WHO Prequalification Programme WHO PUBLIC ASSESSMENT REPORT (WHOPAR)

[MA169 trade name]^{*}

Amodiaquine (as hydrochloride) 150 mg dispersible tablets + Pyrimethamine/Sulfadoxine 25 mg/500 mg dispersible tablets

[MA169 trade name], manufactured at Universal Corporation Limited, Kikuyu, Kenya, was included in the WHO list of prequalified medicinal products for malaria prevention on 30 July 2023.

[MA169 trade name] is indicated for malaria prevention during the malaria season (seasonal malaria chemoprevention, SMC) in patients aged 1 year to 10 years. Detailed information on the use of this product is described in the summary of product characteristics (SmPC), which can be found in this WHOPAR.

The active pharmaceutical ingredients of [MA169 trade name] are amodiaquine (as hydrochloride), pyrimethamine and sulfadoxine.

The efficacy and safety of amodiaquine (as hydrochloride), pyrimethamine and sulfadoxine are well established based on extensive clinical experience in the malaria prevention.

For details on the uses of this product and for side effects and warnings, see Part 4 of this WHOPAR (summary of product characteristics).

On the basis of data submitted and public information on the use of amodiaquine (as hydrochloride), pyrimethamine and sulfadoxine in malaria prevention, the team of assessors advised that [MA169 trade name] is of acceptable quality, efficacy and safety to allow inclusion of [MA169 trade name] in the list of prequalified medicinal products.

^{*} Trade names are not prequalified by WHO. This is the national medicines regulatory authority's responsibility.

Pyrimethamine/sulfadoxine+amodiaquine (hydrochloride) 25 mg/500 mg + 150 mg dispersible tablets (Universal Corporation Ltd.), MA169

| Initial acceptance | Date | Outcome |
|---|---|---------|
| Status on PQ list | 30 July 2023 | listed |
| Quality | 20 January 2023 | MR |
| Bioequivalence | 24 January 2023 | MR |
| Safety, efficacy | NA | NA |
| GMP (re-)inspection | | |
| API | 22 June 2021 | MR* |
| APIs | 24 June 2022 | MR |
| APIs | 28 June 2022 | MR |
| FPP | 17 March 2023 | MR |
| GCP/GLP (re-)inspection | 27 July 2020 | MR* |
| GCP/GLP (re-)inspection | 27 May 2022 | MR |
| API: active pharmaceutical ingredient FPP: finished pharmaceutical product GCP: good clinical practice [quality standard] GLP: good laboratory practice [quality standard] | GMP: good manufacturing practice [quality standard] MR: meets requirements MR*: desk review (based on recent inspection reports) NA: not applicable, not available PQ: prequalification | |

Summary of prequalification status for [MA169 trade name]:

The table represents the status of relevant completed activities only.