APPENDIX 5

GUIDELINE ON REGISTRATION OF GENERICS

IMPORTANT NOTES:

This document shall be read in conjunction with the relevant sections of the main guidance document: **Drug Registration Guidance Document (DRGD)**, which is in accordance to the legal requirements of the **Sale of Drugs Act 1952** and the **Control of Drugs and Cosmetics Regulations 1984**.

1. **DEFINITION**

A generic product is a product that is essentially similar to a currently registered product in Malaysia. However, the term generic is not applicable to Biologics.

2. GENERIC APPLICATION

The following categories of product can be processed as generic application provided that it fulfils the definition of a generic product.

(i) Scheduled Poison

(Known as Controlled Medicine/Controlled Poison)

Products containing active ingredients as <u>listed</u> in the First Schedule under Poisons Act 1952.

(ii) Non-Scheduled Poison

(Known as "Over-the-Counter", OTC)

Products containing active ingredients which are <u>not listed</u> in the First Schedule under Poisons Act 1952; and are excluding active ingredients which are categorized under health supplements or natural products or cosmetics.

(a) Full Evaluation

Other than listed at (b) Abridge Evaluation

(b) Abridged Evaluation

which include, but not limited to the following:

- Antiseptics/ skin disinfectants;
- Locally-acting lozenges/ pastilles;
- Topical analgesic/ counter-irritants;
- Topical nasal decongestants;
- Emollient/ demulcent/ skin protectants;
- Keratolytics;
- Anti-dandruff;
- •Oral care:
- Anti-acne;
- Medicated plasters/ patch/ pad; and
- Topical antibacterial.

Medicinal Gas

For medicinal gases classified as generic products, please refer to Directive No. 8, 2021 and <u>Guideline on Registration of Medicinal Gases</u>.

Reference:

• Directive No. 8, 2021, NPRA.600-1/9/13 (18): Direktif Berkenaan Pengukuhan Pelaksanaan Kawalan Regulatori Ke Atas Produk-Produk Gas Perubatan dan Penggunaan Guideline on Registration of Medicinal Gases (11 February 2021)

3. SUBMISSION OF APPLICATION

Applicants are advised to refer to **Section A (5. Application Procedures)** of the DRGD for further explanation.

4. EVALUATION TIMELINE FOR GENERIC APPLICATION

Table 1: Evaluation Timeline for Generic Application

No.	Product Category	Evaluation Timeline			
(A)	Full Evaluation				
	Generic (Schedule Poison)	210 working days			
	Generic (Non-Schedule Poison)	210 working days			
(B)	Abridged Evaluation				
	Generic (Non-Schedule Poison)				
	(i) Single active ingredient	116 working days			
	(ii) Two (2) or more active ingredients	136 working days			

5. REQUIREMENTS FOR GENERIC APPLICATION

Please refer to the following Appendices supplemented together with the DRGD for further information, where applicable:

Appendix 9	Fees
Appendix 11	Regulatory Control of Active Pharmaceutical Ingredients (APIs)
Appendix 12	Priority Review
Appendix 13	Designation and Registration of Orphan Medicines
Appendix 14	Evaluation Routes
Appendix 15	Requirements for Full Evaluation and Abridged Evaluation
Appendix 16	Bioequivalence (BE) Requirements
Appendix 17	Product Names Not Permitted To Be Registered
Appendix 18	List of Permitted, Prohibited and Restricted Substances
Appendix 19	General Labelling Requirements
Appendix 19A	Prohibited Visual/ Graphics/ Statements on Label
Appendix 20	Specific Labelling Requirements
Appendix 21	Special Conditions for Registration of a Particular Product or Group of Products

Appendix 22	Educational Materials		
Appendix 23	Patient Dispensing Pack for Pharmaceutical Products		
Appendix 24	Appeal		
Appendix 25	Guideline for the Submission of Protocol of Analysis (POA)		
Appendix 26	Guideline for the Submission of Analytical Method Validation (AMV) Documents		
Appendix 27	Inspection		
Appendix 32	Explanatory Notes for Repackers		

6. REFERENCES FOR GENERIC APPLICATION

Applicants are also advised to refer to <u>NPRA's website</u> for the latest registration requirements. In addition, other relevant and latest international guidelines e.g. by EMA, USFDA and ICH should also be referred to complement the ASEAN Guidelines and the DRGD as appropriate.

7. OTHERS

7.1 Classification of products containing Glucosamine, Chondroitin and Methylsulphonylmethane (MSM)

No.	Product		Product Category	Route of Evaluation	Condition on Product Indication	Remark
		As single active ingredient	ОТС	Full evaluation	As adjuvant therapy for osteoarthritis	Products containing glucosamine in combination with
1.	Products containing Glucosamine	As combination with Chondroitin and/ or MSM	ОТС	Full evaluation	As adjuvant therapy for osteoarthritis	other health supplement ingredients are only allowed to be registered for therapeutic purposes and NOT allowed to be registered as Health Supplement Product.

No.	Product		Product Category	Route of Evaluation	Condition on Product Indication	Remark
2.	Products containing Chondroitin	As single ingredient OR In combination with other supplement ingredients	Health supplement	Abridged Evaluation	No therapeutic claims are allowed	-
3.	Products containing MSM	As single ingredient OR In combination with other supplement ingredients As combination with Chondroitin	Health supplement Health supplement	Abridged Evaluation Abridged Evaluation	No therapeutic claims are allowed No therapeutic claims are allowed	-

References: Circulars

(i) Bil. (66) dlm BPFK/02/5/1.3

Produk yang Mengandungi Glucosamine dan Chondroitin (14 November 2006)

(ii) *Bil.* (20) *dlm.BPFK/PPP/01/03*

Produk yang mengandungi Glucosamine, Chondroitin dan Methylsulfonylmethane (MSM) (31 December 2008)

7.2 Classification of products containing combination of vitamin and/or mineral

- (i) Products containing a combination of vitamin and/or mineral are classified as Health Supplements. Please refer to <u>Appendix 6</u>: Guideline on Registration of Health Supplements for daily limit and registration requirements.
- (ii) For product containing a combination of vitamin and/or mineral with therapeutic indication:
 - (a) Product classification is required to determine the category of the said product as different regulatory requirements may apply. Applicant may submit a classification form, which can be downloaded from the NPRA website for classification of product category.
 - (b) Data/references to support the proposed combination and strength of active ingredients, dosage form, indication and dosing/posology will be required.
 - (c) Other supporting documents deemed necessary shall be submitted upon request to support the efficacy and safety of the product for the proposed indication.
 - (d) Approval status (for the same indication) together with the classification of the product in DCA reference countries (United Kingdom, Sweden, France, United States of America, Australia, Canada, Japan and Switzerland) is required.