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Message from the Chairman 主席獻辭

In 2019, the Pharmacy and Poisons Board ("the Board") remained steadfast in its commitment to protecting and promoting public health through the regulation of pharmaceutical products, pharmacy trade and pharmacy profession. We are pleased to see the remarkable achievements in the year.

To tackle the global public health concern of antimicrobial resistance ("AMR") and with reference to the "Hong Kong Strategy and Action Plan on Antimicrobial Resistance (2017 - 2022)", the Board has actively implemented various measures for strengthening regulation on over-thecounter purchase of prescription-only antimicrobials. These measures include special inspections against authorized sellers of poisons ("ASPs") by the Drug Office of the Department of Health, conducting more test purchases of antimicrobials against retailers and enhancement of disciplinary actions against licensees who have been convicted of offences related to sale of antimicrobials. With the enhanced measures and support from the pharmacy profession and the pharmacy trade, the overall supply of antimicrobials to ASPs decreased from 18.4% in 2016 to 8.0% in 2018. In addition, the Board took the initiative to strengthen the control of certain antimicrobials which were used to be over-thecounter medicines by re-classifying them as prescription drugs. These antimicrobials include nifuratel, nifuroxazide, nitrofural, nitrofurantoin and nitroxoline, the new sales restrictions of which would take effect from 18 October 2020.

In view of the high risks associated with the rapid scientific development of advanced therapy products ("ATPs"), consideration was given to introducing a dedicated regulatory framework for controlling ATPs under the Pharmacy and Poisons Ordinance (Cap. 138, Laws of Hong Hong) ("the Ordinance"). After completion of public consultation on the regulation of ATPs in 2018, the Board has been supporting the Government to work on the legislative amendment of the Ordinance in order to enhance regulation of ATPs. Legislative proposals were made to the Panel on Health Services of the Legislative Council in April 2019 and the Pharmacy and Poisons (Amendment) Bill 2019 was introduced and first read at the Legislative Council on 30 October 2019. It is anticipated that the legislative amendments would provide Hong Kong with a regulatory framework for ATPs which is on a par with the international practice for better protection of public health.

In order to ensure that marketed pharmaceutical products are safe and of good quality, the Board has continuously made reference to the international regulatory practice and requirements and imposed various

二零一九年,香港藥劑業及毒藥管理局(下稱「管理局」)繼續克盡厥職, 透過規管藥劑製品、藥劑行業及藥劑 專業,致力保障和促進公眾健康。年 內有關工作成效卓著,令人欣悦。

為應對抗菌素耐藥性此一全球關注的 公共衞生議題,並參照《香港抗菌素 耐藥性策略及行動計劃(2017-2022)》, 管理局已積極推行多項措施,藉此加 強監管在沒有處方下售賣抗菌素的情 况。有關措施包括由衞生署藥物辦公 室對獲授權毒藥銷售商進行特別巡查 和對零售商進行更多的抗菌素試買行 動,以及加強紀律處分以懲處因干犯 與售賣抗菌素相關罪行而被定罪的持 牌人士。透過上述加強措施及藥劑專 業和藥劑行業的支持,該等銷售商供 應的抗菌素佔整體供應的比例由二零 一六年的 18.4% 下降至二零一八年的 8.0%。此外,為加強對抗菌素的管制, 管理局主動將若干原屬非處方藥物的 抗菌素改列為處方藥物,當中包括硝 呋太爾、硝呋齊特、呋喃西林、呋喃 妥因及硝羥喹啉。新訂的銷售管制將 於二零二零年十月十八日生效。

先進療法製品的科學發展迅速,鑑於 此類製品存在的高風險,管理局考慮 在香港法例第 138 章《藥劑業及毒藥 條例》(下稱「《條例》」)增設專屬的 規管框架以管制先進療法製品。有關 規管此類製品的公眾諮詢於二零一八 年完成後,管理局協助政府着手進行 修訂《條例》的工作,以期加強對先進 療法製品的規管。二零一九年四月, 政府已將有關修訂法例建議提交立法 會衞生事務委員會,並於二零一九年 十月三十日將《2019年藥劑業及毒藥 (修訂)條例草案》提交立法會進行首 讀。《條例》經修訂後,將使香港具有 與國際做法看齊的先進療法製品規管 框架,以加強保障公眾健康。

enhancement measures in 2019. With effect from 11 May 2019, the sales restrictions of pharmaceutical products containing desflurane, isoflurane and sevoflurane, which are used as general anesthetics, have been strengthened by classifying them as prescription drugs. Besides, starting from 1 October 2019, all registered non-sterile pharmaceutical products have been required to meet the acceptance criteria for microbiological standards prescribed in the pharmacopeias throughout their shelf life. The content of Class 1 elemental impurities (i.e. arsenic, cadmium, mercury and lead) in registered pharmaceutical products should not exceed the permitted daily exposure specified in the Guideline for Elemental Impurities promulgated by the International Council for Harmonisation of Technical Requirements for Pharmaceuticals for Human Use (ICH Q3D). In addition, the Board endorsed in October 2019 to tighten the sales control of skin antiseptic products containing chlorhexidine for human or animal use as pharmaceutical products unless otherwise stated, or except when (a) they are clearly labelled in English and Chinese for washing hands only (or equivalent); or (b) chlorhexidine is used as preservative or antimicrobials in cosmetic products. The tightened control would come into effect on 8 July 2020.

As for the development of the pharmacist profession, in response to the Government's Report of the Strategic Review on Healthcare Manpower Planning and Professional Development, the Board has set up the Task Force on Professional Development of Pharmacists to collect views from the stakeholders and make recommendations on the implementation of continuing professional education / continuing professional development. Opportunity will also be taken to collect views from the profession on the suggestion of setting up a pharmacy council in the long run.

It is only through the collective efforts of all the stakeholders that the Board can discharge its statutory functions efficiently. I am deeply grateful for the unwavering support and valuable contribution of all Members of the Board and the pharmacy community.

Dr CHAN Hon-yee, Constance, JP Chairman Pharmacy and Poisons Board 為確保在市場出售的藥劑製品安全及 優質,管理局於二零一九年繼續參照 國際規管慣例和規定,推行多項加強 措施。管理局由二零一九年五月十一 日起把含有地氟烷、異氟烷及七氟烷 而用作全身麻醉藥的藥劑製品改列為 處方藥物,藉此加強對相關藥劑製品 的銷售管制。此外,由二零一九年十 月一日起,所有非無菌的註冊藥劑製 品在整個保質期內必須符合藥典訂明 的微生物限度標準。註冊藥劑製品中 的第1類元素雜質(即砷、鎘、汞和 鉛)的含量不得超出國際醫藥法規協 調會議頒布的《元素雜質指南》(英文 簡稱為ICH Q3D) 所訂明的每日允許暴 露量。此外,管理局於二零一九年十 月同意收緊銷售規管,將含氯己定及 用於人或動物皮膚的消毒產品歸類為 藥劑製品(除另有説明外),或除非(a) 該產品清楚用中文及英文標示為只作 洗手用途(或同等標示);或(b)該氯 己定是用作化妝產品的防腐劑或抗菌 劑。收緊銷售規管的措施於二零二零 年七月八日生效。

在藥劑師專業的發展方面,為回應政府的《醫療人力規劃和專業發展策略檢討報告》,管理局成立了藥劑師專業發展專案組 (Task Force on Professional Development of Pharmacists),向持份者收集意見,同時就如何落實持續專業發展提出建議。管理局亦會藉此機會,收集業界對成立藥劑師管理局此項長遠建議的意見。

全賴各方持份者同心協力,管理局方 能順利履行其法定職能。各管理局成 員與藥劑業界一直支持管理局的工 作,堅定不移,貢獻良多,本人謹此 向各位衷心致謝。

> 藥劑業及毒藥管理局主席 陳漢儀醫生, JP

Introduction 引言

This annual report covers the calendar year 2019. Through this report, the Pharmacy and Poisons Board ("the Board") aims to keep all registered pharmacists and interested parties in the pharmacy profession posted on the functions and work of the Board during the year. A brief summary of the work of the Pharmacy and Poisons Appeal Tribunal ("the Appeal Tribunal") established under section 30 of the Pharmacy and Poisons Ordinance is also included.

In order to provide readers a quick reference, the description of the functions of the Board, its committees and the Appeal Tribunal has been simplified. Readers interested in more specific details of the statutory functions of these bodies should refer to the relevant provisions under the Pharmacy and Poisons Ordinance and its subsidiary legislation.

All inquiries with regard to this report or to the Board in general can be addressed to:

The Pharmacy and Poisons Board Secretariat 1/F, Shun Feng International Centre 182 Queen's Road East Wanchai, Hong Kong

Facsimile: (852) 2527 2277 Telephone: (852) 2527 8418 E-mail address: ppb@dh.gov.hk Website: www.ppbhk.org.hk 這份年報載錄藥劑業及毒藥管理局 (「管理局」)在二零一九年的工作。 管理局希望透過這份年報,使所有註 冊藥劑師以及藥劑業有關人士及團體 了解管理局的職能及在這年內的工作; 同時亦扼要介紹根據《藥劑業及毒藥 條例》第30條成立的藥劑業及毒藥上 訴審裁處(「上訴審裁處」)的工作。

為使讀者可以更容易掌握有關內容, 年報內對管理局及其轄下的委員會和 上訴審裁處的職能的描述以精簡為旨。 讀者如希望對這些組織的法定職能有 更深入的認識,請查閱《藥劑業及毒藥 條例》及其附屬法例的有關條文。

所有有關本年報或管理局的查詢,請 聯絡:

香港灣仔皇后大道東 182 號順豐國際中心1樓 藥劑業及毒藥管理局秘書處

圖文傳真: (852) 2527 2277 電話: (852) 2527 8418 電郵地址: ppb@dh.gov.hk 網址: www.ppbhk.org.hk

Membership and Functions of the Board 管理局的成員及職能



Dr CHAN Hon-yee, Constance, JP (Chairman) 陳漢儀醫生, JP (主席)



Dr SIN Wai-mei, Della, JP 單慧媚博士,JP



Mr CHAN Ling-fung, Frank 陳凌峯先生



Dr CHIU Pui-yin, Amy, JP 趙佩燕醫生,JP



Mr WONG Wai-hung, Geoffrey (Legal Adviser) 黃惠鴻先生 (法律顧問)



Professor WONG Chi-kei, lan 黃志基教授



Professor ZUO Zhong, Joan 左中教授



Mrs CHENG, Mary Catherine 鄭陳佩華女士



Mr WONG Hing-mang, Matthew 黃興孟先生



Mr YAU Fuk-loi, Rico 邱福來先生



Dr SO Yui-chi 蘇睿智醫生

1. Membership

Members of the Board are appointed by the Chief Executive. Each term is for a period of not more than three years. Members may be re-appointed. The current membership is as follows:

ex officio members

- (a) the Director of Health (Chairman);
- (b) the Government Chemist;
- (c) the Assistant Director of Health in the Drug Office of the Department of Health;
- (d) a medical officer in the Department of Health;
- (e) a legal adviser;
- (f) a full-time teaching staff of pharmacology of The University of Hong Kong;
- (g) a full-time teaching staff of pharmacology of The Chinese University of Hong Kong;
- (h) three registered pharmacists (not being public officers) nominated by the Pharmaceutical Society of Hong Kong; and
- (i) a registered medical practitioner (not being a public officer) nominated by the Hong Kong Medical Association.

The membership of the Board as at 31 December 2019 was as follows:

- (a) Dr CHAN Hon-yee, Constance, JP (Chairman)
- (b) Dr SIN Wai-mei, Della, JP
- (c) Mr CHAN Ling-fung, Frank
- (d) Dr CHIU Pui-yin, Amy, JP
- (e) Mr WONG Wai-hung, Geoffrey (Legal Adviser)
- (f) Professor WONG Chi-kei, lan
- (g) Professor ZUO Zhong, Joan
- (h) Mrs CHENG, Mary Catherine Mr WONG Hing-mang, Matthew Mr YAU Fuk-loi, Rico
- (i) Dr SO Yui-chi

Secretary

Miss WONG So-san, Suzanne

1. 成員

管理局的成員由行政長官委任,每屆 任期不多於三年,可以再獲委任。現 任成員包括:

- (a) 衞生署署長(主席);
- (b) 政府化驗師;

虽然 成員

- (c) 衞生署藥物辦公室的 衞生署助理署長;
- (d) 一名衞生署醫生;
- (e) 一名法律顧問;
- (f) 一名香港大學藥理學全職教員;
- (g) 一名香港中文大學藥理學全職教員;
- (h) 三名經香港藥學會提名的註冊藥 劑師(非公職人員);及
- (i) 一名經香港醫學會提名的註冊醫 生(非公職人員)。

在二零一九年十二月三十一日,管理局的成員計有:

- (a) 陳漢儀醫生, JP(主席)
- (b) 單慧媚博士, JP
- (c) 陳凌峯先生
- (d) 趙佩燕醫生, JP
- (e) 黃惠鴻先生(法律顧問)
- (f) 黃志基教授
- (g) 左中教授
- (h) 鄭陳佩華女士 黃興孟先生 邱福來先生
- (i) 蘇睿智醫生

秘書

黄素珊女士

2. Functions

The Board is established under section 3 of the Pharmacy and Poisons Ordinance to carry out the following functions in accordance with the provisions of the same Ordinance and its subsidiary legislation:

- (a) registration of pharmacists, including the prescription of training required for registration, the organization of registration examinations and the issue of certificates of registration and annual practising certificates;
- (b) discipline of pharmacists, through inquiry into their conduct by Disciplinary Committees appointed for the purpose;
- (c) licensing and regulatory control of retail traders of pharmaceutical products (authorized sellers of poisons and listed sellers of poisons), conducting inspections and test purchases and initiating prosecution of offences and, for authorized sellers of poisons, inquiring into their conduct by Disciplinary Committees appointed for the purpose;
- (d) licensing and regulatory control of wholesale dealers and manufacturers of pharmaceutical products;
- (e) regulatory control of the selling, purchasing, compounding and dispensing of pharmaceutical products; and
- (f) registration and classification of pharmaceutical products.

The Board is assisted by seven committees. They meet regularly to consider and decide on policies and actions in relation to the conduct of the above functions. The decisions of the Board and its committees are carried out jointly by the Secretariat of the Board and the Drug Office of the Department of Health.

2. 職能

管理局根據《藥劑業及毒藥條例》第3 條成立,執行該條例及其附屬法例規 定的下述職能:

- (a) 處理藥劑師註冊事宜,包括訂明 註冊所須的訓練、主辦註冊考試、 簽發註冊證明書及週年執業證明 書等;
- (b) 委出紀律委員會,調查藥劑師的 行為操守,並懲處被裁定行為不 當的藥劑師;
- (c) 規管及簽發藥劑製品零售商(獲授權毒藥銷售商及列載毒藥銷售商) 牌照。有關工作包括進行巡查及試買行動、提出檢控及委出紀律委員會調查獲授權毒藥銷售商的經營手法等:
- (d) 規管及簽發藥劑製品批發商和製 造商牌照;
- (e) 規管藥劑製品的銷售、購買、合成和配發事宜;及
- (f) 處理藥劑製品的註冊和分類事宜。

管理局轄下設有七個委員會。這些委員會定期舉行會議,就執行上述職能 審議和制定政策及行動計劃。管理局 及委員會的決定則由管理局秘書處及 衞生署藥物辦公室執行。

Membership and Functions of the Committees 管理局委員會的成員及職能

To assist the Board in performing its functions, the following seven committees are established under various provisions of the Pharmacy and Poisons Ordinance:

管理局根據《藥劑業及毒藥條例》內 相關的條文成立了下述七個委員會, 協助管理局執行職能:

(1) Examination Committee

(i) Membership as at 31 December 2019

Professor WONG Chi-kei, Ian (Chairman)

Dr SIN Wai-mei, Della, JP

Mr CHAN Ling-fung, Frank

Dr NG Ping-sum, Sammy

Ms CHU Kwok-pui, Jody

Mr LAM Fung-shing, Edwin

Dr LAM Tai-ning, Teddy

Dr WONG Siu-ming, Raymond

Ms TANG Suk-man, Alice (Secretary)

(ii) Functions

The Examination Committee is established under section 8(3) of the Pharmacy and Poisons Ordinance to:

- (a) advise the Board on matters regarding the registration of pharmacists and the training requirements and the examinations for registration;
- (b) draw up and review the syllabuses of the registration examinations;
- (c) appoint panels to assist in setting question papers and marking answer scripts for the registration examinations;
- (d) oversee the setting and marking of examination papers;
- (e) prepare and conduct the registration examinations;
- review the results of registration examinations and make recommendations regarding applicants' eligibility for registration to the Board for consideration;
- (g) consider complaints and unusual circumstances arising from applications for registration or examinations, and make recommendations to the Board for consideration; and
- (h) keep under review the standard of the registration examinations.

(1)考試委員會

(i) 截至二零一九年十二月三十一日 的成員名單

黃志基教授(主席)

單慧媚博士,JP

陳凌峯先生

吳秉琛醫生

朱幗珮女士

林豐盛先生

林泰寧博士

王紹明醫生

鄧淑雯女士(秘書)

(ii) 職能

考試委員會根據《藥劑業及毒藥條例》第8(3)條成立,負責:

- (a) 就有關藥劑師註冊、註冊的 訓練要求和考試的事宜向管 理局提供意見;
- (b) 制定及檢討註冊考試的範圍;
- (c) 委聘小組設定註冊試題及評 閱試卷;
- (d) 監督試卷設定及評卷工作;
- (e) 籌備及主辦註冊考試;
- (f) 覆核註冊考試的成績,並向 管理局就申請人的註冊資格 提交建議;
- (g) 調查註冊或考試申請的投訴 及異常情況,並提交建議供 管理局考慮;及
- (h) 檢討註冊考試的水平。

(2) Pharmacy and Poisons (Listed Sellers of Poisons) Committee

(i) Membership as at 31 December 2019

Mr CHAN Ling-fung, Frank (Chairman)

Ms AU YEUNG Kar-wai, Terese

Mr FONG Wing-kai, Guy

Ms LEUNG Chau-yung, Catherine

Mr MUI Cheuk-nang, Kenny

Ms TANG Mui-fun

Mr YAU Fook-wing, Edward William

Mr LEE Hi-fung, Lees (Secretary)

(ii) Functions

The Pharmacy and Poisons (Listed Sellers of Poisons) Committee is established to consider and approve applications for listed sellers of poisons under regulation 24A of the Pharmacy and Poisons Regulations.

(3) Pharmacy and Poisons (Wholesale Licences) Committee

(i) Membership as at 31 December 2019

Mr CHAN Ling-fung, Frank (Chairman)

Mr CHENG Kit-man, MH

Mr CHEUNG Yiu-kwong, Alex

Mr CHIU Kwok-leung, Philip

Mr WONG Chi-yin, Andrew

Mr WONG Hing-man

Ms TSE Po-yiu, Blouie (Secretary)

(2)藥劑業及毒藥(列載毒藥銷售商)委員會

(i) 截至二零一九年十二月三十一日 的成員名單

陳凌峯先生(主席)

歐陽嘉慧女士

方永佳先生

梁秋容女士

梅卓能先生

鄧梅芬女士

邱福榮先生

李浠鋒先生(秘書)

(ii) 職能

藥劑業及毒藥(列載毒藥銷售商) 委員會負責審批根據《藥劑業及 毒藥規例》第24A條提出的列載毒 藥銷售商牌照申請。

(3)藥劑業及毒藥(批發牌照) 委員會

(i) 截至二零一九年十二月三十一日 的成員名單

陳凌峯先生(主席)

鄭結文先生,MH

張耀光先生

趙國亮先生

黃志賢先生

王慶文先生

謝寶瑤女士(秘書)

In accordance with regulation 26 of the Pharmacy and Poisons Regulations, the Pharmacy and Poisons (Wholesale Licences) Committee is established to:

- (a) consider and approve applications for wholesale dealer licence, subject to any conditions it thinks fit to impose; and
- (b) revoke a wholesale dealer licence, suspend a wholesale dealer licence for a specified period, issue warning letter(s) to the licensed wholesale dealer or vary a condition of the wholesale dealer licence in the circumstances specified in regulation 26 of the Pharmacy and Poisons Regulations.

(4) Pharmacy and Poisons (Manufacturers Licensing) Committee

(i) Membership as at 31 December 2019

Mr CHAN Ling-fung, Frank (Chairman)

Dr CHENG Celine Heung-kwan

Dr CHEUNG Yan-ting, Kara

Professor LEE Wai-yip, Thomas

Dr MAK Yin-fong

Mr TSE Kin-on, Andrew

Dr WONG Sai-yin, Samson

Dr WONG Yiu-chung

Dr YEUNG Shu-ying, Ken

Mr YEUNG Yee-fai, Raphael (Secretary)

(ii) Functions

The Pharmacy and Poisons (Manufacturers Licensing) Committee is established to carry out the following functions in accordance with the Pharmacy and Poisons Regulations:

 (a) consider and approve applications for licence to manufacture pharmaceutical products, subject to any conditions it thinks fit to impose;

(ii) 職能

藥劑業及毒藥(批發牌照)委員會根據《藥劑業及毒藥規例》第26條,執行下列職能:

- (a) 在委員會認為適宜施加的條件的規限下,審議及批准批發商牌照的申請;及
- (b) 在《藥劑業及毒藥規例》第 26 條指明的情況下,撤銷批發商 牌照、在訂明期間內暫時吊 銷批發商牌照、向有關持牌 批發商發出警告信或更改施 加於批發商牌照的牌照條件。

(4)藥劑業及毒藥(製造商牌照)委員會

(i) 截至二零一九年十二月三十一日 的成員名單

陳凌峯先生(主席)

鄭香郡博士

張欣庭博士

李偉業教授

麥燕芳博士

謝建安先生

黄世賢醫生

黃耀松博士

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楊樹英博士

楊義輝先生(秘書)

(ii) 職能

藥劑業及毒藥(製造商牌照)委員會根據《藥劑業及毒藥規例》,執 行下列職能:

(a) 在委員會認為適宜施加的條件的規限下,審議及批准藥劑製品製造牌照的申請;

- (b) revoke a licence to manufacture pharmaceutical products, suspend a licence to manufacture pharmaceutical products for a specified period, issue warning letter(s) to the licensed manufacturer or vary a condition of the licence to manufacture pharmaceutical products in the circumstances specified in regulation 29 of the Pharmacy and Poisons Regulations;
- (c) consider and approve applications for registration as authorized person or renewal of registration as authorized person, subject to any conditions it thinks fit to impose; and
- (d) cancel the registration as authorized person, suspend the registration as authorized person for a specified period, issue warning letter(s) to the registered authorized person or vary a condition of the registration as authorized person in the circumstances specified in regulation 30F of the Pharmacy and Poisons Regulations.
- (5) Pharmacy and Poisons (Registration of Pharmaceutical Products and Substances: Certification of Clinical Trial/Medicinal Test) Committee
- (i) Membership as at 31 December 2019

Mr CHAN Ling-fung, Frank (Chairman)

Professor CHIM Chor-sang

Dr HO Chin-ho, Kenny

Dr HO King-man

Professor LOONG Ho-fung, Herbert

Dr NG Fook-hong

Professor TO Kin-wah, Kenneth

Dr TO Kwong-yuk

Mr WONG Yik-cheong, Anthony

Mr YIM Tsz-kok, Michael (Secretary)

- (b) 在《藥劑業及毒藥規例》第29 條指明的情況下,撤銷藥劑製品製造牌照或在指明期間內 暫時吊銷藥劑製品製造牌照、 向有關持牌製造商發出警告 信或更改施加於藥劑製品製 造牌照的牌照條件;
- (c) 在委員會認為適宜施加的條件的規限下,審議及批准註冊為獲授權人的註冊申請或續期申請;及
- (d) 在《藥劑業及毒藥規例》第 30F條指明的情況下,取消獲 授權人的註冊或在指明的期 間內暫時吊銷獲授權人的註 冊、向有關已註冊為獲授權 人發出警告信或更改註冊為 獲授權人所施加的註冊條件。
- (5)藥 劑 業 及 毒 藥 (藥 劑 製品及物質註冊:臨牀 試驗及藥物測試證明書) 委員會
- (i) 截至二零一九年十二月三十一日 的成員名單

陳凌峯先生(主席)

詹楚生教授

何展豪獸醫

何景文醫生

龍浩鋒教授

吳福康醫生

杜健華教授

杜光旭博士

黄益昌先生

嚴子閣先生(秘書)

The Pharmacy and Poisons (Registration of Pharmaceutical Products and Substances: Certification of Clinical Trial/Medicinal Test) Committee is established to carry out the following functions in accordance with the Pharmacy and Poisons Regulations:

- (a) consider new or renewal applications for registration of pharmaceutical products or substances, and issue registration certificates subject to any conditions it thinks fit to impose;
- (b) deregister a pharmaceutical product or substance, suspend the registration of a pharmaceutical product or substance for a specified period, issue warning letter(s) to the holder of a registration certificate or vary a condition of the registration of pharmaceutical products or substances;
- (c) consider applications for approval to change any of the registrable particulars of a pharmaceutical product or substance;
- (d) consider applications for conducting a clinical trial on human beings or a medicinal test on animals, and issue a clinical trial certificate or medicinal test certificate, subject to any conditions it thinks fit to impose; and
- (e) cancel a clinical trial certificate or medicinal test certificate, suspend a clinical trial certificate or medicinal test certificate for a specified period, issue warning letter(s) to the holder of the certificate or vary a condition of the certificate.

(6) Poisons Committee

(i) Membership as at 31 December 2019

Dr SIN Wai-mei, Della, JP (Chairman)

Mr CHAN Ling-fung, Frank

Mrs CHENG, Mary Catherine

Dr SO Yui-chi

Professor WONG Chi-kei, lan

Mr WONG Hing-mang, Matthew

Ms TANG Suk-man, Alice (Secretary)

(ii) 職能

藥劑業及毒藥(藥劑製品及物質註冊:臨牀試驗及藥物測試證明書)委員會根據《藥劑業及毒藥規例》,執行下列職能:

- (a) 在委員會認為適宜施加的條件 的規限下,審議藥劑製品或物 質的新註冊申請或續期註冊申 請以及簽發註冊證明書;
- (b) 撤銷藥劑製品或物質的註冊、在 指明期間內暫時吊銷藥劑製品或 物質的註冊、向有關註冊證明書 持有人發出警告信或更改施加於 藥劑製品或物質的註冊條件;
- (c) 審議有關更改藥劑製品或物質 註冊詳情的申請;
- (d) 在委員會認為適宜施加的條件 的規限下,審議有關對人類進 行臨牀試驗或對動物進行藥物 測試的申請以及簽發臨牀試驗 證明書或藥物測試證明書;及
- (e) 取消臨牀試驗證明書或藥物測 試證明書、在指明期間內暫時 吊銷臨牀試驗證明書或藥物測 試證明書、向有關證明書的持 有人發出警告信或更改施加於 證明書的條件。

(6)毒藥委員會

(i) 截至二零一九年十二月三十一日 的成員名單

單慧媚博士,JP(主席)

陳凌峯先生

鄭陳佩華女士

蘇睿智醫生

黄志基教授

黃興孟先生

鄧淑雯女士(秘書)

The Poisons Committee is set up under section 31 of the Pharmacy and Poisons Ordinance to advise the Board on the classification and distribution of poisons in Part 1 and Part 2 of the Poisons List and matters relating to the control of poisons and pharmaceutical products, including:

- (a) the classification of pharmaceutical products pending registration; and
- (b) the review of the classification of pharmaceutical products regulated under the Pharmacy and Poisons Regulations.

(7) Pharmacy Internship Training Committee

(i) Membership as at 31 December 2019

Professor ZUO Zhong, Joan (Chairman)

Mr CHAN Ling-fung, Frank

Mr CHENG Kit-man, MH

Mr CHEUNG Yiu-kwong, Alex

Ms HSUEH Cheung-mei, Vivian

Mr LAM Fung-shing, Edwin

Dr LAU Sze-ngar, Grace

Dr LEUNG Pak-heng, George

Mr LOK Wing-huen, Winham

Dr NG Ping-sum, Sammy

Professor TO Kin-wah, Kenneth

Dr TO Kwong-yuk

Mr WONG Ka-kin, Andy

Ms TANG Suk-man, Alice (Secretary)

(ii) 職能

毒藥委員會根據《藥劑業及毒藥條例》第31條成立,就各種毒藥在毒藥表第1部及第2部中的分類及分配,以及有關管制毒藥及藥劑製品的事宜,向管理局提供意見。有關事宜包括:

- (a) 有待註冊的藥劑製品的分類; 及
- (b) 檢討根據《藥劑業及毒藥規例》管制的藥劑製品的分類。

(7)藥劑師實習培訓委員會

(i) 截至二零一九年十二月三十一日 的成員名單

左中教授(主席)

陳凌峯先生

鄭結文先生,MH

張耀光先生

薛長薇女士

林豐盛先生

劉思雅博士

梁栢行博士

駱永煊先生

吳秉琛醫生

杜健華教授

杜光旭博士

黄家健先生

鄧淑雯女士(秘書)

The Pharmacy Internship Training Committee is set up under the Board to:

- (a) assist the Board in the registration of internship training institutions and preceptors;
- (b) assist the Board in drawing up the criteria for the approval of the content of the preceptor's quarterly appraisal forms and the intern's annual assessment forms proposed by the different training institutions and in implementing the criteria, and to establish sub-committees for these purposes where necessary;
- (c) assist the Board in drawing up the criteria for the evaluation of the preceptor's quarterly appraisals and the intern's annual assessments and in implementing the criteria, and to establish sub-committees for these purposes where necessary;
- (d) advise the Board on matters pertaining to pharmacy internship training;
- (e) liaise with internship training institutions and with preceptors on matters pertaining to internship training as necessary; and
- (f) carry out such other functions connected with internship training as may be permitted or assigned to the Committee by the Board.

(ii) 職能

藥劑師實習培訓委員會由管理局 成立,負責:

- (a) 協助管理局處理實習培訓機構 及導師註冊事宜;
- (b) 協助管理局制訂準則用以批核 由不同培訓機構提交的導師所 用的季度評核表格及實習人員 所用的年度評核表格,以及執 行這些準則並按需要設立小組 委員會;
- (c) 協助管理局制訂準則用以審核 導師提交的季度評核表格及實 習人員提交的年度評核表格, 以及執行這些準則並按需要設 立小組委員會;
- (d) 就有關藥劑師實習培訓的事宜 向管理局提供意見;
- (e) 按需要與實習培訓機構及導師 緊密聯絡;及
- (f) 執行管理局所容許並賦予的有 關實習培訓的其他職能。

The Work of the Board and its Committees 管理局及其委員會的工作

(1) Registration of Pharmacists

Any person intending to practise as a pharmacist in Hong Kong must first be registered with the Board. To be eligible for registration, an applicant must be able to meet the qualification, examination and training requirements specified by the Board.

(i) Qualification

An applicant must satisfy either one of the following two criteria:

- (a) holds a pharmacy degree awarded by a recognized university in Hong Kong; or
- (b) non-local applicant must have completed his/her tertiary education of not less than three full-time academic years or equivalent in pharmacy, and be registered or be professionally qualified to be registered as a pharmacist normally in the country in which he/she has completed his/her studies in pharmacy.

(ii) Examination

An applicant who possesses the qualification in (i)(b) above must also pass the Board's registration examinations in three subjects, namely Pharmacy Legislation in Hong Kong, Pharmacy Practice and Pharmacology.

The Examination Committee conducted two registration examinations in June and December 2019. A total of 70 applicants cumulatively passed all the three subjects in 2019.

The results of these two registration examinations are shown in **Table 1**. Figures for 2015 to 2019 are also included for comparison purpose.

(iii) Training

Applicants holding a pharmacy degree awarded by a recognized university in Hong Kong are required to undergo Board-approved training for one year before they can be registered as pharmacists.

(1)藥劑師的註冊

擬於香港執業的藥劑師必須向管理局 註冊。申請人必須具備管理局規定的 資格、考試成績及實習履歷,方符合 資格註冊。

(i) 資格

申請人必須符合下述其中一項條件:

- (a) 具備香港認可大學頒授的藥劑 學學位;或
- (b) 在本港以外地區完成不少於三個完整學年或相等的藥劑學課程,並已在其完成學業的地區註冊為藥劑師;或取得註冊為藥劑師的專業資格。

(ii) 考試

符合上述 (i)(b) 項要求的申請人, 必須通過由管理局舉辦的三個科 目的註冊考試,包括香港藥劑法 例、藥劑執業及藥理學。

考試委員會在二零一九年分別在 六月及十二月舉辦了兩次註冊考 試。同年共有70人累積取得全部 三科合格的成績。

表 1 列出該兩次註冊考試的成績, 以及二零一五年至二零一九年的 有關數字,以供比較。

(iii) 實習

持有香港認可大學頒授的藥劑學學位的申請人,在獲准成為註冊藥劑師前,須接受管理局認可的實習訓練,為期一年。

Applicants holding a recognized pharmacy degree awarded elsewhere should have had an aggregate of relevant preregistration training and post-registration experience of not less than one year. An applicant with less than one year's training and experience may be permitted to take the registration examinations but, upon passing all parts thereof, will be required to undergo an appropriate period of make-up training specified by the Board.

(iv) Registration

Upon registration, the Secretary of the Board will issue to a registered pharmacist a certificate of registration.

The Secretary is also responsible for keeping a register of pharmacists in which particulars of all pharmacists registered in Hong Kong are entered. The register is open for inspection by the general public. A copy of the register is also published in the Gazette once every 12 months.

(v) Practising Certificates

All practising pharmacists must obtain a practising certificate annually in accordance with section 10A of the Pharmacy and Poisons Ordinance. A total of 3 001 registered pharmacists were issued with practising certificates for 2019. Statistical data regarding registration of pharmacists and breakdown of fresh registration, removal from the register, and re-registration of pharmacists for 2015 to 2019 are shown in **Tables 2 and 3**.

(vi) Discipline of Pharmacists

Inquiries into the conduct of registered pharmacists are made by Disciplinary Committees appointed under section 15 of the Pharmacy and Poisons Ordinance. A registered pharmacist found guilty of misconduct will be subject to disciplinary sanctions, ranging from censure, written warning, to removal from the register of pharmacists for a specified period of time. A more detailed account of the set-up and work of Disciplinary Committees is given in pages 24 to 26 of this report.

Statistical data regarding disciplinary actions taken by the Board in respect of registered pharmacists in 2015 to 2019 are shown in **Tables 4, 5 and 6**.

持有其他地方頒發的認可藥劑學 學位的申請人,他的註冊前實習 訓練及取得註冊後的工作經驗 合共不可少於一年。訓練及經驗 合共少於一年的申請人亦可獲得 批准參加註冊考試,惟通過全部 考試後,須接受一段管理局認可 的補償實習。

(iv) 註冊

一經註冊,管理局秘書會向註冊 藥劑師發出註冊證明書。

管理局秘書負責備存一份藥劑師名冊,詳列所有在香港註冊的藥劑師的個人資料,並公開予市民查閱。該名冊每十二個月在憲報刊登一次。

(v) 執業證明書

所有執業藥劑師必須根據《藥劑業及毒藥條例》第10A條的規定取得週年執業證明書。共有3001位註冊藥劑師獲發2019年的執業證明書。表2及3列出二零一五年至二零一九年有關藥劑師註冊,以及新註冊、刪除註冊及重新註冊的分項數字的統計資料。

(vi) 藥劑師的紀律事宜

管理局根據《藥劑業及毒藥條例》第 15 條的規定,委出紀律委守納定,調查註冊藥劑的行為操師的行為內方為不當的註冊藥劑的註一藥劑的主體受紀律制裁,包括被譴責劑的接受紀律制定的時期內從藥賣會的名冊上除名。有關紀律委員會的組成及工作詳情,可參閱本年報 24 至 26 頁。

表 4、5 及 6 詳列管理局在二零 一五年至二零一九年對註冊藥劑 師採取紀律行動的統計數字。

(2) Licensing and Regulatory Control of Retail Traders (Including Authorized Sellers of Poisons and Listed Sellers of Poisons)

(i) Authorized Sellers of Poisons: Licensing

An authorized seller of poisons ("ASP"), commonly known as "pharmacy" or "dispensary", is a business authorized to sell poisons included in Part 1 of the Poisons List, by or under the supervision of a registered pharmacist. Any person wishing to operate as an ASP shall apply to the Board for registration of the premises where the retail sale of poisons is to be conducted. Besides, ASPs are also authorized to conduct retail sale of poisons included in Part 2 of the Poisons List at registered premises. If the Board is satisfied that the requirements stipulated in section 13(4) of the Pharmacy and Poisons Ordinance are complied with, the application will be granted, subject to the payment of the prescribed fee.

The name, the certificate of registration and a notice setting out the attending hours of the duty pharmacist are required to be displayed in a conspicuous location in the premises of an ASP. The ASP may also display a logo prescribed under section 13A of the Pharmacy and Poisons Ordinance.

ASPs must apply for renewal of registration annually. All records of an ASP will be taken into account when the Board assesses its application for renewal of registration of premises each year. If the ASP is not considered a fit and proper person to continue the retail business of poisons, the application will be rejected.

There were 649 ASPs registered in Hong Kong as at the end of 2019. Statistical data regarding the total number of ASPs, applications for registration and renewal of registration of premises of ASPs in 2015 to 2019 are shown in **Tables 7 and 8**.

(2)零售商(包括獲授權毒藥 銷售商及列載毒藥銷售商) 的發牌及規管工作

(i) 獲授權毒藥銷售商:發牌工作

獲授權毒藥銷售商須在其處所的當 眼處展示其當值藥劑師的姓名、註冊證明書及工作時間的告示。銷售商亦可以展示根據《藥劑業及毒藥條例》第13(A)條訂明的標識。

獲授權毒藥銷售商須每年向管理局申請註冊續期。管理局在審理獲授權毒藥銷售商的週年處所註冊續期申請時,會考慮有關銷售商的全部記錄。假如管理局認為實授權毒藥銷售商並不適宜繼續經營毒藥零售業務,管理局將拒絕其申請。

截至二零一九年年終,香港共有649名獲授權毒藥銷售商。表7及8詳列二零一五年至二零一九年獲授權毒藥銷售商的總數、處所註冊申請及註冊續期申請的統計數字。

(ii) Authorized Sellers of Poisons: Discipline

The premises registered with the Board are subject to inspection by pharmacist inspectors of the Department of Health. Test purchases to check illegal activities involving controlled drugs or unregistered pharmaceutical products are conducted and prosecutions are instituted against offenders.

Any act of alleged misconduct will be subject to inquiry by Disciplinary Committees appointed by the Board. If found guilty of misconduct, the ASP will be subject to disciplinary sanctions, ranging from written warning, variation on the conditions relating to the registration of premises to disqualification from being an ASP for a specified period of time.

Eleven inquiries were held in 2019 and all of the ASPs concerned were found guilty of misconduct. Six ASPs were issued with written warning whilst five others were disqualified from being an ASP for a period of time.

For minor infringement, if the pharmacist of the ASP concerned is directly involved in the case, the Board may direct the proprietor / director and duty pharmacist of the ASP to be interviewed by the Assistant Director (Drug) of the Department of Health and the Secretary of the Board to give them verbal cautions. On the other hand, verbal caution may be given to the director / proprietor in the presence of the pharmacist when the pharmacist is not involved in the case. A total of six such interviews were held in 2019.

The number of inspections and test purchases conducted by pharmacist inspectors against ASPs in 2015 to 2019 is shown in **Table 9**.

Statistical data regarding disciplinary cases handled by the Board in respect of ASPs in 2015 to 2019 are given in **Tables 10, 11, 12** and 12A.

(ii) 獲授權毒藥銷售商:紀律事宜

衛生署的藥劑師督察會巡查已經 向管理局註冊的銷售商處所。署 方亦會派員抽樣進行試買,偵查 涉及受管制藥物或未經註冊藥劑 製品的違法活動,並檢控違法者。

管理局會委出紀律委員會就任何 不當行為展開研訊。銷售商如被 裁定犯有不當行為,將會受到紀 律制裁,由書面警告、更改處所 註冊條件,以至在指定期間被取 消銷售商資格。

在二零一九年,管理局舉行了11次紀律研訊,全部涉事的獲授權毒藥銷售商被裁定犯有不當行為。紀律委員會向其中六名獲授權毒藥銷售商發出書面警告,五名獲授權毒藥銷售商則被取消銷售商資格一段時間。

表 9 列出二零一五年至二零一九年由藥劑師督察對獲授權毒藥銷售商進行巡查及試買的數字。

表 10、11、12 及 12A 詳 列 二 零 一五年至二零一九年管理局處理 有關獲授權毒藥銷售商的紀律個 案的統計數字。

(iii) Listed Sellers of Poisons: Licensing

A listed seller of poisons ("LSP"), commonly known as a medicine company, is a person approved to conduct the retail sale of poisons included in Part 2 of the Poisons List subject to the provision of the Pharmacy and Poisons Ordinance. Any person wishing to operate as a LSP should apply for his name to be entered onto the list of LSPs kept by the Board. On behalf of the Board, the Pharmacy and Poisons (Listed Sellers of Poisons) Committee issues licences to LSPs.

There were 4 295 LSPs as at the end of 2019. The number of licensed LSPs in 2015 to 2019 is shown in **Table 13**. Statistical data regarding applications for LSP licences in these five years are shown in **Table 14**.

(iv) Listed Sellers of Poisons: Discipline

Like ASPs, LSPs are also subject to inspection by pharmacist inspectors but unlike ASPs, no disciplinary inquiries by Disciplinary Committees are held to inquire into the conduct of a LSP. If a LSP is convicted of any offence under the Pharmacy and Poisons Ordinance, the Antibiotics Ordinance, the Dangerous Drugs Ordinance, the Trade Descriptions Ordinance, or the LSP has contravened the Code of Practice or licensing conditions, his case will be submitted to the Board for consideration. His name will be removed or suspended for a period specified by the Board from the list of LSPs if the Board considers him not a fit and proper person to continue the retail business of Part 2 poisons. For minor infringement, the Board may issue a written warning to the LSP concerned.

The number of inspections and test purchases conducted by pharmacist inspectors on LSPs in 2015 to 2019 is shown in **Table 15**. Statistical data regarding disciplinary cases handled by the Board in respect of LSPs in these five years are given in **Tables 16, 17 and 17A**.

(iii) 列載毒藥銷售商:發牌工作

列載毒藥銷售商一般稱為藥行, 是根據《藥劑業及毒藥條例》的規 定,獲准經營毒藥表內第2部毒藥 等售業務的人士。擬成為列載毒 銷售商的人士,可向管理局申 其姓名載入管理局備存的列載 其姓名載內。藥劑業及 載毒藥銷售商) 委員會會代 電 員 簽發牌照予列載毒藥銷售商。

截至二零一九年年終,香港共有 4 295 名列載毒藥銷售商。表 13 列出二零一五年至二零一九年列 載毒藥銷售商的總數。表 14 列出 在上述五年申請發牌的統計數字。

(iv) 列載毒藥銷售商:紀律事宜

衞生署藥劑師督察同樣地會巡查列 載毒藥銷售商的處所。但是,管理 局不會因調查列載毒藥銷售商的經 營手法而召開紀律研訊,這點與處 理有關獲授權毒藥銷售商的紀律事 宜的方法不同。假如有列載毒藥銷 售商被裁定干犯任何《藥劑業及毒 藥條例》、《抗生素條例》、《危險 藥物條例》、《商品説明條例》或違 反其《執業守則》或發牌條件,有 關個案將直接呈交管理局考慮。管 理局假如認為涉案的列載毒藥銷售 商並不適宜繼續經營第2部毒藥零 售業務,便會把該列載毒藥銷售商 的姓名從列載毒藥銷售商名單上刪 除或在指明的期間內暫時吊銷其名 列該名單內的資格。至於輕微的違 法行為,管理局可向有關的列載毒 藥銷售商發出書面警告。

表 15 列出二零一五年至二零一九年由藥劑師督察對列載毒藥銷售商進行巡查以及試買的數字。表 16、17 及 17A 詳列管理局在上述五年處理有關列載毒藥銷售商的紀律個案的統計數字。

(3) Licensing and Regulatory Control of Wholesalers and Manufacturers

(i) Wholesale Dealers of Pharmaceutical Products

Subject to the provisions of the Pharmacy and Poisons Regulations, any person wishing to deal in wholesale and/or import / export of poisons and/or pharmaceutical products should apply to the Pharmacy and Poisons (Wholesale Licences) Committee for an annual wholesale dealer licence.

A licensed wholesale dealer must keep a record of all transactions involving poisons included in Part 1 of the Poisons List or any pharmaceutical product. Sales of poisons are restricted to authorized persons only.

There were 770 holders of wholesale dealer licence as at the end of 2019. Statistical data for 2015 to 2019 are shown in **Table 18**.

(ii) Manufacturers of Pharmaceutical Products

Any person wishing to manufacture any pharmaceutical product should apply to the Pharmacy and Poisons (Manufacturers Licensing) Committee for a licence annually.

Manufacturers are subject to the provisions of the Pharmacy and Poisons Regulations. They are required to label the container of each pharmaceutical product with the appropriate particulars, such as the active constituents or ingredients of the product, the quantitative particulars of these constituents or ingredients, and the name and address of the manufacturer. They are also required to take adequate steps to ensure that all personnel involved in the manufacturing or packing of pharmaceutical products do not contaminate or infect the products.

It is the duty of every manufacturer to test each batch of raw material intended to be used in the manufacture of pharmaceutical products to ensure identity and purity, and to test its finished form to ensure identity and potency. A system of control that will enable the rapid and complete recall of any product from sale to the public should be put in place.

A manufacturer should also ensure that the premises and the fittings and machinery in such premises meet an appropriate standard in respect of temperature, humidity, cleanliness and hygiene, and that a set of records regarding the manufacture of pharmaceutical products is properly kept.

(3)批發商及製造商的發牌及 規管工作

(i) 藥劑製品批發商

除《藥劑業及毒藥規例》另有規定外,任何人如欲經營毒藥及/或藥劑製品批發及/或進/出口,均須向藥劑業及毒藥(批發牌照)委員會申請一年期的批發商牌照。

持牌的批發商須備存所有涉及毒藥表第1部所列毒藥或所有藥劑製品的交易記錄,而銷售對象只限於獲授權人士。

截至二零一九年年終,香港共有770名批發商牌照持有人。表18列出二零一五年至二零一九年的統計數字。

(ii) 藥劑製品製造商

任何人如欲製造任何藥劑製品, 每年均須向藥劑業及毒藥(製造 商牌照)委員會申請牌照。

製造商必須遵守《藥劑業及毒藥規例》的規定,他們須在等額等人藥劑等人藥劑等人藥劑等,他們須在時期,他們須當的標識等,一個一個人。 一個人。 一個人 一個一。 一個一

每名製造商必須測試擬用於製造藥劑製品的每一批原料,確保原料的本質及純度;及測試製成品,以確保其本質及效力。製造商亦須設立一套管理制度,以便能向市場迅立地完全回收任何正在銷售的產品。 製造商同時須確保其廠房以及、 製造商同時須確保其廠房,以及機器符合溫度、濕度 裝置及機器符合溫度、濕度 素及衛生的標準,以及備存一套有關生產藥劑製品的記錄。 The manufacture of pharmaceutical products must be supervised by a registered pharmacist or persons with such other qualifications as approved by the Board. A manufacturer must also ensure that at least one authorized person is employed to be responsible for ensuring and certifying that the pharmaceutical products are manufactured in accordance with the Pharmaceutical Inspection Co-operation Scheme Good Manufacturing Practice ("PIC/S GMP") Guide and registration requirements.

There were 70 holders of a manufacturer's licence as at the end of 2019, and all of them were required to comply with the GMP Guide with effect from 1 October 2015. Among these 70 holders, 47 holders of them were only authorized to conduct secondary packaging of pharmaceutical products. Statistical data for 2015 to 2019 are given in **Table 19**.

(4) Registration and Classification of Pharmaceutical Products

(i) Registration of Pharmaceutical Products

In accordance with regulation 36 of the Pharmacy and Poisons Regulations, any person wishing to sell, offer for sale, distribute or possess any pharmaceutical product or substance, shall register the product or substance with the Pharmacy and Poisons (Registration of Pharmaceutical Products and Substances: Certification of Clinical Trial/Medicinal Test) Committee.

In considering an application for registration of a pharmaceutical product, the Committee will take into account the safety, efficacy and quality of the subject product. In dealing with an application manufactured outside Hong Kong, the Committee may require the applicant to take any or all of the following actions:

- (a) produce an undertaking to permit the Committee to inspect the manufacturing premises;
- (b) produce a declaration that the subject product is manufactured in accordance with the requirements imposed by or under the law of the country concerned; and
- (c) pay a fee as representing the expenditure incurred by or on behalf of the Committee in carrying out an inspection at the manufacturing premises.

(4)藥劑製品的註冊及分類

(i) 藥劑製品的註冊

根據《藥劑業及毒藥規例》第36條的規定,任何人如欲銷售、要約出售、分銷或管有任何藥劑製品或物質,均須將有關製品或物質向藥劑業及毒藥(藥劑製品及物質註冊:臨牀試驗及藥物測試證明書)委員會註冊。

在決定是否批准某一藥劑製品申請註冊時,委員會會考慮該藥品的安全程度、效能及素質。處理在香港境外製造的產品的申請時,委員會可能要求申請人出示下列其中一份或全部文件:

- (a) 准許委員會視察其生產廠房的 承諾書;
- (b) 承諾該產品是遵照有關國家的 法律或根據法律施加的任何規 定而製造的聲明書;及
- (c) 繳付由委員會釐定的費用,該 筆費用相當於委員會或其代表 在視察生產廠房時所招致或相 當可能招致的開支。

A registration certificate will be issued on registration, and will be subject to any conditions the Committee thinks fit to impose. The applicant will also be advised of the classification of the product.

There were 16 186 registered pharmaceutical products in Hong Kong as at the end of 2019. The number of registered pharmaceutical products as at the end of the years 2015 to 2019 is shown in **Table 20**.

(ii) Classification of Pharmaceutical Products

On the advice of the Poisons Committee, the Board determines and reviews the classification and distribution of pharmaceutical products in the Poisons List and regulates their control by posing further restrictions on their sales through the provision of Schedules 1 and 3 of the Pharmacy and Poisons Regulations. The classification of pharmaceutical products in Schedule 10, i.e. Poisons List, and restrictions on sales under the two schedules are:

Classification Restriction(s) on sale They can be sold only by (a) Part 1 Poisons: Poisons included in authorized sellers of poisons Part 1 of Schedule 10, i.e. under the supervision of registered Poisons List pharmacists. They should also be stored in retail premises in a locked receptacle in a part of the premises to which customers do not have access. (b) Schedule 1 Poisons: They can be sold only by Poisons included in authorized sellers of poisons Part 1 of Schedule 10, i.e. under the supervision of registered Poisons List, and Schedule pharmacists and after entry in the 1 to the Pharmacy and poisons book stating the particulars Poisons Regulations of the sale. (c) Schedule 3 Poisons: They can be sold only by authorized Poisons included in sellers of poisons under the Part 1 of Schedule 10, i.e. supervision of registered pharmacists Poisons List, and Schedule with the authority of a prescription 3 to the Pharmacy and from a registered medical practitioner, Poisons Regulations registered dentist or registered veterinary surgeon. (d) Part 2 Poisons: They can be sold by listed sellers Poisons included in of poisons and authorized sellers Part 2 of Schedule 10, i.e. of poisons without the supervision

of registered pharmacists.

一經註冊,申請者會獲發註冊證明書,並獲告知產品的分類。

截至二零一九年年終,香港共有16 186 種已註冊的藥劑製品。 表20 列出截至二零一五年至二零 一九年年終的註冊藥劑製品數字。

(ii) 藥劑製品的分類

就毒藥委員會的建議,管理局會 決定及檢討藥劑製品在毒藥表內 的分類及分配,並透過《藥劑業 及毒藥規例》附表1和附表3,進 一步規管藥劑製品的銷售。藥劑 製品在附表10,即毒藥表,內的 各種不同分類及在附表1和附表3 內的銷售規管分述如下:

分類 銷售的限制

- (a) 第 1 部毒藥:在註冊藥劑師監督 附 表 10 ,即 毒藥表,第 1 銷售商銷售。這類 部所列毒藥 毒藥必須存放在上 鎖的盛器內,而盛 器則須存放在處所 內顧客不准進入的 地方。
- (b)附表 1 毒藥: 在註冊 藥劑師監同時列於附督下,由獲授權表 10,即毒毒藥銷售商銷售,藥表,第 1 部及《藥劑業及將銷售詳情記錄毒藥規例》附在毒藥冊中。表 1 的毒藥
- (c) 附表 3 毒藥: 須由註冊醫生、註 同時 列於 附冊牙醫或註冊獸醫 表 10, 即 毒處方授權,並在註 藥表,第 1 部冊藥劑師監督下, 及《藥劑業及 由獲授權毒藥銷售 毒藥規例》附商銷售。 表 3 的毒藥
- (d)第2部毒藥:無須藥劑師監督, 附表10,即由列載毒藥銷售 毒藥表,第2商或獲授權毒藥 部所列毒藥 銷售商銷售。

Poisons List

Regulatory provisions in other related areas are contained in Schedules 2 and 4 to 7 to the Pharmacy and Poisons Regulations:

Schedule	Provisions
Schedule 2	providing for certain articles to be exempted from some of the provisions of the Pharmacy and Poisons Ordinance and the Pharmacy and Poisons Regulations
Schedule 4	setting out the statement of particulars as to proportion of poisons in certain cases
Schedule 5	prescribing the labelling requirements for certain poisons
Schedule 6	listing out certain poisons which are exempted from labelling provisions when sold or supplied in certain circumstances
Schedule 7	listing out certain poisons which are required to be specially labelled for transport

Classification and distribution in Schedule 10, i.e. Poisons List, and imposition of control through the various schedules were made through amendments to the Pharmacy and Poisons Regulations. Amendments proposed by the Board and approved by the Legislative Council in 2019 are shown in **Table 21**.

《藥劑業及毒藥規例》附表 2、附表 4至 7 詳列對下述其他方面的規管:

附表	內容
附表 2	列舉豁免受《藥劑業及毒藥條例》及《藥劑業及毒藥 規例》一些條文規限的某 些物品
附表 4	詳列在某些情況下有關毒 藥比例的詳情説明
附表 5	説明對某些毒藥的標籤要求
附表 6	列出在某些情況下銷售或 供應則無須加上標籤的某 些毒藥
附表 7	列出為運輸而須特別加上 標籤的某些毒藥

管理局透過修訂《藥劑業及毒藥規例》,將藥劑製品在附表 10,即毒藥表內分類和分配,並透過多個附表對藥劑製品施加規管。立法會在二零一九年批准管理局就藥劑製品分類對《藥劑業及毒藥規例》作出的修訂列載於表 21。

Membership and Functions of the Disciplinary Committee 紀律委員會的成員及職能

(1) Membership

A Disciplinary Committee consists of the following persons:

- (a) a Chairman, who is the medical officer in the Department of Health appointed by the Chief Executive under section 3(2)(e) of the Pharmacy and Poisons Ordinance as a member of the Board;
- (b) two registered pharmacists (not being public officers) nominated by the Pharmaceutical Society of Hong Kong; and
- (c) a legal adviser appointed by the Chief Executive.

As at 31 December 2019, the Chairman of the Disciplinary Committee was Dr CHIU Pui-yin, Amy, JP, Controller, Regulatory Affairs of the Department of Health. Registered pharmacists who had served as members in 2019 included:

Mr CHAN Yat-ming

Ms CHAN Yin-yin, Ivy

Mr CHEN Wen-ben, Benny

Ms CHEUNG Oi-ling

Ms CHEW Leng-leng

Mr LEE Pak-hei

Mr LEE Siu-to

Mr LEUNG Kwong-hei, Kenneth

Ms MOK Lai-fong

Mr NG Wing-yan

Mr NG Yu-chau, Patrick

Mr SUNG Ming-tat, Dick

Ms TAM Hi

Ms TSANG Sheung-chee, Sandra

Mr WONG Chi-ming

Mr WONG Kwong-cheung, Aaron

(1)成員

紀律委員會的成員包括下列人士:

- (a) 一名根據《藥劑業及毒藥條例》第 3(2)(e) 條由行政長官委任為管理 局成員的衞生署醫生,並由其出 任主席;
- (b) 兩名由香港藥學會提名的註冊藥 劑師(非公職人員);及
- (c) 一名由行政長官委任的法律顧問。

衛生署規管事務總監趙佩燕醫生,JP 是紀律委員會在二零一九年十二月 三十一日的主席。曾在二零一九年出 任成員的註冊藥劑師包括:

陳日明先生

陳妍賢女士

陳文斌先生

張靄玲女士

周凌綾女士

李伯熙先生

李兆濤先生

梁廣熙先生

莫麗芳女士

吳榮恩先生

吳如就先生

沈明達先生

譚起女士

曾尚智女士

黄志明先生

黄庸長先生

(2) Functions

In accordance with section 15 of the Pharmacy and Poisons Ordinance, a Disciplinary Committee is appointed by the Board for the purpose of disciplinary inquiry if:

- (a) a complaint is received by the Board regarding the conduct of a registered pharmacist or an employee of a registered pharmacist, or it appears to the Board that a registered pharmacist has contravened a code of conduct applicable to the registered pharmacist;
- (b) a complaint is received by the Board regarding the conduct of an authorized seller of poisons ("ASP") or an employee, officer or partner of an ASP, or it appears to the Board that an ASP has contravened a code of practice applicable to the ASP;
- (c) any of the persons mentioned in (a) or (b) above, is convicted of an offence under:
 - the Pharmacy and Poisons Ordinance, the Dangerous Drugs Ordinance, the Antibiotics Ordinance, the Undesirable Medical Advertisements Ordinance; or
 - (ii) section 52, 54 or 61 of the Public Health and Municipal Services Ordinance or section 7, 7A or 9 of the Trade Descriptions Ordinance;
- (d) it appears to the Board that a condition imposed under section 13 of the Pharmacy and Poisons Ordinance in respect of the registration of any premises of an ASP has been contravened; or
- (e) it otherwise appears necessary or desirable to the Board to inquire into the conduct of any of the persons mentioned in (a) or (b) above.

In respect of a registered pharmacist or an employee of a registered pharmacist, the Disciplinary Committee may, at the conclusion of an inquiry:

- (a) censure the registered pharmacist;
- (b) issue a warning letter to the registered pharmacist; or
- (c) remove his name from the register of pharmacists and not to re-enter it thereon for such period as the Disciplinary Committee directs.

(2)職能

根據《藥劑業及毒藥條例》第15條, 管理局委出紀律委員會就下列情況召 開紀律研訊:

- (a) 當管理局接到有關某註冊藥劑師 或其僱員的行為操守的投訴,或 管理局覺得某藥劑師已違反適用 於該藥劑師的《行為守則》;
- (b) 當管理局接到有關某獲授權毒藥 銷售商、其僱員、高級人員或合 夥人的行為操守的投訴;或管理 局覺得某獲授權毒藥銷售商已 違反適用於該銷售商的《執業守 則》;
- (c) 當上述 (a) 或 (b) 項所述的任何人 士被裁定干犯:
 - (i) 《藥劑業及毒藥條例》、《危險 藥物條例》、《抗生素條例》或 《不良廣告(醫藥)條例》所訂 罪行;或
 - (ii) 《公 眾 衞 生 及 市 政 條 例》第 52、54 或 61 條或《商品説明條 例》第 7、7A 或 9 條所訂罪行;
- (d) 當管理局覺得根據《藥劑業及毒藥 條例》第13條就某獲授權毒藥銷 售商的處所的註冊而施加的某條 件,遭人違反;或
- (e) 當管理局在其他情況下,覺得有需要或適宜就任何在 (a) 或 (b) 段所述的人的行為操守進行研訊。

如研訊是就某註冊藥劑師或其僱員而 進行,紀律委員會可在研訊完結時:

- (a) 譴責該藥劑師;
- (b) 向該藥劑師發出警告信;或
- (c) 在紀律委員會指示的期間內,將 該藥劑師的姓名從藥劑師名冊中 刪除。

As for an ASP or an employee, officer or partner of an ASP, the Disciplinary Committee may at the conclusion of an inquiry direct that:

- (a) the ASP be disqualified, for such period as may be specified in the direction, from being an ASP;
- (b) any or all of the premises of that ASP be removed from the register of premises, either until the expiry of the certificate of registration issued to that ASP in respect of the premises, or for a shorter period as may be specified in the direction;
- (c) variations be made to the conditions relating to the registration of any or all of the premises of that ASP; or
- (d) a warning letter be served on that ASP.

At the conclusion of a disciplinary inquiry, the direction of the Disciplinary Committee against a registered pharmacist or an ASP takes effect immediately if the Disciplinary Committee considers it in the public interest to bring the direction into immediate effect. In other cases, the direction takes effect on the date specified by the Disciplinary Committee if no appeal has been lodged before the expiry of the period for lodging an appeal. If an appeal has been lodged, the direction takes effect on the date on which the appeal is finally determined.

The Disciplinary Committee may, subject to any conditions it thinks fit to impose, suspend for a period not exceeding three years (suspension period) the operation of a direction to remove a pharmacist's name from the register of pharmacists, disqualify a person from being an ASP, or remove any or all of the premises of an ASP from the register of premises so that the direction takes effect only if a condition so imposed is contravened during the suspension period.

The Disciplinary Committee may, subject to any appeal, cause its decision in any inquiry to be published in the Gazette, with or without an account of the proceedings. An appeal against any direction of the Disciplinary Committee shall be made, within 28 days after receipt of notice of direction, to the Court of First Instance.

Statistical figures on the results of disciplinary inquiries into the conduct of registered pharmacists and ASPs are given in **Tables 5 and 11** respectively. There was no appeal to the Court of First Instance in 2019.

至於獲授權毒藥銷售商或其僱員、高級人員或合夥人,紀律委員會可在研訊完結時作出下列指示:

- (a) 在某一指定的期間內,取消該銷售商的獲授權毒藥銷售商的資格;
- (b) 從處所註冊記錄冊中刪除該銷售 商的任何或全部處所,直至向該 銷售商發出的有關處所註冊證明 書的有效期屆滿,或為期一段在 該項指示指明較短的時間;
- (c) 更改該銷售商的任何或全部處所 的註冊條件;或
- (d) 向該獲授權毒藥銷售商送達警告信。 紀律委員會在研訊完結時,如認為其就 某註冊藥劑師或某獲授權毒藥銷售商 作出的指示即時生效是合乎公眾利益, 可指示即時生效;或在其他情況下如沒 有上訴在限期屆滿前提出,則於紀律委 員會指明的日期生效或如有上訴提出, 則於該上訴獲最終裁定的日期生效。

紀律委員會可在適宜施加的條件的規限下,暫緩執行其作出將某藥劑師的姓名從藥劑師名冊中除去、取消某或售商的獲授權毒藥銷售商的任何或內方。 將某獲授權毒藥銷售商的任何或內方。 處所註冊紀錄冊中除去的資格。 處所說冊紀錄冊中除去的,令到是 是有如此施加的條件在暫緩期內遭違 反,該指示才會生效。

如有關人士不提出上訴,紀律委員會 便可安排將其指令在憲報刊登,並可 刊登或不刊登有關研訊程序的報告。 有關人士欲就紀律委員會作出的指令 提出上訴,須於收到指示通知書的 二十八日內,向原訟法庭提出。

表 5 及 11 分別詳載有關註冊藥劑師 及獲授權毒藥銷售商的紀律研訊結果 的統計數字。二零一九年,原訟法庭 沒有收到任何上訴申請。

Membership and Functions of the Pharmacy and Poisons Appeal Tribunal

藥劑業及毒藥上訴審裁處的成員及職能

(1) Membership

The Tribunal consists of the following members who are appointed by the Chief Executive in accordance with section 30(2) of the Pharmacy and Poisons Ordinance:

- (a) a legally qualified person who shall be the chairman of the Tribunal;
- (b) a registered medical practitioner;
- (c) a registered pharmacist;
- (d) a person qualified in pharmacology;
- (e) a person nominated by the Director of Health from a panel consisting of persons nominated by pharmacists' associations;
- (f) a person nominated by the Director of Health from a panel consisting of persons nominated by pharmaceutical industry associations; and
- (g) a person nominated by the Director of Health from a panel consisting of persons nominated by the retail pharmaceutical trade associations.

The membership of the Tribunal as at 31 December 2019 was as follows:

Name	Membership
Mr WONG Kai-ming, Stewart, S.C.	Chairman
Dr TSANG Ho-fai, Thomas	Member
Dr LAM Ka-wing, Jenny	Member
Professor CHAN Yan-keung, Thomas, BBS, JP	Member
Miss CHU Man-wa	Panel Member
Miss LEUNG Sik-yin, McShirley	Panel Member
Miss TANG Choi-hung	Panel Member
Mr CHAN Cho-sun	Panel Member
Mr WONG Kwok-kai	Panel Member
Mr WOO Pui-hong, Christopher	Panel Member
Mr CHENG Kin-tung, Johnny	Panel Member
Mr PANG Hok-ming	Panel Member
Mr YEUNG Chi-fat	Panel Member

(1)成員

審裁處包括下列根據《藥劑業及毒藥條例》第30(2)條由行政長官委任的人士:

- (a) 一名具備法律專業資格的人士, 並由其出任審裁處主席;
- (b) 一名註冊醫生;
- (c) 一名註冊藥劑師;
- (d) 一名具備藥理學資格的人士;
- (e) 一名由藥劑師組織提名組成的小 組的成員,並為衞生署署長提名 的人士;
- (f) 一名由藥劑業組織提名組成的小 組的成員,並為衞生署署長提名 的人士;及
- (g) 一名由藥劑零售業組織提名組成 的小組的成員,並為衞生署署長 提名的人士。

在二零一九年十二月三十一日,審裁 處的成員如下:

姓名	成員
黃繼明先生,SC	主席
曾浩輝醫生	委員
林嘉穎博士	委員
陳恩強教授,BBS,JP	委員
朱敏華女士	小組委員
梁錫燕女士	小組委員
鄧彩紅女士	小組委員
陳祖新先生	小組委員
黄國佳先生	小組委員
胡培康先生	小組委員
鄭健東先生	小組委員
彭鶴鳴先生	小組委員
楊志發先生	小組委員

(2) Functions

The Pharmacy and Poisons Appeal Tribunal, established under section 30 of the Pharmacy and Poisons Ordinance, hears and determines the following matters:

- (a) any appeal against a decision of the Board in respect of application for registration of premises or application for renewal of registration of premises of an authorized seller of poisons;
- (b) any appeal against a direction of the Board in respect of suspension or removal of the name of a listed seller of poisons ("LSP") from the list of LSPs; and
- (c) any appeal against a decision of a committee of the Board, excluding the Disciplinary Committee.

No appeal was heard from 2015 to 2019.

(2)職能

藥劑業及毒藥上訴審裁處根據《藥劑業及毒藥條例》第30條成立,負責聆訊和裁定下列事宜:

- (a) 就管理局對獲授權毒藥銷售商的 處所註冊申請或處所註冊續期申 請的決定而提出的上訴;
- (b) 就管理局對從列載毒藥銷售商名 單中暫時吊銷或刪除列載毒藥銷 售商資格的決定而提出的上訴; 及
- (c) 就管理局屬下的委員會的決定提出的上訴,惟紀律委員會的決定 除外。

由二零一五年至二零一九年上訴審裁處沒有研訊上訴個案。

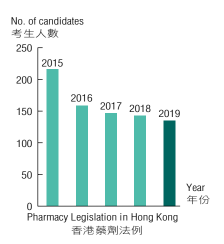
Table 表 1

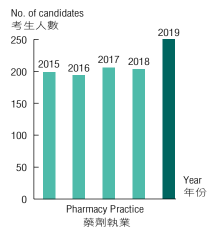
Results of the Registration Examinations 註冊考試成績

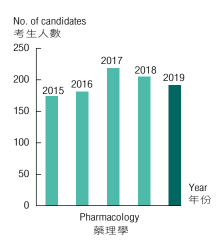
Year	_	Pharmacy Legislation in Hong Kong 香港藥劑法例			Pharmacy Practice Pharmacology 藥劑執業 藥理學			y	
年份	No. sat 參加人數	No. passed 合格人數	passing % 合格率	No. sat 參加人數	No. passed 合格人數	passing % 合格率	No. sat 參加人數	No. passed 合格人數	passing % 合格率
2015	216	87	40.3	199	57	28.6	174	38	21.8
2016	159	68	42.8	194	46	23.7	181	44	24.3
2017	147	71	48.3	206	67	32.5	219	55	25.1
2018	143	78	54.5	203	52	25.6	205	92	44.9
2019	135	77	57.0	250	82	32.8	192	71	37.0

Number of candidates sitting each examination subject

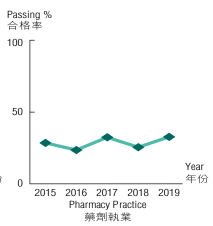
每科考試的考生人數







Passing percentage in each examination subject 每科考試的合格率



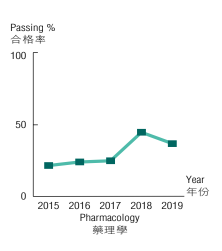


Table 表 2

Number of Registered Pharmacists in Hong Kong 香港註冊藥劑師人數

Year 年份	2015	2016	2017	2018	2019
Number of registered pharmacists as at end of year 截至年終的註冊藥劑師人數	2 504	2 659	2 753	2 890	3 001

Number of registered pharmacists as at end of year 截至年終的註冊藥劑師人數

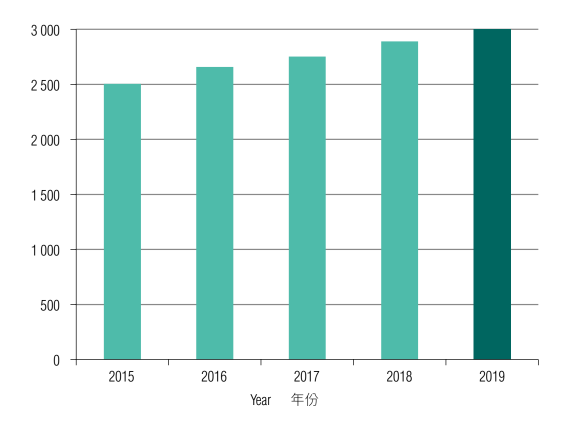


Table 表 3

Breakdown of Fresh Registration, Removal from the Register and Re-registration of Pharmacists

新註冊、刪除註冊及重新註冊的分項數字

Year 年份	2015	2016	2017	2018	2019
Fresh registration (Non-local graduates) 新註冊 (非本地畢業)	75	67	36	78	55
Fresh registration (Local graduates) 新註冊 (本地畢業)	51	98	79	80	75
Removal from the register* 刪除註冊 *	14	12	25	22	21
Re-registration 重新註冊	2	2	4	1	2
Net increase 淨增長	114	155	94	137	111

^{*} excluding orders by the Disciplinary Committee

不包括紀律委員會的指令

Table 表 4

Disciplinary Actions against Registered Pharmacists Taken by the Pharmacy and Poisons Board

藥劑業及毒藥管理局對註冊藥劑師採取的紀律行動

Disciplinary actions taken 採取的紀律行動	Number of cases 個案數目						
大中文ロッポレ1手1 J 里J	2015	2016	2017	2018	2019		
Formal disciplinary actions (i.e. Inquiry by Disciplinary Committee) 正式紀律行動 (即由紀律委員會進行紀律研訊)	1	1	0	0	1		

Table 表 5

Results of Disciplinary Inquiries into Registered Pharmacists 對註冊藥劑師進行紀律研訊的結果

Findings of the Disciplinary Committee	Number of cases 個案數目						
紀律委員會的裁決	2015	2016	2017	2018	2019		
Charge dismissed 指控不成立	0	1	0	0	0		
Guilty of the charge 指控成立	1	0	0	0	1		
Directions of the Disciplinary Committee 紀律委員會的指示							
Censure 譴責	0	0	0	0	1		
Removed from the register for a period of time 由名冊除名一段時間	1	0	0	0	0		

Table 表 6

Disciplinary Cases regarding Registered Pharmacists Handled by the Pharmacy and Poisons Board

藥劑業及毒藥管理局處理有關註冊藥劑師的紀律個案

Nature of offences*	Number of counts 次數					
個案性質	2015	2016	2017	2018	2019	
(1) Supplying Part 1 poison(s) from unlicensed premises 從沒有牌照的處所供應第 1 部毒藥	1	0	0	0	0	
(2) Failing to keep proper record of Part 1 poison(s) 沒有備存第 1 部毒藥的適當記錄	1	0	0	0	0	
(3) Possession of substance to which the Antibiotics Ordinance applies 管有《抗生素條例》適用的物質	1	0	0	0	0	
(4) Possession for sale or for any purpose of trade or manufacture goods to which a forged trade description was applied 管有應用虛假商品説明的貨品作售賣或任何商業或製造用途	1	0	0	0	0	
(5) Professional misconduct 專業失德	0	1	0	0	0	
(6) Selling Schedule 3 poison(s) without the authority of a prescription 在沒有處方授權的情況下出售附表 3 毒藥	0	0	0	0	1	
(7) Failing to keep proper record of antibiotics 沒有備存抗生素的適當記錄	0	0	0	0	1	

^{*} Some cases involve multiple nature of offences 部份個案涉及多個罪行

Table 表 7

Number of Authorized Sellers of Poisons in Hong Kong 香港的獲授權毒藥銷售商數目

Year 年份	2015	2016	2017	2018	2019
Number of authorized sellers of poisons as at end of year 截至年終的獲授權毒藥銷售商數目	607	604	614	641	649

Number of authorized sellers of poisons as at end of year 截至年終的獲授權毒藥銷售商數目

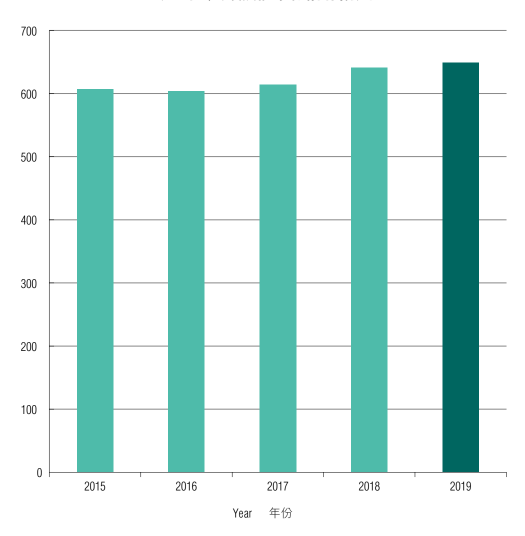


Table 表 8

Applications for Registration of Premises of Authorized Sellers of Poisons 獲授權毒藥銷售商的處所註冊申請

Year 年份	2015	2016	2017	2018	2019
Number of applications for registration of premises approved 接納處所註冊申請的數目	34	29	37	50	51
Number of applications for registration of premises rejected 拒絕處所註冊申請的數目	0	0	0	0	0
Number of applications for renewal of registration of premises rejected 拒絕處所註冊的續期申請數目	1	0	1	0	0

Table 表 9

Regulatory Control of Authorized Sellers of Poisons 獲授權毒藥銷售商的規管

Year 年份	2015	2016	2017	2018	2019
Number of inspections conducted 巡查數目	1 214	1 209	1 220	1 212	1 305
Number of test purchases conducted 試買數目	4 136	3 955	4 329	4 194	4 101

Table 表 10

Disciplinary Actions against Authorized Sellers of Poisons Taken by the Pharmacy and Poisons Board

藥劑業及毒藥管理局對獲授權毒藥銷售商採取的紀律行動

Disciplinary actions taken		Number of cases 個案數目					
採取的紀律行動	2015	2016	2017	2018	2019		
Formal disciplinary actions (i.e. Inquiry by Disciplinary Committee) 正式紀律行動 (即由紀律委員會進行紀律研訊)	13	12	9	11	11		
Informal disciplinary actions (i.e. Verbal caution by representatives of the Board) 非正式紀律行動 (即由管理局代表給予口頭警告)	18	18	8	5	6		
The authorized seller of poisons ceased operation before action taken by the Board 該銷售商在管理局採取紀律行動前已經結業	1	1	0	1	1		
Total 總數	32	31	17	17	18		

Table 表 11

Results of Disciplinary Inquiries into Authorized Sellers of Poisons 對獲授權毒藥銷售商進行紀律研訊的結果

Findings of the Disciplinary Committee	Number of cases 個案數目								
紀律委員會的裁決	2015	2016	2017	2018	2019				
Charge dismissed 指控不成立	0	0	0	0	0				
Guilty of the charge 指控成立	13	12	9	11 *	11				
Directions of the Disciplinary Committee 紀律委員會的指示									
Issue of written warning 發出書面警告	4	4	0	3	6				
Disqualified from being an authorized seller of poisons for a period of time 取消銷售商資格一段時間	9	8	9	7	5				

^{*} The decision of the Disciplinary Committee against one authorized seller of poisons was reversed by the Court of First Instance. 原訟法庭推翻了紀律委員會對一名獲授權毒藥銷售商的裁決。

Table 表 12

Disciplinary Inquiries into Authorized Sellers of Poisons Handled by the Pharmacy and Poisons Board

藥劑業及毒藥管理局處理有關獲授權毒藥銷售商的紀律研訊個案

Nature of offences*		Number of counts (percentage) 次數 (百份比)					
個案性質	2015	2016	2017	2018	2019		
(1) Sale of Part 1/Part 2 poison(s) without label/proper label 銷售沒有標籤/沒有妥善標籤的第1部 第2部毒藥	龙 4 (10%)	0 (0%)	0 (0%)	1 (4.17%)	0 (0%)		
(2) Sale of Part 1 poison(s) without the supervision of registered pharmacist / proper supervision 在沒有註冊藥劑師監督 / 適當監督的情況下銷售第 1 部毒藥	5	4 (12.9%)	6 (20.69%)	4 (16.66%)	5 (10%)		
(3) Sale of Schedule 3 poison(s) without the authority a prescription 在沒有處方授權的情況下出售附表 3 毒藥	of 9 (22.5%)	5 (16.13%)	7 (24.13%)	3 (12.5%)	12 (24%)		
(4) Sale of antibiotics without the authority of a prescription 在沒有處方授權的情況下銷售抗生素	0 (0%)	0 (0%)	2 (6.9%)	0 (0%)	5 (10%)		
(5) Possession of poison(s) included in Part 1 of the Poisons List 管有毒藥表第 1 部所列任何毒藥	1 (2.5%)	3 (9.68%)	0 (0%)	1 (4.17%)	0 (0%)		
(6) Possession of unregistered pharmaceutical product(s) 管有未經註冊藥劑製品	4 (10%)	0 (0%)	3 (10.34%)	3 (12.5%)	6 (12%)		
(7) Failing to store Schedule 1 poison(s) in a receptacle sole for that purpose 沒有將附表 1 的毒藥貯存於專門的盛器內	1 (2.5%)	0 (0%)	0 (0%)	0 (0%)	0 (0%)		
(8) Possession for sale or for any purpose of trade manufacture goods to which a false trade description was applied 為售賣或任何商業或製造用途而管有應用虛作商品説明的貨品	on 2 (5%)	3 (9.68%)	0 (0%)	1 (4.17%)	0 (0%)		
(9) Possession for sale or for any purpose of trade manufacture goods to which a forged trade mawas applied 為售賣或任何商業或製造用途而管有應用偽資商標的貨品	·k 2 (5%)	1 (3.23%)	2 (6.9%)	3 (12.5%)	2 (4%)		
(10) Selling substance(s) to which the Antibiotics Ordinand applies 售賣《抗生素條例》適用的物質	0 (0%)	4 (12.9%)	0 (0%)	0 (0%)	0 (0%)		

Table 表 12 (Con't) 續

Nature of offences* 個案性質		Number of counts (percentage) 次數 (百份比)						
	2015	2016	2017	2018	2019			
(11) Failing to store poison(s) properly	9	2	2	4	3			
沒有適當地貯存毒藥	(22.5%)	(6.45%)	(6.9%)	(16.66%)	(6%)			
(12) Illegal sale of unregistered pharmaceutical product(s)	2	0	0	1	2			
非法銷售未經註冊的藥劑製品	(5%)	(0%)	(0%)	(4.17%)	(4%)			
(13) Supplying/Offering to supply false trade description goods	0	3	0	0	0			
供應 / 要約供應虛假商品説明的貨品	(0%)	(9.68%)	(0%)	(0%)	(0%)			
(14) Failing to seek prior approval from the Pharmacy and Poisons Board for change in the ownership/directorship or person-in-charge of the authorized seller of poisons 獲授權毒藥銷售商未事先獲得藥劑業及毒藥管理局的批准,更改了擁有人/董事或負責人	1 (2.5%)	0 (0%)	0 (0%)	0 (0%)	1 (2%)			
(15) Selling goods to which a forged trade mark was applied	0	6	3	2	12			
出售應用偽造商標的貨品	(0%)	(19.35%)	(10.34%)	(8.33%)	(24%)			
(16) Failing to ensure that all the keys of the lockable receptacle in the dispensing area were kept by the registered pharmacist 未能確保配藥室可上鎖盛器的所有鎖匙由註冊藥劑師保管	0 (0%)	0 (0%)	1 (3.45%)	0 (0%)	0 (0%)			
(17) Failing to provide written order(s) in relation to purchasing	0	0	4	0	0			
of controlled medicines 未能提供有關訂購受管制藥物的書面訂單	(0%)	(0%)	(3.45%)	(0%)	(0%)			
(18) Failing to provide sales invoice(s) for controlled medicines	0	0	1	0	0			
未能提供有關受管制藥物的銷售發票	(0%)	(0%)	(3.45%)	(0%)	(0%)			
(19) Failing to keep proper record of psychotropic substances in psychotropic substances book 未能在精神藥物記錄冊內保存精神藥物有關的妥善記錄	0 (0%)	0 (0%)	1 (3.45%)	0 (0%)	0 (0%)			
(20) Trafficking in dangerous drug	0	0	0	1	0			
版運危險藥物	(0%)	(0%)	(0%)	(4.17%)	(0%)			
(21) Failing to maintain proper record of antibiotics	0	0	0	0	2			
沒有備存合適的抗生素紀錄	(0%)	(0%)	(0%)	(0%)	(4%)			

^{*} Some cases involve multiple nature of offences 部份個案涉及多個罪行

Table 表 12A

Disciplinary Inquiries into Authorized Sellers of Poisons in 2019 2019 年有關獲授權毒藥銷售商的紀律研訊個案

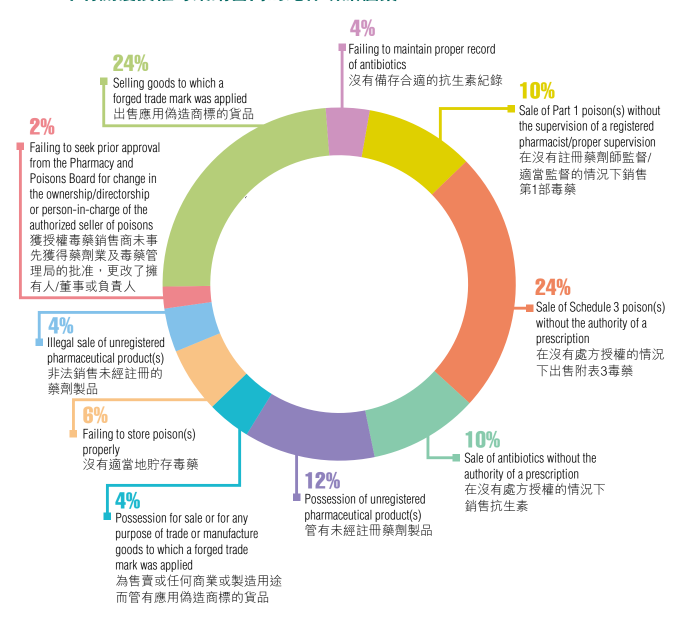


Table 表 13

Number of Listed Sellers of Poisons in Hong Kong 香港的列載毒藥銷售商數目

Year 年份	2015	2016	2017	2018	2019
Number of listed sellers of poisons as at end of year 截至年終的列載毒藥銷售商數目	4 012	3 937	3 937	3 937	4 295

Number of listed sellers of poisons as at end of year 截至年終的列載毒藥銷售商數目

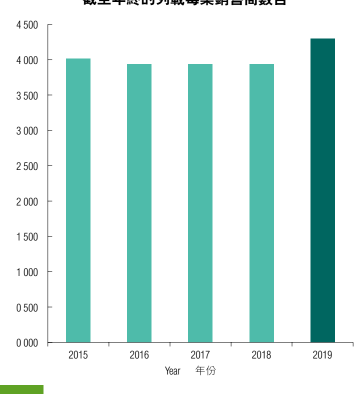


Table 表 14

Applications for Licensing as Listed Sellers of Poisons 申請列載毒藥銷售商牌照

Year 年份	2015	2016	2017	2018	2019
Number of applications approved 接納牌照申請數目	277	231	258	264	624
Number of applications rejected 拒絕牌照申請數目	2	2	0	0	1

Table 表 15

Regulatory Control of Listed Sellers of Poisons 列載毒藥銷售商的規管

Year 年份	2015	2016	2017	2018	2019
Number of inspections conducted 巡查數目	7 977	7 956	7 874	7 814	8 323
Number of test purchases conducted 試買數目	3 008	4 021	3 229	3 350	3 353

Table 表 16

Disciplinary Actions against Listed Sellers of Poisons Taken by the Pharmacy and Poisons Board

藥劑業及毒藥管理局對列載毒藥銷售商採取的紀律行動

Disciplinary actions taken 採取的紀律行動	Number of cases 個案數目						
採取的紀律行動	2015	2016	2017	2018	2019		
Removal from the list of listed sellers of poisons 從列載毒藥銷售商名單除名	4	4	0	0	1		
Issue of written warning 發出書面警告	6	3	4	2	5		
Suspension of name from the list of listed sellers of poisons for a specified period of time 暫時吊銷名列列載毒藥銷售商名單內的資格一段時間	0	3	9	4	6		
The listed seller of poisons ceased operation before action taken by the Board 該銷售商在管理局採取紀律行動前已經結業	1	0	0	0	0		
Total 總數	11	10	13	6	12		

Table 表 17

Disciplinary Cases regarding Listed Sellers of Poisons Handled by the Pharmacy and Poisons Board

藥劑業及毒藥管理局處理有關列載毒藥銷售商的紀律個案

Nature of offences *	Number of counts (percentage) 次數 (百份比)						
個案性質	2015	2016	2017	2018	2019		
(1) Sale of Part 1 poison(s) without the supervision of a registered pharmacist/proper supervision 在沒有註冊藥劑師監督 / 適當監督的情況下銷售第 1 部毒藥	1 (5.26%)	4 (10.81%)	1 (3.45%)	1 (8.33%)	2 (7.14%)		
(2) Sale of Schedule 3 poison(s) without the authority of a prescription 在沒有處方授權的情況下出售 附表 3 毒藥	0 (0%)	3 (8.11%)	1 (3.45%)	0 (0%)	2 (7.14%)		
(3) Possession of Part 1 poison(s) 管有第 1 部毒藥	4 (21.05%)	6 (16.22%)	8 (27.58%)	4 (33.34%)	3 (10.72%)		
(4) Possession of unregistered pharmaceutical product(s) 管有未經註冊藥劑製品	0 (0%)	1 (2.7%)	4 (13.79%)	4 (33.34%)	6 (21.43%)		
(5) Possession of substance(s) to which the Antibiotics Ordinance applies 管有《抗生素條例》適用的物質	0 (0%)	1 (2.7%)	3 (10.34%)	1 (8.33%)	2 (7.14%)		
(6) Possession for sale or for any purpose of trade or manufacture goods to which a forged trade mark was applied 為售賣或任何商業或製造用途而管有應用偽造商標的貨品	3 (15.79%)	8 (21.63%)	1 (3.45%)	0 (0%)	2 (7.14%)		
(7) Possession for sale or for any purpose of trade or manufacture goods to which a false trade description was applied 為售賣或任何商業或製造用途而管有應用虛假商品説明的貨品	2 (10.53%)	0 (0%)	1 (3.45%)	0 (0%)	0 (0%)		

Table 表 17(Con't) 續

Nature of offences *	Number of counts (percentage) 次數 (百份比)						
個案性質	2015	2016	2017	2018	2019		
(8) Sale of unregistered pharmaceutical product(s) 售賣未經註冊藥劑藥品	2	0	1	0	2		
	(10.53%)	(0%)	(3.45%)	(0%)	(7.14%)		
(9) Selling goods to which a forged trade mark was applied 售賣應用偽造商標的貨品	1 (5.26%)	7 (18.92%)	1 (3.45%)	0 (0%)	4 (14.3%)		
(10) Applying a false trade description on goods in the course of any trade or business	3	1	1	0	1		
在營商過程或業務運作中將虛假商品説明應用於貨品	(15.79%)	(2.7%)	(3.45%)	(0%)	(3.57%)		
(11) Unlawful sale of Part 2 poison(s)	3	4	4	0	2		
非法銷售第 2 部毒藥	(15.79%)	(10.81%)	(13.79%)	(0%)	(7.14%)		
(12) Illegal use of restricted title	0 (0%)	1	0	0	1		
非法使用名銜		(2.7%)	(0%)	(0%)	(3.57%)		
(13) Illegal use of restricted logo	0 (0%)	1	1	0	1		
非法展示標籤		(2.7%)	(3.45%)	(0%)	(3.57%)		
(14) Sale of Part 1 poison(s)	0 (0%)	0	1	0	0		
銷售第 1 部毒藥		(0%)	(3.45%)	(0%)	(0%)		
(15) Sale of Schedule 1 Chinese herbal medicine without prescription 沒有按照處方銷售附表 1 中藥材	0 (0%)	0 (0%)	1 (3.45%)	0 (0%)	0 (0%)		
(16) Failing to store poison(s) properly 沒有適當地貯存毒藥	0 (0%)	0 (0%)	0 (0%)	1 (8.33%)	0 (0%)		
(17) Engaging in relation to a consumer in a commercial practice that is a misleading omission 就消費者作出屬誤導性遺漏的營業行為	0 (0%)	0 (0%)	0 (0%)	1 (8.33%)	0 (0%)		

^{*} Some cases involve multiple nature of offences 部份個案涉及多個罪行

Table 表 17A

Disciplinary Cases regarding Listed Sellers of Poisons in 2019 2019 年有關列載毒藥銷售商的紀律個案

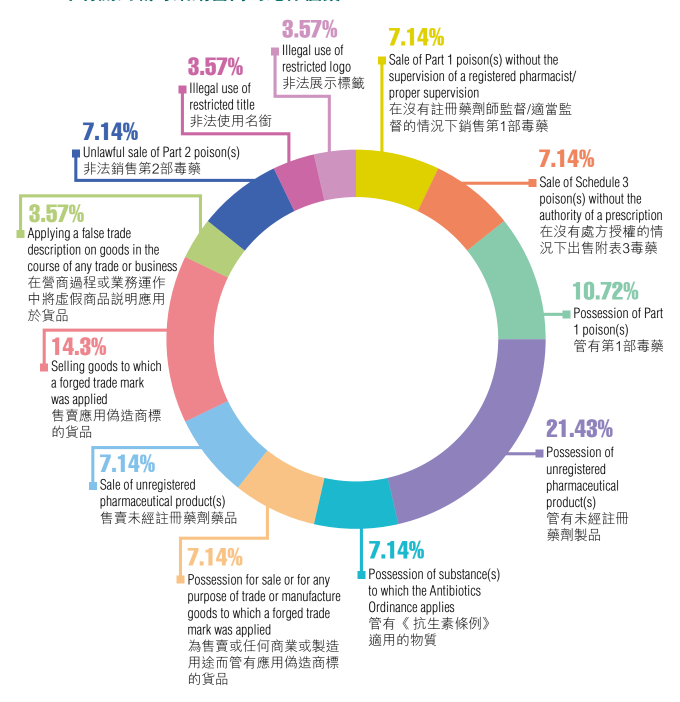


Table 表 18

Issue of Wholesale Dealer Licences 批發商牌照的簽發

Year 年份	2015	2016	2017	2018	2019
Number of holders of wholesale dealer licences as at end of year 截至年終的批發商牌照持有人的數目	797	779	773	770	770
Number of wholesale dealer licences revoked/ suspended 撤銷或吊銷批發商牌照的數目	1	1	3	1	2

Note: With effect from 6 February 2015, the wholesale dealer licence was introduced to replace the wholesale poisons licence and certificate of registration as an importer/exporter of pharmaceutical products. Holders of valid wholesale poisons licence or certificate of registration of importers and exporters are regarded as licensed wholesale dealers until the expiry of their licences or certificates.

註 :由 2015 年 2 月 6 日起,批發商牌照已推出,以取代毒藥批發牌照及進/出口商註冊證明書。持有有效期毒藥批發 牌照或進口商及出口商註冊證明書的人士,在其牌照或註冊證明書失效前,將會被視為持牌批發商。

Table 表 19

Issue of Manufacturer's Licences for Pharmaceutical Products藥劑製品製造商牌照的簽發

Year 年份	2015	2016	2017	2018	2019
Number of holders of manufacturer's licences as at end of year 截至年終的製造商牌照持有人的數目	80	72	72	71	70
Number of holders of manufacturer's licences only authorized to conduct secondary packaging of pharmaceutical products 截至年終只獲授權從事藥劑製品外包裝操作的製造商牌照持有人數目	57	49	48	48	47
Number of manufacturer's licences revoked/ suspended 撤銷或吊銷製造商牌照的數目	0	0	0	0	0

Note: With effect from 1 October 2015, all licensed manufacturers were required to fully comply with the PIC/S GMP.

註 :由 2015年10月1日起,所有持牌製造商必須完全符合國際醫藥品稽查協約組織的生產質量管理規範指引。

Table 表 20

Registration of Pharmaceutical Products

藥劑製品的註冊

Year 年份	2015	2016	2017	2018	2019
Number of registered pharmaceutical products as at end of year 截至年終的註冊藥劑製品數目	19 486	18 584	18 120	17 323	16 186

Number of registered pharmaceutical products as at end of year 截至年終的註冊藥劑製品數目

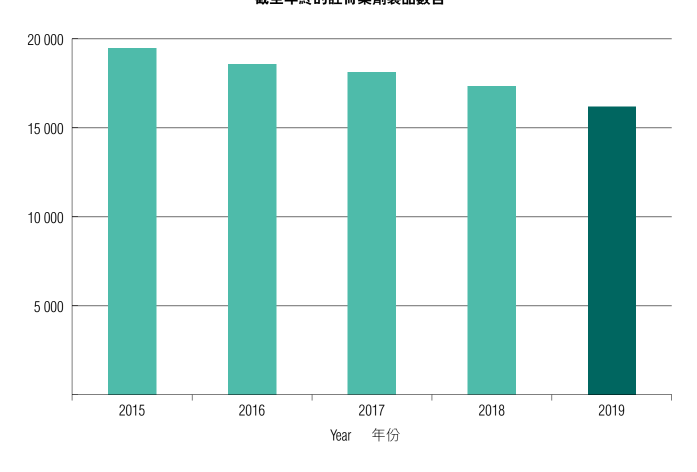


Table 表 21

Amendments to Schedules 1, 3 and 10 to the Pharmacy and Poisons Regulations in 2019 2019 年在《藥劑業及毒藥規例》附表 1、3 及 10 作出的修訂

New Substances Added 加入的新物質						
1.	Abemaciclib; its salts	阿貝西利; 其鹽類				
2.	Baloxavir; its salts; its esters and ethers; their salts	Baloxavir:其鹽類;其酯類及醚類; 它們的鹽類				
3.	Brexpiprazole; its salts	布瑞哌唑;其鹽類				
4.	Brigatinib; its salts	布格替尼;其鹽類				
5.	Cannabidiol; its salts; when contained in pharmaceutical products	大麻二酚;其鹽類; 但限於包含在藥劑製品內者				
6.	Dacomitinib; its salts	Dacomitinib;其鹽類				
7.	Efinaconazole; its salts	艾非康唑;其鹽類				
8.	Add sub-item "Epoprostenol" in the item "Prostaglandins, the following and their derivatives"	於"前列腺素類如下及它們的衍生物" 項目中加入"依前列醇"分項				
9.	Erenumab	依瑞奈人單抗				
10.	Ertugliflozin; its salts	艾托格列淨;其鹽類				
11.	Gemtuzumab ozogamicin	奧加米星吉妥組單抗				
12.	Add sub-item"Indigo carmine" in the item "Pharmaceutical products for human parenteral administration containing the following or their salts, as active ingredients, except in mixture with insulin"	於"供注射入人體的藥劑製品,並包含 (作為有效成分)以下物質或它們的鹽類, 但與胰島素的混合物除外"項目中加入 "Indigo carmine"分項				
13.	Isavuconazole; its salts; its derivatives; their salts	艾沙康唑;其鹽類;其衍生物;它們的鹽類				
14.	Larotrectinib; its salts	拉羅替尼;其鹽類				
15.	Latanoprostene bunod; its salts	拉坦前列烯酯;其鹽類				
16.	Letermovir; its salts; its esters; their salts	來特莫韋;其鹽類;其酯類;它們的鹽類				
17.	Lorlatinib; its salts	洛拉替尼; 其鹽類				
18.	Neratinib; its salts	奈拉替尼; 其鹽類				
19.	Nicardipine; its salts	尼卡地平;其鹽類				
*20.	Nifuratel; its salts	硝呋太爾;其鹽類				

Table 表 21 (Con't) 續

*21.	Nifuroxazide; its salts	硝呋齊特;其鹽類			
*22.	Nitrofural; its salts	呋喃西林;其鹽類			
*23.	Nitrofurantoin; its salts	呋喃妥因;其鹽類			
*24.	Nitroxoline; its salts	硝羥喹啉;其鹽類			
25.	Omega-3 fatty acids; their salts; their esters; when contained in pharmaceutical products intended to be used for the treatment of hypertriglyceridaemia	奧米加-3 脂肪酸;其鹽類;其酯類;但限於包含在擬用於治療高甘油三酯血症的藥劑製品內者			
[#] 26.	Pentoxifylline; its salts	己酮可可鹼;其鹽類			
*27.	Piracetam; its salts	吡拉西坦;其鹽類			
28.	Risankizumab	利生奇組單抗			
29.	Semaglutide	司美魯肽			
30.	Sodium zirconium cyclosilicate	Sodium zirconium cyclosilicate			
31.	Sugammadex; its salts; its esters; their salts	舒更葡糖;其鹽類;其酯類;它們的鹽類			
Others 其他					
1.	Item "Etidronic acid; its salts", after "salts" - Add "; when contained in pharmaceutical products"	"羥乙磷酸;其鹽類"項目,在"類"之後- 加入";但限於包含在藥劑製品內者"			
*2.	Item relating to "Pharmaceutical products for human parenteral administration" - Repeal "Piracetam"	關乎 "供注射入人體的藥劑製品" 的項目 - 廢除 "吡拉西坦"			
#3.	Item "Pharmaceutical products for human parenteral administration containing the following or their salts, as active ingredients, except in mixture with insulin" - Repeal sub-item "Pentoxifylline"	"供注射入人體的藥劑製品,並包含(作為有效成分)以下物質或它們的鹽類,但在與胰島素的混合物內者除外"項目 - 廢除 "己酮可哥堿"分項			
4.	Chinese text, item "供注射入人體的藥劑製品,並包含(作為有效成分)以下物質或它們的鹽類,但與胰島素的混合物除外"- Repeal "與胰島素的混合物" Substitute "在與胰島素的混合物內者"	中文文本"供注射入人體的藥劑製品·並包含(作為有效成分)以下物質或它們的鹽類·但與胰島素的混合物除外"項目- 廢除 "與胰島素的混合物" 代以 "在與胰島素的混合物內者"			

^{*} These amendments will come into operation on 18 October 2020. 這些修訂自 2020 年 10 月 18 日起實施。

[#] These amendments will come into operation on 13 December 2020. 這些修訂自 2020 年 12 月 13 日起實施。

