FDA/SMC/SMD/VGU/18/0192

Distribution list attached

21st May 2018

Dear Healthcare Professional,

POTENTIAL FOR INCREASED LONG-TERM RISKS WITH CLARITHROMYCIN IN PATIENTS WITH HEART DISEASE

The Food and Drugs Authority (FDA) wishes to bring to your attention a potential increased risk of heart problems or death that can occur years later in patients with heart diseases who are prescribed with clarithromycin.

This information followed the recent review by the United States Food and Drugs Administration of the results of a 10-year follow-up study\(^1\) of patients with coronary heart disease and from a large clinical trial\(^2\) that first observed this safety issue.

Meanwhile, the Food and Drugs Authority (FDA) had received two individual case safety reports of cardiovascular related adverse drug reaction described as palpitations to clarithromycin; both patients fully recovered and were also on other medications.

The FDA has informed marketing authorization holders of clarithromycin containing products registered by the Authority to add the new warning about this increased risk of death in patients with heart disease to the product labels.

Prescribers should therefore be aware of these significant risks and weigh the benefits and risks of clarithromycin before prescribing it to any patient, particularly in patients with heart disease and even for short periods, and consider using other available antibiotics. Prescribers are also to advise patients with heart disease of the signs and symptoms of cardiovascular problems, regardless of the medical condition for they are being treated with clarithromycin.

Clarithromycin is a semi-synthetic macrolide antimicrobial for oral use and registered by the FDA for the following indications when caused by susceptible bacteria: Bacterial pharyngitis, Mild to moderate community acquired pneumonia, Acute bacterial sinusitis (adequately diagnosed), Acute exacerbation of chronic bronchitis, Skin infections and


soft tissue infections of mild to moderate severity, and in appropriate combination with antibacterial therapeutic regimens and an appropriate ulcer healing agent for the eradication of *Helicobacter pylori* in patients with *H. pylori* associated ulcers.

The FDA will like to advice healthcare professionals to educate patients on the possible side effect of all medicines including clarithromycin and report these to the FDA by completing the Adverse Reaction Reporting Form or online using the link http://adr.fdaghana.gov.gh/ or call 024 431 0297 or send an email to drug.safety@fdaghana.gov.gh.

Yours faithfully,

[Signature]

DELESE A. A. DARKO (MRS)
CHIEF EXECUTIVE OFFICER