



## PUBLIC HEALTH UNIT INFECTION PREVENTION AND CONTROL LAPSE REPORT

## **Initial Report**

Premise/facility under investigation (name and address)	Callander Bay Dental – A Dawson Family Practice 299 Main Street North, Callander, ON POH 1H0
Type of premise/facility: (e.g., clinic, personal services setting)	Dental Office
Date Board of Health became aware of IPAC lapse	March 19, 2025
Date IPAC lapse was linked to the premise/facility	March 19, 2025
Date of Initial Report posting	April 3, 2025
Date of Initial Report update(s) (if applicable)	Not applicable
How the IPAC lapse was identified	Callander Bay Dental – A Dawson Family Practice
Summary Description of the IPAC Lapse	Improperly reprocessed reusable instruments used on clients.

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- 345 Oak Street West, North Bay, ON P1B 2T2
- 1-800-563-2808 705-474-1400
- **705-474-8252**
- 90 Bowes Street, Suite 201, Parry Sound, ON P2A 2L7
- 1-800-563-2808 705-746-5801
- **a** 705-746-2711



## **IPAC Lapse Investigation**

Did the IPAC lapse involve a member of a regulatory college? If yes, was the issue referred to the regulatory college? Were other stakeholders notified	Yes. • College of Dental Hygienists of Ontario • Royal College of Dental Surgeons of Ontario April 3, 2025 Yes
Concise description of the corrective action required.	The dental clinic identified and notified the impacted clients about the IPAC lapse.
Please provide further details/steps	<ol> <li>Along with the remedial measures already in place at the dental clinic, sterile packages must be inspected at the point of use to ensure they are not unsealed, damaged, wet, or visibly soiled, and that Chemical Indicators have not failed.</li> </ol>
	<ol> <li>Reprocessing areas shall be made up of distinct and separate work areas based on the service provided and generally include areas for:</li> </ol>
	Receiving contaminated devices;
	<ul> <li>Decontamination of medical devices;</li> </ul>
	High-level disinfection if applicable;
	<ul> <li>Preparation and packaging;</li> </ul>
	Sterilization and storage.
	3. The reprocessing area must have a one-way workflow of instruments with clear separation of dirty and clean sides to prevent contamination. A barrier (i.e., plexiglass) shall be installed to separate the decontamination area from the clean preparation and packaging area.
	4. The electronic record for load record requirements shall be inspected at the conclusion of each cycle by the operator of the sterilizer. The operator shall confirm that the process parameters were met either by initialing the report (if printed) or by means of a password-protected entry (if electronic).

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If you have any further questions, please contact:		
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