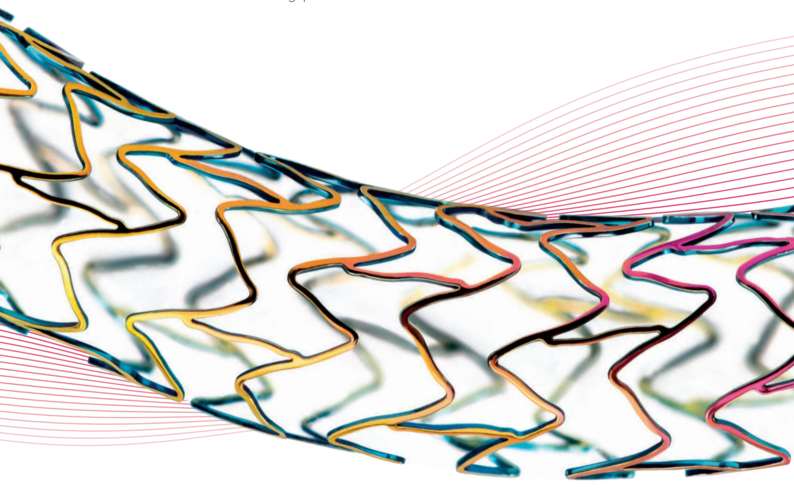
# Orsiro®

Ultrathin struts. Outstanding patient outcomes.





Outstanding patient outcomes



Highly deliverable



Ultrathin 60\* µm struts



# **Orsiro**

Ultrathin struts§. Outstanding patient outcomes¢.

### **Outstanding patient outcomes**

Improving patient outcomes, year after year\*
BIOFLOW-V (n = 1.334) the FDA pivotal trial<sup>1,2,3,4,5</sup>

Significant differences in TLF observed at year 1 and 2 were maintained and further increased at year 3 (8.6% vs. 14.4%, p = 0.003), driven by significant differences in TV-MI (5.5% vs. 10.1%, p = 0.004) and Ischemia-driven TLR (3.4% vs. 6.9%, p = 0.008) that favor Orsiro over Xience.

#### TLF and components at 12, 24 and 36 Months





<sup>§</sup> As characterized with respect to strut thickness in Bangalore et al. Meta-analysis.



<sup>♦</sup> Based on investigator's interpretation of BIOFLOW-V primary endpoint results.

<sup>\*</sup>Compared to Xience, based on three consecutive years.

<sup>\*</sup>p-values for 36-m frequentist analysis (see supplemental material).

<sup>•</sup>vs. Xience, based on 36-m frequentist analysis (see supplemental material).



Strut thickness in perspective<sup>12</sup>

Orsiro BIOTRONIK CoCr-SES

0

60 μm\*

Synergy Boston Scientific PtCr-EES



74 μm

Ultimaster Terumo CoCr-SES



80 µm

Resolute Onyx<sup>13,14</sup>
Medtronic
CoNi-ZES



81 µm

Xience Family Abbott CoCr-EES



81 µm

Promus
Boston Scientific
PtCr-EES



81 µm

**BioMatrix** Biosensors 316L-BES



120 µm

\* ø 2.25 – 3.0 mm

### Ultrathin 60 µm struts

#### Improved outcomes start in the early phase

43% lower in-hospital MI rate<sup>‡</sup> vs. Xience<sup>1</sup>

48 hours

Thinner struts mean less vessel injury<sup>8</sup>



**30 days**<sup>∆</sup> 80.4% strut coverage<sup>9</sup>



**90 days**<sup>∆</sup> 98.7% strut coverage<sup>9</sup>

#### Thinner struts make the difference

Ultrathin vs. second generation DES in a large scale meta-analysis including more than 11,000 patients<sup>10,11</sup>

16%

Relative risk reduction in TLF at 12 months RR (95% CI) 0.84 (0.72, 0.99)

- <sup>‡</sup> Driven by peri-procedural MI events (<48 hours). In-hospital rate may include events > 48 hours.
- <sup>a</sup> 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.



#### Superiority in STEMI<sup>6</sup>

BIOSTEMI (n=1,300) is the first RCT demonstrating superiority between two contemporary DES

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

 $\frac{40}{0}$ 

6% Xience 41% Lower risk\* of TLF with Orsiro in STEMI

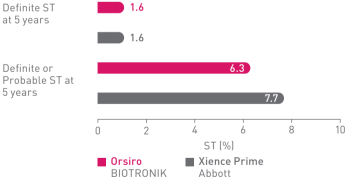
Rate Ratio (95% BCI\*\*): 0.59, (0.37-0.94), **Posterior Probability of Superiority: 98.6%** 

Bayesian ITT Population®

#### Long-term safety

In the randomized, all-comers BIOSCIENCE trial (n= 2,119)7

Orsiro shows numerically equal or lower Stent Thrombosis (ST) in complex patients in comparison to Xience.



<sup>\*</sup>Compared to Xience, BIOTRONIK data on file based on the Rate Ratio of 0.59.

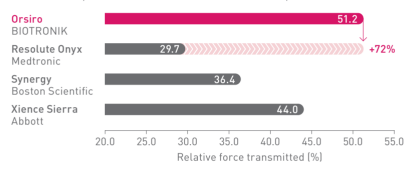
<sup>\*\*</sup>BCI: Bayesian Credibility Interval.

<sup>&</sup>quot;n= 1,300 newly enrolled STEMI patients including 407 patients from the BIOSCIENCE STEMI subgroup used as prior information.

## Highly deliverable

#### Better push

Transmits up to 72% more force from hub to tip. 15



#### Easier cross

Up to 79% less force needed to successfully cross demanding anatomies. 15



#### Lower crossing profile

Improved acute performance – up to 7% lower crossing profile.<sup>15</sup>





# Indicated for discrete de novo stenotic lesions and in-stent restenotic lesions.\*

Technical Data	Stent											
		Stent material			Coba	Cobalt chromium, L-605						
		Passive coating			proB	proBIO (Amorphous Silicon Carbide)						
Active coating					BIOli	BIOlute bioabsorbable Poly-L-Lactide (PLLA) eluting a limus drug						
Drug dose				1.4 μ	1.4 μg/mm²							
	Strut thickness				ø 2.2	ø 2.25 - 3.0 mm: 60 μm (0.0024"); ø 3.50 - 4.0 mm: 80 μm (0.0031")						
		Delivery s										
		Catheter type				Rapid exchange						
Recommended guide catheter					5F (m	5F (min. I.D. 0.056")						
		Lesion entry profile			0.017	0.017"						
	Guide wire diameter			0.014	0.014"							
	Usable catheter length			140 c	140 cm							
Balloon material				Semi	Semi crystalline polymer material							
	Coating (distal shaft)					Hydrophilic coating						
		Marker bands			Two	Two swaged platinum-iridium markers						
	Proximal shaft diameter				2.0F	2.0F						
	Distal shaft diameter				2.6F:	2.6F: ø 2.25 - 3.5 mm; 2.8F: ø 4.0 mm						
	Nominal pressure (NP)				8 atm	8 atm						
Rated burst pressure (RBP)					16 at	16 atm						
Compliance Chart		Balloon diameter x length (mm)										
		ø 2.25 x 9	ø 2.25 x 9-40		ø 2.7	ø 2.75 × 9-40		9-40	ø 3.50 × 9-40 ø 4		00 × 9-40	
Nominal Pressure (NP)	atm**	8	8		8		8		8		8	
	ø (mm)	2.25	2.50		2.75		3.00	3.50		4.00	4.00	
Rated Burst Pressure (RBP)	atm**	16	16		16		16	16		16		
	ø (mm)	2.50	2.77		3.05		3.33		3.88	4.44	4	
Ordering Information		Stent ø (mm)		r <mark>length 140</mark> ngth (mm)	cm					**1 atı	m = 1.013 bar	
		1	9	13	15	18	22	26	30	35	40	
		2.25	364469	364475	364481	364487	364499	364505	364511	391234	391238	
		2.50	364470	364476	364482	364488	364500	364506	364512	391235	391239	
		2.75	364471	364477	364483	364489	364501	364507	364513	391236	391240	
		3.00	364472	364478	364484	364490	364502	364508	364514	391237	391241	
		3.50	364473	364479	364485	364491	364503	364509	364515	391018	391020	
		4.00	364474	364480	364486	364492	364504	364510	364516	391019	391021	

1. Kandzari Detal. Ultrathin, bioresorbable polymersirolimus-eluting stents versus thin, durable polymer everolimus-eluting stents in patients undergoing coronary revascularisation [BIOFLOW V]: a randomised trial. Lancet. 2017 Oct 21; 39(110165]:1843-1852; 2. 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; 3. Kandzari D et al. Ultrathin bioresorbable polymer sirolimus-eluting stents versus thin durable polymer everolimus-eluting stents. Journal of the American College of Cardiology. 2018 Dec 17;72[25]:3287-97; 4. Kandzari D et al. J Am Coll Cardiol. Cardiovasc Interven. 2020, doi: 10.1016/j. jcin.2020.02.019; 5. Kandzari D et al. J Am Coll Cardiol. Cardiovasc Interven. 2020, doi: 10.1016/j. jcin.2020.02.019; 5. Kandzari D et al. J Am Coll Cardiol. Cardiovasc Interven. 2020, doi: 10.1016/j. jcin.2020.02.019; 5. Kandzari D et al. J Am Coll Cardiol. Cardiovasc Interven. 2020, doi: 10.1016/j. jcin.2020.02.019; 5. Kandzari D et al. J Am Coll Cardiol. Cardiovasc Interven. 2020, doi: 10.1016/j. jcin.2020.02.019; 5. Kandzari D et al. J Am Coll Cardiol. Cardiovasc Interven. 2020, doi: 10.1016/j. jcin.2020.02.019; 5. Kandzari D et al. J Am Coll Cardiol. Cardiovasc Interven. 2020, doi: 10.1016/j. jcin.2020.02.019; 5. Kandzari D et al. J Am Coll Cardiol. Cardiovasc Interven. 2020, doi: 10.1016/j. jcin.2020.02.019; 5. Kandzari D et al. J Am Coll Cardiol. Cardiovasc Interven. 2020, doi: 10.1016/j. jcin.2020.02.019; 5. Kandzari D et al. J Am Coll Cardiol. 2016 Delymer sirolimus-eluting stents versus durable polymer everolimus-eluting stents in patients with 5T-sepanent elevation myocardial infarction infarction infarction propendix. Intervention of the BloSCIENCE randomised trial. Supplemental Material, 6. Iglebia as ingle bluing propendix plantal. 2018/j. 10.1016/S0140-6736[18]31715-X

Target Lesion Failure (TLF), Target Lesion Revascularization (TLR), Target Vessel Myocardial Infarction (TV-MI), Stent Thrombosis (ST).

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<sup>\*</sup>Indication as per IFU.