

## Steps before prequalification

### I. BACKGROUND INFORMATION ON THE PROCEDURE

#### 1. Submission of the dossier

The company Guilin Pharmaceutical Co., Ltd submitted in 2019 an application for [MA154 trade name]\* (MA154) to be assessed with the aim of including [MA154 trade name] in the list of prequalified medicinal products for the treatment of malaria.

[MA154 trade name] was assessed according to the ‘*Procedure for Assessing the Acceptability, in Principle, of Pharmaceutical Products for Purchase by United Nations Agencies*’ by the team of WHO assessors. The assessors are senior experts, mainly from national authorities, invited by WHO to participate in the prequalification assessment process.

#### 2. Steps taken in the evaluation of the product

|                   |  |
|-------------------|--|
| May 2018          | The manufacturer of the APIs was inspected for compliance with WHO requirements for GMP.                                   |
| October 2018      | The manufacturer of the FPP was inspected for compliance with WHO requirements for GMP.                                    |
| March 2019        | During the meeting of the assessment team the safety and efficacy data were reviewed and further information was requested |
| April 2019        | The applicant’s response letter was received.  |
| May 2019          | The safety and efficacy data were reviewed and found to comply with the relevant WHO requirements.                         |
| May and July 2019 | During the meetings of the assessment team the quality data were reviewed and further information was requested.           |
| November 2019     | The applicant’s response letter was received.  |
| November 2019     | During the meeting of the assessment team the additional quality data were reviewed and further information was requested. |
| March 2020        | The applicant’s response letter was received.  |
| April 2020        | The additional quality data were reviewed and further information was requested.   |
| May 2020          | The applicant’s response letter was received.  |
| May 2020          | During the meeting of the assessment team the additional quality data were reviewed and further information was requested. |
| July 2020         | The applicant’s response letter was received.  |
| July 2020         | During the meeting of the assessment team the additional quality data were reviewed and further information was requested. |
| September 2020    | The applicant’s response letter was received.  |
| September 2020    | During the meeting of the assessment team the additional quality data were reviewed and further information was requested. |
| November 2020     | The applicant’s response letter was received.  |

\* Trade names are not prequalified by WHO. This is the national medicines regulatory authority’s responsibility.

|                              |   |
|------------------------------|---|
| November 2020 and March 2021 | During the meetings of the assessment team the additional quality data were reviewed and further information was requested. |
| March 2021                   | The applicant's response letter was received.   |
| March 2021                   | During the meeting of the assessment team the additional quality data were reviewed and further information was requested.  |
| March 2021                   | The applicant's response letter was received.   |
| April 2021                   | The quality data were reviewed and found to comply with the relevant WHO requirements.                                      |
| April 2021                   | Product dossier accepted (quality assurance)  |
| 13 April 2021                | [MA154 trade name] was included in the list of prequalified medicinal products.   |

## II. GENERAL CONDITIONS FOR THE PREQUALIFICATION

### 1. Manufacturer and Inspection status

#### Manufacturer of the finished product and responsible for batch release

Guilin Pharmaceutical Co., Ltd  
Oral Solid Dosage Workshop I  
No. 43, Qilidian Road, Guilin 541004  
Guangxi  
China

#### Inspection status

The sites inspected were found to be in compliance with WHO requirements for GMP.  
Not inspected for GCP/GLP since a biowaiver applies.

### 2. (Advice on) Conditions or restrictions regarding supply and use

Medicinal product subject to medical prescription.

Further information is available at:

<https://extranet.who.int/pqweb/medicines/prequalified-lists/finished-pharmaceutical-products>