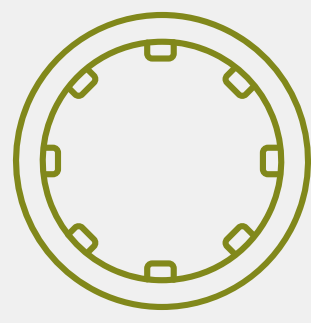




Exceptional
deliverability^{1,2}



Covered single
stent design



Designed to save lives
when seconds count³

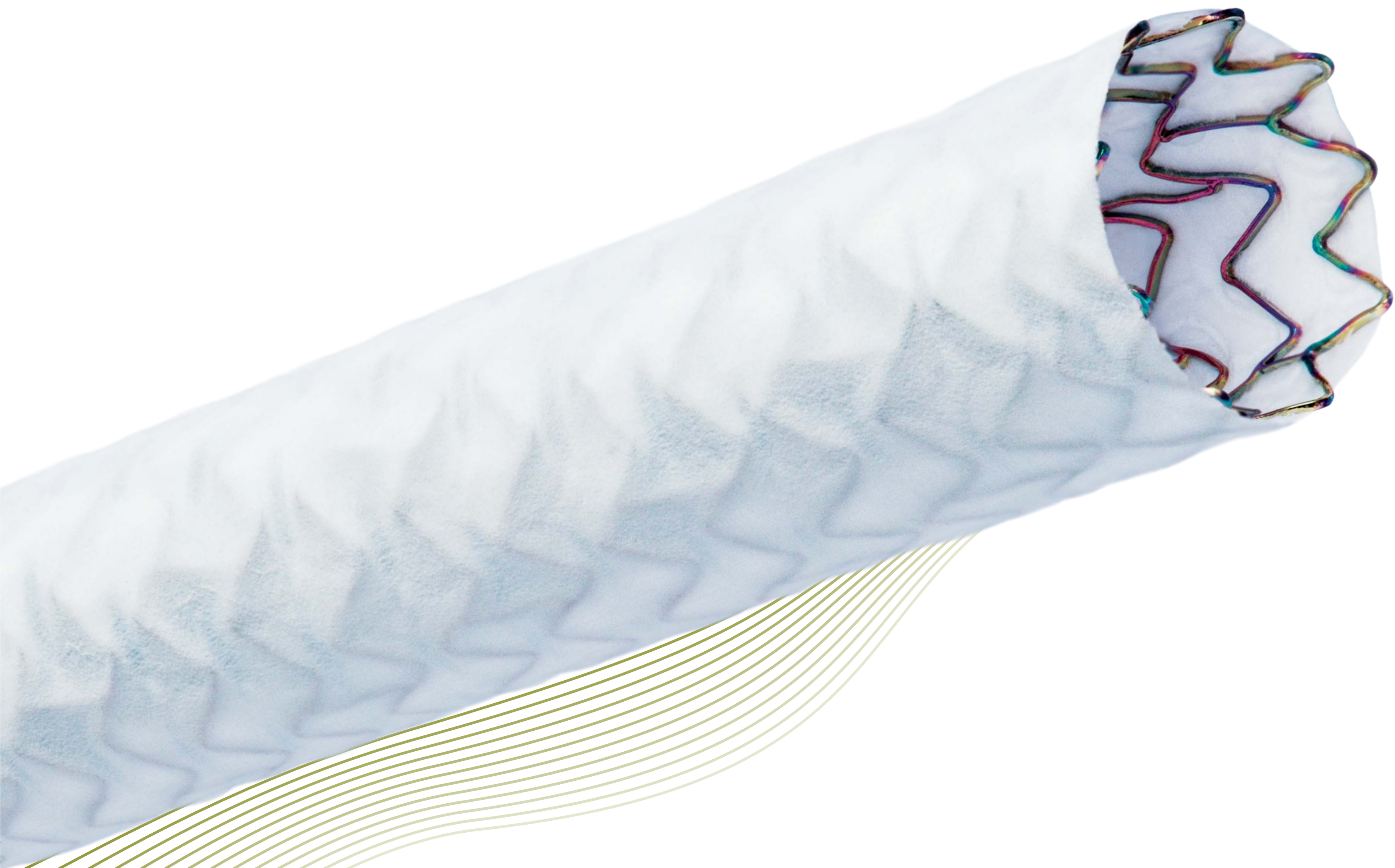


Technical data /
ordering info

Vascular Intervention // **Coronary**
Covered Coronary Stent System

 **BIOTRONIK**
excellence for life

PK Papyrus



Humanitarian Device. Authorized by Federal law for use in the treatment of acute perforations of native coronary arteries and coronary bypass grafts in vessels 2.5 to 5.0 mm in diameter. The effectiveness of this device for this use has not been demonstrated.

PK Papyrus

Designed to deliver more like a conventional stent^{1,2}

Superior design for exceptional deliverability^{1,2}

Lowest crossing profile¹

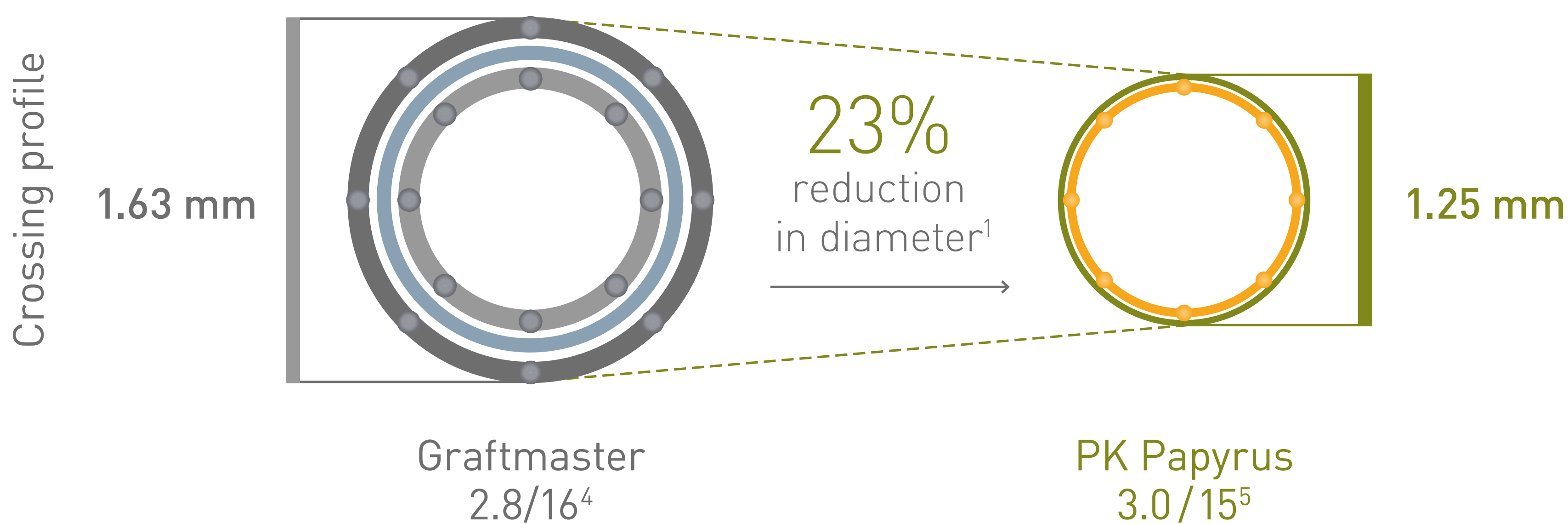
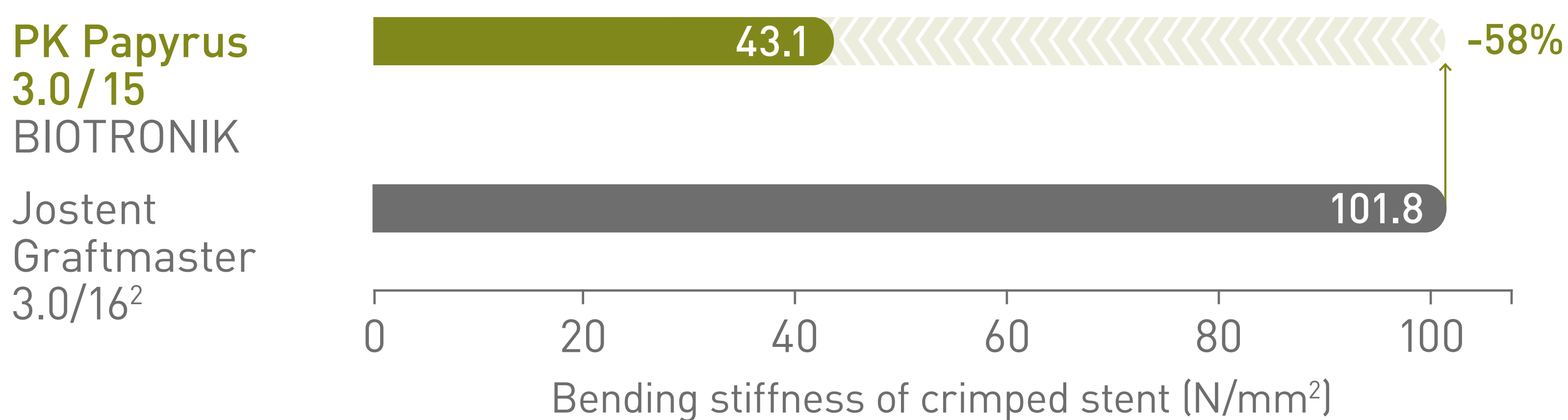


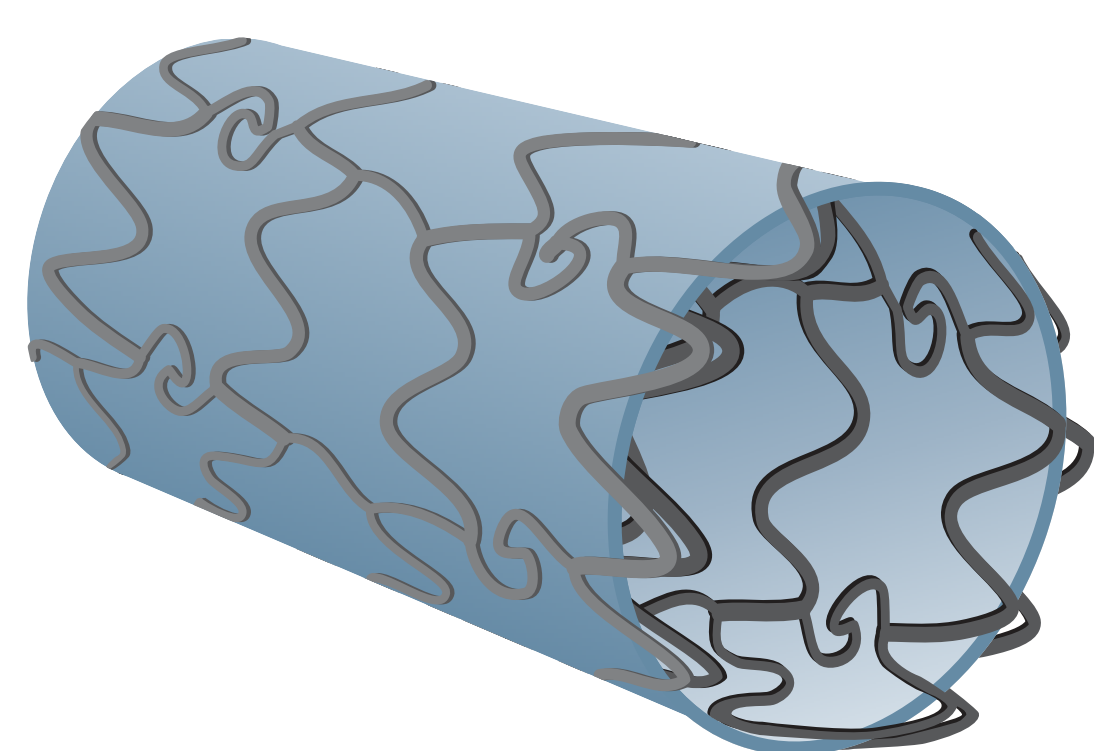
Illustration depicts crimped devices prior to inflation

Superior flexibility²

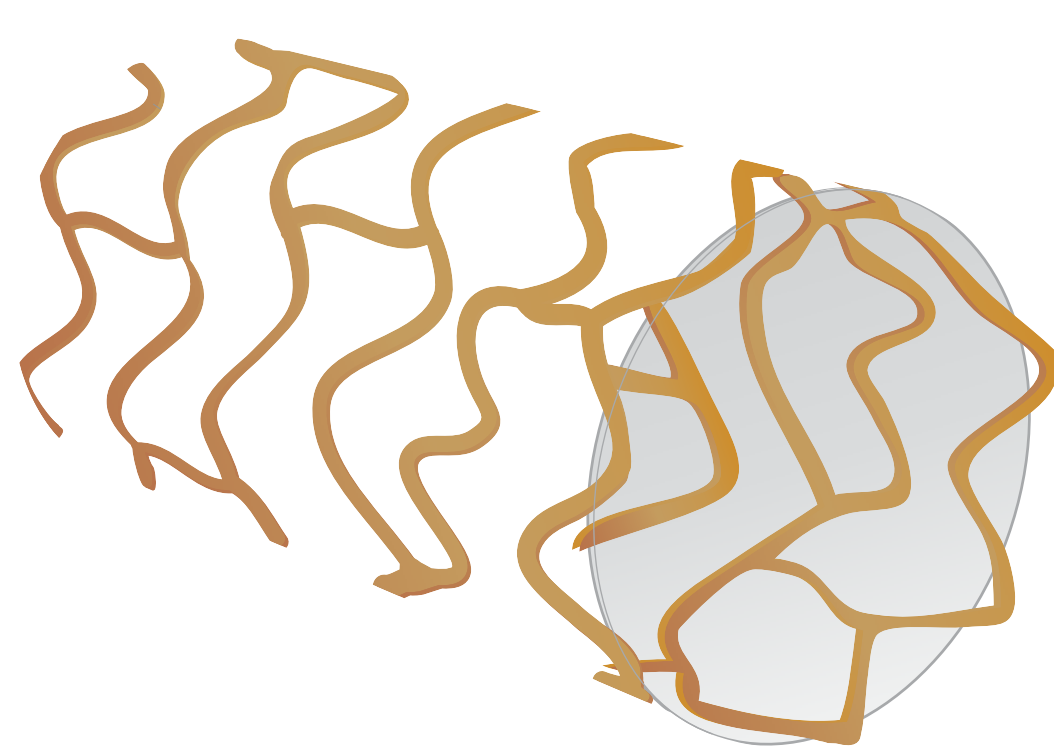


Covered single stent design

- BIOTRONIK's ultrathin strut stent platform (Cobalt Chromium).
- Highly elastic membrane capable of sealing coronary artery perforations.⁶



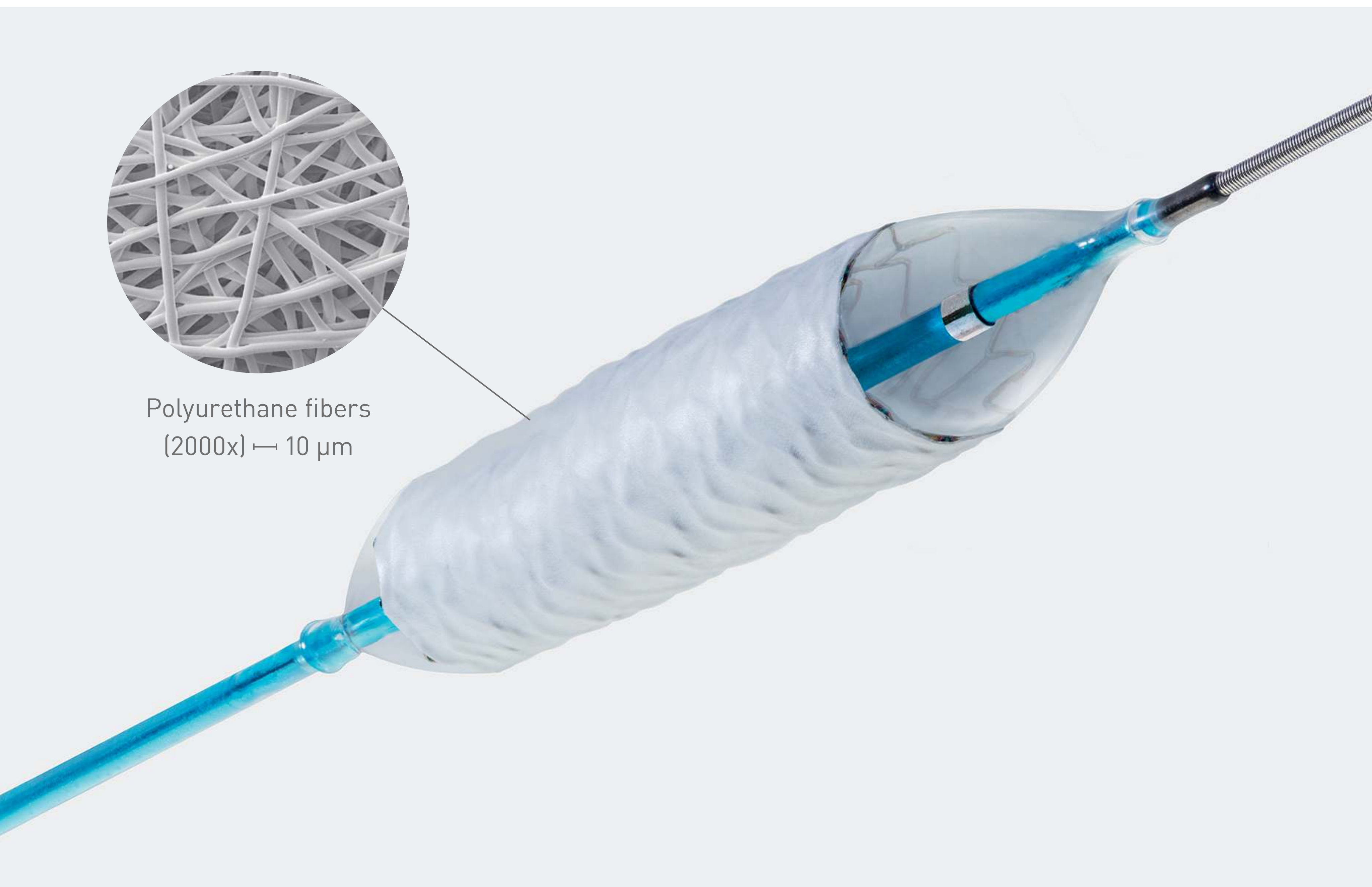
Jostent Graftmaster
Traditional sandwich stent design



PK Papyrus
Covered single stent design

The only
5F
compatibility*

58%
greater flexibility²



Time to expand your options with **PK Papyrus**.

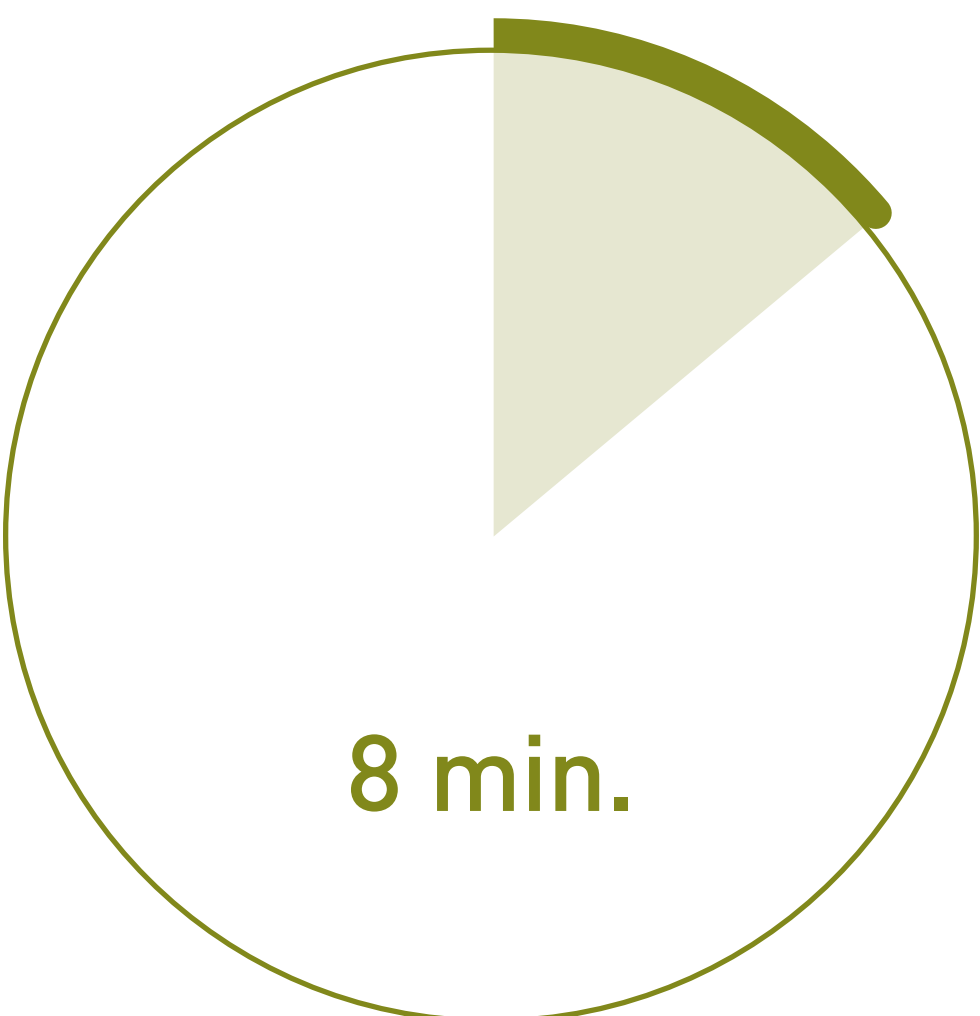
Broadest range of sizes on the US market⁷

- The only 5F compatible* coronary covered stent with the broadest range of sizes.⁷ For main sizes – no need for guide catheter upgrade.*
- **First** FDA approved 2.5 mm diameter.

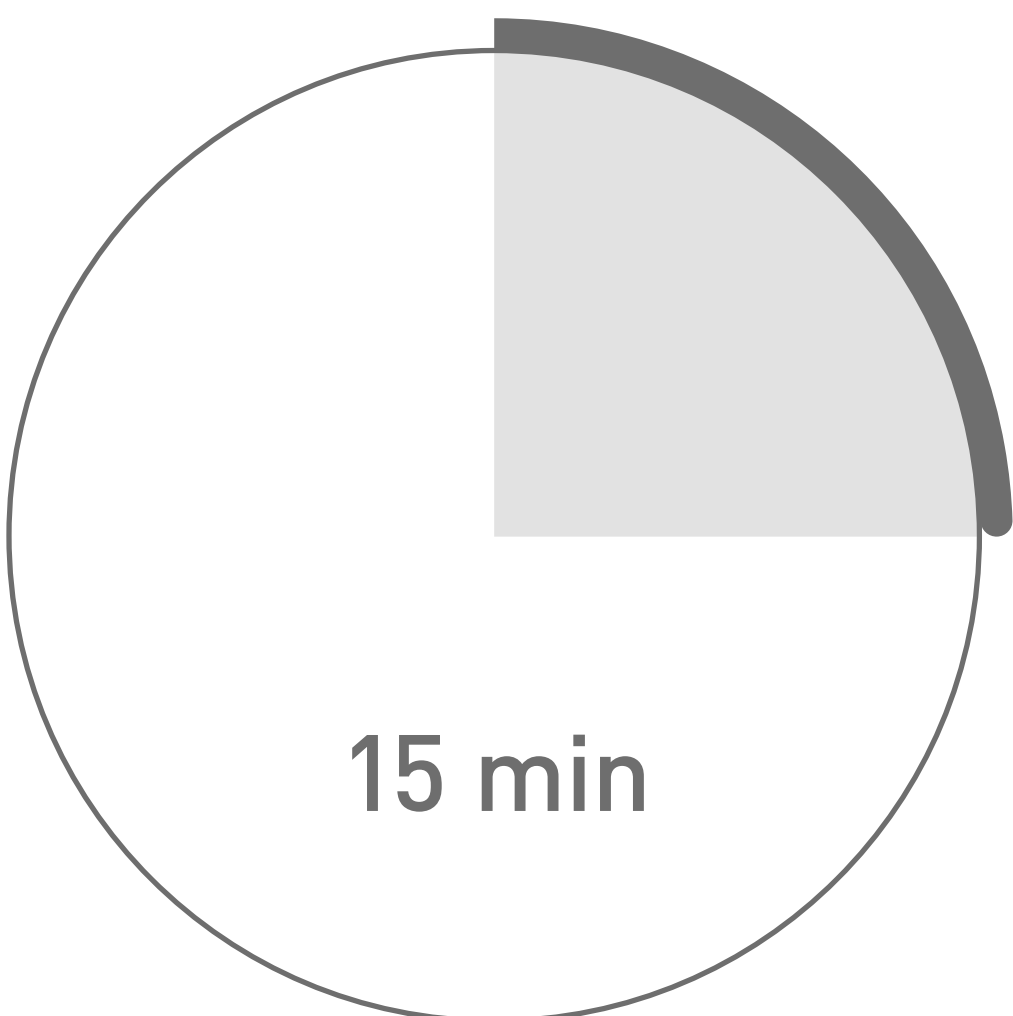
Designed to save lives when seconds count³

Shorter time to deliver⁶

Single center, retrospective investigation of 61 patients treated with covered coronary stents.⁸



PK Papyrus
n = 22



Jostent Graftmaster
n = 39

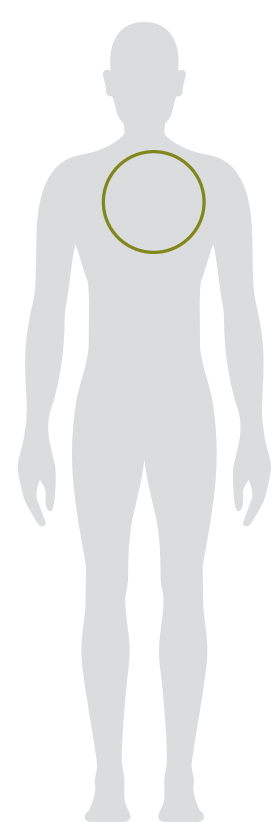
Median Time to deliver (p=0.001)

*5F compatible for Ø2.5-4.0 mm; 6F for Ø4.5-5.0 mm.

PK Papyrus

Indicated for the treatment of acute perforations of native coronary arteries and coronary bypass grafts in vessels 2.5 to 5.0 mm in diameter*

Vascular
Intervention
Coronary



Technical Data	Stent	
	Stent cover material	Non-woven, electrospun polyurethane
	Stent material	Cobalt chromium (L-605) with proBIO (amorphous silicon carbide) coating
	Maximum stent expansion diameter	ø 2.5 - 3.0 mm: 3.50 mm; ø 3.5 - 4.0 mm: 4.65 mm; ø 4.5 - 5.0 mm: 5.63 mm
	Delivery system	
	Guide wire diameter	0.014"
	Usable catheter length	140 cm
	Recommended guide catheter	ø 2.5 - 4.0 mm: 5F (min. I.D.** 0.056"); ø 4.5 - 5.0 mm: 6F (min. I.D.** 0.070")
	Nominal pressure (NP)	ø 2.5 - 3.5 mm: 8 atm; ø 4.0 - 5.0 mm: 7 atm
	Rated burst pressure (RBP)	ø 2.5 - 4.0 mm: 16 atm; ø 4.5 - 5.0 mm: 14 atm
**I.D. = Inner Diameter		

Compliance Chart	Inflation pressure	Stent inner diameter (mm)					
	atm	2.5	3.0	3.5	4.0	4.5	5.0
Nominal pressure (NP)	7	-	-	-	4.01	4.55	4.93
Nominal pressure (NP)	8	2.52	2.99	3.53	4.14	4.69	5.09
	9	2.59	3.07	3.63	4.26	4.82	5.23
	10	2.65	3.15	3.71	4.35	4.91	5.34
	11	2.70	3.21	3.77	4.43	4.99	5.43
	12	2.74	3.26	3.82	4.49	5.06	5.50
	13	2.77	3.30	3.86	4.54	5.11	5.56
Rated burst pressure (RBP)	14	2.80	3.34	3.90	4.59	5.16	5.61
	15	2.83	3.37	3.93	4.63	-	-
Rated burst pressure (RBP)	16	2.86	3.40	3.96	4.67	-	-

Ordering Information	Stent ø (mm)	Catheter length 140 cm Stent length (mm)		
		15	20	26
5F	2.5	434887	434893	-
	3.0	434888	434894	434899
	3.5	434889	434895	434900
	4.0	434890	434896	434901
6F	4.5	434891	434897	434902
	5.0	434892	434898	434903

1. Compared to Graftmaster 2.8 /16 (BIOTRONIK data on file); 2. Compared to Jostent Graftmaster 3.0/16 (BIOTRONIK data on file); 3. Broad range of sizes available on the US market; 4. Data obtained from Graftmaster Coronary Stent Graft System Brochure 11/13/12; 5. PK Papyrus 3.0/15: (BIOTRONIK data on file); 6. Hernandez-Enriquez M, et al. Outcomes after use of covered stents to treat coronary artery perforations. Comparison of old and new-generation covered stents. J Interv Cardiol. 2018; 1-7; 7. Compared to Graftmaster based on the broader range of sizes available on the US market ; 8. Population is representative of real world interventional practice and was not a randomized prospective clinical trial. Jostent and Graftmaster are registered trademarks of the Abbott Group of Companies.

*Indication as per IFU.

Information on devices manufactured at companies other than BIOTRONIK was gathered from multiple sources. However, it has not been verified by the vendors and we cannot guarantee its accuracy.

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