



Class III Medical Device Licence Evaluation Report

Administrative Information

<i>Application Information</i>			
Application number	Licence Name	Licence Number	Risk Class
Application Type	Licence Type	Manufacturer	Company ID

<i>Scientific Review Information</i>		
Reviewed by	Review Division:	Review Division Manager:
Date Review Assigned: Click here to enter a date.	Date Review Completed: Click here to enter a date.	

This report contains Confidential Business Information (CBI) that should be redacted: Yes/No



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1 Background Information

Application Bundle Information	
Primary Application	
Secondary Applications	

1.1 Device Description

1.1.1 General Description

Reviewer's Discussion

1.1.2 Modifications

Reviewer's Discussion

1.2 Design Philosophy

Reviewer's Discussion

1.3 Indications and/or Intended Use, and Contraindications

This application includes changes to device indications: Yes/No

Reviewer's Discussion

2 Device Labels, Package Labelling and Documentation

2.1 Labels & Documentation



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Reviewer's Discussion

2.2 Warnings, Precautions and Limitations

Reviewer's Discussion

3 Marketing History/Regulatory Status

3.1 Canadian

This device has been released via SAP: Yes/No

This device has been authorised via ITA: Yes/No

Reviewer's Discussion

3.2 International

Reviewer's Discussion

3.3 Incident Reports

Reviewer's Discussion

4 Safety and Effectiveness Studies

4.1 List of standards

Reviewer's Discussion



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4.2 Preclinical studies

4.2.1 Physical/ Mechanical/ Bench Testing

Reviewer's Discussion

4.2.2 Shelf Life Studies for the Product

Reviewer's Discussion

4.2.3 Software Verification and Validation

Version Tested:

Reviewer's Discussion

4.2.4 Biocompatibility Tests

Reviewer's Discussion

4.2.5 Animal Studies

Reviewer's Discussion

4.3 Clinical studies

Reviewer's Discussion

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4.4 Sterilization, Packaging and Shelf-life Validation Studies

4.4.1 Sterilization/Validation/Re-sterilization/Bioburden

Reviewer's Discussion

4.4.2 Sterilant/Residual Toxicity Test

Reviewer's Discussion

4.4.3 Pyrogen Test

Reviewer's Discussion

4.4.4 Packaging and Packaging Stability

Reviewer's Discussion

4.5 Literature Studies & Bibliography

Reviewer's Discussion

5 Conclusion

[Start/Review Quality Report Card \(You will need to click OK in the popup after clicking here\)](#)

Note: Changes cannot be made once the feedback results are concurred by SH.

6 Recommendation

Acceptance for licensing

Acceptance of the following amendments:

Application no.

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GRP TMP_0001 v6.5



***Class III Medical Device Licence
Evaluation Report***

Acceptance with the following conditions:

Refusal:

Request for additional information:

DO NOT DELETE – Ensure recommendation falls above this line



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Executive Summary – DRAFT

<Paste Executive Summary here>