Access

Peel-Away Introducer

14 cm Sheath Introducer Kit 5 F-16 F

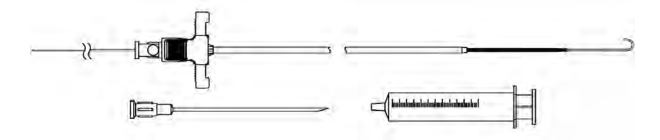
Product Highlights

- Proprietary materials improve insertion characteristics and reduce vessel trauma
- Close tolerance extrusion and proprietary tipping process improves tracking on a guidewire
- Reliable peeling characteristics
- Sheath can be totally occluded without kinking to prevent air inspiration
- Di-Lock[™] feature secures dilator in sheath during insertion

Ordering Information

Contents: Peel-Away Sheath, Di-Lock[™] Dilator, 12 cc syringe, 18 ga. XTW Needle, and 50 cm Guidewire with 3 mm "J" (5 units per box)

| Reorder Number | French Size | Maximum Guidewire Diameter (in) | Sheath Usable Length (cm) |
|-----------------------|----------------|------------------------------------|------------------------------|
| 405100 | 5 | .038 | 14 |
| 4051 <mark>0</mark> 4 | 6 | .038 | 14 |
| 405108 | 7 | .038 | <u>14</u> |
| 405112 | 8 | .038 | 14 |
| 405116 | 9 | .038 | 14 |
| 405118 | 9.5 | .038 | 14 |
| 4051 <mark>20</mark> | 10 | .038 | 14 |
| 405122 | 10.5 | .038 | 14 |
| 405124 | 11 | .038 | 14 |
| 405128 | 12 | .038 | 14 |
| 405132 | 13 | .038 | 14 |
| 405136 | 14 | .038 | 14 |
| 405144 | 16 | .038 | 14 |



U.S. Patent Number 5,098,392











EU Technical Documentation Assessment Certificate (MDR)

Pursuant to Regulation (EU) 2017/745 on Medical Devices, Annex IX Chapter II (Implantable Class IIb Devices and Class III Devices)

No. G70 014607 0256 Rev. 00

| Manufacturer: | Abbott Medical 15900 Valley View Court Sylmar CA 91342 USA |
|-------------------------------|--|
| SRN Manufacturer: | US-MF-000010383 |
| Authorized Representative: | Abbott Medical The Corporate Village, Da Vincilaan 11 Box F1, 1935 Zaventem, BELGIUM |

The Certification Body of TÜV SÜD Product Service GmbH certifies that the manufacturer has drawn up and presented a Technical Documentation according to Annex II and III of the Regulation (EU) 2017/745 on medical devices. Details on devices covered by the Technical Documentation are described on the following page(s). The Report referenced below summarises the result of the assessment and includes reference to relevant CS, harmonized standards and test reports. The technical documentation assessment included an assessment of the clinical evaluation assessment. The conformity assessment has been carried out according to Annex IX chapter II of this regulation with a positive result. Changes to the approved device, where such changes could affect the safety and performance of the device or the conditions prescribed for use of the device, shall require approval from the notified body TÜV SÜD Product Service GmbH. In order to place the devices on the market with CE-marking, an EU Quality Management System Certificate pursuant to Annex IX chapters I and III is necessary in addition to this EU Technical Documentation Assessment Certificate. All applicable requirements of the testing and certification regulation of TÜV SÜD Group have to be complied with. For details and certificate validity see: www.tuvsud.com/ps-cert?q=cert:G70 014607 0256 Rev. 00

Report No.: 713234168 Valid from: 2023-03-08 Valid until: 2028-03-07

Christoph Dicks Head of Certification/Notified

Body

Issue date: 2023-03-08

Page 1 of 2 TÜV SÜD Product Service GmbH is Notified Body with identification no. 0123 TÜV SÜD Product Service GmbH • Certification Body • Ridlerstraße 65 • 80339 Munich • Germany







EU Technical Documentation Assessment Certificate (MDR)

Pursuant to Regulation (EU) 2017/745 on Medical Devices, Annex IX Chapter II (Implantable Class IIb Devices and Class III Devices)

No. G70 014607 0256 Rev. 00

| Classification: | Class III |
|-------------------|--|
| Device Group: | C0503 - CARDIOVASCULAR INTRODUCER SHEATHS, PEEL- AWAY |
| Basic UDI-DI: | 5415067PLA0001FY |
| Intended Purpose: | The Peel Away Introducers are intended to provide a transvenous conduit for the introduction of cardiac leads and catheters into the venous vascular system. |
| Device(s): | Peel-Away Introducer. Article numbers: 405104, 405108, 405112, 405116, 405118, 405119, 405120, 405122, 405124, 405128, 405129, 405136, 405144, 405145, 405146, 405147, 405149, 405153, 405154, 405254, 405269, 405270, 405404, 405408, 405412, 405416, 405418, 405420, 405422, 405424, 405428 |

The validity of this certificate ./. depends on conditions and/or is limited to the following:

| Rev. | Dated | Report |
|------|------------|-----------|
| 00 | 2023-03-08 | 713234168 |

Description Initial issuance



0081901 Rev. A

Declaration of Conformity

| Manufacturer: | Abbott Medical |
|---|---|
| Manufacturer SRN: | US-MF-000010383 |
| Address: | 15900 Valley View Court Sylmar, CA 91342 United States of America |
| Manufacturing Site(s): | Abbott Medical 5050 Nathan Lane Plymouth, Minnesota 55442 United States of America |
| European Authorized Representative: | Abbott Medical The Corporate Village Da Vincilaan 11 Box F1 1935 Zaventem, Belgium |
| European Authorized Representative SRN: | BE-AR-000008744 |

This declaration of conformity is issued under the sole responsibility of the manufacturer.

| Device Name(s): | Introducers |
|---|---|
| Product Trade Name(s): | Peel-Away Introducer |
| Model Number(s): | 405104, 405108, 405112, 405116, 405118, 405119, 405120, 405122, 405124, 405128, 405129, 405136, 405144, 405145, 405146, 405147, 405149, 405153, 405154, 405254, 405269, 405270, 405404, 405408, 405412, 405416, 405418, 405420, 405422, 405424, 405428 |
| Intended Purpose: | The Peel-Away Introducers are intended to provide a transvenous conduit for the introduction of cardiac leads and catheters into the venous vascular system. |
| Risk Classification: | Class III as per EU MDR 2017/745 per Annex VIII |
| Signature: Digitally signed by CANANCX Date: 2023.05.11 12:48:06 -07'00' | May 11, 2023 |
| Colleen Canan Divisional Vice President, Global Regulatory Affairs, Cardiac Rhythm Management | Issue Date On behalf of Abbott Medical, signed at Sylmar, CA |
| 8136 MDR Declaration of Conformity Template Rev H | Page 1 of 3 |

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0081901 Rev. A

MDR Declaration of Conformity

| Risk Classification Rationale: | Annex VIII, Rule 6, 3 rd indent |
|--------------------------------|---|
| EMDN Code(s): | C0503 - Cardiovascular Introducing Sheaths, peel-away |
| GMDN Code: | 17846 |
| Basic UDI-DI: | 5415067PLA0001FY |

The products described in this declaration are in conformity with all applicable EU harmonized legislation, including:

• Regulation (EU) 2017/745, and the applicable General Safety & Performance Requirements in Annex 1

| Common Specifications Applied: | Not Applicable. No common specifications are available for this type of device |
|-----------------------------------|---|
| Notified Body: | TÜV SÜD Product Services GmbH Ridlerstraße 65 80339 Munich Germany |
| | |
| Supporting Certificate(s): | EU Technical Documentation Assessment Certificate (MDR) No: G70 014607 0256 Rev. 00 Expiration Date: 2028-03-07 EU Quality Management System Certificate (MDR) No: G12 014607 0255 Rev. 02 Expiration Date: 2027-08-14 |
| Original CE Mark Date: | May 16, 2013 (MDD) |
| Conformity Assessment: | EU MDR 2017/745, Annex IX |

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0081901 Rev. A

MDR Declaration of Conformity

The products in the attached Declaration of Conformity Product List are approved under EC Certificate No: G70 014607 0256 Rev. 00.

Declaration of Conformity Product List

| Model No. | Product Trade Name | UDI-DI |
|-----------|----------------------------------|----------------|
| 405104 | Peel-Away Introducer, 6F 14CM | 05415067040855 |
| 405104 | Peel-Away Introducer, 7F 14CM | 05415067040879 |
| 405100 | Peel-Away Introducer, 8F 14CM | 05415067040893 |
| 405112 | Peel-Away Introducer, 9F 14CM | 05415067040916 |
| 405118 | Peel-Away Introducer, 91 14CM | 05415067040930 |
| 405119 | Peel-Away Introducer, 3:51 14CM | 05415067040954 |
| 405120 | Peel-Away Introducer, 0.51 14CM | 05415067040978 |
| 405122 | Peel-Away Introducer, 10.5F 14CM | 05415067040992 |
| 405122 | Peel-Away Introducer, 10.51 14CM | 05415067041012 |
| 405128 | Peel-Away Introducer, 111 140M | 05415067041036 |
| 405129 | Peel-Away Introducer, 8F 14CM | 05415067041050 |
| 405136 | Peel-Away Introducer, 01 14CM | 05415067041074 |
| 405144 | Peel-Away Introducer, 14F 14CM | 05415067041098 |
| 405145 | Peel-Away Introducer, 8F CM | 05415067041111 |
| 405146 | Peel-Away Introducer, 8.5F CM | 05415067041135 |
| 405147 | Peel-Away Introducer, 9F CM | 05415067041159 |
| 405149 | Peel-Away Introducer, 10F CM | 05415067041173 |
| 405153 | Peel-Away Introducer, 7F CM | 05415067041197 |
| 405154 | Peel-Away Introducer, 7F 14CM | 05415067041210 |
| 405254 | Peel-Away Introducer, 9F 23CM | 05415067041258 |
| 405269 | Peel-Away Introducer, 7F 23CM | 05415067041272 |
| 405270 | Peel-Away Introducer, 8F 23CM | 05415067041296 |
| 405404 | Peel-Away Introducer, 6F 14CM | 05415067041319 |
| 405408 | Peel-Away Introducer, 7F 14CM | 05415067041333 |
| 405412 | Peel-Away Introducer, 8F 14CM | 05415067041357 |
| 405416 | Peel-Away Introducer, 9F 14CM | 05415067041371 |
| 405418 | Peel-Away Introducer, 9.5F 14CM | 05415067041395 |
| 405420 | Peel-Away Introducer, 10F 14CM | 05415067041418 |
| 405422 | Peel-Away Introducer, 10.5F 14CM | 05415067041432 |
| 405424 | Peel-Away Introducer, 11F 14CM | 05415067041456 |
| 405428 | Peel-Away Introducer, 12F 14CM | 05415067041470 |

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CERTIFICATE



This is to certify that



SANTE INTERNATIONAL S.A.

Str. Mantuleasa nr. 33, Sector 2 023961 Bucuresti Romania

has implemented and maintains a Quality Management System.

Scope:

Import, trade and storage of medical and laboratory equipment, disinfectants, laboratory reagents, cardiovascular surgery devices, service for medical and laboratory equipment. Consulting for state and private medical units.

Through an audit, documented in a report, it was verified that the management system fulfills the requirements of the following standard:

ISO 9001 : 2015

| Certificate registration no. | 497269 QM15 |
|------------------------------|-------------|
| Valid from | 2021-06-16 |
| Valid until | 2024-06-15 |
| Date of certification | 2021-06-16 |



DQS GmbH

Markus Bleher Managing Director







Annex to certificate Registration No. 497269 QM15

SANTE INTERNATIONAL S.A.

Str. Mantuleasa nr. 33, Sector 2 023961 Bucuresti Romania

Location

075906 Sante International SA Sos. Mihai Bravu nr. 7, bl. P37-P37A, sector 2 021303 Bucuresti Romania

497270 Sante International SA Str. Pupitrului, nr. 81, sect. 3 033036 Bucuresti Romania

31050285 Sante International SA Calea Ghirodei, nr. 36 300327 Timisoara Romania

31050284 Sante International SA Calea Dorobantilor, nr. 111 400609 Cluj-Napoca Romania

31050283 Sante International SA Str. Lascar Catargi, nr. 37 700107 Iasi Romania Scope

Import, trade and storage of medical and laboratory equipment, disinfectants, laboratory reagents, cardiovascular surgery devices. Consulting for state and private medical units.

Storage of medical and laboratory equipment, disinfectants, laboratory reagents,cardiovascular surgery devices, service for medical and laboratory equipment. Consulting for state and private medical units.

Trade of medical and laboratory equipment, disinfectants, laboratory reagents, cardiovascular surgery devices, service for medical and laboratory equipment. Consulting for state and private medical units.

Trade of medical and laboratory equipment, disinfectants, laboratory reagents, cardiovascular surgery devices, service for medical and laboratory equipment. Consulting for state and private medical units.

Trade of medical and laboratory equipment, disinfectants, laboratory reagents, cardiovascular surgery devices, service for medical and laboratory equipment. Consulting for state and private medical units.



This annex (edition:2021-06-16) is only valid in connection with the above-mentioned certificate.