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ORDIN DE PLATA NR.: 91                                TIP.DOC. 1 :
                                DATA EMITERII:21 decembrie 2021 :
=====:
PLATITI: 200000-00                                LEI: Doua Sute Mii lei 00 bani :
:
:
=====:
PLATITOR: (R) S.C. "OXIVI          CONTUL DE PLATI/CODUL IBAN :
T-MED" S.R.L.                                MD44ML000000002251729503 :
                                CODUL FISCAL :1007600044280 / :
:
:
=====:
PRESTATORUL PLATITOR                                CODUL BANCII:
BC"Moldindconbank"S.A. suc."Invest" Chisinau        :MOLDMD2X329:
=====:
BENEFICIAR (R) Centrul pen          CONTUL DE PLATI/CODUL IBAN :
tru Achizi?ii Publice Central MD23TRPCCC518430B01859AA :
izate in Sanatate                                CODUL FISCAL :1016601000212 / :
:
:
=====:
PRESTATORUL BENEFICIAR                                CODUL BANCII:
Ministerul Finantelor - Trezoreria de Stat          :TREZMD2X :
=====:
DESTINATIA PLATII:/P102/200000,00 Garan?:          TIPUL TRANSFERULUI :
ia pentru oferta la procedura de achizi?:          NORMAL/URGENT :N:
ie publica nr.ocds-b3wdpl-MD-16373279033:          :
23 din 21.12.21 Achizi?ionarea consumabi:          :
lelor conf. necesita?ilor IMSP SCR ?i IM:          :
SP INN pentru anul 2022                            :
: L.S. :
=====:
                                CODUL TRANZACTIEI:101:
DATA PRIMIRII:21/12/2021                            : SEMNATURILE
DATA EXECUTARII:                                    : EMITENTULUI
-----:
CONDUCTOR:Web Kojevnikov Dmitrii
MIIGfAYJKoZiHvcNAQcCoIIGbTCCBmkCAQExCzAJBgUrDgMCGGUAMAsGCSqGSIB3:
DQEHAAcCBUIUwggSBMIIDaaADAgECAhNHAACEjCA/4xcrKCbFAAAAAISMMA0GCSqG:
SIB3DQEBcWUAMCIxIDAeBgNVBAMTF0NFULQxLUNBLU1vbGRpbmRjb25iYW5rMB4X:
DTIwMDMxNjA4NDUwM1oXDTIzMDMxNjA4NTUwM1owgBgxCzAJBgNVBAYTAklEMRow:
YDVQOIEExFSZSXB1YmXpY2EgTW9sZG92YTERMA8GA1UEBxMIQ2hpc2luYXUxZjZAV :
-----:
                                (semnatura electronica)
CONTABIL-SEF:Web Kojevnikov Dmitrii
MIIGfAYJKoZiHvcNAQcCoIIGbTCCBmkCAQExCzAJBgUrDgMCGGUAMAsGCSqGSIB3:
DQEHAAcCBUIUwggSBMIIDaaADAgECAhNHAACEjCA/4xcrKCbFAAAAAISMMA0GCSqG:
SIB3DQEBcWUAMCIxIDAeBgNVBAMTF0NFULQxLUNBLU1vbGRpbmRjb25iYW5rMB4X:
DTIwMDMxNjA4NDUwM1oXDTIzMDMxNjA4NTUwM1owgBgxCzAJBgNVBAYTAklEMRow:
YDVQOIEExFSZSXB1YmXpY2EgTW9sZG92YTERMA8GA1UEBxMIQ2hpc2luYXUxZjZAV :
-----:
L.S.                                (semnatura electronica)
CONDUCTOR:
                                (semnatura manuala)
CONTABIL-SEF:
                                (semnatura manuala)
SEMNATURA PRESTATORUL          L.S.
:
MOTIVUL REFUZULUI          : L.S.
-----:

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**CERTIFICAT**  
**privind lipsa sau existența restanțelor față de bugetul public național**

Nr.  
№ **A2121595**

din  
от **16.12.2021**

**1. Destinația / Назначение**

AGENȚIA ACHIZIȚII PUBLICE

**2. Date despre contribuabil / Информация о налогоплательщике**

Denumirea Наименование	Codul fiscal / Numărul de identificare Фискальный код / Идентификационный номер
S.C. OXIVIT-MED S.R.L.	1007600044280
Adresa sediului de bază (strada, numărul) Адрес основного месторасположения (улица, номер)	Codul - Denumirea localității Код - Наименование населенного пункта
Decebal bd. nr.82 of.90	0110-SEC.BOTANICA

**3. Atestarea lipsei sau existenței restanțelor conform datelor Sistemului Informațional Automatizat /  
Подтверждение отсутствия или наличия недоимки согласно данных Информационной автоматизированной системы**

La data emiterii prezentului certificat restanța față de bugetul public național constituie/ На дату выдачи данной справки недоимка перед национальным публичным бюджетом составляет: **0,00 lei/лей.**

**4. Valabil pînă la / Действителен до 31.12.2021**

**5. Autentificarea Serviciului Fiscal de Stat / Подтверждение Государственной налоговой службы**

ȘEF DDE BOTANICA

Funcția/Dолжность

L.Ș/ М.П.

Executor: Ginga  
Numele și prenumele/Фамилия и имя



ANA STOICOV

Numele și prenumele/Фамилия и имя

Este extras din Sistemul Informațional al SFS SIA „Contul curent al contribuabilului”// 16.12.2021 ora 10:44:50  
cu aplicarea prevederilor pct. 82-83 Ordin IFPS nr.400 din 14.03.2014 (Monitorul Oficial 72-77/399, 28.03.2014)

NOTA (0,00)

Nr. 12101-504

18.03.2016

**CERTIFICAT  
PRIVIND EXISTENTA CONTURILOR CURENTE**

Prin prezentul, **BC „Mobiasbancă – Groupe Societe Generale” S.A.**, codul băncii (BIC): **MOBBMD22**, confirmă că compania **OXIVIT-MED SRL**, cod fiscal (IDNO) **1007600044280**, deține următoarele conturi curente la BC "Mobiasbancă-Groupe Societe Generale" S.A., Filiala. 1 Stejaur :

1. **MDL - 2224710SV23488147100; IBAN- MD09MO2224ASV23488147100**
2. **EUR - 2224710SV22227957100; IBAN- MD17MO2224ASV22227957100**
3. **USD - 2224710SV22214937100; IBAN- MD86MO2224ASV22214937100**

Certificatul este emis în baza cererii întreprinderii: Oxivit-Med SRL.

  
Dumitru Popa  
Director filială „Stejaur”



Executor : Mariana Guzun  
Tel: 022 812 614

Filiala Nr. 1 „Stejaur”  
Bd. Ștefan cel Mare și Sfânt 196  
MD-2004, Chișinău, Moldova  
Cod MOBBMD22  
Cont de corespondență 35213892  
la Centrul de Decontări al BNM

Tel. +373 22 81 26 15  
Fax. +373 22 81 26 15  
www.mobiasbanca.md

BC „Mobiasbancă – Groupe Société Générale” SA  
Capital Social: 100 000 000 MDL  
Număr de înregistrare de stat - 1002600006089  
Sediul Central:  
bd. Ștefan cel Mare și Sfânt 81a  
MD-2012, Chișinău, Moldova

REPUBLICA



MOLDOVA

# CERTIFICAT DE ÎNREGISTRARE

**Societatea Comercială "OXIVIT-MED" S.R.L.**  
ESTE ÎNREGISTRATĂ LA CAMERA ÎNREGISTRĂRII DE STAT

*Numărul de identificare de stat - codul fiscal*  
**1007600044280**

*Data înregistrării*

**30.07.2007**

*Data eliberării*

**30.07.2007**

**Bordeianu Tatiana, registrator de stat**

*Funcția, numele, prenumele persoanei  
care a eliberat certificatul*

*semnătura*

MD 0067985





**I.P. "AGENȚIA SERVICII PUBLICE"**

Departamentul înregistrare și licențiere a unităților de drept

**EXTRAS**  
**din Registrul de stat al persoanelor juridice**

nr. 8871 din 05.05.2021

Denumirea completă: **Societatea Comercială «OXIVIT-MED» S.R.L.**

Denumirea prescurtată: **S.C. «OXIVIT-MED» S.R.L.**

Forma juridică de organizare: **Societate cu Răspundere Limitată.**

Numărul de identificare de stat și codul fiscal: **1007600044280.**

Data înregistrării de stat: **30.07.2007.**

Sediul: **MD-2032, bd. Decebal, 82, ap.(of.) 90, mun. Chișinău, Republica Moldova.**

Modul de constituire: **nou creată.**

Obiectul principal de activitate:

- 1 Importul, fabricarea, comercializarea, asistența tehnică și (sau) reparația dispozitivelor medicale și (sau) a opticii;**
- 2 Comerțul cu ridicata al parfumurilor și produselor cosmetice;**
- 3 Comerțul cu amănuntul al produselor cosmetice și de parfumerie, articolelor de toaletă;**
- 4 Intermedieri pentru vânzarea unui asortiment larg de mărfuri;**
- 5 Alte tipuri de comerț cu amănuntul în magazine nespecializate;**
- 6 Alte tipuri de comerț cu ridicata;**
- 7 Închirierea altor mașini și echipamente.**

Capitalul social: **5400 lei.**

**Administrator: KOJEVNIKOV DMITRII, IDNP 0972305012362,**

Asociați:

**1. KOJEVNIKOV DMITRII , IDNP 0972305012362**

**cota 5400.00 lei, ce constituie 100 %.**

Prezentul extras este eliberat în temeiul art. 34 al Legii nr. 220-XVI din 19 octombrie 2007 privind înregistrarea de stat a persoanelor juridice și a întreprinzătorilor individuali și confirmă datele din Registrul de stat la data de: 05.05.2021.

Specialist coordonator  
tel. 022-207-840

Lazari Aliona



EEI 0354094

# OXIVIT MED

c/f: 1007600044280; adresa: str. Decebal 82-90, or. Chişinău, Republica Moldova  
telefon: + 373 22 808002; fax: + 373 22 808003  
web: www.oxivit-med.com; e-mail: info@oxivit-med.com

## **Lista fondatorilor companiei SRL „Oxivit-Med”**

Nr.	Numele, Prenumele	Codul Personal
1	Kojevnikov Dmitrii	0972305012362

Data predăstării 31.05.2021 09:44:13

Anexa la SNC  
"Prezentarea situațiilor financiare"  
Aprobat de Ministerul Finanțelor  
al Republicii Moldova

**SITUAȚIILE FINANCIARE**  
pentru perioada 01.01.2020 - 31.12.2020

Entitatea: S.C. OXVIT-MED S.R.L.  
Cod CUIU: 40424951  
Cod IDNO: 1007600044280

Sediul:  
MD:  
Raiounul(municipiul): 103\_DOF\_BOTANICA  
Cod CUATM: 0110\_SEC\_BOTANICA  
Strada: Decebal bd. nr.82 of.90

Activitatea principală: 63774\_Comerț cu amănuntul al articolelor medicale și ortopedice, în magazine specializate  
Forma de proprietate: 15\_Proprietate privată  
Forma organizatorico-juridică: 530\_Societate cu răspundere limitată

Date de contact:  
Telefon: +37322808002  
WEB:  
E-mail: oxvit-medical@gmail.com  
Numele și coordonatele al contabilului-șef: DI (dna) Kojevnikov Dmitrii Tel. 069200308

Numărul mediu al salariaților în perioada de gestiune: 4 persoane.

Persoanele responsabile de semnarea situațiilor financiare\* Kojevnikov Dmitrii

Unitatea de măsură: leu

**BILANȚUL**

la 31.12.2020

Anexa 1

Nr. cpt.	Indicatori	Cod rd.	Sold la	
			Începutul perioadei de gestiune	Sfârșitul perioadei de gestiune
1	2	3	4	5
<b>A C T I V</b>				
<b>ACTIVE IMOBILIZATE</b>				
<b>I. Imobilizări necorporale</b>				
1.	Imobilizări necorporale în curs de execuție	010		
2.	Imobilizări necorporale în exploatare, total	020	1787	1137
din care:				
2.1.	concesiuni, licențe și mărci	021	1787	1137
2.2.	drepturi de autor și titluri de protecție	022		
2.3.	programe informatice	023		
2.4.	alte imobilizări necorporale	024		
3.	Fond comercial	030		
4.	Avansuri acordate pentru imobilizări necorporale	040		
	<b>Total imobilizări necorporale</b> (rd.010 + rd.020 + rd.030 + rd.040)	050	1787	1137
<b>II. Imobilizări corporale</b>				
1.	Imobilizări corporale în curs de execuție	060		
2.	Terenuri	070		
3.	Mijloace fixe, total	080	2234	9980
din care:				
3.1.	clădiri	081		
3.2.	construcții speciale	082		
3.3.	mașini, utilaje și instalații tehnice	083		9235
3.4.	mijloace de transport	084		

A.	3.5. inventar și mobilier	085			
	3.6. alte mijloace fixe	086	2234	745	
	4. Resurse minerale	090			
	5. Active biologice imobilizate	100			
	6. Investiții imobiliare	110			
	7. Avansuri acordate pentru imobilizări corporale	120			
	<b>Total imobilizări corporale</b> (rd.060 + rd.070 + rd.080 + rd.090 + rd.100 + rd.110 + rd.120)	130	2234	9980	
	<b>III. Investiții financiare pe termen lung</b>				
	1. Investiții financiare pe termen lung în părți nefiliate	140			
	2. Investiții financiare pe termen lung în părți afiliate, total	150			
din care:					
2.1. acțiuni și cote de participație deținute în părțile afiliate	151				
2.2. Împrumuturi acordate părților afiliate	152				
2.3. Împrumuturi acordate aferente intereselor de participație	153				
2.4. alte investiții financiare	154				
<b>Total investiții financiare pe termen lung</b> (rd.140 + rd.150)	160				
<b>IV. Creațe pe termen lung și alte active imobilizate</b>					
1. Creațe comerciale pe termen lung	170				
2. Creațe ale părților afiliate pe termen lung	180				
Inclusiv: creațe aferente intereselor de participație	181				
3. Alte creațe pe termen lung	190				
4. Cheltuieli anticipate pe termen lung	200				
5. Alte active imobilizate	210				
<b>Total creațe pe termen lung și alte active imobilizate</b> (rd.170 + rd.180 + rd.190 + rd.200 + rd.210)	220				
<b>TOTAL ACTIVE IMOBILIZATE</b> (rd.050 + rd.130 + rd.160 + rd.220)	230	4021	11117		
<b>ACTIVE CIRCULANTE</b>					
<b>I. Stocuri</b>					
1. Materiale și obiecte de mică valoare și scurtă durată	240	434	617		
2. Active biologice circulante	250				
3. Producția în curs de execuție	260				
4. Produse și mărfuri	270	8318982	6895348		
5. Avansuri acordate pentru stocuri	280				
<b>Total stocuri</b> (rd.240 + rd.250 + rd.260 + rd.270 + rd.280)	290	8319416	6895965		
<b>II. Creațe curente și alte active circulante</b>					
1. Creațe comerciale curente	300	4555544	17423930		
2. Creațe ale părților afiliate curente	310				
Inclusiv: creațe aferente intereselor de participație	311				
3. Creațe ale bugetului	320	491901	1593996		
4. Creațele ale personalului	330		1452		
5. Alte creațe curente	340				
6. Cheltuieli anticipate curente	350	8082	6076		
7. Alte active circulante	360	1601496	3786977		
<b>Total creațe curente și alte active circulante</b> (rd.300 + rd.310 + rd.320 + rd.330 + rd.340 + rd.350 + rd.360)	370	6667023	22812431		
<b>III. Investiții financiare curente</b>					
1. Investiții financiare curente în părți nefiliate	380				
2. Investiții financiare curente în părți afiliate, total	390				
din care:					
2.1. acțiuni și cote de participație deținute în părțile afiliate	391				
2.2. Împrumuturi acordate părților afiliate	392				
2.3. Împrumuturi acordate aferente intereselor de participație	393				

B.	din care:			
	2.1. Împrumuturi din emisiunea de obligațiuni	721		
	Inclusiv: Împrumuturi din emisiunea de obligațiuni convertibile	722		
	2.2. alte împrumuturi pe termen scurt	723		
	3. Datorii comerciale curente	730	6729208	15784405
	4. Datorii față de părțile afiliate curente	740		
	Inclusiv: datorii aferente intereselor de participație	741		
	5. Avansuri primite curente	750	336114	349631
	6. Datorii față de personal	760	73257	116957
	7. Datorii privind asigurările sociale și medicale	770		
8. Datorii față de buget	780	300		
9. Datorii față de proprietari	790			
10. Venituri anticipate curente	800			
11. Alte datorii curente	810	27875	33023	
<b>TOTAL DATORII CURENTE</b> (rd.710 + rd.720 + rd.730 + rd.740 + rd.750 + rd.760 + rd.770 + rd.780 + rd.790 + rd.800 + rd.810)	820	7166754	16284016	
<b>PROVIZIOANE</b>				
1. Provizioane pentru beneficiile angajaților	830			
2. Provizioane pentru garanții acordate cumpărătorilor/clientșilor	840			
3. Provizioane pentru impozite	850			
4. Alte provizioane	860			
<b>TOTAL PROVIZIOANE</b> (rd.830 + rd.840 + rd.850 + rd.860)	870			
<b>TOTAL PASIVE</b> (rd.620 + rd.700 + rd.820 + rd.870)	880	28730633	41305620	

**SITUAȚIA DE PROFIT ȘI PIERDERE**  
de la 01.01.2020, până la 31.12.2020

Anexa 2

Indicatori	Cod rd.	Perioada de gestiune	
		precedenta	curenta
1	2	3	4
Venituri din vânzări, total	010	56125772	61054881
din care:			
venituri din vânzarea produselor și mărfurilor	011	56125772	61054881
venituri din prestarea serviciilor și executarea lucrărilor	012		
venituri din contracte de construcție	013		
venituri din contracte de leasing	014		
venituri din contracte de microfinanțare	015		
alte venituri din vânzări	016		
Costul vânzărilor, total	020	44624941	50207602
din care:			
valoarea contabilă a produselor și mărfurilor vândute	021	44624941	50207602
costul serviciilor prestate și lucrărilor executate terților	022		
costuri aferente contractelor de construcție	023		
costuri aferente contractelor de leasing	024		
costuri aferente contractelor de microfinanțare	025		
alte costuri aferente vânzărilor	026		
<b>Profit brut (pierdere brută)</b> (rd.010 - rd.020)	030	11500831	10847279
Alte venituri din activitatea operațională	040	1954484	1967064
Cheltuieli de distribuție	050	46711	68333
Cheltuieli administrative	060	668046	995848
Alte cheltuieli din activitatea operațională	070	161844	34858
<b>Rezultatul din activitatea operațională: profit (pierdere)</b> (rd.030 + rd.040 - rd.050 - rd.060 - rd.070)	080	12578714	11715304

C.	2.4. alte investiții financiare în părți afiliate	394			
	<b>Total investiții financiare curente</b> (rd.380 + rd.390)	400			
	<b>IV. Numerar și documente bănești</b>	410	13740173	11586107	
	<b>TOTAL ACTIVE CIRCULANTE</b> (rd.290 + rd.370 + rd.400 + rd.410)	420	28726612	41294503	
	<b>TOTAL ACTIVE</b> (rd.230 + rd.420)	430	28730633	41305620	
	<b>P A S I V</b>				
	<b>CAPITAL PROPRIU</b>				
	<b>I. Capital social și neînregistrat</b>				
	1. Capital social	440	5400	5400	
	2. Capital nevărsat	450	( )	( )	
3. Capital neînregistrat	460				
4. Capital retras	470	( )	( )		
5. Patrimoniul primit de la stat cu drept de proprietate	480				
<b>Total capital social și neînregistrat</b> (rd.440 + rd.450 + rd.460 + rd.470 + rd.480)	490	5400	5400		
<b>II. Prime de capital</b>					
<b>III. Rezerve</b>					
1. Capital de rezervă	510				
2. Rezerve statutare	520				
3. Alte rezerve	530				
<b>Total rezerve</b> (rd.510 + rd.520 + rd.530)	540				
<b>IV. Profit (pierdere)</b>					
1. Corecții ale rezultatelor anilor precedenți	550	X			
2. Profit nerepartizat (pierdere neacoperită) al anilor precedenți	560	21481849	14484236		
3. Profit net (pierdere netă) al perioadei de gestiune	570	X	10455338		
4. Profit utilizat al perioadei de gestiune	580	X	( )		
<b>Total profit (pierdere)</b> (rd.550 + rd.560 + rd.570 + rd.580)	590	21481849	24939574		
<b>V. Rezerve din reevaluare</b>					
600					
<b>VI. Alte elemente de capital propriu</b>					
610					
<b>TOTAL CAPITAL PROPRIU</b> (rd.490 + rd.500 + rd.540 + rd.600 + rd.610)	620	21487249	24944974		
<b>DATORII PE TERMEN LUNG</b>					
1. Credite bancare pe termen lung	630				
2. Împrumuturi pe termen lung	640	76630	76630		
din care:					
2.1. Împrumuturi din emisiunea de obligațiuni	641				
Inclusiv: Împrumuturi din emisiunea de obligațiuni convertibile	642				
2.2. alte împrumuturi pe termen lung	643	76630	76630		
3. Datorii comerciale pe termen lung	650				
4. Datorii față de părțile afiliate pe termen lung	660				
Inclusiv: datorii aferente intereselor de participație	661				
5. Avansuri primite pe termen lung	670				
6. Venituri anticipate pe termen lung	680				
7. Alte datorii pe termen lung	690				
<b>TOTAL DATORII PE TERMEN LUNG</b> (rd.630 + rd.640 + rd.650 + rd.660 + rd.670 + rd.680 + rd.690)	700	76630	76630		
<b>DATORII CURENTE</b>					
1. Credite bancare pe termen scurt	710				
2. Împrumuturi pe termen scurt, total	720				

Venituri financiare, total	090			1752762
din care:				
venituri din interese de participare	091			
inclusiv: veniturile obținute de la părțile afiliate	092			
venituri din dobânzi	093			
inclusiv: veniturile obținute de la părțile afiliate	094			
venituri din alte investiții financiare pe termen lung	095			
inclusiv: veniturile obținute de la părțile afiliate	096			
venituri aferente ajustărilor de valoare privind investițiile financiare pe termen lung și curente	097			
venituri din ieșirea investițiilor financiare	098			
venituri aferente diferențelor de curs valutar și de sumă	099			1752762
Cheltuieli financiare, total	100		20827	1580853
din care:				
cheltuieli privind dobânzile	101			
inclusiv: cheltuielile aferente părților afiliate	102			
cheltuieli aferente ajustărilor de valoare privind investițiile financiare pe termen lung și curente	103			
cheltuieli aferente ieșirii investițiilor financiare	104			
cheltuieli aferente diferențelor de curs valutar și de sumă	105		20827	1580853
<b>Rezultatul: profit (pierdere) financiar(ă)</b> (rd.090 - rd.100)	110		-20827	171909
Venituri cu active immobilizate și excepționale	120			
Cheltuieli cu active immobilizate și excepționale	130			
<b>Rezultatul din operațiuni cu active immobilizate și excepționale: profit (pierdere)</b> (rd.120 - rd.130)	140			
<b>Rezultatul din alte activități: profit (pierdere)</b> (rd.110 + rd.140)	150		-20827	171909
<b>Profit (pierdere) până la impozitare</b> (rd.080 + rd.150)	160		12557887	11887213
Cheltuieli privind impozitul pe venit	170		1543063	1431875
<b>Profit net (pierdere netă) al perioadei de gestiune</b> (rd.160 - rd.170)	180		11014824	10455338

**SITUAȚIA MODIFICĂRILOR CAPITALULUI PROPRIU**  
de la până la

Anexa 3

Nr. din	Indicatori	Cod rd	Sold la începutul perioadei de gestiune	Majorări	Diminuări	Sold la sfârșitul perioadei de gestiune
1	2	3	4	5	6	7
	<b>Capital social și nelregistrat</b>					
	1. Capital social	010				
	2. Capital nevărsat	020	( )	( )	( )	( )
	3. Capital nelregistrat	030				
I.	4. Capital retras	040	( )	( )	( )	( )
	5. Patrimoniul primit de la stat cu drept de proprietate	050				
	<b>Total capital social și nelregistrat</b> (rd.010 + rd.020 + rd.030 + rd.040 + rd.050)	060				
II.	<b>Prime de capital</b>	070				
	<b>Rezerve</b>					
	1. Capital de rezervă	080				
III.	2. Rezerve statutare	090				
	3. Alte rezerve	100				
	<b>Total rezerve</b> (rd.080 + rd.090 + rd.100)	110				
	<b>Profit (pierdere)</b>					
	1. Corecții ale rezultatelor anilor precedenți	120	X			

IV.	2. Profit nerepartizat (pierdere neacoperită) al anilor precedenți	130				
	3. Profit net (pierdere netă) al perioadei de gestiune	140	X			
	4. Profit utilizat al perioadei de gestiune	150	X	( )	( )	( )
	<b>Total profit (pierdere)</b> (rd.120 + rd.130 + rd.140 + rd.150)	160				
V.	<b>Rezerve din reevaluare</b>	170				
VI.	<b>Alte elemente de capital propriu</b>	180				
	<b>Total capital propriu</b> (rd.060 + rd.070 + rd.110 + rd.160 + rd.170 + rd.180)	190				

**SITUAȚIA FLUXURILOR DE NUMERAR**  
de la până la

Anexa 4

Indicatori	Cod rd	Perioada de gestiune	
		precedentă	curentă
1	2	3	4
<b>Fluxuri de numerar din activitatea operațională</b>			
Încasări din vânzări	010		
Plăți pentru stocuri și servicii procurate	020		
Plăți către angajați și organe de asigurare socială și medicală	030		
Dobânzi plătite	040		
Plata impozitului pe venit	050		
Alte încasări	060		
Alte plăți	070		
<b>Fluxul net de numerar din activitatea operațională</b> (rd.010 - rd.020 - rd.030 - rd.040 - rd.050 + rd.060 - rd.070)	080		
<b>Fluxuri de numerar din activitatea de investiții</b>			
Încasări din vânzarea activelor immobilizate	090		
Plăți aferente intrărilor de active immobilizate	100		
Dobânzi încasate	110		
Dividende încasate	120		
inclusiv: dividende încasate din străinătate	121		
Alte încasări (plăți)	130		
<b>Fluxul net de numerar din activitatea de investiții</b> (rd.090 - rd.100 + rd.110 + rd.120 ± rd.130)	140		
<b>Fluxuri de numerar din activitatea financiară</b>			
Încasări sub formă de credite și împrumuturi	150		
Plăți aferente rambursării creditelor și împrumuturilor	160		
Dividende plătite	170		
inclusiv: dividende plătite nereșidenților	171		
Încasări din operațiuni de capital	180		
Alte încasări (plăți)	190		
<b>Fluxul net de numerar din activitatea financiară</b> (rd.150 - rd.160 - rd.170 + rd.180 ± rd.190)	200		
<b>Fluxul net de numerar total</b> (± rd.080 ± rd.140 ± rd.200)	210		
Diferențe de curs valutar favorabile (nefavorabile)	220		
<b>Sold de numerar la începutul perioadei de gestiune</b>	230		
<b>Sold de numerar la sfârșitul perioadei de gestiune</b> (± rd.210 ± rd.220 + rd.230)	240		

**Documente atașate - Notă explicativă (fișierul pdf)**

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**Расписка 2**

Респондент

Фискальный код: 1007600044280, наименование: S.C. OXIVIT-MED S.R.L.

Предоставил отчет: RSE1\_21

На фискальный период: A/2020

Дата предоставления: 31.05.2021

Временная метка отчёта зарегистрированного в Информационной Системе НБС : 31.05.2021 10:20:49

National Bureau of Statistics (NBS) received the electronic version of the report, sent by you. The data provided is verified by NBS.



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## Расписка

Респондент

Фискальный код: 1007600044280, наименование: S.C. OXIVIT-MED S.R.L.

Предоставил отчёт: RSF1\_21

На фискальный период: A/2020

Дата предоставления: 31.05.2021

Временная метка отчёта зарегистрированного в Системе Электронной Отчётности и отправленного в Информационную Систему БНС : 31.05.2021 09:44:13

# AORTIC PERIPHERAL AND VENOUS PRODUCT CATALOGUE

AORTIC



PERIPHERAL

VENOUS

# AORTIC CONTENTS

---

## STENT GRAFTS

Endurant™ II/IIIs



Talent™ Occluder



Valiant™ Navion™



Valiant™ Captivia™



## ENDOANCHOR™ SYSTEMS

Heli-Fx™ / Heli-Fx™ TAA



## ANCILLARY

Sentrant™



Reliant™



TourGuide™



# PERIPHERAL CONTENTS 1/3

## DRUG COATED BALLOONS

IN.PACT™ Admiral™



IN.PACT™ Pacific™



## STENT SYSTEMS

Protégé™ Rx™



VisiPro™



Protégé™ GPS™



EverFlex™



EverFlex™ with Entrust™  
Delivery System



Paramount Mini™ GPS™



Hippocampus™



IntraStent™ LD



## PTA BALLOONS

Admiral™ Xtreme™



EverCross™



Fortrex™



Pacific™ Plus



Pacific™ Extreme



Submarine™ Rapido



Amphirion™ Deep



NanoCross™ Elite



RapidCross™



Chocolate™



BACK

NEXT



# PERIPHERAL CONTENTS 2/3

## DIRECTIONAL ATHERECTOMY

HawkOne™



TurboHawk™



SilverHawk™



## EMBOLIC PROTECTION DEVICES

Mo.Ma™ Ultra



SpiderFX™



## CROSSING CATHETERS

TrailBlazer™  
Support Catheter



TrailBlazer™  
Angled Support Catheter



## CTO DEVICES

Viance™



Enteer™



## CATHETERS

Piton™ GC



Rebar™



## THROMBUS MANAGEMENT

Cragg-McNamara™



MicroMewi™



ProStream™



BACK

NEXT



# PERIPHERAL CONTENTS 3/3

## GUIDEWIRES

Nitrex™



Babywire™



AqWire™



Wholey™



Kitewire™ Deep



## SNARES

Amplatz GooseNeck™  
Snare Kit



Amplatz GooseNeck™  
MicroSnare Kit



## Y-CONNECTORS

Bigeasy™



Sequel™



## VASCULAR EMBOLIZATION

Onyx™



Onyx™ 34L



Onyx™ Mixer



Onyx™ Syringe Catheter  
Interface Adapter



1ml Luer-Lock  
Injection Syringe



Concerto™ Helix/3D



I.D. Instant Detacher



MVP™



BACK

NEXT



# VENOUS CONTENTS

---

## ABRE™ VENOUS STENT

Abre™



## CLOSUREFAST™ PROCEDURE

ClosureFast™



ClosureRFS™



ClosureRFG™



## PROCEDURE ACCESSORIES

Procedure Packs



Tumescent Infiltration Pump



Ultrasound



## VENASEAL™ SYSTEMS

VenaSeal™



BACK

# AORTIC



AORTIC

PERIPHERAL

VENOUS



# STENT GRAFTS



AORTIC



PERIPHERAL

VENOUS

# Endurant™ II/IIIs

## AAA Stent Graft System

### Features\*

#### Complete conformability, optimal seal

- M-shaped proximal stents provide wall apposition and a short sealing zone
- Suprarenal stent anchor pins provide secure fixation
- Limb stent and stent spacing reduce kinking

#### Total control, consistent precision

- Tip capture mechanism allows for precise positioning adjustments
- Back-end thumb wheel provides controlled release of the suprarenal stent and anchor pins
- Improved radiopacity provides increased visibility†
- Four proximal markers assist in accurate deployment
- E-shaped marker assists with A/P orientation

#### Durable build, dependable performance

- Ultra-high molecular weight polyethylene sutures are three times stronger than surgical sutures
- High-density multifilament polyester graft material provides low porosity
- Electropolished nitinol stents improve fatigue resistance

#### Expanded anatomical customization with Endurant™ IIIs

- Endurant™ IIIs complements the Endurant™ II AAA Stent Graft System

#### Low profile, easy access

- Low profile and hydrophilic coating enhance access and trackability
- Flexible, kink-resistant delivery system facilitates stent graft delivery

\*Test data on file at Medtronic. Bench test results may not be indicative of clinical performance.

†Contralateral gate marker



# Endurant™ II/IIs

## AAA Stent Graft System

### ENDURANT™ II PRODUCT CODE DESCRIPTION

ET	B	F	23	13	C	124	EE	18
----	---	---	----	----	---	-----	----	----

Catheter Outer Diameter

Delivery System  
EE - Endurant™ II

Total Covered Length

Distal Design  
C - Closed Web

Distal Graft Diameter

Proximal Graft Diameter

Proximal Design

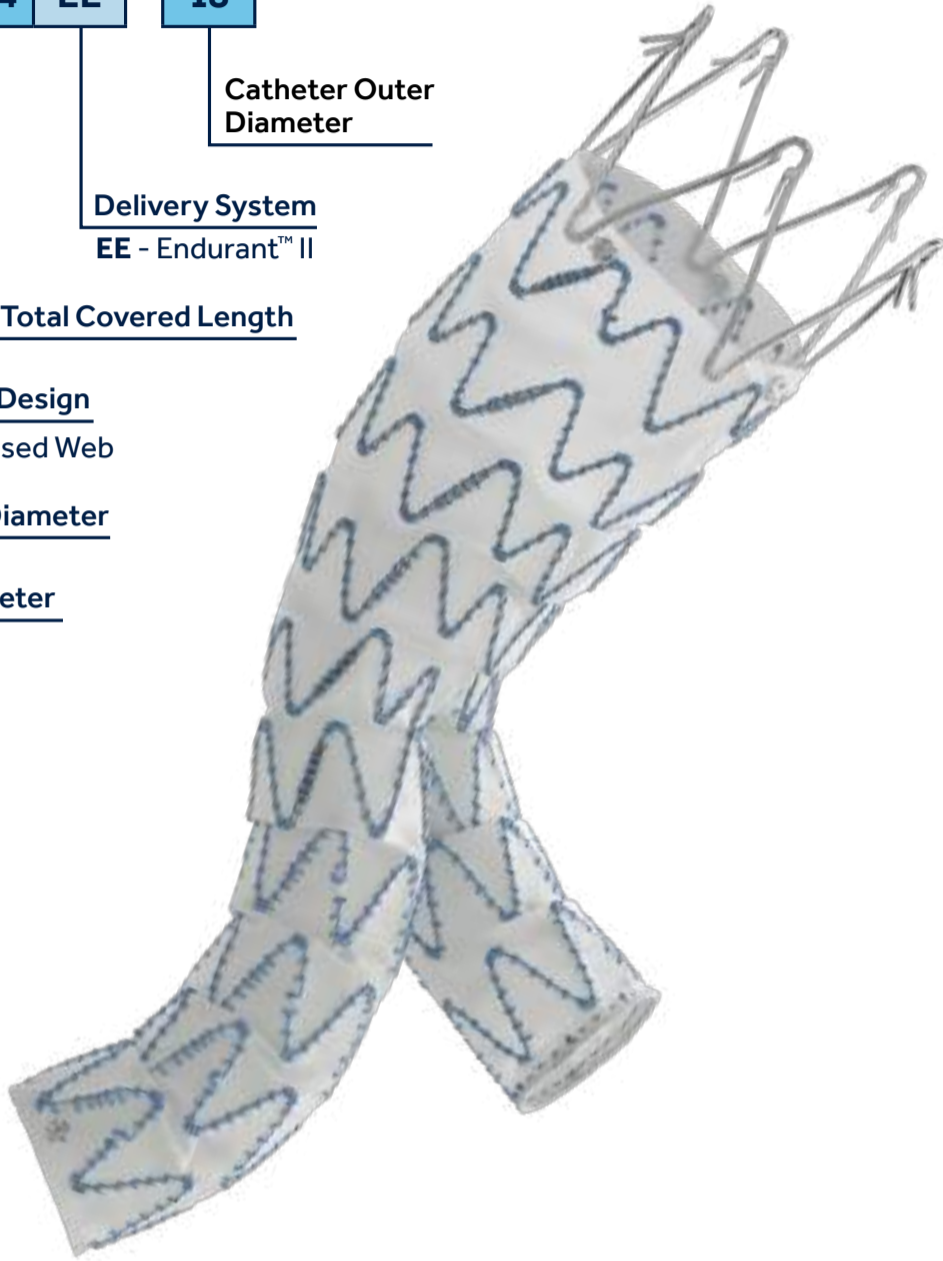
F - FreeFlo  
W - Open Web

Device Configuration

B - Bifurcated  
L - Contralateral Iliac Limb  
E - Iliac Extension  
C - Aortic Extension (Cuff)  
T - Abdominal Tube  
U - Aorto-Uni-Iliac (AUI)

Product Name

ET - Endurant™ II  
ES - Endurant™ IIs



### ENDURANT™ IIs BIFURCATIONS

Product Code

	Proximal Graft Diameter (mm)	Distal Graft Diameter (mm)	Distal Design	Total Covered Length (mm)	Delivery System	Catheter Outer Diameter (F)
ESBF	23	14	C	103	EE	18
ESBF	25	14	C	103	EE	18
ESBF	28	14	C	103	EE	18
ESBF	32	14	C	103	EE	20
ESBF	36	14	C	103	EE	20

# Endurant™ II/IIs

## AAA Stent Graft System

### ENDURANT™ II BIFURCATIONS

Product Code						
	Proximal Graft Diameter (mm)	Distal Graft Diameter (mm)	Distal Design	Total Covered Length (mm)	Delivery System	Catheter Outer Diameter (F)
ETBF	23	13	C	124	EE	18
ETBF	23	13	C	145	EE	18
ETBF	23	13	C	166	EE	18
ETBF	23	16	C	124	EE	18
ETBF	23	16	C	145	EE	18
ETBF	23	16	C	166	EE	18
ETBF	25	13	C	124	EE	18
ETBF	25	13	C	145	EE	18
ETBF	25	13	C	166	EE	18
ETBF	25	16	C	124	EE	18
ETBF	25	16	C	145	EE	18
ETBF	25	16	C	166	EE	18
ETBF	28	13	C	124	EE	18
ETBF	28	13	C	145	EE	18
ETBF	28	13	C	166	EE	18
ETBF	28	16	C	124	EE	18
ETBF	28	16	C	145	EE	18
ETBF	28	16	C	166	EE	18
ETBF	28	20	C	124	EE	18
ETBF	28	20	C	145	EE	18
ETBF	28	20	C	166	EE	18
ETBF	32	16	C	124	EE	20
ETBF	32	16	C	145	EE	20
ETBF	32	16	C	166	EE	20
ETBF	32	20	C	124	EE	20
ETBF	32	20	C	145	EE	20
ETBF	32	20	C	166	EE	20
ETBF	36	16	C	145	EE	20
ETBF	36	16	C	166	EE	20
ETBF	36	20	C	145	EE	20
ETBF	36	20	C	166	EE	20

# Endurant™ II/IIIs

## AAA Stent Graft System

### LIMBS\*

Product Code								
	Proximal Graft Diameter (mm)	Distal Graft Diameter (mm)	Distal Design	Total Covered Length (mm)	Delivery System	Catheter Outer Diameter (F)	Total Contralateral Covered Length with EII / EIIIs Bifurcated†	Total Ipsilateral Covered Length with EIIIs Bifurcated‡
ETLW	16	10	C	82	EE	14	136	155
ETLW	16	10	C	93	EE	14	147	166
ETLW	16	10	C	124	EE	14	178	177–197
ETLW	16	10	C	156	EE	16	210	209–229
ETLW	16	10	C	199	EE	16	253	252–272
ETLW	16	13	C	82	EE	14	136	155
ETLW	16	13	C	93	EE	14	147	166
ETLW	16	13	C	124	EE	14	178	177–197
ETLW	16	13	C	156	EE	16	210	209–229
ETLW	16	13	C	199	EE	16	253	252–272
ETLW	16	16	C	82	EE	14	136	135–155
ETLW	16	16	C	93	EE	14	147	146–166
ETLW	16	16	C	124	EE	14	178	177–197
ETLW	16	16	C	156	EE	16	210	209–229
ETLW	16	16	C	199	EE	16	253	252–272
ETLW	16	20	C	82	EE	16	136	155
ETLW	16	20	C	93	EE	16	147	166
ETLW	16	20	C	124	EE	16	178	177–197
ETLW	16	20	C	156	EE	16	210	209–229
ETLW	16	20	C	199	EE	16	253	252–272
ETLW	16	24	C	82	EE	16	136	155
ETLW	16	24	C	93	EE	16	147	166
ETLW	16	24	C	124	EE	16	178	177–197
ETLW	16	24	C	156	EE	16	210	209–229
ETLW	16	24	C	199	EE	16	253	252–272
ETLW	16	28	C	82	EE	16	136	155
ETLW	16	28	C	93	EE	16	147	166
ETLW	16	28	C	124	EE	16	178	177–197
ETLW	16	28	C	156	EE	16	210	209–229
ETLW	16	28	C	199	EE	16	253	252–272

\* The limb mates with the AUI stent graft on the ipsilateral side.

† These calculations assume the minimum 30 mm overlap between the bifurcated stent graft and the contralateral iliac limb per the Endurant™ II Stent Graft System Instructions For Use (IFU). When using the 124 mm length bifurcated stent graft, subtract 10 mm from total contralateral covered length with Bifurcated.

‡ The 3 – 5 stent overlap is available only with select limbs. Please refer to the Instructions For Use for more information. The contralateral iliac limb per the Endurant™ II Stent Graft System Instructions For Use. When using the 124 mm length bifurcated stent graft, subtract 10 mm from total contralateral covered length with Bifurcated.

# Endurant™ II/IIs

## AAA Stent Graft System

### ILIAC EXTENSIONS

Product Code						
	Proximal Graft Diameter (mm)	Distal Graft Diameter (mm)	Distal Design	Total Covered Length (mm)	Delivery System	Catheter Outer Diameter (F)
ETEW	10	10	C	82	EE	14
ETEW	13	13	C	82	EE	14
ETEW	20	20	C	82	EE	16
ETEW	24	24	C	82	EE	16
ETEW	28	28	C	82	EE	18

### AORTIC EXTENSIONS

Product Code						
	Proximal Graft Diameter (mm)	Distal Graft Diameter (mm)	Distal Design	Total Covered Length (mm)	Delivery System	Catheter Outer Diameter (F)
ETCF	23	23	C	49	EE	18
ETCF	25	25	C	49	EE	18
ETCF	28	28	C	49	EE	18
ETCF	32	32	C	49	EE	20
ETCF	36	36	C	49	EE	20

### ABDOMINAL TUBES

Product Code						
	Proximal Graft Diameter (mm)	Distal Graft Diameter (mm)	Distal Design	Total Covered Length (mm)	Delivery System	Catheter Outer Diameter (F)
ETTF	23	23	C	70	EE	18
ETTF	25	25	C	70	EE	18
ETTF	28	28	C	70	EE	18
ETTF	32	32	C	70	EE	20
ETTF	36	36	C	70	EE	20

### AUI

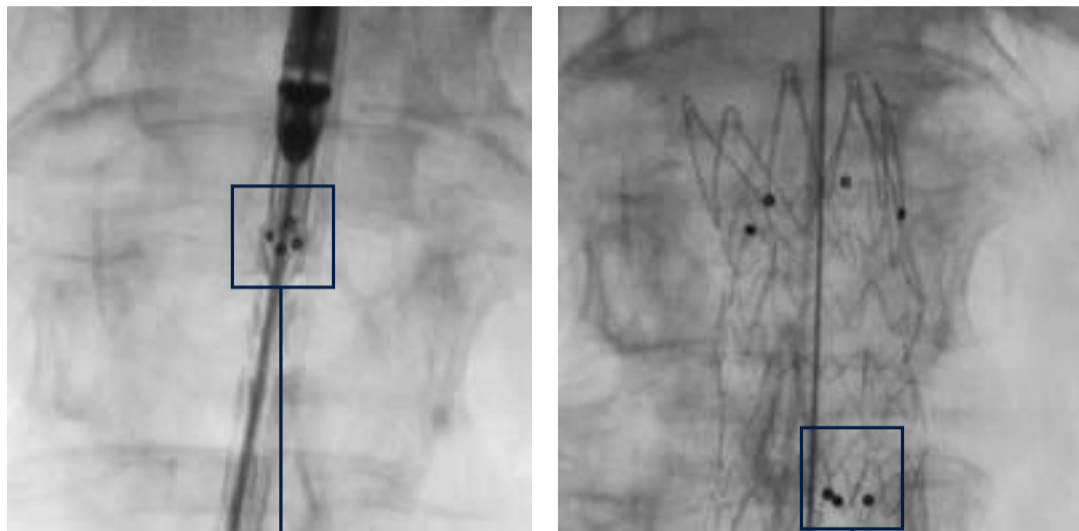
Product Code						
	Proximal Graft Diameter (mm)	Distal Graft Diameter (mm)	Distal Design	Total Covered Length (mm)	Delivery System	Catheter Outer Diameter (F)
ETUF	23	14	C	102	EE	18
ETUF	25	14	C	102	EE	18
ETUF	28	14	C	102	EE	18
ETUF	32	14	C	102	EE	20
ETUF	36	14	C	102	EE	20

# Endurant™ II/IIIs

## AAA Stent Graft System

### PLACEMENT AND SIZING GUIDELINES

Use the proximal radiopaque markers to position the top edge of the graft material.



RADIOPAQUE MARKERS

  
E-SHAPED MARKER  
ASSISTS  
WITH A / P  
ORIENTATION

For the contralateral side: The radiopaque markers at the proximal limb should be aligned with the radiopaque markers at the flow divider of the Endurant™ II or Endurant™ IIIs bifurs.

For the ipsilateral side: Depending on the limb configuration used, the radiopaque markers at the proximal end of the limb should be aligned to the distal radiopaque marker on the ipsilateral leg or the flow divider marker of the Endurant™ IIIs bifur.

Select limbs will allow a 3-5 stent overlap adjustment during the case.

Please refer to the Instructions for Use for more information as needed.

# Endurant™ II/IIs

## AAA Stent Graft System

### ENDURANT™ II/IIS STENT GRAFT SIZING GUIDELINES

Proximal Aortic Diameter (mm)		Proximal Stent Graft Size
Standard EVAR*	ChEVAR†	
19 - 20	n/a	23
21 - 22	19 - 20	25
23 - 25	21 - 23	28
26 - 28	24 - 26	32
29 - 32	27 - 30	36

Distal Iliac Diameter (mm)	Distal Stent Graft Size
8 - 9	10
10 - 11	13
12 - 14	16
15 - 18	20
19 - 22	24
23 - 25	28

Each Endurant™ II / IIs AAA Stent Graft System must be ordered in a size that is appropriate to fit the patient's anatomy.

Proper sizing of the Endurant™ II AAA Stent Graft System is the responsibility of the physician.

The above suggestions for stent graft diameters are based on vessel inner wall measurements.

\*EVAR: Bifurs, Cuff, AUI, Tube configuration

†Limb, Iliac Extension configuration



# Talent™ Occluder

## Endovascular Occluder



### Occluder Stent Graft

#### Double spring configuration

- Securely anchors in the iliac artery to seal the lumen and to prevent retrograde blood flow

#### Expand your options

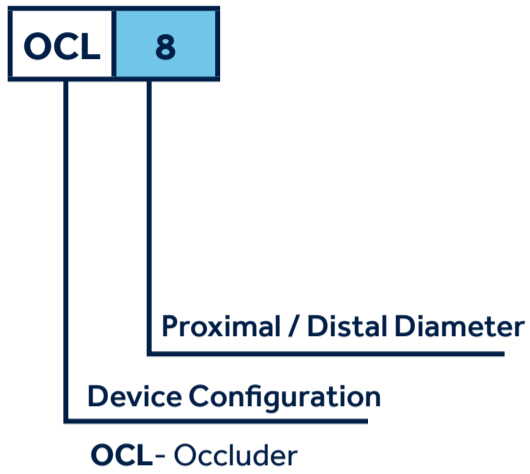
- The Occluder is a less invasive option to surgical ligation and is reported to have better results than coil embolization\*

\*Kato, et al. Use of a self-expanding vascular occluder for embolization during endovascular aortic aneurysm repair. JVIR 8:27-33, 1997.

# Talent™ Occluder

## Endovascular Occluder

### TALENT OCCLUDER / PRODUCT CODE DESCRIPTION



Product Code	Proximal / Distal Diameter (mm)	Total Length (mm)	Catheter Diameter (F)
OCL 8	8	31	17.5
OCL 10	10	31	17.5
OCL 12	12	31	17.5
OCL 14	14	33	17.5
OCL 16	16	33	17.5
OCL 18	18	33	17.5
OCL 20	20	35	17.5
OCL 22	22	35	17.5
OCL 24	24	35	17.5



### OCCLUDER SYSTEM COMPONENT

Native Vessel (mm)	Suggested Stent Graft Diameter (mm)	Oversizing (mm)
19-20	24	4-5
18	22	4
16-17	20	3-4
14-15	18	3-4
13	16	3
11-12	14	2-3
9-10	12	2-3
7-8	10	2-3
6	8	2

# Valiant Navion™

## Thoracic Stent Graft System

AORTIC

### The freedom to do more

Low profile, easy-to-use delivery system designed for expanded access with smooth navigation

### Features

#### Delivery System

- Tip Capture: for controlled delivery and deployment on both FreeFlo and CoveredSeal configurations
- Designed for simplified navigation: flexible, kink-resistant hydrophilic-coated catheter
- Shorter tapered tip designed to decrease vessel impact

#### Stent Graft

- Multi-filament thoracic graft material based on Endurant™ stent graft yarn designed for flexibility and superior permeability resistance
- Aligned stent peaks and valleys designed for increased flexibility throughout the stent graft
- Increased distance between stents designed to optimize migration resistance and conformability

#### Proven Platforms

- Leverages proven design of the Valiant™ and Endurant™ stent graft system platforms
- 100K+ thoracic and 300K abdominal patients treated\*
- Over 20 years of endovascular experience with deep clinical history



PERIPHERAL

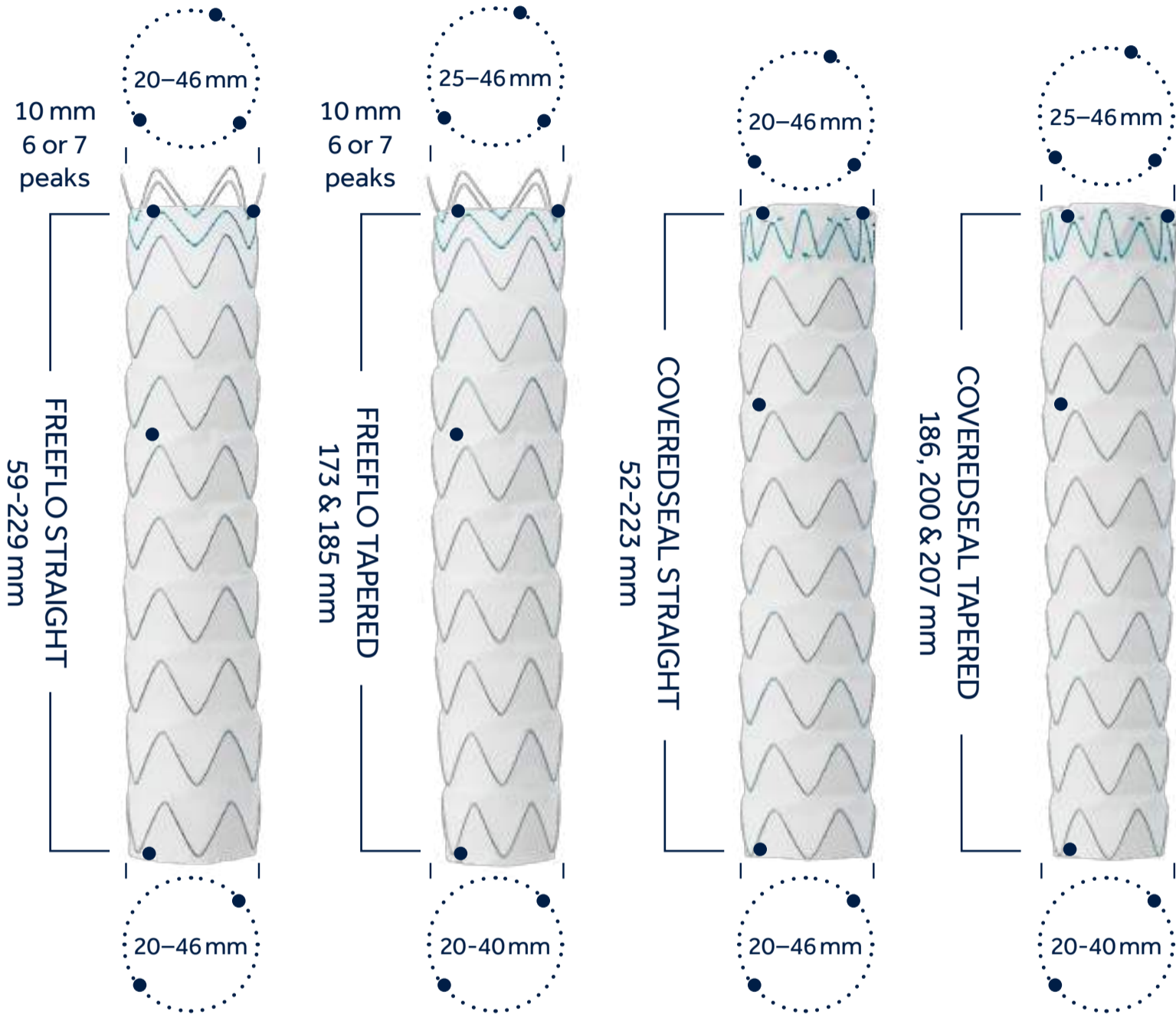
VENOUS

\* Data on file at Medtronic.

# Valiant Navion™

## Thoracic Stent Graft System

### COMPONENT GUIDE



### DISTINCT RADIOPAQUE MARKER

- Spherical RO Marker

# Valiant Navion™

## Thoracic Stent Graft System

Each Valiant Navion™ thoracic stent graft device must be ordered in a size appropriate to fit the patient's anatomy. Proper sizing of the Valiant Navion™ thoracic stent graft is the responsibility of the physician.

### **ANEURYSMS OR PENETRATING ULCERS (PAU):**

Oversize the aortic portion of the stent graft by 3 to 7 mm, as appropriate for the patient.

The following table is provided as a guideline:

### **ANEURYSMS OR PAU**

Native Vessel (mm)	Suggested Stent Graft Diameter (mm)	Oversizing (mm)
16	20	4
17	20	3
18	22	4
19	22	3
20	25	5
21	25	4
22	25	3
23	28	5
24	28	4
25	28	3
26	31	5
27	31	4
28	31	3
28	34	6
29	34	5
30	34	4
31	34	3
30	37	7
31	37	6
32	37	5
33	37	4
33	40	7
34	40	6
35	40	5
36	40	4
36	43	7
37	43	6
38	43	5
39	43	4
39	46	7
40	46	6
41	46	5
42	46	4

### DISSECTION

Native Vessel (mm)	Suggested Stent Graft Diameter (mm)	Oversizing (mm)
19	20	1
20	22	2
21	22	1
22	22	0
23	25	2
24	25	1
25	25	0
26	28	2
27	28	1
28	28	0
29	31	2
30	31	1
31	34	3
32	34	2
33	34	1
34	37	3
35	37	2
36	37	1
37	40	3
38	40	2
39	40	1
39	43	4
40	43	3
41	43	2
42	43	1
42	46	4
43	46	3
44	46	2
45	46	1

#### DISSECTION:

Do not oversize the stent graft more than 10% of the healthy aorta nominal diameter. The following table is provided as a guideline.

#### FOR ADDITIONAL SECTIONS:

When the stent graft junction is located within the aneurysmal sac or is not supported by tissue, 6 mm oversizing between the primary component and additional section is recommended. In the case when a 20 mm stent graft is used as an outside component, the diameter of the inside component should be oversized by 5 mm relative to the outside component.

When the stent graft junction is supported by tissue (e.g., dissections), the stent graft should be oversized relative to the supporting native vessel.

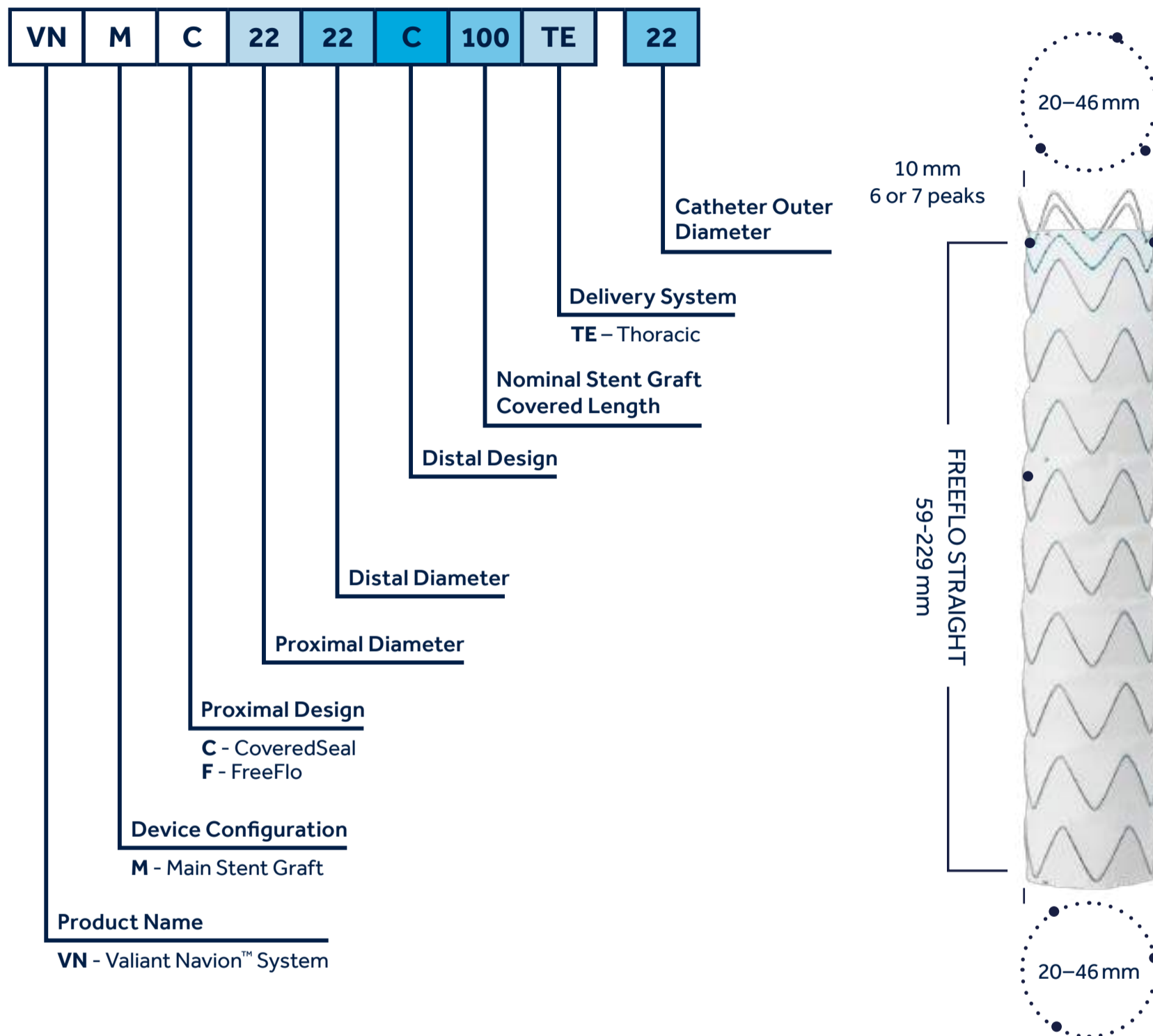
### BLUNT THORACIC AORTIC INJURY

Native Vessel (mm)	Suggested Stent Graft Diameter (mm)	Oversizing (mm)
16	20	4
17	20	3
18	22	4
19	22	3
20	22	2
20	25	5
21	25	4
22	25	3
23	25	2
23	28	5
24	28	4
25	28	3
26	28	2
26	31	5
27	31	4
28	31	3
29	31	2
28	34	6
29	34	5
30	34	4
31	34	3
32	34	2
30	37	7
31	37	6
32	37	5
33	37	4
34	37	3
35	37	2
33	40	7
34	40	6
35	40	5
36	40	4
37	40	3
38	40	2
36	43	7
37	43	6
38	43	5
39	43	4
40	43	3
41	43	2
39	46	7
40	46	6
41	46	5
42	46	4
43	46	3
44	46	2

# Valiant Navion™

## Thoracic Stent Graft System

### VALIANT NAVION™ SYSTEM PRODUCT CODE DESCRIPTION



# Valiant Navion™

## Thoracic Stent Graft System

### FREEFLO STRAIGHT

Product Code						
	Proximal Graft Diameter (mm)	Distal Graft Diameter (mm)		Stent Graft Covered Length (mm)		Catheter Outer Diameter (Fr)
VNMF	20	20	C	96	TE	18
VNMF	22	22	C	96	TE	18
VNMF	22	22	C	185	TE	18
VNMF	25	25	C	96	TE	18
VNMF	25	25	C	185	TE	18
VNMF	28	28	C	97	TE	20
VNMF	28	28	C	174	TE	20
VNMF	31	31	C	97	TE	20
VNMF	31	31	C	174	TE	20
VNMF	31	31	C	229	TE	20
VNMF	34	34	C	59	TE	20
VNMF	34	34	C	97	TE	20
VNMF	34	34	C	174	TE	20
VNMF	34	34	C	229	TE	20
VNMF	37	37	C	59	TE	20
VNMF	37	37	C	97	TE	20
VNMF	37	37	C	174	TE	20
VNMF	37	37	C	229	TE	20
VNMF	40	40	C	62	TE	22
VNMF	40	40	C	103	TE	22
VNMF	40	40	C	183	TE	22
VNMF	40	40	C	223	TE	22
VNMF	43	43	C	62	TE	22
VNMF	43	43	C	103	TE	22
VNMF	43	43	C	183	TE	22
VNMF	43	43	C	223	TE	22
VNMF	46	46	C	62	TE	22
VNMF	46	46	C	103	TE	22
VNMF	46	46	C	183	TE	22
VNMF	46	46	C	223	TE	22

AORTIC

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VENOUS



# Valiant Navion™

## Thoracic Stent Graft System

### FREEFLO TAPERED

Product Code						Catheter Outer Diameter (Fr)
VNMF	Proximal Graft Diameter (mm)	Distal Graft Diameter (mm)	C	Stent Graft Covered Length (mm)	TE	
VNMF	28	22	C	173	TE	20
VNMF	31	25	C	173	TE	20
VNMF	34	28	C	173	TE	20
VNMF	37	31	C	173	TE	20
VNMF	40	34	C	185	TE	22
VNMF	43	37	C	185	TE	22
VNMF	46	40	C	185	TE	22



# Valiant Navion™

## Thoracic Stent Graft System

### COVEREDSEAL STRAIGHT

Product Code						
VNMC	Proximal Graft Diameter (mm)	Distal Graft Diameter (mm)	C	Stent Graft Covered Length (mm)	TE	Catheter Outer Diameter (Fr)
VNMC	20	20	C	94	TE	18
VNMC	22	22	C	94	TE	18
VNMC	22	22	C	180	TE	18
VNMC	25	25	C	94	TE	18
VNMC	25	25	C	180	TE	18
VNMC	28	28	C	90	TE	20
VNMC	28	28	C	182	TE	20
VNMC	31	31	C	90	TE	20
VNMC	31	31	C	182	TE	20
VNMC	31	31	C	223	TE	20
VNMC	34	34	C	52	TE	20
VNMC	34	34	C	90	TE	20
VNMC	34	34	C	182	TE	20
VNMC	34	34	C	223	TE	20
VNMC	37	37	C	52	TE	20
VNMC	37	37	C	90	TE	20
VNMC	37	37	C	182	TE	20
VNMC	37	37	C	223	TE	20
VNMC	40	40	C	55	TE	22
VNMC	40	40	C	95	TE	22
VNMC	40	40	C	175	TE	22
VNMC	40	40	C	218	TE	22
VNMC	43	43	C	55	TE	22
VNMC	43	43	C	95	TE	22
VNMC	43	43	C	175	TE	22
VNMC	43	43	C	218	TE	22
VNMC	46	46	C	55	TE	22
VNMC	46	46	C	95	TE	22
VNMC	46	46	C	175	TE	22
VNMC	46	46	C	218	TE	22



# Valiant Navion™

## Thoracic Stent Graft System

### COVEREDSEAL TAPERED

Product Code						
	Proximal Graft Diameter (mm)	Distal Graft Diameter (mm)		Stent Graft Covered Length (mm)		Catheter Outer Diameter (Fr)
VNMC	25	20	C	186	TE	18
VNMC	28	22	C	207	TE	20
VNMC	31	25	C	207	TE	20
VNMC	34	28	C	207	TE	20
VNMC	37	31	C	207	TE	20
VNMC	40	34	C	200	TE	22
VNMC	43	37	C	200	TE	22
VNMC	46	40	C	200	TE	22



# Valiant™ Captivia™

## TAA Stent Graft System

### Features

#### Conformability delivered

- Sinusoidal shape and placement of nitinol springs provide flexibility and conformability
- Super-elastic nitinol springs exert active radial force to enhance seal and conformability

#### Confidence in control

- Tip capture provides controlled deployment and precise placement in the thoracic aorta
- Tip capture release handle provides simple turn-and-pull motion to release proximal stents

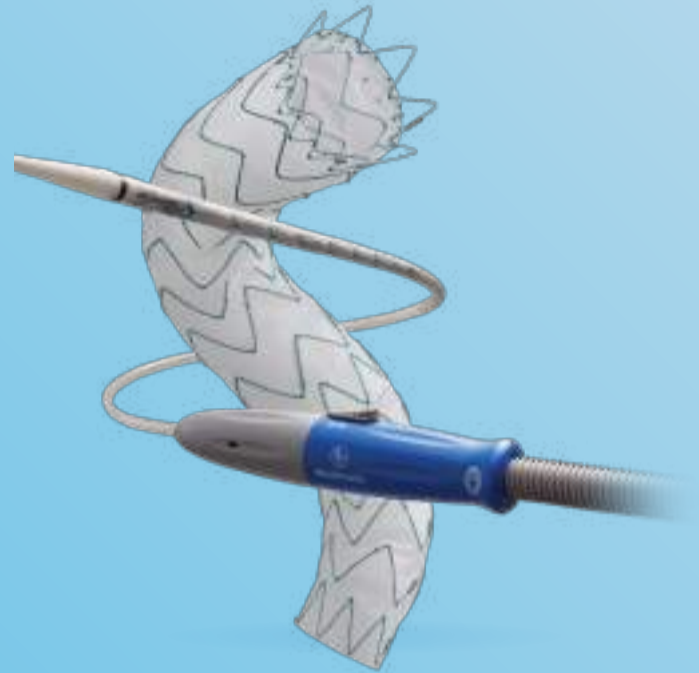
#### Advanced design\*

- Proximal 8-Peak FreeFlo configuration evenly distributes radial force over multiple apices
- Platinum iridium Figur8 markers provide high visibility and assist deployment
- Broad selection of proximal and distal components treats a variety of patients

#### Optimized access

- Crossing profile is similar to or lower than other thoracic stent grafts
- Hydrophilic coating facilitates stent graft delivery
- Easy three-step deployment process

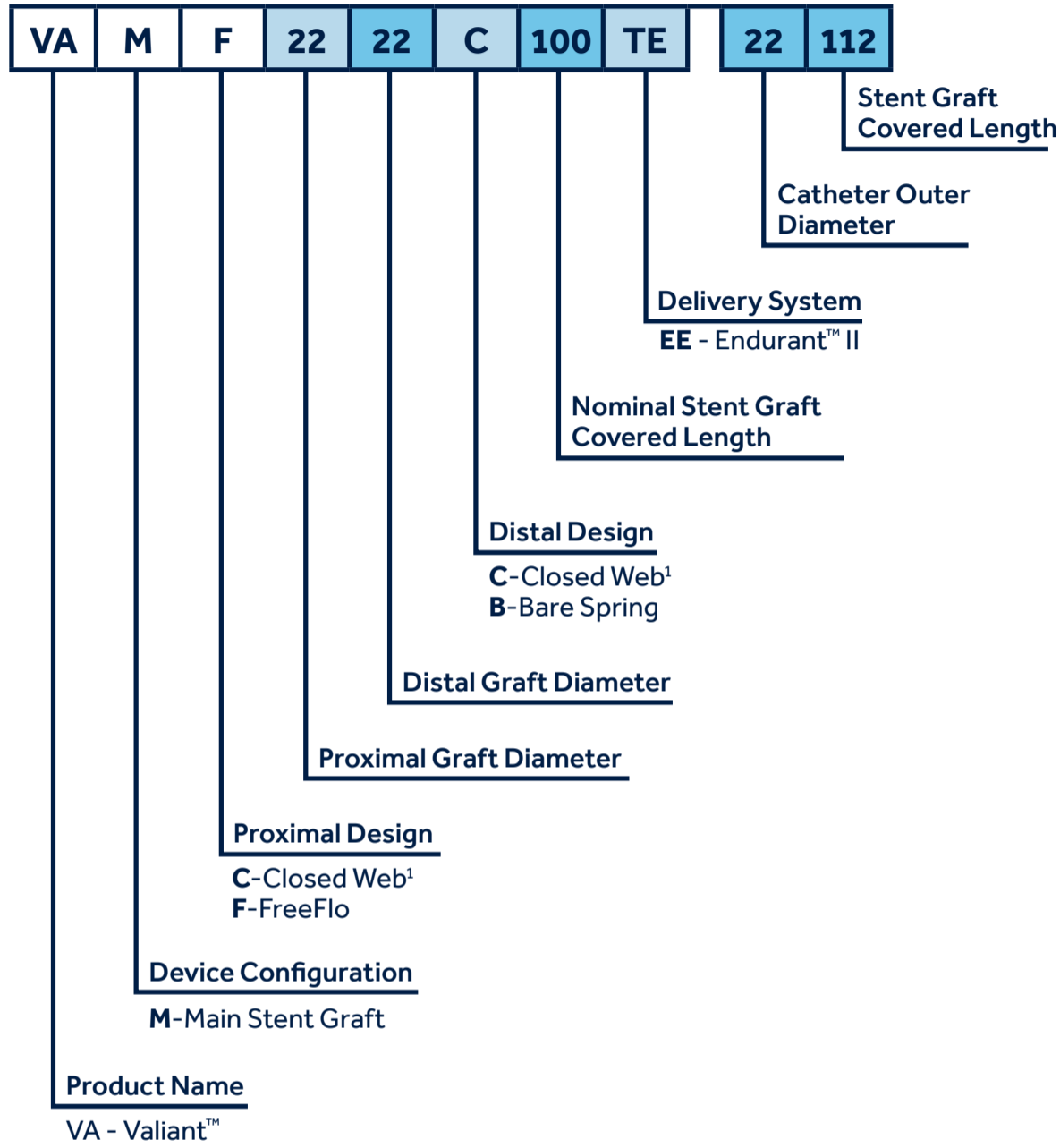
\* Test data on file at Medtronic. Bench test results may not be indicative of clinical performance.



# Valiant™ Captivia™

## TAA Stent Graft System

### VALIANT™ CAPTIVIA™ PRODUCT CODE DESCRIPTION



# Valiant™ Captivia™

## TAA Stent Graft System

### PROXIMAL FREEFLO STRAIGHT

Product Code							
	Proximal Graft Diameter (mm)	Distal Graft Diameter (mm)	Distal Design			Catheter Outer Diameter (F)	Stent Graft Covered Length (mm)
VAMF	22	22	C	100	TE	22	112
VAMF	24	24	C	100	TE	22	112
VAMF	26	26	C	100	TE	22	112
VAMF	28	28	C	100	TE	22	117
VAMF	30	30	C	100	TE	22	117
VAMF	32	32	C	100	TE	22	117
VAMF	34	34	C	100	TE	24	107
VAMF	36	36	C	100	TE	24	107
VAMF	38	38	C	100	TE	24	107
VAMF	40	40	C	100	TE	24	107
VAMF	42	42	C	100	TE	25	112
VAMF	44	44	C	100	TE	25	112
VAMF	46	46	C	100	TE	25	112
VAMF	22	22	C	150	TE	22	152
VAMF	24	24	C	150	TE	22	152
VAMF	26	26	C	150	TE	22	152
VAMF	28	28	C	150	TE	22	157
VAMF	30	30	C	150	TE	22	157
VAMF	32	32	C	150	TE	22	157
VAMF	34	34	C	150	TE	24	167
VAMF	36	36	C	150	TE	24	167
VAMF	38	38	C	150	TE	24	167
VAMF	40	40	C	150	TE	24	167
VAMF	42	42	C	150	TE	25	157
VAMF	44	44	C	150	TE	25	157
VAMF	46	46	C	150	TE	25	162
VAMF	30	30	C	200	TE	22	192
VAMF	32	32	C	200	TE	22	192
VAMF	34	34	C	200	TE	24	212
VAMF	36	36	C	200	TE	24	207
VAMF	38	38	C	200	TE	24	207
VAMF	40	40	C	200	TE	24	212
VAMF	42	42	C	200	TE	25	207
VAMF	44	44	C	200	TE	25	212
VAMF	46	46	C	200	TE	25	212

# Valiant™ Captivia™

## TAA Stent Graft System

### PROXIMAL FREEFLO TAPERED



Product Code							
	Proximal Graft Diameter (mm)	Distal Graft Diameter (mm)	Distal Design			Catheter Outer Diameter (F)	Stent Graft Covered Length (mm)
VAMF	26	22	C	150	TE	22	152
VAMF	28	24	C	150	TE	22	157
VAMF	30	26	C	150	TE	22	157
VAMF	32	28	C	150	TE	22	157
VAMF	34	30	C	150	TE	24	167
VAMF	36	32	C	150	TE	24	167
VAMF	38	34	C	150	TE	24	167
VAMF	40	36	C	150	TE	24	167
VAMF	42	38	C	150	TE	25	157
VAMF	44	40	C	150	TE	25	157
VAMF	46	42	C	150	TE	25	162

	Proximal Spring (mm)	Spring #2 (mm)	Spring #3 (mm)	Spring #4 (mm)	Spring #5 (mm)	Spring #6 (mm)	Spring #7 (mm)	Spring #8 (mm)	Spring #9 (mm)
	8-Peak	5-Peak	5-Peak	5-Peak	5-Peak	5-Peak	5-Peak	5-Peak	8-Peak
<b>40x36</b>	40	40	40	38	38	38	36	36	36
<b>38x34</b>	38	38	38	36	36	36	34	34	34
<b>36x32</b>	36	36	36	34	34	34	32	32	32
<b>34x30</b>	34	34	34	32	32	32	30	30	30
<b>32x28</b>	32	32	32	30	30	30	28	28	28
<b>30x26</b>	30	30	30	28	28	28	26	26	26
<b>28x24</b>	28	28	28	26	26	26	24	24	24
<b>26x22</b>	26	26	26	24	24	24	22	22	22

	Proximal Spring (mm)	Spring #2 (mm)	Spring #3 (mm)	Spring #4 (mm)	Spring #5 (mm)	Spring #6 (mm)	Spring #7 (mm)	Spring #8 (mm)	N/A
	8-Peak	5-Peak	5-Peak	5-Peak	5-Peak	5-Peak	5-Peak	8-Peak	
<b>46x42</b>	46	46	46	44	44	44	42	42	
<b>44x40</b>	44	44	44	42	42	42	40	40	
<b>42x38</b>	42	42	42	40	40	40	38	38	

# Valiant™ Captivia™

## TAA Stent Graft System

### CLOSED WEB STRAIGHT



Product Code							
	Proximal Graft Diameter (mm)	Distal Graft Diameter (mm)	Distal Design			Catheter Outer Diameter (F)	Stent Graft Covered Length (mm)
VAMC	22	22	C	100	TE	22	105
VAMC	24	24	C	100	TE	22	105
VAMC	26	26	C	100	TE	22	105
VAMC	28	28	C	100	TE	22	110
VAMC	30	30	C	100	TE	22	110
VAMC	32	32	C	100	TE	22	110
VAMC	34	34	C	100	TE	24	100
VAMC	36	36	C	100	TE	24	100
VAMC	38	38	C	100	TE	24	100
VAMC	40	40	C	100	TE	24	100
VAMC	42	42	C	100	TE	25	105
VAMC	44	44	C	100	TE	25	105
VAMC	46	46	C	100	TE	25	105
VAMC	22	22	C	150	TE	22	145
VAMC	24	24	C	150	TE	22	145
VAMC	26	26	C	150	TE	22	145
VAMC	28	28	C	150	TE	22	150
VAMC	30	30	C	150	TE	22	150
VAMC	32	32	C	150	TE	22	150
VAMC	34	34	C	150	TE	24	160
VAMC	36	36	C	150	TE	24	160
VAMC	38	38	C	150	TE	24	160
VAMC	40	40	C	150	TE	24	160
VAMC	42	42	C	150	TE	25	150
VAMC	44	44	C	150	TE	25	150
VAMC	46	46	C	150	TE	25	155
VAMC	30	30	C	200	TE	22	185
VAMC	32	32	C	200	TE	22	185
VAMC	34	34	C	200	TE	24	205
VAMC	36	36	C	200	TE	24	200
VAMC	38	38	C	200	TE	24	200
VAMC	40	40	C	200	TE	24	205
VAMC	42	42	C	200	TE	25	200
VAMC	44	44	C	200	TE	25	205
VAMC	46	46	C	200	TE	25	205



# Valiant™ Captivia™

## TAA Stent Graft System

### CLOSED WEB TAPERED



Product Code							
	Proximal Graft Diameter (mm)	Distal Graft Diameter (mm)	Distal Design			Catheter Outer Diameter (F)	Stent Graft Covered Length (mm)
VAMC	26	22	C	150	TE	22	150
VAMC	28	24	C	150	TE	22	150
VAMC	30	26	C	150	TE	22	150
VAMC	32	28	C	150	TE	22	150
VAMC	34	30	C	150	TE	24	160
VAMC	36	32	C	150	TE	24	160
VAMC	38	34	C	150	TE	24	160
VAMC	40	36	C	150	TE	24	160
VAMC	42	38	C	150	TE	25	150
VAMC	44	40	C	150	TE	25	150
VAMC	46	42	C	150	TE	25	155

	Proximal Spring (mm)	Spring #2 (mm)	Spring #3 (mm)	Spring #4 (mm)	Spring #5 (mm)	Spring #6 (mm)	Spring #7 (mm)	Spring #8 (mm)
	8-Peak	5-Peak	5-Peak	5-Peak	5-Peak	5-Peak	5-Peak	8-Peak
26x22	26	26	26	24	24	24	22	22
28x24	28	28	28	26	26	26	24	24
30x26	30	30	30	28	28	28	26	26
32x28	32	32	32	30	30	30	28	28
34x30	34	34	34	32	32	32	30	30
36x32	36	36	36	34	34	34	32	32
38x34	38	38	38	36	36	36	34	34
40x36	40	40	40	38	38	38	36	36

	Proximal Spring (mm)	Spring #2 (mm)	Spring #3 (mm)	Spring #4 (mm)	Spring #5 (mm)	Spring #6 (mm)	Spring #7 (mm)	N/A
	8-Peak	5-Peak	5-Peak	5-Peak	5-Peak	5-Peak	8-Peak	
42x38	42	42	42	40	40	40	38	
44x40	44	44	44	42	42	42	40	
46x42	46	46	46	44	44	44	42	

# Valiant™ Captivia™

## TAA Stent Graft System

### DISTAL BARE SPRING STRAIGHT



Product Code							
	Proximal Graft Diameter (mm)	Distal Graft Diameter (mm)	Distal Design			Catheter Outer Diameter (F)	Stent Graft Covered Length (mm)
VAMC	22	22	B	100	TE	22	112
VAMC	24	24	B	100	TE	22	112
VAMC	26	26	B	100	TE	22	112
VAMC	28	28	B	100	TE	22	117
VAMC	30	30	B	100	TE	22	117
VAMC	32	32	B	100	TE	22	117
VAMC	34	34	B	100	TE	24	107
VAMC	36	36	B	100	TE	24	107
VAMC	38	38	B	100	TE	24	107
VAMC	40	40	B	100	TE	24	107
VAMC	42	42	B	100	TE	25	112
VAMC	44	44	B	100	TE	25	112
VAMC	46	46	B	100	TE	25	112

# Valiant™ Captivia™

## TAA Stent Graft System

Each Valiant™ Thoracic Stent Graft with Captivia™ Delivery System must be ordered in a size appropriate to fit the patient's anatomy. Proper sizing of the Valiant™ Captivia™ Thoracic Stent Graft Delivery System is the responsibility of the physician.

### **ANEURYSMS, PENETRATING ULCERS, AND TRAUMATIC RUPTURES:**

Oversize the aortic portion of the stent graft by 3 to 5 mm as appropriate for the patient. The following table is provided as a guideline:

Native Vessel (mm)	Suggested Stent Graft Diameter (mm)	Oversizing (mm)
18	22	4
19	22	3
20	24	4
21	24	3
22	26	4
24	26	3
25	28	4
25	30	5
26	30	4
27	30	3
27	32	5
28	32	4
29	32	3
29	34	5
30	34	4
31	34	3
31	36	5
32	36	4
33	38	5
34	38	4
35	40	5
36	40	4
37	42	5
38	42	4
39	44	5
40	44	4
41	46	5
42	46	4

# Valiant™ Captivia™

## TAA Stent Graft System

### DISSECTION:

Do not oversize the stent graft by more than 10% of the healthy aorta nominal diameter. The following table is provided as a guideline:

Native Vessel (mm)	Suggested Stent Graft Diameter (mm)	Oversizing (mm)
20	22	2
21	22	1
22	24	2
23	24	1
24	26	2
25	26	1
26	28	2
27	28	1
28	30	2
29	32	3
30	32	2
31	34	3
32	34	2
33	36	3
34	36	2
35	38	3
36	38	2
37	40	3
38	40	2
39	42	3
40	42	2
40	44	4
41	44	3
42	44	2
42	46	4
43	46	3
44	46	2

### FOR ADDITIONAL SECTIONS:

When the stent graft junction is located within the aneurismal sac or is not supported by tissue, a 4 mm oversizing between the primary component and additional section is recommended. When the stent graft junction is supported by tissue (e.g., dissections), recommendations listed above.

# ENDOANCHOR™ SYSTEMS

AORTIC

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# Heli-Fx™ / Heli-Fx™ TAA

## AAA/TAA EndoAnchor™ System

Tailor seal and fixation in your primary and revision Tevar cases

### Stability of a surgical anastomosis

- Helical EndoAnchor™ implant designed to provide the stability of a surgical anastomosis

### Enhanced sealing and fixation

- Enhances the inherent sealing and fixation mechanisms of an endograft

### Simplified revisions

- Simplifies revision surgery for endograft migration and Type I endoleak

### Precise and accurate placement

- Steerable guide for precise and accurate EndoAnchor™ implant placement

### Intuitive and controlled deployment

- Motorized, intuitive controls for precise placement of EndoAnchor™ implants

### High visibility

- Excellent system and EndoAnchor™ implant radiopacity



Primary implantation of EndoAnchor™ implant with an Endurant™ bifurcated endograft in complex proximal neck anatomy.



# Heli-Fx™ EndoAnchor™

## AAA EndoAnchor™ System

### RECOMMENDED HELI-FX™ GUIDE SELECTION

Aortic Inner Diameter (mm)	Deflected Tip Reach (mm)
18 - 28	22
28 - 32	28

### ORDER INFORMATION

AAA Components	Product Catalogue Number
Heli-FX™ Guide, 22 mm	SG-64
Heli-FX™ Guide, 28 mm	HG-16-62-28
Heli-FX™ Applier and EndoAnchor™ Cassette (Contains 10 EndoAnchors)	SA-85

### TECHNICAL SPECIFICATION

EndoAnchor™ Implant	Heli-FX™ Guide	Heli-FX™ Applier
3.0 mm diameter x 4.5 mm length	Two deflection-tip lengths address varying neck diameters	Battery-operated delivery device with visual and audio feedback
0.5 mm diameter MP35N-LT wire thickness	Unique radiopaque tip markers for 3D orientation	One-touch auto-loading of EndoAnchors
Atraumatic conical tip (similar to SH1 needle)	16 F OD	Two-stage EndoAnchor deployment allows placement confirmation and repositioning
Crossbar feature prevents over-penetration	62 cm working length	86 cm working length
	0.035" guidewire compatible	

# Heli-Fx™ TAA

## TAA EndoAnchor™ System

### RECOMMENDED HELI-FX™ GUIDE SELECTION

Aortic Inner Diameter (mm)	Deflected Tip Reach (mm)
18-28	22
28-38	32
38-42	42

### ORDER INFORMATION

TAA Components	Product Catalogue Number
Heli-FX™ Guide, 22 mm	HG-18-90-22
Heli-FX™ Guide, 32 mm	HG-18-90-32
Heli-FX™ Guide, 42 mm	HG-18-90-42
Heli-FX™ Applier and EndoAnchor™ Cassette (contains 10 EndoAnchors)	HA-18-114

### TECHNICAL SPECIFICATIONS

EndoAnchor™ Implant	Heli-FX™ Guide	Heli-FX™ Applier
3.0 mm diameter x 4.5 mm length	Three deflected tip reach lengths address varying neck diameters	Battery-operated delivery device with visual and audio feedback
0.5 mm diameter MP35N-LT wire thickness	Unique radiopaque tip markers for 3D orientation	One-touch auto-loading of EndoAnchors
Atraumatic conical tip (similar to SH1 needle)	18 F OD	Two-stage EndoAnchor deployment allows placement confirmation and repositioning
Crossbar feature prevents over-penetration	90 cm working length	114 cm working length
	0.035" guidewire compatible	



# ANCILLARY

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# Sentrant™

## Introducer Sheath with Hydrophilic Coating

### EnsureSeal technology†

- Optimal hemostasis versus competitive sheaths

### Complex ready design

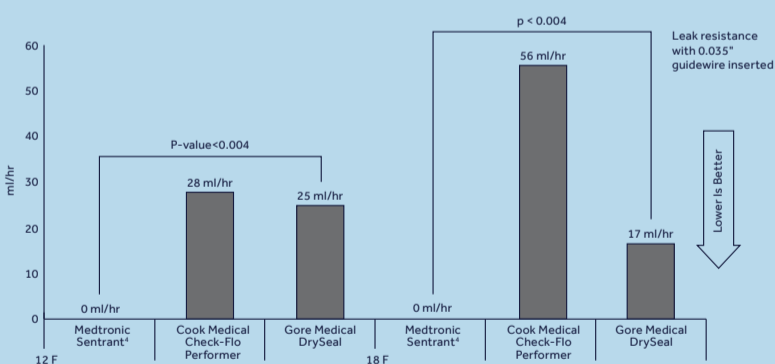
- Hydrophilic coating with flexibility for easier tracking through tortuous and calcified iliacs
- Coil reinforced for kink resistance

### Improved procedural confidence

- Radiopaque marker for easy visibility
- Dilator locking feature secures desired position

†Bench Test Data on file at Medtronic. Test data not indicative of clinical performance. Bench Test compared Cook Check-Flo™™ Performer™™ and Gore DrySeal 12F and 18F to Sentrant™™ 12F and 18F.

### Superior leak resistance versus Cook Check-Flo® Performer® and Gore Dryseal†



\*Bench Test Data on file at Medtronic. Test data not indicative of clinical performance. Bench Test compared Cook Check-Flo Performer and Gore DrySeal 12F and 18F to Sentrant™ 12F and 18F.

†Medtronic Sheath had zero leakage on 0.035" guidewire.



## Introducer Sheath With Hydrophilic Coating

### ORDER INFORMATION

Product Catalogue Number	Inner Diameter Size (F)	Usable Length (cm)
SENSH1228W	12	28
SENSH1428W	14	28
SENSH1628W	16	28
SENSH1828W	18	28
SENSH2028W	20	28
SENSH2228W	22	28
SENSH2428W	24	28
SENSH2628W	26	28

### THE CHOICE FOR SUPERIOR HEMOSTASIS

- 1 RADIOPAQUE MARKER BAND
- 2 HYDROPHILIC COATING
- 3 REINFORCED COILED TUBING
- 4 GUIDEWIRE DIAMETER: 0.035"
- 5 LOCKING MECHANISM ON DILATOR HANDLE



# Reliant™

## Stent Graft Balloon Catheter

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Multiple purposes, single solution

### Versatile design

- Widest range of inflation diameters in a single low profile balloon able to treat thoracic to iliacs (10 mm–46 mm)<sup>2</sup>
- Compatible with 12 F sheath

### Reliable performance

- Consistent inflation and deflation time
- Stable expansion with minimum balloon overhang to reduce risk of vessel trauma
- Dependable expansion even after multiple inflations and deflations<sup>3</sup>

### Improved conformability

- Expands kinks and smooths creases from the graft material
- Compliant balloon expansion

### Clinical uses include:

- For use in thoracic, abdominal and iliac endograft procedures
- Endograft molding
- Apposition of endografts in seal zones
- Temporary aortic occlusion

<sup>2</sup> Reliant™ Stent Graft Balloon Catheter Instructions For Use, Cook Coda™™ Balloon Catheter Instructions For Use, Gore Tri-Lobe™™ Instructions For Use.

<sup>3</sup> Data on file Medtronic. Maximum of 20 inflations, deflations.

### BALLOON INFLATION TABLE\*

#### 46 MM BALLOON

Diameter (mm)	Volume (cc)
10	3
20	9
30	19
40	41
46*	60

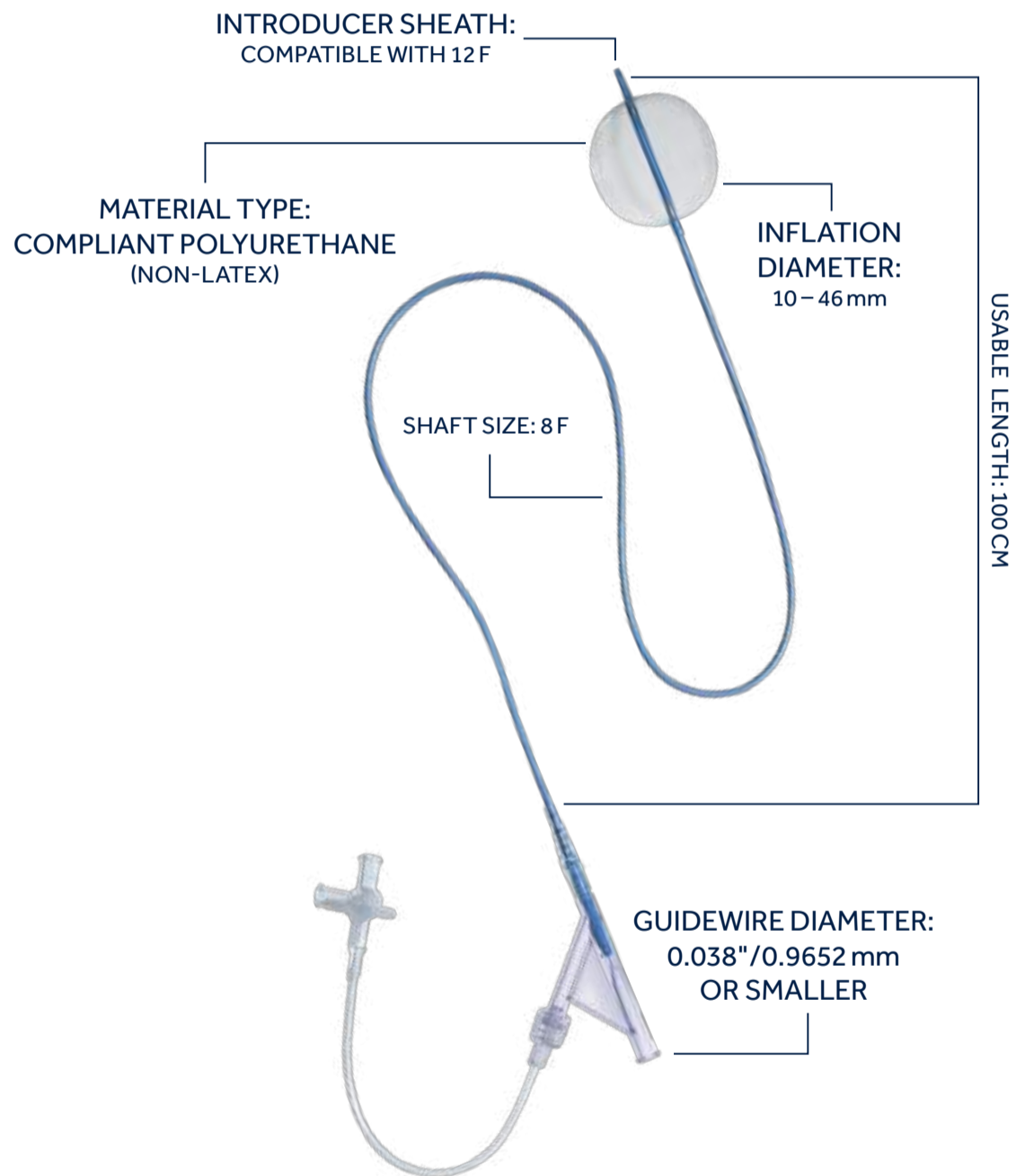
\* CAUTION: This table is only a guide. Balloon expansion should be carefully monitored under fluoroscopy. Do not exceed maximum inflation diameter (46 mm). Rupture of balloon may occur.



## Stent Graft Balloon Catheter

### ORDER INFORMATION

Product Catalogue Number	Inflation Diameter (mm)	Shaft Size (F)	Usable Length (cm)	Sheath Compatibility (F)
REL46	10-46	8	100	12



# TourGuide™

## Steerable Sheath

### Dilator and tip†

- Facilitates sheath trackability, enabling the TourGuide™ Sheath to advance through indicated anatomy

### Advanced catheter technology

- Provides the strength and conformability needed for delivering various interventional devices

### Curve retention†

- Maintains desired deflection angle, providing a stable platform for delivering devices to the desired destination

### Radiopaque tip†

- Provides excellent visualization during positioning, enabling accurate vessel access and potentially less manipulation

### Hemostatic seal and flush port

- Maintains hemostasis and allows for manual fluid injection

### Safety in mind†

- May reduce overall procedure time by minimizing multiple exchanges associated with different catheter selections

### 180° tip deflection†

- Physician-directed to deliver diagnostic and therapeutic devices to a wide variety of vessel take offs and difficult anatomical areas

### Three working lengths

- Designed to access indicated areas to perform interventions

### Inner diameter compatibility

- For use with interventional devices

### Dilator

- With French size and guidewire indicator†

†Bench Test Data on file at Medtronic. Test data not indicative of clinical performance.

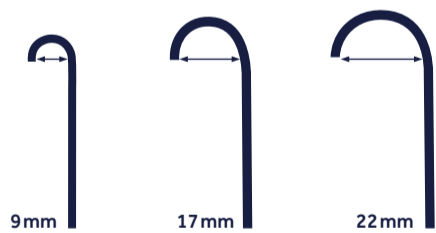


### ORDER INFORMATION

Product Catalogue Number	Inner Diameter Size (F)	Useable Length (cm)	Deflection Length @ 180° (mm)	Outer Diameter Size (F)
TG0654509	6.5	45	9	8.5
TG0654517	6.5	45	17	8.5
TG0655509	6.5	55	9	8.5
TG0655517	6.5	55	17	8.5
TG0659009	6.5	90	9	8.5
TG0704509	7.0	45	9	9.5
TG0704517	7.0	45	17	9.5
TG0705509	7.0	55	9	9.5
TG0705517	7.0	55	17	9.5
TG0709009	7.0	90	9	9.5
TG0854517	8.5	45	17	12
TG0854522	8.5	45	22	12
TG0855517	8.5	55	17	12
TG0855522	8.5	55	22	12
TG0859017	8.5	90	17	12

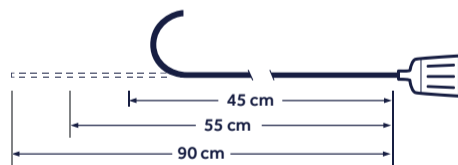
#### ADJUSTABLE TIP DEFLECTION

May reduce overall procedure time by minimizing multiple exchanges associated with different catheter selections



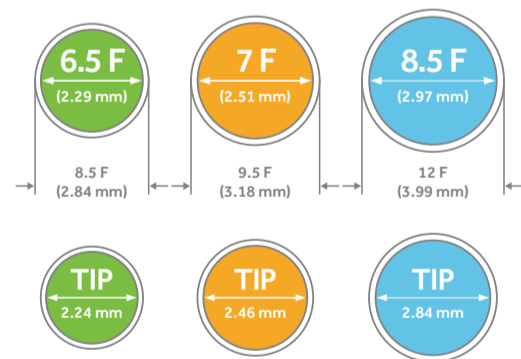
#### THREE WORKING LENGTHS

Can access most anatomic areas to perform peripheral interventions



#### INNER DIAMETER COMPATIBILITY

For use with peripheral interventional devices



#### DILATOR

With French size and guidewire indicator

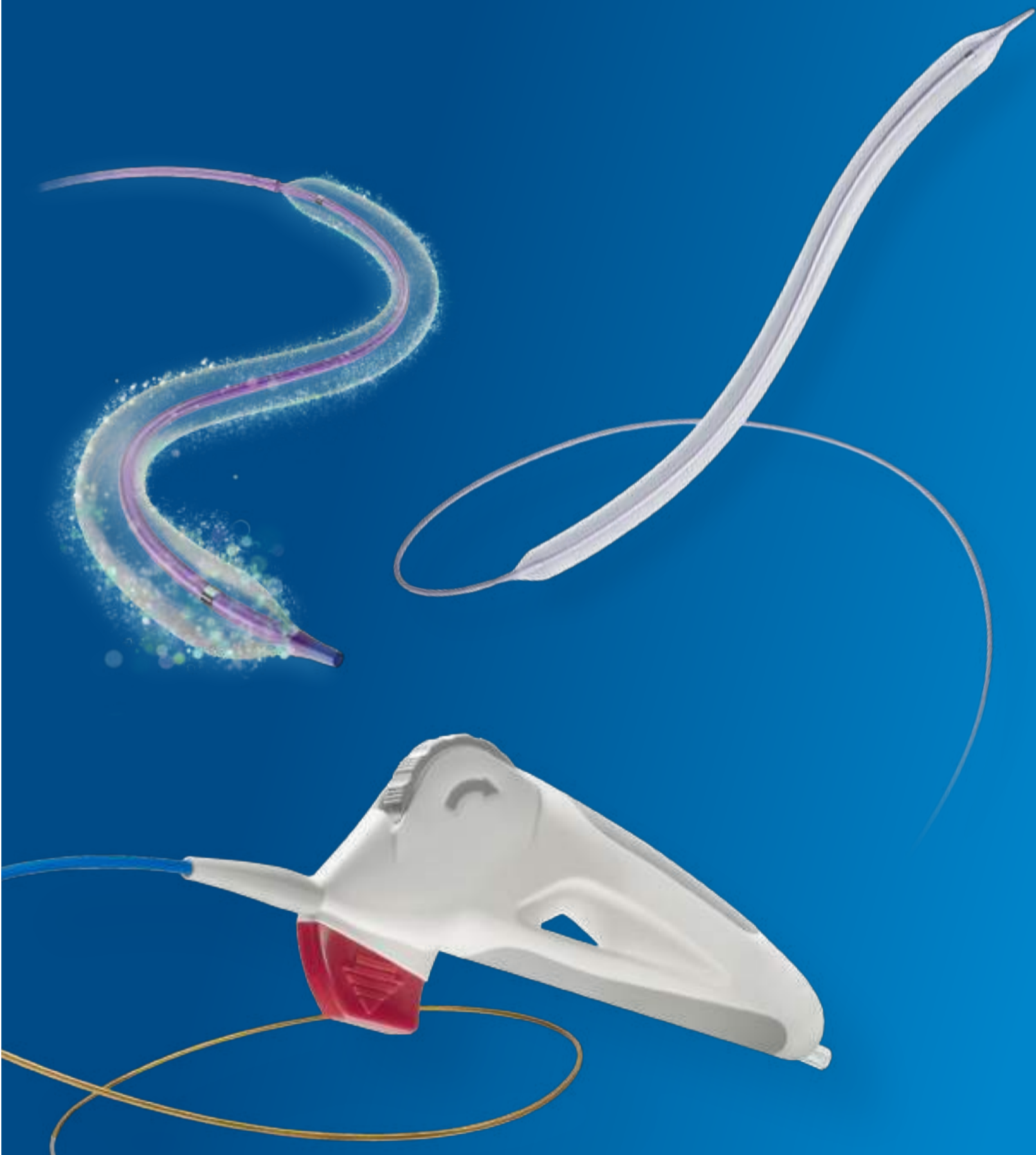


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# DRUG COATED BALLOONS

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# IN.PACT™ Admiral™

Paclitaxel-eluting PTA Balloon Catheter 0.035"



## TECHNICAL SPECIFICATIONS

Catheter design	Over the Wire (OTW)
Balloon coating	FreePac – Paclitaxel and Urea (Excipient)
Usable shaft lengths	40, 80 and 130 cm
Introducer sheath compatibility	5, 6, 7 and 9F
Max. recommended guidewire	0.035" depending on balloon size
Nominal pressure	8 atm

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# IN.PACT™ Admiral™

## Paclitaxel-eluting PTA Balloon Catheter 0.035"

### DURABLE

IN.PACT™ Admiral™ drug-eluting balloon demonstrates best-in-class clinical outcomes with durable performance through 3 years.

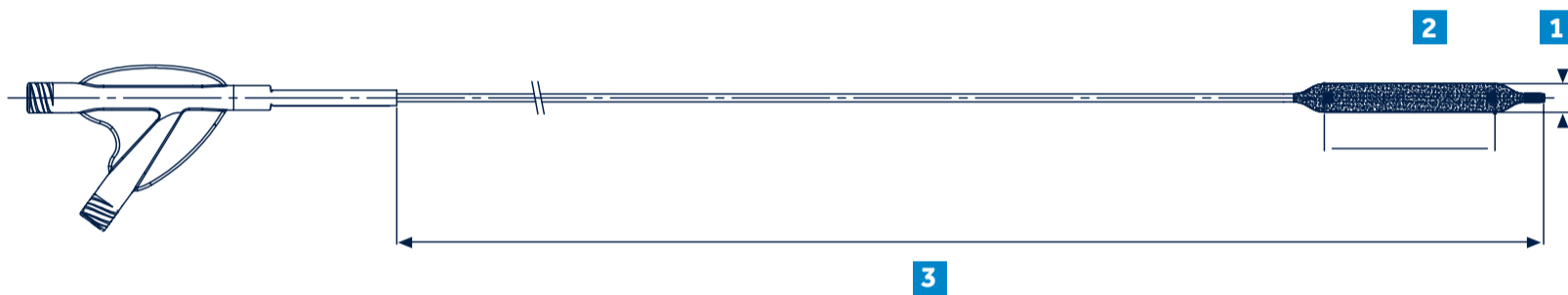
### SAFE

IN.PACT™ Admiral™ drug-eluting balloon has an excellent safety profile, with superior results relative to PTA.

### CONSISTENT

IN.PACT™ Admiral™ drug-eluting balloon demonstrates positive, consistent outcomes across trials, complex patients and lesion subgroups.

- 1** Balloon Diameter      **2** Balloon Length      **3** Usable Length



# IN.PACT™ Admiral™

## Paclitaxel-eluting PTA Balloon Catheter 0.035"

### ORDER INFORMATION

Product Catalogue Number			Balloon Diameter (mm)	Balloon Length (mm)	Recommended Introducer Sheath (F)	RBP (atm)
Usable Length 40 cm	Usable Length 80 cm	Usable Length 130 cm				
SBI 040 040 04P	SBI 040 040 08P	SBI 040 040 13P	4	40	5	14
SBI 040 060 04P	SBI 040 060 08P	SBI 040 060 13P	4	60	5	14
SBI 040 080 04P	SBI 040 080 08P	SBI 040 080 13P	4	80	5	14
-	SBI 040 120 08P	SBI 040 120 13P	4	120	5	14
-	SBI 040 150 08P	SBI 040 150 13P	4	150	5	14
SBI 050 040 04P	SBI 050 040 08P	SBI 050 040 13P	5	40	6	14
SBI 050 060 04P	SBI 050 060 08P	SBI 050 060 13P	5	60	6	14
SBI 050 080 04P	SBI 050 080 08P	SBI 050 080 13P	5	80	6	14
-	SBI 050 120 08P	SBI 050 120 13P	5	120	6	14
-	SBI 050 150 08P	SBI 050 150 13P	5	150	6	14
SBI 060 040 04P	SBI 060 040 08P	SBI 060 040 13P	6	40	6	14
SBI 060 060 04P	SBI 060 060 08P	SBI 060 060 13P	6	60	6	14
SBI 060 080 04P	SBI 060 080 08P	SBI 060 080 13P	6	80	6	14
-	SBI 060 120 08P	SBI 060 120 13P	6	120	6	14
-	SBI 060 150 08P	SBI 060 150 13P	6	150	6	14
SBI 070 040 04P	SBI 070 040 08P	SBI 070 040 13P	7	40	7	14
SBI 070 060 04P	SBI 070 060 08P	SBI 070 060 13P	7	60	7	14
SBI 070 080 04P	SBI 070 080 08P	SBI 070 080 13P	7	80	7	14
SBI 080 040 04P	SBI 080 040 08P	SBI 080 040 13P	8	40	7	10
SBI 080 060 04P	SBI 080 060 08P	SBI 080 060 13P	8	60	7	10
SBI 080 080 04P	SBI 080 080 08P	SBI 080 080 13P	8	80	7	10
SBI 090 040 04P	SBI 090 040 08P	SBI 090 040 13P	9	40	7	10
SBI 090 060 04P	SBI 090 060 08P	SBI 090 060 13P	9	60	7	10
SBI 090 080 04P	SBI 090 080 08P	SBI 090 080 13P	9	80	7	10
SBI 100 040 04P	SBI 100 040 08P	SBI 100 040 13P	10	40	7	9
SBI 120 040 04P	SBI 120 040 08P	SBI 120 040 13P	12	40	9	9

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# IN.PACT™ Admiral™

Paclitaxel-eluting PTA Balloon Catheter 0.035"

Balloon Lengths (mm)

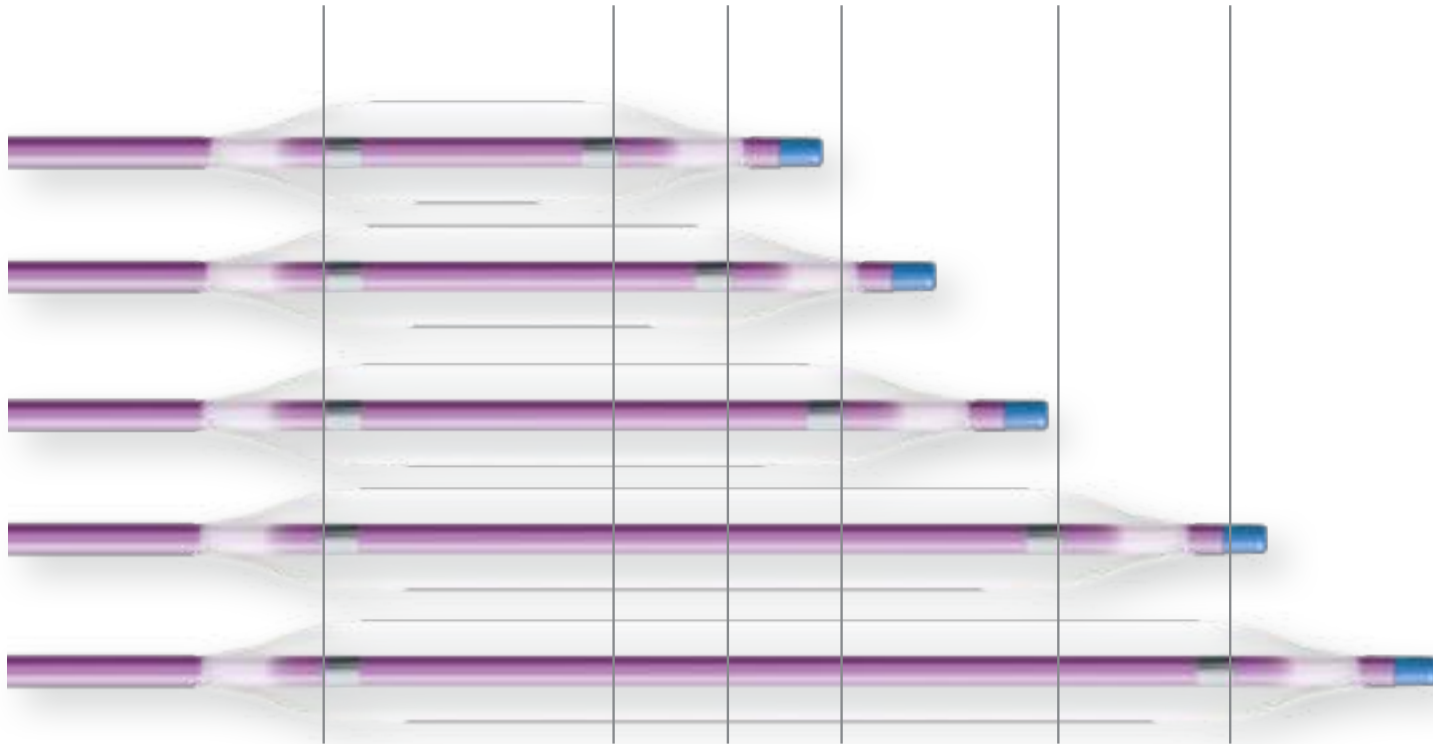
40

60

80

120

150



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# IN.PACT™ Pacific™

Paclitaxel-eluting PTA Balloon Catheter 0.018"



## UNPARALLELED IN PERIPHERAL TECHNICAL SPECIFICATIONS

Catheter design	Over the Wire (OTW)
Balloon coating	FreePac – Paclitaxel and Urea (Excipient)
Usable shaft lengths	90 and 130 cm
Introducer sheath compatibility	5 - 6 F depending on balloon size
Max. recommended guidewire	0.018"
Nominal pressure	7 atm

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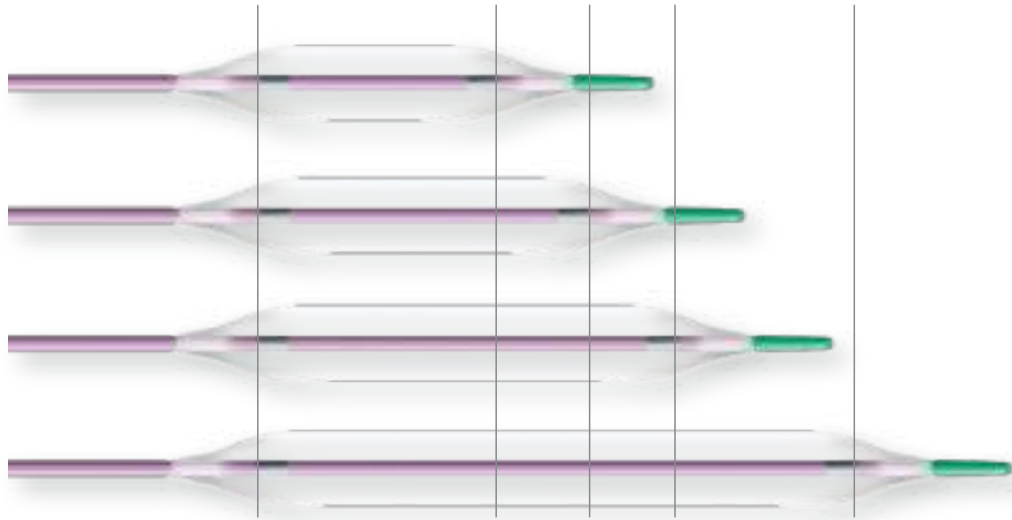
VENOUS

# IN.PACT™ Pacific™

## Paclitaxel-eluting PTA Balloon Catheter 0.018"

Balloon Lengths (mm)

40 60 80 120



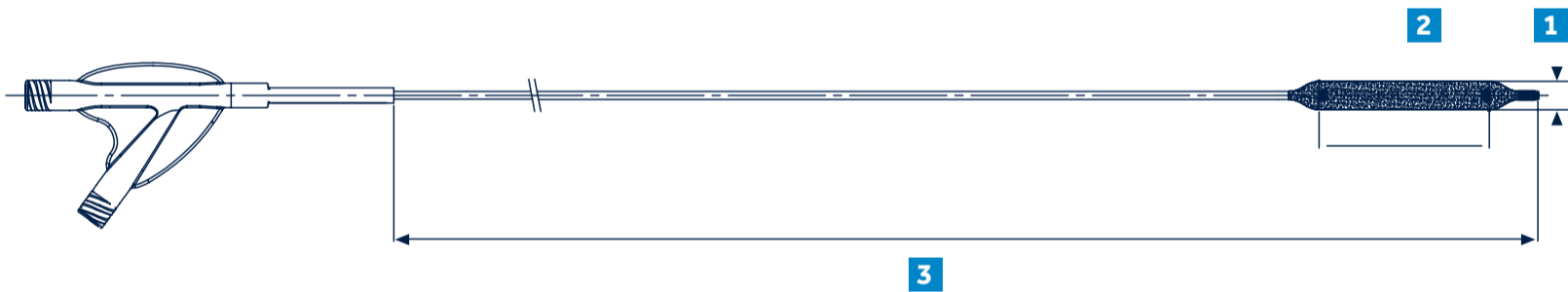
Balloon Diameters  
4, 5, 6, 7mm



**1** Balloon Diameter

**2** Balloon Length

**3** Usable Length



## Paclitaxel-eluting PTA Balloon Catheter 0.018"

### ORDER INFORMATION

Product Catalogue Number		Balloon Diameter (mm)	Balloon Length (mm)	Recommended Introducer Sheath (F)	RBP (atm)
Usable Length 90 cm	Usable Length 130 cm				
PCF 040 040 09P	PCF 040 040 13P	4	40	5	20
PCF 040 060 09P	PCF 040 060 13P	4	60	5	14
PCF 040 080 09P	PCF 040 080 13P	4	80	5	14
PCF 040 120 09P	PCF 040 120 13P	4	120	5	14
PCF 050 040 09P	PCF 050 040 13P	5	40	5	20
PCF 050 060 09P	PCF 050 060 13P	5	60	5	14
PCF 050 080 09P	PCF 050 080 13P	5	80	5	14
PCF 050 120 09P	PCF 050 120 13P	5	120	5	14
PCF 060 040 09P	PCF 060 040 13P	6	40	5	16
PCF 060 060 09P	PCF 060 060 13P	6	60	5	14
PCF 060 080 09P	PCF 060 080 13P	6	80	5	14
PCF 060 120 09P	PCF 060 120 13P	6	120	5	14
PCF 070 040 09P	PCF 070 040 13P	7	40	6	12
PCF 070 060 09P	PCF 070 060 13P	7	60	6	12
PCF 070 080 09P	PCF 070 080 13P	7	80	6	12
PCF 070 120 09P	PCF 070 120 13P	7	120	6	12

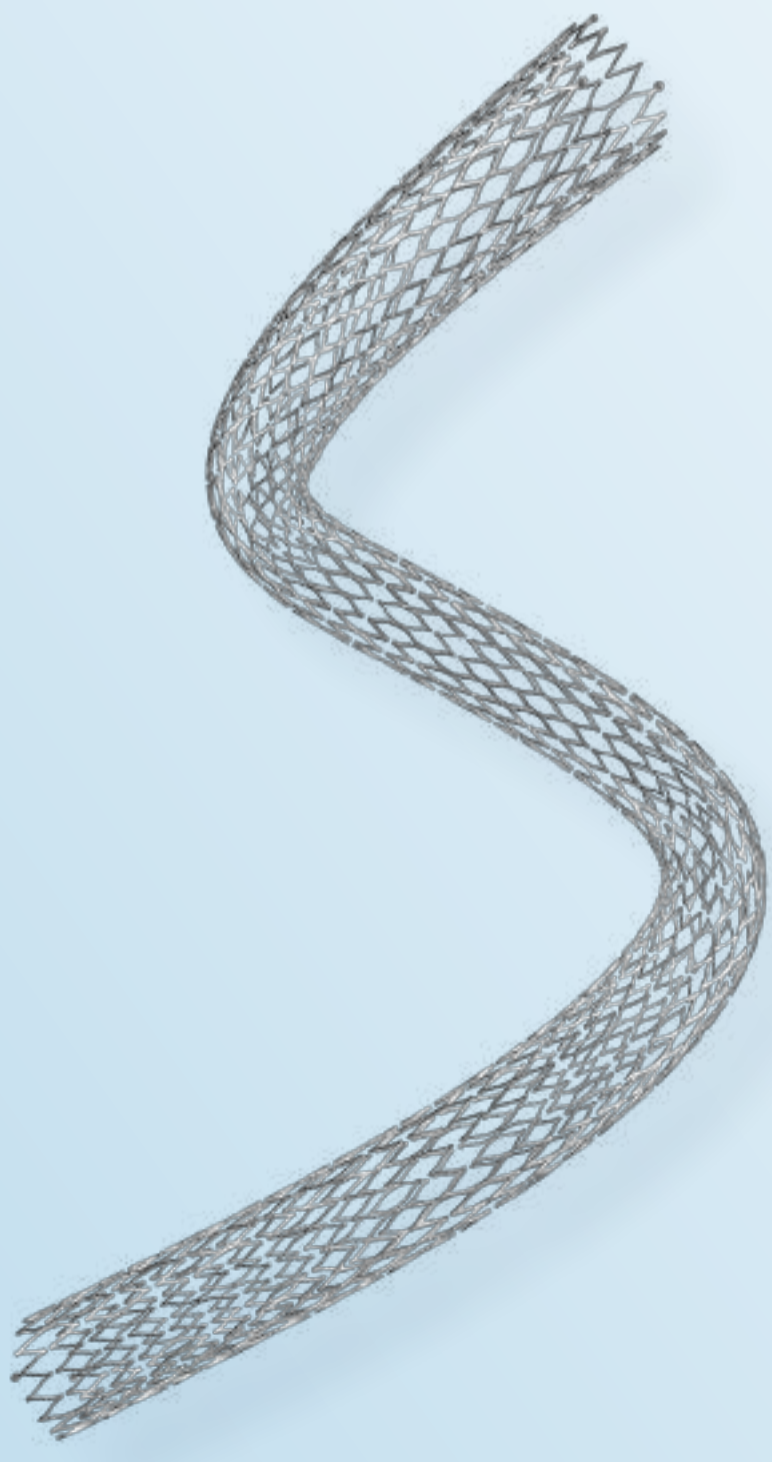


# STENT SYSTEMS

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# Protégé™ Rx™

## Carotid Self-Expanding Stent System

The Protégé™ RX™ stent is the next generation stent designed for the anatomy of the carotid artery.

Protégé™ RX™ provides control and accurate placement for carotid interventions.

### Predictable Deployment

- Proprietary EX.P.R.T.™ release technology essentially eliminates premature deployment or jumping
- No stent shortening
- Unique anatomically designed tapered stent for better fit in the carotid bifurcation
- 0.014" rapid exchange catheter with 6 F low crossing profile and flexible atraumatic tip
- Radiopaque marker on catheter clearly indicates tapered location for precise positioning

### Visible Results

- Tantalum GPS™ markers enhance visibility for precise positioning and result confirmation
- Cell design produces expansion force that resists compression while providing excellent wall apposition
- Straight and tapered options for customized fit in carotid vessels

Protégé RX 6 Fr/0.014" Catheter Length 135cm

### Each system includes:

One stent and delivery catheter system



## Carotid Self-Expanding Stent System

### ORDER INFORMATION

Product Catalogue Number	Stent Dimensions		Recommended Sheath Size (F)	Recommended Guidewire (inch)	Crossing Profile (inch)
Catheter Length 135cm	Diameter (mm)	Length (mm)			
<b>TAPERED</b>					
SEPX-8-6-30-135	8x6	30	6	0.014	0.078
SEPX-8-6-40-135	8x6	40	6	0.014	0.078
SEPX-10-7-30-135	10x7	30	6	0.014	0.078
SEPX-10-7-40-135	10x7	40	6	0.014	0.078
<b>STRAIGHT</b>					
SEPX-6-20-135	6	20	6	0.014	0.078
SEPX-7-20-135	7	20	6	0.014	0.078
SEPX-8-20-135	8	20	6	0.014	0.078
SEPX-9-20-135	9	20	6	0.014	0.078
SEPX-10-20-135	10	20	6	0.014	0.078
SEPX-6-30-135	6	30	6	0.014	0.078
SEPX-7-30-135	7	30	6	0.014	0.078
SEPX-8-30-135	8	30	6	0.014	0.078
SEPX-9-30-135	9	30	6	0.014	0.078
SEPX-10-30-135	10	30	6	0.014	0.078
SEPX-6-40-135	6	40	6	0.014	0.078
SEPX-7-40-135	7	40	6	0.014	0.078
SEPX-8-40-135	8	40	6	0.014	0.078
SEPX-9-40-135	9	40	6	0.014	0.078
SEPX-10-40-135	10	40	6	0.014	0.078
SEPX-6-60-135	6	60	6	0.014	0.078
SEPX-7-60-135	7	60	6	0.014	0.078
SEPX-8-60-135	8	60	6	0.014	0.078
SEPX-9-60-135	9	60	6	0.014	0.078
SEPX-10-60-135	10	60	6	0.014	0.078

INDICATIONS: The Protégé™ Rx™ is indicated for use in the iliac or subclavian arteries in the palliative treatment of malignant neoplasms in the biliary tree. It is also indicated for treatment of stenoses of the common carotid artery (CCA), internal carotid artery (ICA) and carotid bifurcation.

## Balloon-Expandable Peripheral Stent System

- Broad offering of 6 Fr-compatible 0.035" balloon-expandable stent with radiopaque marker technology for optimized visibility.
- Low crossing profile.
- Minimal shortening for placement confidence.

VisiPro™ catheter lengths  
80 cm and 135 cm

**Each system includes:**

One stent and delivery catheter system



### COMPLIANCE CHART

Diameter (mm)	Inflation Pressure (atm)				
	8	9	10	11	12
5.0	5.00 <sup>1</sup>	5.09	5.16	5.22	5.28 <sup>2</sup>
6.0	6.00 <sup>1</sup>	6.11	6.22	6.31	6.39 <sup>2</sup>
7.0			7.00 <sup>1</sup>	7.09	7.17 <sup>2</sup>
8.0			8.00 <sup>1</sup>	8.15	8.26 <sup>2</sup>
9.0			9.00 <sup>1</sup>	9.15	9.28 <sup>2</sup>
10.0			10.00 <sup>1</sup>	10.11	10.21 <sup>2</sup>

<sup>1</sup>Diameter at Nominal Pressure

<sup>2</sup>Diameter at Rated Burst Pressure

## Balloon-Expandable Peripheral Stent System

### ORDER INFORMATION

Product Catalogue Number		Stent dimensions		Balloon Length (mm)	Recommended Sheath Size (F)	Recommended Guidewire (inch)	Crossing Profile (inch)
Catheter Length 80 cm	Catheter Length 135 cm	Diameter (mm)	Length (mm)				
PXP35-05-12-080							
PXP35-05-17-080	PXP35-05-17-135	5.0	17	20	6	0.035	0.079
PXP35-05-27-080	PXP35-05-27-135	5.0	27	30	6	0.035	0.079
PXP35-05-37-080	PXP35-05-37-135	5.0	37	40	6	0.035	0.079
PXP35-05-57-080	PXP35-05-57-135	5.0	57	60	6	0.035	0.079
PXP35-06-12-080		6.0	12	15	6	0.035	0.079
PXP35-06-17-080	PXP35-06-17-135	6.0	17	20	6	0.035	0.079
PXP35-06-27-080	PXP35-06-27-135	6.0	27	30	6	0.035	0.079
PXP35-06-37-080	PXP35-06-37-135	6.0	37	40	6	0.035	0.081
PXP35-06-57-080	PXP35-06-57-135	6.0	57	60	6	0.035	0.083
PXP35-07-12-080		7.0	12	15	6	0.035	0.079
PXP35-07-17-080	PXP35-07-17-135	7.0	17	20	6	0.035	0.079
PXP35-07-27-080	PXP35-07-27-135	7.0	27	30	6	0.035	0.079
PXP35-07-37-080	PXP35-07-37-135	7.0	37	40	6	0.035	0.081
PXP35-07-57-080	PXP35-07-57-135	7.0	57	60	6	0.035	0.083
PXP35-08-17-080	PXP35-08-17-135	8.0	17	20	6	0.035	0.083
PXP35-08-27-080	PXP35-08-27-135	8.0	27	30	6	0.035	0.083
PXP35-08-37-080	PXP35-08-37-135	8.0	37	40	6	0.035	0.083
PXP35-08-57-080	PXP35-08-57-135	8.0	57	60	6	0.035	0.084
PXP35-09-17-080	PXP35-09-17-135	9.0	17	20	7	0.035	0.088
PXP35-09-27-080	PXP35-09-27-135	9.0	27	30	7	0.035	0.088
PXP35-09-37-080	PXP35-09-37-135	9.0	37	40	7	0.035	0.088
PXP35-09-57-080	PXP35-09-57-135	9.0	57	60	7	0.035	0.088
PXP35-10-17-080	PXP35-10-17-135	10.0	17	20	7	0.035	0.092
PXP35-10-27-080	PXP35-10-27-135	10.0	27	30	7	0.035	0.092
PXP35-10-37-080	PXP35-10-37-135	10.0	37	40	7	0.035	0.092
PXP35-10-57-080	PXP35-10-57-135	10.0	57	60	7	0.035	0.092

Specifications Nominal  
Balloon Expandable Peripheral Stent System is indicated for use in the iliac, renal or subclavian arteries, as well as malignant biliary use.

# Protégé™ GPS™

## Self-Expanding Stent System

The Protégé™ GPS™ stent system gives control for precise stent placement avoiding jumping of the stent through the EX.P.R.T.™ retention system.

### Compact delivery

- Diameters up to 14 mm
- Full line is 6 F compatible

### Precision

- Proprietary EX.P.R.T.™ deployment system secures the stent to eliminate premature deployment or “jumping”
- Tantalum GPS markers enhance visibility for easier, precise positioning

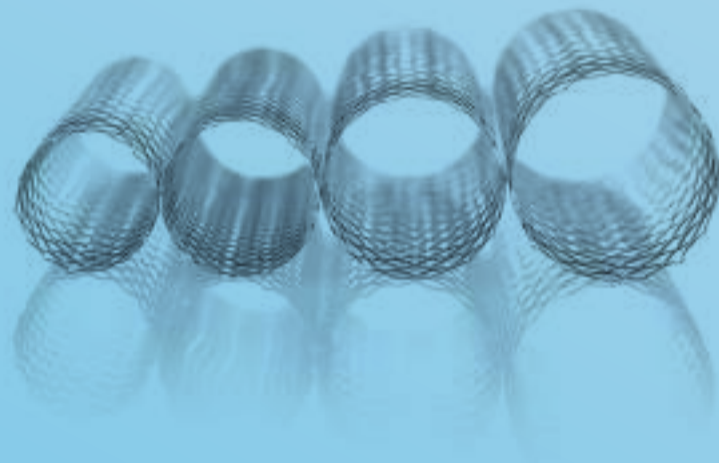
### Radial strength and flexibility

- Designed for radial strength without sacrificing flexibility

### Each system includes:

One stent and delivery catheter system

Protégé GPS catheter lengths 80 cm and 120 cm



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### ORDER INFORMATION

Product Catalogue Number		Stent Dimensions		Recommended		Recommended Guidewire (inch)	Crossing Profile (inch)
Catheter Length 80 cm	Catheter Length 120 cm	Diameter (mm)	Length (mm)	Lumen Size (mm)	Sheath Size (F)		
SERP65-09-20-80	SERP65-09-20-120	9	20	7.5 - 8.5	6	0.035	0.079
SERP65-09-30-80	SERP65-09-30-120	9	30	7.5 - 8.5	6	0.035	0.079
SERP65-09-40-80	SERP65-09-40-120	9	40	7.5 - 8.5	6	0.035	0.079
SERP65-09-60-80	SERP65-09-60-120	9	60	7.5 - 8.5	6	0.035	0.079
SERP65-09-80-80	SERP65-09-80-120	9	80	7.5 - 8.5	6	0.035	0.079
SERP65-10-20-80	SERP65-10-20-120	10	20	8.5 - 9.5	6	0.035	0.079
SERP65-10-30-80	SERP65-10-30-120	10	30	8.5 - 9.5	6	0.035	0.079
SERP65-10-40-80	SERP65-10-40-120	10	40	8.5 - 9.5	6	0.035	0.079
SERP65-10-60-80	SERP65-10-60-120	10	60	8.5 - 9.5	6	0.035	0.079
SERP65-10-80-80	SERP65-10-80-120	10	80	8.5 - 9.5	6	0.035	0.079
SERP65-12-20-80	SERP65-12-20-120	12	20	9.5 - 11.0	6	0.035	0.079
SERP65-12-30-80	SERP65-12-30-120	12	30	9.5 - 11.0	6	0.035	0.079
SERP65-12-40-80	SERP65-12-40-120	12	40	9.5 - 11.0	6	0.035	0.079
SERP65-12-60-80	SERP65-12-60-120	12	60	9.5 - 11.0	6	0.035	0.079
SERP65-12-80-80	SERP65-12-80-120	12	80	9.5 - 11.0	6	0.035	0.079
SERP65-14-20-80	SERP65-14-20-120	14	20	11.5 - 13.0	6	0.035	0.079
SERP65-14-30-80	SERP65-14-30-120	14	30	11.5 - 13.0	6	0.035	0.079
SERP65-14-40-80	SERP65-14-40-120	14	40	11.5 - 13.0	6	0.035	0.079
SERP65-14-60-80	SERP65-14-60-120	14	60	11.5 - 13.0	6	0.035	0.079
SERP65-14-80-80	SERP65-14-80-120	14	80	11.5 - 13.0	6	0.035	0.079

Specifications Nominal

INDICATIONS: The Protégé™ GPS™ stent is indicated for use in the iliac or subclavian arteries and malignant biliary use.

Indications, contraindications, warnings and instructions for use can be found in the product labeling supplied with each device.

Protected under one or more of the following: US Patent 6,814,746; 6,749,627; 6,623,518; 6,623,491; 6,558,415; 6,358,274; 6,132,460; D458,679. Non-US Patent pending.

Product availability and/or specifications subject to change.

# EverFlex™

## Self-Expanding Stent System

The EverFlex™ self-expanding peripheral stent system is a self-expanding nitinol stent system.

The spiral-cell interconnecting design significantly improves flexibility and vessel conformability, without sacrificing radial strength. Excellent wall apposition and compression resistance is provided by the three-wave peak design.

### **Spiral cell connection**

- Peak-to-peak connection nodes to disperse force uniformly

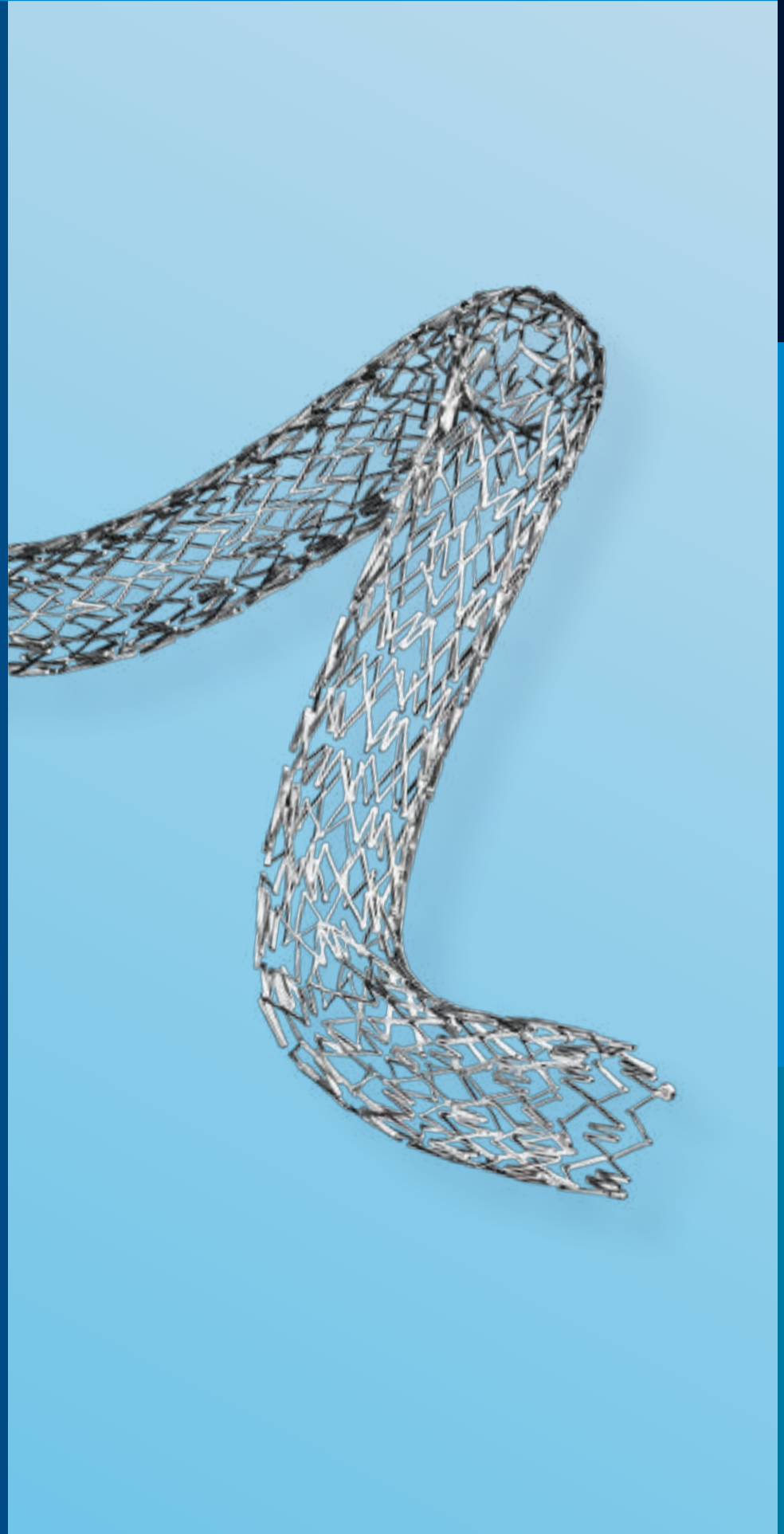
### **Flexible design**

- Improves fracture resistance and restores vessel patency

### **Three-wave peak design**

- Designed to resist compression and provide wall apposition

EverFlex™ catheter lengths: 80 cm and 120 cm



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## Self-Expanding Stent System

### ORDER INFORMATION

Product Catalogue Number		Stent Dimensions		Recommended		Recommended Guidewire (inch)	Crossing Profile (inch)
Catheter Length 80 cm	Catheter Length 120 cm	Diameter (mm)	Length (mm)	Lumen Size (mm)	Sheath Size (F)		
PRP35-05-020-080	PRP35-05-020-120	5	20	3.5 - 4.5	6	0.035	0.079
PRP35-05-030-080	PRP35-05-030-120	5	30	3.5 - 4.5	6	0.035	0.079
PRP35-05-040-080	PRP35-05-040-120	5	40	3.5 - 4.5	6	0.035	0.079
PRP35-05-060-080	PRP35-05-060-120	5	60	3.5 - 4.5	6	0.035	0.079
PRP35-05-080-080	PRP35-05-080-120	5	80	3.5 - 4.5	6	0.035	0.079
PRP35-05-100-080	PRP35-05-100-120	5	100	3.5 - 4.5	6	0.035	0.079
PRP35-05-120-080	PRP35-05-120-120	5	120	3.5 - 4.5	6	0.035	0.079
PRP35-05-150-080	PRP35-05-150-120	5	150	4.5 - 5.5	6	0.035	0.079
PRP35-06-020-080	PRP35-06-020-120	6	20	4.5 - 5.5	6	0.035	0.079
PRP35-06-030-080	PRP35-06-030-120	6	30	4.5 - 5.5	6	0.035	0.079
PRP35-06-040-080	PRP35-06-040-120	6	40	4.5 - 5.5	6	0.035	0.079
PRP35-06-060-080	PRP35-06-060-120	6	60	4.5 - 5.5	6	0.035	0.079
PRP35-06-080-080	PRP35-06-080-120	6	80	4.5 - 5.5	6	0.035	0.079
PRP35-06-100-080	PRP35-06-100-120	6	100	4.5 - 5.5	6	0.035	0.079
PRP35-06-120-080	PRP35-06-120-120	6	120	4.5 - 5.5	6	0.035	0.079
PRP35-06-150-080	PRP35-06-150-120	6	150	4.5 - 5.5	6	0.035	0.079
-	PRP35DR-06-200-120	6	200	4.5 - 5.5	6	0.035	0.079
PRP35-07-020-080	PRP35-07-020-120	7	20	5.5 - 6.5	6	0.035	0.079
PRP35-07-030-080	PRP35-07-030-120	7	30	5.5 - 6.5	6	0.035	0.079
PRP35-07-040-080	PRP35-07-040-120	7	40	5.5 - 6.5	6	0.035	0.079
PRP35-07-060-080	PRP35-07-060-120	7	60	5.5 - 6.5	6	0.035	0.079
PRP35-07-080-080	PRP35-07-080-120	7	80	5.5 - 6.5	6	0.035	0.079
PRP35-07-100-080	PRP35-07-100-120	7	100	5.5 - 6.5	6	0.035	0.079
PRP35-07-120-080	PRP35-07-120-120	7	120	5.5 - 6.5	6	0.035	0.079
PRP35-07-150-080	PRP35-07-150-120	7	150	5.5 - 6.5	6	0.035	0.079
-	PRP35DR-07-200-120	7	200	5.5 - 6.5	6	0.035	0.079
PRP35-08-020-080	PRP35-08-020-120	8	20	6.5 - 7.5	6	0.035	0.079
PRP35-08-030-080	PRP35-08-030-120	8	30	6.5 - 7.5	6	0.035	0.079
PRP35-08-040-080	PRP35-08-040-120	8	40	6.5 - 7.5	6	0.035	0.079
PRP35-08-060-080	PRP35-08-060-120	8	60	6.5 - 7.5	6	0.035	0.079
PRP35-08-080-080	PRP35-08-080-120	8	80	6.5 - 7.5	6	0.035	0.079
PRP35-08-100-080	PRP35-08-100-120	8	100	6.5 - 7.5	6	0.035	0.079
PRP35-08-120-080	PRP35-08-120-120	8	120	6.5 - 7.5	6	0.035	0.079
PRP35-08-150-080	PRP35-08-150-120	8	150	6.5 - 7.5	6	0.035	0.079
-	PRP35DR-08-200-120	8	200	6.5 - 7.5	6	0.035	0.079

INDICATIONS: The EverFlex™ self-expanding peripheral stent system is indicated for use in common iliac, external iliac, superficial femoral, proximal popliteal, and subclavian arteries.

# EverFlex™ with Entrust™ Delivery System

## Self-Expanding Stent System



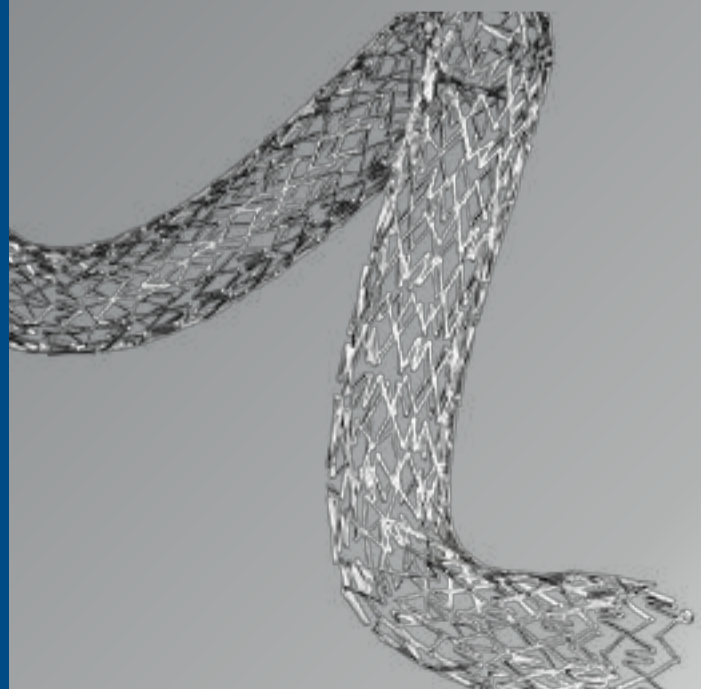
The Entrust™ delivery system is a one-handed, triaxialstent delivery system with a low 5 F profile.

This low profile was achieved without compromising the design of the EverFlex™ stent or the 0.035" guidewire compatibility.

The device was engineered specifically for control and accuracy based on physician feedback provided during extensive interviews and procedural observations.

EverFlex™ stent: The DURABILITY II study proves strong stent performance with a 60% primary patency at 3 years.

EverFlex™ Entrust™ catheter lengths: 80 cm, 120 cm, 150 cm.



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# EverFlex™ with Entrust™ Delivery System

## Self-Expanding Stent System

### ORDER INFORMATION

Product Catalogue Number			Stent Dimensions (Unconstrained)		Size Compatibility		Guidewire Acceptance (inch)
Catheter Length 80 cm	Catheter Length 120 cm	Catheter Length 150 cm	Diameter (mm)	Length (mm)	Vessel Size (mm)	Sheath / Guide (F)	
EVX35-05-020-080	EVX35-05-020-120	EVX35-05-020-150	5	20	3.5 - 4.5	5	0.035
EVX35-05-040-080	EVX35-05-040-120	EVX35-05-040-150	5	40	3.5 - 4.5	5	0.035
EVX35-05-060-080	EVX35-05-060-120	EVX35-05-060-150	5	60	3.5 - 4.5	5	0.035
EVX35-05-080-080	EVX35-05-080-120	EVX35-05-080-150	5	80	3.5 - 4.5	5	0.035
EVX35-05-100-080	EVX35-05-100-120	EVX35-05-100-150	5	100	3.5 - 4.5	5	0.035
EVX35-05-120-080	EVX35-05-120-120	EVX35-05-120-150	5	120	3.5 - 4.5	5	0.035
EVX35-05-150-080	EVX35-05-150-120	EVX35-05-150-150	5	150	3.5 - 4.5	5	0.035
EVX35-06-020-080	EVX35-06-020-120	EVX35-06-020-150	6	20	4.5 - 5.5	5	0.035
EVX35-06-040-080	EVX35-06-040-120	EVX35-06-040-150	6	40	4.5 - 5.5	5	0.035
EVX35-06-060-080	EVX35-06-060-120	EVX35-06-060-150	6	60	4.5 - 5.5	5	0.035
EVX35-06-080-080	EVX35-06-080-120	EVX35-06-080-150	6	80	4.5 - 5.5	5	0.035
EVX35-06-100-080	EVX35-06-100-120	EVX35-06-100-150	6	100	4.5 - 5.5	5	0.035
EVX35-06-120-080	EVX35-06-120-120	EVX35-06-120-150	6	120	4.5 - 5.5	5	0.035
EVX35-06-150-080	EVX35-06-150-120	EVX35-06-150-150	6	150	4.5 - 5.5	5	0.035
EVX35-07-020-080	EVX35-07-020-120	EVX35-07-020-150	7	20	5.5 - 6.5	5	0.035
EVX35-07-040-080	EVX35-07-040-120	EVX35-07-040-150	7	40	5.5 - 6.5	5	0.035
EVX35-07-060-080	EVX35-07-060-120	EVX35-07-060-150	7	60	5.5 - 6.5	5	0.035
EVX35-07-080-080	EVX35-07-080-120	EVX35-07-080-150	7	80	5.5 - 6.5	5	0.035
EVX35-07-100-080	EVX35-07-100-120	EVX35-07-100-150	7	100	5.5 - 6.5	5	0.035
EVX35-07-120-080	EVX35-07-120-120	EVX35-07-120-150	7	120	5.5 - 6.5	5	0.035
EVX35-07-150-080	EVX35-07-150-120	EVX35-07-150-150	7	150	5.5 - 6.5	5	0.035
EVX35-08-020-080	EVX35-08-020-120	EVX35-08-020-150	8	20	6.5 - 7.5	5	0.035
EVX35-08-040-080	EVX35-08-040-120	EVX35-08-040-150	8	40	6.5 - 7.5	5	0.035
EVX35-08-060-080	EVX35-08-060-120	EVX35-08-060-150	8	60	6.5 - 7.5	5	0.035
EVX35-08-080-080	EVX35-08-080-120	EVX35-08-080-150	8	80	6.5 - 7.5	5	0.035
EVX35-08-100-080	EVX35-08-100-120	EVX35-08-100-150	8	100	6.5 - 7.5	5	0.035
EVX35-08-120-080	EVX35-08-120-120	EVX35-08-120-150	8	120	6.5 - 7.5	5	0.035
EVX35-08-150-080	EVX35-08-150-120	EVX35-08-150-150	8	150	6.5 - 7.5	5	0.035

INDICATIONS: The stent is indicated for use in occlusions, lesions at high risk for abrupt closure or threatened closure following percutaneous transluminal angioplasty (PTA), or lesions believed to be at high risk for restenosis following PTA in the common iliac, external iliac, superficial femoral, proximal popliteal, or subclavian arteries. Stenting is intended to improve and maintain artery luminal diameter.

# Paramount Mini™ GPS™

## Balloon-Expandable Peripheral Stent System

The Paramount Mini™ GPS™ is a pre-mounted renal stent line with tantalum markers on a balloon catheter delivery system. The devices are compatible with 5 and 6 F introducers and 0.014" and 0.018" Guidewires.

**Each kit includes:**

One stent and delivery catheter system

Paramount mini catheter length 80cm



### COMPLIANCE CHART

ParaMount Mini™ GPS™ Diameter (mm)	Inflation Pressure (atm)			
	9	10	11	12
5.0	4.96	5.04 <sup>1</sup>	5.12	5.20 <sup>2</sup>
6.0	5.78	5.88 <sup>1</sup>	5.98	6.08 <sup>2</sup>

<sup>1</sup>Diameter at Nominal Pressure

<sup>2</sup>Diameter at Rated Burst Pressure

INDICATIONS: The ParaMount Mini™ GPS™ Stent System is indicated for use in the renal artery, as well as malignant biliary use.

# Paramount Mini™ GPS™

## Balloon-Expandable Peripheral Stent System

### ORDER INFORMATION

Product Catalogue Number	Expanded Stent Size		Balloon Length (mm)	Usable Length (cm)	Rated Burst Pressure (atm)	Nominal Burst Pressure (atm)	Recommended Guide / Catheter Sheath Size (inch)	Recommended Guidewire (inch)	Crossing Profile (inch)
	Diameter (mm)	Length (mm)							
PMP4-5-14-80	5.0	14	17	80	12	10	6 / 5	0.014	0.062
PMP4-5-18-80	5.0	18	20	80	12	10	6 / 5	0.014	0.062
PMP4-5-21-80	5.0	21	24	80	12	10	6 / 5	0.014	0.062
PMP4-6-14-80	6.0	14	17	80	12	10	7 / 6	0.014	0.066
PMP4-6-18-80	6.0	18	20	80	12	10	7 / 6	0.014	0.066
PMP4-6-21-80	6.0	21	24	80	12	10	7 / 6	0.014	0.066
PMP4-7-14-80	7.0	14	17	80	12	10	7 / 6	0.014	0.070
PMP4-7-18-80	7.0	18	20	80	12	10	7 / 6	0.014	0.070
PMP4-7-21-80	7.0	21	24	80	12	10	7 / 6	0.014	0.070
PMP8-5-14-80	5.0	14	17	80	12	10	6 / 5	0.018	0.062
PMP8-5-18-80	5.0	18	20	80	12	10	6 / 5	0.018	0.062
PMP8-5-21-80	5.0	21	24	80	12	10	6 / 5	0.018	0.062
PMP8-6-14-80	6.0	14	17	80	12	10	6* / 5	0.018	0.066
PMP8-6-18-80	6.0	18	20	80	12	10	6* / 5	0.018	0.066
PMP8-6-21-80	6.0	21	24	80	12	10	6* / 5	0.018	0.066
PMP8-7-14-80	7.0	14	17	80	12	10	7 / 6	0.018	0.070
PMP8-7-18-80	7.0	18	20	80	12	10	7 / 6	0.018	0.070
PMP8-7-21-80	7.0	21	24	80	12	10	7 / 6	0.018	0.070

Specifications Nominal, 6 F=0.070" I.D.

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# Hippocampus™

## Renal RX Stent System 0.014"

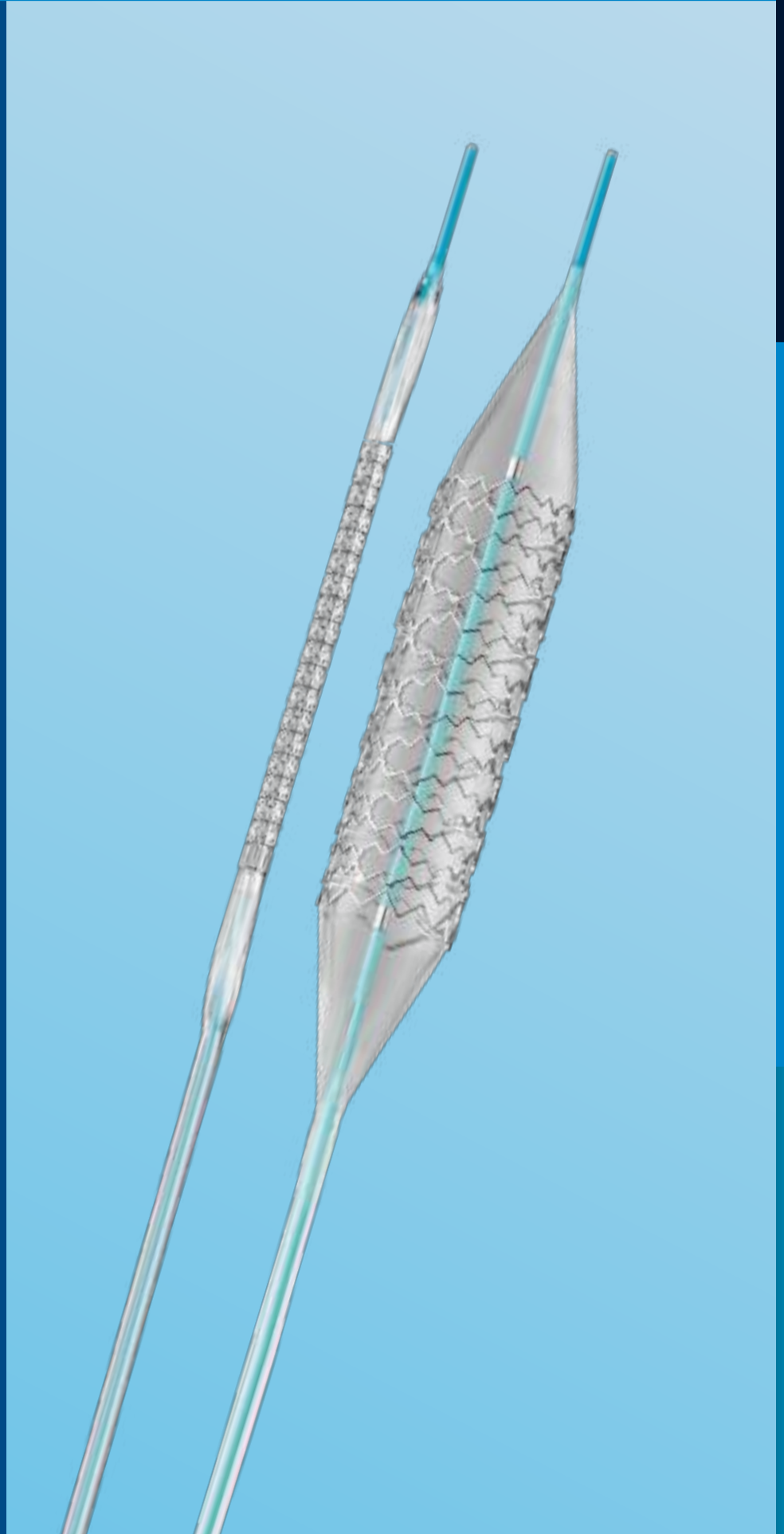
### TECHNICAL SPECIFICATIONS

#### STENT

Stent design	Closed cell
Stent material	Stainless Steel
Stent inner diameter	4.0, 5.0, 5.5, 6.0, 6.5, 7.0 mm
Stent length	10, 15, 20, 24 mm
Strut thickness / width	165 / 110 µm

#### STENT DELIVERY SYSTEM

Catheter design	RX (Rapid Exchange)
Shaft diameter prox./ dist.	2.3 F / 3.5 F
Usable shaft length	80 and 145 cm
Recommended guidewire	0.014"
Introducer sheath compatibility	5 F
Guiding catheter compatibility	6 F (> 0.066")
Nominal Pressure	8 bar



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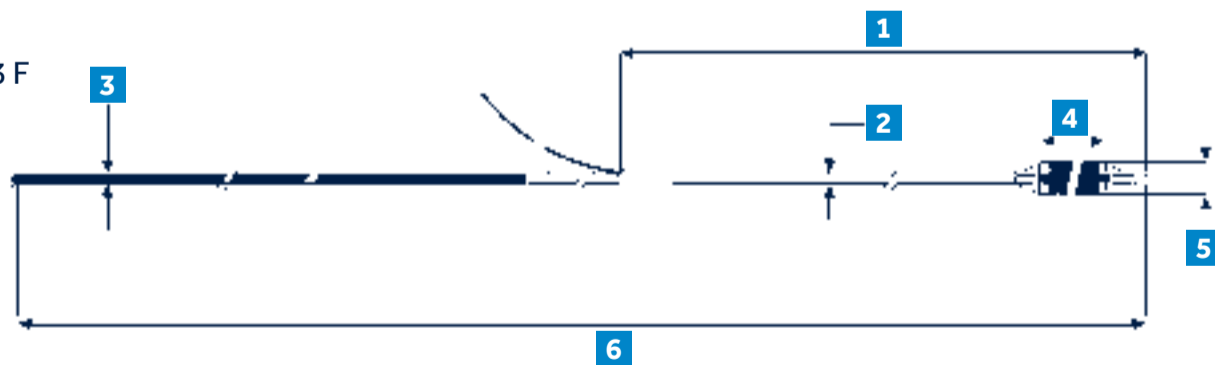
### ORDER INFORMATION

Product Catalogue Number		Stent inner Ø (mm)	Stent length (mm)	RBP (bar)
Usable length 80 cm	Usable length 145 cm			
IHP040 100 080	IHP040 100 145	4.0	10	15
IHP040 150 080	IHP040 150 145	4.0	15	15
IHP040 200 080	IHP040 200 145	4.0	20	15
IHP050 100 080	IHP050 100 145	5.0	10	15
IHP050 150 080	IHP050 150 145	5.0	15	15
IHP050 200 080	IHP050 200 145	5.0	20	15
IHP050 240 080	IHP050 240 145	5.0	24	15
IHP055 100 080	IHP055 100 145	5.5	10	15
IHP055 150 080	IHP055 150 145	5.5	15	15
IHP055 200 080	IHP055 200 145	5.5	20	15
IHP060 100 080	IHP060 100 145	6.0	10	14
IHP060 150 080	IHP060 150 145	6.0	15	14
IHP060 200 080	IHP060 200 145	6.0	20	14
IHP060 240 080	IHP060 240 145	6.0	24	14
IHP065 150 080	IHP065 150 145	6.5	15	14
IHP065 200 080	IHP065 200 145	6.5	20	14
IHP070 150 080	IHP070 150 145	7.0	15	14
IHP070 200 080	IHP070 200 145	7.0	20	14
IHP070 240 080	IHP070 240 145	7.0	24	14

- 1 RX section, 15 cm
- 2 Distal shaft diameter 3.5 F
- 3 Proximal shaft diameter 2.3 F



- 4 Stent length
- 5 Stent inner diameter
- 6 Usable shaft length



# IntraStent™ LD

## Large Diameter Stents

The IntraStent™ LD stent family of large-lumen stainless steel stents has been designed to supply a larger diameter device with the flexibility, strength, coverage, and profile normally associated with smaller diameter stents.

**Three models are available:**

### **IntraStent™ DoubleStrut™ LD peripheral stent**

- Four-cell design for balanced radial strength and flexibility
- Low profile, 8 F introducer sheath
- Best choice for flexibility

### **IntraStent™ Mega™ LD peripheral stent**

- Five-cell design for increased radial strength over the DoubleStrut LD Stent
- 9 F sheath compatibility
- Best choice for strength and flexibility

### **IntraStent™ Max™ LD peripheral stent**

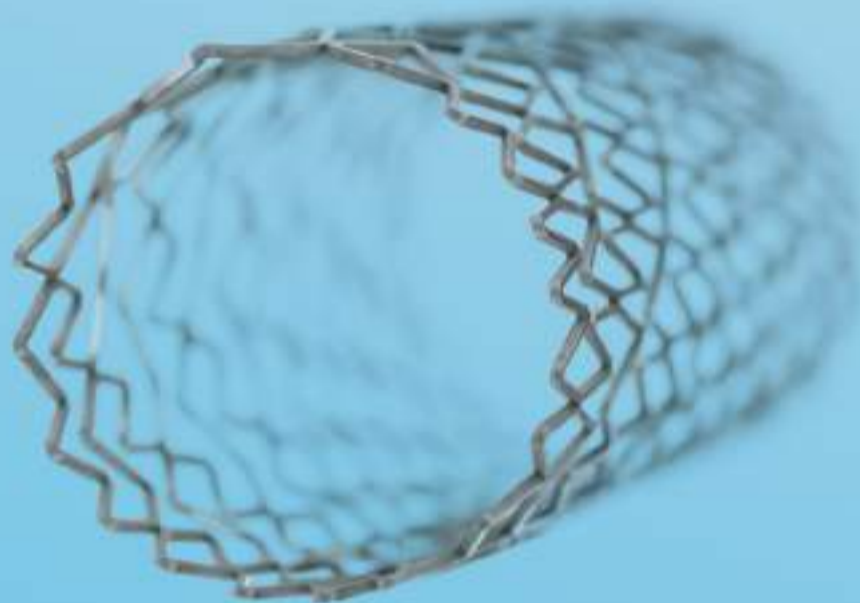
- Six-cell design for maximum strength and coverage
- 11 F sheath compatibility
- Best for strength

### **More choices**

- The LD series of stents is available in a variety of designs to accommodate the unique needs of your patients

### **Innovative design**

- Flexibility, strength and coverage delivered in ways not found in other large lumen stents
- Minimal shortening after expansion up to 12 mm
- Rounded edges minimize potential for lumen trauma





# IntraStent™ LD

## Large Diameter Stents

### INTRASTENT™ LD STENT FAMILY

Product Catalogue Number	Un-expanded Stent Size		Expanded Stent Size	
	Diameter (mm)	Length (mm)	Diameter (mm)	Length (mm)
<b>Intrastent™ LD Doublestrut™</b>				
90-1504-000	3.8	16.0	9, 10, 11, 12	16.0
90-1504-001	3.8	26.0	9, 10, 11, 12	26.0
90-1504-002	3.8	36.0	9, 10, 11, 12	36.0
90-1504-003	3.8	56.0	9, 10, 11, 12	56.0
90-1504-004	3.8	76.0	9, 10, 11, 12	76.0
<b>Intrastent™ LD Mega™</b>				
90-2336-000	3.8	16.0	9, 10, 11, 12	16.0
90-2336-001	3.8	26.0	9, 10, 11, 12	26.0
90-2336-002	3.8	36.0	9, 10, 11, 12	36.0
<b>Intrastent™ LD Max™</b>				
90-2337-000	4.5	16.0	12	16.0
90-2337-001	4.5	26.0	12	26.0
90-2337-002	4.5	36.0	12	36.0

Specifications Nominal

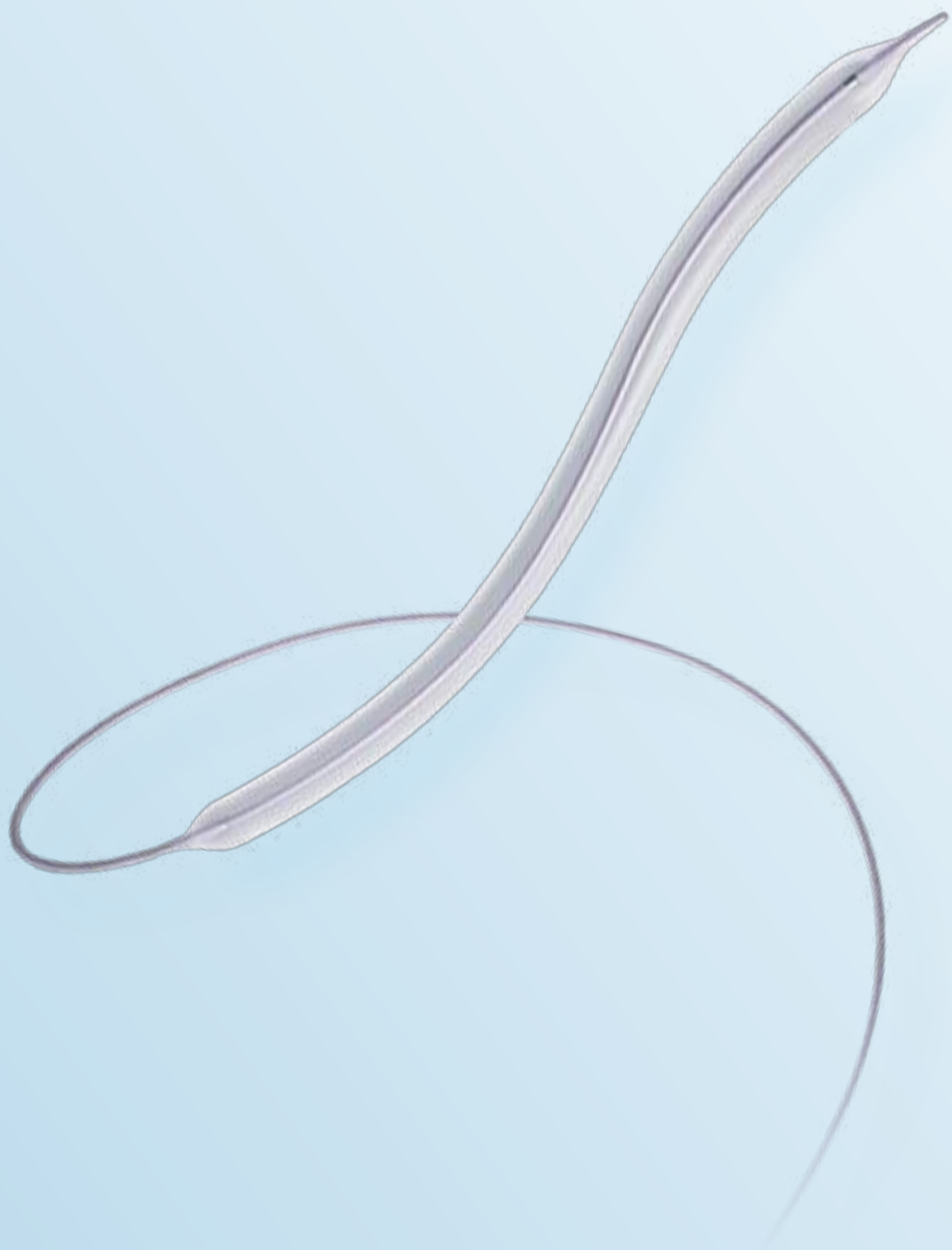
INDICATIONS: The IntraStent™ LD Double Strut™, IntraStent™ LD Mega™ and the IntraStent™ LD Max™ stents are indicated for use in iliac and subclavian arteries. The IntraStent™ LD Double Strut™ is also indicated for malignant biliary use.

### INTRASTENT™ LD STENT MEGA™ AND LD MAX™ STENT EXPANSION CHART

Stent Expanded Diameter (mm)	IntraStent™ LD Mega™ Stent Lengths (mm)			IntraStent™ LD Max™ Stent Lengths (mm)		
	16	26	36	16	26	36
9	16.0	26.0	36.0	16.0	26.0	36.0
10	16.0	26.0	36.0	16.0	26.0	36.0
12	16.0	26.0	36.0	16.0	26.0	36.0
14	14.0	24.0	34.0	15.5	25.5	35.5
16	13.0	22.5	32.5	15.0	25.0	35.0
18	12.0	21.5	31.0	14.5	24.5	34.5
20				14.0	24.0	34.0
22				13.5	23.0	33.0
25				13.0	22.0	32.0

Stent was expanded in a single increment. Stepped expansion will result in less shortening of the stent.

# PTA BALLOONS



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# Admiral™ Xtreme™

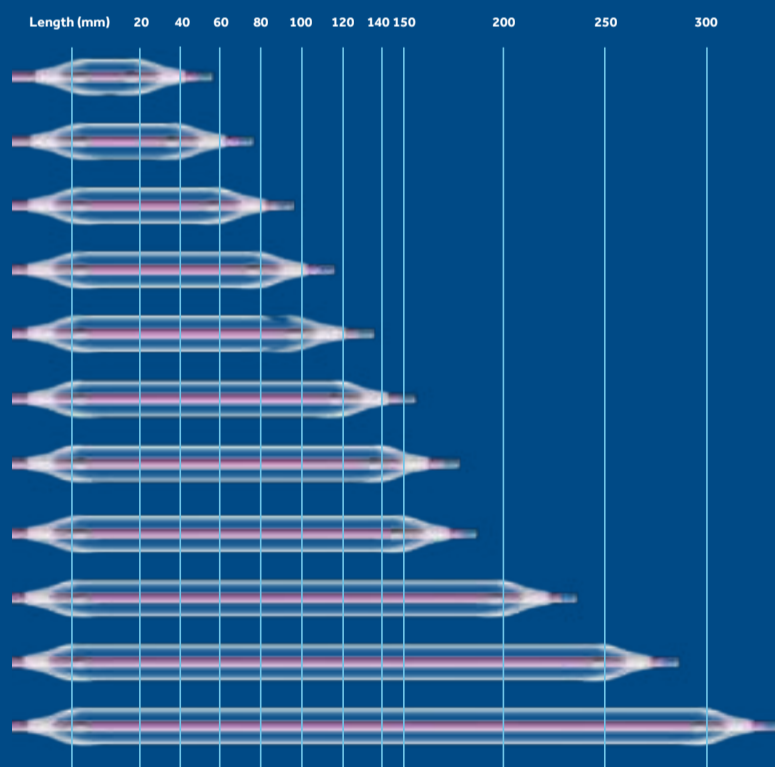
PTA Balloon Catheter OTW 0.035"

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## CROSS LESIONS WITH CONFIDENCE

### TECHNICAL SPECIFICATIONS

Catheter design	Over the Wire (OTW)
Balloon coating	LFC Hydrophilic
Balloon marker	2 swaged (zero profile) Platinum Iridium
Shaft diameter	5 F – 5.3 F
Usable shaft lengths	80 cm, 130 cm, 150 cm
Introducer sheath compatibility	5 F – 6 F – 7 F
Guidewire compatibility	0.035"



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# Admiral™ Xtreme™

## PTA Balloon Catheter OTW 0.035"

### ORDER INFORMATION

Ref. N° Usable Length 80 cm	Ref. N° Usable Length 130 cm	Ref. N° Usable Length 150 cm	Balloon Ø (mm)	Balloon Length (mm)	Recom. Introducer Sheath (F)	RBP (Bar)
SBI030020080	SBI030020130		3	20	5	18
SBI030040080	SBI030040130		3	40	5	18
SBI030080080	SBI030080130		3	80	5	18
SBI030100080	SBI030100130		3	100	5	18
SBI030120080	SBI030120130		3	120	5	18
SBI040020080	SBI040020130	SBI040020150	4	20	5	18
SBI040040080	SBI040040130	SBI040040150	4	40	5	18
SBI040060080	SBI040060130	SBI040060150	4	60	5	18
SBI040080080	SBI040080130	SBI040080150	4	80	5	18
SBI040100080	SBI040100130	SBI040100150	4	100	5	18
SBI040120080	SBI040120130	SBI040120150	4	120	5	18
SBI040150080	SBI040150130	SBI040150150	4	150	5	14
SBI040200080	SBI040200130	SBI040200150	4	200	5	14
SBI040250080	SBI040250130		4	250	5	14
SBI040300080	SBI040300130		4	300	5	14
SBI050020080	SBI050020130	SBI050020150	5	20	5	17
SBI050040080	SBI050040130	SBI050040150	5	40	5	17
SBI050060080	SBI050060130	SBI050060150	5	60	5	17
SBI050080080	SBI050080130	SBI050080150	5	80	5	15
SBI050100080	SBI050100130	SBI050100150	5	100	5	15
SBI050120080	SBI050120130	SBI050120150	5	120	5	15
SBI050150080	SBI050150130	SBI050150150	5	150	5	14
SBI050200080	SBI050200130	SBI050200150	5	200	5	14
SBI050250080	SBI050250130		5	250	5	14
SBI050300080L	SBI050300130L		5	300	5	14
SBI060020080	SBI060020130	SBI060020150	6	20	5	17
SBI060040080	SBI060040130	SBI060040150	6	40	5	17
SBI060060080	SBI060060130	SBI060060150	6	60	5	17
SBI060080080	SBI060080130	SBI060080150	6	80	5	15
SBI060100080	SBI060100130	SBI060100150	6	100	5	15
SBI060120080	SBI060120130	SBI060120150	6	120	5	15
SBI060150080	SBI060150130	SBI060150150	6	150	5	12
SBI060200080L	SBI060200130L	SBI060200150	6	200	5	12
SBI060250080L	SBI060250130L		6	250	5	12
SBI060300080L	SBI060300130L		6	300	5	12
SBI070020080	SBI070020130	SBI070020150	7	20	5	16
SBI070040080	SBI070040130	SBI070040150	7	40	5	16
SBI070060080	SBI070060130	SBI070060150	7	60	5	14
SBI070080080	SBI070080130	SBI070080150	7	80	5	14
SBI070100080	SBI070100130	SBI070100150	7	100	5	12
SBI070120080L	SBI070120130L	SBI070120150	7	120	5	12
SBI070150080L	SBI070150130L	SBI070150150	7	150	5	12
SBI070200080L	SBI070200130L	SBI070200150	7	200	5	12
SBI070250080	SBI070250130		7	250	6	12
SBI080020080	SBI080020130		8	20	6	14
SBI080040080	SBI080040130		8	40	6	14
SBI080060080	SBI080060130		8	60	6	11
SBI080080080	SBI080080130		8	80	6	11
SBI090020080	SBI090020130		9	20	6	14
SBI090040080	SBI090040130		9	40	6	14
SBI090060080	SBI090060130		9	60	6	11
SBI090080080	SBI090080130		9	80	6	11
SBI100020080	SBI100020130		10	20	6	11
SBI100040080	SBI100040130		10	40	6	11
SBI120020080	SBI120020130		12	20	7	11
SBI120040080	SBI120040130		12	40	7	11

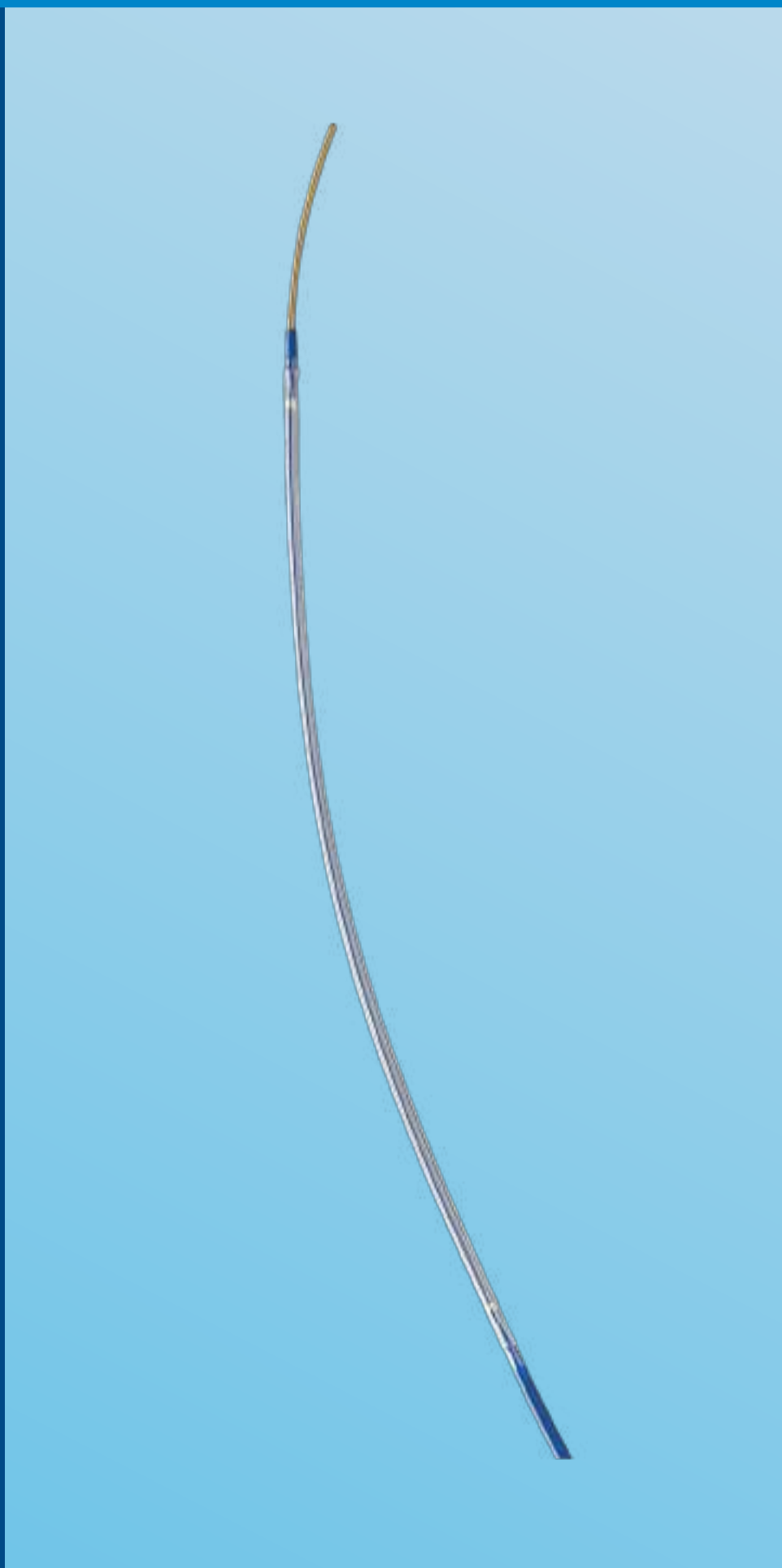
# EverCross™

## OTW PTA Dilatation Catheter 0.035"

EverCross™ 0.035" PTA balloon is an over-the-wire, 0.035" balloon catheter that features a bevel 360° tip for smooth tip to wire tracking. EverCross™ nylon folds, extending the length of the balloon, were engineered for superior rewrap, facilitating multiple inflations and insertions. Each system includes: One PTA balloon catheter and one compliance chart.

### TECHNICAL SPECIFICATIONS

Catheter design	Over-the-wire
Useable catheter lengths	40, 80 and 135cm
Introducer sheath compatibility	5, 6 F
Guidewire compatibility	0.035"



AORTIC

PERIPHERAL

VENOUS

## OTW PTA Dilatation Catheter 0.035"

### ORDER INFORMATION

Product Catalogue Number			Balloon Diameter (mm)	Balloon Length (mm)	Nominal Pressure (atm)	Rated Burst Pressure (atm)	Recommended Introducer Sheath (Fr)
Usable Shaft Length 135cm	Usable Shaft Length 80cm	Usable Shaft Length 40cm					
AB35W03020135	AB35W03020080	-	3.0	20	10	20	5
AB35W03030135	AB35W03030080	-	3.0	30	10	20	5
AB35W03040135	AB35W03040080	-	3.0	40	10	20	5
AB35W03060135	AB35W03060080	-	3.0	60	10	20	5
AB35W03080135	AB35W03080080	-	3.0	80	10	20	5
AB35W03100135	AB35W03100080	-	3.0	100	10	20	5
AB35W03120135	AB35W03120080	-	3.0	120	10	20	5
AB35W03150135	AB35W03150080	-	3.0	150	10	20	5
AB35W03200135	AB35W03200080	-	3.0	200	10	20	5
AB35W04020135	AB35W04020080	-	4.0	20	10	20	5
AB35W04030135	AB35W04030080	-	4.0	30	10	20	5
AB35W04040135	AB35W04040080	-	4.0	40	10	20	5
AB35W04060135	AB35W04060080	-	4.0	60	10	20	5
AB35W04080135	AB35W04080080	-	4.0	80	10	20	5
AB35W04100135	AB35W04100080	-	4.0	100	10	20	5
AB35W04120135	AB35W04120080	-	4.0	120	10	20	5
AB35W04150135	AB35W04150080	-	4.0	150	10	20	5
AB35W04200135	AB35W04200080	-	4.0	200	10	20	5
AB35W05020135	AB35W05020080	AB35W05020040	5.0	20	10	18	5
AB35W05030135	AB35W05030080	AB35W05030040	5.0	30	10	18	5
AB35W05040135	AB35W05040080	AB35W05040040	5.0	40	10	18	5
AB35W05060135	AB35W05060080	AB35W05060040	5.0	60	10	18	5
AB35W05080135	AB35W05080080	AB35W05080040	5.0	80	10	18	5
AB35W05100135	AB35W05100080	-	5.0	100	10	18	5
AB35W05120135	AB35W05120080	AB35W05120040	5.0	120	10	16	5
AB35W05150135	AB35W05150080	-	5.0	150	10	16	5
AB35W05200135	AB35W05200080	-	5.0	200	10	16	5
AB35W06020135	AB35W06020080	AB35W06020040	6.0	20	8	14	5
AB35W06030135	AB35W06030080	-	6.0	30	8	14	5

## OTW PTA Dilatation Catheter 0.035"

### ORDER INFORMATION

Product Catalogue Number			Balloon Diameter (mm)	Balloon Length (mm)	Nominal Pressure (atm)	Rated Burst Pressure (atm)	Recommended Introducer Sheath (Fr)
Usable Shaft Length 135cm	Usable Shaft Length 80cm	Usable Shaft Length 40cm					
AB35W06040135	AB35W06040080	AB35W06040040	6.0	40	8	14	5
AB35W06060135	AB35W06060080	-	6.0	60	8	14	5
AB35W06080135	AB35W06080080	AB35W06080040	6.0	80	8	14	5
AB35W06100135	AB35W06100080	-	6.0	100	8	14	5
AB35W06120135	AB35W06120080	AB35W06120040	6.0	120	8	12	5
AB35W06150135	AB35W06150080	-	6.0	150	8	12	5
AB35W06200135	AB35W06200080	-	6.0	200	8	11	6
AB35W07020135	AB35W07020080	AB35W07020040	7.0	20	7	14	5
AB35W07030135	AB35W07030080	-	7.0	30	7	14	5
AB35W07040135	AB35W07040080	AB35W07040040	7.0	40	7	14	5
AB35W07060135	AB35W07060080	AB35W07060040	7.0	60	7	14	6
AB35W07080135	AB35W07080080	-	7.0	80	7	14	6
AB35W07100135	AB35W07100080	-	7.0	100	7	14	6
AB35W07120135	AB35W07120080	-	7.0	120	7	10	6
AB35W07150135	AB35W07150080	-	7.0	150	7	10	6
AB35W07200135	AB35W07200080	-	7.0	200	7	10	6
AB35W08020135	AB35W08020080	AB35W08020040	8.0	20	7	14	6
AB35W08030135	AB35W08030080	-	8.0	30	7	14	6
AB35W08040135	AB35W08040080	AB35W08040040	8.0	40	7	14	6
AB35W08060135	AB35W08060080	AB35W08060040	8.0	60	7	14	6
AB35W08080135	AB35W08080080	-	8.0	80	7	14	6
AB35W09020135	AB35W09020080	-	9.0	20	7	12	6
AB35W09030135	AB35W09030080	-	9.0	30	7	12	6
AB35W09040135	AB35W09040080	-	9.0	40	7	12	6
AB35W09060135	AB35W09060080	-	9.0	60	7	12	6
AB35W09080135	AB35W09080080	-	9.0	80	7	12	6
AB35W10020135	AB35W10020080	-	10.0	20	7	11	6
AB35W10030135	AB35W10030080	-	10.0	30	7	11	6
AB35W10040135	AB35W10040080	-	10.0	40	7	11	6
AB35W10060135	AB35W10060080	-	10.0	60	7	11	7
AB35W12020135	AB35W12020080	-	12.0	20	7	10	7
AB35W12040135	AB35W12040080	-	12.0	40	7	10	7
AB35W12060135	AB35W12060080	-	12.0	60	7	10	7

INDICATIONS: The EverCross™ 0.035" over-the-wire PTA dilatation catheter is intended to dilate stenoses in the iliac, femoral, ilio-femoral, popliteal, infra-popliteal, renal arteries and for the treatment of obstructive lesions of native or synthetic arteriovenous dialysis fistulae. This device is also indicated for stent post-dilatation in the peripheral vasculature.

### Predictable high-pressure treatment for AV access lesions

The high-pressure Fortrex™ PTA Balloon is the next-generation solution for deliverability, predictability, and procedural efficiency.

#### Deliverability

- Low tip entry profile enables tight tracking on the wire
- Robust, flexible shaft design facilitates successful navigation in tortuous vessels

#### Procedural efficiency

- Rapid deflation time contributes to reduced procedure length
- Balloon material and wall thickness permit reliable balloon rewrap into the sheath

#### Predictability

- Balloon material and design allow for shape retention at higher pressure
- Focal pressure is exerted on the lesion for controlled, targeted treatment



Fortrex™ Balloon  
Pressure is directed toward the lesion



Competitive Design  
Pressure is lost longitudinally



<sup>1</sup> Coriolis-Competitive Cheat Sheet - RE-PV1461.p.6.8-9.

<sup>2</sup> Coriolis-Competitive Cheat Sheet - RE-PV1461.p.7.

<sup>3</sup> Coriolis-Competitive Cheat Sheet - RE-PV1461.p.5.16-17-17-Competitive



## PTA Balloon Catheter OTW 0.035"

### ORDER INFORMATION

Product Catalogue Number			Balloon Diameter (mm)	Balloon Length (mm)	Nominal Pressure (atm)	Rated Burst Pressure (atm)	Recommended Introducer Sheath (F)
135 cm Catheter Length	80 cm Catheter Length	40 cm Catheter Length					
A35HPV04020135	A35HPV04020080	A35HPV04020040	4	20	12	24	6
A35HPV04040135	A35HPV04040080	A35HPV04040040	4	40	12	24	6
A35HPV04080135	A35HPV04080080	A35HPV04080040	4	80	12	24	6
A35HPV04100135	A35HPV04100080	A35HPV04100040	4	100	12	24	6
A35HPV05020135	A35HPV05020080	A35HPV05020040	5	20	12	24	6
A35HPV05040135	A35HPV05040080	A35HPV05040040	5	40	12	24	6
A35HPV05080135	A35HPV05080080	A35HPV05080040	5	80	12	24	6
A35HPV05100135	A35HPV05100080	A35HPV05100040	5	100	12	24	6
A35HPV06020135	A35HPV06020080	A35HPV06020040	6	20	12	24	6
A35HPV06040135	A35HPV06040080	A35HPV06040040	6	40	12	24	6
A35HPV06080135	A35HPV06080080	A35HPV06080040	6	80	12	23	6
A35HPV06100135	A35HPV06100080	A35HPV06100040	6	100	12	23	6
A35HPV07020135	A35HPV07020080	A35HPV07020040	7	20	9	20	6
A35HPV07040135	A35HPV07040080	A35HPV07040040	7	40	9	20	6
A35HPV07080135	A35HPV07080080	A35HPV07080040	7	80	9	20	6
A35HPV07100135	A35HPV07100080	A35HPV07100040	7	100	9	20	6
A35HPV08040135	A35HPV08040080	A35HPV08040040	8	40	9	20	6
A35HPV08080135	A35HPV08080080	A35HPV08080040	8	80	9	19	6
A35HPV08100135	A35HPV08100080	A35HPV08100040	8	100	9	18	6
A35HPV09040135	A35HPV09040080	A35HPV09040040	9	40	9	18	7
A35HPV09080135	A35HPV09080080	A35HPV09080040	9	80	9	18	7
A35HPV10040135	A35HPV10040080	A35HPV10040040	10	40	8	16	7
A35HPV10080135	A35HPV10080080	A35HPV10080040	10	80	8	16	7
A35HPV12040135	A35HPV12040080	A35HPV12040040	12	40	8	14	7
A35HPV12080135	A35HPV12080080	A35HPV12080040	12	80	7	12	7

# Pacific™ Plus

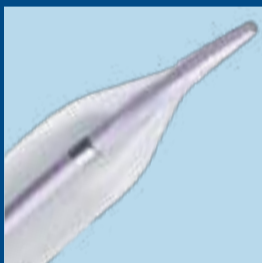
## PTA Balloon Catheter OTW 0.018"

### Versatility for everyday and beyond

- Versatile shaft lengths and guidewire compatibility for more access options
- Proprietary balloon technology allows for great crossability\*
- Improved shaft design for faster deflation\*

### TECHNICAL SPECIFICATIONS

Catheter design	Over-the-wire
Balloon coating	Hydrophilic
Balloon marker	2 swaged, platinum iridium
Shaft diameter	4.0 F
Usable shaft lengths	90, 130 and 180 cm
Introducer sheath compatibility	4.0 F
Guidewire compatibility	0.018"



Tapered tip



\* Six samples of each brand tested, 5 mm x 40 mm. Medtronic data on file. Bench test results may not be indicative of clinical performance.

# Pacific™ Plus

## PTA Balloon Catheter OTW 0.018"

### ORDER INFORMATION

Product Catalogue Number			Balloon Diameter (mm)	Balloon Length (mm)	Recommended Introducer Sheath (F)	Nominal Pressure (atm)	RBP (atm)
Usable Length 90 cm	Usable Length 130 cm	Usable Length 180 cm					
PCE 020 020 090	PCE 020 020 130		2.00	20	4	8	22
PCE 020 040 090	PCE 020 040 130	PCE 020 040 180	2.00	40	4	8	22
PCE 020 060 090	PCE 020 060 130		2.00	60	4	8	22
PCE 020 080 090	PCE 020 080 130	PCE 020 080 180	2.00	80	4	8	22
PCE 020 120 090	PCE 020 120 130	PCE 020 120 180	2.00	120	4	8	22
PCE 020 150 090	PCE 020 150 130	PCE 020 150 180	2.00	150	4	8	22
PCE 025 020 090	PCE 025 020 130		2.50	20	4	8	16
PCE 025 040 090	PCE 025 040 130	PCE 025 040 180	2.50	40	4	8	16
PCE 025 060 090	PCE 025 060 130		2.50	60	4	8	16
PCE 025 080 090	PCE 025 080 130	PCE 025 080 180	2.50	80	4	8	16
PCE 025 120 090	PCE 025 120 130	PCE 025 120 180	2.50	120	4	8	16
PCE 025 150 090	PCE 025 150 130	PCE 025 150 180	2.50	150	4	8	16
PCE 030 020 090	PCE 030 020 130		3.00	20	4	8	16
PCE 030 040 090	PCE 030 040 130	PCE 030 040 180	3.00	40	4	8	16
PCE 030 060 090	PCE 030 060 130		3.00	60	4	8	16
PCE 030 080 090	PCE 030 080 130	PCE 030 080 180	3.00	80	4	8	16
PCE 030 120 090	PCE 030 120 130	PCE 030 120 180	3.00	120	4	8	16
PCE 030 150 090	PCE 030 150 130	PCE 030 150 180	3.00	150	4	8	16
PCE 035 020 090	PCE 035 020 130		3.50	20	4	8	16
PCE 035 040 090	PCE 035 040 130		3.50	40	4	8	16
PCE 035 060 090	PCE 035 060 130		3.50	60	4	8	16
PCE 035 080 090	PCE 035 080 130		3.50	80	4	8	16
PCE 035 120 090	PCE 035 120 130		3.50	120	4	8	16
PCE 035 150 090	PCE 035 150 130		3.50	150	4	8	16
PCE 040 020 090	PCE 040 020 130		4.00	20	4	8	14
PCE 040 040 090	PCE 040 040 130	PCE 040 040 180	4.00	40	4	8	14
PCE 040 060 090	PCE 040 060 130		4.00	60	4	8	14
PCE 040 080 090	PCE 040 080 130	PCE 040 080 180	4.00	80	4	8	14
PCE 040 120 090	PCE 040 120 130	PCE 040 120 180	4.00	120	4	8	14
PCE 050 020 090	PCE 050 020 130		5.00	20	4	8	14
PCE 050 040 090	PCE 050 040 130	PCE 050 040 180	5.00	40	4	8	14
PCE 050 060 090	PCE 050 060 130	PCE 050 060 180	5.00	60	4	8	14
PCE 050 080 090	PCE 050 080 130	PCE 050 080 180	5.00	80	4	8	14
PCE 050 120 090	PCE 050 120 130	PCE 050 120 180	5.00	120	4	8	14
PCE 060 020 090	PCE 060 020 130		6.00	20	4	8	14
PCE 060 040 090	PCE 060 040 130	PCE 060 040 180	6.00	40	4	8	14
PCE 060 060 090	PCE 060 060 130		6.00	60	4	8	14
PCE 060 080 090	PCE 060 080 130	PCE 060 080 180	6.00	80	4	8	14
PCE 060 120 090	PCE 060 120 130	PCE 060 120 180	6.00	120	4	8	14
PCE 070 020 090	PCE 070 020 130		7.00	20	4	8	12
PCE 070 040 090	PCE 070 040 130	PCE 070 040 180	7.00	40	4	8	12
PCE 070 060 090	PCE 070 060 130		7.00	60	4	8	12
PCE 070 080 090	PCE 070 080 130	PCE 070 080 180	7.00	80	5	8	12
PCE 070 120 090	PCE 070 120 130	PCE 070 120 180	7.00	120	5	8	12

# Pacific™ Xtreme™

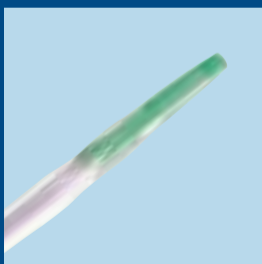
## PTA Balloon Catheter OTW 0.018"

### Versatility for everyday and beyond

- Balloons from 150-300 mm for treating long femoropopliteal lesions
- Strong pushability and kink resistance combined with excellent flexibility\*
- Low profile introducer sheath compatibility for less possible puncture trauma\*\*

### TECHNICAL SPECIFICATIONS

Catheter design	Over-the-wire, Coaxial Shaft
Balloon coating	Hydrophilic
Balloon marker	2 swaged, platinum iridium
Shaft diameter	3.9 – 4.2 F
Usable shaft lengths	90, 130 and 180 cm
Introducer sheath compatibility	4,5 F
Guidewire compatibility	0.018"



Tapered tip



Six-fold balloon

\* Bench test data on file at Medtronic, Inc. Test data not indicative of clinical performance.  
\*\* 4F/5F depending on diameter and length.

# Pacific™ Xtreme

## PTA Balloon Catheter OTW 0.018"

### ORDER INFORMATION

Product Catalogue Number			Balloon Diameter (mm)	Balloon Length (mm)	Recommended Introducer Sheath (F)	Nominal Pressure (atm)	RBP (atm)
Usable Length 90 cm	Usable Length 130 cm	Usable Length 180 cm					
PCF 040 150 090	PCF 040 150 130	PCF 040 150 180	4.00	150	4	6	14
PCF 040 200 090	PCF 040 200 130		4.00	200	4	6	14
PCF 040 250 090	PCF 040 250 130		4.00	250	4	6	14
PCF 040 300 090	PCF 040 300 130		4.00	300	4	6	14
PCF 050 150 090	PCF 050 150 130	PCF 050 150 180	5.00	150	4	6	14
PCF 050 200 090	PCF 050 200 130		5.00	200	4	6	14
PCF 050 250 090	PCF 050 250 130		5.00	250	5	6	14
PCF 050 300 090	PCF 050 300 130		5.00	300	5	6	14
PCF 060 150 090	PCF 060 150 130		6.00	150	5	6	12
PCF 060 200 090	PCF 060 200 130		6.00	200	5	6	12
PCF 060 250 090	PCF 060 250 130		6.00	250	5	6	12
PCF 060 300 090	PCF 060 300 130		6.00	300	5	6	12
PCF 070 150 090	PCF 070 150 130		7.00	150	5	6	12
PCF 070 200 090	PCF 070 200 130		7.00	200	5	6	12
PCF 070 250 090	PCF 070 250 130		7.00	250	5	6	12

# Submarine™ Rapido

## PTA Balloon Catheter RX 0.018"

AORTIC

PERIPHERAL

VENOUS

Low profile with strength and control

### Low profile

Compatible with 6F guiding catheter\*\*\*

- Swaged "zero profile" markers enable easy penetration of the target lesion
- Special 3-folded balloon minimizes the re-wrap profile
- 0.021" tip entry profile to enhance crossability of subocclusive lesions

### Strength and control

For carotid\*\* and renal interventions, a delicate and high performance PTA is required

- Flexitec LP material allows for a wide working range from nominal pressure of 7 bar up to 17 bar RBP
- Controlled balloon compliance for exact sizing and reliable balloon performance

### Shaft design

- Push transmission by homogeneous RX transition and adequate flexibility of the distal part
- Hydrophilic coating for swift and easy navigation

### Size mix

- Broad size mix (including diameters up to 7 mm) makes this catheter appropriate for carotid\* and renal interventions

### TECHNICAL SPECIFICATIONS

Catheter design	Rapid exchange (RX)
Balloon coating	Hydrophilic
Balloon marker	2 swaged, Platinum Iridium
Shaft diameter	Proximal 2.3 F Distal 3.0 – 3.5 F
Usable shaft lengths	135 cm
Introducer sheath compatibility	6,7 F
Guidewire compatibility	0.018"
Nominal pressure	7 atm

\* Bench test data on file at Medtronic. Test data not indicative of actual performance.

\*\* All codes with balloon lengths up to 40mm are certified for Carotid application.

\*\*\* Up to 6mm balloon diameter.



# Submarine™ Rapido

## PTA Balloon Catheter RX 0.018"

### ORDER INFORMATION

Product Catalogue Number Shaft Length 135 cm	Balloon Diameter (mm)	Balloon Length (mm)	RBP (atm)	Usable Shaft Length (cm)	Guiding Catheter Compatibility (F)	Distal Shaft Diameter (F)
SBR 020 020 135	2.00	20	17	135	6	3.0
SBR 020 040 135	2.00	40	17	135	6	3.0
SBR 020 060 135	2.00	60	15	135	6	3.0
SBR 025 020 135	2.50	20	17	135	6	3.0
SBR 025 040 135	2.50	40	17	135	6	3.0
SBR 025 060 135	2.50	60	15	135	6	3.0
SBR 030 020 135	3.00	20	17	135	6	3.0
SBR 030 040 135	3.00	40	17	135	6	3.0
SBR 030 060 135	3.00	60	15	135	6	3.0
SBR 030 080 135	3.00	80	15	135	6	3.0
SBR 035 020 135	3.50	20	17	135	6	3.5
SBR 035 030 135	3.50	30	17	135	6	3.5
SBR 040 020 135	4.00	20	17	135	6	3.5
SBR 040 030 135	4.00	30	17	135	6	3.5
SBR 040 040 135	4.00	40	17	135	6	3.5
SBR 045 020 135	4.50	20	17	135	6	3.5
SBR 045 040 135	4.50	40	17	135	6	3.5
SBR 050 020 135	5.00	20	16	135	6	3.5
SBR 050 030 135	5.00	30	16	135	6	3.5
SBR 050 040 135	5.00	40	16	135	6	3.5
SBR 055 020 135	5.50	20	16	135	6	3.5
SBR 055 030 135	5.50	30	16	135	6	3.5
SBR 055 040 135	5.50	40	16	135	6	3.5
SBR 060 020 135	6.00	20	16	135	6	3.5
SBR 060 030 135	6.00	30	16	135	6	3.5
SBR 060 040 135	6.00	40	16	135	6	3.5
SBR 065 020 135	6.50	20	16	135	7	3.5
SBR 065 030 135	6.50	30	16	135	7	3.5
SBR 065 040 135	6.50	40	16	135	7	3.5
SBR 070 020 135	7.00	20	16	135	7	3.5
SBR 070 030 135	7.00	30	16	135	7	3.5
SBR 070 040 135	7.00	40	16	135	7	3.5

The Submarine Rapido PTA catheter is indicated for PTA in patients with obstructive disease in peripheral arteries (i.e. carotid, supraaortic, ilio-femoral, popliteal, infrapopliteal, and renal arteries).

# Amphirion™ Deep

## Infrapopliteal PTA Balloon Catheter OTW 0.014"

### Easy access to the extremities\*

- Tip profile (0.017")
- LFC hydrophilic coating
- Three-step shaft design to optimize tracking through distal vessels
- Reinforced proximal shaft design for strong pushability
- 4 F compatible in all sizes

### Balloon sizes to accommodate your needs\*

- Size range 1.5–4.0 mm in diameter and 20–210 mm in length
- Long balloons (up to 210 mm) are suited for treatment of extremely diffuse lesions

### Conformable balloon material\*

- Proprietary polymer blend provides wonderful conformability

### Tapered balloon\*

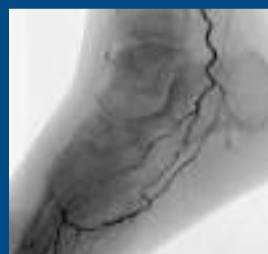
- The 210 mm balloon tapers by 0.5 mm to respect the arterial anatomy

### TECHNICAL SPECIFICATIONS

Catheter design	Over-the-wire
Balloon marker	1/2 swaged, Platinum Iridium
Shaft diameter	Proximal 3.9 F Middle 3.3 F Distal 2.8 F
Introducer sheath compatibility	4 F
Guidewire compatibility	0.014"
Nominal pressure	7 atm



Image courtesy of Dr. Marco Manzi, Italy.



Below the ankle



\* Bench Test Data on file with Medtronic, Inc.  
Test data not indicative of clinical performance

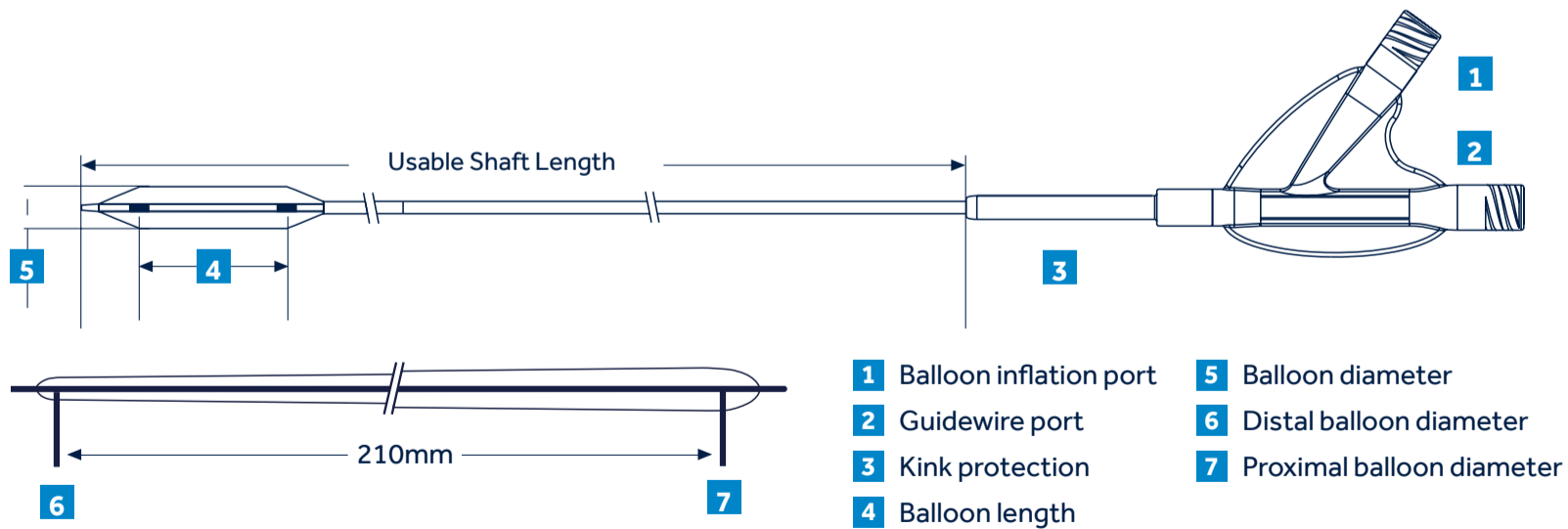


# Amphirion™ Deep

## Infrapopliteal PTA Balloon Catheter OTW 0.014"

### ORDER INFORMATION

Product Catalogue Number		Balloon Diameter (mm)	Balloon Length (mm)	Recommended Introducer Sheath (F)	Nominal Pressure (BAR)	RBP (BAR)
OTW Usable Length 120 cm	OTW Usable Length 150 cm					
AMD 015 020 001	AMD 015 020 151	1.5	20	4	7	14
AMD 015 020 002	AMD 015 020 152	1.5	20	4	7	14
AMD 020 040 002	AMD 020 040 152	2.0	40	4	7	15
AMD 020 080 002	AMD 020 080 152	2.0	80	4	7	14
AMD 020 120 002	AMD 020 120 152	2.0	120	4	7	14
AMD 020 150 002	AMD 020 150 152	2.0	150	4	7	14
AMD 025 040 002	AMD 025 040 152	2.5	40	4	7	16
AMD 025 080 002	AMD 025 080 152	2.5	80	4	7	15
AMD 025 120 002	AMD 025 120 152	2.5	120	4	7	14
AMD 025 150 002	AMD 025 150 152	2.5	150	4	7	14
AMD 030 040 002	AMD 030 040 152	3.0	40	4	7	16
AMD 030 080 002	AMD 030 080 152	3.0	80	4	7	15
AMD 030 120 002	AMD 030 120 152	3.0	120	4	7	14
AMD 030 150 002	AMD 030 150 152	3.0	150	4	7	14
AMD 035 040 002	AMD 035 040 152	3.5	40	4	7	16
AMD 035 080 002	AMD 035 080 152	3.5	80	4	7	15
AMD 035 120 002	AMD 035 120 152	3.5	120	4	7	14
AMD 035 150 002	AMD 035 150 152	3.5	150	4	7	14
AMD 040 040 002	AMD 040 040 152	4.0	40	4	7	16
AMD 040 080 002	AMD 040 080 152	4.0	80	4	7	15
AMD 040 120 002	AMD 040 120 152	4.0	120	4	7	14
AMD 040 150 002	AMD 040 150 152	4.0	150	4	7	14
AMD 225 210 002	AMD 225 210 152	2.0/2.5	210	4	7	14
AMD 253 210 002	AMD 253 210 152	2.5/3.0	210	4	7	14
AMD 335 210 002	AMD 335 210 152	3.0/3.5	210	4	7	14
AMD 354 210 002	AMD 354 210 152	3.5/4.0	210	4	7	14



# NanoCross™ Elite

## OTW PTA Dilatation Catheter 0.014"

NanoCross™ Elite, the next generation 0.014" PTA balloon, with its 360° beveled tip provides smooth transition from wire to tip. The SlimTec™ balloon-folding process is designed to provide the lowest 0.014" crossing profile.

### TECHNICAL SPECIFICATIONS

Catheter design	Over-the-wire
Useable catheter lengths	90 and 150 cm
Introducer sheath compatibility	4 F
Guidewire compatibility	0.014"



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# NanoCross™ Elite

## OTW PTA Dilatation Catheter 0.014"

### ORDER INFORMATION

Product Catalogue Number		Balloon Diameter (mm)	Balloon Length (mm)	Nominal Pressure (atm)	Pressure (atm)	Introducer Sheath (F)
90 cm Catheter Length	150 cm Catheter Length					
AB14W015020090	AB14W015020150	1.5	20	8	14	4
AB14W015040090	AB14W015040150	1.5	40	8	14	4
AB14W020020090	AB14W020020150	2	20	8	14	4
AB14W020040090	AB14W020040150	2	40	8	14	4
AB14W020060090	AB14W020060150	2	60	8	14	4
AB14W020080090	AB14W020080150	2	80	8	14	4
AB14W020100090	AB14W020100150	2	100	8	14	4
AB14W020120090	AB14W020120150	2	120	8	14	4
AB14W020150090	AB14W020150150	2	150	8	14	4
AB14W020210090	AB14W020210150	2	210	8	14	4
AB14W025020090	AB14W025020150	2.5	20	8	14	4
AB14W025040090	AB14W025040150	2.5	40	8	14	4
AB14W025060090	AB14W025060150	2.5	60	8	14	4
AB14W025080090	AB14W025080150	2.5	80	8	14	4
AB14W025100090	AB14W025100150	2.5	100	8	14	4
AB14W025120090	AB14W025120150	2.5	120	8	14	4
AB14W025150090	AB14W025150150	2.5	150	8	14	4
AB14W025210090	AB14W025210150	2.5	210	8	14	4
AB14W030020090	AB14W030020150	3	20	8	14	4
AB14W030040090	AB14W030040150	3	40	8	14	4
AB14W030060090	AB14W030060150	3	60	8	14	4
AB14W030080090	AB14W030080150	3	80	8	14	4
AB14W030100090	AB14W030100150	3	100	8	14	4
AB14W030120090	AB14W030120150	3	120	8	14	4
AB14W030150090	AB14W030150150	3	150	8	14	4
AB14W030210090	AB14W030210150	3	210	8	14	4
AB14W035020090	AB14W035020150	3.5	20	8	14	4
AB14W035040090	AB14W035040150	3.5	40	8	14	4
AB14W035060090	AB14W035060150	3.5	60	8	14	4
AB14W035080090	AB14W035080150	3.5	80	8	14	4
AB14W035100090	AB14W035100150	3.5	100	8	14	4
AB14W035120090	AB14W035120150	3.5	120	8	14	4
AB14W035150090	AB14W035150150	3.5	150	8	14	4
AB14W035210090	AB14W035210150	3.5	210	8	14	4
AB14W040020090	AB14W040020150	4	20	8	14	4
AB14W040040090	AB14W040040150	4	40	8	14	4
AB14W040060090	AB14W040060150	4	60	8	14	4
AB14W040080090	AB14W040080150	4	80	8	14	4
AB14W040100090	AB14W040100150	4	120	8	14	4
AB14W040120090	AB14W040120150	4	100	8	14	4
AB14W040150090	AB14W040150150	4	150	8	14	4
AB14W040210090	AB14W040210150	4	210	8	14	4
AB14W050020090	AB14W050020150	5	20	8	14	5
AB14W050040090	AB14W050040150	5	40	8	14	5
AB14W050060090	AB14W050060150	5	60	8	14	5
AB14W050080090	AB14W050080150	5	80	8	14	5
AB14W050100090	AB14W050100150	5	100	8	14	5
AB14W050120090	AB14W050120150	5	120	8	14	5
AB14W050150090	AB14W050150150	5	150	8	14	5
AB14W050200090	AB14W050200150	5	200	8	14	5
AB14W060020090	AB14W060020150	6	20	8	14	5
AB14W060040090	AB14W060040150	6	40	8	14	5
AB14W060060090	AB14W060060150	6	60	8	14	5
AB14W060080090	AB14W060080150	6	80	8	14	5
AB14W060100090	AB14W060100150	6	100	8	14	5
AB14W060120090	AB14W060120150	6	120	8	14	5
AB14W060150090	AB14W060150150	6	150	8	14	5
AB14W060200090	AB14W060200150	6	200	8	14	6

INDICATIONS: The NanoCross™ Elite 0.014" Over-the-Wire PTA Dilatation Catheter is intended to dilate stenoses in the iliac, femoral, ilio-femoral, popliteal, infra-popliteal, and renal arteries and for the treatment of obstructive lesions of native or synthetic arteriovenous dialysis fistulae. Each system includes: One PTA balloon catheter, one compliance chart and one balloon folding tool.

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# RapidCross™

## PTA Balloon Dilatation Catheter RX 0.014"

RapidCross™ 0.014 PTA Balloon Catheter was developed exclusively for below the knee treatment. Every detail, from the 0.017" tip entry profile to the Rapid Exchange Port construction was thoughtfully designed for exceptional performance.

### Low profile

- For ease in crossing

### Dual middle marker bands\*

- For enhanced visualization and accuracy

### Tapered long balloon\*\*

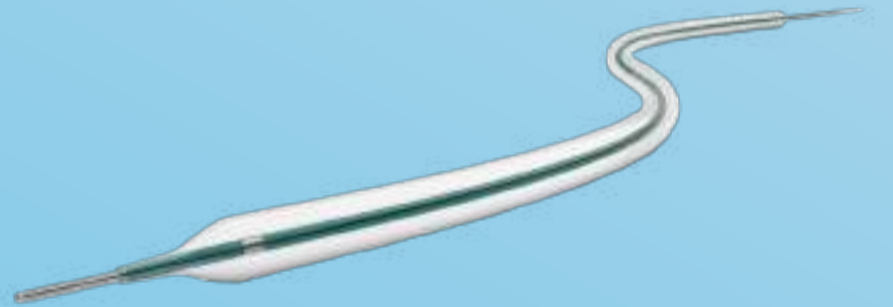
- Tailored to distal tibial and pedal anatomy

### Maximum inner lumen

- For 2-3 times faster deflation

### Proprietary balloon coating

- Ensures sustained lubricity for crossing challenging lesions



\* Dual middle marker bands available on 150 mm and 210 mm balloon lengths

\*\* Tapered balloon available in 210 mm balloon length

### ORDER INFORMATION

Product Catalogue Number		Balloon Diameter (mm)	Balloon Length (mm)	Nominal Pressure (atm)	Rated Burst Pressure (atm)	Recommended Introducer Sheath (F)
Catheter Shaft Length 90 cm	Catheter Shaft Length 170 cm					
A14BX020020090	A14BX020020170	2.0	20	8	14	4
A14BX020040090	A14BX020040170	2.0	40	8	14	4
A14BX020060090	A14BX020060170	2.0	60	8	14	4
A14BX020080090	A14BX020080170	2.0	80	8	14	4
A14BX020100090	A14BX020100170	2.0	100	8	14	4
A14BX020120090	A14BX020120170	2.0	120	8	14	4
A14BX020150090	A14BX020150170	2.0	150	8	14	4
A14BX020210090	A14BX020210170	2.0 proximal / 1.5 distal	210	8	14	4
A14BX025020090	A14BX025020170	2.5	20	8	14	4
A14BX025040090	A14BX025040170	2.5	40	8	14	4
A14BX025060090	A14BX025060170	2.5	60	8	14	4
A14BX025080090	A14BX025080170	2.5	80	8	14	4
A14BX025100090	A14BX025100170	2.5	100	8	14	4
A14BX025120090	A14BX025120170	2.5	120	8	14	4
A14BX025150090	A14BX025150170	2.5	150	8	14	4
A14BX025210090	A14BX025210170	2.5 proximal / 2.0 distal	210	8	14	4
A14BX030020090	A14BX030020170	3	20	8	14	4
A14BX030040090	A14BX030040170	3	40	8	14	4
A14BX030060090	A14BX030060170	3	60	8	14	4
A14BX030080090	A14BX030080170	3	80	8	14	4
A14BX030100090	A14BX030100170	3	100	8	14	4
A14BX030120090	A14BX030120170	3	120	8	14	4
A14BX030150090	A14BX030150170	3	150	8	14	4
A14BX030210090	A14BX030210170	3.0 proximal / 2.5 distal	210	8	14	4
A14BX035020090	A14BX035020170	3.5	20	8	14	4
A14BX035040090	A14BX035040170	3.5	40	8	14	4
A14BX035060090	A14BX035060170	3.5	60	8	14	4
A14BX035080090	A14BX035080170	3.5	80	8	14	4
A14BX035100090	A14BX035100170	3.5	100	8	14	4
A14BX035120090	A14BX035120170	3.5	120	8	14	4
A14BX035150090	A14BX035150170	3.5	150	8	14	4
A14BX035210090	A14BX035210170	3.5 proximal / 3.0 distal	210	8	14	4
A14BX040020090	A14BX040020170	4	20	8	14	4
A14BX040040090	A14BX040040170	4	40	8	14	4
A14BX040060090	A14BX040060170	4	60	8	14	4
A14BX040080090	A14BX040080170	4	80	8	14	4
A14BX040100090	A14BX040100170	4	100	8	14	4
A14BX040120090	A14BX040120170	4	120	8	14	4
A14BX040150090	A14BX040150170	4	150	8	14	4
A14BX040210090	A14BX040210170	4.0 proximal / 3.5 distal	210	8	14	4

INDICATIONS: The RapidCross™ rapid exchange PTA balloon dilatation catheter is intended to dilate stenoses in the iliac, femoral, iliofemoral, popliteal, infrapopliteal, and renal arteries, and for the treatment of obstructive lesions of native or synthetic arteriovenous dialysis fistulae.

# Chocolate™

## PTA Balloon

Minimize vessel trauma, dissections and the need for bailout stenting above or below the knee with the Chocolate™ PTA balloon. The balloon's unique nitinol constraining structure creates pillows and grooves that provide a predictable, uniform and atraumatic dilatation.

- Minimize vessel trauma
- Reduce dissections
- Decrease bailout stenting

### TECHNICAL SPECIFICATIONS

Catheter design	Over-the-wire
Useable catheter lengths	120, 135 and 150 cm
Introducer sheath compatibility	5, 6 F
Guidewire compatibility	0,014" and 0,018"



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## PTA Balloon

### Braided Catheter Shaft

- Designed to provide robust pushability to reach and cross lesions

### Grooves

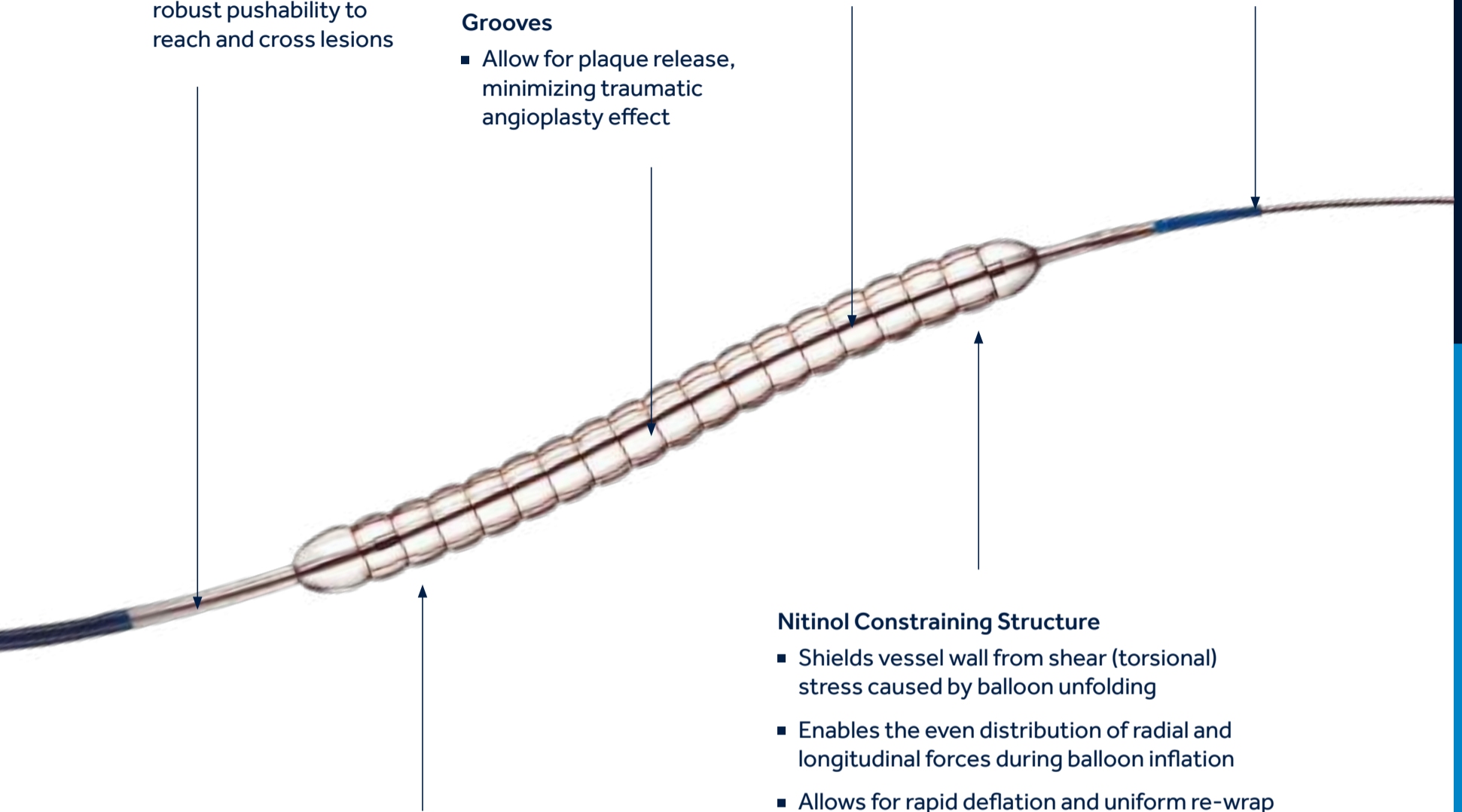
- Allow for plaque release, minimizing traumatic angioplasty effect

### Pillows

- Provide predictable vessel dilation without cutting or scoring

### Tapered Tip

- Enables lower entry profile for optimal lesion access

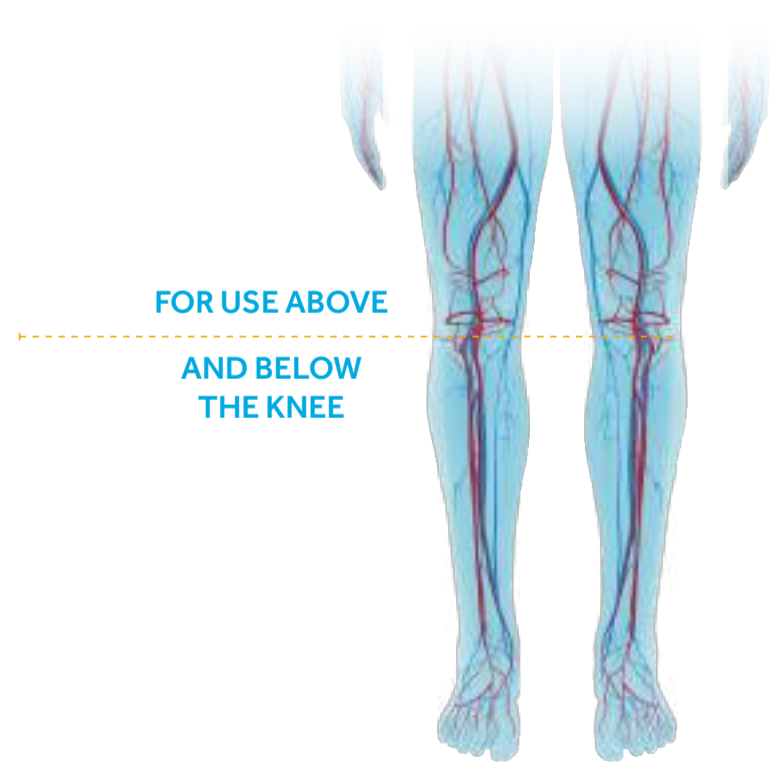
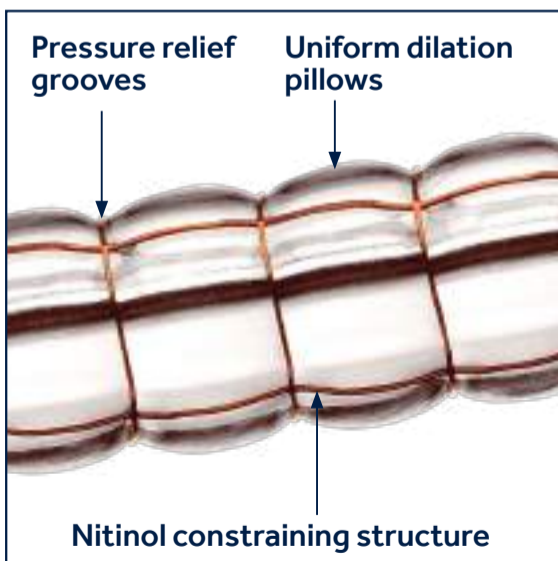


### Nitinol Constraining Structure

- Shields vessel wall from shear (torsional) stress caused by balloon unfolding
- Enables the even distribution of radial and longitudinal forces during balloon inflation
- Allows for rapid deflation and uniform re-wrap

### Nylon, Semi-Compliant Balloon

- Allows for optimal balloon pillow formation



### ORDER INFORMATION

Product Catalogue Number	Balloon Diameter (mm)	Balloon Length (mm)	Catheter Length (cm)	Guidewire (in)	Introducer Sheath (F)	Nominal Pressure (atm)	Rated Burst Pressure (atm)
CE1415025040 OTW	2.5	40	150	0.014"	5	9	14
CE1415025080 OTW		80					
CE1415025120 OTW		120					
CE1415030040 OTW	3.0	40					
CE1415030080 OTW		80					
CE1415030120 OTW		120					
CE1413535040 OTW	3.5	40	135	0.014"	5	9	14
CE1413535080 OTW		80					
CE1413535120 OTW		120					
CE1413540040 OTW	4.0	40					
CE1413540080 OTW		80					
CE1413540120 OTW		120					
CE1812050040 OTW	5.0	40	120	0.018"	6	6	12
CE1812050080 OTW		80					
CE1812050120 OTW		120					
CE1812060040 OTW	6.0	40					
CE1812060080 OTW		80					
CE1812060120 OTW		120					

**Important Information:** Indications, contraindications, warnings and instructions for use can be found in the product labeling supplied with each device.

**Indications for Use:** The Chocolate™ PTA Balloon Catheter is intended for balloon dilatation of lesions in the peripheral vasculature, including the iliac, femoral, ilio-femoral, popliteal, infra-popliteal, and renal arteries.

**CAUTION:** Federal (USA) law restricts these products for sale by or on the order of a physician.

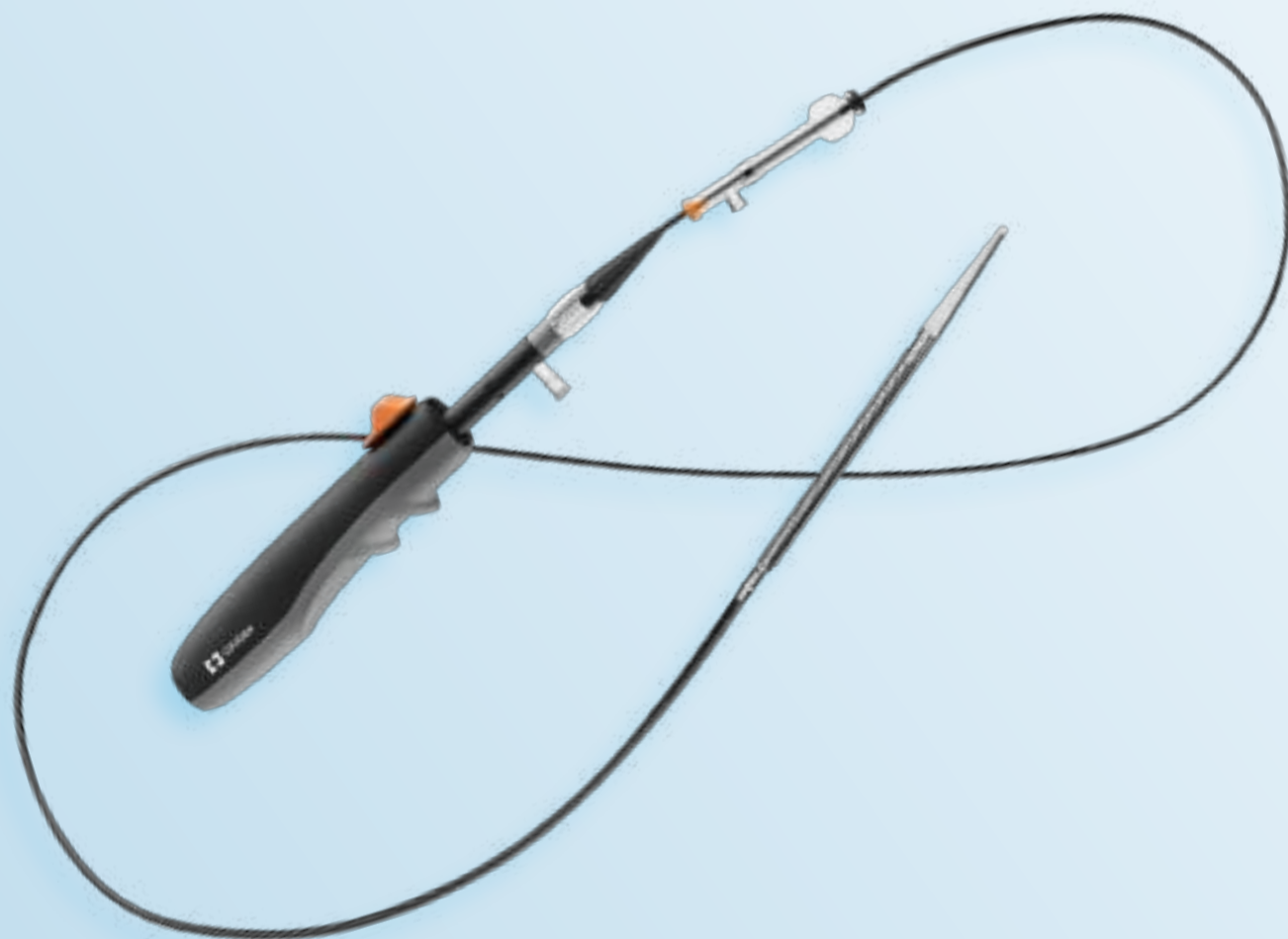


# DIRECTIONAL ATHERECTOMY

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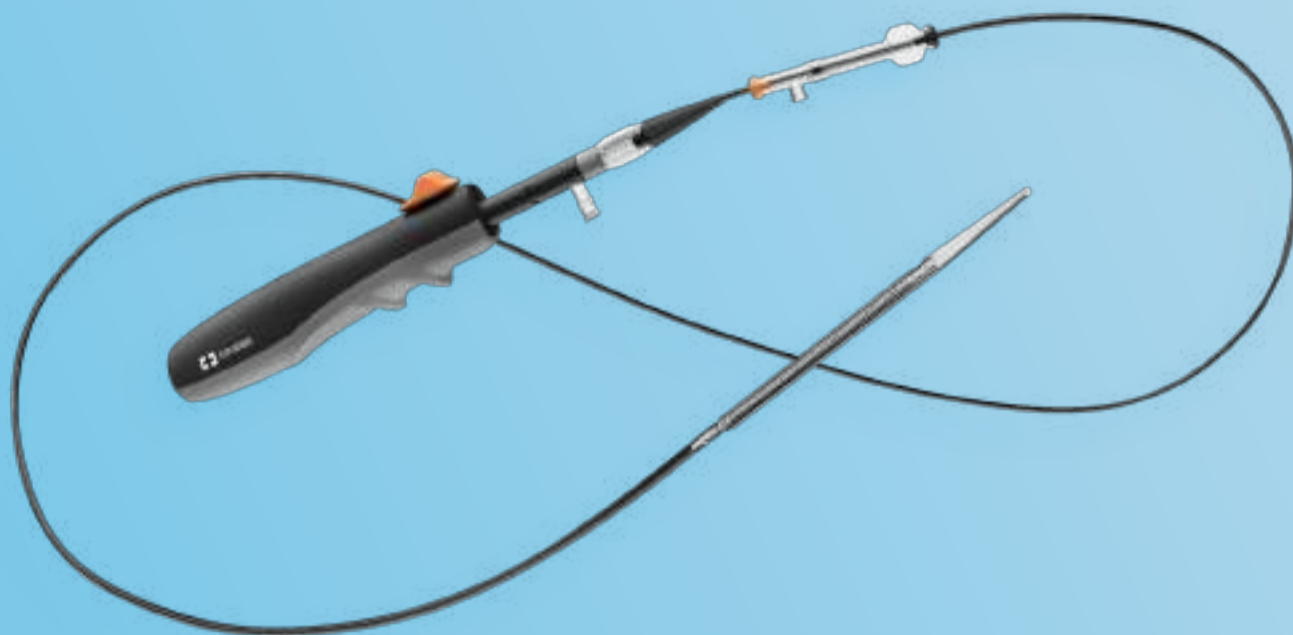
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# HawkOne™

## Directional Atherectomy System



One device for above and below the knee available in 6 F and 7 F sizes

Just as the name implies, the all-new HawkOne™ Directional Atherectomy System is one device that treats all morphologies<sup>1</sup>, including severe calcium, and offers procedural efficiency with enhanced cutting, crossing, and cleaning capabilities.\*

One device that:

- Treats all morphologies
- Offers procedural efficiency
- Restores blood flow in PAD patients

\* Comparison and claims in reference to the TurboHawk™ High Efficiency Cutter.

<sup>1</sup> HawkOne™ Bench Performance Verification Testing (RE-PV13728); Calcified Cutting Efficiency Bench Validation Data (RE-PV13729)



### CUTTING BLADE

Four contoured cutting blades engage and treat all atherosclerotic morphologies.



### DRIVE SHAFT

A four layered, counter-wound design efficiently transmits power, offering a **25%** improvement\* in torsional performance.



### JOG

Optimized\* for improved engagement in calcified lesions with no increase in cut depth.



### CUTTER DRIVER

Ergonomically redesigned to effectively treat all atherosclerotic plaque.



### DISTAL TIP

A tapered, radiopaque distal tip provides enhanced\* deliverability and visualization under angiography.

## Directional Atherectomy System

Unlike orbital, laser, or rotational atherectomy, the HawkOne™ Directional Atherectomy System, with its directional cutting design, offers the greatest versatility when treating PAD. Whether your atherectomy goal is to maximize luminal gain<sup>2</sup>, to create in-line flow, or to target eccentric circumferential disease, the HawkOne™ is your go-to choice.

### ORDER INFORMATION

#### HAWKONE™ DIRECTIONAL ATHERECTOMY SYSTEM

	Model Name	Product Catalogue Number	Vessel Diameter (mm)	Sheath Compatibility (F)	Crossing Profile (mm)	Working Length <sup>3</sup> (cm)	Effective Length <sup>4</sup> (cm)	Tip Length (cm)	Max. cut Length (mm)
6 F	HawkOne™ M (KIT)**	H1-M-6FKIT	3.0-7.0	6	2.2	135	129	5.9	40
	HawkOne™ M <sup>+</sup>	H1-M-INT	3.0-7.0	6	2.2	135	129	5.9	40
	HawkOne™ S (KIT)**	H1-S-6FKIT	2.0-4.0	6	2.2	151	145	5.9	40
	HawkOne™ S <sup>+</sup>	H1-S-INT	2.0-4.0	6	2.2	151	145	5.9	40
7 F	HawkOne™ LS Standard tip (KIT)**	H1-LS-7FKIT	3.5 - 7.0	7	2.6	114	107	6.6	50
	HawkOne™ LS Standard tip <sup>+</sup>	H1-LS-INT	3.5 - 7.0	7	2.6	114	107	6.6	50
	HawkOne™ LX Extended tip (KIT)**	H1-LX-7FKIT	3.5 - 7.0	7	2.6	114	104	9.6	75
	HawkOne™ LX Extended tip <sup>+</sup>	H1-LX-INT	3.5 - 7.0	7	2.6	114	104	9.6	75

### ATHERECTOMY SYSTEMS

		Directional	Orbital	Laser	Rotational
Plaque modification	Maximize lumen gain	X			
	Restore in-line flow	X	X	X	X
Lesion morphology	Treat severe calcium	X	X		X
	Treat soft-moderate plaque	X		X	X
Plaque distribution	Target eccentric disease	X			
	Target circumferential disease	X	X	X	X

Max guidewire is 0.014" for HawkOne™ device.

\*\* This catalogue number includes the HawkOne™ and the Cutter Driver.

\* Cutter driver H1-14550 needs to be ordered separately with this catalogue number.

<sup>2</sup> During Definitive LE clinical trial physicians were able to achieve technical success defined by debulking to < 30% stenosis.

Reference DEFINITIVE LE clinical trial: James F. McKinsey, MD, Thomas Zeller, MD, Krishna Rocha-Singh, MD, Michael R. Jaff, DO, and Lawrence A. Garcia, MD. Lower Extremity Revascularization Using Directional Atherectomy:

12 Month Prospective Results of the DEFINITIVE LE Study. JACC: Cardiovascular Interventions 7 (2014) pp. 923-933. 10.1016/j.jcin.2014.05.006.

<sup>3</sup> HawkOne™ Working Length – Distal end of pre-loaded flush tool, in the proximal position, to the distal end of tip.

<sup>4</sup> HawkOne™ Effective Length – Distal end of pre-loaded flush tool, in the proximal position, to the proximal end of cutter window.

# TurboHawk™

## Peripheral Plaque Excision System

Key features of the TurboHawk™ device

### Cutter selection

The TurboHawk™ device has two cutter options to choose from depending on the procedural need and lesion morphology.

- High-efficiency cutter – tackles soft-to-moderately calcified lesions
- Smooth cutter – treats soft-to-mild calcification

### Drive shaft

The counter-wound drive shaft transmits power more efficiently to the cutting blade.

### Micro Efficient Compression (MEC)™ technology

Tiny, laser-drilled holes in the nose cone allow excess fluid to escape so physicians are able to capture more plaque with each pass of the cutting blade, potentially reducing the number of insertions and procedure time.

- 45% increase in tissue collection capacity with MEC technology

### Dual catheter jog

The bend in the catheter enhances contact between the cutting blade and lesion, collecting more plaque with each pass.

### Distal flush tool

The distal flush tool effectively cleans and flushes plaque from the device with increased pressure.

### Tapered tip

The low-profile tip of the TurboHawk™ small-vessel catheter allows the device to maneuver through tortuous anatomies and challenging lesions with greater ease.

### Catheter alignment marker

This feature easily aligns the nose cone with the distal flush tool for faster cleaning.



## Peripheral Plaque Excision System

### ORDER INFORMATION

Model Name	Product Catalogue Number	Vessel Diameter (mm)	Sheath Compatibility (F)	Crossing Profile (inch)	Working Length <sup>1</sup> (cm)	Effective Length <sup>2</sup> (cm)	Tip Length (cm)	Max Cut Length (mm)
LS-C Super Cutter Large Vessel Standard Calcium Tip	THS-LS-C	3.5 - 7.0	7	0.105 (2.7 mm)	110	104	6.0	50
LS-M Smooth Cutter Large Vessel Standard Tip	THS-LS-M	3.5 - 7.0	7 / 8	0.105 (2.7 mm)	110	104	6.0	50
LX-C Super Cutter Large Vessel Xtended Calcium Tip	THS-LX-C	3.5 - 7.0	7	0.105 (2.7 mm)	113	104	9.0	75
LX-M Smooth Cutter Large Vessel Xtended Tip	THS-LX-M	3.5 - 7.0	7 / 8	0.105 (2.7 mm)	113	104	9.0	75
SX-C High Efficiency Cutter Small Vessel Xtended Calcium Tip	THS-SX-C	2.0 - 4.0	6	0.085 (2.2 mm)	135	129	5.9	40
SS-C High Efficiency Cutter Small Vessel Standard Calcium Tip	THS-SS-C	2.0 - 4.0	6	0.085 (2.2 mm)	133	129	3.9	20
SS-CL High Efficiency Cutter Small Vessel Standard Calcium Tip Long Catheter	THS-SS-CL	2.0 - 4.0	6	0.085 (2.2 mm)	149	145	3.9	20

Cutter driver FG02550 needs to be ordered with each TurboHawk™ device

INDICATIONS: The TurboHawk™ Peripheral Plaque Excision System is intended for use in atherectomy of the peripheral vasculature. The TurboHawk™ Catheter is not intended for use in the coronary, carotid, iliac or renal vasculature.

# SilverHawk™

## Peripheral Plaque Excision System

Our first-generation SilverHawk™ device treats PAD by removing soft-to-mild plaque buildup in leg arteries.

SilverHawk™ technology uses a directional cutting blade to shave plaque from the vessel—maximizing luminal gain. The plaque is captured in the nose cone of the device and safely removed from the vessel.

The SilverHawk™ device is backed by the landmark DEFINITIVE LE Clinical Study.



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## Peripheral Plaque Excision System

### ORDER INFORMATION

Model Name	Product Catalogue Number	Vessel Diameter (mm)	Sheath Compatibility (F)	Crossing Profile (inch)	Working Length <sup>1</sup> (cm)	Effective Length <sup>2</sup> (cm)	Tip Length (cm)	Max Cut Length (mm)
LS-M Large Vessel Standard Tip	P4052	4.5 - 7.0	7 / 8	0.105 (2.7 mm)	110	104	6.0	50
LX-M Large Vessel Xtended Tip	P4055	4.5 - 6.5	7 / 8	0.105 (2.7 mm)	113	104	6.0	75
MS-M Medium Vessel Standard Tip	P4056	3.5 - 5.0	7 / 8	0.105 (2.7 mm)	110	104	6.0	50
SXL Small Vessel Xtra Long Tip	P4033	3.0 - 3.5	7	0.095 (2.4 mm)	136	129	7.2	50
SS+ Small Vessel Standard Tip	P4030	3.0 - 3.5	7	0.090 (2.3 mm)	135	132	2.6	15
EXL Extra Small Vessel Xtra Long Tip	P4044	2.0 - 3.0	6	0.080 (2.0 mm)	135	129	6.0	15
ES+ Extra Small Vessel Standard Tip	P4034	2.0 - 2.5	6	0.075 (1.9 mm)	135	132	2.2	10
DS Distal Vessel Standard Tip	P4028	1.5 - 2.0	6	0.077 (1.9 mm)	135	132	2.6	10

Cutter driver FG02550 needs to be ordered with each SilverHawk™ device

INDICATIONS: The SilverHawk™ Peripheral Plaque Excision System is intended for use in atherectomy of the peripheral vasculature. The SilverHawk™ Catheter is not intended for use in the coronary, carotid, iliac or renal vasculature.

<sup>1</sup>Working Length - distal end of strain relief to the distal end of tip.

<sup>2</sup>Effective Length - distal end of strain relief to the distal end of the cutter window.

# EMBOLIC PROTECTION DEVICES

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# Mo.Ma™ Ultra

## Cerebral Protection Device

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Full-time protection and control

### Guide-catheter technology

- Provides excellent trackability, support and stability for ease of lesion crossing and accurate stent deployment

### Working channel exit port distal to CCA balloon

- Provides lesion access and effective, efficient aspiration of debris\*

### Radiopaque markers

- Centrally located in each balloon for precise positioning and orientation

### Optimal device selection

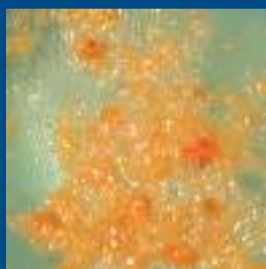
- Wires, stents and balloons

### High-capture efficiency

- Removal of all sizes of debris\*\*

### TECHNICAL SPECIFICATIONS

Balloon material	Compliant elastomeric rubber
Balloon marker distance	6 cm*
Distal shaft profile	5 F*
Recommended guidewire	0.035" (0.89 mm)
Balloon occlusion range	Up to 13 mm (CCA prox. balloon) Up to 6 mm* (ECA dist. balloon)



All sizes of debris are captured

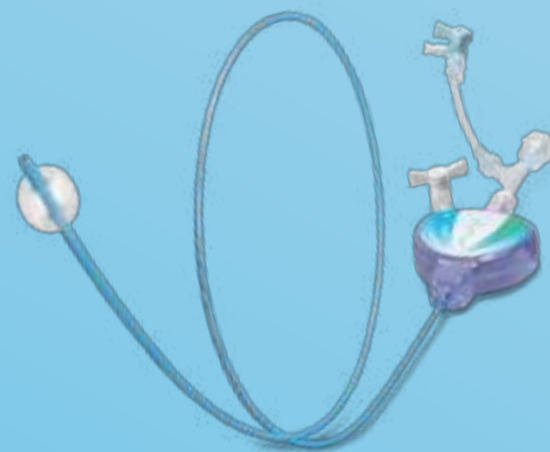
\* Double-Occlusion Balloon System only

\*\*Bench test data on file at Medtronic, Inc. Test data not indicative of clinical performance.



### DOUBLE-OCCLUSION BALLOON SYSTEM\*\*

Utilizes highly-compliant, elastomeric balloons that provide atraumatic flow suspension and stability



### MONO-OCCLUSION BALLOON SYSTEM

In case of occlusion of the external carotid artery (ECA), the system utilizes one highly-compliant, elastomeric balloon in the common carotid artery (CCA), that provides atraumatic flow suspension and stability.

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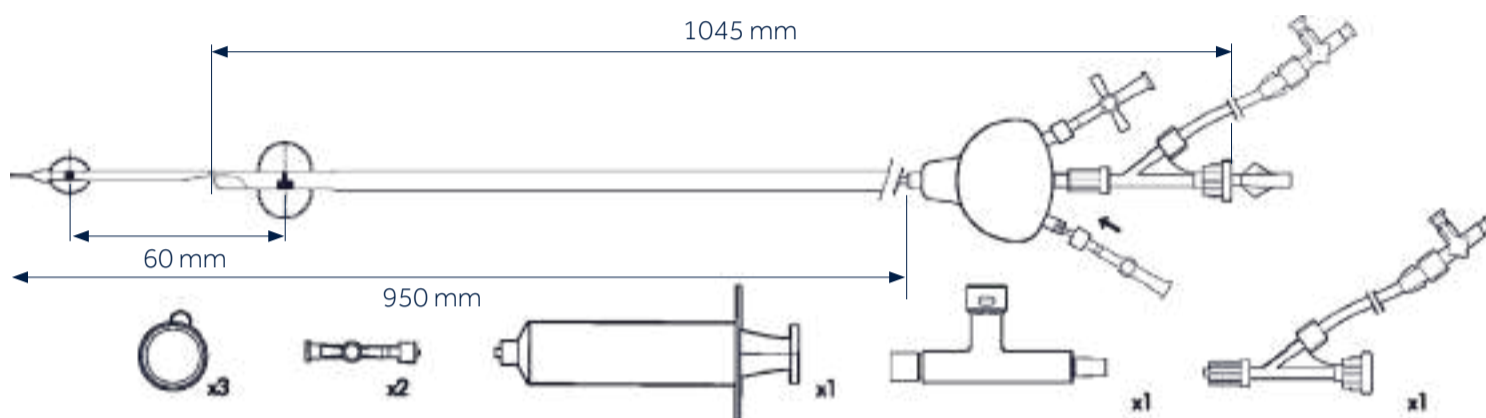
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# Mo.Ma™ Ultra

## Cerebral Protection Device

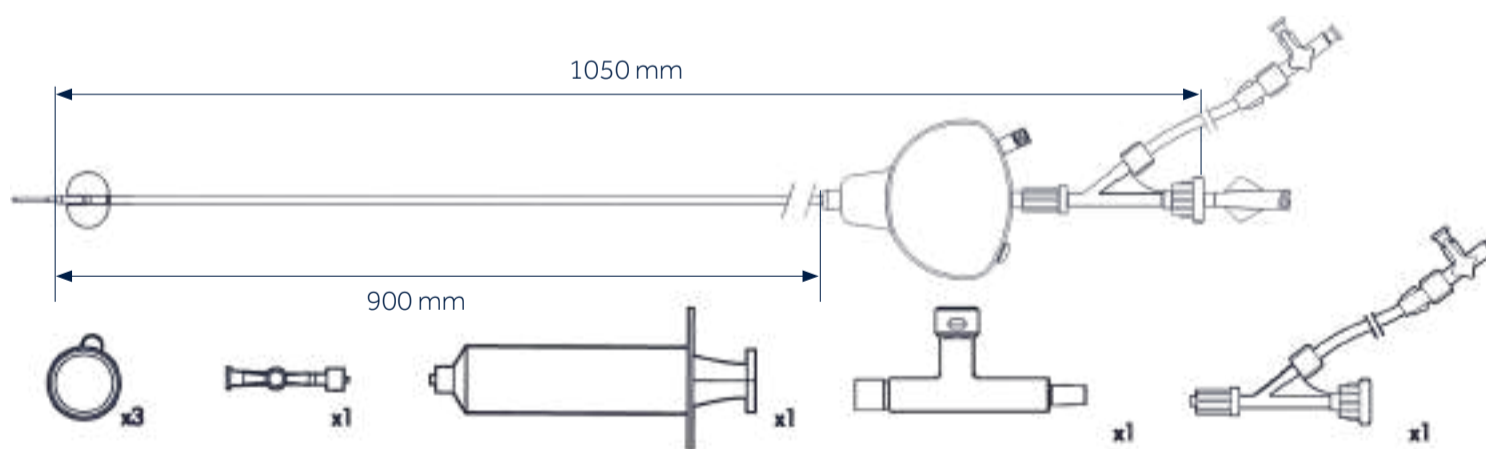
### ORDER INFORMATION

Product Catalogue Number Double-Balloon	Minimum Sheath Size	Inner Diameter Of The Working Channel
MOM0130068X5	8 F	0.069" / 1.76 mm
MOM0130069X6	9 F	0.083" / 2.12 mm



### ORDER INFORMATION

Product Catalogue Number Mono-Balloon	Minimum Sheath Size	Inner Diameter Of The Working Channel
MOM0130008X5	8 F	0.069" / 1.76 mm



# SpiderFX™

## Embolic Protection System

The SpiderFX™ System is the only embolic protection device that works with any 0.014" or 0.018" guidewire of choice to cross the most challenging lesions.

The device offers enhanced visibility due to the nitinol frame with gold / tungsten marker. The extensive product portfolio permits treatment within a range of vessel sizes from 2 mm to 7 mm. The SpiderFX™ System is compatible with a guide catheter / sheath minimum ID of 0.066" ( typically a 6 F guide catheter or 5 F access / long sheath ). Check catheter manufacturer information for size compatibility.

The use of the SpiderFX™ device is strongly associated with:

- Lower costs
- Shorter inpatient hospital stays
- Lower ICU utilization rate
- Shorter OR times

### Basket design

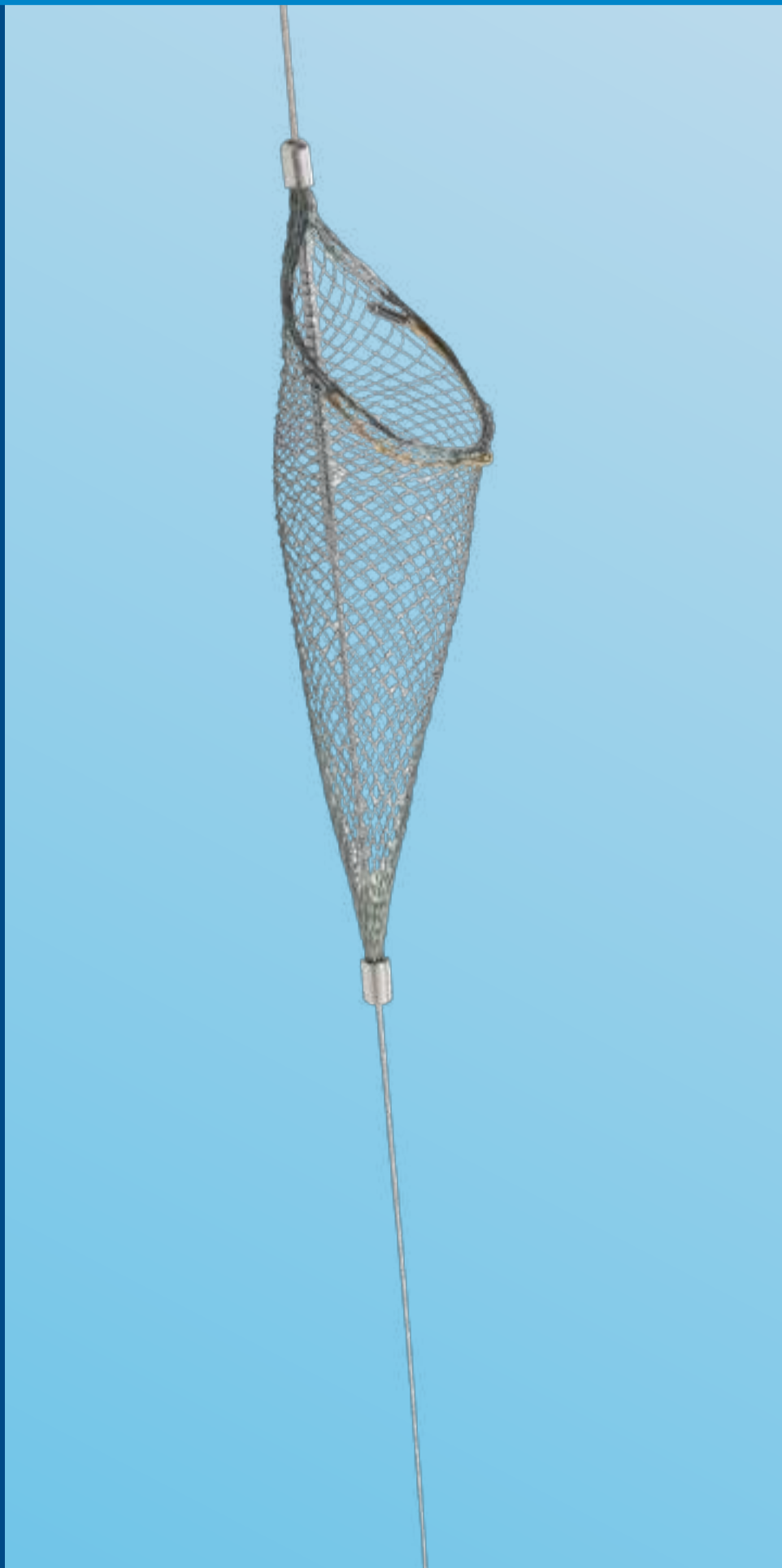
- The unique braided nitinol filter conforms to the vessel wall and maintains full-wall apposition during the intervention. Flow is directed into the filter's conical design, effectively capturing debris while maintaining blood flow

### Visible markers

- A gold tungsten loop around the mouth of the filter, along with radiopaque markers, allows for precise positioning and verification of apposition before proceeding with the intervention

### Wire movement

- The capture wire (available in 190 cm and 320 cm lengths) rotates and moves longitudinally, independent of the filter, for enhanced stability during the procedure
- The SpiderFX™ device is available in a variety of sizes (3 – 7 mm) for optimal fit and apposition in a range of vessels



### ORDER INFORMATION

Product Catalogue Number (1/Box)	Capture Wire				Delivery Catheter Cross Profile (F)	Recovery Catheter Diameter (F)	Guide Catheter Sheath Minimum ID (inch)
	Filter Size (mm)	Target Vessel Size (mm)	Wire Length OTW/RX (cm)	Wire Diameter (inch / mm)			
SPD2-030-190	3.0	2.0–3.0	190	0.014 / 0.36	3.2	4.2	0.066
SPD2-030-320	3.0	2.0–3.0	320 / 190	0.014 / 0.36	3.2	4.2	0.066
SPD2-040-190	4.0	3.1–4.0	190	0.014 / 0.36	3.2	4.2	0.066
SPD2-040-320	4.0	3.1–4.0	320 / 190	0.014 / 0.36	3.2	4.2	0.066
SPD2-050-190	5.0	4.1–5.0	190	0.014 / 0.36	3.2	4.2	0.066
SPD2-050-320	5.0	4.1–5.0	320 / 190	0.014 / 0.36	3.2	4.2	0.066
SPD2-060-190	6.0	4.5–6.0	190	0.014 / 0.36	3.2	4.2	0.066
SPD2-060-320	6.0	4.5–6.0	320 / 190	0.014 / 0.36	3.2	4.2	0.066
SPD2-070-190	7.0	5.5–7.0	190	0.014 / 0.36	3.2	4.2	0.066
SPD2-070-320	7.0	5.5–7.0	320 / 190	0.014 / 0.36	3.2	4.2	0.066

#### INDICATIONS:

The SpiderFX™ Embolic Protection System provides distal embolization protection during general vascular use, including peripheral, coronary, and carotid interventions.

# CROSSING CATHETERS

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# TrailBlazer™

## Support Catheter

TrailBlazer™ support catheter is a single lumen over the wire support catheter with a low-profile, tapered tip.

Three platinum / iridium markers are embedded between the two layers of this seamless catheter.

TrailBlazer™ is designed for increased pushability for crossing tight stenoses and occlusions.

**Each box includes:**

Five catheters in single sterile pouches.



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### ORDER INFORMATION

Product Catalogue Number (5/Box)	Guidewire Compatibility (inch)	Usable Catheter Length (cm)	Space Between Radiopaque Markers (mm)	Minimum Guide Catheter (F)	Minimum Introducer Sheath (F)
SC-035-065	0.035	65	50	6	5
SC-035-090	0.035	90	50	6	5
SC-035-135	0.035	135	50	6	5
SC-035-150	0.035	150	50	6	5
SC-018-090	0.018	90	15	5	4
SC-018-135	0.018	135	15	5	4
SC-018-150	0.018	150	15	5	4
SC-014-135	0.014	135	15	5	4
SC-014-150	0.014	150	15	5	4

INDICATIONS: The TrailBlazer™ Support Catheters are percutaneous, single lumen catheters designed for use in the peripheral vascular system.

TrailBlazer™ is intended to guide and support a guidewire during access of the vasculature, allow for wire exchanges, and provide a conduit for the delivery of saline solutions or diagnostic contrast agents.

# TrailBlazer™

## Angled Support Catheter



### **Braided catheter**

Stainless steel braid provides robust pushability and kink resistance

### **Ultra low profile tip**

Tapered design provides low lesion entry to aid in crossing

### **1:1 torque**

Reliable rotational control

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### ORDER INFORMATION

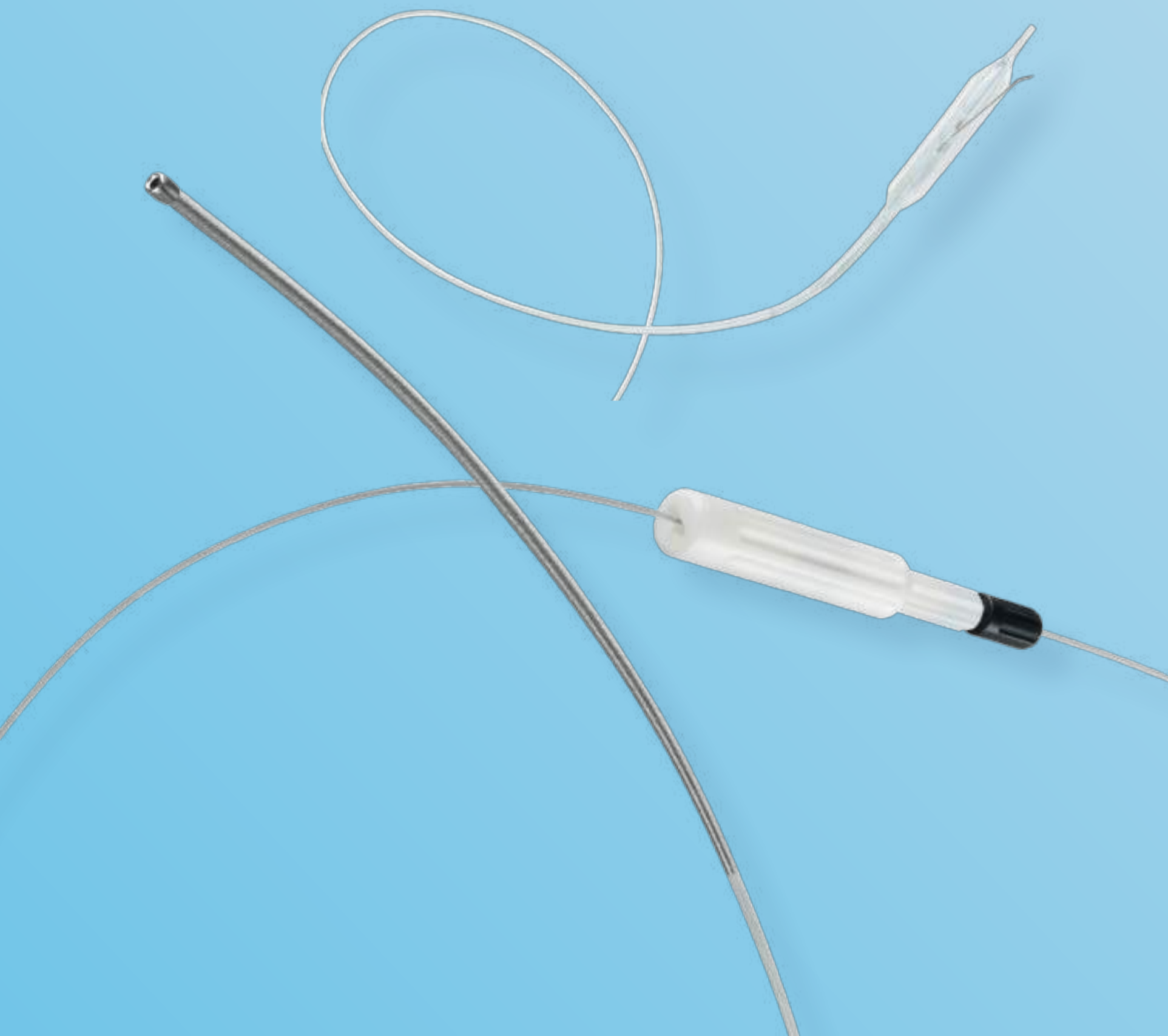
	Minimum Guide Sheath (F)	Minimum Introducer Sheath (F)	Marker Band Space (mm)	Size Outer Diameter (in)	Product Catalogue Number			
					65 cm	90 cm	135 cm	150 cm
0.014"	5	4	15	0.030" (2.3 F)		ASC-014-090	ASC-014-135	ASC-014-150
0.018"	5	4	15	0.034" (2.6 F)		ASC-018-090	ASC-018-135	ASC-018-150
0.035"	5	4	50	0.050"	ASC-035-065	ASC-035-090	ASC-035-135	ASC-035-150

# CTO DEVICES

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# Viance™

## Crossing Catheter

A precision instrument designed to quickly and safely deliver a guidewire via the true lumen, the Viance™ crossing catheter puts the control of crossing where it belongs: in your hands.

Providing an effective frontline option for CTOs, the Viance™ crossing catheter enables you to utilize a proactive technique to cross total occlusions via the true lumen.

### **Low profile atraumatic tip**

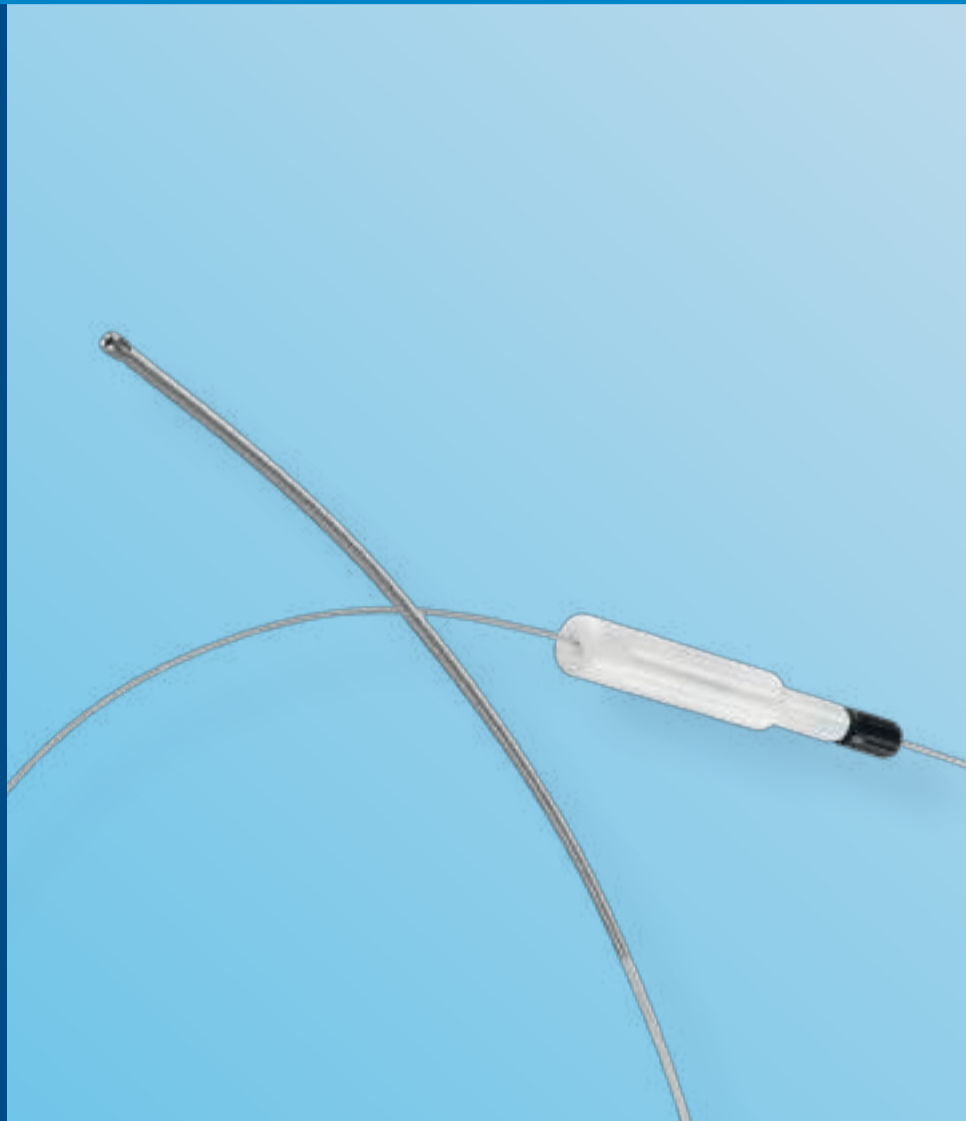
Designed for smooth crossing and minimizes risk of perforation.

### **Multi-coiled wire shaft**

Provides 1:1 torque.

### **Fast-spin torque handle**

Allows for tactile, self-controlled spinning motion enabling the Viance™ catheter tip to find its way through the lesion.



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## Crossing Catheter

### ORDER INFORMATION

Product Catalogue Number	Description	Working Length (cm)	Guidewire Compatibility (inch)	Crossing Profile (max inch)	Sheath Compatibility
VNC-FX-150	Flexible	150	0.014	0.038	5 F
VNC-SD-150	Standard	150	0.014	0.038	5 F

INDICATIONS: Viance™ Crossing Catheter is intended for use with a guidewire to access discrete regions of the peripheral vasculature.

# Enteer™

## Re-Entry Catheter

The Enteer re-entry system, consisting of the catheter and guidewire, gives you intuitive control to reliably target the true lumen from the subintimal channel above or below the knee.

The system requires no capital equipment. It's designed to be nothing less than a precise extension of your own expert hand.

### **Flat shaped self-orienting balloon**

The Enteer™ catheter's unique balloon design self-orientates towards the true lumen within the subintimal space when inflated.

### **180° and offset exit ports**

Offset exit ports are located on each side of the device allowing the Enteer™ guidewire to re-enter the correct port into the true lumen.

### **OTW 0.014" & 0.018" guidewire compatible**

Allows for flexibility during your case and minimizes guidewire exchanges.



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## Re-Entry Catheter

### ORDER INFORMATION

Product Catalogue Number	Balloon Size (W x H x L mm)	Working Length (cm)	Guidewire Compatibility (inch)	Crossing Profile (max inch)	Sheath Compatibility
ENB-375-20-135	3.75 x 1.5 x 20	135	≤ 0.018	0.066	5 F
ENB-275-20-150	2.75 x 1.0 x 20	150	≤ 0.018	0.066	5 F

INDICATIONS: Enteer™ Re-entry Catheter is indicated for directing, steering, controlling, and supporting a guidewire in order to access discrete regions of the peripheral vasculature.

### ORDER INFORMATION

Product Catalogue Number	Product	Description
ENW-FX-014-300	Enteer™ Guidewire	0.014" x 300 xm Flexible
ENW-SD-014-300	Enteer™ Guidewire	0.014" x 300 cm Standard
ENW-SF-014-300	Enteer™ Guidewire	0.014" x 300 cm Stiff

# CATHETERS

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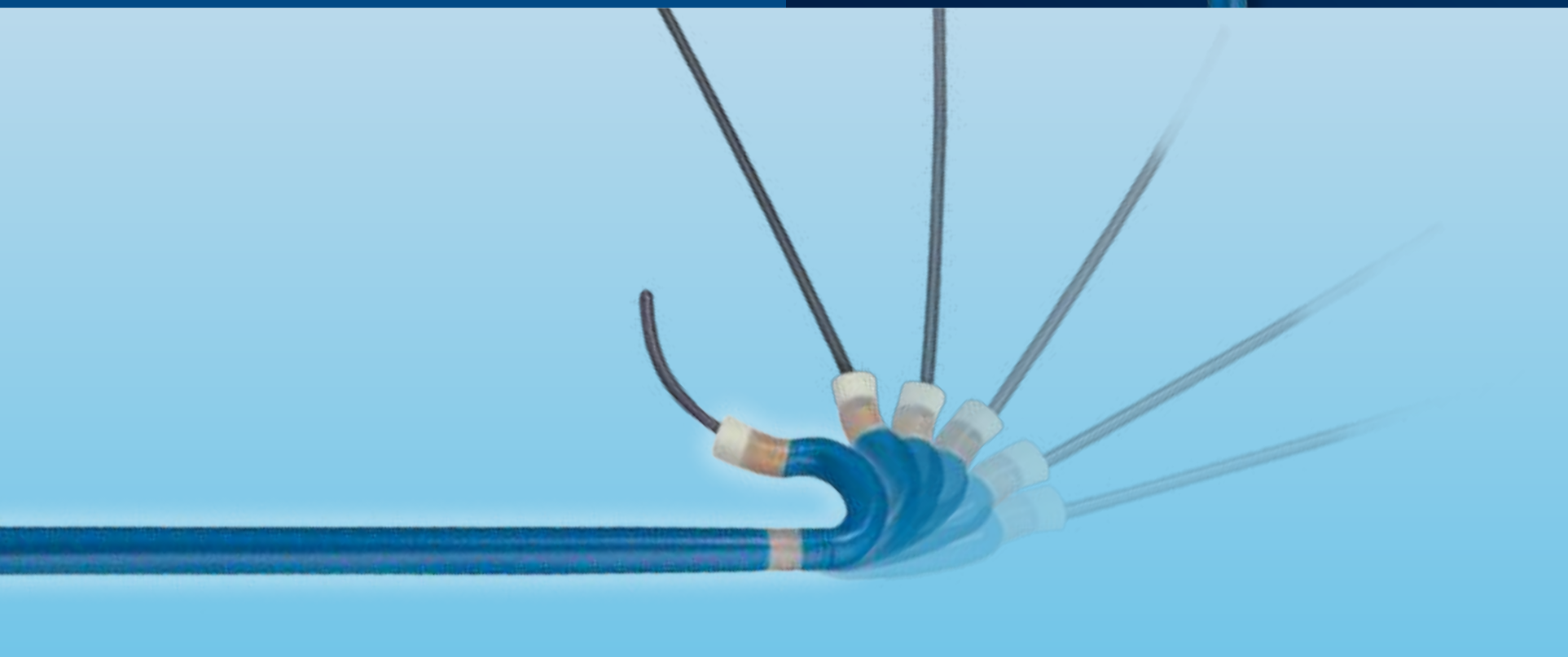
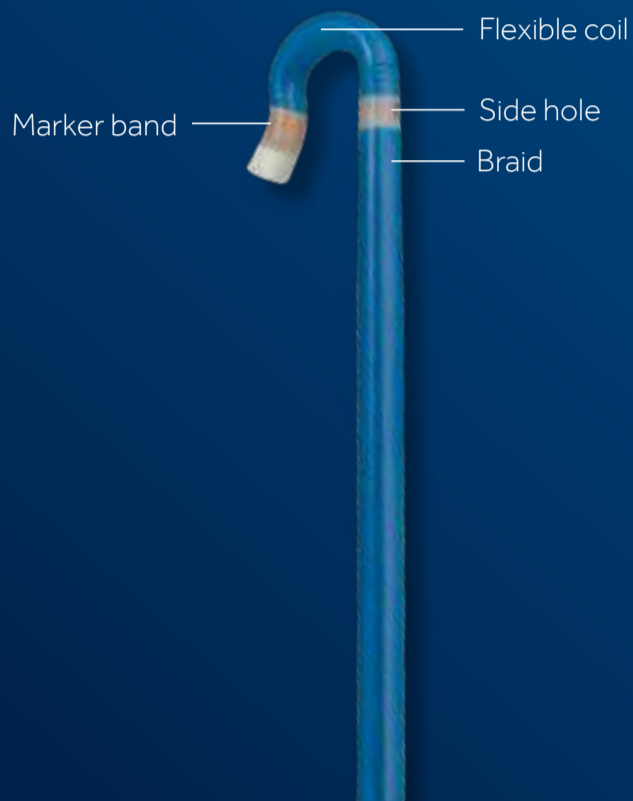
# Piton™ GC

## Carotid Guide Catheter

### ACCESSING THE FUTURE OF CAS INTERVENTION

#### TECHNICAL SPECIFICATIONS

Outer diameter	8 F (0.104" / 0.264 mm)
Inner diameter	5 F (0.073" / 0.186 mm)
Total catheter length	91 cm
Usable catheter length	85 cm
Guidewire outer diameter	max. 2 x 0.035"



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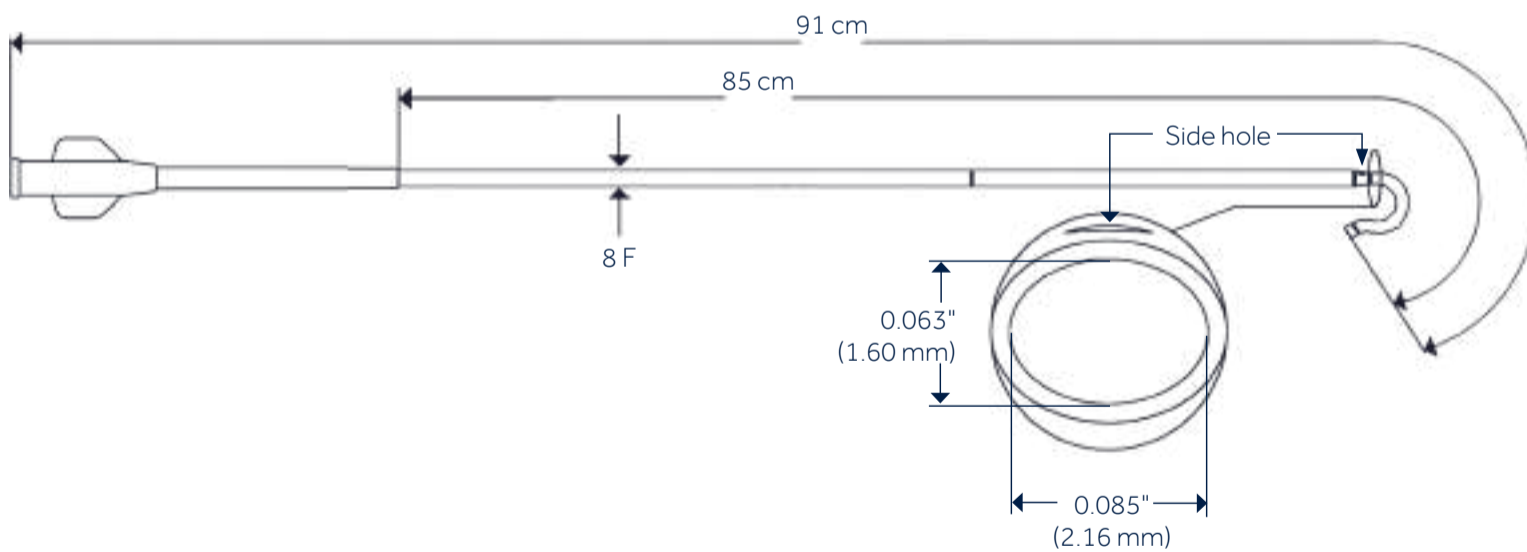
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## Carotid Guide Catheter

### ORDER INFORMATION

Product Catalogue Number	Usable Length (cm)	Tip Curve
PTN8SC063085	85 cm	Small



# Rebar™

## Reinforced Microcatheter



The Rebar™ reinforced microcatheter is an end hole single-lumen catheter designed to be introduced via a steerable guidewire into the vasculature.

The catheter has a semi-rigid proximal shaft which transitions into the flexible distal shaft to facilitate the advancement of the catheter into the anatomy.

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## Reinforced Microcatheter

### ORDER INFORMATION

Product Name	Product Catalogue Number	Proximal OD/ Distal OD (F)	Distal ID (inch)	Total Length (cm)	Usable Length (cm)	Maximum Guidewire (inch)
Rebar™ -18	105-5081-130	2.7 / 2.4	0.021	137	130	0.018
Rebar™ -18	105-5083-153	2.7 / 2.4	0.021	160	153	0.018
Rebar™ -27	105-5082-145	2.8 / 2.8	0.027	150	145	0.021

### TECHNICAL SPECIFICATIONS

Stainless steel reinforced microcatheter	Lubricious hydrophilic outer coating	Steam-shapeable tips
This microcatheter is DMSO compatible and therefore optimized for delivering the Onyx™ Liquid Embolic System	Less effort required to track through tortuous vessels	The tip of the catheter can be steam-shaped using the mandrel provided
High resistance to kinking when maneuvering around tight bends	Smooth movement when navigating through vasculature	Onyx™ Liquid Embolic System compatible
Radiopaque marker at the distal end facilitates fluoroscopic visualization		

# THROMBUS MANAGEMENT

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# Cragg-McNamara™

## Valved Infusion Catheters

The Cragg-McNamara™ Valved Infusion Catheter is a single lumen infusion catheter with a valved tip that allows infusion without the need of a tip-occluding guidewire.

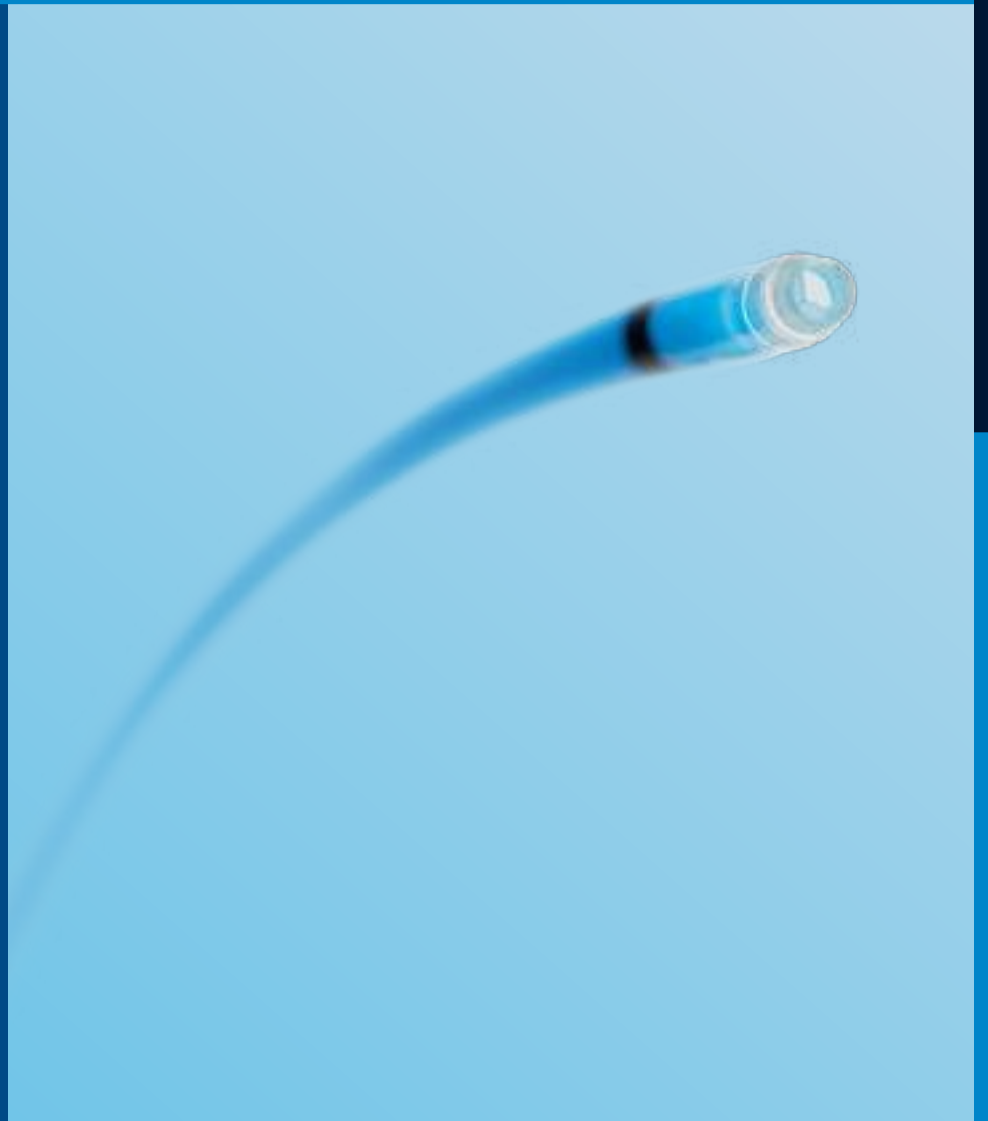
### Large infusion lumen

Only 5 F Cragg-McNamara™ catheters give you the option to infuse without a guidewire in place, nearly doubling the infusion lumen area.

### Streamlined patient care

Infuse overnight, without a guidewire in place which eliminates the risk of guidewire movement for simplified patient care.

Treat without a tip-occluding guidewire and sidearm adapter, which may contribute to cost savings.



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### ORDER INFORMATION

Product Catalogue Number (1 / Box)	Diameter (F)	Usable Length (cm)	Infusion Length (cm)	Recommended Guidewire (inch)
41032-01	4	40	10	0.035
41033-01	4	40	20	0.035
41034-01	4	65	5	0.035
41035-01	4	65	10	0.035
41036-01	4	65	20	0.035
41037-01	4	100	5	0.035
41038-01	4	100	10	0.035
41039-01	4	100	20	0.035
41040-01	4	135	5	0.035
41041-01	4	135	10	0.035
41042-01	4	135	20	0.035
41043-01	5	40	5	0.038
41044-01	5	40	10	0.038
41045-01	5	40	20	0.038
41046-01	5	65	5	0.038
41047-01	5	65	10	0.038
41048-01	5	65	20	0.038
41049-01	5	100	5	0.038
41050-01	5	100	10	0.038
41051-01	5	100	20	0.038
41052-01	5	100	30	0.038
41053-01	5	100	40	0.038
41054-01	5	100	50	0.038
41055-01	5	135	5	0.038
41056-01	5	135	10	0.038
41057-01	5	135	20	0.038
41058-01	5	135	30	0.038
41059-01	5	135	40	0.038
41060-01	5	135	50	0.038

INDICATIONS: The Cragg-McNamara™ Infusion Catheter is indicated for use in the controlled, selective infusion of physician-specified pharmacological agents or radiopaque contrast media into the general vasculature.

# MicroMewi™

## Multiple Sidehole Infusion Catheters

The MicroMewi™ multiple sidehole infusion catheters feature radiopaque platinum markers providing fluoroscopic visualization for precise catheter placement.

Flexible and trackable distal catheter segment allows access to tortuous anatomy.

### Design Details

- Radiopaque platinum markers provide for exceptional visualization and precise catheter placement
- The flexible and trackable distal catheter segment permits easy navigation through tortuous anatomy for above- or below-the knee applications
- **Multiple sideholes** permit direct infusion into a thrombosed segment



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## Multiple Sidehole Infusion Catheters

### ORDER INFORMATION

Product Catalogue Number (1/Box)	Diameter (F)	Usable Length (cm)	Infusion Length (cm)	Recommended Guidewire (inch)
41063-01	2.9	150	5	0.018
41064-01	2.9	150	10	0.018
41066-01	2.9	180	5	0.018
41067-01	2.9	180	10	0.018

INDICATIONS: The MicroMewi™ multiple sidehole infusion catheter is indicated for use in the controlled, selective infusion of physician-specified pharmacological agents or radiopaque contrast media into general vasculature.



# ProStream™

## Multiple Sidehole Infusion Wires

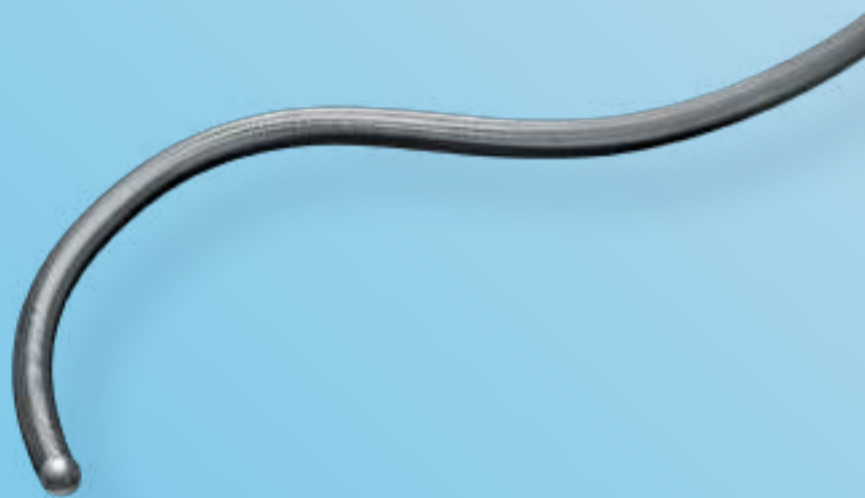
The ProStream™ multiple sidehole infusion wires are constructed with an integral core wire, stainless steel coil and an outer Teflon™ layer.

The wires are available in a wide variety of sidehole infusion lengths.

The ProStream™ multiple sidehole infusion wires can be used coaxially through 5 F infusion catheters.

### Unique Core Wire Design

- Radiopaque markers
- Eliminates the need for a separate core wire
- Provides torqueability for added control
- Is small enough to reach distal vessels alone or when used inside a 5 F coaxial system
- Enhances visualization and permits precise placement proximal and distal to the infusion length



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## Multiple Sidehole Infusion Wires

### ORDER INFORMATION

Product Catalogue Number (1 / Box)	Usable Length (cm)	Infusion Length (cm)	Diameter (inch)
41272-01	145	6	0.035
41273-01	145	9	0.035
41274-01	145	12	0.035
41276-01	175	6	0.035
41277-01	175	9	0.035
41278-01	175	12	0.035

INDICATIONS: The ProStream™ Multiple Sidehole Infusion Wire is indicated for use in the controlled, selective infusion of physician-specified pharmacological agents or radiopaque contrast media into general vasculature.

# GUIDEWIRES

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# Nitrex™

## Guidewires

The Nitrex™ Guidewires are constructed of a solid nitinol core offering nitinol kink-resistance and 1:1 torque.

All models feature a silicone coating and a gold-plated tungsten coil for enhanced radiopacity. The guidewires also come in a variety of sizes and angles.

**Each box includes:**

Three guidewires in carrying hoop. Torque devices included on 0.014" and 0.018" wire sizes.



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### ORDER INFORMATION

Product Catalogue Number (3 / Box)	Diameter (inch)	Length (cm)	Tip Style	Tip Length (cm)	Tip Shape	Tip Angle
<b>0.014"</b>						
N140801	0.014	80	INT	5	Angle	15°
N141802	0.014	180	INT	5	Angle	15°
N143001	0.014	300	INT	5	Angle	15°
<b>0.018"</b>						
N180601	0.018	60	INT	5	Straight	0
N180603	0.018	60	INT	7	Straight	0
N180801	0.018	80	STD	2	Straight	0
N180802	0.018	80	INT	5	Angle	15°
N181804	0.018	180	STD	2	Straight	0
N181805	0.018	180	INT	5	Angle	15°
N181806	0.018	180	FLOP	20	Angle	15°
N183001	0.018	300	STD	2	Straight	0
N183002	0.018	300	INT	5	Angle	15°
<b>0.025"</b>						
N251801	0.025	180	INT	8	Angle	15°
N251802	0.025	180	STD	2	Straight	0
N252601	0.025	260	INT	8	Angle	15°

INDICATIONS: The 0.014" (0.36 mm) and 0.018" (0.46 mm) diameter Nitrex™ Guidewires are intended for use in the peripheral and coronary vasculature. The 0.025" (0.64 mm) and 0.035" (0.89 mm) diameter Nitrex™ nitinol Guidewires are indicated for use in the peripheral vasculature.

ABBREVIATIONS:

INT: Intermediate - STD: Standard - FLOP: Floppy Indications

### ORDER INFORMATION

Product Catalogue Number (3 / Box)	Diameter (inch)	Length (cm)	Tip Style	Tip Length (cm)	Tip Shape	Tip Angle
<b>0.035" FLEXIBLE SHAFT</b>						
N351451	0.035	145	INT	15	Straight	0
N351452	0.035	145	INT	15	Angle	45°
N351803	0.035	180	INT	15	Straight	0
N352601	0.035	260	INT	15	Angle	45°
N354001	0.035	400	INT	15	Straight	0
<b>0.035" STIFF SHAFT</b>						
N350801	0.035	80	INT	9	Straight	0
N351453	0.035	145	FLOP	14	Angle	45°
N351455	0.035	145	FLOP	14	Straight	0
N351454	0.035	145	INT	9	Straight	0
N351804	0.035	180	INT	9	Straight	0
N351805	0.035	180	STD	4	Angle	45°
N352602	0.035	260	FLOP	14	Straight	0
N352604	0.035	260	INT	9	Straight	0
N352603	0.035	260	STD	4	Angle	45°
N353001	0.035	300	INT	9	Straight	0
N354002	0.035	400	INT	9	Straight	0

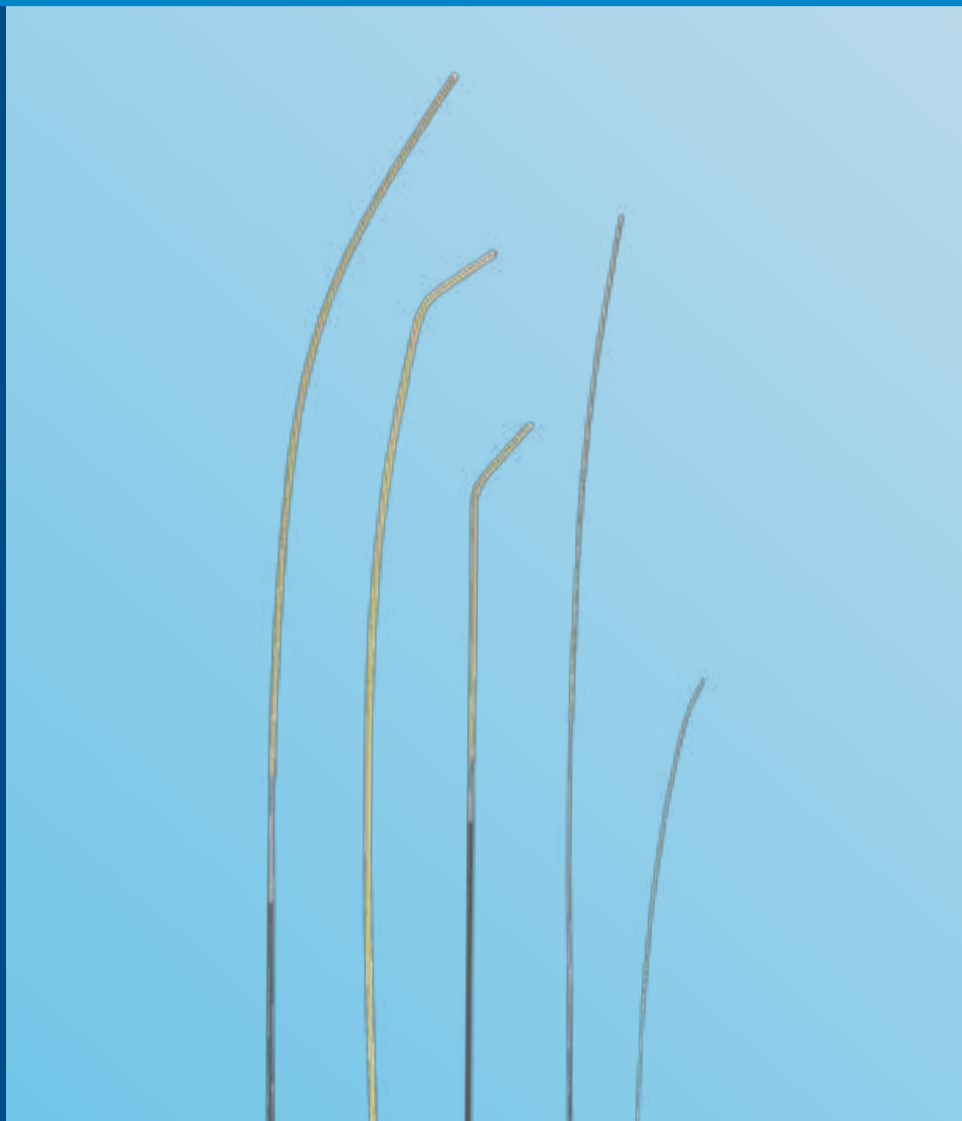
# Babywire™

## Double-Ended Nitinol Guidewire

The Babywire™ Double-Ended Nitinol Guidewires assist the placement of IV Catheters and exchange of small vessel arterial / venous lines.

**Each box includes:**

Ten wires



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## Double-Ended Nitinol Guidewire

### ORDER INFORMATION

Product Catalogue Number (10 / Box)	Diameter (inch)	Length (cm)
BW1200	0.012	18
BW1201	0.012	50

INDICATIONS: The Babywire™ Guidewire is intended for assisting the placement of initial catheters and/or exchange in the small vessel anatomy. The Babywire™ Guidewire is compatible with a 24 - gauge needle or 2.0 F catheter.



# AqWire™

## Guidewire

The AqWire™ guidewire combines the lubricity of a hydrophilic coating with the durability and kink resistance of a solid nitinol core.

**Each box includes:**

Three hydrophilic guidewires.

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### ORDER INFORMATION

Product Catalogue Number (3/Box)	Diameter (inch)	Length (cm)	Body Type	Tip Angle
<b>0.018"</b>				
A181501	0.018	150	Standard	0
A181502	0.018	150	Standard	45°
A181801	0.018	180	Standard	0
A181802	0.018	180	Standard	45°
A182601	0.018	260	Standard	0
A182602	0.018	260	Standard	45°
<b>0.035" STANDARD BODY</b>				
A351501	0.035	150	Standard	0
A351502	0.035	150	Standard	45°
A351801	0.035	180	Standard	0
A351802	0.035	180	Standard	45°
A352601	0.035	260	Standard	0
A352602	0.035	260	Standard	45°
<b>0.035" STIFF BODY</b>				
A351503	0.035	150	Stiff	0
A351504	0.035	150	Stiff	45°
A351803	0.035	180	Stiff	0
A351804	0.035	180	Stiff	45°
A352603	0.035	260	Stiff	0
A352604	0.035	260	Stiff	45°

INDICATIONS: contraindications, warnings and instructions for use can be found in the product labeling supplied with each device.

# Wholey™

## Guidewire System 0.035"

The Wholey™ Guidewire System provides enhanced torqueability and lubricity, allowing interventionalists to approach challenging cases with confidence.

**Each box includes:**

Three hydrophilic coated guidewires.

**Steer**

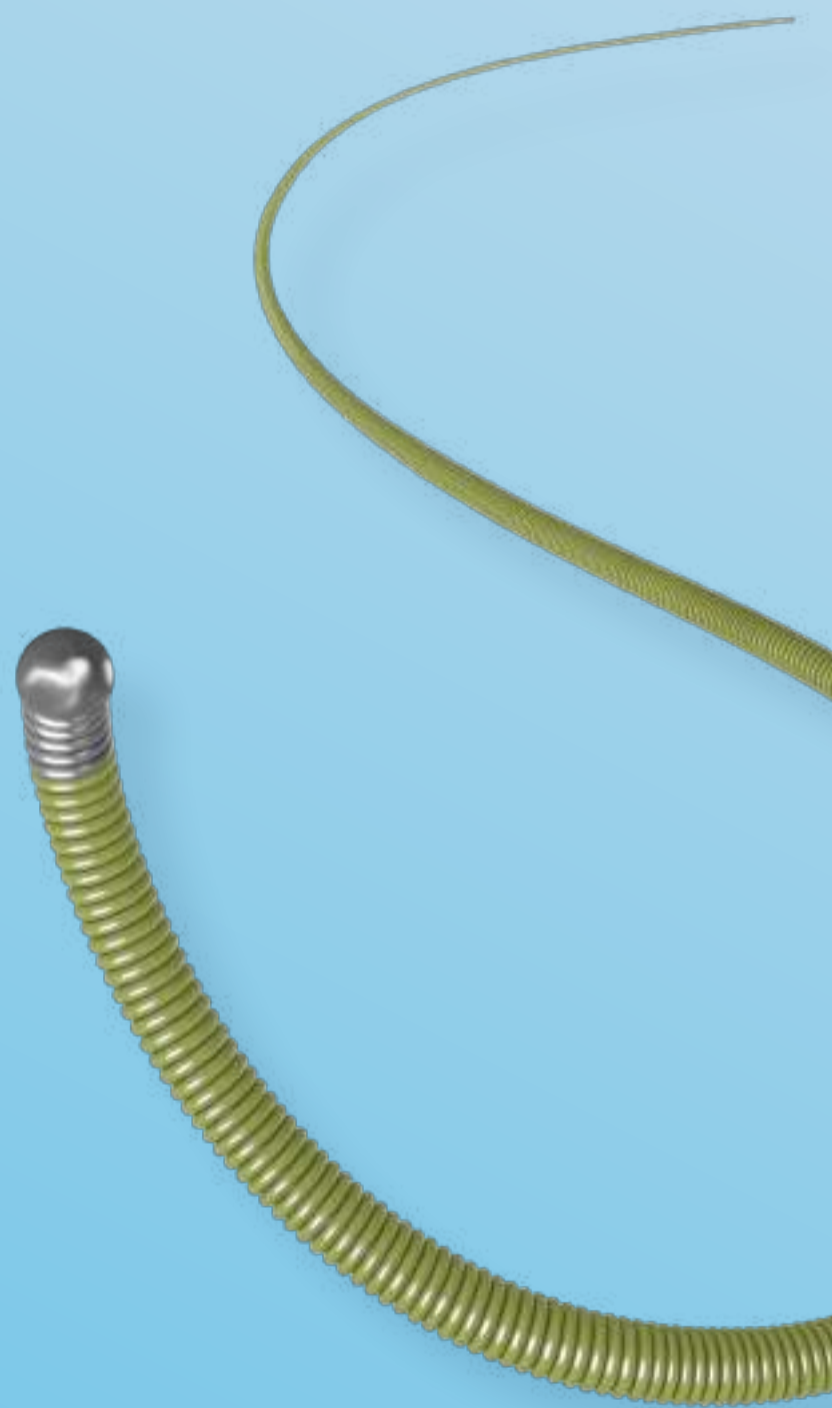
One-to-one torque for precise navigation in tortuous anatomy.

**Slide**

Proprietary precoating technology provides consistent coating over entire coil.

**See**

Platinum tungsten coil tip offers high visibility to assist with accurate placement.



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## Guidewire System 0.035"

### ORDER INFORMATION

Product Catalogue Number	Description	Stiffness Profile	Tip Style	Outer Diameter (inch)	Length (cm)	Quantity
WWFS35145	Floppy tip, extension compatible	Floppy	Straight / shapeable	0.035	145	3/pkg
WWFS35175	Floppy tip, extension compatible	Floppy	Straight / shapeable	0.035	175	3/pkg
WWFS35260	Floppy tip, exchange length	Floppy	Straight / shapeable	0.035	260	3/pkg
WWFS35300	Floppy tip, exchange length	Floppy	Straight / shapeable	0.035	300	3/pkg
WWIJ35145	Modified J tip, extension compatible	Intermediate	Modified J / shapeable	0.035	145	3/pkg
WWIJ35175	Modified J tip, extension compatible	Intermediate	Modified J / shapeable	0.035	175	3/pkg
WWIJ35260	Modified J tip, exchange length	Intermediate	Modified J / shapeable	0.035	260	3/pkg
WWIJ35300	Modified J tip, exchange length	Intermediate	Modified J / shapeable	0.035	300	3/pkg
WWSS35145	Standard tip, extension compatible	Standard	Straight / shapeable	0.035	145	3/pkg
WWSS35175	Standard tip, extension compatible	Standard	Straight / shapeable	0.035	175	3/pkg
WWSS35260	Standard tip, exchange length	Standard	Straight / shapeable	0.035	260	3/pkg
WWSS35300	Standard tip, exchange length	Standard	Straight / shapeable	0.035	300	3/pkg
WWES35001	Extension system	Standard	Straight / shapeable	0.035	155	3/pkg

INDICATIONS: The Wholey™ guidewire system is intended to facilitate the placement and exchange of interventional devices during diagnostic or therapeutic interventional procedures. The guidewire can be torqued to facilitate navigation through tortuous arteries and/or avoid unwanted side branches.

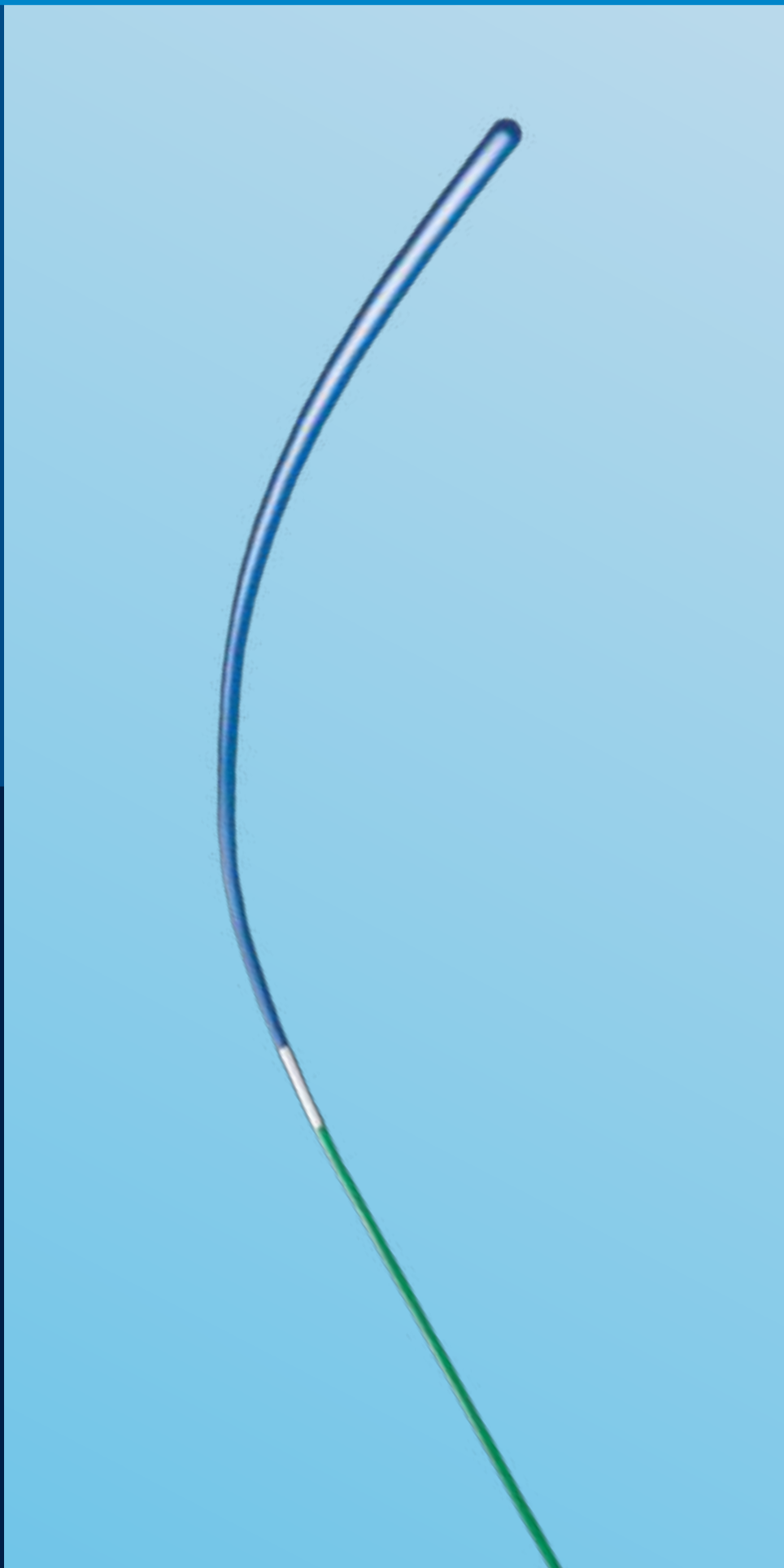
# Kitewire™ Deep

Peripheral Guidewire 0.014"

SMOOTH TRACKABILITY FOR LONG DIFFUSE

## TECHNICAL SPECIFICATIONS

Guidewire outer diameter	0.014"
Coating of the guidewire coil	Polymer and hydrophilic
Total length of the guidewire	195, 250 and 300 cm
Support	Extra support
Radiopaque coil length	3 cm platinum alloy spring coil
Tip shape	Straight, shapeable
Tip stiffness variations	Standard and intermediate



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# Kitewire™ Deep

## Peripheral Guidewire 0.014"

### ORDER INFORMATION

Product Catalogue Number	Usable Length (cm)	Tip Stiffness
KTD 195 INT 14S	195	Intermediate
KTD 195 STD 14S	195	Standard
KTD 250 INT 14S	250	Intermediate
KTD 250 STD 14S	250	Standard
KTD 300 INT 14S	300	Intermediate
KTD 300 STD 14S	300	Standard

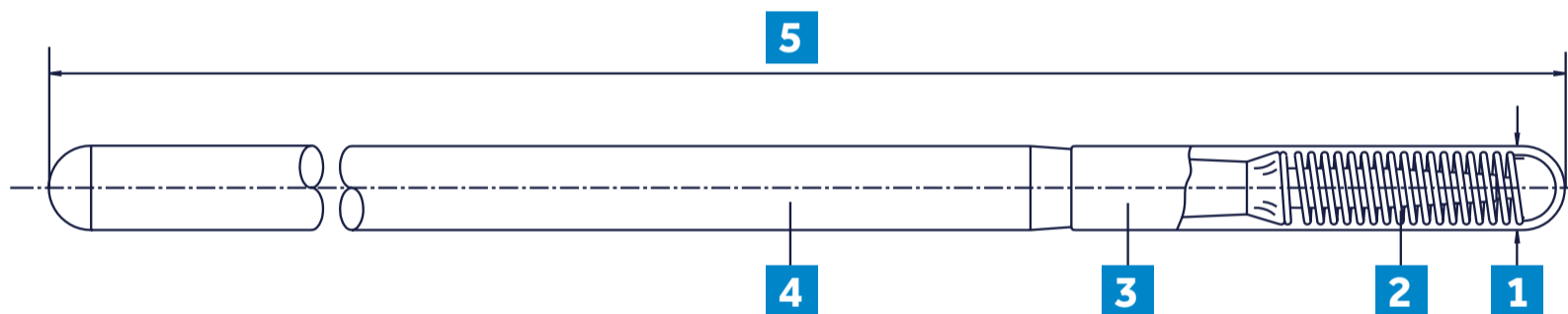
1 Outer diameter

3 Platinum coil

5 Polymer jacket

2 Stainless steel

4 Usable length



# SNARES

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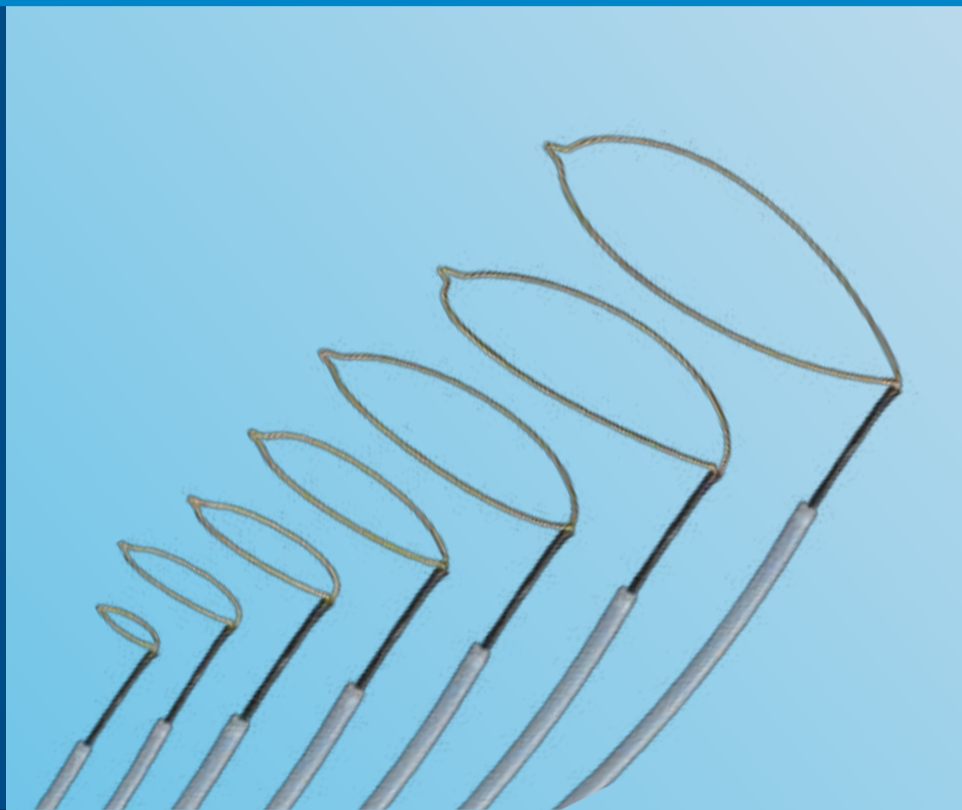
# Amplatz GooseNeck™

## Snare Kit

Engineered for precise retrieval and manipulation, the Amplatz GooseNeck™ Snares and Microsnares (for small vessel applications) feature a highly radiopaque snare loop that is 90° to shaft of the snare. Other features include a nitinol shaft for kink resistance and gold tungsten loop for enhanced visualization.

**Each kit includes:**

One snare, one snare catheter, one introducer and one torque device.



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# Amplatz GooseNeck™

## Snare Kit

### ORDER INFORMATION

Product Catalogue Number (1/box)	Catheter o.d (F)	Catheter Length (cm)
MC4000	4	102
MC4001	4	48
MC6000	6	102
MC6001	6	48

**Each kit includes:**

One snare, one snare catheter, one introducer and one torque device.

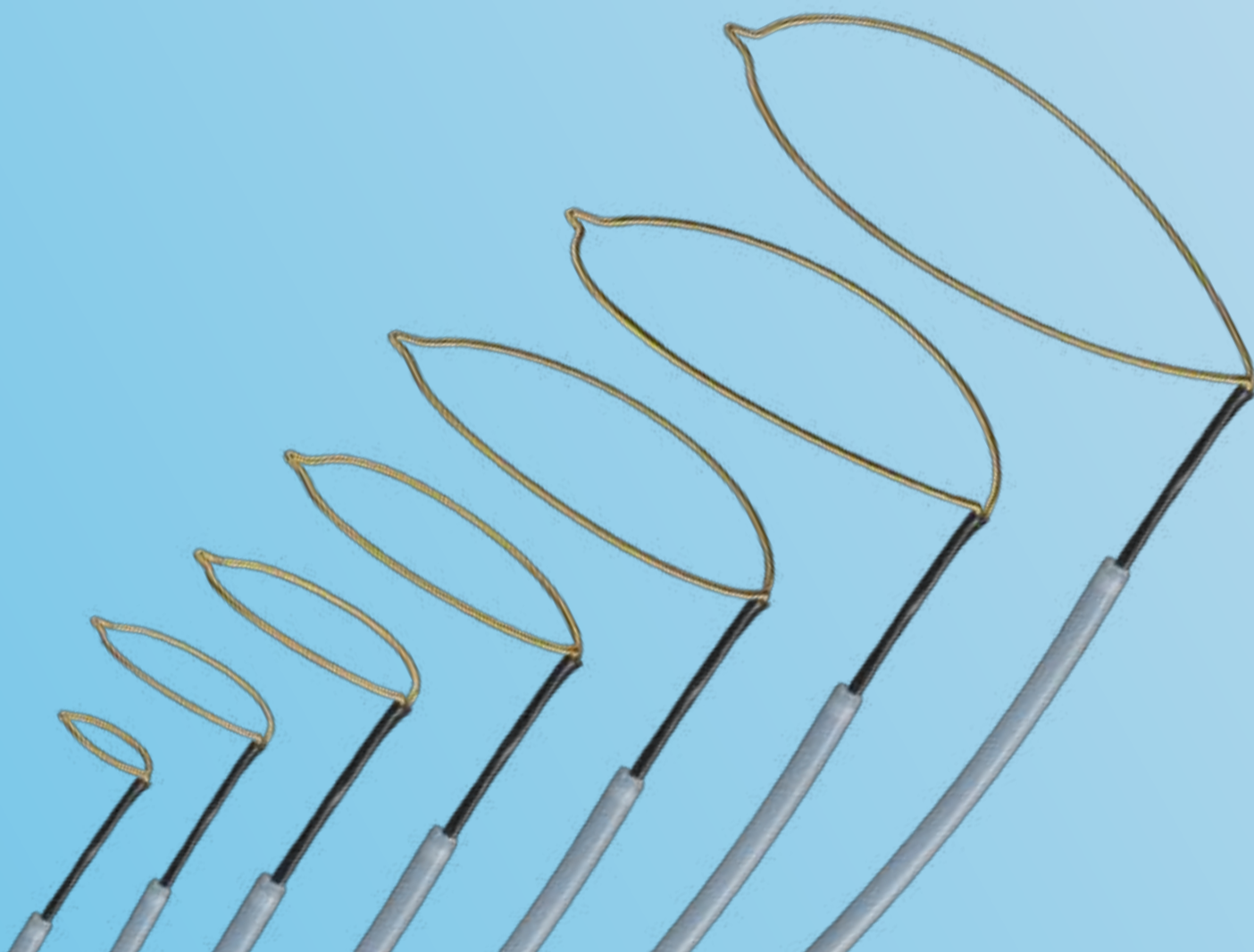
### ORDER INFORMATION

Product Catalogue Number (1/box)	Loop Diameter (mm)	Snare Length (cm)	Catheter Size (F)	Catheter Length (cm)
GN500	5	120	4	102
GN1000	10	120	4	102
GN1001	10	65	4	48
GN1500	15	120	6	102
GN2000	20	120	6	102
GN2501	25	65	6	48
GN2500	25	120	6	102
GN3000	30	120	6	102
GN3500	35	120	6	102

INDICATIONS: The Amplatz GooseNeck™ Snare is intended for use in the cardiovascular system or hollow viscus to retrieve and manipulate foreign objects. Manipulation procedures include indwelling venous catheter, fibrin sheath stripping, and central venous access venipuncture procedure assistance.

# Amplatz GooseNeck™

MicroSnare Kit



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# Amplatz GooseNeck™

## MicroSnare Kit

### ORDER INFORMATION

Product Catalogue Number (1/box)	Loop Diameter (mm)	Snare Length (cm)	Catheter Size Distal-Proximal (F)	Catheter Length (cm)
SK200	2	175	2.3 - 3	150
SK201	2	200	2.3 - 3	175
SK400	4	175	2.3 - 3	150
SK401	4	200	2.3 - 3	175
SK700	7	175	2.3 - 3	150
SK701	7	200	2.3 - 3	175

INDICATIONS: The Amplatz GooseNeck™ Snare is intended for use in the retrieval and manipulation of atraumatic foreign bodies located in the coronary and peripheral cardiovascular system and the extra-cranial neurovascular anatomy.

Each kit includes: one snare, one snare catheter, one introducer, and one torque device

# Y-CONNECTORS

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# Bigeasy™

## Rotating Y-connector

### ORDER INFORMATION

Product Catalogue Number (5/package)	Description
MVA100	2-way Adjustable Valve

# Sequel™

## Rotating Double Y-connector

### ORDER INFORMATION

Product Catalogue Number (5/package)	Description
MVA200	2-way Adjustable Valve

# VASCULAR EMBOLIZATION

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# Onyx™

## Liquid Embolic System

The Onyx™ Liquid Embolic System is an EVOH co-polymer designed to provide complete occlusion in a controlled embolization procedure, achieving clinical success across a variety of applications.



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# Liquid Embolic System

## ORDER INFORMATION

Product Catalogue Number	Onyx™ Formulation
105-7200-060	Onyx™ 18 kit 1.5 ml
105-7200-080	Onyx™ 34 kit 1.5 ml

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INDICATIONS: Embolization of lesions in the peripheral vasculature, including endoleaks, arteriovenous malformations, portal veins, bleeding, and tumors.



# Onyx™ 34L

## Liquid Embolic System

Onyx™ 34L Liquid Embolic System has less tantalum compared to the current version of Onyx™ Liquid Embolic System 34.

Clinical Benefit: Less streak artifacts on CT with a good visibility during injection. Available in 1.5 ml and 6 ml vials.



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# Onyx™ 34L

## Liquid Embolic System

### ORDER INFORMATION

Product Catalogue Number	Onyx™ Formulation
105-7315-080	Onyx™ 34L kit 1.5 ml
105-7360-080	Onyx™ 34L kit 6.0 ml

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INDICATIONS: Embolization of lesions in the peripheral vasculature, including endoleaks, arteriovenous malformations, portal veins, bleeding and tumors.

# Onyx™

## Accessories

### Vial mixer

It contains four spaces for preparation of Onyx™ 1.5 ml vial and two spaces for preparation of Onyx™ 6 ml vial simultaneously.

### Syringe catheter interface adapter

This device is an Onyx™ Syringe Catheter Interface Adapter and DMSO compatible adapter used to provide an interface between a Covidien 1 ml syringe and the 1.5 F UltraFlow HPC / 1.5 F Marathon™ and Apollo™ microcatheter during an Onyx™ embolization.

### 1ml Luer-lock injection syringe



# Onyx™ Mixer

## Vial Mixer

### ORDER INFORMATION

Product Catalogue Number	Information
103-1205-002	240 V
103-1205-100	New mixer attachment

# Onyx™

## Syringe Catheter Interface Adapter

### ORDER INFORMATION

Product Catalogue Number	Capacity (ml)	Syringes/ Box
103-1207	1	20

# 1ml Luer-Lock Injection Syringe

### ORDER INFORMATION

Product Catalogue Number	Capacity (ml)	Syringes/ Box
103-1203	1	10

INDICATIONS: The Onyx™ Mixer facilitates proper suspension of the Onyx™ tantalum for better visualization prior to use. The proximal end of the Onyx™ Syringe Catheter Interface Adapter incorporates a standard ISO, female luer design to facilitate connection to the syringe. The distal end is designed specifically to fit the hub of the 1.5 F UltraFlow™ HPC / Marathon™ 1.5 F and Apollo™ microcatheter.

# Concerto™ Helix/3D

## Detachable Coil System

The Concerto™ Detachable Coil System is a stretch-resistant, detachable coil that can be repositioned easily prior to detachment and uses enlaced microfilament technology called LatticeFX™.

### Softness with smooth navigation

- Soft coils track easily through tortuous anatomy to access distal locations\*
- Soft distal pusher reduces microcatheter kickback during deployment\*

### Reliable deployment

- Fully resheathable, after complete or partial deployment, and easily repositionable\*
- Coil detaches instantaneously with proven reliability\*

### Optimal framing

- Conformable 3D shape with excellent stability\*
- Designed to create a complex frame for filling\*

### Enhanced thrombogenicity

- Fibers increase thrombogenicity of the coil compared to bare metal equivalents†
- Nylon and PGLA fiber system features the unique LatticeFX™ technology which promotes thrombosis response†



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\*Internal data on file

# Concerto™ Helix/3D

## Detachable Coil System

### ORDER INFORMATION

Product Catalogue Number (1 / box)	Description	Diameter (mm)	Length (cm)	Min. Microcatheter Inner Diameter (inch)
<b>HELIX</b>				
NV-2-4-Helix	Concerto™ Nylon Helical	2	4	0.0165
NV-2-6-Helix	Concerto™ Nylon Helical	2	6	0.0165
NV-2-8-Helix	Concerto™ Nylon Helical	2	8	0.0165
NV-3-4-Helix	Concerto™ Nylon Helical	3	4	0.0165
NV-3-8-Helix	Concerto™ Nylon Helical	3	8	0.0165
NV-4-8-Helix	Concerto™ Nylon Helical	4	8	0.0165
NV-4-10-Helix	Concerto™ Nylon Helical	4	10	0.0165
NV-5-15-Helix	Concerto™ Nylon Helical	5	15	0.021
NV-5-20-Helix	Concerto™ Nylon Helical	5	20	0.021
NV-6-20-Helix	Concerto™ Nylon Helical	6	20	0.021
NV-7-30-Helix	Concerto™ Nylon Helical	7	30	0.021
NV-8-30-Helix	Concerto™ Nylon Helical	8	30	0.021
NV-9-30-Helix	Concerto™ Nylon Helical	9	30	0.021
NV-10-30-Helix	Concerto™ Nylon Helical	10	30	0.021
PV-12-30-Helix	Concerto™ PGLA Helical	12	30	0.021
PV-14-30-Helix	Concerto™ PGLA Helical	14	30	0.021
PV-16-40-Helix	Concerto™ PGLA Helical	16	40	0.021
PV-18-40-Helix	Concerto™ PGLA Helical	18	40	0.021
PV-20-50-Helix	Concerto™ PGLA Helical	20	50	0.021
<b>3D</b>				
PV-2-2-3D	Concerto™ PGLA 3D	2	2	0.0165
PV-2-4-3D	Concerto™ PGLA 3D	2	4	0.0165
PV-2-6-3D	Concerto™ PGLA 3D	2	6	0.0165
PV-3-4-3D	Concerto™ PGLA 3D	3	4	0.0165
PV-3-6-3D	Concerto™ PGLA 3D	3	6	0.0165
PV-3-8-3D	Concerto™ PGLA 3D	3	8	0.0165
PV-4-8-3D	Concerto™ PGLA 3D	4	8	0.0165
PV-4-10-3D	Concerto™ PGLA 3D	4	10	0.0165
PV-4-12-3D	Concerto™ PGLA 3D	4	12	0.0165
PV-5-15-3D	Concerto™ PGLA 3D	5	15	0.0165
PV-6-20-3D	Concerto™ PGLA 3D	6	20	0.0165
PV-7-30-3D	Concerto™ PGLA 3D	7	30	0.0165
PV-8-30-3D	Concerto™ PGLA 3D	8	30	0.0165
PV-9-30-3D	Concerto™ PGLA 3D	9	30	0.0165
PV-10-30-3D	Concerto™ PGLA 3D	10	30	0.0165
PV-12-40-3D	Concerto™ PGLA 3D	12	40	0.021
PV-14-40-3D	Concerto™ PGLA 3D	14	40	0.021
PV-16-40-3D	Concerto™ PGLA 3D	16	40	0.021
PV-18-40-3D	Concerto™ PGLA 3D	18	40	0.021

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# I.D. Instant Detacher

Detacher for Concerto™ Detachable Coil System,  
One detacher required per procedure.

**Each box contains:**  
Five instant detachers.



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# I.D. Instant Detacher

## ORDER INFORMATION

Product Catalogue Number	Number/ Box
ID-1-5	5



# MVP™

## Microvascular Plug System

The MVP™ Microvascular Plug System is indicated to obstruct or reduce the rate of blood flow in the peripheral vasculature.



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## Microvascular Plug System

### ORDER INFORMATION

Product Catalogue Number	Recommended Vessel Size	MVP™ Device Outer Diameter Unconstrained	MVP™ Device Length Unconstrained	Delivery Wire Length	Minimum Recommended Catheter Dimensions	Recommended Microcatheter
MVP-3Q	1.5 – 3.0 mm	5.3 mm	12 mm	180 cm	ID ≥ 0.021"	150 cm
MVP-5Q	3.0 – 5.0 mm	6.5 mm	12 mm	180 cm	ID ≥ 0.027"	150 cm
MVP-7Q	5.0 – 7.0 mm	9.2 mm	16 mm	165 cm	ID ≥ 0.035" <b>AND</b> OD ≥ 4F	120 cm
MVP-9Q	7.0 – 9.0 mm	13 mm	18 mm	165 cm	ID ≥ 0.038" <b>AND</b> OD ≥ 5F	120 cm

# VENOUS



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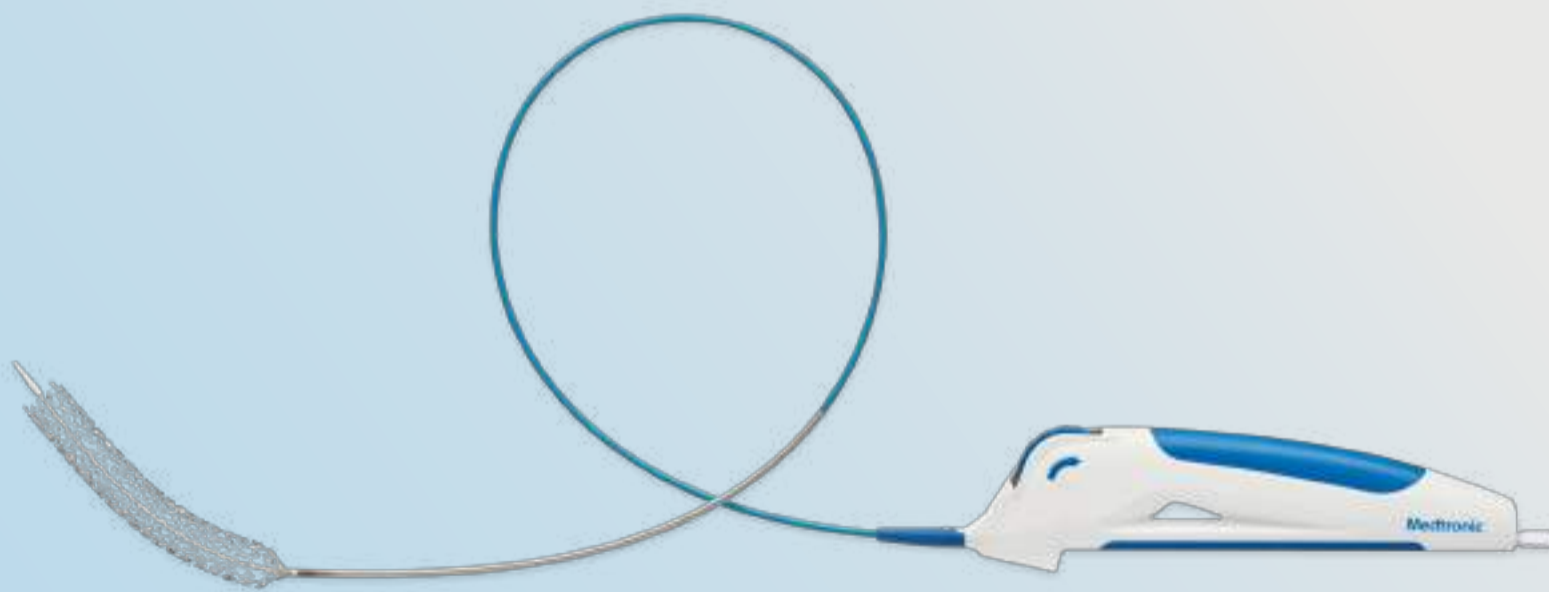
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# ABRE™ VENOUS STENT

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# Abre™

## Venous Self-expanding Stent System

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The stent made for the unique challenges of deep venous disease.

### **The Abre™ System size difference**

- A full range of sizes specifically tailored for the iliofemoral venous profile
- A consistent 9 F delivery system across our full matrix for a simplified procedure
- Catheter length of 90 cm that supports all three primary access sites and can be used with a standard length guidewire

## Venous Self-expanding Stent System

### ORDER INFORMATION

Product Catalogue Number	Stent Diameter (mm)	Stent Length (mm)	Catheter Length (cm)	Estimated Anatomic Vessel Diameter (mm)
AB9G10040090	10	40	90	7.5-9.5
AB9G10060090	10	60	90	7.5-9.5
AB9G10080090	10	80	90	7.5-9.5
AB9G10100090	10	100	90	7.5-9.5
AB9G10120090	10	120	90	7.5-9.5
AB9G10150090	10	150	90	7.5-9.5
AB9G12060090	12	60	90	9.5-11.5
AB9G12080090	12	80	90	9.5-11.5
AB9G12100090	12	100	90	9.5-11.5
AB9G12120090	12	120	90	9.5-11.5
AB9G12150090	12	150	90	9.5-11.5
AB9G14060090	14	60	90	11.5-13.5
AB9G14080090	14	80	90	11.5-13.5
AB9G14100090	14	100	90	11.5-13.5
AB9G14120090	14	120	90	11.5-13.5
AB9G14150090	14	150	90	11.5-13.5
AB9G16060090	16	60	90	13.5-15.5
AB9G16080090	16	80	90	13.5-15.5
AB9G16100090	16	100	90	13.5-15.5
AB9G16120090	16	120	90	13.5-15.5
AB9G16150090	16	150	90	13.5-15.5
AB9G18060090	18	60	90	15.5-17.5
AB9G18080090	18	80	90	15.5-17.5
AB9G18100090	18	100	90	15.5-17.5
AB9G18120090	18	120	90	15.5-17.5
AB9G18150090	18	150	90	15.5-17.5
AB9G20060090	20	60	90	17.5-19.0
AB9G20080090	20	80	90	17.5-19.0
AB9G20100090	20	100	90	17.5-19.0
AB9G20120090	20	120	90	17.5-19.0
AB9G20150090	20	150	90	17.5-19.0

# CLOSUREFAST™ PROCEDURE

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# ClosureFast™

## Endovenous Radiofrequency Ablation (RFA) Catheter



The ClosureFast™ Catheter uses radiofrequency energy to precisely and effectively treat patients suffering from chronic venous insufficiency (CVI).

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### ORDER INFORMATION

Product Catalogue Number	Product Names	Working Length (cm)	Heating Element Length (cm)	Compatible Guidewire
CF7-7-60	7 F ClosureFast™ catheter	60	7	0.025"
CF7-7-100	7 F ClosureFast™ catheter	100	7	0.025"
CF7-3-60	7 F ClosureFast™ 3 cm catheter*	60	3	0.025"

\*ClosureRFG™ software version 4.4.0 or higher is required.

# ClosureRFS™

## Endovenous Radiofrequency Stylet for Ablation of Perforators

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The ClosureRFS™ Stylet is the only endovenous radiofrequency ablation device indicated for the treatment of incompetent perforator veins for patients suffering from chronic venous insufficiency (CVI).

# ClosureRFS™

## Endovenous Radiofrequency Stylet for Ablation of Perforators

### ORDER INFORMATION

Product Catalogue Number	Product Names	French Size (F)	Working Length (cm)	Compatible Guidewire
RFS2-6-12	ClosureRFS™ stylet	6	12	0.035"

# ClosureRFG™

## Radiofrequency Generator



The ClosureRFG™ Generator delivers radiofrequency energy to the ClosureFast™ Catheter and ClosureRFS™ Stylet with real time monitoring, automatic delivery adjustments and alarms to continuously meet the preset parameters for safe and effective treatment of Chronic Venous Insufficiency (CVI).

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# ClosureRFG™

## Radiofrequency Generator

### ORDER INFORMATION

Product Catalogue Number*	Coaxial Dilator	Voltage
RFG3	ClosureRFG™ Radiofrequency Generator	Universal (100-240 V)

Product Catalogue Number	Product Name
RM55-079-01	EU power cord for ClosureRFG™ Radiofrequency Generator
RM55-081-01	UK power cord for ClosureRFG™ Radiofrequency Generator

\*Contact your local Medtronic representative to verify which generator version is approved for use in your country.

# PROCEDURE ACCESSORIES

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# Procedure Packs

Procedure packs provide the accessory materials for the ClosureFast™ and / or ClosureRFS™ procedures while standardising procedure preparation and inventory management.



EU-CPP-7F  
ClosureFast™ Procedure Pack w/out tumescent tubing

# Procedure Packs

## ClosureFast™ and ClosureRFS™ Stylet Procedure Accessories

### ORDER INFORMATION

Product Catalogue Number	Product Names
EU-CLF-KITAC-TP	ClosureFast™ Procedure Pack with tumescent tubing
EU-CPP-7F	ClosureFast™ Procedure Pack without tumescent tubing
EU-CLF-USC	EndoVenous Procedure Pack Ultrasound Probe cover



EU-CLF-KITAC-TP  
ClosureFast™ Procedure Pack with tumescent tubing



EU-CLF-USC  
endoVenous Procedure Pack Ultrasound Probe cover  
ClosureFast™ Catheter and ClosureRFS™ Stylet not included in the procedure packs.



EU-CPP-7F  
ClosureFast™ Procedure Pack w/out tumescent tubing



# Tumescent

## Infiltration Pump

Tumescent infiltration technique can be delivered with less effort, faster and with higher patient comfort via a Tumescent Pump and infiltration kit.

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# Tumescent

## Infiltration Pump

### ORDER INFORMATION

Product Catalogue Number	Product Names
4187	Tumescent Pump w/foot pedal
1501COV	Foot Pedal for tumescent pump

Product Catalogue Number	Product Name	Quantity
6022COV	Tumescent tubing set	10 units (package)

Product Catalogue Number	Product Name	Quantity
4180	DP30 Pump w/o pedal	1

# Ultrasound

## Probe Cover

The Ultrasound probe cover ensures probe sterility while facilitating probe handling.



# Ultrasound

## Probe Cover

### ORDER INFORMATION

Product Catalogue Number	Product Name	Quantity
EU-CLF-USC	Ultrasound probe cover	10 units (package)



EU-CLF-USC  
Ultrasound Probe cover

# VENASEAL™ PROCEDURE

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# VenaSeal™

## Closure System

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The VenaSeal™ Closure System is a non-thermal, non-tumescent, non-sclerosant procedure that uses a proprietary medical adhesive to treat symptomatic venous reflux.



# VenaSeal™

## Closure System

### ORDER INFORMATION

Product Catalogue Number	Product Name	Quantity
SP-101	VenaSeal™ Closure System	5 units (package)



AORTIC

PERIPHERAL

VENOUS

# Medtronic

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[medtronic.eu](https://www.medtronic.eu)



# OBTURA™ TECHNICAL SPECIFICATIONS

Parameter	Specification
Device Size (F)	6 and 8
Device Total Length (mm)	205 ± 10
Device Effective Length(mm)	155 ± 10
Complete Degradation Period (Days)	90
Device Size Compatibility	6F device compatible with 6F & 7F Introducer Sheath 8F device compatible with 8F & 9F Introducer Sheath

## OBTURA™ ORDERING INFORMATION

Obtura™ Vascular Closure Device 6F	OBT6F
Obtura™ Vascular Closure Device 8F	OBT8F

Obtura™ is not approved & not available for sale in USA.  
Obtura™ is a registered trademark of Meril Life Sciences Pvt. Ltd.

CE  
1434

More to Life

### Meril's Global Presence

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W www.merillife.com

OBTURA™ VASCULAR CLOSURE DEVICE (M.L.S./2020)67/CE/GBAL

# Assured Closure. Delivered!



# Obtura™

Vascular Closure Device

CE

# Obtura™

Vascular Closure Device

Leading towards innovation Obtura™ Vascular Closure Device is an effective mechanical system for femoral artery puncture site closure.

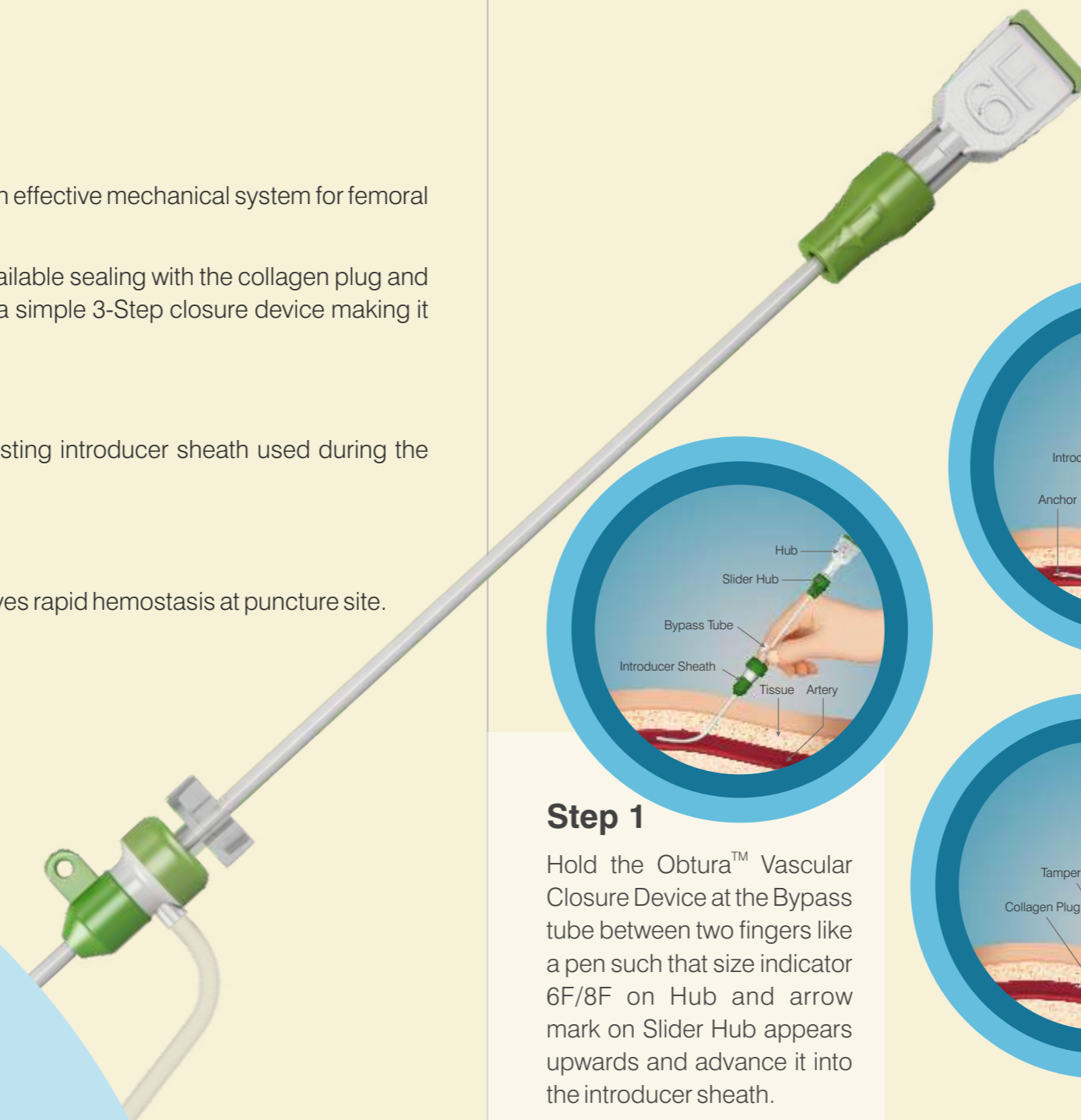
Incorporated with the Potent sealing system, it provides an unassailable sealing with the collagen plug and efficiently designed anchor. Obtura™ Vascular Closure Device is a simple 3-Step closure device making it easy to use and yielding rapid hemostasis.

## NO exchange of introducer sheath

Specially designed Obtura™ is compatible with the standard existing introducer sheath used during the interventional procedures leading to less blood loss.

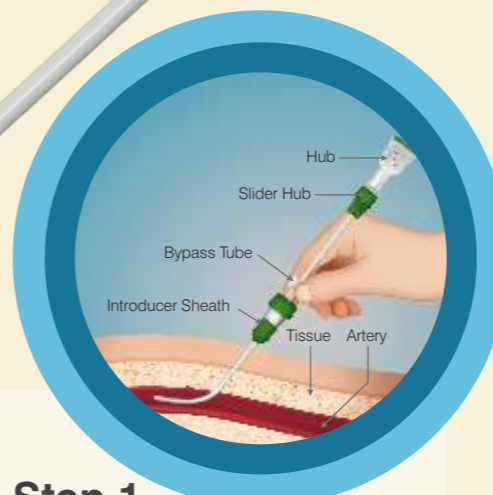
## Rapid Hemostasis

Efficient Anchor-Collagen plug sandwiches the arteriotomy and gives rapid hemostasis at puncture site.



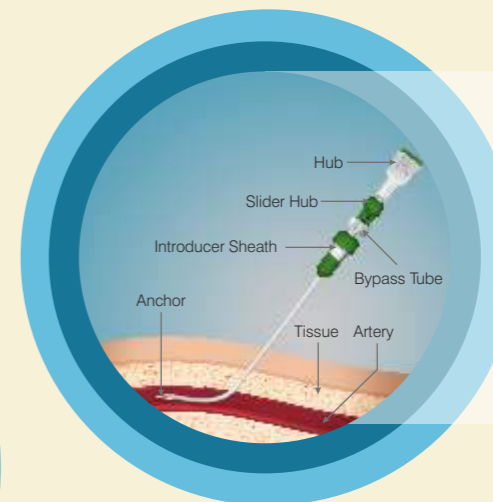
## Less procedural time

With only 3 simple steps Obtura™ saves procedural time which leads to less blood loss and early patient ambulation.



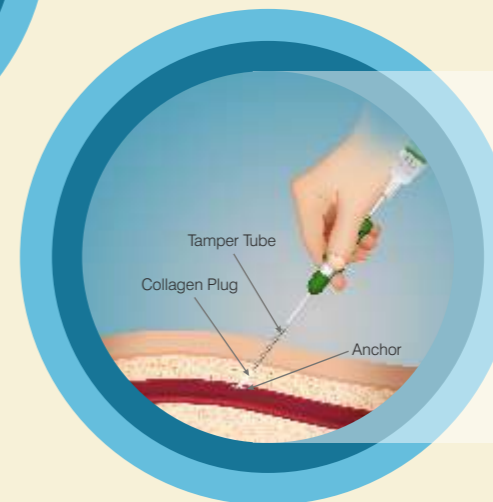
### Step 1

Hold the Obtura™ Vascular Closure Device at the Bypass tube between two fingers like a pen such that size indicator 6F/8F on Hub and arrow mark on Slider Hub appears upwards and advance it into the introducer sheath.



### Step 2

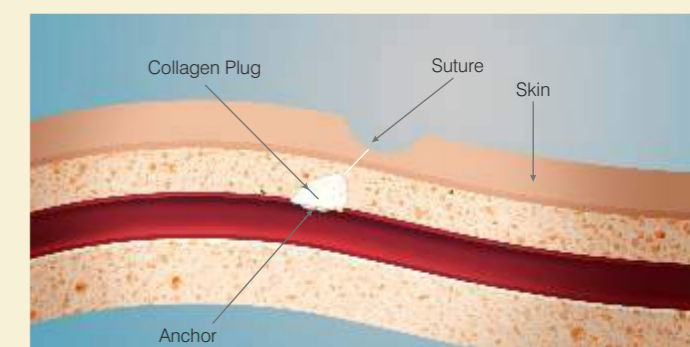
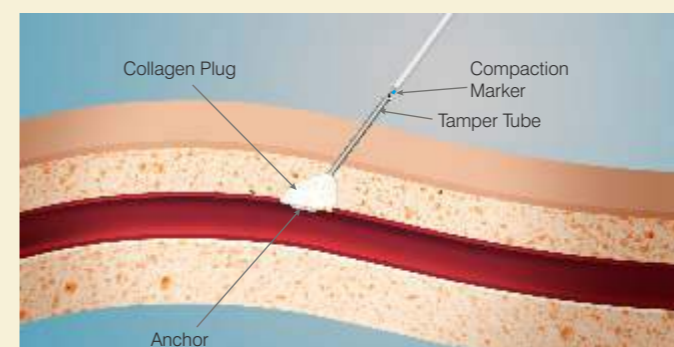
Holding the Slider Hub with one hand, pull the hub until the black marker appears.



### Step 3

Hold the three components Hub - Slider Hub - Hub of introducer sheath in the palm and retract the entire assembly along with the introducer sheath. Keep a tension on suture.

## Deployment Result



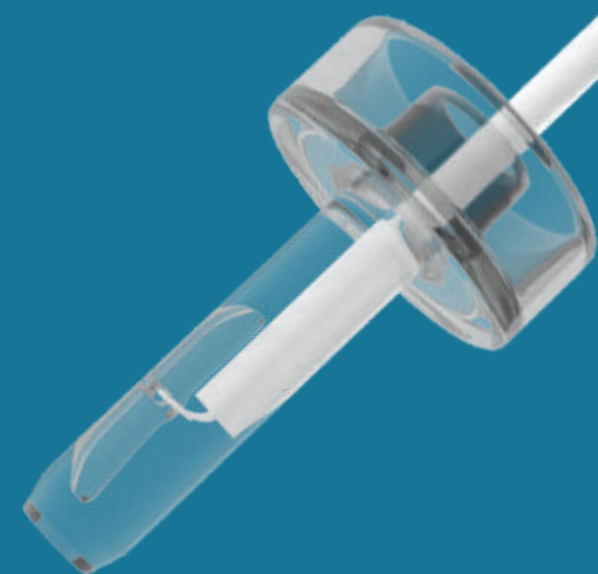
Keeping a tension on suture, push the tamper tube towards puncture site simultaneously (push-pull motion) and cut the excess suture below the locking tube. Once the anchor and collagen plug are deployed, rapid hemostasis is achieved. No blood oozing out from the site of puncture indicates a complete sealing.

## 90 days absorption

Obtura™ facilitates patient friendly sealing with the implant deployed at the puncture site which is completely absorbed within 90 days.

## Easy to use

Ergonomically designed Obtura™ is easy to use with minimal assistance and higher efficiency.



## Ghid de selecție

### Fir de ghidaj Synchro™

Cod produs	Descriere	lungime totală	Segment distal	ID proximal/distal
M00269010	Lungime activă, <b>Moale</b>	200cm	35cm	0.14x (0.30mm)
M00269110	Lungime activă, <b>Moale, Preformat</b>	200cm	35cm	0.14x (0.30mm)
M00269410	Lungime activă, <b>Standard</b>	200cm	35cm	0.14x (0.30mm)
M00269420	Lungime activă, <b>Standard, Preformat</b>	200cm	35cm	0.14x (0.30mm)
M00269510	Lungime activă, <b>Moale</b>	300cm	35cm	0.14x (0.30mm)
M00269520	Lungime activă, <b>Moale, Preformat</b>	300cm	35cm	0.14x (0.30mm)
M00269510	Lungime activă, <b>Standard</b>	300cm	35cm	0.14x (0.30mm)
M00269520	Lungime activă, <b>Standard, Preformat</b>	300cm	35cm	0.14x (0.30mm)

### Fir de ghidaj Original Synchro™-14

Cod produs	Descriere	lungime totală	Segment distal	ID proximal/distal
M00129010	Lungime activă	200cm	35cm	0.14x (0.30mm)
M00129020	Lungime activă	200cm	40cm	0.14x (0.30mm)
M00129110	Lungime activă	300cm	35cm	0.14x (0.30mm)
M00129120	Lungime activă	300cm	40cm	0.14x (0.30mm)
M00129410	Tipuri activă	200cm	35cm	0.14x (0.30mm)

### Fir de ghidaj Synchro-10

Cod produs	Descriere	lungime totală	Segment distal	ID proximal/distal
M0016010	Lungime activă	200cm	35cm	0.10x (0.25mm) / 0.20x (0.50mm)
M0016030	Lungime activă	200cm	35cm	0.10x (0.25mm) / 0.20x (0.50mm)

### Fir de ghidaj Synchro™ Neuro

A se vedea prospectul din ambalaj pentru indicații complete, contraindicații, avertismente și instrucțiuni de utilizare.

#### INDICAȚII PENTRU UTILIZARE

Seriile de fir de ghidaj Synchro Neuro este destinată utilizării neurovasculare. Pot fi utilizate pentru introducerea și poziționarea selectivă a cateterilor și a altor dispozitive intervenționale în neurovasculatură.

Acest dispozitiv trebuie utilizat de către medicii care au primit în formare și proceduri pentru utilizare, antrenament.

### ACEST DOCUMENT ESTE DESTINAT

PENTRU UTILIZAREA DE CĂTRE PROFESIONIȘTI DIN DOMENIUL SĂNĂȚII.

Un medic trebuie să se bazeze întotdeauna pe propria sa judecată clinică profesională atunci când decide dacă să utilizeze un anumit produs atunci când tratează un anumit pacient. Stryker nu poate fi considerat medicul și recomandă ca medicii să fie instruiți și să utilizeze un produs în conformitate cu instrucțiunile din prospectul produsului. Informațiile și instrucțiunile sunt disponibile în limbajul engleză și franceză pe site-ul produsului Stryker. Medicul trebuie să consulte întotdeauna prospectul produsului, eticheta produsului și / sau instrucțiunile de utilizare înainte de a folosi orice produs Stryker. Este posibil ca produsele să nu fie disponibile pe toate piețele, deoarece disponibilitatea produselor este supusă acțiunilor de reglementare și / sau medicale de pe fiecare piață în parte. Contactați reprezentantul Stryker dacă aveți întrebări legate de disponibilitatea produselor Stryker din zona dvs. Produsele Stryker enumerate mai sus sunt marcate CE în conformitate cu Directiva 93/42 / CEE privind dispozitivele medicale.



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Data emiterii: IAN/2024  
EX\_EH\_1L

**stryker**  
Neurovascular



**Synchro™**  
GUIDEWIRES

*Design avansat pentru acces*



# Synchro™

Transformate pentru acces  
GUIDEWIRES

- Proiectate pentru controlul cuplului de strângere
- Destinate pentru stabilitate și flexibilitate fiabile
- Disponibile atât cu vârf modelabil, cât și preformat
- Prezentate în opțiuni cu vârf Standard și Moale

Firul principal proximal rotund, din oțel inoxidabil, oferă adaptabilitate și stabilitate acolo unde contează.



Vârf bobină din aliaj platină-tungsten

Bobina din aliaj platină-tungsten asigură vizualizarea fluoroscopică pentru lungimea distală de 10 cm în firele de ghidaj Synchro™ și 15 cm în firele de ghidaj Synchro-14.

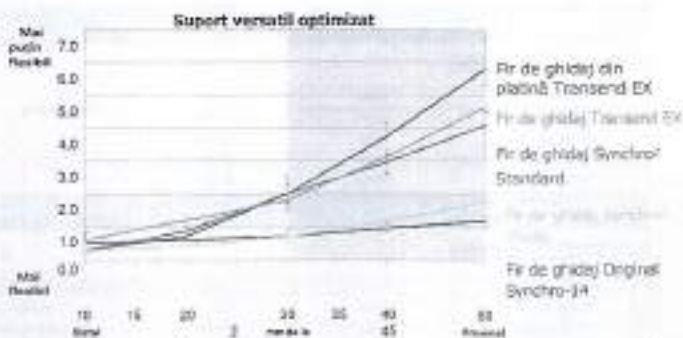
Fir principal distal din bandă plată

Îmbunătățește memoria formei la firele de ghidaj Synchro™.

## Profile pt. suport distal\*

**Fire de ghidaj moi**  
Synchro™ cu Floppy Body  
Comparabile cu firele de ghidaj originale Synchro-14

**Fire de ghidaj**  
Synchro™ cu corp standard  
Comparabile cu firele de ghidaj Transend™ EX și firele din platină Transend EX

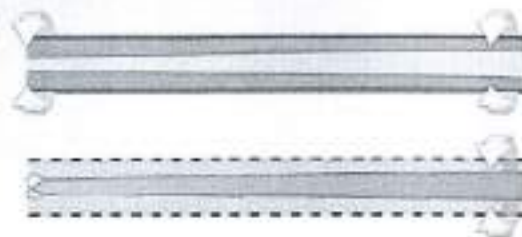


## Tehnologia recunoscută a firelor de ghidaj Synchro™

distal

proximal

Transfer cuplu de strângere



**Fire de ghidaj Synchro**

Mare parte din cuplul aplicat la capătul proximal este transmis prin structura exterioră microfabricată care se extinde de-a lungul lungimii până la vârf.

**Fire de ghidaj Convenționale**

Cuplul este transmis numai de la sfârșitul interior și se diminuează de-a lungul lungimii, pe măsură ce firul de bază se îndoaie.

**Hipotub distal microfabricat din Nitinol**

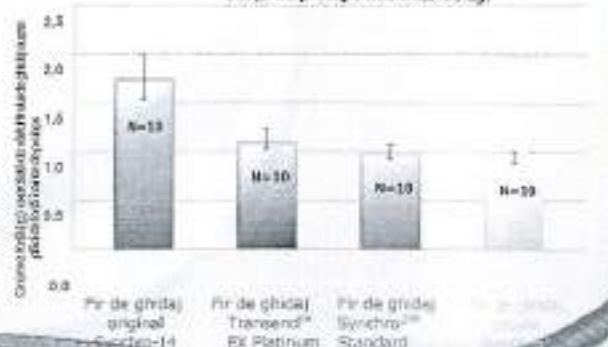
Transmite eficient cuplul de strângere de la capătul proximal la vârf distal și este proiectat pentru a oferi opțiuni de vârf standard și moale.



## Varietate de vârfuri atraumate moi

**Arbore proximal cu invelț PTFE**  
Arborele proximal prezintă un invelț PTFE proiectat pentru a îmbunătăți urmărirea și manipularea firelor de ghidaj în microscatozi.

## Forță de prolaps mai mică de 2gf



Informațiile la care s-a referit în acest document sunt de natură informativă și nu trebuie utilizate ca singura sursă de informații pentru a lua decizii medicale sau de altă natură.

TRADUCĂTOR AUTORIZAT  
ENGLEZĂ - ROMÂNĂ  
Aut. Nr. 104  
BĂDEA ANA-MARIA

\* Datele prezentate în acest document sunt de natură informativă și nu trebuie utilizate ca singura sursă de informații pentru a lua decizii medicale sau de altă natură. Informațiile la care s-a referit în acest document sunt de natură informativă și nu trebuie utilizate ca singura sursă de informații pentru a lua decizii medicale sau de altă natură.

\*\*\*\*\*

Subsemnata, **Ana-Maria Badea**, interpret și traducător autorizat pentru limba engleză în temeiul Autorizației nr. 7194 din data de 22.05.2003, eliberată de Ministerul Justiției din România, certific exactitatea traducerii efectuate din limba engleză în limba română, că textul prezentat a fost tradus complet, fără omisiuni, și că, prin traducere, înscrisului nu i-au fost denaturate conținutul și sensul.

**INTERPRET ȘI TRADUCĂTOR AUTORIZAT**

Ana-Maria Badea



TRANSFORMING  
TOTAL  
TREATMENT

NEUROVASCULAR



## Product Catalogue

Neurovascular 2020 Edition

**Medtronic**  
Further, Together



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# HEMORRHAGIC STROKE SOLUTIONS

## ONYX™ LIQUID EMBOLIC SYSTEM



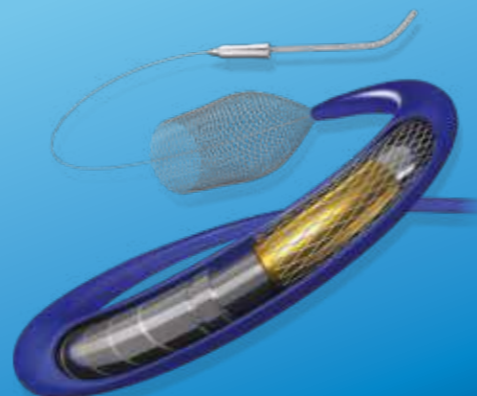
ONYX™



AXIUM™



PIPELINE™ FLEX



### Onyx™ Liquid Embolic System

Onyx™ liquid embolic system is an EVOH co-polymer designed to provide full penetration and complete packing for the embolization of vascular lesions.

PRODUCT CATALOGUE NO.	ONYX™ FORMULATION
105-7000-060	Onyx™ 18
105-7000-065	Onyx™ 20
105-7000-080	Onyx™ 34

INDICATIONS: Embolization of lesions in the peripheral and neurovasculature, including arteriovenous malformations and hypervascular tumors.

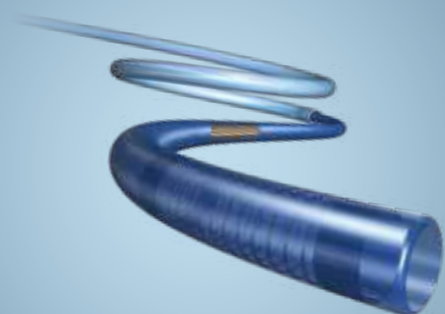
### Onyx™ Liquid Embolic System Syringe Catheter Interface Adapter

The micro therapeutics' Onyx™ liquid embolic system syringe - catheter interface adapter is intended for use as an accessory to the Onyx™ liquid embolic system and the Marathon™/Apollo™ during Onyx™ liquid embolic system injection for the embolization of brain arteriovenous malformations.

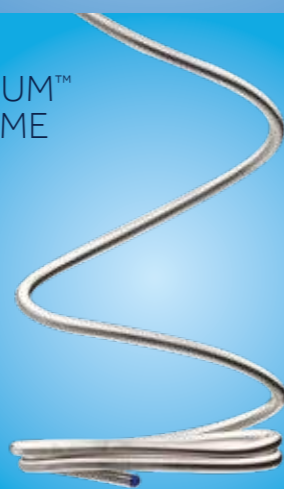
PRODUCT CATALOGUE NO.
103-1207

INDICATIONS: The proximal end of the adapter incorporates a standard ISO, female luer design to facilitate connection to the syringe. The distal end is designed specifically to fit the hub of Marathon™ and Apollo™ micro catheter 1.5F.

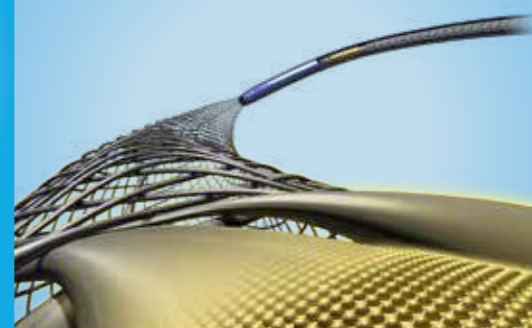
PHENOM™ MICRO  
CATHETER



AXIUM™  
PRIME



PIPELINE™ FLEX  
WITH SHIELD  
TECHNOLOGY™



### Onyx™ Liquid Embolic System Mixer

The Onyx™ liquid embolic system mixer (shaker) is packaged one unit per box. It contains four spaces for preparation of Onyx™ liquid embolic system vials simultaneously.

PRODUCT CATALOGUE NO.	VOLTAGE
103-1205-002	240

Onyx™ liquid embolic system mixer is an accessory to the Onyx™ liquid embolic system that allows proper suspension of the Onyx™ liquid embolic system tantalum for better visualization prior to use.

### Echelon™ Syringe Adapter (ESA)

The Echelon™ syringe adapter was created to address an immediate need of physicians for an additional fully compatible micro catheter option for use with the Onyx™ liquid embolic system. The ESA establishes complete compatibility between Onyx™ liquid embolic system and Echelon™ micro catheter by:

- 1) Reducing dead space in the catheter hub, which prevents Onyx™ liquid embolic system and DMSO mixing in the hub.
- 2) Simplifying the preparation process. By increasing the compatibility in these ways, the ESA significantly improves the visualization of Onyx™ liquid embolic system under fluoroscopy.

PRODUCT CATALOGUE NO.	UNITS
103-5095	20

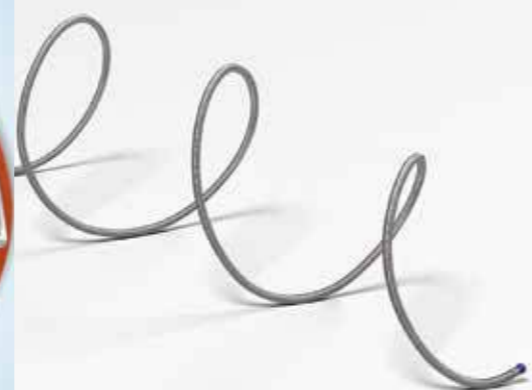
SOLITAIRE™  
AB



AXIUM™ FC



AXIUM™ ES





# AXIUM™ DETACHABLE COIL SYSTEMS



# AXIUM™ PRIME DETACHABLE COILS



The Axium™ detachable coil system provides an elegant solution that addresses challenges by optimising coil delivery, deployment, and detachment. Its progressive coil diameter system vastly refines your coil selection process.

The Axium™ Prime detachable coil system provides balanced softness through the progressive coil diameter concept. Axium™ Prime soft and extra soft coils are optimal for small and amorphic aneurysms as well as for filling small spaces during the coiling procedure.

## Axium™ 3D

PRODUCT CATALOGUE NO.	DIA. (MM)	LENGTH (CM)	COIL O.D. (IN)	O.D. (MM)	VOL. (MM)
QC-2-2-3D	2	2	0.0115	0.2921	1.34
QC-2-4-3D	2	4	0.0115	0.2921	2.68
QC-2-6-3D	2	6	0.0115	0.2921	4.02
QC-2.5-2-3D	2.5	2	0.0115	0.2921	1.34
QC-2.5-4-3D	2.5	4	0.0115	0.2921	2.68
QC-2.5-6-3D	2.5	6	0.0115	0.2921	4.02
QC-2.5-8-3D	2.5	8	0.0115	0.2921	5.36
QC-3-4-3D	3	4	0.0115	0.2921	2.68
QC-3.5-6-3D	3.5	6	0.0115	0.2921	4.02
QC-3.5-12-3D	3.5	12	0.0115	0.2921	8.04
QC-3.5-15-3D	3.5	15	0.0115	0.2921	10.05
QC-3-6-3D	3	6	0.0115	0.2921	4.02
QC-3-8-3D	3	8	0.0115	0.2921	5.36
QC-3-10-3D	3	10	0.0115	0.2921	6.7
QC-4-6-3D	4	6	0.0125	0.3175	4.75
QC-4-8-3D	4	8	0.0125	0.3175	6.33
QC-4-10-3D	4	10	0.0125	0.3175	7.92
QC-4-12-3D	4	12	0.0125	0.3175	9.5
QC-5-8-3D	5	8	0.0125	0.3175	6.33
QC-5-10-3D	5	10	0.0125	0.3175	7.92
QC-5-15-3D	5	15	0.0125	0.3175	11.88
QC-6-10-3D	6	10	0.0125	0.3175	7.92

PRODUCT CATALOGUE NO.	DIA. (MM)	LENGTH (CM)	COIL O.D. (IN)	O.D. (MM)	VOL. (MM)
QC-6-15-3D	6	15	0.0125	0.3175	11.88
QC-6-20-3D	6	20	0.0125	0.3175	15.83
QC-7-15-3D	7	15	0.0135	0.3429	13.85
QC-7-20-3D	7	20	0.0135	0.3429	18.47
QC-7-30-3D	7	30	0.0135	0.3429	27.7
QC-8-15-3D	8	15	0.0135	0.3429	13.85
QC-8-20-3D	8	20	0.0135	0.3429	18.47
QC-8-30-3D	8	30	0.0135	0.3429	27.7
QC-9-20-3D	9	20	0.0135	0.3429	18.47
QC-9-30-3D	9	30	0.0135	0.3429	27.7
QC-10-20-3D	10	20	0.0135	0.3429	18.47
QC-10-30-3D	10	30	0.0135	0.3429	27.7
QC-12-30-3D	12	30	0.0145	0.3683	31.96
QC-12-40-3D	12	40	0.0145	0.3683	42.61
QC-14-30-3D	14	30	0.0145	0.3683	31.96
QC-14-40-3D	14	40	0.0145	0.3683	42.61
QC-16-40-3D	16	40	0.0145	0.3683	42.61
QC-18-40-3D	18	40	0.0145	0.3683	42.61
QC-20-50-3D	20	50	0.0145	0.3683	53.27
QC-22-50-3D	22	50	0.0145	0.3683	53.27
QC-25-50-3D	25	50	0.0145	0.3683	53.27

## Axium™ Helix

PRODUCT CATALOGUE NO.	DIA. (MM)	LENGTH (CM)	COIL O.D. (IN)	O.D. (MM)	VOL. (MM)
QC-1.5-1-HELIX	1.5	1	0.0115	0.2921	0.67
QC-1.5-2-HELIX	1.5	2	0.0115	0.2921	1.34
QC-1.5-3-HELIX	1.5	3	0.0115	0.2921	2.01
QC-1.5-4-HELIX	1.5	4	0.0115	0.2921	2.68
QC-2-1-HELIX	2	1	0.0115	0.2921	0.67
QC-2-2-HELIX	2	2	0.0115	0.2921	1.34
QC-2-3-HELIX	2	3	0.0115	0.2921	2.01
QC-2-4-HELIX	2	4	0.0115	0.2921	2.68
QC-2-6-HELIX	2	6	0.0115	0.2921	4.02
QC-2-8-HELIX	2	8	0.0115	0.2921	5.36
QC-2.5-2-HELIX	2.5	2	0.0115	0.2921	1.34
QC-2.5-4-HELIX	2.5	4	0.0115	0.2921	2.68
QC-2.5-6-HELIX	2.5	6	0.0115	0.2921	4.02
QC-2.5-8-HELIX	2.5	8	0.0115	0.2921	5.36
QC-3-4-HELIX	3	4	0.0115	0.2921	2.68
QC-3-6-HELIX	3	6	0.0115	0.2921	4.02
QC-3-8-HELIX	3	8	0.0115	0.2921	5.36
QC-4-8-HELIX	4	8	0.0125	0.3175	6.33
QC-4-10-HELIX	4	10	0.0125	0.3175	7.92
QC-4-12-HELIX	4	12	0.0125	0.3175	9.5

PRODUCT CATALOGUE NO.	DIA. (MM)	LENGTH (CM)	COIL O.D. (IN)	O.D. (MM)	VOL. (MM)
QC-5-15-HELIX	5	15	0.0125	0.3175	11.88
QC-5-20-HELIX	5	20	0.0125	0.3175	15.83
QC-6-20-HELIX	6	20	0.0125	0.3175	15.83
QC-7-20-HELIX	7	20	0.0135	0.3429	18.47
QC-7-30-HELIX	7	30	0.0135	0.3429	27.7
QC-8-20-HELIX	8	20	0.0135	0.3429	18.47
QC-8-30-HELIX	8	30	0.0135	0.3429	27.7
QC-9-20-HELIX	9	20	0.0135	0.3429	18.47
QC-9-30-HELIX	9	30	0.0135	0.3429	27.7
QC-10-20-HELIX	10	20	0.0135	0.3429	18.47
QC-10-30-HELIX	10	30	0.0135	0.3429	27.7
QC-12-30-HELIX	12	30	0.0145	0.3683	31.96
QC-12-40-HELIX	12	40	0.0145	0.3683	42.61
QC-14-30-HELIX	14	30	0.0145	0.3683	31.96
QC-14-40-HELIX	14	40	0.0145	0.3683	42.61
QC-16-30-HELIX	16	30	0.0145	0.3683	31.96
QC-16-40-HELIX	16	40	0.0145	0.3683	42.61
QC-18-40-HELIX	18	40	0.0145	0.3683	42.61
QC-20-40-HELIX	20	40	0.0145	0.3683	42.61
QC-20-50-HELIX	20	50	0.0145	0.3683	53.27

INDICATIONS: Axium™ detachable coil systems are indicated for the endovascular embolization of intracranial aneurysms and other neurovascular abnormalities, such as arteriovenous malformations and arteriovenous fistulae.

## Axium™ Prime Super Soft 3D

PRODUCT CATALOGUE NO.	DIA. (MM)	LENGTH (CM)	COIL O.D. (IN)	O.D. (MM)	VOL. (MM)
APB-4-6-3D-SS	4	6	0.0115	0.2921	4.02
APB-4-8-3D-SS	4	8	0.0115	0.2921	5.36
APB-4-10-3D-SS	4	10	0.0115	0.2921	6.7
APB-4-12-3D-SS	4	12	0.0115	0.2921	8.04
APB-5-8-3D-SS	5	8	0.0115	0.2921	5.36
APB-5-10-3D-SS	5	10	0.0115	0.2921	6.7
APB-5-15-3D-SS	5	15	0.0115	0.2921	10.05
APB-6-10-3D-SS	6	10	0.0115	0.2921	6.7
APB-6-15-3D-SS	6	15	0.0115	0.2921	10.05
APB-6-20-3D-SS	6	20	0.0115	0.2921	13.4

## Axium™ Prime Super Soft Helix

PRODUCT CATALOGUE NO.	DIA. (MM)	LENGTH (CM)	COIL O.D. (IN)	O.D. (MM)	VOL. (MM)
APB-4-6-HX-SS	4	6	0.0115	0.2921	4.02
APB-4-8-HX-SS	4	8	0.0115	0.2921	5.36
APB-4-10-HX-SS	4	10	0.0115	0.2921	6.7
APB-4-12-HX-SS	4	12	0.0115	0.2921	8.04
APB-5-10-HX-SS	5	10	0.0115	0.2921	6.7
APB-5-15-HX-SS	5	15	0.0115	0.2921	10.05
APB-5-20-HX-SS	5	20	0.0115	0.2921	13.4
APB-6-12-HX-SS	6	12	0.0115	0.2921	8.04
APB-6-20-HX-SS	6	20	0.0115	0.2921	13.4

## Axium™ Prime Extra Soft 3D

PRODUCT CATALOGUE NO.	DIA. (MM)	LENGTH (CM)	COIL O.D. (IN)	O.D. (MM)	VOL. (MM)
APB-1-2-3D-ES	1	2	0.0108	0.2743	1.34
APB-1-3-3D-ES	1	3	0.0108	0.2743	2.01
APB-1-4-3D-ES	1	4	0.0108	0.2743	2.68
APB-1.5-2-3D-ES	1.5	2	0.0108	0.2743	1.34
APB-1.5-3-3D-ES	1.5	3	0.0108	0.2743	2.01
APB-1.5-4-3D-ES	1.5	4	0.0108	0.2743	2.68
APB-2-2-3D-ES	2	2	0.0108	0.2743	1.34
APB-2-3-3D-ES	2	3	0.0108	0.2743	2.01
APB-2-4-3D-ES	2	4	0.0108	0.2743	2.68
APB-2.5-4-3D-ES	2.5	4	0.0108	0.2743	2.68
APB-2.5-6-3D-ES	2.5	6	0.0108	0.2743	4.02
APB-3-4-3D-ES	3	4	0.0108	0.2743	2.68
APB-3-6-3D-ES	3	6	0.0108	0.2743	4.02
APB-3-8-3D-ES	3	8	0.0108	0.2743	5.36
APB-3.5-6-3D-ES	3.5	6	0.0108	0.2743	4.02
APB-3.5-8-3D-ES	3.5	8	0.0108	0.2743	5.36
APB-3.5-10-3D-ES	3.5	10	0.0108	0.2743	6.7

INDICATIONS: Axium™ and Axium™ Prime detachable coil systems are indicated for the endovascular embolization of intracranial aneurysms and other neurovascular abnormalities, such as arteriovenous malformations and arteriovenous fistulae.

## Axium™ Prime Extra Soft Helix

PRODUCT CATALOGUE NO.	DIA. (MM)	LENGTH (CM)	COIL O.D. (IN)	O.D. (MM)	VOL. (MM)
APB-1-1-HX-ES	1	1	0.0108	0.2743	0.67
APB-1-2-HX-ES	1	2	0.0108	0.2743	1.34
APB-1-3-HX-ES	1	3	0.0108	0.2743	2.01
APB-1.5-1-HX-ES	1.5	1	0.0108	0.2743	0.67
APB-1.5-2-HX-ES	1.5	2	0.0108	0.2743	1.34
APB-1.5-3-HX-ES	1.5	3	0.0108	0.2743	2.01
APB-1.5-4-HX-ES	1.5	4	0.0108	0.2743	2.68
APB-2-1-HX-ES	2	1	0.0108	0.2743	0.67
APB-2-2-HX-ES	2	2	0.0108	0.2743	1.34
APB-2-3-HX-ES	2	3	0.0108	0.2743	2.01
APB-2-4-HX-ES	2	4	0.0108	0.2743	2.68
APB-2-6-HX-ES	2	6	0.0108	0.2743	4.02
APB-2-8-HX-ES	2	8	0.0108	0.2743	5.36
APB-2.5-3-HX-ES	2.5	3	0.0108	0.2743	2.01
APB-2.5-4-HX-ES	2.5	4	0.0108	0.2743	2.68
APB-2.5-6-HX-ES	2.5	6	0.0108	0.2743	4.02
APB-2.5-8-HX-ES	2.5	8	0.0108	0.2743	5.36
APB-3-4-HX-ES	3	4	0.0108	0.2743	2.68
APB-3-6-HX-ES	3	6	0.0108	0.2743	4.02
APB-3-8-HX-ES	3	8	0.0108	0.2743	5.36
APB-3-10-HX-ES	3	10	0.0108	0.2743	6.7

## I.D. Instant Detacher

One detacher required per procedure.

PRODUCT CATALOGUE NO.	NUMBER BY BOX
ID-1-5	5

INDICATIONS: The I.D instant detacher is intended for use with all versions of the Axium™ detachable coil systems: Axium™, Axium™ Prime and Axium™ MicroFX™.

# AXIUM™ PRIME DETACHABLE FRAMING COILS



# AXIUM™ MICROFX™ DETACHABLE COIL SYSTEMS



## Axium™ Prime Frame 3D

PRODUCT CATALOGUE NO.	DIA. (MM)	LENGTH (CM)	O.D. (MM)	VOL. (MM)
FC-3-6-3D	3	6	0.0115 (0.29mm)	4.02
FC-3-8-3D	3	8	0.0115 (0.29mm)	5.36
FC-3-10-3D	3	10	0.0115 (0.29mm)	6.70
FC-3.5-6-3D	3.5	6	0.0115 (0.29mm)	4.02
FC-3.5-8-3D	3.5	8	0.0115 (0.29mm)	5.36
FC-3.5-10-3D	3.5	10	0.0115 (0.29mm)	6.70
FC-4-6-3D	4	6	0.0125 (0.32mm)	4.75
FC-4-8-3D	4	8	0.0125 (0.32mm)	6.33
FC-4-10-3D	4	10	0.0125 (0.32mm)	7.92
FC-4-12-3D	4	12	0.0125 (0.32mm)	9.50
FC-4-15-3D	4	15	0.0125 (0.32mm)	11.88
FC-5-8-3D	5	8	0.0125 (0.32mm)	6.33
FC-5-10-3D	5	10	0.0125 (0.32mm)	7.92
FC-5-15-3D	5	15	0.0125 (0.32mm)	11.88
FC-5-20-3D	5	20	0.0125 (0.32mm)	15.83
FC-6-10-3D	6	10	0.0125 (0.32mm)	7.92
FC-6-15-3D	6	15	0.0125 (0.32mm)	11.88
FC-6-20-3D	6	20	0.0125 (0.32mm)	15.83
FC-6-25-3D	6	25	0.0125 (0.32mm)	19.79
FC-7-12-3D	7	12	0.0135 (0.34mm)	11.08
FC-7-15-3D	7	15	0.0135 (0.34mm)	13.85
FC-7-20-3D	7	20	0.0135 (0.34mm)	18.47
FC-7-30-3D	7	30	0.0135 (0.34mm)	27.70
FC-8-15-3D	8	15	0.0135 (0.34mm)	13.85
FC-8-20-3D	8	20	0.0135 (0.34mm)	18.47
FC-8-30-3D	8	30	0.0135 (0.34mm)	27.70
FC-9-20-3D	9	20	0.0135 (0.34mm)	18.47
FC-9-30-3D	9	30	0.0135 (0.34mm)	27.70
FC-10-20-3D	10	20	0.0135 (0.34mm)	18.47
FC-10-30-3D	10	30	0.0135 (0.34mm)	27.70
FC-10-40-3D	10	40	0.0135 (0.34mm)	36.94
FC-12-30-3D	12	30	0.0145 (0.37mm)	31.96
FC-12-40-3D	12	40	0.0145 (0.37mm)	42.61
FC-12-50-3D	12	50	0.0145 (0.37mm)	53.27
FC-14-30-3D	14	30	0.0145 (0.37mm)	31.96
FC-14-40-3D	14	40	0.0145 (0.37mm)	42.61
FC-14-50-3D	14	50	0.0145 (0.37mm)	53.27
FC-16-40-3D	16	40	0.0145 (0.37mm)	42.61
FC-16-50-3D	16	50	0.0145 (0.37mm)	53.27
FC-18-40-3D	18	40	0.0145 (0.37mm)	42.61
FC-18-50-3D	18	50	0.0145 (0.37mm)	53.27
FC-20-50-3D	20	50	0.0145 (0.37mm)	53.27
FC-22-50-3D	22	50	0.0145 (0.37mm)	53.27
FC-25-50-3D	25	50	0.0145 (0.37mm)	53.27

INDICATIONS: The Axium™ and Axium™ Prime detachable coils are not intended for all patients and may not be the appropriate treatment for all clinical scenarios. See Instructions for Use for a complete list of warnings, precautions, and contraindications. The Axium™ Prime detachable coil system is indicated for the endovascular embolization of intracranial aneurysms and other neurovascular abnormalities, such as arteriovenous malformations and arteriovenous fistulae. The Axium™ Prime detachable coils are also indicated for arterial and venous embolizations in the peripheral vasculature.

The Axium™ detachable coil system utilizes an enlaced microfilament technology called LatticeFX™ and provides an ideal balance of softness, stability and volume through the progressive coil diameter and a single complete coil line.

## Axium™ MicroFX™ PGLA 3D

PRODUCT CATALOGUE NO.	DIA. (MM)	LENGTH (CM)	COIL O.D. (IN)	O.D. (MM)	VOL. (MM)
PC-2-2-3D	2	2	0.0115	0.314159	1.34
PC-2-4-3D	2	4	0.0115	0.314159	2.68
PC-2-6-3D	2	6	0.0115	0.314159	4.02
PC-3-4-3D	3	4	0.0115	0.314159	2.68
PC-3-6-3D	3	6	0.0115	0.314159	4.02
PC-3-8-3D	3	8	0.0115	0.314159	5.36
PC-4-6-3D	4	6	0.0125	0.314159	4.75
PC-4-8-3D	4	8	0.0125	0.314159	6.33
PC-4-10-3D	4	10	0.0125	0.314159	7.92
PC-4-12-3D	4	12	0.0125	0.314159	9.5
PC-5-8-3D	5	8	0.0125	0.314159	6.33
PC-5-10-3D	5	10	0.0125	0.314159	7.92
PC-5-15-3D	5	15	0.0125	0.314159	11.88
PC-6-10-3D	6	10	0.0125	0.314159	7.92
PC-6-15-3D	6	15	0.0125	0.314159	11.88
PC-6-20-3D	6	20	0.0125	0.314159	15.83

PRODUCT CATALOGUE NO.	DIA. (MM)	LENGTH (CM)	COIL O.D. (IN)	O.D. (MM)	VOL. (MM)
PC-7-15-3D	7	15	0.0135	0.314159	13.85
PC-7-20-3D	7	20	0.0135	0.314159	18.47
PC-7-30-3D	7	30	0.0135	0.314159	27.7
PC-8-15-3D	8	15	0.0135	0.314159	13.85
PC-8-20-3D	8	20	0.0135	0.314159	18.47
PC-8-30-3D	8	30	0.0135	0.314159	27.7
PC-9-20-3D	9	20	0.0135	0.314159	18.47
PC-9-30-3D	9	30	0.0135	0.314159	27.7
PC-10-20-3D	10	20	0.0135	0.314159	18.47
PC-10-30-3D	10	30	0.0135	0.314159	27.7
PC-12-30-3D	12	30	0.0145	0.314159	31.96
PC-12-40-3D	12	40	0.0145	0.314159	42.61
PC-14-30-3D	14	30	0.0145	0.314159	31.96
PC-14-40-3D	14	40	0.0145	0.314159	42.61
PC-16-40-3D	16	40	0.0145	0.314159	42.61
PC-18-40-3D	18	40	0.0145	0.314159	42.61

## Axium™ MicroFX™ PGLA Helix

PRODUCT CATALOGUE NO.	DIA. (MM)	LENGTH (CM)	COIL O.D. (IN)	O.D. (MM)	VOL. (MM)
PC-2-1-HELIX	2	1	0.0115	0.314159	0.67
PC-2-2-HELIX	2	2	0.0115	0.314159	1.34
PC-2-3-HELIX	2	3	0.0115	0.314159	2.01
PC-2-4-HELIX	2	4	0.0115	0.314159	2.68
PC-2-6-HELIX	2	6	0.0115	0.314159	4.02
PC-2-8-HELIX	2	8	0.0115	0.314159	5.36
PC-3-4-HELIX	3	4	0.0115	0.314159	2.68
PC-3-6-HELIX	3	6	0.0115	0.314159	4.02
PC-3-8-HELIX	3	8	0.0115	0.314159	5.36
PC-4-8-HELIX	4	8	0.0125	0.314159	6.33
PC-4-10-HELIX	4	10	0.0125	0.314159	7.92
PC-4-12-HELIX	4	12	0.0125	0.314159	9.5

PRODUCT CATALOGUE NO.	DIA. (MM)	LENGTH (CM)	COIL O.D. (IN)	O.D. (MM)	VOL. (MM)
PC-5-15-HELIX	5	15	0.0125	0.314159	11.88
PC-5-20-HELIX	5	20	0.0125	0.314159	15.83
PC-6-20-HELIX	6	20	0.0125	0.314159	15.83
PC-7-20-HELIX	7	20	0.0135	0.314159	18.47
PC-7-30-HELIX	7	30	0.0135	0.314159	27.7
PC-8-20-HELIX	8	20	0.0135	0.314159	18.47
PC-8-30-HELIX	8	30	0.0135	0.314159	27.7
PC-9-20-HELIX	9	20	0.0135	0.314159	18.47
PC-9-30-HELIX	9	30	0.0135	0.314159	27.7
PC-10-20-HELIX	10	20	0.0135	0.314159	18.47
PC-10-30-HELIX	10	30	0.0135	0.314159	27.7

## Axium™ MicroFX™ Nylon Helix

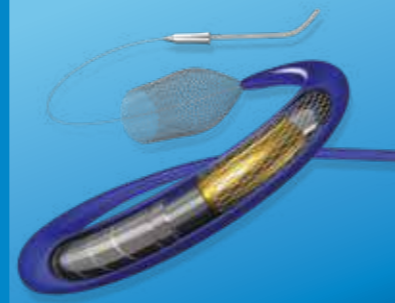
PRODUCT CATALOGUE NO.	DIA. (MM)	LENGTH (CM)	COIL O.D. (IN)	O.D. (MM)	VOL. (MM)
NC-2-1-HELIX	2	1	0.0115	0.2921	0.67
NC-2-2-HELIX	2	2	0.0115	0.2921	1.34
NC-2-3-HELIX	2	3	0.0115	0.2921	2.01
NC-2-4-HELIX	2	4	0.0115	0.2921	2.68
NC-2-6-HELIX	2	6	0.0115	0.2921	4.02
NC-2-8-HELIX	2	8	0.0115	0.2921	5.36

PRODUCT CATALOGUE NO.	DIA. (MM)	LENGTH (CM)	COIL O.D. (IN)	O.D. (MM)	VOL. (MM)
NC-3-4-HELIX	3	4	0.0115	0.2921	2.68
NC-3-6-HELIX	3	6	0.0115	0.2921	4.02
NC-3-8-HELIX	3	8	0.0115	0.2921	5.36
NC-4-8-HELIX	4	8	0.0125	0.3175	6.33
NC-4-10-HELIX	4	10	0.0125	0.3175	7.92

INDICATIONS: Axium™ and Axium™ MicroFX™ detachable coil systems are indicated for the endovascular embolization of intracranial aneurysms and other neurovascular abnormalities, such as arteriovenous malformations and arteriovenous fistulae.

# PIPELINE™ FLEX EMBOLIZATION DEVICE

PIPELINE™ IS OPTIMAL DELIVERED WITH PHENOM™



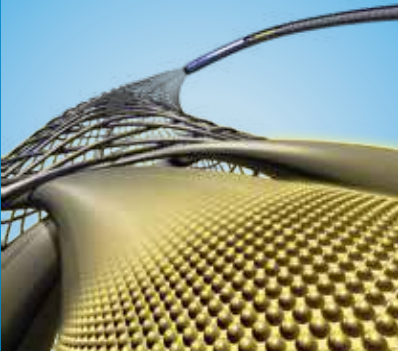
The Pipeline™ Flex embolization device redefines the scope of treatment for large, giant, wide-necked, failed-treatment, and fusiform aneurysms by reconstructing the parent artery and restoring its natural course with or without the use of adjunctive embolic devices.

PRODUCT CATALOGUE NO.	DIAMETER (MM)	LENGTH (MM)
PED-250-10	2.5	10
PED-250-12	2.5	12
PED-250-14	2.5	14
PED-250-16	2.5	16
PED-250-18	2.5	18
PED-250-20	2.5	20
PED-300-10	3	10
PED-300-12	3	12
PED-300-14	3	14
PED-300-16	3	16
PED-300-18	3	18
PED-300-20	3	20
PED-300-25	3	25
PED-300-30	3	30
PED-300-35	3	35
PED-350-12	3.5	12
PED-350-14	3.5	14
PED-350-16	3.5	16
PED-350-18	3.5	18
PED-350-20	3.5	20
PED-350-25	3.5	25
PED-350-30	3.5	30
PED-350-35	3.5	35
PED-400-10	4	10
PED-400-12	4	12
PED-400-14	4	14
PED-400-16	4	16
PED-400-18	4	18
PED-400-20	4	20
PED-400-25	4	25
PED-400-30	4	30
PED-400-35	4	35
PED-450-12	4.5	12
PED-450-14	4.5	14
PED-450-16	4.5	16
PED-450-18	4.5	18
PED-450-20	4.5	20
PED-450-25	4.5	25
PED-450-30	4.5	30
PED-450-35	4.5	35
PED-500-14	5	14
PED-500-16	5	16
PED-500-18	5	18
PED-500-20	5	20
PED-500-25	5	25
PED-500-30	5	30
PED-500-35	5	35

INDICATIONS: The Pipeline™ Flex embolization device is intended for endovascular embolization of cerebral aneurysms. The Pipeline™ Flex embolization device redefines the scope of treatment for large, giant, wide-necked, failed-treatment, and fusiform aneurysms by reconstructing the parent artery and restoring its natural course with or without the use of adjunctive embolic devices.

# PIPELINE™ FLEX EMBOLIZATION DEVICE WITH SHIELD TECHNOLOGY™

PIPELINE™ IS OPTIMAL DELIVERED WITH PHENOM™



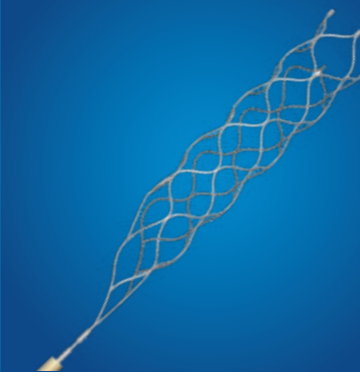
The Pipeline™ Flex embolization device with Shield Technology™ redefines the scope of treatment for large, giant, wide-necked, failed-treatment, and fusiform aneurysms by reconstructing the parent artery and restoring its natural course with or without the use of adjunctive embolic devices. Shield Technology™ reduces the material thrombogenicity of Pipeline™ device as per the Thrombogram test.

PRODUCT CATALOGUE NO.	DIAMETER (MM)	LENGTH (MM)
PED2-250-10	2.5	10
PED2-250-14	2.5	14
PED2-250-16	2.5	16
PED2-250-20	2.5	20
PED2-275-12	2.75	12
PED2-275-14	2.75	14
PED2-275-16	2.75	16
PED2-275-18	2.75	18
PED2-275-20	2.75	20
PED2-300-10	3	10
PED2-300-12	3	12
PED2-300-14	3	14
PED2-300-16	3	16
PED2-300-18	3	18
PED2-300-20	3	20
PED2-300-25	3	25
PED2-300-30	3	30
PED2-300-35	3	35
PED2-325-10	3.25	10
PED2-325-12	3.25	12
PED2-325-14	3.25	14
PED2-325-16	3.25	16
PED2-325-18	3.25	18
PED2-325-20	3.25	20
PED2-325-25	3.25	25
PED2-325-30	3.25	30
PED2-325-35	3.25	35
PED2-350-10	3.5	10
PED2-350-12	3.5	12
PED2-350-14	3.5	14
PED2-350-16	3.5	16
PED2-350-18	3.5	18
PED2-350-20	3.5	20
PED2-350-25	3.5	25
PED2-350-30	3.5	30
PED2-350-35	3.5	35
PED2-375-10	3.75	10
PED2-375-12	3.75	12
PED2-375-14	3.75	14
PED2-375-16	3.75	16
PED2-375-18	3.75	18
PED2-375-20	3.75	20
PED2-375-25	3.75	25
PED2-375-30	3.75	30
PED2-375-35	3.75	35

PRODUCT CATALOGUE NO.	DIAMETER (MM)	LENGTH (MM)
PED2-400-10	4	10
PED2-400-12	4	12
PED2-400-14	4	14
PED2-400-16	4	16
PED2-400-18	4	18
PED2-400-20	4	20
PED2-400-25	4	25
PED2-400-30	4	30
PED2-400-35	4	35
PED2-425-10	4.25	10
PED2-425-12	4.25	12
PED2-425-14	4.25	14
PED2-425-16	4.25	16
PED2-425-18	4.25	18
PED2-425-20	4.25	20
PED2-425-25	4.25	25
PED2-425-30	4.25	30
PED2-425-35	4.25	35
PED2-450-10	4.5	10
PED2-450-12	4.5	12
PED2-450-14	4.5	14
PED2-450-16	4.5	16
PED2-450-18	4.5	18
PED2-450-20	4.5	20
PED2-450-25	4.5	25
PED2-450-30	4.5	30
PED2-450-35	4.5	35
PED2-475-10	4.75	10
PED2-475-12	4.75	12
PED2-475-14	4.75	14
PED2-475-16	4.75	16
PED2-475-18	4.75	18
PED2-475-20	4.75	20
PED2-475-25	4.75	25
PED2-475-30	4.75	30
PED2-475-35	4.75	35
PED2-500-10	5	10
PED2-500-12	5	12
PED2-500-14	5	14
PED2-500-16	5	16
PED2-500-18	5	18
PED2-500-20	5	20
PED2-500-25	5	25
PED2-500-30	5	30
PED2-500-35	5	35

INDICATIONS: The Pipeline™ Flex embolization device with Shield Technology™ is intended for endovascular embolization of cerebral aneurysms.

# SOLITAIRE™ AB NEUROVASCULAR REMODELING DEVICE



The Solitaire™ AB neurovascular remodeling device is the first fully deployable and retrievable device of its kind. Solitaire™ AB is a self-expanding stent that is designed for bridging the neck of aneurysms to support the coil mass. It can be delivered and deployed by a single operator. The Solitaire™ AB neurovascular remodeling device is electrolytically detached using the Solitaire™ AB detachment system.

PRODUCT CATALOGUE NO.	VESSEL DIAMETER RANGE (MM)	DEVICE DIAMETER (MM)	MINIMUM MICROCATETER I.D. (IN)	DISTAL MARKERS	PROXIMAL MARKERS
SAB-3-20	2.2-3.0	3	0.021	3	1
SAB-3-30	2.2-3.0	3	0.021	3	1
SAB-4-15	3.0-4.0	4	0.021	3	1
SAB-4-20	3.0-4.0	4	0.021	3	1
SAB-4-30	3.0-4.0	4	0.021	3	1
SAB-4-40	3.0-4.0	4	0.021	3	1
SAB-5-20	4.0-5.0	5	0.027	4	1
SAB-5-30	4.0-5.0	5	0.027	4	1
SAB-5-40	4.0-5.0	5	0.027	4	1
SAB-6-20	5.0-6.0	6	0.027	4	1
SAB-6-30	5.0-6.0	6	0.027	4	1

INDICATIONS: The Solitaire™ AB neurovascular remodeling device is designed for use as an adjunctive device in the treatment of intracranial aneurysms.

## Vessel Diameter Sizing Chart

PRODUCT CATALOGUE NO.	USEABLE LENGTH (MM)					TOTAL LENGTH (MM)				
	VESSEL DIAMETER (MM)					VESSEL DIAMETER (MM)				
	2.2	3	4	5	6	2.2	3	4	5	6
SAB-3-20	24.2	21.7	-	-	-	32.2	31.1	-	-	-
SAB-3-30	36.6	32.1	-	-	-	44.8	41.7	-	-	-
SAB-4-15	-	17.6	15.6	-	-	-	27.7	27.3	-	-
SAB-4-20	-	22.5	20.6	-	-	-	33.1	32.1	-	-
SAB-4-30	-	33.1	31.1	-	-	-	43.5	42.3	-	-
SAB-4-40	-	44.3	40.2	-	-	-	54.2	51.6	-	-
SAB-5-20	-	-	23.2	20.1	-	-	-	33.6	32.6	-
SAB-5-30	-	-	32.4	29.1	-	-	-	42.9	41.8	-
SAB-5-40	-	-	42.1	38.3	-	-	-	52.4	50.9	-
SAB-6-20	-	-	-	19.6	17.9	-	-	-	32.7	32.3
SAB-6-30	-	-	-	30.9	28.3	-	-	-	43.9	42.8

## Solitaire™ AB Detachment System

The Solitaire™ AB detachment system is a battery operated device working with detachment cables designed to initiate and control the detachment of the Solitaire™ AB neurovascular remodeling device.

### Detachment Box

PRODUCT CATALOGUE NO.	DESCRIPTION
NDS-2X	Solitaire™ Detachment Box

### Cable Set

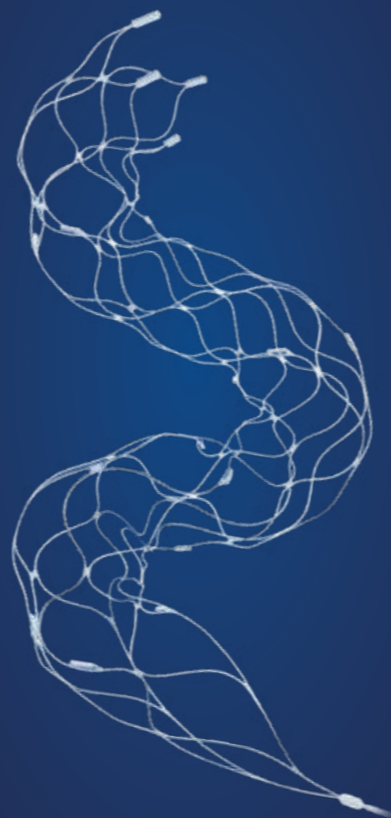
PRODUCT CATALOGUE NO.	LENGTH (M)	QUANTITY
CSS-2.75-1X	2.75	1
CSS-2.75-5X	2.75	5

# ACUTE ISCHEMIC STROKE SOLUTIONS

CELLO™ BALLOON GUIDE CATHETER



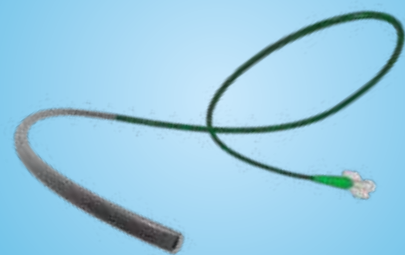
SOLITAIRE™ X REVASCULARIZATION DEVICE



FAMILY OF MICRO CATHETERS



REACT™ 68/71 FAMILY OF ASPIRATION CATHETERS



AVIGO™ GUIDEWIRE



RIPTIDE™ ASPIRATION SYSTEM



## React™ Aspiration Catheters for Acute Ischemic Stroke

The React™ 68 Catheter and the React™ 71 Catheter feature a coil and braid design along with end-to-end Nitinol construction — easing navigation to the M1 and M2 segments. Combined with the Riptide™ Aspiration System, these catheters are designed to revascularize patients experiencing acute ischemic stroke.

PRODUCT CATALOGUE NO.	ID	MAX OD	WORKING LENGTH
REACT-68	0.068"	0.083"	132 cm
REACT-71	0.071"	0.0855"	132 cm

INDICATIONS: The React™ 68 and React™ 71 Catheter is indicated for the introduction of interventional devices into the peripheral and neuro vasculature.

## Riptide™ Aspiration System

The Riptide™ Aspiration System is designed to remove occlusive thrombus from the cerebral vasculature using continuous aspiration. The catheter is connected to the Riptide™ Aspiration Pump through the aspiration tubing. The Intermediate Tubing connects the Riptide™ Collection Canister to the Riptide™ Aspiration Pump. The Riptide™ Aspiration System in this configuration may be used to aspirate thrombus from the occluded vessel.

PRODUCT NAME	CFNS	PRODUCT NAME	CFNS	VOLUME
Riptide™ Aspiration Pump	MAP-1000EU	Riptide™ Collection Canister with Intermediate Tubing	MAC-1200	1200ml

PRODUCT NAME	CFNS	INNER DIAMETER	TUBING LENGTH	DISTAL LENGTH
Riptide™ Large Bore Aspiration Tubing	MAT-110-110	1.1"	110"	7"

## Solitaire™ X Revascularization Device

The Solitaire™ X revascularization device is the latest generation of Solitaire family of devices, featuring Parametric™ design, a unique overlapping stent retriever-based technology, restores blood flow and retrieves clots from occluded blood vessels in the brain for patients experiencing acute ischemic stroke (AIS) due to a large vessel occlusion (LVO).

PRODUCT CATALOGUE NO.	REC. VESSEL DIAMETER (MM)	MINIMUM MICRO CATHETER I.D. (IN)	PUSH WIRE LENGTH (CM)	STENT DIA. (MM)	TOTAL LENGTH (MM)	PROXIMAL MARKER TO DISTAL MARKER LENGTH (MM)	DISTANCE FROM DISTAL TIP TO FLUOROSAFE MARKER (CM)	RADIOPAQUE MARKERS		RADIOPAQUE STENT MARKERS SPACING (MM)
								DIST.	PROX.	
SFR4-4-20-05	2.0 - 4.0	0.021	200	4.0	20.0	31.0	<130	3	1	5
SFR4-4-40-10	2.0 - 4.0	0.021	200	4.0	40.0	50.0	<130	3	1	10
SFR4-6-24-06	2.0 - 5.5	0.021	200	6.0	24.0	37.0	<130	4	1	6
SFR4-6-40-10	2.0 - 5.5	0.021	200	6.0	40.0	47.0	<130	4	1	10

INDICATIONS:

The Solitaire™ X Revascularization Device is indicated for use to restore blood ow in the neurovasculature by removing thrombus for the treatment of acute ischemic stroke to reduce disability in patients with a persistent, proximal anterior circulation, large vessel occlusion, and smaller core infarcts who have rst received intravenous tissue plasminogen activator (IV t-PA). Endovascular therapy with the device should be started within 6 hours of symptom onset. The Solitaire™ X Revascularization Device is indicated to restore blood ow by removing thrombus from a large intracranial vessel in patients experiencing ischemic stroke within 8 hours of symptom onset. Patients who are ineligible for IV t-PA or who fail IV t-PA therapy are candidates for treatment. The Solitaire™ X Revascularization Device is indicated for use to restore blood ow in the neurovasculature by removing thrombus for the treatment of acute ischemic stroke to reduce disability in patients with a persistent, proximal anterior circulation, large vessel occlusion of the internal carotid artery (ICA) or middle cerebral artery (MCA)-M1 segments with smaller core infarcts (<70 cc by CTA or MRA, <25 cc by MR-DWI). Endovascular therapy with the device should start within 6-16 hours of time last seen well in patients who are ineligible for intravenous tissue plasminogen activator (IV t-PA) or who fail IV t-PA therapy.

## Solitaire™ Platinum Revascularization Device

The Solitaire™ Platinum Revascularization Device is the next generation into the Solitaire™ family which has an enhanced visibility that provides feedback during the treatment of stroke patients. The Solitaire™ Platinum Revascularization Device is designed to restore blood flow in patients experiencing ischemic stroke due to large intracranial vessel occlusion. The device is designed for use in the neurovasculature such as the internal carotid artery, M1 and M2 segments of the middle cerebral artery, basilar, and the vertebral arteries.

PRODUCT CATALOGUE NO.	REC. VESSEL DIAMETER (MM)	MINIMUM MICRO CATHETER I.D. (IN)	PUSH WIRE LENGTH (CM)	STENT DIA. (MM)	TOTAL LENGTH (MM)	PROXIMAL MARKER TO DISTAL MARKER LENGTH (MM)	DISTANCE FROM DISTAL TIP TO FLUOROSAFE MARKER (CM)	RADIOPAQUE MARKERS		RADIOPAQUE STENT MARKERS SPACING (MM)
								DIST.	PROX.	
SRD3-4-20-10	2.0-4.0	0.021	180	4.0	20.0	31.0	<130	3	1	10
SRD3-4-40-10	2.0-4.0	0.021	180	4.0	40.0	50.0	<130	3	1	10
SRD3-6-20-10	3.0-5.5	0.027	180	6.0	20.0	31.0	<130	4	1	10
SRD3-6-40-10	3.0-5.5	0.027	180	6.0	40.0	47.0	<130	4	1	10

INDICATIONS: The Solitaire™ Platinum Revascularization Device is intended to restore blood flow by removing thrombus from a large intracranial vessel in patients experiencing ischemic stroke within 8 hour of symptom onset. Patients who are ineligible for intravenous tissue plasminogen activator (IV t-PA) or who fail IV t-PA therapy are candidates for treatment.

# NEURO RADIOLOGY ACCESS AND DELIVERY DEVICES

## BALLOONS



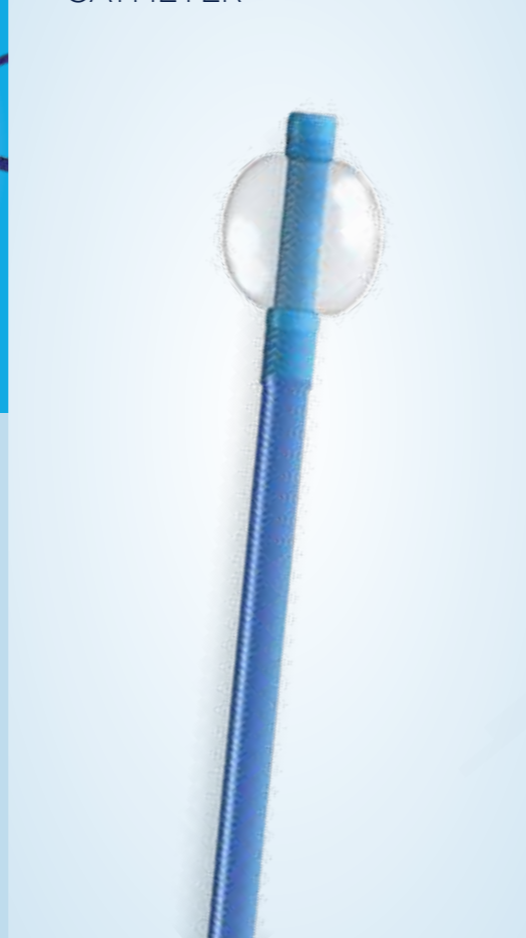
### BALLOONS



### GUIDEWIRES



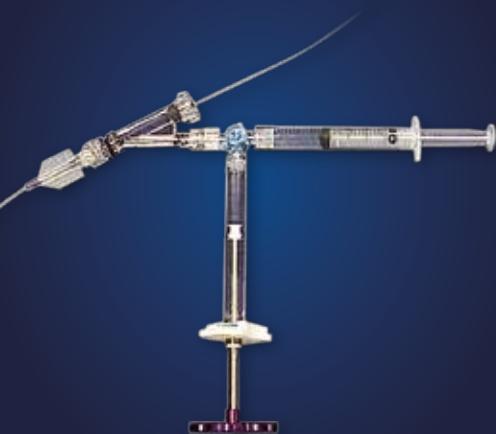
### BALLOON GUIDE CATHETER



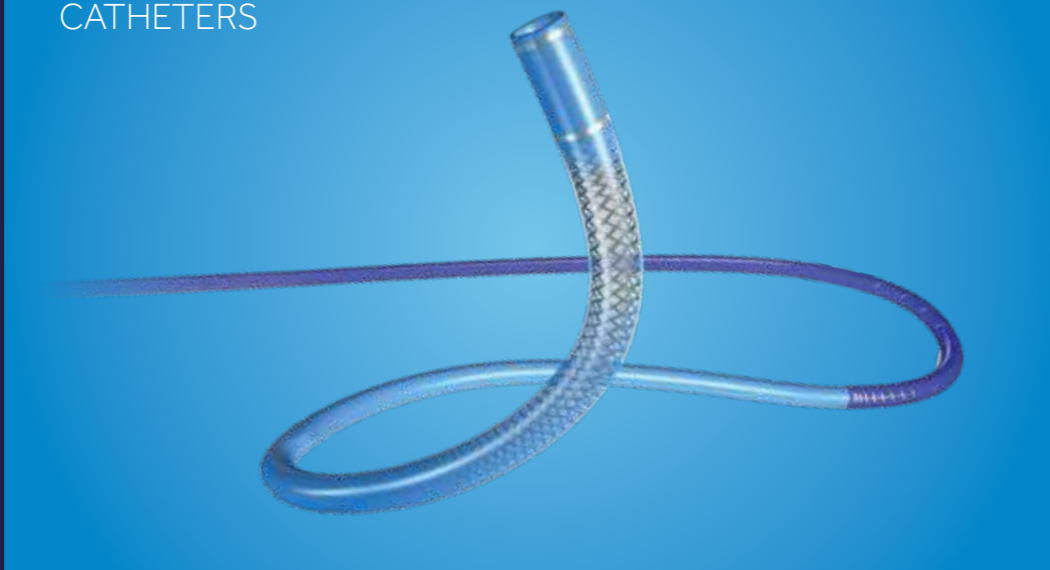
### INTRACRANIAL CATHETERS



### ACCESSORIES



### MICRO CATHETERS



A single lumen design balloon catheter with a complete line of sizes of compliant and super compliant neurovascular balloons for the optimal support of neurovascular procedures. Hyper balloons are DMSO compatible.

### HyperForm™ Occlusion Balloon System Line Extension

The HyperForm™ occlusion balloon system is a single lumen balloon catheter that requires the insertion of the 0.010" guidewire to occlude the central lumen to allow inflation of the balloon. When the distal 10 cm platinum coil tip of the guidewire is advanced to or past the catheter tip, it occludes the inflation holes allowing the balloon to inflate through catheter sideholes.

PRODUCT CATALOGUE NO.	PROXIMAL O.D. (F)	DISTAL O.D. (F)	USABLE LENGTH (CM)	BALLOON DIAMETER (MM)	BALLOON LENGTH (MM)	TIP LENGTH (MM)
104-4370	2.8	2.2	150	3	7	2
104-4153	2.8	2.2	150	3	15	2
104-4470	2.8	2.5	150	4	7	2
104-4415	2.8	2.5	150	4	15	2
104-4420	2.8	2.5	150	4	20	2
104-4770	2.8	3	150	7	7	2
104-4715	2.8	3	150	7	15	2

INDICATIONS: The HyperForm™ occlusion balloon catheter is designed for the use in the blood vessels of the peripheral and neurovasculature where temporary occlusion is desired. The HyperForm™ occlusion balloon catheter offers a vessel selective technique of temporary vascular occlusion which is useful in selectively stopping or controlling blood flow.

### HyperGlide™ Occlusion Balloon System

The HyperGlide™ occlusion balloon system is a single lumen balloon catheter that requires the insertion of the 0.010" guidewire to occlude the central lumen to allow inflation of the balloon. When the distal 10 cm platinum coil tip of the guidewire is advanced to or past the catheter tip, it occludes the inflation holes allowing the balloon to inflate through catheter sideholes. All systems packaged with an X-pedion™ hydrophilic guidewire (103-0605-200).

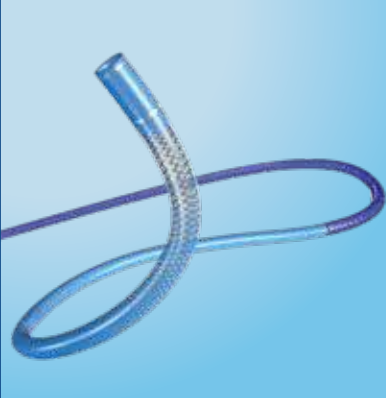
PRODUCT CATALOGUE NO.	PROXIMAL O.D. (F)	DISTAL O.D. (F)	USABLE LENGTH (CM)	BALLOON DIAMETER (MM)	BALLOON LENGTH (MM)	TIP LENGTH (MM)
104-4310	2.8	2.2	150	3	10	4
104-4315	2.8	2.2	150	3	15	4
104-4113	2.8	2.2	150	4	10	4
104-4112	2.8	2.2	150	4	15	4
104-4127	2.8	2.2	150	4	20	4
104-4132	2.8	2.2	150	4	30	4
104-4515	2.8	2.2	150	5	15	4
104-4520	2.8	2.2	150	5	20	4
104-4530	2.8	2.2	150	5	30	4

INDICATIONS: The HyperGlide™ occlusion balloon catheter is designed for the use in the blood vessels of the peripheral and neurovasculature where temporary occlusion is desired. The HyperGlide™ occlusion balloon catheter offers a vessel selective technique of temporary vascular occlusion which is useful in selectively stopping or controlling blood flow.

# GUIDEWIRES



# MICRO CATHETERS



## Avigo™ Hydrophilic Guidewire

The Avigo™ 0.014" hydrophilic guidewire offers ultimate support for tracking robust systems, crossing clot, and maintaining catheter position.

PRODUCT CATALOGUE NO.	DIAMETER (IN)	TOTAL LENGTH (CM)	COIL LENGTH (CM)	TIP SHAPE
103-0606-200	0.014	205	5	Straight

INDICATIONS: The Avigo™ hydrophilic guidewire is indicated for general intravascular use to aid in the selective placement of catheters in the peripheral and cerebral vasculature during diagnostic and/or therapeutic procedures. The device is not intended for use in the coronary arteries.

## Mirage™ 0.008" Hydrophilic Guidewire

The Mirage™ hydrophilic guidewire is a stainless steel guidewire with a radiopaque, platinum distal coil. The guidewire is hydrophilically coated on the distal portion.

PRODUCT CATALOGUE NO.	DIAMETER (IN)	TOTAL LENGTH (CM)	COIL LENGTH (CM)
103-0608	0.008	200	10

INDICATIONS: The Mirage™ hydrophilic guidewire is indicated for general intravascular use to aid in the selective placement of catheters in the peripheral, visceral and cerebral vasculature during diagnostic and/or therapeutic procedures.

## X-pedion™ Hydrophilic Guidewire

The X-pedion™ hydrophilic guidewire is a stainless steel guidewire with a radiopaque, platinum distal coil. The guidewire is hydrophilically coated on the distal portion.

PRODUCT CATALOGUE NO.	DIAMETER (IN)	TOTAL LENGTH (CM)	COIL LENGTH (CM)
103-0605-200	0.010	200	10

INDICATIONS: The X-pedion™ hydrophilic guidewire is indicated for general intravascular use to aid in the selective placement of catheters in the peripheral, visceral and cerebral vasculature during diagnostic and/or therapeutic procedures.

## Marathon™ Micro Catheter

Marathon™ is a flow directed micro catheter with a proximal stainless steel coil for great proximal support and a nitinol distal braiding for high kink resistance for optimised delivery of Onyx™ liquid embolic system.

PRODUCT CATALOGUE NO.	OUTER DIAMETER (FR)	DISTAL INNER DIAMETER (IN)	TOTAL LENGTH (CM)	USABLE LENGTH (CM)	DISTAL LENGTH (CM)	MAX. GUIDEWIRE (IN)
105-5056	2.7-1.5	0.013	170	165	25	0.01

INDICATIONS: The Marathon™ micro catheter is intended to access peripheral and neurovasculature for the controlled selective infusion of physician-specified therapeutic agents such as embolization catheter materials and diagnostic materials such as contrast media.

## Apollo™ Onyx™ Delivery Micro Catheter

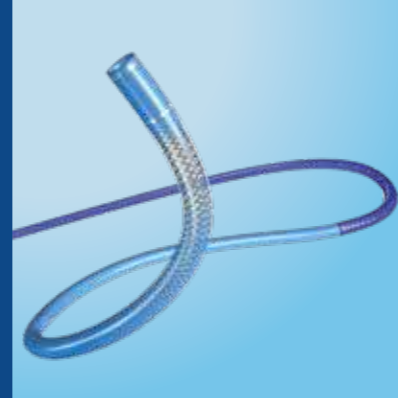
Apollo™ Onyx™ delivery micro catheter is designed with a stainless steel proximal coil for high kink resistance and distal nitinol braiding for reinforcement.

PRODUCT CATALOGUE NO.	DISTAL OUTER DIAMETER (FR)	PROXIMAL DIAMETER (FR)	INNER DIAMETER (IN)	TOTAL LENGTH (CM)	TIP LENGTH (CM)	TIP SHAPE	WIRE COMPATIBILITY (IN)
105-5095-000	1.5	2.7	0.013	165	1.5	Straight	≤ 0.01
105-5096-000	1.5	2.7	0.013	165	3	Straight	≤ 0.01
105-5097-000	1.5	2.7	0.013	165	5	Straight	≤ 0.01

INDICATIONS: Apollo™ Onyx™ delivery micro catheter is intended to access the neurovasculature for the controlled selective infusion of physician-specified therapeutic such as embolization materials and diagnostic materials such as contrast media.

A torque device to assist in guidewire manipulation and a guidewire introducer to ease the introduction of the guidewire into the catheter hub and/or hemostatic valve is provided in the sterile pouch of every guidewire.

# MICRO CATHETERS



## Echelon™ Micro Catheter

The Echelon™ micro catheter is a steam shapeable micro catheter with variable nitinol braiding and a flexible distal tip for optimal device delivery. The Echelon™ micro catheter is DMSO compatible.

PRODUCT NAME	PRODUCT CATALOGUE NO.	OUTER DIAMETER (FR)	DISTAL INNER DIAMETER (IN)	TOTAL LENGTH (CM)	USABLE LENGTH (CM)	MAX GUIDEWIRE (IN)	TIP LENGTH (MM)	TIP SHAPE
Echelon™ 10	105-5091-150	2.1-1.7	0.017	155	150	0.014	-	Straight
	145-5091-150	2.1-1.7	0.017	155	150	0.014	2.5	45°
	190-5091-150	2.1-1.7	0.017	155	150	0.014	5	90°
Echelon™ 14	105-5092-150	2.4-1.9	0.017	155	150	0.014	-	Straight
	145-5092-150	2.4-1.9	0.017	155	150	0.014	2.5	45°
	190-5092-150	2.4-1.9	0.017	155	150	0.014	5	90°

INDICATIONS: The Echelon™ micro catheter is intended to access peripheral and neurovasculature for the controlled selective infusion of physician-specified therapeutic agents such as embolization materials and of diagnostic materials such as contrast media.

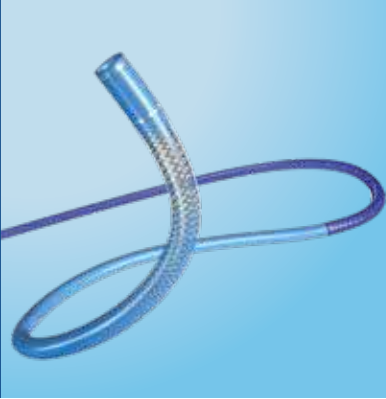
## Marksman™ Micro Catheter

All Marksman™ micro catheters are specially designed for the optimal and safe delivery of the Pipeline™ embolization device except Marksman™ 160 micro catheter specially designed for Stroke. Proximally the design is offering a stainless steel coil and a stainless steel braiding for optimised handling and support. The Marksman™ catheter has a high kink resistant stainless steel braiding for a smooth device delivery.

PRODUCT CATALOGUE NO.	OUTER DIAMETER DISTAL/PROXIMAL (FR)	INNER DIAMETER (IN)	WORKING LENGTH (CM)	DISTAL FLEXIBLE LENGTH (CM)
FA-55105-1015	2.8/3.2	0.027	105	10
FA-55135-1030	2.8/3.2	0.027	135	10
FA-55150-1030	2.8/3.2	0.027	150	10
FA-55160-1030	2.8/3.2	0.027	160	10

INDICATIONS: The Marksman™ micro catheter is intended for the introduction of interventional and infusion of diagnostic or therapeutic agents into the neuro, peripheral, and coronary vasculature.

# MICRO CATHETERS



## Rebar™ Reinforced Micro Catheter

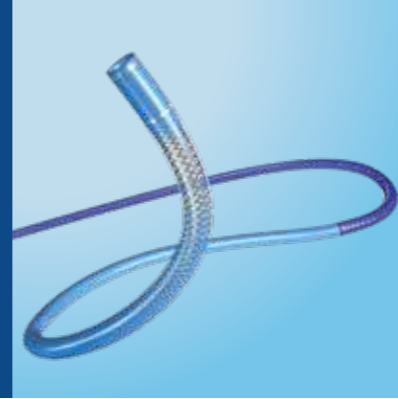
Rebar™ micro catheter is a stainless steel coil reinforced micro catheter offering high kink resistance for optimal device delivery. Rebar™ micro catheter is compatible with Onyx™ liquid embolic system.

PRODUCT NAME	PRODUCT CATALOGUE NO.	OUTER DIAMETER (FR)	DISTAL INNER DIAMETER (IN)	TOTAL LENGTH (CM)	USABLE LENGTH (CM)	MAX GUIDEWIRE (IN)	MARKER BANDS
Rebar™ 10	105-5078-153	2.3-1.7	0.015	158	153	0.012	1
	105-5078-153C	2.3-1.7	0.015	158	153	0.012	2
Rebar™ 18	105-5081-153	2.7-2.4	0.021	158	153	0.018	2
Rebar™ 27	105-5082-130	2.8-2.8	0.027	135	130	0.021	1

INDICATIONS: The Rebar™ micro catheter is intended for the controlled selective infusion of physician-specified therapeutic agents or contrast media into the vasculature of the peripheral and neuro anatomy.



# PHENOM™ FAMILY OF CATHETERS



# INTRACRANIAL CATHETERS



Phenom™ 017 Catheter: select your tip shape for optimal coil delivery

PRODUCT CATALOGUE NO.	SIZE	DISTAL O.D. (IN)	PROXIMAL O.D. (IN)	CATHETER I.D. (IN)	WORKING LENGTH (CM)	COIL LENGTH (CM)	DISTAL LENGTH (CM)	TIP SHAPE	TIP MARKERS
FG11150-0615-2S	17	0.024	0.029	0.017	150	15	6	Straight	2
FG11150-0615-2J	17	0.024	0.029	0.017	150	15	6	J Curve	2
FG11150-0615-2X	17	0.024	0.029	0.017	150	15	6	45 Curve	2
FG11150-0615-2R	17	0.024	0.029	0.017	150	15	6	90 Curve	2

Phenom™ 021 Catheter: Purposefully designed for Pipeline™

PRODUCT CATALOGUE NO.	SIZE	DISTAL O.D. (IN)	PROXIMAL O.D. (IN)	CATHETER I.D. (IN)	WORKING LENGTH (CM)	COIL LENGTH (CM)	DISTAL LENGTH (CM)	TIP SHAPE	TIP MARKERS
FG13150-0615-S2	21	0.024	0.029	0.021	150	15	6	Straight	2

Phenom™ 021 Catheter: Ultimate Delivery Platform for Solitaire™

Phenom™ 160 cm microcatheters are the ultimate delivery platform for stent deployment. The soft and rounded distal tip is designed to reduce the risk of vessel trauma. The flexibility of the distal shaft allows the catheters to track well in tortuous anatomy while maintaining stability during stent delivery. The lumen is strategically engineered to prevent ovalization and provide durability for delivering stents with minimal friction.

PRODUCT CATALOGUE NO.	SIZE	DISTAL O.D. (IN)	PROXIMAL O.D. (IN)	CATHETER I.D. (IN)	WORKING LENGTH (CM)	COIL LENGTH (CM)	DISTAL LENGTH (CM)	TIP SHAPE	TIP MARKERS
FG13160-0615-1S	21	2.3	2.6	0.021	160	15	6	Straight	1

Phenom™ 027 Catheter: Purposefully designed for Pipeline™

Phenom™ 027 microcatheter was precisely engineered for Pipeline™ Flex Embolization Device with Shield Technology, to achieve a symbiotic function. Designed to reach the aneurysm and secure access, thereby enhancing trackability and an inner lumen engineered to prevent ovalization for a fully controlled and smooth delivery. Robust proximal pushability from composite shaft with optimal flexibility for complex procedures.

PRODUCT CATALOGUE NO.	SIZE	DISTAL O.D. (IN)	PROXIMAL O.D. (IN)	CATHETER I.D. (IN)	WORKING LENGTH (CM)	COIL LENGTH (CM)	DISTAL LENGTH (CM)	TIP SHAPE	TIP MARKERS
FG15150-0615-1S	27	0.036	0.040	0.027	150	15	6	Straight	1
FG15150-0630-1S	27	0.036	0.040	0.027	150	30	6	Straight	1

Navien™ A+ Intracranial Support Catheter

Navien™ intracranial support catheter with a variable pitch nitinol coil offering a very flexible and supportive catheter with minimal ovalization for controlled device delivery. Optimal aspiration and durability for AIS interventions.

PRODUCT CATALOGUE NO.	OUTER DIAMETER (FR/IN)	INNER DIAMETER (IN)	TOTAL LENGTH (CM)	TIP SHAPE	WIRE COMPATIBILITY (IN)
RFXA058-105-08	5/0.07 max	0.058	105	Straight	0.035/0.038
RFXA058-115-08	5/0.07 max	0.058	115	Straight	0.035/0.038
RFXA058-125-08	5/0.07 max	0.058	125	Straight	0.035/0.038
RFXA058-130-08	5/0.070 max	0.058	130	Straight	0.035/0.038
RFXA072-95-08	6/0.084 max	0.072	95	Straight	0.035/0.038
RFXA072-95-08MP	6/0.084 max	0.072	95	Multi-Purpose 25°	0.035/0.038
RFXA072-105-08	6/0.084 max	0.072	105	Straight	0.035/0.038
RFXA072-115-08	6/0.084 max	0.072	115	Straight	0.035/0.038
RFXA072-125-08	6/0.084 max	0.072	125	Straight	0.035/0.038
RFXA072-105-08MP	6/0.084 max	0.072	105	Multi-Purpose 25°	0.035/0.038
RFXA072-115-08MP	6/0.084 max	0.072	115	Multi-Purpose 25°	0.035/0.038
RFXA072-125-08MP	6/0.084 max	0.072	125	Multi-Purpose 25°	0.035/0.038
RFXA072-130-08	6/0.084 max	0.072	130	Straight	0.035/0.038
RFXA072-130-08MP	6/0.084 max	0.072	130	Multi-Purpose 25°	0.035/0.038

Distal Flexible Length for all sizes = 8cm

INDICATIONS: The Navien™ intracranial catheter is indicated for the introduction of interventional/diagnostic devices into the peripheral, coronary, and neuro vasculature. The Navien™ intracranial catheter is also indicated for the removal/aspiration of fresh, soft emboli and thrombi from selected blood vessels in the arterial system, including neurovasculature.

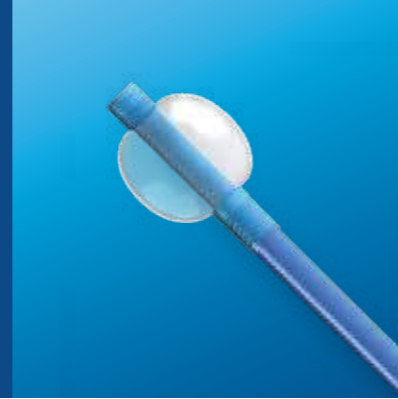
Phenom™ Plus Catheter

With features such as an elite composite shaft, proprietary inner lining, and a rounded distal tip, the PHENOM™ FAMILY OF CATHETERS are the ultimate delivery platform for coil and stent delivery. Different clinical scenarios require variations in levels of support and navigation. With Phenom™ catheters, we provide various catheter specifications to meet your needs.

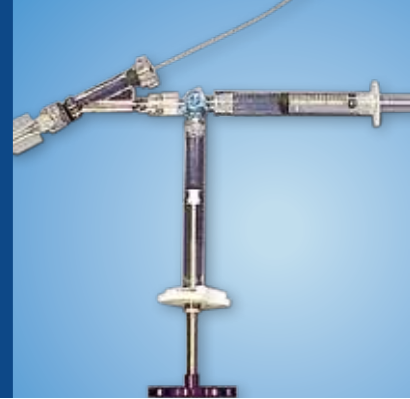
PRODUCT CATALOGUE NO.	SIZE	DISTAL O.D. (IN)	PROXIMAL O.D. (IN)	CATHETER I.D. (IN)	WORKING LENGTH (CM)	COIL LENGTH (CM)	DISTAL LENGTH (CM)	TIP SHAPE	TIP MARKERS
FG19105-0630-1S	45	0.055	0.061	0.0445	105	30	6	Straight	1
FG19120-1030-1S	45	0.055	0.061	0.0445	120	30	10	Straight	1

INDICATIONS: Phenom™ catheters are intended for the introduction of interventional devices and infusion of diagnostic or therapeutic agents into the neuro, peripheral, and coronary vasculatures.

# BALLOON GUIDE CATHETERS



# ACCESSORIES



## Cello™ Balloon Guide Catheter

The Cello™ balloon guide catheter is indicated for use in facilitating the insertion and guidance of intravascular catheters into a selected blood vessel in the peripheral and neuro vascular systems. The balloon provides temporary vascular occlusion during these and other angiographic procedures.

PRODUCT NAME	PRODUCT CATALOGUE NO.	CONFORMABLE SHEATH (F)	OUTER DIAMETER (IN)	INNER DIAMETER (IN)	TIP LENGTH (MM)	BALLOON LENGTH (MM)	EFFECTIVE LENGTH (CM)	TOTAL LENGTH (CM)
Cello 6F+	1610060	7	6F+ (0.079)	0.051	3	7	95	103
Cello 7F+	1610070	8	7F+ (0.094)	0.067	3	7	95	103
Cello 8F	1610080	8	8F (0.106)	0.075	3	10	95	103
Cello 9F	1610090	9	9F (0.114)	0.085	3	10	92	100

INDICATIONS: The Cello™ balloon guide catheter is intended to temporarily block blood flow by expanding a balloon inside blood vessels during operations, such as: urgent hemostasis, hemostasis for surgery, perfusion of blood to peripheral vessel and arterial injection for chemotherapy.

## Cadence™ Precision Injector

Cadence™ precision injector syringe with threaded plunger.

PRODUCT CATALOGUE NO.	CAPACITY (ML)	SYRINGES/BOX
103-0304	1	5

INDICATIONS: The Cadence™ precision injector is intended for the delivery of fluids.

## 1ml Luer-Lock Injection Syringe

PRODUCT CATALOGUE NO.	CAPACITY (ML)	SYRINGES/BOX
103-1203	1	10

INDICATIONS: The Luer-Lock injector syringe is intended for the delivery of fluids.



See the device manual for detailed information regarding the instructions for use, indications, contraindications, warnings, precautions, and potential adverse events. For further information, contact your local Medtronic representative and/or consult the Medtronic website at [medtronic.eu](http://medtronic.eu).

## Reference

<sup>1</sup> Cathera Document ML-0001.A Phenom is a trademark of Cathera, Inc.

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