

Protected B when completed

New class III medical device licence application form

(disponible en français)

Before completing this form, you must consult the document *Guidance for Industry – How to Complete the Application for a New Medical Device Licence* (available on the website).

1. Name of the device (as it appears on the	he label)						
2. Manufacturer information (as it appear	ars on the label)						
Contact Name and Title:					Comp	any	ID (if known):
Company Name:							
Telephone:		Fax:					
E-mail:							
Street:			ı	Suite	:	P.(O. Box:
City:	Province/State:		Country:				Postal/Zip Code:
3. Regulatory correspondent informat	ion	Sam	ne as Manu	facturer		Othe	er (specify below)
Contact Name and Title:					Comp	any	ID (if known):
Company Name:							
Telephone:		Fax:					
E-mail:				r			
Street:			T	Suite	:	P.0	O. Box:
City:	Province/State:		Country:				Postal/Zip Code:



4. Invoicing information	tion Same as Manufacturer Same as Regulatory Correspon		respondent	Oth	ther (specify below)		
Contact Name and Title:	tact Name and Title:		Comp	Company ID (if known):			
Company Name:							
Telephone:			Fax:				
E-mail:			<u> </u>				
Street:					Suite:	P.C	O. Box:
City:	Province	Province/State: Country:					Postal/Zip Code:
5. Quality management systen	n certificate (ensu	re that ce	tificate is attache	ed)			
Quality Management System Cert	ificate Number:	Name o	of Registrar:				
6. Attestations							
the items checked above and declapplication and in any attached do I, as a senior official of the manufaction information and documents set out Where a person is named in Item Minister on my behalf. I further autapplication to the person named in	ocumentation is actering acturer named in It in Part 1, section 3 of this application thorize the Medica	curate ar tem 2 of t n 32(3) of on, I herel al Devices	nd complete. this application the <i>Medical D</i> o by authorize th	, hereby a evices Reg at person	ttest that I am gulations. to submit this	also appli	providing the cation to the
Name:							
Title:							
Signature:			Date (yyyy/m	ım/dd):			
7. Purpose/intended use of de or represented. (Note: failure to suppl							

8. Licence application type (check one only)					
Single device	Test kit		Medical device group		
System	Medical device family		Medical device group fan		ımily
9. Place of use					
Is this device sold for home use?	Yes No	Is this device an IVDD?		Yes	No
Is this device used at a point of care, such as (In Vitro Diagnostic Devices [IVDD] ONLY)	a pharmacy, be	edside, or healthcare profession	al's office?	Yes	No
10. Medical devices containing drugs					
10.1 Non-IVD devices containing drugs					
If the device contains a drug and is not an IVDD, in and complete the information listed below. If the drug (DEL) number of the company from where the drug	rug does not have				
Brand/Trade Name of Drug:				DIN/NPN:	
Active Ingredient(s):					
Drug Manufacturer:					
DEL Number:					
10.2 IVDD Test Kits containing controlle	ed substances				
If this device is an IVDD test kit containing a substances Act, complete the section below.		in Schedule I, II, III, or IV of the	: Controlled I	Drugs and	
Is this an IVDD Test Kit containing a controlle	ed substance?			Yes	No
Test Kit Number (T.K. Number):					
Please note: The manufacturer will need to contact the	Office of Controlled	Substances to obtain a T.K. Number if of	one has not yet	been issued.	
11. Device history					
Has this device been previously authorized for Special Access provisions of the <i>Medical Dev</i>			ing or	Yes	No
If yes, provide the authorization number or the	e device identifi	cation number:			

12. Identifier of device: Include an identifier for each device or medical devi w/w of Di (2-Ethyl hexyl) Pthalate (DEHP) or is manufactured from raw materia	ice group listed and indica als containing or derived fr	te (by a che om Bispher	ck mark) if iol A (BPA	it contains ≥ 0.1%).
Name of device, components, parts and/or accessories as per product label	Identifier for device (bar code, catalogue, model or part number)	DEHP	BPA	Preferred Name Code (for Health Canada use only)

Notice to Industry – Licensing Requirements of Interdependent Medical Devices (April 30, 2002) available on the website. (Filenced medical devices, refer to: www.mdall.ca)	or a complete lis	st of
Name of compatible device	cence Numbe	er
14. List of recognized standards complied with in the manufacture of the device: Please answer "Yes" to one, and only one, of the following.		
The medical devices subject to this application conform with Recognized Standards as set out in the <i>Guidance Document on Recognition and Use of Standards under the Medical Devices Regulations</i> , which is available on the website.	Yes	No
If yes, I am including with this application Declarations of Conformity that the medical devices comply w Recognized Standards:	ith the followi	ing
The medical devices subject to this application DO NOT conform with Recognized Standards but meet an equivalent or better standard.	Yes	No
If yes, I am including detailed information proving that the devices meet the following equivalent or bette	er standards:	
The medical devices subject to this application DO NOT conform with Recognized Standards, NOR		
do they meet an equivalent or better standard, but I am including detailed information as evidence		

15. Priority review: The following section should be completed by manufacturers that wish to request priority review for their application in order to ensure a timely review of critical new medical devices for serious, life threatening, or severely debilitating diseases or conditions. Manufacturers will be notified at screening acceptance whether their application was accepted for priority review.

Is priority review requested for this application?

Yes

No

Priority review is being requested for the subject devices as it is intended for the diagnosis or treatment of a serious, life-threatening or severely debilitating disease or condition and there is substantial clinical evidence that the medical device:

Provides effective treatment or diagnosis of a disease or condition for which no medical device is currently licensed in Canada.

Provides significant risk-benefit improvement over existing therapeutic or diagnostic devices for a disease or condition that is not adequately managed by existing products marketed in Canada.

Responds to an unmet urgent healthcare need.

If a priority review is requested, a rationale must be included with this application form. The rationale should be an executive summary (10 pages or less) consisting of the following information:

- A synopsis of the clinical evidence establishing that the device provides effective treatment or diagnosis of a disease or condition for which no medical device is currently licensed in Canada or for which existing devices are inadequate.
- A brief description of the disease or condition and the role of the device.
- A summary of the risk-analysis for the device and/or the improvements it represents over existing products.
- **16. Review documents:** Indicate which documents listed below are included as attachments to this application. For details regarding content and format, you are requested to consult the *Guidance Document Preparation of a Premarket Review Document for Class III and Class IV Device Licence Applications* (available on the website).

Executive Summary

Table of Contents

Background, which includes Device Description, Design Philosophy, and Marketing History

Summary of Safety and Effectiveness Studies, which includes List of Standards, Method of Sterilization, Summary of Studies, and Bibliography

Near Patient Diagnostic Device Testing Results (if applicable)

Labelling Material

17. Fees

Please indicate that the Medical Device Licence Application Fee Form has been included with this application form.

Licence application disclosure request

As you are aware, Health Canada is striving to add transparency to the medical device review process. One area we would like to address is the requests from interested parties regarding whether or not a licence application has been received by the Medical Devices Directorate (MDD).

The purpose of this form is to request your signed authorization - in advance - if we receive such a request, to disclose the date on which a licence application has been received by the MDD. No other information would be supplied.

Please indicate your consent by completing this form and sending it with your application for a new medical device licence, or any time after a licence has been granted.

Disclosure Statement:

In the case where the MDD has received requests concerning the status of the new licence application, amendment application, or fax-back application for (enter device name)

from interested parties,

this certifies that (enter the manufacturer's name)

has **no objection** to the disclosure to the requester, by the MDD, of the date when an application for the device entered above, has been received by the MDD.

this certifies that (enter the manufacturer's name)

objects to the disclosure to the requester, by the MDD, of the date when an application for the device entered above, has been received by the MDD.

In accordance with the Access to Information Act, confidential, third party information will not be disclosed without your expressed consent.

Manufacturer's authorized signing official	

Application forms should be sent to:

Bureau of Licensing Services Medical Devices Directorate Health Canada 11 Holland Avenue Address Locator: 3002A Ottawa, Ontario K1A 0K9

Phone: (613) 957-7285 Facsimile: (613) 957-6345

E-mail: hc.devicelicensing-homologationinstruments.sc@canada.ca