

Ordering Information

DK Score™ Pro		Balloon Length				
		10mm	15mm	20mm	25mm	30mm
Balloon Diameter	1.75mm	CSL-1710	CSL-1715	CSL-1720	CSL-1725	CSL-1730
	2.00mm	CSL-2010	CSL-2015	CSL-2020	CSL-2025	CSL-2030
	2.25mm	CSL-2210	CSL-2215	CSL-2220	CSL-2225	CSL-2230
	2.50mm	CSL-2510	CSL-2515	CSL-2520	CSL-2525	CSL-2530
	2.75mm	CSL-2710	CSL-2715	CSL-2720	CSL-2725	CSL-2730
	3.00mm	CSL-3010	CSL-3015	CSL-3020	CSL-3025	CSL-3030
	3.25mm	CSL-3210	CSL-3215	CSL-3220	CSL-3225	CSL-3230
	3.50mm	CSL-3510	CSL-3515	CSL-3520	CSL-3525	CSL-3530
	3.75mm	CSL-3710	CSL-3715	CSL-3720	CSL-3725	CSL-3730
	4.00mm	CSL-4010	CSL-4015	CSL-4020	CSL-4025	CSL-4030

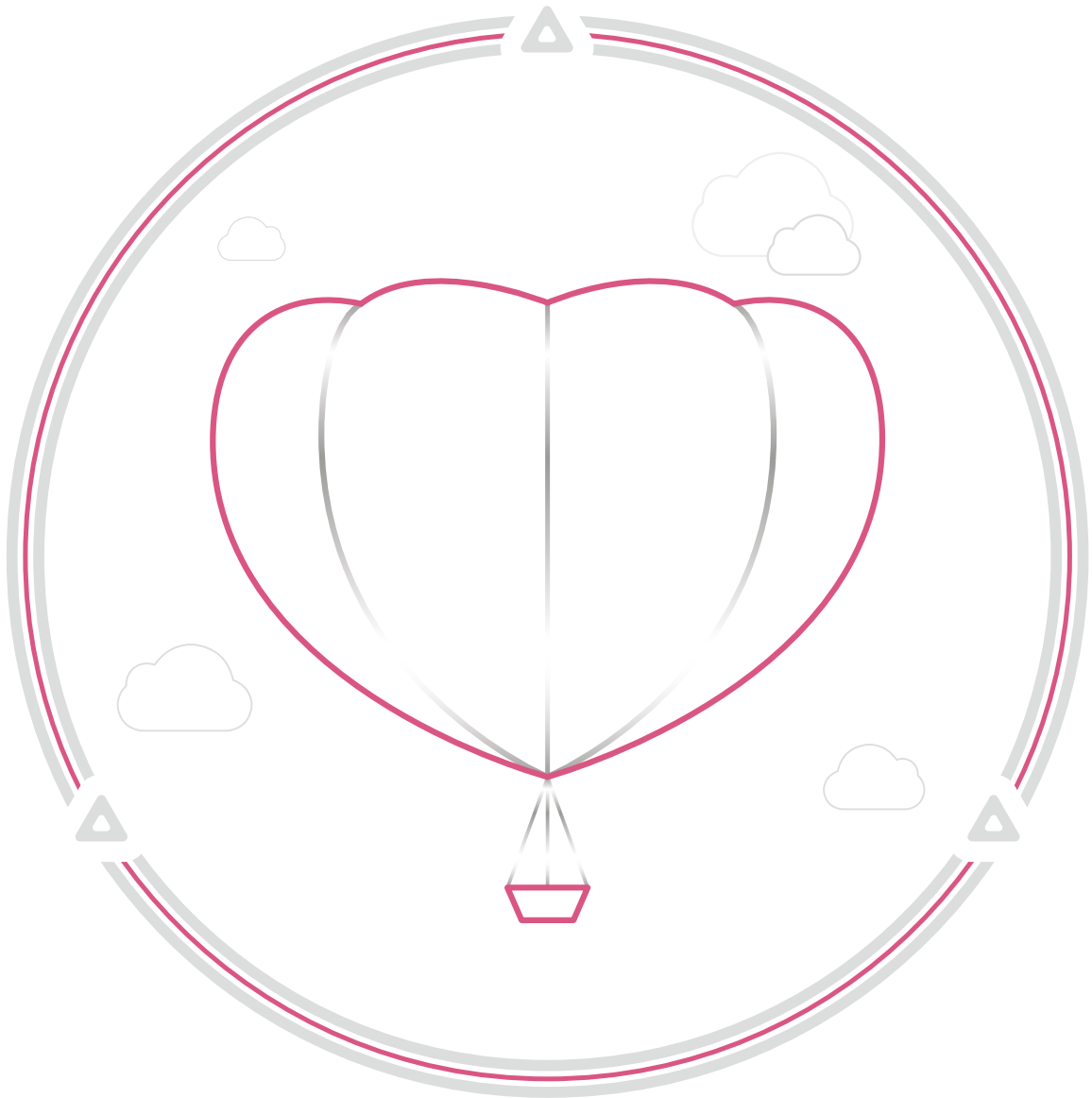
Technical Specifications

Catheter Design	Rapid Exchange (Rx)	Tip Entry Profile	0.016"
Nominal Pressure (NP)	10atm	Guidewire Compatibility	0.014"
Rated Burst Pressure (RBP)	20atm	Guiding Catheter	6F(0.066")

DK Score™ Pro

Coronary Scoring Balloon Dilatation Catheter

CE: M.2025.MDR.1062



RBP 20atm

Triangular Scoring Elements

Patented Nitinol Coiled Wire Technology



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DK Score™ Pro

Precision · Flexibility · Safe
Redefining vascular care with confidence

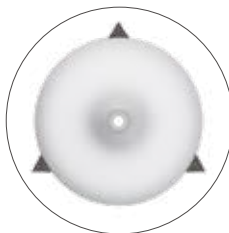
Patented Nitinol Coiled
Wire Technology

Ensures superior crossability
even in tortuous vessels



Triangular Scoring
Elements

Exceptional scoring accuracy
without compromising on patient safety



Distributed 120° Scoring Element
Focused, uniform & controlled dilatation

Proven to Deliver High Focused Force



DK Score™ Pro
Maximum stress 2.782 MPa

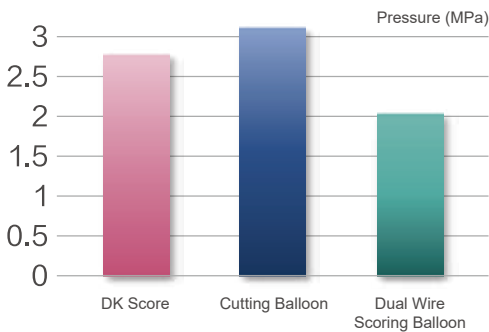


Cutting Balloon
Maximum stress 3.109 MPa



Dual Wire Scoring Balloon
Maximum stress 2.036 MPa

DK Score™ Pro has focused force
transmission that is comparable
to cutting balloons.



Clinical Trial Result

Study conclusion: Acute Lumen Gain and Device Procedural Success of DK Score Scoring balloon is better than Non-Slip Element Dilatation Catheter.

Study Design: DK Score vs. NSE for comparison
Clinical Trials.gov Identifier: NCT05250193



200
Patients
1:1 RCT

