

# Medical Device Roundtable: Post-Market Surveillance Obligations

November 18, 2021

From medical device manufacturers and importers to hospitals and health professionals, in this webinar, the panel of regulatory and legal experts covered the following:

- 2021 updates to the Medical Devices Regulations on post-market surveillance obligations of medical device license holders;
- Reporting obligations to Health Canada on adverse incidents involving medical devices sold and imported into Canada; and
- Use of post-market surveillance regulatory disclosure in medical device litigation and cross-border issues.

# **Our Speakers:**

- Keegan Boyd, Partner, BLG
- Don Boyer, President, BOYER@RegulatorySolns
- Edona Vila, Partner, BLG

# Watch the webinar

By

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