Frequently Asked Questions and Answers on
Phase 2 Requirement of Bioavailability and Bioequivalence Studies

1. What is the timeline for the Phase 2 requirement of bioavailability and bioequivalence (BABE) studies for the registration of generic drugs?

   A. New applications for registration received before 1 August 2016
      For all new applications for registration of the 38 Critical Dose Drugs / Narrow Therapeutic Range Drugs (NTRD) appended below received before 1 August 2016 but have not been completed for registration before 1 August 2017, the applicants must satisfy the Phase 2 requirement for BABE studies before the approval of the applications.

   B. New applications for registration received on or after 1 August 2016
      With effect from 1 August 2016, all new applications for registration of the 38 Critical Dose Drugs / NTRD appended below must include evidence to satisfy the Phase 2 requirement for BABE studies. Otherwise, the applications will not be accepted for evaluation.

   C. Renewal applications of registered generic drugs
      With effect from 1 August 2017, all renewal applications for registration of the 38 Critical Dose Drugs / NTRD appended below must include evidence to satisfy the Phase 2 requirement for BABE studies. Otherwise, the registration of the generic drugs will not be renewed.

38 Critical Dose Drugs / Narrow Therapeutic Range Drugs

- Acetohexamide
- Aminophylline
- Aprindine
- Chloramphenicol
- Choline theophylline
- Clindamycin
- Clonidine
- Cyclosporine
- Digitoxin
- Digoxin
- Isoetharine
- Isoprenaline
- Levodopa and Carbidopa
- Levothyroxine
- Lithium
- Metaproterenol
- Methotrexate
- Minoxidil
- Phenobarbital
- Prazosin
Diprophylline  Procainamide  
Disopyramide  Proxyphylline  
Ethynyl Estradiol  Quinidine  
Flecainide  Sirolimus  
Glibenclamide  Tacrolimus  
Gliclazide  Theophylline  
Glybuzole  Tolazamide  
Glyclopymide  Tolbutamide  
Guanethidine  Warfarin

2. What dose form will be affected by the Phase 2 requirement of BABE studies?

The Phase 2 requirement of BABE studies for the 38 Critical Dose Drugs / NTRD is applicable to only oral dose form except (i) clonidine which would be applicable to both oral dose form and transdermal patch; and (ii) isoetharine which would be applicable to only inhalation aerosol.

3. What is the bioequivalence acceptance range applicable for Critical Dose Drugs / Narrow Therapeutic Range Drugs?

The current World Health Organization (WHO) guidelines on BABE registration requirements¹, including the bioequivalence acceptance range of 90.00-111.11% will be applicable for 6 antiepileptic drugs subject to the Phase 1 requirement for BABE studies, namely carbamazepine, clonazepam, divalproex, phenytoin, valproates and zonisamide belonging to Narrow Therapeutic Range Drugs (NTRD) and the 38 Critical Dose Drugs / NTRD subject to the Phase 2 requirement of BABE studies according to the following timeline:

(i) 1 January 2018 for new applications and including applications for change of registered particulars of registered pharmaceutical products; and
(ii) 1 January 2023 for renewal applications.

For the current WHO guidelines on BABE registration requirements, please visit the following website:


4. Are there any arrangements to exempt BABE studies for some registered pharmaceutical products when their comparators cannot be identified nor available?

In order to address the concerns on the difficulties to identify comparators of some registered pharmaceutical products for BABE studies and to preserve the continuous supply of these registered products for the interests of patients, the following arrangements are provided for registered products affected by the Phase 2 requirement of BABE studies:

A. Exemption will be applicable:

   for the renewal of registered products which have been registered with the Pharmacy and Poisons Board before 1 August 2016 if the applicants could provide evidence to prove that no comparators could be identified according to the current WHO guidelines on BABE registration requirements nor available in the market;

B. Exemption will not be applicable:

   (i) when there are any changes in manufacturers of the registered products mentioned in 4A above, pharmacokinetic and/or pharmacodynamic studies will be required unless justified; and

   (ii) when there is any addition of active ingredient(s) of the registered products mentioned in 4A above, new drug applications and including clinical studies will be required.