

Unique[†] size mix with 48 mm length and 2.0 mm diameter

Ordering Information

	PRO ^x								PRO 48
Balloon ø (mm)	References by Length and Diameter								
	8 mm	12 mm	15 mm	18 mm	23 mm	28 mm	33 mm	38 mm	48 mm
2.00	1076200-08	1076200-12	1076200-15	1076200-18	1076200-23	1076200-28	—	—	—
2.25	1076225-08	1076225-12	1076225-15	1076225-18	1076225-23	1076225-28	—	—	—
2.50	1076250-08	1076250-12	1076250-15	1076250-18	1076250-23	1076250-28	1076250-33	1076250-38	1017250-48
2.75	1076275-08	1076275-12	1076275-15	1076275-18	1076275-23	1076275-28	1076275-33	1076275-38	1017275-48
3.00	1076300-08	1076300-12	1076300-15	1076300-18	1076300-23	1076300-28	1076300-33	1076300-38	1017300-48
3.50	1076350-08	1076350-12	1076350-15	1076350-18	1076350-23	1076350-28	1076350-33	1076350-38	1017350-48
4.00	1076400-08	1076400-12	1076400-15	1076400-18	1076400-23	1076400-28	1076400-33	1076400-38	—

Stent Specifications⁴

Stent Design	MULTI-LINK, 3-3-3, non-linear link	
Stent Material	L-605 Cobalt Chromium	
Drug	Everolimus	
Drug Dose	100 µg/cm ²	
Polymer	Fluorinated Copolymer	
Strut Thickness	0.0032"	
Maximum Expansion Diameter	Size (mm)	Max Expansion (mm)
	2.00–2.50	3.25
	2.75–3.00	3.75
	3.50–4.00	4.50

Stent Delivery System Specifications⁴

Nominal Pressure	10 atm (11 atm for 48 mm)		
Rated Burst Pressure	18 atm		
Shaft Measurements	Proximal	Mid-Shaft	Distal
		Min: 0.035"	Min: 0.032"
		Max: 0.028"	Max: 0.038"
GW Notch Width, Average	0.033"		
Balloon Material	Multilayer Pebax®		
Crossing Profile	0.0439" (3.0 x 48 mm)		
	0.0425" (3.0 x 18 mm)		
Tip Entry Profile	0.017"		
Working Catheter Length	145 cm		

The XIENCE product was designed as an overall system, with each system component contributing to its clinical outcomes. The system includes: the thin-strut CoCr, multi-generation MULTI-LINK, with its flexible ring and 3-link stent design; advanced delivery system; novel everolimus compound; the multi-layer coating technologies, utilizing a primer and co-polymer, previously known for cardiovascular implants and for having excellent mechanical properties.

[†] Among Major DES Brands.

References: **1.** Based on patient numbers from various Abbott and non-Abbott-sponsored trials. Data on file at Abbott Vascular. **2.** Trials registered on www.clinicaltrials.gov as of Aug. 3, 2011. **3.** Data on file at Abbott Vascular. **4.** All tests performed by and data on file at Abbott Vascular.

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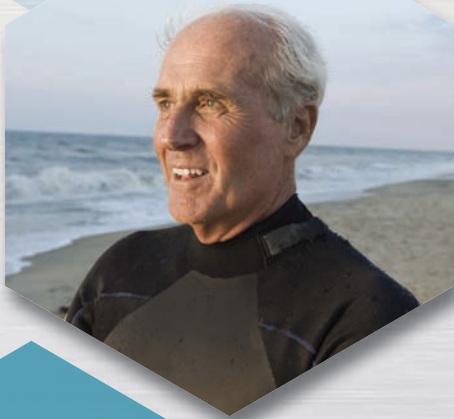
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The most deliverable
XIENCE ever.



XIENCE
Safety*

*Based on ST rates from SPIRIT II, SPIRIT III and SPIRIT IV.

The most deliverable XIENCE ever.

Pushable

Cross challenging anatomy with ease

- Smooth and reduced transitions across the entire system for optimal force transfer
- Hypotube skive transition for exceptional support

Skive

Trackable

Track smoothly through tortuous anatomy

- Fully integrated tip continues to guide wire notch with no transitions for excellent tracking
- Ultra low distal seal technology for outstanding crossability



XIENCE Safety*

More than
62,000
Patients
Studied¹

100+
RCTs and
Registries²

Over
6 million
Patients³

Flexible

Deliver with confidence

- Flexible multilayered balloon with flatter compliance for superior deliverability



*Based on ST rates from SPIRIT II, SPIRIT III and SPIRIT IV.