Vascular Intervention // Coronary // Orsiro

BIOFLOW-VI

9-month angiographic and 1 year clinical follow-up for Orsiro in China

Conclusions

- The non-inferiority of Orsiro to Xience Prime was confirmed concerning In-Stent Late Lumen Loss (LLL) at 9 months¹
- Low Target Lesion Failure² (TLF) rates at 12 months in both arms with no statistically significant differences between the individual composites¹
- No definite/probable Stent Thrombosis³ (ST) was reported up to 12 months

The TLF rates are comparable to other clinical trials in Western and Asian populations¹

Study design

Prospective, multi-center, non-inferiority, 1:1 randomized controlled trial comparing Orsiro to Xience in 445 patients with de novo lesions

Endpoints

Primary endpoint

In-Stent LLL at 9 months

Secondary endpoints

- TLF at 12 months
- ST



Annual clinical follow-up up to 60-month

<mark>Orsiro</mark> n = 220	Xience Prime n = 220	p-value
59.1 ± 8.5	58.4 ± 8.6	0.41
27.3%	35.5%	0.06
54.5%	56.8%	0.63
38.2%	42.3%	0.38
27.2%	26.3%	0.98
9.1%	9.5%	0.87
3.2%	0.9%	0.18
57.3%	55.4%	0.50
	Orsiro $n = 220$ 59.1 ± 8.5 27.3% 54.5% 38.2% 27.2% 9.1% 3.2% 57.3%	Orsiro $n = 220$ Xience Prime $n = 220$ 59.1 ± 8.5 58.4 ± 8.6 27.3% 35.5% 54.5% 56.8% 38.2% 42.3% 27.2% 26.3% 9.1% 9.5% 3.2% 0.9% 57.3% 55.4%

Lesion characteristics	<mark>Orsiro</mark> n = 257	Xience Prime n = 259	p-value
Moderate/severe tortuosity	43.4%	43.6%	1.00
Bifurcation lesion	18.4%	20.1%	0.90
Thrombus	0.4%	0.4%	1.00
ACC-AHA Lesion Class B2/C	77.4%	78.8%	0.85

Procedural results	Orsiro	Xience Prime	p-value
Lesion success**	98.8%	99.2%	1.00
Device success ⁺	98.9%	98.9%	1.00
Procedural success§	96.8%	97.7%	0.56

- * 3 withdrawal of consent in Orsiro group, 2 withdrawal of consent in Xience group; did not receive device treatment
- ** Lesion Success: ≤30% residual stenosis of target lesion using any percutaneous method
- ⁺ Device Success: <30% residual stenosis of target lesion using the assigned study stent only
- ⁺⁺ Data shown as mean ± SD
- § Procedural Success: <30% residual stenosis of target lesion using the assigned study stent only without occurrence of in hospital major adverse cardiac event (composite of all death, Q-wave or non-Q-wave MI and clinically-driven TLR)



TLF rates at 12 months¹



TLF components	Orsiro	Xience Prime	p-value
Cardiac death	0.5%	0.0%	0.50
ΤΥ ΜΙ	1.8%	0.9%	0.45
CD-TLR	0.0%	0.9%	0.50
CABG	0.0%	0.5%	1.00

TLF rates⁴ out to 12 months¹



- ⁺ Xience was the comparator for all the trials except for BIO-RESORT in which Resolute Integrity was the comparator
- [‡] TLF components as per respective definitions in trials

Principal investigator

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1. Yang Y, et al. BIOFLOW VI : BIOTRONIK-Safety and Clinical PerFormance of the Drug ELuting Orsiro Stent in the Treatment of Subjects with de novo Coronary Artery Lesions – VI, Presented at: CIT 2018; March 23, 2018; Suzhou, China; ClinicalTrials.gov: NCT02870985; 2. Composite of cardiac death, target vessel Q-wave or non-Q wave Myocardial Infarction (MI), Emergent Coronary Artery Bypass Graft (CABG), clinically driven Target Lesion Revascularization (TLR); 3. ST as per ARC definition; 4. Numbers represented are Kaplan–Meier estimates. Xience is a registered trademark of Abbott Cardiovascular Systems Inc.

Orsiro is a trademark or registered trademark of the BIOTRONIK Group of Companies.

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

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