

LEGISLATIVE COUNCIL BRIEF

Pharmacy and Poisons Ordinance (Cap. 138)

PHARMACY AND POISONS (AMENDMENT) (NO. 5) REGULATION 2017

INTRODUCTION

The Pharmacy and Poisons Regulations (Cap. 138A) (“the Regulations”) was made under section 29 of the Pharmacy and Poisons Ordinance (Cap. 138) (“the Ordinance”). The Pharmacy and Poisons (Amendment) (No. 5) Regulation 2017 (“the Amendment Regulation”) at **Annex A** is to amend the First, Third and Tenth Schedules to the Regulations.

JUSTIFICATIONS

General Background

2. Under section 29(1B) of the Ordinance, the Pharmacy and Poisons Board (“the Board”) set up under section 3 of the Ordinance is empowered to make regulations to amend the Poisons List; or any list of articles or substances in the Regulations, subject to section 31 of the Ordinance and to the approval of the Secretary for Food and Health.

Proposal of the Pharmacy and Poisons Board

3. Arising from applications for registration of a pharmaceutical product, the Board proposes adding the following substance to Division A of the First Schedule, Division A of the Third Schedule and Division A of Part I of the Poisons List set out in the Tenth Schedule to the Regulations:

(a) Netupitant; its salts

4. Details of the above substance are set out at **Annex B**. The Board considers the proposed amendments appropriate in view of the potency, toxicity and potential side effects of the substance.

THE AMENDMENT REGULATION

5. The Amendment Regulation is to add the above drug (in paragraph 3) to the relevant Schedules to the Regulations.

LEGISLATIVE TIMETABLE

6. The legislative timetable will be –

Publication in the Gazette	13 October 2017
Date of Commencement	13 October 2017

IMPLICATIONS OF THE PROPOSAL

7. The proposal will impose appropriate control on pharmaceutical products consisting of the above substance so that they can be sold in the market upon fulfillment of the relevant regulations.

ENQUIRY

8. For any enquiries, please contact Mr Lam Fong-tat James, Assistant Secretary for Food and Health at 3509 8956.

Food and Health Bureau
October 2017

**Pharmacy and Poisons (Amendment) (No. 5) Regulation
2017**

(Made by the Pharmacy and Poisons Board under section 29(1B) of the Pharmacy and Poisons Ordinance (Cap. 138) subject to the approval of the Secretary for Food and Health)

1. Pharmacy and Poisons Regulations amended

The Pharmacy and Poisons Regulations (Cap. 138 sub. leg. A) are amended as set out in sections 2, 3 and 4.

2. Schedule 1 amended (substances to which certain restrictions with respect to the sale, supply, labelling and storage apply under regulations 3, 5, 6, 22 and 24)

Schedule 1, Division A, after item “Nesiritide”—

Add

“Netupitant; its salts”.

3. Schedule 3 amended (substances required by regulation 9 to be sold by retail only upon a prescription given by a registered medical practitioner, registered dentist or registered veterinary surgeon)

Schedule 3, Division A, after item “Nesiritide”—

Add

“Netupitant; its salts”.

4. Schedule 10 amended (Poisons List)

Schedule 10, section 2, Table, Part 1, Division A, after item “Nesiritide”—

Add

“Netupitant; its salts”.



Chairman,
Pharmacy and Poisons Board

4 October 2017

Explanatory Note

This Regulation amends the Pharmacy and Poisons Regulations (Cap. 138 sub. leg. A) to add Netupitant and its salts (*Netupitant*) to—

- (a) Division A of its Schedule 1;
- (b) Division A of its Schedule 3; and
- (c) Division A of Part 1 of the Poisons List set out in its Schedule 10.

2. The amendments relate to requirements concerning the sale, supply, labelling and storage of Netupitant. Main effects of the amendments include—

- (a) that the sale, by retail, of Netupitant—
 - (i) may only be effected on registered premises of an authorized seller of poisons by a registered pharmacist or in the presence and under the supervision of a registered pharmacist; and
 - (ii) may only be effected on and in accordance with a prescription by a registered medical practitioner, registered dentist or registered veterinary surgeon; and
- (b) that Netupitant, if stored in retail premises, must be stored properly in a part of the premises to which customers are not permitted access.

Pharmacy and Poisons (Amendment) (No.5) Regulation 2017

Supplementary Information to the Legislative Council

《2017年藥劑業及毒藥（修訂）（第5號）規例》

提交立法會的補充資料

<p>Netupitant; its salts</p>	<p>Part 1 of the Tenth Schedule, First and Third Schedules poison</p>	<p>This drug is used in adults and in combination with palonosetron for the prevention of acute and delayed nausea and vomiting associated with highly emetogenic cisplatin-based and moderately emetogenic cancer chemotherapy.</p> <p>Side effects include headache, constipation and fatigue.</p> <p>Its use should be decided by a doctor based on the patient's conditions.</p>
<p>奈妥匹坦；其鹽類</p>	<p>附表十的第一部，附表一及附表三毒藥</p>	<p>此藥物與帕洛諾司瓊聯合使用於成年患者，預防與高致吐性順鉑基礎及中度致吐性癌症化療相關的急性和延遲性噁心和嘔吐。</p> <p>副作用包括頭痛、便秘和疲勞。</p> <p>使用此藥物與否，須由醫生按病人情況決定。</p>