



PHARMACY AND POISONS BOARD
HONG KONG
香港藥劑業及毒藥管理局

Your Ref. :
貴處檔號

Our Ref. : DH DO PRIE/1-55/1
本局檔號

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圖文傳真

C/O Drug Office
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15th December 2020

To: Certificate holders of
registered pharmaceutical products

Dear Sir / Madam,

New Warnings for Hydrochlorothiazide Products

On 7th December 2020, the Pharmacy and Poisons (Registration of Pharmaceutical Products and Substances: Certification of Clinical Trial/Medicinal Test) Committee (the Committee) considered the latest new warnings for hydrochlorothiazide-containing products by the drug regulatory authorities of the United States, Canada, European Union, United Kingdom, Australia and Singapore, and decided that the sales pack labels and / or package inserts of registered pharmaceutical products containing hydrochlorothiazide should include the following new safety information (or equivalent):

“Special warnings and precautions for use

Non-melanoma skin cancer

An increased risk of non-melanoma skin cancer (NMSC) [basal cell carcinoma (BCC) and squamous cell carcinoma (SCC)] with increasing cumulative dose of hydrochlorothiazide exposure has been observed in two epidemiological studies based on the Danish National Cancer Registry.

Photosensitizing actions of hydrochlorothiazide could act as a possible mechanism for NMSC.

Patients taking hydrochlorothiazide should be informed of the risk of NMSC and advised to regularly check their skin for any new lesions and promptly report any suspicious skin lesions. Possible preventive measures such as limited exposure to sunlight and UV rays and, in case of exposure, adequate protection should be advised to the patients in order to minimize the risk of skin cancer. Suspicious skin lesions should be promptly examined potentially including histological examinations of biopsies. The use of hydrochlorothiazide may also need to be reconsidered in patients who have experienced previous NMSC."

You are therefore required to review and revise, if necessary, the sales pack labels and / or package inserts of the concerned products registered by your company to ensure that the products comply with the above new requirements. The revised sales pack labels and / or package inserts should be submitted to the Committee for approval within 2 months from the date of this letter. Failing to comply with the above requirements may result in de-registration of the products or registration not renewed by the Committee.

If you have any enquiries on the above issue, please contact Ms. Queenie CHAN at 3974 4147.

Yours faithfully,



(T.K. YIM)
Secretary,

Pharmacy and Poisons (Registration of
Pharmaceutical Products and Substances:
Certification of Clinical Trial/Medicinal Test)
Committee

c.c. 7-15/3, Product Files

TK/QC