Invitation for Using the Pharmaceuticals Licence Application and Movement Monitoring System (Phase II)

Drug Office will roll out the second phase of Pharmaceuticals Licence Application and Movement Monitoring System (“PLAMMS II”) to extend the scope of import/export licence processing to unregistered pharmaceutical products for other purposes.

In order to ensure smooth transition from manual to electronic submission, the implementation of electronic submission will be conducted in two stages according to the following timeline, and manual submission will still be accepted during the transition period until further notice:

(i) From **30 September 2019** onwards, all applications for import and export licences of registered pharmaceutical products and removal licence for dangerous drugs; and

(ii) From **30 December 2019** onwards, all applications for import and export licences/certificates of the following products or substances –

a. Registered pharmaceutical products;

b. Dangerous drugs (other than removal licence for dangerous drugs);

c. Unregistered pharmaceutical products for the treatment of particular patients/animals by a registered medical practitioner/dentist/veterinary surgeon;

d. Pharmaceutical products for the purpose of clinical trials/medicinal test; and

e. Pharmaceutical products or substances imported by a pharmaceutical manufacturer for the purpose of manufacture of pharmaceutical products.

If you are not an existing user of PLAMMS, you are cordially invited to register as a user in order to be able to use PLAMMS II for online submission of application for import/export licences of pharmaceutical products, dangerous drugs or psychotropic substances. For details of new account registration, please refer to our website: