablet 30 mg/240 mg
     tablet 40 mg/320 mg

3. Artemisinin-based single-ingredient formulations
   - Artemether, oily injection 20 mg/ml; 40 mg/ml; 80 mg/ml; 100 mg/ml
   - Artesunate, powder for injection 30 mg; 60 mg; 120 mg (vial)
   - Artesunate, suppositories 50 mg; 100 mg; 200mg
   - Artesunate, tablet¹ 25 mg; 50 mg; 100 mg

¹ Artesunate tablets to be used only in combination with Mefloquine
4. Combination antimalarial medicines in co-blistered formulations, preferably dispersible
   - Amodiaquine+Sulfadoxine/Pyrimetamine
either
   tablet 75 mg+250 mg/12.5 mg
tablet 150 mg+500 mg/25 mg
   or
   - Amodiaquine+Sulfadoxine/Pyrimetamine
either
   tablet 76.5 mg+250 mg/12.5 mg
tablet 153 mg+500 mg/25 mg

5. Other antimalarial medicines
   - Mefloquine tablet 250 mg
   - Primaquine base 2.5 mg tablets
   - Primaquine base 3.75 mg tablets
   - Primaquine base 5 mg tablets (scored)
   - Primaquine base 7.5 mg scored tablets (scored)
   - Primaquine base 15 mg tablets (scored)
   - Sulfadoxine/Pyrimetamine tablets 250 mg/12.5 mg (scored); 500 mg/25 mg (scored)

Product presentations which support adherence to treatment and rational drug use are strongly encouraged.

ARTICLE 3. HOW TO SUBMIT AN EXPRESSION OF INTEREST

In order to submit an expression of interest for product evaluation, the manufacturer must send the required documentation, arranged according to the information provided in the Procedures & Fees section of PQTm’s website at https://extranet.who.int/prequal

ARTICLE 4. QUALITY ASSESSMENT PROCEDURE FOLLOWING SUBMISSION OF AN EXPRESSION OF INTEREST BY A MANUFACTURER

The quality assessment is undertaken to evaluate whether the pharmaceutical product being evaluated meets the requirements recommended by WHO, and is manufactured in compliance with good manufacturing practices (GMP).

The procedure established by WHO for quality assessment incorporates:
- general understanding of the production and quality control activities of the manufacturer;
- assessment of product data and information on safety, efficacy and quality submitted by the manufacturer, including product formulation, manufacture and test data and results;
- assessment of the manufacturing site's adherence to GMP, and its consistency in production and quality control of starting materials, with specific emphasis on active pharmaceutical ingredients, and finished product;
- assessment of clinical testing units or organizations (i.e. parties performing one or more clinical trials with the product) for compliance with good clinical practices and good laboratory practices, as appropriate;
- random sampling and testing of medicines supplied.
Previous evaluation conducted by the relevant national medicines regulatory authority (NMRA) may be taken into account during the evaluation conducted by WHO, provided that the NMRA has expertise in the product area. If appropriate, the relevant NMRA may be invited to collaborate with WHO on the quality assessment. Any manufacturer who submits a product for evaluation, is therefore encouraged to authorize its NMRA to discuss relevant product files with WHO representatives, during assessments and inspections, if required (subject to appropriate confidentiality provisions, if necessary).

Once WHO is satisfied that quality assessment has been completed for the manufacturer of the relevant starting materials, the finished pharmaceutical product, and the clinical testing units, and that the product meets WHO recommended standards, the product (as produced at the specified manufacturing site) is added to the WHO List of Prequalified Medicinal Products.

**ARTICLE 5. REFERENCES AND FURTHER INFORMATION**

For further information on the WHO Prequalification Team: medicines, please visit the PQTm website at: https://extranet.who.int/prequal. Should you have any questions relating to the procedure for responding to an EOI, please write to the WHO Prequalification Team - Medicines at its email address: prequal@who.int. Your question(s) will be directed to the prequalification team member who can best advise you.