





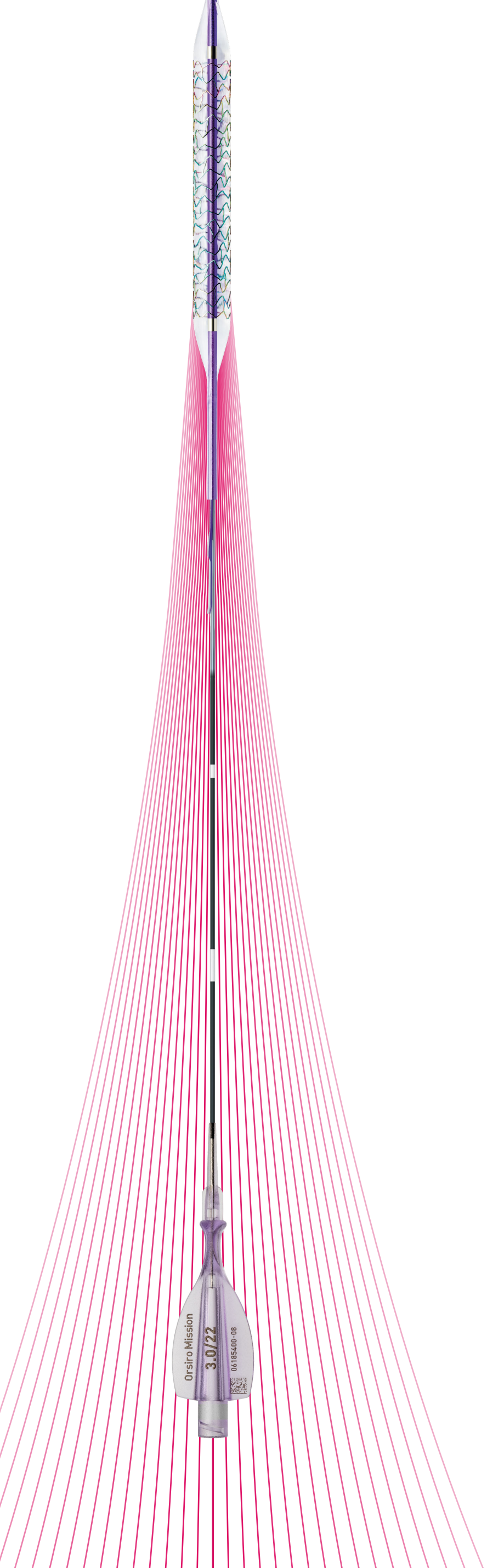


Vascular Intervention // Coronary
Drug-Eluting Stent System



Orsiro® Mission des

Even better deliverability for the outstanding Orsiro DES



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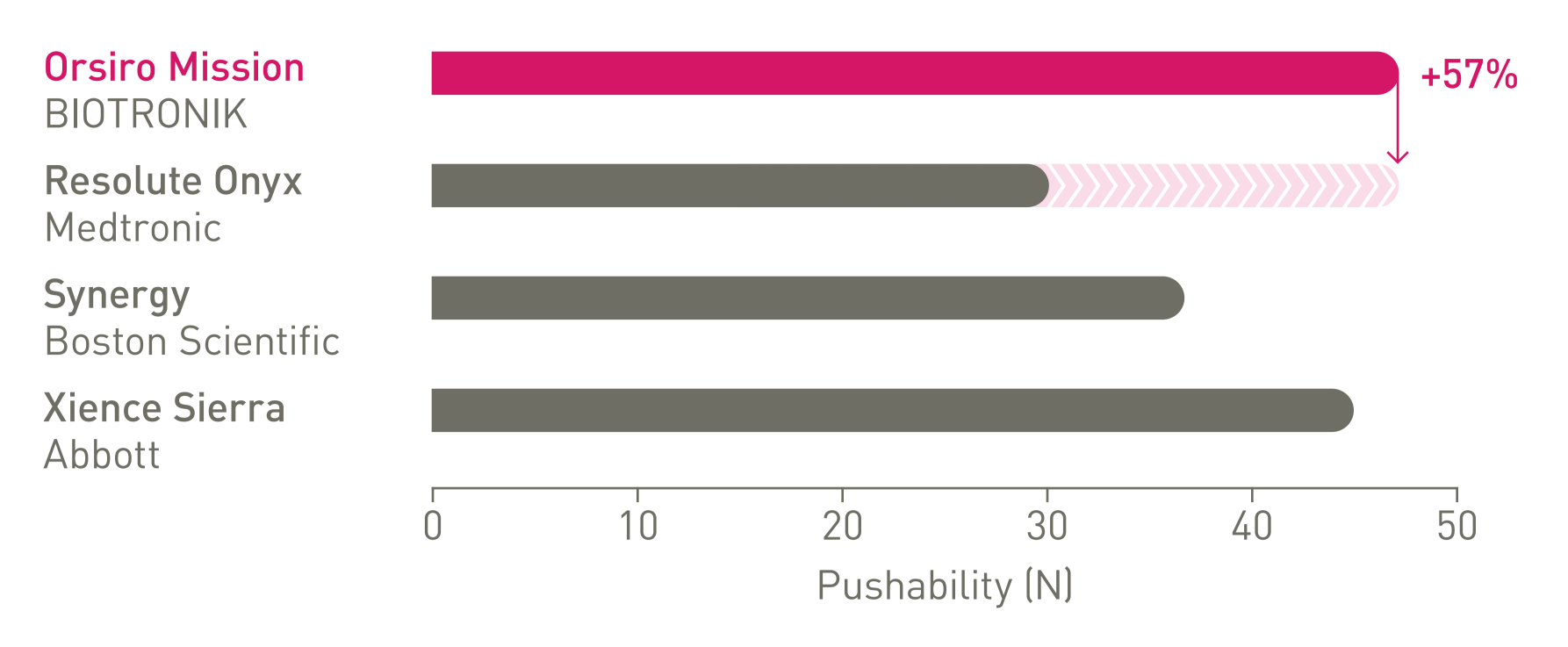
Orsiro Mission des

Even better deliverability for the outstanding Orsiro DES

The next level of deliverability¹

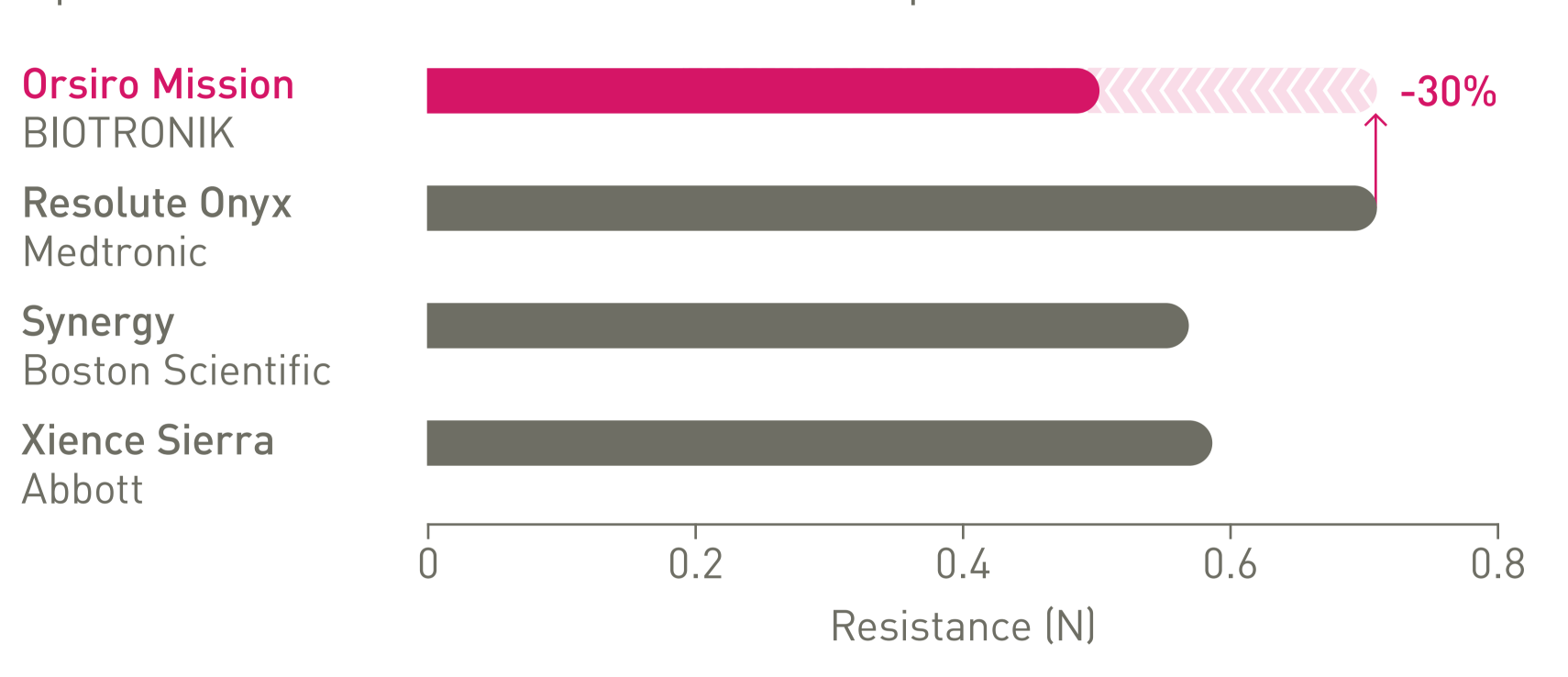
1st in Push⁴

Transmitting up to 57% more force from hub to tip.



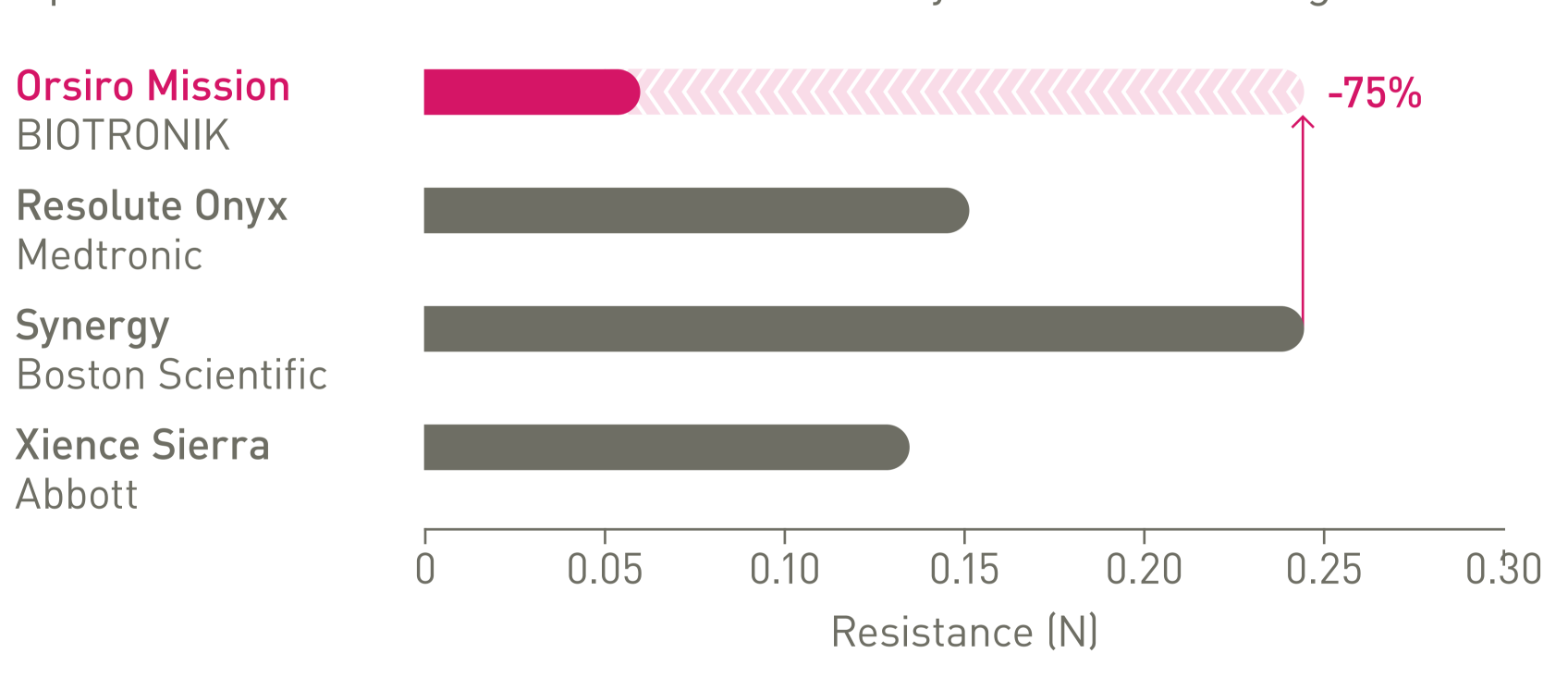
1st in Track⁴

Up to 30% less force needed to follow the path to the lesion.



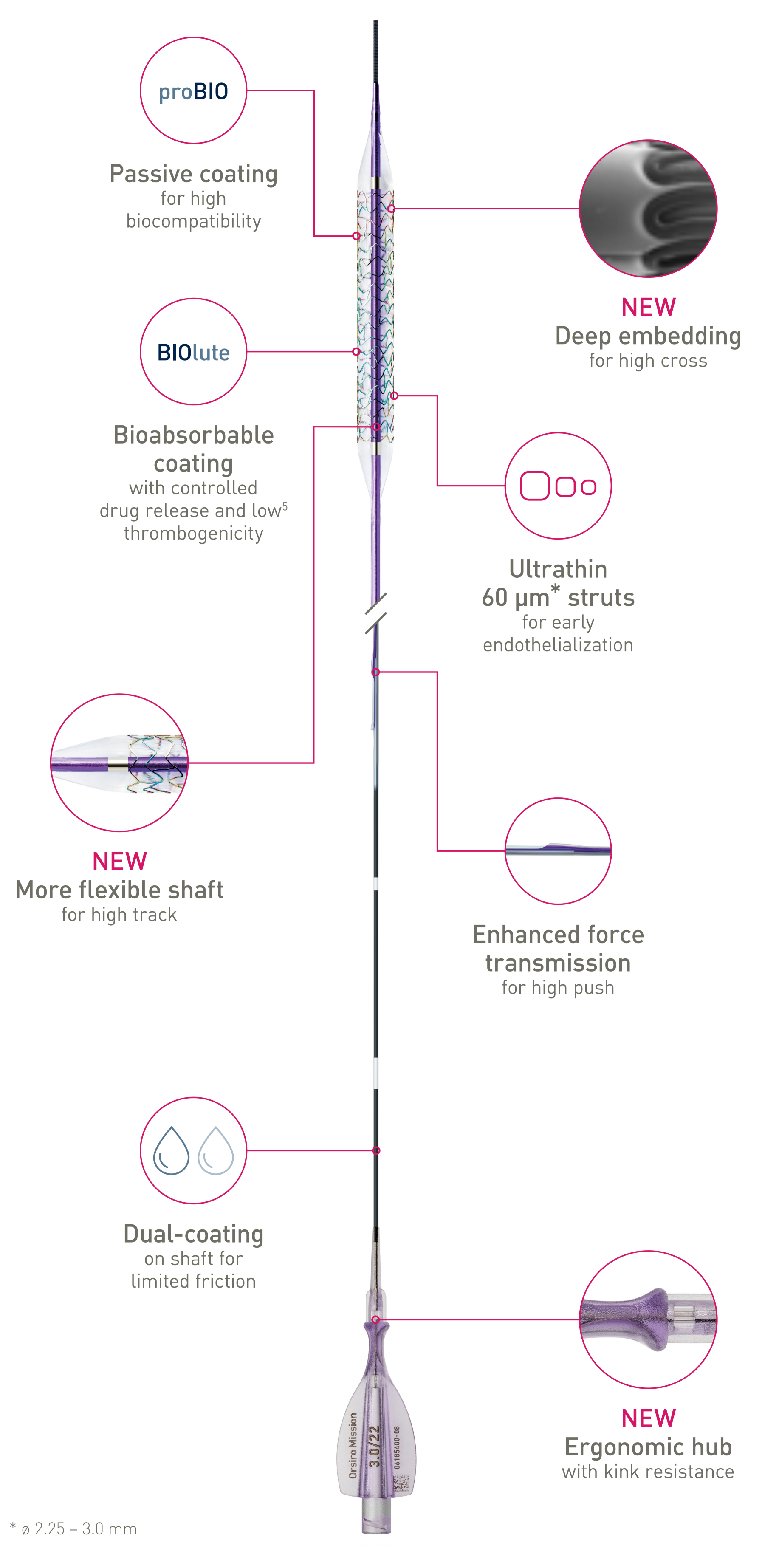
1st in Cross⁴

Up to 75% less force needed to successfully cross demanding anatomies.











Strut thickness in perspective⁶

Orsiro BIOTRONIK CoCr-SES



60 μm*

Synergy Boston Scientific PtCr-EES



74 µm

Ultimaster

Terumo CoCr-SES



80 µm

Resolute Onyx^{7,8} Medtronic

CoNi-ZES



81 µm

Xience Family Abbott

CoCr-EES



 $81 \mu m$

Promus Boston Scientific PtCr-EES



81 µm

BioMatrixBiosensors 316L-BES



120 µm

* ø 2.25 – 3.0 mm

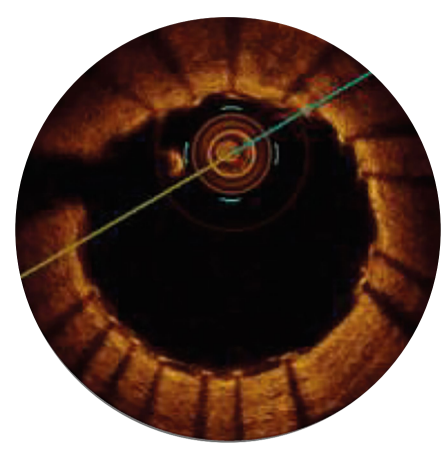
Ultrathin struts²

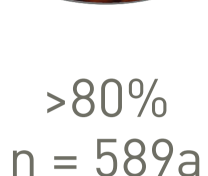
For early endothelialization

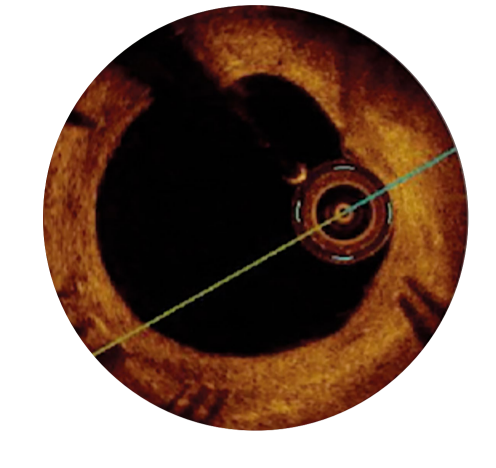
Strut coverage⁹
30 days[△]

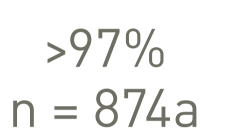
Strut coverage⁹
90 days^Δ

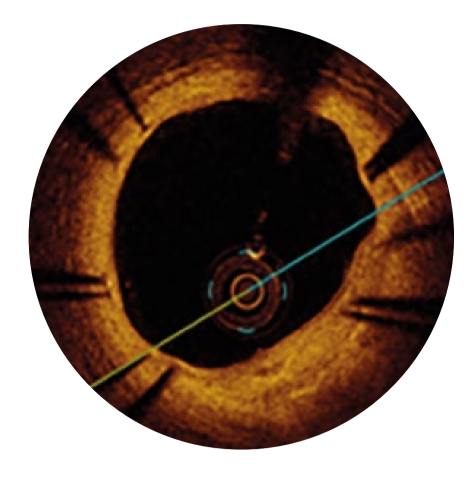












>98%n = 1,130a

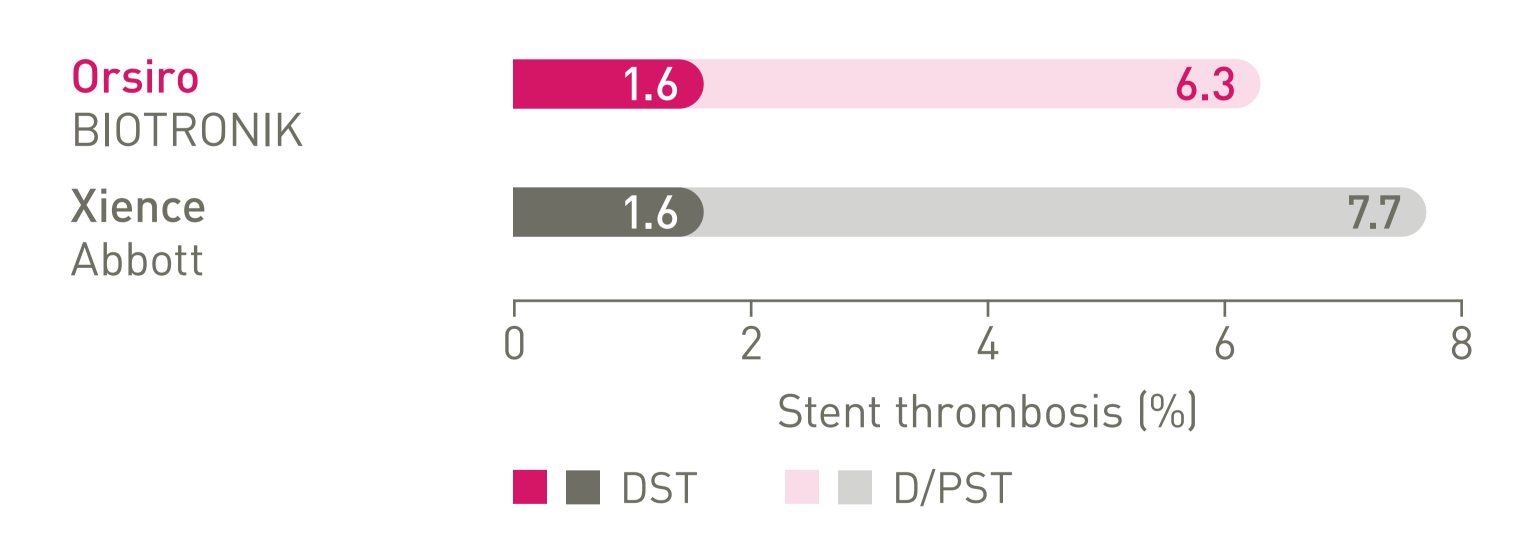
Immature tissue coverage

HEALING PROGRESS

Tissue maturation and full coverage

Long-term safety

Low definite Stent Thrombosis (ST) out to 5 years BIOSCIENCE, all-comers RCT (n= 2,119)¹⁰



DST – Definite Stent Thrombosis D/PST – Definite/Probable Stent Thrombosis



[△]Images: Secco G et al. Time-related changes in neointimal tissue coverage following a new generation SES implantation: an OCT observational study. Presented at: euro PCR, May 20, 2014; Paris, France.

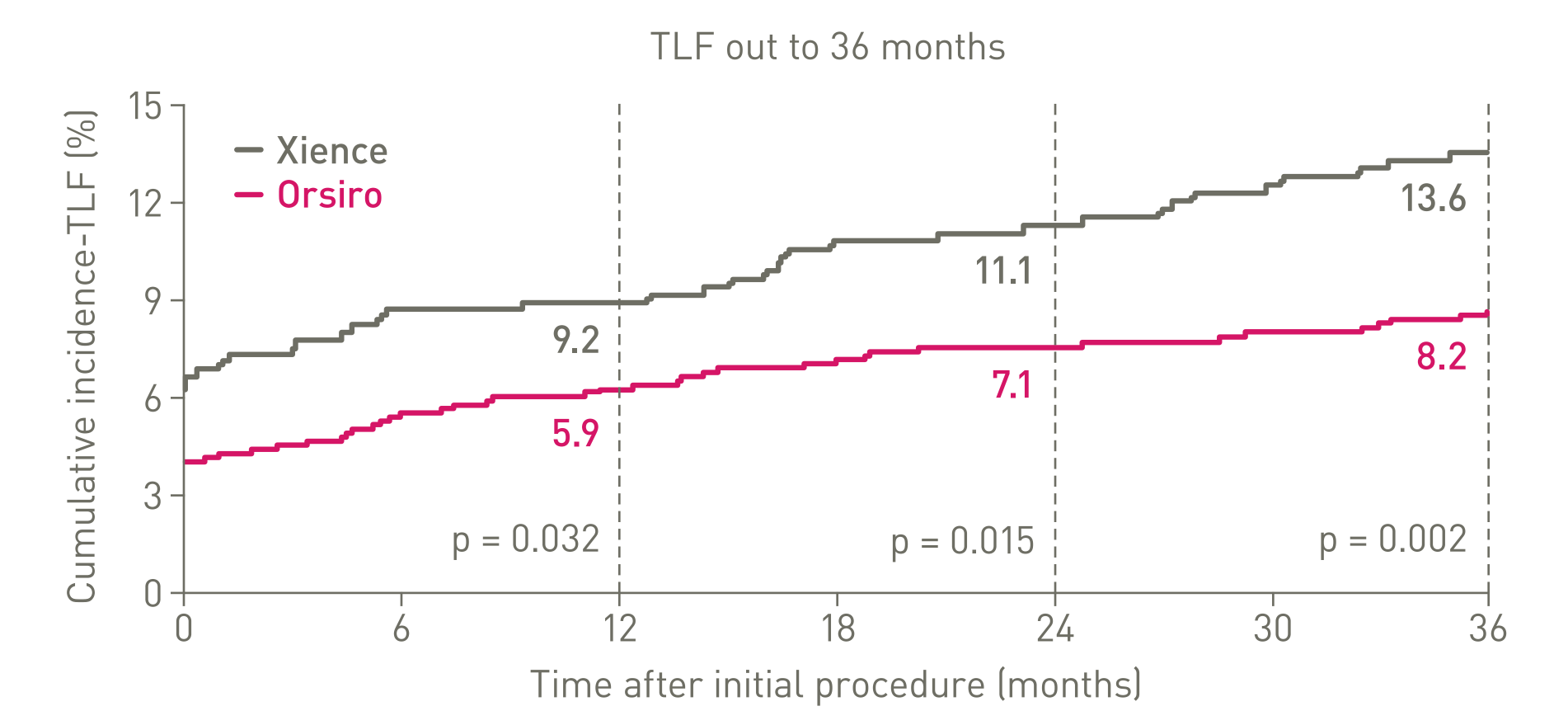
Clinical data conducted with Orsiro, Orsiro Mission's predecessor device can be used to illustrate Orsiro Mission clinical outcomes.



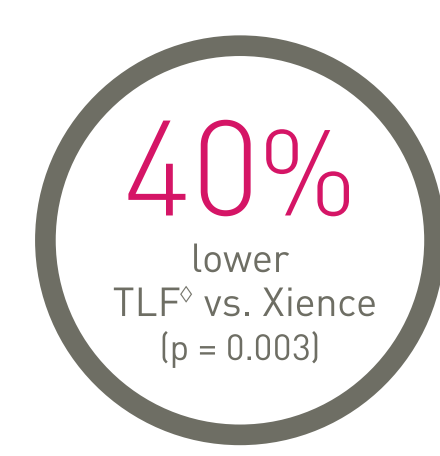
Outstanding patient outcomes³

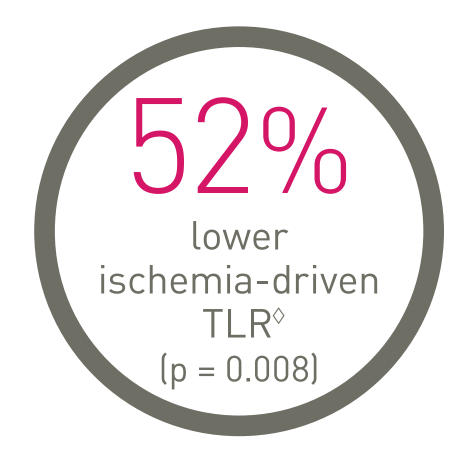
Clinically proven Orsiro DES^{11, 12, 13, 14}

BIOFLOW-V, FDA pivotal trial (n = 1,334)



55,000 patients enrolled





Orsiro Mission is indicated for complex patients and lesions, including:*



BIOSTEMI (n=1,300)

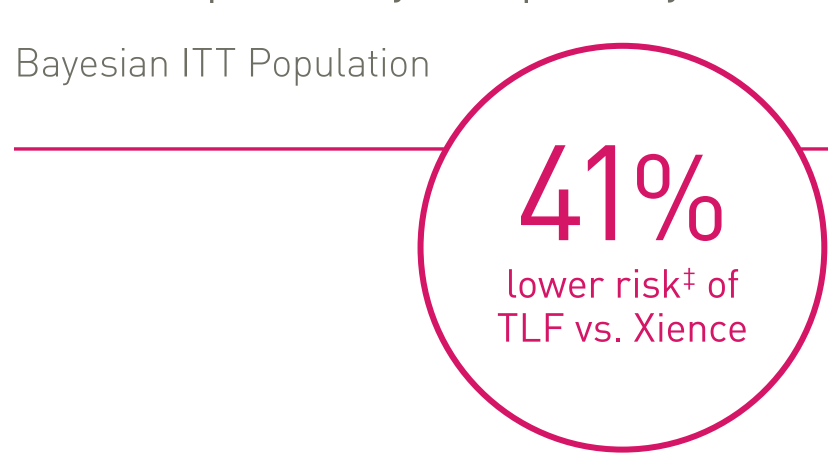
Superiority in STEMI. The first RCT demonstrating superiority between two contemporary DES.¹⁵

Orsiro is superior to Xience in STEMI patients undergoing primary PCI with respect to Target Lesion Failure (TLF) rate at 12 months

40/0 **Orsiro**

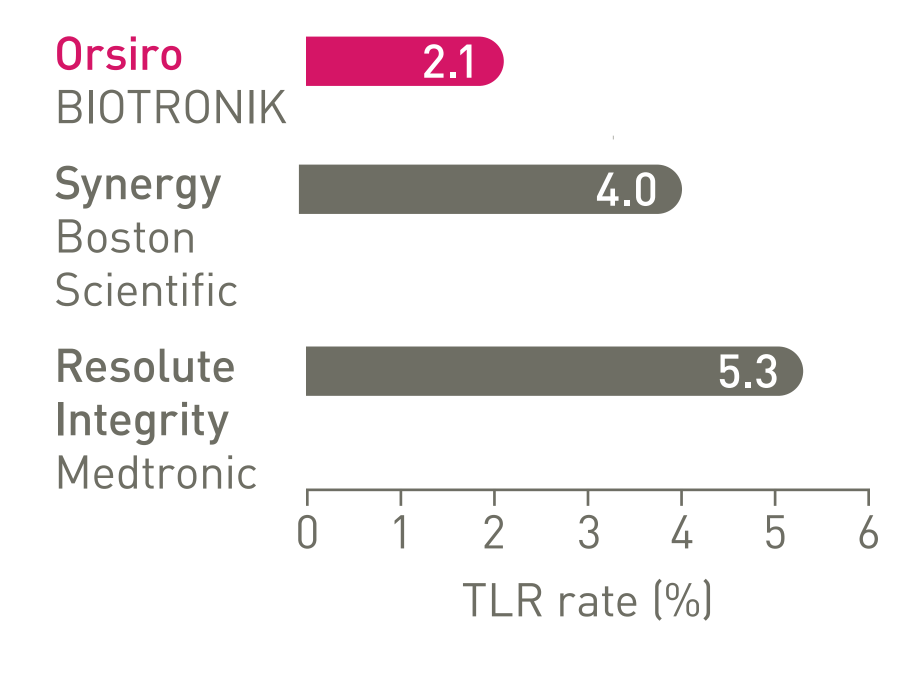
60/o Xience

Rate Ratio (95% BCI**): 0.59, (0.37-0.94) Posterior probability of Superiority: 98.6%



BIO-RESORT Small Vessels (n=1,506)

Target Lesion Revascularization (TLR) rate at 3 yrs.¹⁶





Based on 36-m frequentist analysis.

^{*}As per IFU: ACS – Acute Coronary Syndrome; STEMI – ST-Elevation Myocardial Infarction; DM – Diabetes Mellitus.

HBR – High Bleeding Risk; B2C – Complex Lesions; SV – Small Vessels; MVD – Multi-Vessel Disease.

^{**}BCI: Bayesian Credibility Interval.

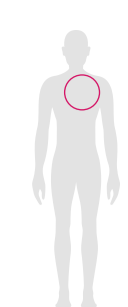
[‡]Based on a Rate Ratio 0.59.

Orsiro® Mission Des

The Orsiro Mission Sirolimus-Eluting Coronary Stent System

is a drug-eluting balloon-expandable stent pre-mounted on

Vascular Intervention Coronary



a rapid-exchange PTCA catheter delivery system.

Indication

Orsiro Mission is indicated for improving coronary luminal diameter in patients with
symptomatic ischemic heart disease due to discrete de-novo stenotic lesions and
in-stent restenotic lesions (length ≤ 40 mm) in the native coronary arteries with a
reference vessel diameter of 2.25 mm to 4.0 mm including the following patient and
lesion subsets:

Acute Coronary Syndrome (ACS) ST-Elevation Myocardial Infarction (STEMI) Diabetes Mellitus (DM) Complex Lesions (B2/C) High Bleeding Risk (HBR)

Long Lesions (LL) (e.g. ≥ 20 mm) Small Vessels (SV) (e.g. ≤ 2.75 mm) Multi-Vessel Disease (MVD) Male/Female Old Patients (e.g. > 65 y)

Technical Data

Stent

Stent material	Cobalt chromium, L-605			
Strut thickness	ø 2.25 – 3.0 mm: 60 μm (0.0024"); ø 3.50 – 4.0 mm: 80 μm (0.0031")			
Passive coating	proBIO (Amorphous Silicon Carbide)			
Active coating	BIOlute bioabsorbable Poly-L-Lactide (PLLA) eluting a limus drug			
Drug dose	1.4 μg/mm²			

Dalivany avetan

Delivery system	
Catheter type	Rapid exchange
Recommended guide catheter	5F (min. I.D. 0.056")
Guide wire diameter	0.014"
Usable catheter length	140 cm
Balloon material	Semi crystalline polymer material
Coating (Distal shaft)	Hydrophilic
Coating (Proximal shaft)	Hydrophobic
Marker bands	Two swaged platinum-iridium markers
Lesion entry profile	0.017"
Distal shaft diameter	2.7F: ø 2.25 – 3.0 mm; 2.9F: ø 3.5 – 4.0 mm
Proximal shaft diameter	2.0F
Nominal pressure (NP)	10 atm
Rated burst pressure (RBP)	16 atm

Storage

otor age					
Use Before Date (UBD)	24 months				
Temperature	Between 15°C (59°F) and 25°C (77°F), short term excursions between 10°C (50°F) and 40°C (104°F) are allowed				

Ordering	Information

Stent ø (mm)	Stent Length (mm)								
	9	13	15	18	22	26	30	35	40
2.25	419101	419107	419113	419119	419125	419131	419137	419143	419149
2.5	419102	419108	419114	419120	419126	419132	419138	419144	419150
2.75	419103	419109	419115	419121	419127	419133	419139	419145	419151
3.0	419104	419110	419116	419122	419128	419134	419140	419146	419152
3.5	419105	419111	419117	419123	419129	419135	419141	419147	419153
4.0	419106	419112	419118	419124	419130	419136	419142	419148	419154

1. In comparison to Xience Sierra, Resolute Onyx and Synergy for bench tests on pushability, trackability and crossability, BIOTRONIK data on file; 2. As characterized with respect to strut thickness in Bangalore et al. Meta-analysis; 3. Based on investigator's interpretation of BIOFLOW-V primary endpoint result; 4. BIOTRONIK data on file; 5. Per investigators' interpretation of preclinical studies with Orsiro as mentioned in Cassese et al. J Thorac Dis 2018;10(2):688-692; 6. Stefanini GG et al. Coronary stents: novel developments. Heart. 2014 Jul 1;100(13):1051-61; 7. Low AF. Stent platform for procedural success: Introducing the Continuous Sinusoidal & Core Wire Technologies. Presented at: AsiaPCR; 22-24 January, 2015; Singapore, Singapore; 8. Tolentino A. Evolving DES Strategy: Biodegradable Polymer vs. Bioabsorbable Scaffold. Presented at: Cardiovascular Nurse/Technologist Symposium; June 17, 2016; New York, USA; 9. Secco G et al. Time-related changes in neointimal tissue coverage of a novel Sirolimus eluting stent: Serial observations with optical coherence tomography. Cardiovascular Revascularization Medicine 17.1 (2016): 38-43; 10. Pilgrim Tet al. 5-year outcomes of the BIOSCIENCE randomised trial. Supplementary appendix; Lancet 2018; published online Aug 28. http://dx.doi. org/10.1016/S0140-6736(18)31715-X; 11. Kandzari D, et al. BIOFLOW-V: A Prospective Randomized Multicenter Study to Assess the Safety and Effectiveness of the Orsiro Sirolimus Eluting Coronary Stent System in the Treatment Of Subjects With up to Three De Novo or Restenotic Coronary Artery Lesions Science. Presentation at ESC 2017; 12. Kandzari D et al. Ultrathin Bioresorbable Polymer Sirolimus-Eluting Stents versus Thin Durable Polymer Everolimus-Eluting Stents: Journal of American College of Cardiology (2018), doi: https://doi.org/10.1 016/j.jac c.2018. 09.019; 13. Kandzari D et al. J Am Coll Cardiol. Cardiovasc Interven. 2020, doi: 10.1016/j.jcin.2020.02.019; 14. Kandzari D et al. J Am Coll Cardiol. Cardiovasc Interven. 2020. Supplemental Material; 15. Iglesias JF et al. Biodegradable polymer sirolimus-eluting stents versus durable polymer everolimus-eluting stents in patients with ST-segment elevation myocardial infarction (BIOSTEMI): a single-blind, prospective, randomised superiority trial; Lancet, September, 2019; 16. Buiten R et al. Outcomes in patients treated with thin-strut, very thin-strut, or ultrathin-strut drug-eluting stents in small coronary vessels – A prespecified analysis of the randomized BIO-RESORT trial; JAMA Cardiol. Published online May 21, 2019. doi:10.1001/jamacardio.2019.1776; ClinicalTrials.gov: NCT01674803. Orsiro, Orsiro Mission, proBIO and BIOlute are trademarks or registered trademarks of the BIOTRONIK Group of Companies. Synergy and Promus are trademarks or registered trademarks of the Boston Scientific group of companies. Resolute, Resolute Onyx and Integrity are trademarks or registered trademarks of the Medtronic group of companies. Xience and Xience Sierra are trademarks or registered trademarks of the Abbott group of companies. Ultimaster is a trademark or registered trademark of the Terumo group of companies. BioMatrix is a trademark or registered trademark of the Biosensors International Group.

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