Pharmacy and Poisons Board of Hong Kong Pharmacy and Poisons Ordinance (Cap. 138)

Guidelines for Application for Change of Particulars of
Wholesale Dealer Licence (Cap. 138 Pharmacy and Poisons Ordinance)/
Antibiotics Permit (Cap. 137 Antibiotics Ordinance)/
Wholesale Dealer's Licence to Supply Dangerous Drugs (Cap. 134 Dangerous Drugs Ordinance)

Should any Wholesale Dealer Licence/ Antibiotic Permit/ Wholesale Dealer's Permit holder wish to apply change of any particular(s); they shall submit application by writing to the Pharmacy and Poisons (Wholesale Licences) Committee (hereafter as 'the Committee') and/or Drug Office Licensing and Compliance Division Wholesalers Regulatory Unit (hereafter as 'Wholesalers Regulatory Unit') well in advance. The applied change of particulars shall be valid upon the Wholesale Dealer Licence holder obtained approval from 'the Committee' and/or 'Wholesalers Regulatory Unit'.

The licence holder must maintain the business of wholesale and storage of Poisons/Pharmaceutical Products according to the approved terms and condition under the respective licence(s) or 'permit' until further applied changes approved by the 'the Committee' and/or 'Wholesalers Regulatory Unit'. Under "Cap. 138 Pharmacy and Poisons Ordinance", 'the Committee' may revoke a Wholesale Dealer Licence or suspend it for a period it thinks fit, issue a warning letter, or vary a condition of the licence, if, in the Committee's opinion, the licensed wholesale dealer has contravened a condition of the licence or any of the regulations provided by the "Pharmacy and Poisons Ordinance" or "Antibiotic Ordinance" or "Dangerous Drugs Ordinance" Regulations, a "Code of Practice for Holder of Wholesale Dealer Licence", and/or has been convicted of a drug-related offence.

I. Application requirements

- 1. The applicant must be the licence holder (the holder's proprietor/ partner(s)/ director(s), person in charge of poisons and pharmaceutical products (hereafter as 'PIC of PP/Poisons') or deputy person in charge of poisons and pharmaceutical products (hereafter as 'DPIC of PP/Poisons'). If it is necessary to appoint an authorized person to handle the application, please attach an authorization letter signed by the license holder (refer to Appendix 12);and
- 2. The new applied change of particulars shall comply with the licensing requirements.
- 3. General requirements for personnel:
 - The licence holder shall notify 'the Committee' in writing of any change in its proprietor, partner(s) or director(s) within one month from the date of change.
 - The licence holder shall obtain approval from 'the Committee' and/or 'Wholesalers Regulatory Unit' prior to any change of 'PIC of PP/Poisons', 'DPIC of PP/Poisons person and/or 'PIC of Dangerous Drugs' and 'the Committee' and/or 'Wholesalers Regulatory Unit' shall not approve the change unless it considers the person nominated fit and proper.
 - Applicant must nominate a person-in-charge of poisons and pharmaceutical products ("PIC"), whom will be subjected to approval by the Pharmacy and Poisons (Wholesale Licences) Committee ("the Committee"). The nominated person must be a fit and proper person and also possess adequate knowledge to carry on trade related to the pharmaceutical industry. The nomination of a nominated person who is already a PIC for another holder of Wholesale Dealer Licence would normally not be considered.
- 4. General requirements for premises:
 - Only companies occupying commercial premises or industrial buildings would be considered;
 - Companies occupying ground floor or retail premises would normally not be considered;
 - Companies operating in secretarial or accountancy service holding companies would not be considered;
 - Companies sharing premises with another holder of Wholesale Dealer Licence would require a written explanation¹; and
 - If there is no storage facility within the business premises, the company must maintain adequate lockable storage facilities at another premises, and provide a written explanation on why storage facility cannot be provided within the business address of the premises.
- 5. There must be adequate lockable storage facilities with appropriate temperature and humidity for keeping antibiotics/ poisons/ dangerous drugs/ pharmaceutical products within the premises. If there is no storage facility within the premises, the company must maintain adequate lockable storage facilities at another premises, and provide a written explanation on why storage facility cannot be provided within the business address of the premises, provide details of the store, routine maintenance and monitoring. Application with storage facilities outside the premises are subjected to consideration and approval by 'the Committee' on a case by case basis. If the application involved handling of Part I Dangerous Drugs, lockable receptacle designated for storage of Part I Dangerous Drugs must be made available. Detailed requirements on the storage facilities are set out in the "Code of Practice for Holder of Wholesale Dealer Licence".

¹ The written explanation must be supported by relevant and sufficient reasons to the satisfaction of the Pharmacy and Poisons (Wholesale Licence) Committee. Each case will be considered on a case-by-case basis and at the discretion of the Committee.

II. Application procedures

How to obtain application forms

1. Application Form for Change of Particulars for Wholesale Dealer Licence/ Antibiotics Permit/ Wholesale Dealer's Licence to Supply Dangerous Drugs (hereafter as 'COP Application Form') can be obtained free of charge from:

Licensing and Compliance Division, Drug Office, Department of Health, Room 2001-2002, 20/F., Dah Sing Financial Centre

248 Queen's Road East, Wan Chai, Hong Kong Monday to Friday
9:00 a.m. to 1:00 p.m.
2:00 p.m. to 5:45 p.m.
(up to 6:00 p.m. on Monday)
(Closed on Saturdays, Sundays
& Public Holidays)

2. 'COP Application Form' can also be download from the Drug Office official website: (https://www.drugoffice.gov.hk/eps/do/en/pharmaceutical_trade/guidelines_forms/useful_guidelines_forms.html)

Submission of documents or information

Applicants are required to submit the following information:

1. A fully completed 'COP Application form'; and

- 2. Supporting documents in relation to the change of particulars. It is unnecessary to submit repeated supporting document(s) for different particular(s) of change; and
- 3. <u>If the application only involves license cancellation, certified copy and/or license refund</u>, the applicant only needs to complete the relevant appendix.
- 4. Applicant(s) may be required to submit original(s) with his/her signature and company chop for their supporting document(s).

How to submit application

Applicants may submit the application forms, the relevant information and documents via the following ways:

- (i) Mail to Licensing and Compliance Division, Drug Office, Department of Health by post or registered mail (the date shown on the post stamp will be taken as the submission date); or
- (ii) Lodge to the Licensing and Compliance Division, Drug Office, Department of Health in person during office hours.

III. Application results

If the change application <u>involved revise the terms and conditions on licence(s) and/or permit(s)</u>, the applicant will receive a demand note for payment of update of license. Upon the receipt of the prescribed fee, the applicant will be informed to present the original licence in person or by a representative on his/her behalf, to the 'Wholesalers Regulatory Unit' to complete necessary procedures; If the change application do not involved revise the terms and conditions on licence(s) and/or permit(s), the applicant will receive a written notification by 'Wholesalers Regulatory Unit' on behalf of 'the Committee' if the application is approved. If the application is rejected or required further revise that the applicant will still be notified by email or via phone call.

IV. Prescribed fee and methods of payment

The fee for change of particulars application per licence is HK\$155. The Licensing and Compliance Division, Drug Office of the Department of Health will issue a General Demand Note to the applicant. The applicant could make payment according to the payment methods stated in the General Demand Note.

V. Enquiries

Further enquiries regarding the change of particulars as specified in the licence(s) and/or permit(s) or on the content of these guidelines can be made by calling the enquiry hotline, email or post to the 'Wholesalers Regulatory Unit':

Enquiry Hotline: 3107 2194

Enquiry Email: enquirywru@dh.gov.hk

Address: Room 2001-2002, 20/F., Dah Sing Financial Centre 248 Queen's Road East, Wan Chai, Hong

Kong

VI. Notes

Applicants and their employees or agents must not offer an advantage as defined in the Prevention of Bribery Ordinance (Cap. 201) to any government officer or members of statutory organisations (including but not limited to the Pharmacy and Poisons Board and its Committees) in connection with their applications or while having dealings of any kind with government departments or statutory organisations.

<u>Checklist for Change of Particulars of</u> <u>Wholesale Dealer Licence (Cap. 138 Pharmacy and Poisons Ordinance)/</u> Antibiotics Permit (Cap. 137 Antibiotics Ordinance)/

Wholesale Dealer's Licence to Supply Dangerous Drugs (Cap. 134 Dangerous Drugs Ordinance)

Content of Change of Particulars Checklist:

No.	Change of Particulars Details	Submission of Supporting Documents (Refer to Page 6-23)
Com	pany Information	
A	(i) Change of Company Name (Incorporated	'COP Form' + 'COP Checklist Details' (1.a), (2.a), (2.b)
	Company Only)	
	(ii) Change of Company Name (Partnership	'COP Form' + 'COP Checklist Details' (1.a), (2.c)
	Company Only)	
	(iii) Change of Company Name (Sole	'COP Form' + 'COP Checklist Details' (1.a), (2.e)
	Proprietorship Company Only)	
Perso		
В	(i) Change or Addition of Director(s)	'COP Form' + 'COP Checklist Details' (3), (4), (5), (6), (9), (10)
_	(ii) Deletion of Director(s)	'COP Form' + 'COP Checklist Details' (3), (4), (5)
C	(i) Change of Partner(s)	'COP Form' + 'COP Checklist Details' (2.c), (5), (6), (9), (10)
	(ii) Deletion of Partner(s)	'COP Form' + 'COP Checklist Details' (2.c), (5)
D	Change of Sole Proprietor	'COP Form' + 'COP Checklist Details' (2.e), (5), (6), (9), (10)
E	Change of PIC of PP/Poisons	'COP Form' + 'COP Checklist Details' (6), (9), (10)
F	(i) Change or Addition of DPIC of PP/Poisons	'COP Form' + 'COP Checklist Details' (6), (9), (10)
	(ii) Deletion of DPIC of PP/Poisons	'COP Form'
\mathbf{G}	(i) Change or Addition of PIC of Dangerous	'COP Form' + 'COP Checklist Details' (7), (11)
	Drugs Pt. I	(GOD F
	(ii) Deletion of Addition PIC of Dangerous	'COP Form'
TT	Drugs Pt. I #	(CORE 1: (CORCL 11: (R + 11.1(C) (0) (10)
H	(i) Change or Addition of PIC of Dangerous	'COP Form' + 'COP Checklist Details' (6), (9), (10)
	Drugs Pt. II	'COP Form'
	(ii) Deletion of Addition PIC of Dangerous Drugs Pt. II #	*COP Form*
I	Addition of Locum Pharmacist to handle	'COP Form' + 'COP Checklist Details' (8), (11)
I	"Dangerous Drugs Pt. I"	COF FORM + COF CHECKHST Details (8), (11)
Addr	ess / Storage **	
J	(i) Change of Premises Address ⁶ (Room	'COP Form' + 'COP Checklist Details' (1.a), (13.a), (14.b),
J	Temperature Storage)	$(15.b), (16)^{\land}, (17)^{\land}, (18)^{\land}, (19)^{\land}$
	(ii) Change of Premises Address ⁶ (Room	'COP Form' + 'COP Checklist Details' (1.a), (13.a), (14.b),
	Temperature + Cold Chain Storage)	(15.b), (16)^, (17)^, (18)^, (19)^, (20)^
K	Updates of Layout within Approved Premises	Storage at Premises Address ⁶ : 'COP Form' + 'COP Checklist
	Address ⁶ /Additional Warehouse ⁷ with Storage	Details' (14.a), (14.b)
	Area unchanged	Storage at Additional Warehouse ⁷ : 'COP Form' + 'COP
		Checklist Details' (14.c), (14.d)
L	(i) Within Approved Premises Address ⁶ /	Storage at Premises Address ⁶ : 'COP Form' + 'COP Checklist
M	Additional Warehouse ⁷ :	Details' (14.a), (14.b), (15.a), (15.b), (16)\(^\), (17)\(^\), (18)\(^\), (19)\(^\)
0		Storage at Additional Warehouse ⁷ : 'COP Form' + 'COP
	- Change of Storage Room/Facilities Location,	Checklist Details' (14.c), (14.d), (15.a), (15.b), (16)\(^\), (17)\(^\),
	Layout, Shape or Size (Room Temperature	(18)^, (19)^
	Storage)	
	(ii) Within Approved Premises Address ⁶ /	Storage at Premises Address ⁶ : 'COP Form' + 'COP Checklist
	Additional Warehouse ⁷ :- Change or Addition of	Details' (14.a), (14.b), (15.a), (15.b), (16)\hat^, (17)\hat^, (18)\hat^, (19)\hat^,
	Store Room/Facilities or;	(20)^
	- Change of Storage Room/Facilities Location,	Storage at Additional Warehouse ⁷ : 'COP Form' + 'COP
	Layout, Shape or Size (Room Temperature	Checklist Details' (14.c), (14.d), (15.a), (15.b), (16)^, (17)^,
	Storage + Cold Chain Storage)	(18)^, (19)^, (20)^

#(Should maintain at least 1 DD PIC)

%(Should maintain at least 1 storage facility)

² MD: This licence only authorizes the holder to deal in medical devices containing poisons.

^{^(}Not applicable for Wholesale Dealer Licence with 'NM'¹, 'MD'² or 'NT'³ condition)

¹ NM: This licence only authorizes the holder to deal in non-medicinal poisons.

³ NT: The licence holder has to notify the Pharmacy and Poisons (Wholesale Licences) Committee ("the Committee") and to provide storage facilities for pharmaceutical products in accordance with Section 2 of the Code of Practice before it may handle pharmaceutical products.

⁶ Premises Address: The address stated in the <u>same business registration certificate number</u> as registered when applying for the license.

⁷ Additional Warehouse: <u>Any address other than</u> that stated on the <u>same business registration certificate number as registered</u> when applying for the license.

Content of Change of Particulars Checklist (Cont'):

No.		e of Particulars Details	Submission of Supporting Documents (Refer to Page 6-23)				
		rage (Cont') ×	Submission of Supporting Documents (telef to Fage v 20)				
N	(i) With - Chang Conditi	nin Approved Store Room / Facilities: ge of Layout with not affect the Storage on	Storage at Premises Address ⁶ : 'COP Form' + 'COP Checklist Details' (14.a), (14.b), (15.a), (15.b) Storage at Additional Warehouse ⁷ : 'COP Form' + 'COP Checklist Details' (14.c), (14.d), (15.a), (15.b) Storage at Premises Address ⁶ : 'COP Form' + 'COP Checklist				
		thin Approved Store Room / Facilities: on of Additional Store Room / Facilities	Details' (14.b), (15.b) Storage at Additional Warehouse ⁷ : 'COP Form' + 'COP Checklist Details' (14.d), (15.b)				
Q Additional Warehouse ⁷ : - Change or Addition of Pharr Refrigerator / Cold Room / Fr		ge or Addition of Pharmaceutical	Storage at Premises Address ⁶ : 'COP Form' + 'COP Checklist Details' (14.b), (15.b), (19)^, (20)^ Storage at Additional Warehouse ⁷ : 'COP Form' + 'COP Checklist Details' (14.d), (15.b), (19)^, (20)^ Storage at Premises Address ⁶ : 'COP Form' + 'COP Checklist				
	- Deleti Room /	nal Warehouse ⁷ : on of Pharmaceutical Refrigerator / Cold Freezer	Details' (14.b), (15.b) <u>Storage at Additional Warehouse</u> ⁷ : 'COP Form' + 'COP Checklist Details' (14.d), (15.b)				
Other			(CODE 11 (COD CL. 11 (D. 21 (21)				
R	Change	or Addition of Transaction Record Format	'COP Form' + 'COP Checklist Details' (21)				
S	With NC ⁴ and NM ¹ or MD ² Condition	(i) Remove NM¹ or MD² Condition) (To Allow Pharmaceutical Products/Poisons Trade in Room Temperature Storage condition) (ii) Remove NM¹ and NC⁴ or MD² and	Storage at Premises Address ⁶ : 'COP Form' + 'COP Checklist Details' (12.a), (14.b), (15.b), (16), (17), (18), (19) Storage at Additional Warehouse ⁷ : 'COP Form' + 'COP Checklist Details' (12.a), (14.d), (15.b), (16), (17), (18), (19), (25), (26) Storage at Premises Address ⁶ : 'COP Form' + 'COP Checklist				
	M ¹ or	NC ⁴ Condition) (To Allow Pharmaceutical Products/Poisons Trade in Room Temperature Storage + Cold Chain Storage)Room Temperature Storage + Cold Chain Storage condition)	Checklist Details' (12.a), (14.d), (15.b), (16), (17), (18), (19), (20), (25), (26)				
	With IE ⁵ Condition	(iii) Remove IE ⁵ Condition) (To Allow Pharmaceutical Products/Poisons Trade not bound to Import for Re-export only)	Storage at Premises Address ⁶ : 'COP Form' + 'COP Checklist Details' (12.b) Storage at Additional Warehouse ⁷ : 'COP Form' + 'COP Checklist Details' (12.b)				
	Condition (iv) Remove NT ³ Condition) (To Allow Pharmaceutical Products/Poisons Trade in Room Temperature Storage)	Storage at Premises Address ⁶ : 'COP Form' + 'COP Checklist Details' (12.a), (14.b), (15.b), (16), (17), (18), (19) Storage at Additional Warehouse ⁷ : 'COP Form' + 'COP Checklist Details' (12.a), (14.d), (15.b), (16), (17), (18), (19), (25), (26)					
	and NC ⁴	(v) Remove NT ³ and NC ⁴ Condition) (To Allow Pharmaceutical Products/Poisons Trade in Cold Chain Storage condition)	Storage at Premises Address ⁶ : 'COP Form' + 'COP Checklist Details' (12.a), (14.b), (15.b), (16), (17), (18), (19), (20) Storage at Additional Warehouse ⁷ : 'COP Form' + 'COP Checklist Details' (12.a), (14.d), (15.b), (16), (17), (18), (19), (20), (25), (26)				
	With NC ⁴ Condition	(vi) Change of Licence Condition (Remove NC ⁴) (To Allow Pharmaceutical Products/Poisons Trade in Cold Chain Storage condition)	Storage at Premises Address ⁶ : 'COP Form' + 'COP Checklist Details' (12.a), (14.b), (15.b), (16), (17), (18), (19), (20) Storage at Additional Warehouse ⁷ : 'COP Form' + 'COP Checklist Details' (12.a), (14.d), (15.b), (16), (17), (18), (19), (20), (25), (26)				

%(Should maintain at least 1 storage facility)

^{^(}Not applicable for Wholesale Dealer Licence with 'NM'¹, 'MD'² or 'NT'³ condition)

¹ NM: This licence only authorizes the holder to deal in non-medicinal poisons.

² MD: This licence only authorizes the holder to deal in medical devices containing poisons.

³ NT: The licence holder has to notify the Pharmacy and Poisons (Wholesale Licences) Committee ("the Committee") and to provide storage facilities for pharmaceutical products in accordance with Section 2 of the Code of Practice before it may handle pharmaceutical products.

⁴NC: The licence holder must not handle pharmaceutical products that require cold chain management.

⁵IE: This licence only authorizes the holder to carry on the business of importing poisons/pharmaceutical products for reexport purpose.

^{6.} Premises Address: The address stated in the <u>same business registration certificate number</u> as registered when applying for the license.

7. Additional Warehouse: <u>Any address other than</u> that stated on the <u>same business registration certificate number as registered</u> when applying for the license.

Content of Change of Particulars Checklist (Cont'):

No.	Change of Particulars Details	Submission of Supporting Documents (Refer to Page 6-23)				
Othe	rs (Cont')					
T	Cancellation of Licence	'COP Checklist Details' (23)				
U	(i) Change/Addition of Additional Warehouse ⁷	'COP Form' + 'COP Checklist Details' (1.b), (13.b), (14.d),				
	outside Premises Address ⁶ (Room Temperature	$(15.b), (16)^{\wedge}, (17)^{\wedge}, (18)^{\wedge}, (19)^{\wedge}, (25), (26)$				
	Storage) ** Storage Storage					
	(ii) Change/Addition of Additional Warehouse ⁷	'COP Form' + 'COP Checklist Details' (1.b), (13.b), (14.d),				
	outside Premises Address ⁶ (Room Temperature	$(15.b), (16)^{\land}, (17)^{\land}, (18)^{\land}, (19)^{\land}, (20)^{\land}, (25), (26)$				
	+ Cold Chain Storage) ※					
	(iii) Change of Premises Address with storage	'COP Form' + 'COP Checklist Details' (1.a), (13.a), (14.b)				
	facility at approved Additional Warehouse7					
	<mark>only ※</mark>					
	(iv) Change of Premises Address with storage	'COP Form' + 'COP Checklist Details' (1.a), (1.b), (13.a),				
	facility at unapproved Additional Warehouse ⁷	$(13.b), (14.b), (14.d), (15.b), (16)^{\land}, (17)^{\land}, (18)^{\land}, (19)^{\land}, (25),$				
	only <mark>%</mark>	(26), $((20)^{\wedge}$ should be provided if cold chain storage				
		involved)				
	(v) Apply for Certified True Copy	'COP Checklist Details' (24)				
	(vi) Apply for Overpayment Claim	'COP Checklist Details' (27)				
	(vii) Other changes not applicable to Item A-U(i-vi)	Please contact Drug Office 'Wholesale Regulatory Unit'				

%(Should maintain at least 1 storage facility)

^{^(}Not applicable for Wholesale Dealer Licence with 'NM'¹, 'MD'² or 'NT'³ condition)

^{1.} NM: This licence only authorizes the holder to deal in non-medicinal poisons.
2. MD: This licence only authorizes the holder to deal in medical devices containing poisons.

^{3.} NT: The licence holder has to notify the Pharmacy and Poisons (Wholesale Licences) Committee ("the Committee") and to provide storage facilities for pharmaceutical products in accordance with Section 2 of the Code of Practice before it may handle pharmaceutical products.

⁶ Premises Address: The address stated in the <u>same business registration certificate number</u> as registered when applying for the license. ^{7.} Additional Warehouse: Any address other than that stated on the same business registration certificate number as registered when applying for the license.

Application Form for Change of Particulars of Wholesale Dealer Licence (Cap. 138 Pharmacy and Poisons Ordinance)/ Antibiotics Permit (Cap. 137 Antibiotics Ordinance)/ Wholesale Dealer's Licence to Supply Dangerous Drugs (Cap. 134 Dangerous Drugs Ordinance)

(*) represent must fill items									
* Name of Business:									
* Application for Change for Licence (Licence number format: 1/2A/1234):									
	Wholesale Dealer Licenc	e (WDL));		Licence no: /2A/				
☐ Antibiotics Permit (AP); Licence no:/1A/									
☐ Wholesale Dealer's Licence to Supply Dangerous Drugs (Part I); Licence no:									
	☐ Wholesale Dealer's Licence to Supply Dangerous Drugs (Part II); Licence no:/5A/								
_	SI CD 4 I D 4	•1 4							
	Change of Particulars Det			D 1 .		- · ·			
	8	Change	Add	Delete	8 \	Expected			
Details (Refer to Page 3-5) with signed and company stamped if needed) Effective Da									
	mpany Information				N.				
A	Company's Name				Name:				
Pe	rsonnel <mark>#(Should maintain i</mark>	at least 1	DD P	<u>(C)</u>					
В	Director (s)				Name:				
					(e.g. CHAN Tai Man change to LAM Yat Yut/ Add LAM				
					Yat Yut/Delete CHAN Tai Man)				
C	Partner (s)				Name:				
					(CYANETIN I I I I I I I I I I I I I I I I I I				
					(e.g. CHAN Tai Man change to LAM Yat Yut/ Add LAM				
Ъ	C 1 D : 4		/		Yat Yut/Delete CHAN Tai Man) Name:				
D	Sole Proprietor				Name.				
					(e.g. CHAN Tai Man change to LAM Yat Yut)				
Е	Person-in-Charge of		/		Name:				
L	Poisons and				1,000				
	Pharmaceutical				(e.g. CHAN Tai Man change to LAM Yat Yut)				
	Products				Reason of change:				
	Troducts				☐Resign ☐Retire ☐Position Change ☐Others:				
F	D / D '			/	Name:				
Г	Deputy Person-in- Charge of of Poisons				Name.				
	and Pharmaceutical				(e.g. CHAN Tai Man change to LAM Yat Yut/ Add LAM				
	Products				Yat Yut/Delete CHAN Tai Man)				
	Products				Reason of change: ☐ Resign ☐ Retire ☐ Position Change				
					Others:				
G	Person-in-Charge of		П		Name:				
U	Dangerous Drugs Pt. I #				rame.				
	Dangerous Drugs I t. I #				(e.g. CHAN Tai Man change to LAM Yat Yut/ Add LAM				
					Yat Yut/Delete CHAN Tai Man) Reason of change:				
					Resign □Retire □Position Change				
					Others:				
Н	Person-in-Charge of				Name:				
11	Dangerous Drugs Pt.II								
	#				(e.g. CHAN Tai Man change to LAM Yat Yut/ Add LAM				
	_				Yat Yut/Delete CHAN Tai Man)				
					Reason of change:				
					\square Resign \square Retire \square Position Change				
					□Others:				
I	Locum Pharmacist of				Period Covered				
	Dangerous Drugs				From:				
		/		/	To:				

Change of Particulars Details (Cont')*: * Change of Particulars Change Add Delete Details of Change (Provide details in written Expected with signed and company stamped if needed) **Effective Date Details (Refer to Page 3-5)** Address / Storage * (Should maintain at least 1 storage facility) Premises Address⁶ (with Address: storage facility) Premises Layout (storage area unchanged) Store Room Location L Locate at: (e.g. Change of Store Room) Layout of approved Store П Locate at: Room with structural (e.g. Extend or Minimize the Store Room Area) change (e.g. shape, size) Layout of approved Store Locate at: Room/ Facility without structural and Storage (e.g. Change of storing "Quarantined", "Released", Condition Change "Returned", "Recalled", "Rejected" Area) Storage Facility (Room П П Locate at: Temperature) П П Pharmaceutical Grade П Locate at: Refrigerator Cold Room/Pharmaceutical Locate at: Q Grade Freezer **Others** Transaction Record Format Licensing Condition Licence Condition: $\square NM^1 \square MD^2 \square NT^3 \square NC^4 \square \overline{IE^5}$ Others (if item A - T is not applicable) (e.g. Change/Addition of Additional Warehouse⁶ outside the Premises Address⁶; Move of storage facilities from Premises Address⁶ to Additional Warehouse⁶, etc.) ¹ NM: This licence only authorizes the holder to deal in non-medicinal poisons. ² MD: This licence only authorizes the holder to deal in medical devices containing poisons. ³ NT: The licence holder has to notify the Pharmacy and Poisons (Wholesale Licences) Committee ("the Committee") and to provide storage facilities for pharmaceutical products in accordance with Section 2 of the Code of Practice before it may handle pharmaceutical products. ⁴NC: The licence holder must not handle pharmaceutical products that require cold chain management. ⁵IE: This licence only authorizes the holder to carry on the business of importing poisons/pharmaceutical products for reexport purpose. 6. Premises Address: The address stated in the same business registration certificate number as registered when applying for the license. ⁷ Additional Warehouse: **Any address other than** that stated on the **same business registration certificate number as registered** when applying for the license. * Applicant information for COP application: _ Company Chop: _ Signature: _ Application Date: **Position:**

Company Director/Partner/Sole Proprietor ☐ PIC of PP/Poisons ☐ DPIC of PP/Poisons * If Authorized Person! required for application (if applicable, please sign the Appendix 12): Position: Email address: Telephone Number:

Checklist Details for Change of Particulars of Wholesale Dealer Licence (Cap. 138 Pharmacy and Poisons Ordinance)/

Antibiotics Permit (Cap. 137 Antibiotics Ordinance)/
Wholesale Dealer's Licence to Supply Dangerous Drugs (Cap. 134 Dangerous Drugs Ordinance)

Details of Change of Particulars Checklist:

Details (of Change of Particulars Checklist:					
(1.a)	Copy of Licensee's Updated Business Registration Certificate of the Premises Address ⁶ (within valid date & the					
	Business Registration Certificate Number must be consistent with the approved premises address where the					
	license is applied for)					
(1.b)	Copy of Licensee's Branch Business Registration Certificate or Tenancy Agreement or Logistics Services					
	Agreement of the Additional Warehouse ⁷ (within valid date)					
(2.a)	Copy of Form NNC2 (Notice of Change of Company Name with payment notice from Business Registration Office					
	and its payment receipt)					
(2.b)	Copy of Certification of Incorporation on the Change of Name					
(2.c)	Copy of Form 1(c) from Business Registration Office and its payment receipt					
(2.e)	Copy of Form 1(a) from Business Registration Office and its payment receipt					
(3)	Copy of Form NAR1 of Companies Registry and its payment receipt (within valid date)					
(4)	Copy of Form ND2A of Companies Registry with confirm receive date					
(5)	Lists of Director(s) (Appendix 5) (for All Existing Director(s)/Partner(s)/Sole Proprietor information)					
(6)	Declaration (Appendix 2a) (for New Employed Personnel only)					
(7)	Declaration (Dangerous Drugs (Part I) WDL) (Appendix 6) (for New Employed Personnel only)					
(8)	Declaration (Locum Pharmacist) (Appendix 7) (for New Employed Personnel only)					
(9)	Statement of Relevant Work Experiences (Appendix 2b) (for New Employed Personnel who have related work					
(-)	experiences to other than Existing Application Company trader(s) of western medicines in Hong Kong)					
(10)	Copy of Certifications of the above relevant working experience, e.g. testimonials from previous employer(s) (If					
(10)	having, for New Employed Personnel who have related work experiences to other than Existing Application					
	Company trader(s) of western medicines in Hong Kong)					
(11)	Copy of Annual Practicing Certificate and Valid Certificate of Registration (within valid date, for New					
(11)	Appointed PIC of DD(Pt. I) or Locum Pharmacist only)					
(12.a)	Trading documents (At least 1 set of: Import + Export OR Import + Local Distribution OR Local Distribution +					
(12.4)	Export OR Local Distribution Document) with Product Information :					
	Import:					
	- Quotation from Foreign Seller to Applicant					
	Export:					
	- Quotation from Foreign Purchaser to Applicant					
	- Relevant Document proving the Purchaser in Oversea Country is legally authorized to handle the pharmaceutical					
	products					
	Local Distribution Document (For the applicant who is a product certificate holder of pharmaceutical product):					
	- Submit copy of Certificate of Drug/ Product Registration					
	Local Distribution Document (For the applicant who is NOT a product certificate holder of pharmaceutical					
	product):					
	- Submit copies of agency agreement document(s) from the product certificate holder					
	- Certificate of Drug/ Product Registration					
	<u>Product Information</u> : (e.g. photo(s) of product unit carton, menu(s) or package insert)					
	- Showing ingredient(s) of the products					
	- Suggested dosage					
	- Storage condition					
	(For Licence under NC^4 licensing condition should not handle product with cold chain storage condition under $8^{\circ}C$)					
(12.b)	Trading documents (At least 1 set of: Local Distribution Document) with Product Information:					
, ,	Local Distribution Document (For the applicant who is a product certificate holder of pharmaceutical product):					
	- Submit copy of Certificate of Drug/ Product Registration					
	Local Distribution Document (For the applicant who is NOT a product certificate holder of pharmaceutical					
	product):					
	- Submit copies of agency agreement document(s) from the product certificate holder					
	- Certificate of Drug/ Product Registration					
	<u>Product Information</u> : (e.g. photo(s) of product unit carton, menu(s) or package insert)					
	- Showing ingredient(s) of the products					
	- Suggested dosage					
	- Storage condition					
	(For Licence under NC^4 licensing condition should not handle product with cold chain storage condition under $8^{\circ}C$)					
4.NC: The	e licence holder must not handle pharmaceutical products that require cold chain management.					

NC: The licence holder must not handle pharmaceutical products that require cold chain management.
 Premises Address: The address stated in the <u>same business registration certificate number</u> as registered when applying for the license.
 Additional Warehouse: <u>Any address other than</u> that stated on the <u>same business registration certificate number as registered</u> when applying for the license.

Details of Change of Particulars Checklist (Cont'):

Floor plan of the entire floor where the Premises Address⁶ are located including: - Name and address of applicant's company; - Room number of all units on the same floor (if any) and location of the applicant's company; and - Applicant's signature, date and company chop Floor plan of the entire floor where the Additional Warehouse⁷ outside the Premises Address⁶ are located (13.b)including: - Name and address of applicant's company; - Room number of all units on the same floor (if any) and location of the applicant's company; and - Applicant's signature, date and company chop (14.a)Existing Version Layouts of the Premises Address⁶ including: - Name and address of applicant's company; - Location(s) of all compartments and storage facilities inside the premises (if any) and purpose of each location/room: - Dimensions of all compartments and total area of the premises; and - Applicant's signature, date and company chop (14.b)**Proposed Version Layouts of the Premises Address⁶** including: - Name and address of applicant's company; - Location(s) of all compartments and storage facilities inside the premises (if any) and purpose of each location/room; - Dimensions of all compartments and total area of the premises; and - Applicant's signature, date and company chop Existing Version Layouts of Additional Warehouse⁷ outside the Premises Address⁶ including: (14.c)- Name and address of applicant's company; - Location(s) of all compartments and storage facilities inside the warehouse and purpose of each location/room; - Dimensions of all compartments and total area of the premises; and - Applicant's signature, date and company chop Proposed Version Layouts of Additional Warehouse⁷ outside the Premises Address⁶ including: (14.d)- Name and address of applicant's company; - Location(s) of all compartments and storage facilities inside the warehouse and purpose of each location/room; - Dimensions of all compartments and total area of the premises; and - Applicant's signature, date and company chop (15.a)Existing Version Layouts of the storage facilities including: - Name of applicant's company and address of the storage facility; - Dimensions and/or areas of storage facilities; - Areas for storing "Quarantined", "Released", "Rejected", "Returned" and "Recalled" products^; - Location(s) of air-conditioning outlet(s) and/or air-conditioner(s)^; - Location(s) of pest control device(s)^; - Location(s) of temperature and humidity uniformity assessment^: - Location(s) of shielded window (if any); and - Applicant's signature, date and company chop Proposed Version Layouts of the storage facilities including: (15.b)- Name of applicant's company and address of the storage facility; - Dimensions and/or areas of storage facilities; - Areas for storing "Quarantined", "Released", "Rejected", "Returned" and "Recalled" products^; - Location(s) of air-conditioning outlet(s) and/or air-conditioner(s)^; - Location(s) of pest control device(s)^; - Location(s) of temperature and humidity uniformity assessment[^]; - Location(s) of shielded window (if any); and - Applicant's signature, date and company chop (Not applicable for Wholesale Dealer Licence with 'NM'1, 'MD'2 or 'NT'3 condition)

NM: This licence only authorizes the holder to deal in non-medicinal poisons.

MD: This licence only authorizes the holder to deal in medical devices containing poisons.

NT: The licence holder has to notify the Pharmacy and Poisons (Wholesale Licences) Committee ("the Committee") and to provide storage facilities for pharmaceutical products in accordance with Section 2 of the Code of Practice before it may handle pharmaceutical products.

⁴NC. The licence holder must not handle pharmaceutical products that require cold chain management.

⁶ Premises Address: The address stated in the <u>same business registration certificate number</u> as registered when applying for the license. ⁷ Additional Warehouse: Any address other than that stated on the same business registration certificate number as registered when applying for the license.

Details of Change of Particulars Checklist (Cont'):

(16)	Calibration certificate of the hygrothermometer(s) installed in the proposed storage area (valid date should be							
(10)	covered the Temperature and Humidity Mapping & Daily Record Reports):							
	- Calibration certificate must be issued by the manufacturer or laboratory accredited by HOKLAS or CNAS or							
	Mutual Recognition Arrangement Partners for HOKLAS							
(17)	Temperature and humidity uniformity assessment with a conclusion in the proposed storage area:							
(17)	- Report of the 3- consecutive day (3 time-sections including 'morning', 'afternoon' and 'noon' per each mapping							
	location) recommended for at least 4 corners of the storage areas							
	- Conclude and specify the reason of choosing designated location(s) that will place the temperature and humidity							
	monitor for daily supervision							
	(The suggested assessment method may differ which depends on the Actual Size and Layout of storage area)							
(18)	Daily temperature and humidity monitoring record (should be started after the temperature and humidity							
	uniformity assessment at the designated location(s)chosen for daily monitoring) in the proposed storage area:							
	- For at least 3- consecutive day with 3 time-sections including 'morning', 'afternoon' and 'noon') at selected							
	position(s) inside the storage areas							
	(The suggested assessment method may differ which depends on the Actual Size and Layout of storage area)							
(19)	Latest cleaning and pest control procedures and associated record (specify the items and frequencies of relative							
	procedure) in the proposed cold chain storage area							
(20)	CHECKLIST of Application involving set up of pharmaceutical grade cold room, refrigerator(s) or freezer(s)							
	(Appendix 3)^							
(21)	Copy of Transaction Record Form for Proposed Version							
(23)	Cancellation of Wholesale Dealer Licence Form (Appendix 8)							
(24)	Certified True Copy Application Form (Appendix 9)							
(25)	Storage facilities or additional warehouses for poisons/pharmaceutical products outside the premises							
	(Appendix 4)							
(26)	Written Explanation with Company Letterhead including:							
	- Name and address of applicant's company;							
	- Reason for why storage facility cannot be provided within the business address of the premises;							
	- Provide details of the store, routine monitoring and maintenance;							
	- Applicant's signature, date and company chop							
(27)	Over-Payment Claim Application Form (Appendix 10)							

^{^(}Not applicable for Wholesale Dealer Licence with 'NM'¹, 'MD'² or 'NT'³ condition)



Declaration

, *Mr/ Mrs/ Miss/ Ms	(),
Full Name: (in En	glish – Surname first, then Other Names)	(in Chinese)
*HKID / Passport No.:	hereby	y declare that I *have been
have not been an owner,	a director or an employee of other trader(s	6)# of western medicines in
Hong Kong <u>for the past tl</u>	nree years (i.e. importer/exporter, retailer, w	wholesaler or manufacturer,
regardless whether the trade	er(s) is/are still in business.)	
If so, please list out the rele	evant information in the following table.]	
	,	
	experiences at other# Pharmaceutical Trade	er(s) in Hong Kong in the
past three years:		
Full Name of Company		Period
(in English)	Position Held	(from month/year to
		month/year)
	[[] W/DI 2DIC / domute DIC (if amplicable)]	
	[\Bigcup \text{\text{"WDL \(^2\text{PIC}\) / deputy \(PIC\) (if applicable)]	
	[
	[WDL 2PIC / deputy PIC (if applicable)]	
	[\square ¹ WDL ² PIC / deputy PIC (if applicable)]	
WDL: Wholesale Dealer L	icence deputy) of Poisons / Pharmaceutical Product	· ·
ric. reison-m-enarge (or	deputy) of 1 ofsons / 1 harmaceutical 1 foduct	S
	nation given in this declaration is true,	
ınderstand that making fa	alse declaration will be liable to criminal p	rosecution.
	Signatura	
	Signature :	
	Name of Business :	
	Contact number :	
	E-mail Address :	

Not including the company under this application [Fill in Details as stated on Hong Kong Identity Card / Passport] * Delete as appropriate



Statement of Relevant Working Experiences in Western Medicine Traders

, "IVIT/ IVITS/ IVIISS/ IVIS),				
Full Name: (in E	nglish – Surname first, then Other Names)	(in Chinese)				
*HKID / Passport No.:	hereby	declare that I have the				
ollowing relevant working experiences in Hong Kong western medicine trader(s).						
Details of relevant working	experiences at other Pharmaceutical trade	er(s) in Hong Kong:				
Full Name of Company	Period					
(in English)	Position Held	(from month/year to				
		month/year)				
	[\square ¹ WDL ² PIC / deputy PIC (if applicable)]					
	[\square ¹ WDL ² PIC / deputy PIC (if applicable)]					
	[\square ¹ WDL ² PIC / deputy PIC (if applicable)]					
	[\square ¹ WDL ² PIC / deputy PIC (if applicable)]					
	[\square ¹ WDL ² PIC / deputy PIC (if applicable)]					
WDL: Wholesale Dealer L PIC: Person-in-Charge (or	icence deputy) of Poisons / Pharmaceutical Products	S				
	tion given in this Statement of Relevant Wors is true, correct and complete. I under o criminal prosecution.					
	Signature :					
	Name :					
	Name of Business :					
	Date:					

Not including the company under this application [Fill in Details as stated on Hong Kong Identity Card / Passport] * Delete as appropriate

Appendix 3

Application involving set up of pharmaceutical grade cold room, refrigerator(s) or freezer(s)

Please submit this checklist along with all the following documents, or otherwise we will be unable to process your application. Please provide a written explanation for each of the documents not submitted.

(1) Overview of cold chain equipmen on a separate sheet the details of each	t (if multiple pieces of equipment are involved, please piece of equipment):	list
(a) Type of pharmaceutical grade	facility/equipment:	
☐ Cold room ☐ Refrigerator	☐ Freezer ☐ Others (please specify:	_)
(b) Brand:		
(c) Model number:		
(d) Operating range (°C):		
(e) Exterior dimensions (mm): (Width × Depth × Height)		
(f) Interior dimensions (mm): (Width × Depth × Height)		
(g) Net capacity (liters):		
(h) Temperature uniformity assessment date and brief conclusion:		
(i) Open door test date and brief conclusion:		
(j) Close door / Power failure test date and brief conclusion:		
(k) Mode of remote alarm and alarm settings:		
(l) Back-up power test date and brief conclusion:		
(m) Holding duration of validated cold box:		
(n) Product name, active ingredient(s) and labelled storage condition of cold chain product to be handled:		

 (2) Layout of the cold room / refrigerator(s) / freezer(s) including the following items: (a) Name of applicant's company and the address of storage facility; (b) Dimensions and areas of the cold room / refrigerator(s) / freezer(s); (c) Areas for storing "Quarantined", "Released", "Rejected", "Returned" and "Recalled" products; (d) Location(s) of temperature uniformity assessment ("assessment points"); (e) Signature of the person in charge (PIC) of cold chain, date and company chop
 (3) Valid calibration certificate of each piece of the data logger(s) installed in the cold room / refrigerator(s) / freezer(s): (a) Should demonstrate the data logger(s) are calibrated for the operating range required by the pharmaceutical products stored in the cold room / refrigerator(s) / freezer(s); (b) Must be issued by the manufacturer or a laboratory accredited by HOKLAS or CNAS or Mutual Recognition Arrangement Partners for HOKLAS
 (4) Temperature uniformity assessment report: (a) The interval of the data logger(s) should be set at 1 minute or less; (b) At least 3 assessment points in every refrigerator and freezer, and 4 assessment points in the cold room (please justify the number of assessment points) with not less than 24 hours consecutive record at each point; (c) Procedure, data analysis, conclusion and raw data should be included; (d) Specify which designated location(s) will be used for daily monitoring in the conclusion
 (5) Temperature monitoring record (with at least 3 consecutive days data): (a) Should be started after the temperature uniformity assessment at the designated location(s) chosen for daily monitoring; (b) The interval of the data logger(s) should be set at 1 minute or less
(6) Open door test report:(a) Procedure, data analysis, conclusion and raw data should be included
(7) Close door / Power failure test report:(a) Procedure, data analysis, conclusion and raw data should be included
 (8) Temperature alarm test report: (a) Remote alarm (e.g. SMS/email alert); (b) Door open alarm (if any); (c) Specify the alarm settings and procedures for alarm test; (d) Provide raw data and screenshots of the remote alarm (High/Low alarm and door open alarm)
(9) Alarm sensor calibration certificate or report (unless the alarm is triggered by a calibrated data logger)
(10) Back-up power test report:(a) Procedure, data analysis, conclusion and raw data should be included

(11) Procedures for receipt, storage and delivery	y of cold chain products							
(12) Contingency plan during power failure or t	temperature excursion							
(13) Specification of the cold room / refrigerato	(13) Specification of the cold room / refrigerator(s) / freezer(s)							
☐ (14) Back-up power specification								
product (unless a calibrated data logger is used	 (15) Specification and/or validation report of the cold box to be used for delivery of cold chain product (unless a calibrated data logger is used for temperature monitoring during delivery) (a) For validation report, procedure, data analysis, conclusion and raw data should be included 							
	(16) Product information showing the active ingredient(s), dosage and storage condition of the cold chain product to be handled, e.g. photo(s) of product unit carton or package insert							
	klist and confirm the information and reports e PIC responsible for cold chain management							
☐ All sections of this checklist have been compl	leted with necessary documents attached.							
☐ I confirm the cold chain facility under this approducts.	pplication is suitable for storage of cold chain							
Signature of cold chain PIC :	Company chop:							
Name of cold chain PIC :	Date :							

Remarks: In addition to the documents as stated in this checklist above, other relevant supporting documents/information may be required to substantiate the application. Applications with incomplete submission of documents as stated in this checklist and without a written explanation will not be accepted.

Please observe the contents in relation to cold chain management from the "Code of Practice for Holder of Wholesale Dealer, including but not limited to section 2.12, 3.6 and 3.17., including but not limited to section 2.12, 3.6 and 3.17.



Storage facilities or additional warehouses for poisons/pharmaceutical products outside the premises

(As stated on Business Registration Certificate / Lease Contract / Pharmaceutical Logistics Services Agreement)

			Storage fac itional wa				Storage factitional ward (if applicational contraction)	rehouse 2	
Address of the storage facility or additional warehouse outside the premises (in English)							(п аррпса	oic)	
Total area of storage facility or additional warehouse outside the premises					m ²				m ²
Branch Business R of the applicant (no lease contract or a logistics services a submitted)	pharmaceutical								
Person in charge	Name (in English)								
of the storage	Name (in Chinese)								
facility or additional	HKID number								
warehouse	Position								
outside the premises	Office phone number								
	Mobile number								
	E-mail address								
Lockable storage room (area)					m ²				m ²
Lockable cabin	net (dimensions)	Width	Depth	Height	m	Width	Depth	Height	m
Lockable cold	room (area)				m^2				m^2
Lockable phare refrigerator (d	naceutical grade imensions)	Width	Depth	Height	m	Width	Depth	Height	m
Lockable phare freezer (dimer	naceutical grade nsions)	Width	Depth	Height	m	Width	Depth	Height	m
Written explanation	is required for the follo	wing situat	ion:						
i. Company wii. If there is no be provided	he business	premises, t						nnot	
☐ I understand all	written explanation. applications of storage and approval by the Phar							ected to	
Signature of Person Charge of Business Name of Person-in Charge of Business Position of Person- Charge of Business Name of the busines			-						

COMPANY CHOP

Date:



Director List

Name (in English)	Name (in Chinese)	HKID/Passport No.	Position
(Surname first, then			
Other Names)			
Signature of Ar	unlicant/Authorized De	rson!:	
Signature of Ap	pricani/Aumorized Te	18011 .	
Name of Ap	pplicant/Authorized Pe	rson!:	
Position of Ap	pplicant/Authorized Pe	rson':	
	Name of Day	·	
	Name of Bus	iness:	
	Company (Chop :	
		D.A.	
		Date :	

[All personnel listed in the above table should provide a signed declaration.]
[Fill in Details as stated on Hong Kong Identity Card / Passport]

[If application signed by Authorized Person, please submit Appendix 12]



Declaration (Dangerous Drugs (Part I) WDL)

I, *Mr/ Mrs/ Miss/ Ms	(),
Full Name: (in English – Surname first, then Othe	er Names)	(in Chinese)	
*HKID / Passport No.:	hereby	declare that I *hav	ve been
/ have not been an owner, a director or an employee of ot	her trader(s	<u>)</u> # of western medi	cines in
Hong Kong for the past three years (i.e. importer/exporter	er, retailer, w	holesaler or manuf	acturer,
regardless whether the trader(s) is/are still in business.)			
I declare that the information given in this declaration understand that making false declaration will be liable to			plete. I
Signature:			
Name :			
Name of Business:			
Contact number:			
E-mail Address:			
Data			

Not including the company under this application [Fill in Details as stated on Hong Kong Identity Card / Passport] * Delete as appropriate



Declaration (Locum Pharmacist)

I, *Mr/ Mrs/ Miss/ Ms	(),
Full Name: (in English – Surname first, then Other		(in Chinese)	
*HKID / Passport No.:	hereby	declare that I *hav	ve been
/ have not been an owner, a director or an employee of oth	er trader(s	<u>)</u> # of western medic	cines in
Hong Kong for the past three years (i.e. importer/exporter	r, retailer, w	holesaler or manuf	acturer,
regardless whether the trader(s) is/are still in business.)			
I declare that the information given in this declaratio understand that making false declaration will be liable to			olete. I
Signature:			
Name :			
Name of Business:			
Contact number:			
E-mail Address :			
D. (

Not including the company under this application [Fill in Details as stated on Hong Kong Identity Card / Passport] * Delete as appropriate

Appendix 8

(For reference purpose)

Cancellation of Wholesale Dealer Licence Form

Name of Business:		
Application for Cancellation for Licence (Licence number	r format: 1/	2A/1234):
☐ Wholesale Dealer Licence (WDL);	Licence no:	/2A/
☐ Antibiotics Permit (AP);	Licence no:	/1A/
☐ Wholesale Dealer's Licence to Supply Dangerous Drugs (Part I);	Licence no:	/6A/
\Box Wholesale Dealer's Licence to Supply Dangerous Drugs (Part II);	Licence no:	/5A/
Date of Withdrawal:		
I, *Mr/ Mrs/ Miss/ Ms	(_),
Full Name: (in English - Surname first, then Other		
*HKID / Passport No.:	, the undersi	gned company's director
in the dealing of business relating to any licence restr products / poisons / antibiotics permit / dangerous drugs). If the business, a new application of licence is required.	_	
Contact Person (if different to the undersigned person): Name:	Tel:	
Signature of Director:		
Name of Director:		
Name of Business:		
Contact No.:		
Email Address :		
Date :		

[Fill in Details as stated on Hong Kong Identity Card / Passport]
* Delete as appropriate



Certified True Copy Application Form

Name of Business:			
Application for True Copy for Licence (Licen	nce number format: 1/2	A/1234):	
☐ Wholesale Dealer Licence (WDL);	Licence no:	/2A/	Qty:
☐ Antibiotics Permit (AP);	Licence no:	/1A/	Qty:
\Box Wholesale Dealer's Licence to Supply Dangerous \Box	Orugs (Part I);Licence no:	/6A/	Qty:
\Box Wholesale Dealer's Licence to Supply Dangerous \Box	Orugs (Part II);Licence no: _	/5A/	Qty:
Reason for Apply Certified True Copy: (Tick	the appropriate)		
☐ Not received from the date of mail by 'Who	olesale Regulatory Unit	within 1	month (shall
return to Drug Office when original copy was for	ound)		
☐ Lost <shall for="" hk\$220="" licence="" pay="" per=""></shall>			
\square Extra copy for business purpose (e.g. apply to	ender) <shall for="" hk<="" pay="" td=""><td>\$220 per l</td><td>licence></td></shall>	\$220 per l	licence>
☐ Others (Please specify:)
Signature of Applicant/Authorized Pe	rson!:		
2.8 01 1.pp.1			
Name of Applicant/Authorized Pe	rson! ·		
rame of Applicant/Tutiloff2ed Te			
Position of Applicant/Authorized Pe	rson!		
1 osition of Applicant Authorized 1 c			
Name of Ruc	iness:		
Name of Bus.			
Contact	No. :		
Contact	NO.:		
F 11 A 1	1		
Email Add	dress:		
Company S	tamp :		
	Date :		

[Fill in Details as stated on Hong Kong Identity Card / Passport]
¹[If application signed by Authorized Person, please submit Appendix 12]

Appendix 10

(For reference purpose)

Over-Payment Claim Application Form

Name of Business:		
Application for Refund for Licence (Licence number form	nat: 1/2A/12.	34):
☐ Wholesale Dealer Licence (WDL);	Licence no:	/2A/
☐ Antibiotics Permit (AP);	Licence no:	/1A/
☐ Wholesale Dealer's Licence to Supply Dangerous Drugs (Part I);	Licence no: _	/6A/
☐ Wholesale Dealer's Licence to Supply Dangerous Drugs (Part II);	Licence no: _	/5A/
Payment Type involved the Claim: (Tick the appropriate) ☐ New Application Fee ☐ Renewal Fee ☐ Change of Particulars Fee		
Payment Claim Action: (Tick the appropriate)		
☐ Claim for Refund with below Details:		(2 57
- Receiver's Name:		_ (for Cheque Deposit)
- Amount of Payment Refund: Demand Note Number:		
- Payment Date:		
- Mailing Address:		
☐ Rejected to Claim the Refund		
•		
//CAUTION: Submission of batch applications must be according to relevant information as proof before it will be acce		copy of the payment
as proof of the rank information as proof of for it will be acce	prod.i/	
Signature of Applicant/Authorized Person!:		
Name of Applicant/Authorized Person! :		
Position of Applicant/Authorized Person!:		
Name of Business:		
Contact No.:		
Email :		
Company Chop:		
Company Chop .		
Dete		
Date :		

[Fill in Details as stated on Hong Kong Identity Card / Passport] ^¹[If application signed by Authorized Person, please submit Appendix 12]



Authorization Letter

I, *Mr/ Mrs/ Miss/ Ms	(_),
Full Name: (in English – Surname first, then Other Names		(in Chinese)
*HKID / Passport No.:	, the undersi	gned company's director
hereby authorize (Authorized Person's Name: in English – Surname first, a		
manners to apply for Change of Particulars Application a	ccording to WD	L-COP Form submitted on
including signing and pro (Application Date)	viding all docum	nents relating to this matter.
Signature of Director:		
Name of Director:		
Name of Business:		
Contact No. :		
Email Address:		
Company Chop (Authorized Signature) :		
Date:		

[Fill in Details as stated on Hong Kong Identity Card / Passport]
* Delete as appropriate

Statement of Purposes

Purpose of Collection

- 1. This personal data are provided by licence applicants for the purposes of application for licences under the Pharmacy and Poisons Ordinance, the Antibiotics Ordinance and the Dangerous Drugs Ordinance. The personal data provided will be used by DH for the following purposes:
 - (a) Proof of eligibility for a licence
 - (b) Assessment of whether the applicant is a fit and proper person to be granted a licence
- 2. The provision of personal data is voluntary. If you do not provide sufficient information, we may not be able to prove your eligibility for a licence, or to assess whether you are a fit and proper person to be granted a licence.

Classes of Transferees

3. The personal data you provide are mainly for use within DH and the Pharmacy and Poisons Board. Apart from this, the data may only be disclosed to parties where you have given consent to such disclosure or where such disclosure is allowed under the Personal Data (Privacy) Ordinance.

Access to Personal Data

4. You have a right of access and correction with respect to personal data as provided for in sections 18 and 22 and Principle 6 of Schedule 1 of the Personal Data (Privacy) Ordinance. Your right of access includes the right to obtain a copy of your personal data. A fee may be imposed for complying with a data access request.

Enquiries

5. Enquiries concerning the personal data provided, including the making of access and corrections, should be addressed to:

Senior Pharmacist

Licensing and Compliance Division

Drug Office

Department of Health

Room 2001-2002, 20/F, Dah Sing Financial Centre,

248 Queen's Road East, Wan Chai, Hong Kong.

Telephone Number: 3107 2194