
ODB Regulation

12. (5.1) Clauses (1) (h) and (i) do not apply to a generic line extension drug product if the executive officer is satisfied that the product is safe, therapeutically effective or efficacious, and appropriate for public funding, having regard to,

a) its approval for sale in Canada by Health Canada, and
b) any other information available to the executive officer

A "generic line extension drug product" means a drug product with the same active ingredient or ingredients in the same or similar dosage form as an original product, but in a strength for which no original product exists.

Submission Requirements

For generic line extension products to be considered for funding under the Ontario Drug Benefit (ODB) program, the manufacturer must provide a complete submission that satisfies the following requirements:

1. Cover letter.
2. Evidence that Health Canada has approved the product for sale in Canada including:
   - A copy of the Notice of Compliance (NOC), if applicable;
   - A copy of the most recent Product Monograph approved by Health Canada.
3. A letter authorizing the Executive Officer to gain access to all information with respect to the Drug Product in the possession of Health Canada, the Patented Medicines Pricing Review Board established under section 91 of the Patent Act (Canada), the government of any province or territory in Canada or the Canadian Agency for Drugs and Technologies in Health and authorizing the Executive Officer to disclose any information with respect to the Drug Product in the possession of the Ministry to Health Canada, the Patented Medicine Prices Review Board, the government of a province or territory in Canada or the Canadian Agency for Drugs and Technologies in Health.

The ministry’s template letter is available on the ministry’s website.

4. An estimate of the net costs to the ODB Program in three-year period including:
   - Budget Impact Analysis (report and model; must include an estimate of the net costs to the ODB Program in a three-year period);

The Summary Sheet template is available on the ministry’s website.

5. Submit a proposed drug benefit price for the Drug Product.

6. Confirmation that the manufacturer is able to supply the Drug Product at the proposed drug benefit price in a quantity sufficient to meet the anticipated demand for the Drug Product.

The ministry’s template letter is available on the ministry’s website.

7. The manufacturer must certify in writing that no rebates were provided to persons listed under subsection 11.5(1) of the ODBA with respect to the Drug Product from the time that Health Canada approved the Drug Product for sale in Canada.

The ministry’s template letter is available on the ministry’s website.

8. Provide the scientific evidence upon which Health Canada approved the generic line extension drug product for sale in Canada.

9. Provide the Certified Product Identification Document (CPID) or Master Formula as evidence of formulation proportionality for the submitted strength(s).
If the CPID is not available, the ministry will accept the approved master formulation, dated and signed by a Senior Quality Assurance personnel. The master formulation must provide:

- the list of ingredients used to formulate the drug;
- information about the bulk formulation (granulation or liquid), if applicable;
- information about coating ingredients, if applicable; and information about the finished product expressed in the smallest quantity per unit (e.g., mg/tablet, mg/mL, etc.).

10. Provide justification for the therapeutic need for the new strength.
11. Certification confirming that the Drug Product is not a private label product.

The ministry’s template letter is available on the ministry’s website.

**Third Party Authorization/Business Agreement:**

Where a third party is involved with a submission, a letter should be submitted from both the NOC/DIN holder and the third party confirming the business arrangement between the submitting party and the NOC/DIN holder. Depending on the nature of the relationship, the letter from the NOC/DIN holder must provide an authorization to the third party to assume responsibility for the submission or authorization to submit the product on the NOC/DIN holder's behalf. All parties that have information on file with Health Canada, other provinces and other affiliated groups relating to the product must provide a consent letter allowing communication with these bodies as outlined in paragraph 3 above.

**Format and Organization of Submissions:**

The manufacturer must submit one electronic copy of the submission. For electronic submissions, the ministry will accept CDs, DVDs or USB keys, and they must fulfill the following requirements:

- The documents must be provided in MS Word, Excel or PDF format that is unlocked, searchable and printable.
- Users must have the ability to extract information or combine documents.

Manufacturers may wish to password protect any electronic submissions (i.e., CD, DVD or USB key) made to the ministry. Send the password needed to access the files to the email DrugSubmissions.MOH@ontario.ca ahead of the arrival of the electronic drug submission to the ministry.
Filing of Drug Submissions:
All submissions and any additional related information must be sent to:
Senior Manager
Drug Benefits Management Unit
Drug Programs Policy and Strategy Branch
Drugs and Devices Division
Ministry of Health
3rd Floor, 5700 Yonge Street
Toronto, ON M2M 4K5

Note: Although subsection 12 (5.1) of the ODBA Regulation exempts manufacturers of generic line extension products from specific submission requirements relating to clinical efficacy under clause 12(1)(h), the manufacturer must submit the scientific evidence upon which Health Canada approved the generic line extension drug product, as described in paragraph 8 above. The submission filed under subsection 12(5.1) requires CED review. Manufacturer may refer to Part II-A. of the Ontario Guidelines for more information on the process and timeless for submission going through the CED process.

The ministry expects manufacturers to follow the Guidelines when preparing submissions. The onus is on a manufacturer to provide the ministry with a submission that is complete, accurate and complies with applicable legislative, regulatory and policy requirements. Also, the ministry reserves the right to request additional information at any time during the review process.