

PHARMACY AND POISONS ORDINANCE

藥劑業及毒藥條例

[reg. 29(6)]
[第 29(6)條]

(Chapter 138)

(第 138 章)

FREE SALE CERTIFICATE OF PHARMACEUTICAL PRODUCT

藥劑製品自由銷售證明書

Name and dosage form of product (specify strength):
製品的名稱及劑型(指明劑量):

Name and amount of each active ingredient (as provided by manufacturer):
每種有效成分的名稱及分量(按製造商所提供的資料):

.....
.....
.....

Manufacturer, and/or when applicable, the person responsible for placing the product on the market:
製造商及／或(如適用)負責將該製品推出市場出售的人:

.....
.....

Address(es):
地址:

It is certified that:

現證明:

- * This product has been authorized to be placed on the market for use in Hong Kong.
該製品已獲准推出市場出售以供在香港使用。

Number of permit and date of issue:
許可證編號及發出日期

- * This product has not been authorized to be placed on the market for use in Hong Kong for the
該製品不准推出市場出售以供在香港使用，理由如下:

following reasons:
.....
.....

- * It is also certified that (a) the manufacturing plant in which the product is produced is subject to inspection
現亦證明 (a) 製造該製品的製造廠每隔適當的期間即受到檢查，並且 (b) 就
at suitable intervals, and (b) the manufacturer conforms to requirements for good practices in the manufacture
在原產國家內銷售或分銷或供出口的製品而言，製造商已遵行藥品檢查合作
and quality control, as published by the Pharmaceutical Inspection Co-operation Scheme, in respect of
計劃出版關於製造與素質控制方面的良好做法的規定。
products to be sold or distributed within the country of origin or to be exported.

HONG KONG

香港

(Date)

..... (日期)

..... 代行
for Pharmacy and Poisons Board.
藥劑業及毒藥管理局