

Mandatory Medical Device Problem Reporting Form for Industry

CANADA VIGILANCE - MEDICAL DEVICE PROBLEM REPORTING PROGRAM

If more space is required, please attach additional sheets

Fields required to be completed for updates/final reports are indicated by an $\ensuremath{^{\star}}$

| 3. Reporter File | No. * |
|---------------------------|---|
| 0700979127 | |
| 4. Health Canad | a File No. (if applicable) * |
| | |
| 5. Type of Repo | rt * |
| Preliminary | |
| If "preliminary" | only, anticipated date for the final report: |
| | (YYYY-MM-DD) |
| If "update/final" Canada: | , date the previous report was submitted to Health |
| 2015-09-22 | (YYYY-MM-DD) |
| 6. Date Submitted * | |
| 2020-09-17 | (YYYY-MM-DD) |
| | Importer |
| POLIS, MN, US | MEDTRONIC CANADA ULC 99 HEREFORD STREET, BRAMPTON, ON, CANADA Postcode:L6Y 0R3 Tel:(800)268-5346 Fax:(905) 460-3992 |
| | 106916 |
| | 35 |
| | |
| 5. Details of Incid | dent |
| ı | |
| - i | |
| i 4-4460.0 00.10 | FF04070b - F4 1007b 04 |
| 4cr119c9ce60d6 | 5561979ba51d627b21 |
| | 0700979127 4. Health Canad 5. Type of Repo ☐ Preliminary If "preliminary" If "update/final" Canada: 2015-09-22 6. Date Submitte 2020-09-17 |

A program of MedEffectTM Canada HC Pub.: 110180 (October 2011)



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| C. MEDICAL DEVICE INFORMATION | E. INVESTIGATION INFORMATION |
|--|--|
| 1. Trade/Brand Name * | 1. Investigative Actions and Timeline |
| SPRINT QUATTRO SECURE S | |
| 2. Control/Lot/Serial No. | |
| TAU079011V | |
| 3. Expiration Date | |
| 2013-08-12 | |
| 4. i. Device Classification | |
| □ I □ III □ III ⊠ IV | |
| ii. Device License No. | |
| 77725 | 70b05ba442ec98711e671b2a400bb5c8 |
| iii. Device Identification No | |
| iv. Manufacturer's Medical Device Identifier (catalogue/model no.) | |
| 6935-65 | |
| 5. Software Version | |
| Not Applicable | |
| 6. Age of Device | |
| Manufactured Date: 2011-08-22 | This section only applies for preliminary & final, and final reports |
| 7. How long was the device in use? | 2. Root Cause of Problem |
| Implant Date: 2012-08-31 | |
| 8. Was the device labelled as sterile? | |
| | |
| 9. Availability of device for evaluation | |
| ☐ Destroyed ☐ Returned to Manufacturer/Importer | |
| ⊠ Neither (with explanation) | |
| Still in use. | |
| | 6c6149000d0b3011a1bfae8780871f53 |
| D. COMPLAINANT INFORMATION | |
| 1. Complainant is a: | |
| ☐ Consumer ☐ Health professional ☐ Other | |
| 2. Name of Complainant | |
| 2c578407557f3b3d7f7b0a2a27ba3bb7 | |
| 3. Name of Health Care Facility (if applicable) | |
| cb5aa909aad665c3a421bcb3ede30ed7 | 3. Corrective Actions taken as a result of the investigation |
| 4. Address | 5. Sofredive Actions taken as a result of the investigation |
| 56f1ad8a3231c002724fcb6e697f5b09 | |
| 5. Telephone No. and/or E-mail Address | |
| 88cd85c8c6ba70375176edea3013b49c | 6a0cac3dfdf77d691732d3f69b33fdc3 |
| of the complainant and/or reporter will be protected as personal information under the <i>Privacy Act</i> , under the <i>Access to Information Act</i> in the case of an access to information request. For details with regard to personal information collected under this program, visit the Personal Information Bank; Health Canada; Health Products and Food Branch; Branch Incident Reporting | |

System;HC PPU 088 at: http://infosource.gc.ca/inst/1476/1476-fedemp00-

eng.asp