

NEW PRODUCT CLASSIFICATION APPLICATION

1. Objective:

Product Classification main objective is to **determine whether a product is classified as a medical device product or not** under the Medical Device Act 737.

2. Guidance / Reference for submission:

- i. Guideline of Product Classification Application (insert hyperlink when ready)
- ii. E-Submission Guide for New Application of Product Classification Application (insert hyperlink when ready)

3. Disclaimer:

Please be informed that the invoice issued in relation to this application is valid for a period of one (1) month from the date of issuance. If payment is not received within this timeframe, we will follow up via email to confirm whether you wish to proceed.

Should we receive confirmation to withdraw, or if no response is received within the stipulated follow-up period, the application **may be withdrawn accordingly**.

4. General Process Flow for New Application:

