**Guidance on Application for Registration as Authorized Person**

**Introduction**

Under Regulation 30A of the Pharmacy and Poisons Regulations (Cap. 138A), a licensed manufacturer must ensure that at least one Authorized Person (AP) is employed to be responsible for ensuring and certifying that each batch of the pharmaceutical products has been manufactured and checked in accordance with the Good Manufacturing Practice (GMP) Guide issued by the Pharmacy and Poisons Board of Hong Kong, and that the registrable particulars of each batch of the pharmaceutical products correspond exactly with the registered particulars of the products.

2. For the purpose of making an application for registration as an AP, applicants need to provide evidence to the satisfaction of the Pharmacy and Poisons (Manufacturers Licensing) Committee (“the Committee”) that they possess the required qualifications and experience, and that they are fit and proper persons to be registered as APs. The Committee will assess the supporting evidence with the assistance of the Drug Office of the Department of Health.

**Qualification and experience requirements**

*Authorized Person for Pharmaceutical Manufacturers*

3. An applicant applying for registration as an AP for pharmaceutical manufacturers is required to satisfy either one of the following qualification requirements:
   (a) be a registered pharmacist in Hong Kong; or
   (b) hold a qualification awarded on completion of a course recognized by the Committee.

In addition to possessing the above qualifications, the applicant should have:
   (c) at least 3 years of relevant practical experience in GMP pharmaceutical manufacturing and/or quality control (at managerial or supervisory level) in one or more pharmaceutical manufacturers.

4. The relevant experience must have been acquired in Hong Kong or in a country/region whose regulatory authority is a Participating Authority of the Pharmaceutical Inspection Cooperation Scheme (PIC/S).

5. The 3 years of experience should include at least 1 year preparatory period (which is preceded by at least 2 years of the relevant experience) during which the person is under the supervision and professional guidance of a practising registered AP or an equivalent person in Hong Kong and should assist in exercising the duties of an AP. The applicant should provide evidence that he/she has gained such an experience.

6. For an applicant who has already been practising as an AP or equivalent positions in countries/regions whose regulatory authority is a PIC/S Participating Authority, the applicant should provide evidence that his/her qualification and experience are comparable to the requirements stated in paragraph 3.

*Authorized Person for Manufacturers of Advanced Therapy Products*

7. An applicant applying for registration as an AP for advanced therapy product (ATP) manufacturers should, subjected to conditions that the Committee thinks fit to impose, be:

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1 The definitions of advanced therapy products are given in Annex 1.
(a) a holder of a bachelor’s degree in a discipline such as biotechnology, biomedical engineering, medical laboratory science and other similar disciplines with at least 3 years of working experience at managerial or supervisory level in GMP manufacturing or quality control of ATPs;

(b) a holder of a postgraduate degree of science relevant to cell therapy, gene therapy, regenerative medicines or tissue engineering, or other related sciences with at least 2 years of working experience at managerial or supervisory level in GMP manufacturing or quality control of ATPs;

(c) a holder of a degree of doctor of philosophy (PhD) of science relevant to cell therapy, gene therapy, regenerative medicines or tissue engineering, or other related sciences with at least 1 year of working experience at managerial or supervisory level in GMP manufacturing or quality control of ATPs;

(d) a person who has already been practicing as an AP or equivalent positions for ATP manufacturers in a country/region where the regulatory authority is a PIC/S Participating Authority; or

(e) a holder of a degree of PhD of science relevant to cell therapy, gene therapy, regenerative medicines or tissue engineering, or other related sciences with at least 2 years of post-doctoral working experience in the processing or quality control of cells, genes and tissue engineered products and with evidence of theoretical and practical training in GMP principles related to ATP manufacturing

8. The applicant should also provide evidence that he or she has comprehensive knowledge on the pharmaceutical law and administration in Hong Kong as stated in section 1 of Annex A of the “Guidance on Qualification, Experience and Training Requirements for Authorized Persons and Other Key Personnel of Licensed Manufacturers in Hong Kong”.

**Authorized Person for Secondary Packaging Manufacturers**

9. An applicant applying for registration as an AP solely for certification of the release of pharmaceutical products that have undergone secondary packaging operations is required to satisfy either one of the following qualification and experience requirements:

   (a) post-secondary qualification with:
      – 1 year of experience in GMP pharmaceutical manufacturing and/or secondary packaging of pharmaceutical products; or
      – 6 months of experience in GMP pharmaceutical manufacturing and/or secondary packaging of pharmaceutical products, together with a certified GMP training.

   (b) secondary qualification with:
      – 2 years of experience in GMP pharmaceutical manufacturing and/or secondary packaging of pharmaceutical products; or
      – 1 year of experience in GMP pharmaceutical manufacturing and/or secondary packaging of pharmaceutical products, together with a certified GMP training.

10. Secondary qualification means having:

   (a) Level 2 or equivalent or above in five subjects (including Chinese Language and English Language) in the Hong Kong Diploma of Secondary Education Examination (HKDSEE), or equivalent;

   (b) Level 2 / Grade E# or above in five subjects (including Chinese Language and English Language) in the Hong Kong Certificate of Education Examination (HKCEE), or equivalent; or

2 Registration of AP by the qualification and experience listed in 7(e) is subjected to additional requirement stipulated in section 5.6 of the “Guidance on Qualification, Experience and Training Requirements for Authorized Persons and Other Key Personnel of Licensed Manufacturers in Hong Kong.”
(c) local accredited Higher Diploma, Associate Degree, Foundation Diploma, Yi Jin Diploma, or equivalent.

# ‘Grade E’ in Chinese Language and English Language (Syllabus B) in the HKCEE before 2007 are accepted administratively as comparable to ‘Level 2’ in Chinese Language and English Language in the 2007 HKCEE and henceforth.

For qualifications awarded by granting bodies outside Hong Kong, the Hong Kong Council for Accreditation of Academic and Vocational Qualifications (HKCAAVQ) provides assessment services for the general public, organisations, and government bureaux/departments. The HKCAAVQ offers a professional opinion on whether the totality of the educational qualifications (i.e. the integrated learning outcomes of the highest and terminal educational qualification) of an individual meets the standard of a particular level of qualification in Hong Kong (http://www.hkcaavq.edu.hk/).

11. In granting a Certificate of Registration as AP, the Committee must take into consideration but are not limited to the following:
(a) previous drug-related conviction(s), in particular those have significant impact to the public the public interest, of the applicant; and
(b) previous disciplinary action(s) against the applicant.

Application procedure

12. Application form for registration as an AP can be obtained, free of charge, by downloading from the website http://www.drugoffice.gov.hk or in person during the following hours:

Department of Health, Drug Office, Licensing and Compliance Division, Manufacturing Quality Assurance Unit
Room 2550, 25/F, Wu Chung House, 213 Queen's Road East, Wan Chai, Hong Kong
Tel.: 2961 8162 Fax: 3904 1225
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

13. The completed application form together with the relevant supporting documents should be submitted by post or in person to the above address.

14. In applying for registration, applicants have to produce the original (with photocopies) or notarised copies of the following supporting documents, together with the completed application form:
(a) certificate(s) of the academic qualification and relevant training
(b) certificate(s) of the professional qualification (if applicable)
(c) certification(s) of experience in pharmaceutical manufacturing, quality control or secondary packaging (e.g. testimonial(s) from employers)
(d) certification of the preparatory period under the supervision and guidance of a practising AP (e.g. a letter signed by the AP providing guidance certifying that the applicant had gained such experience)
(e) proofs of identity (Hong Kong identity card or passport).

15. After vetting by the Drug Office, the application will be considered by the Committee. The Committee may impose such conditions relating to the registration as it may think fit.

16. Any applicant aggrieved by a decision made in respect of the applicant may, in the prescribed manner, appeal to the Pharmacy and Poisons Appeal Tribunal against that decision.
17. Any enquiries on matters related to the application should be sent to the Manufacturing Quality Assurance Unit at the above address.

**Certificate of registration**

18. An applicant who is accepted for registration is required to pay a prescribed fee of $1420 for a Certificate of Registration which will be issued when his/her name is added to the Register of Authorized Persons. The certificate issued is valid until the end of the year and has to be renewed every year at a prescribed fee of $1,420.

19. The Committee may cancel or suspend the registration of an AP, or vary the registration condition(s) if, in its opinion, the AP is no longer a fit and proper person, has contravened a condition of the registration, any provision of the Pharmacy and Poisons Regulation or a Code of Practice applicable to AP, or has been convicted of a drug-related offence.

**Notes**

20. These notes are only a general guide and must not be treated as a complete or authoritative statement of the law on any particular case.

21. Copies of the Pharmacy and Poisons Ordinance and its subsidiary legislation may be purchased by calling the Publications Sales Section of Information Services Department at 2537 1910 or by email at puborder@isd.gov.hk. Contents of the relevant legislation may also be found at the Department of Justice’s website http://www.elegislation.gov.hk.
Annex 1: Definitions of Advanced Therapy Product (Proposed)

Advanced Therapy Product
Advanced therapy product means any of the following products that is for human use—

(a) a gene therapy product;
(b) a somatic cell therapy product;
(c) a tissue engineered product.

Gene Therapy Product
Gene therapy product—

(a) means a product—
(1) that contains an active substance containing or consisting of a recombinant nucleic acid that may be used in or administered to human beings with a view to regulating, repairing, replacing, adding or deleting a genetic sequence; and
(2) the therapeutic, prophylactic or diagnostic effect of which relates directly to—
(A) the recombinant nucleic acid sequence it contains; or
(B) the product of genetic expression of that sequence; but
(b) does not include a vaccine against an infectious disease.

Somatic Cell Therapy Product
Somatic cell therapy product means a product that—

(a) contains or consists of any of the following cells or tissues—
(1) cells or tissues that have been subject to substantial manipulation so that their biological characteristics, physiological functions or structural properties relevant for the intended clinical use have been altered;
(2) cells or tissues that are not intended to be used for the same essential functions in their recipient as in their donor; and
(b) is presented as having properties for, or may be used in or administered to human beings with a view to—
(1) treating, preventing or diagnosing a disease; or
(2) restoring, correcting or modifying physiological functions, through the pharmacological, immunological or metabolic action of those cells or tissues.

Tissue Engineered Product
Tissue engineered product—

(a) means a product that—
(1) contains or consists of any of the following cells or tissues—
(A) cells or tissues that have been subject to substantial manipulation so that their biological characteristics, physiological functions or structural properties relevant for the intended regeneration, repair or replacement have been altered;
(B) cells or tissues that are not intended to be used for the same essential functions in their recipient as in their donor; and
(2) is presented as having properties for, or may be used in or administered to human beings with a view to, regenerating, repairing or replacing a human tissue; but
(b) does not include a product that—
(1) contains or consists of exclusively non-viable human or animal cells or tissues; and
(2) does not act principally by pharmacological, immunological or metabolic action.
**Substantial Manipulation**

Substantial manipulation, in relation to cells or tissues, does not include the manipulation processes set out in the Schedule of the Pharmacy and Poisons Ordinance, Cap. 138.

Under the Schedule of Cap. 138, the following manipulation processes are not substantial manipulations—

1. Cutting
2. Grinding
3. Shaping
4. Centrifugation
5. Soaking in antibiotic or antimicrobial solutions
6. Sterilization
7. Irradiation
8. Cell separation, concentration or purification
9. Filtering
10. Lyophilization
11. Freezing
12. Cryopreservation
13. Vitrification
Statement of Purposes

Purpose of Collection

This personal data are provided by applicants for the purposes of application for registration as Authorized Person under the Pharmacy and Poisons Ordinance. The provision of personal data is voluntary. If you do not provide sufficient information, we may not be able to process your application.

Classes of Transferees

2. The personal data you provide are mainly for use within the Department of Health and the Pharmacy and Poisons Board. Moreover, according to the Pharmacy and Poisons Ordinance, part of the information provided, such as names of Authorized Person and addresses, will be entered into a Register for public inspection. 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

3. 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

4. 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 2550, 25/F, Wu Chung House,
   213 Queen's Road East, Wan Chai, Hong Kong.
   Tel: 2961 8028