VISIBLE ADAPTABILITY

DERIVO® Embolisation Device





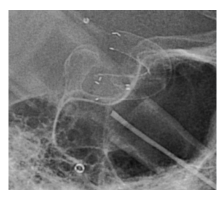


New composite wire concept for outstanding visibility of the DERIVO® contour

Treatment of left saccular ICA aneurysm with DERIVO $^{\circ}$ 5.0 mm x 20 mm



Excellent visibility of DERIVO® contour even in front of dense bone structures. View inside the lumen is possible.

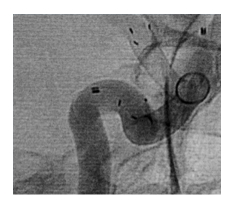


Opening of DERIVO® in tight curve is clearly visible.

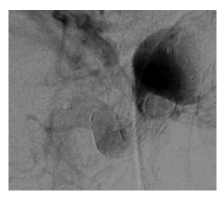
Images by courtesy of: Prof. Reith, Department of Neuroradiology, Saarland University Hospital, Homburg, Germany

Balanced mechanical properties for excellent clinical performance

Treatment of large right ICA aneurysm with DERIVO® 4.0 mm x 30 mm



Perfect wall apposition: DERIVO® contour follows exactly the tortuous shape of the vessel.



Immediate flow diversion effect after DERIVO® placement.



Excellent visibility of fully released DERIVO®.



UNIQUE VISIBILITY

- Completely visible device contour
- Nitinol Composite Wires with Platinum core
- Three Platinum-Iridium X-Ray markers on both ends

BROADEST RANGE

nominal device length from 15 mm – 60 mm, also available in 6 mm σ

- 3D Sizing Support for best flow diversion properties
- Long lengths to avoid telescoping
- Intended vessel diameters from 2.5 mm up to 6 mm

EXCEPTIONAL RELIABILITY

- Secure wall apposition because of flared ends & closed distal ends
- Better corrosion resistance and lower thrombogenicity¹ due to BlueXide® Surface Finishing
- Outstanding flexibility combined with well-balanced radial force

¹ results from in-vitro testings

FLOW - WHERE IT SHOULD BE

Acandis® is using the latest technological developments to ensure a smooth, reliable and precise treatment of intracranial aneurysms with the DERIVO® Embolisation Device.

BlueXide® Surface Finishing

The Acandis® proprietary BlueXide® Surface Finishing
Technology ensures less friction during delivery through the
microcatheter as well as during expansion, making the opening
of the device smooth and reliable. This finishing contributes to
better corrosion resistance which might lead to lower
thrombogenicity.

Nitinol Composite Wires

The entire device consists of Nitinol Composite Wires with Platinum core leading to an outstanding visualisation of the contour and shape of the device under fluoroscopy.

X-Ray Markers

Three Platinum-Iridium X-Ray markers are positioned on each end of the DERIVO® Embolisation Device for an accurate placement.

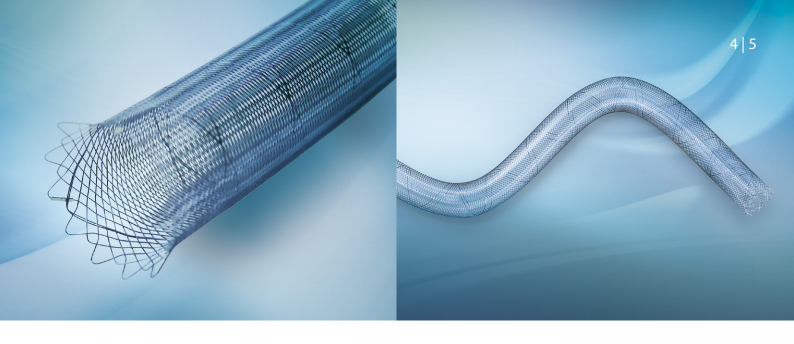
Closed Distal Ends

The closed distal ends of the DERIVO® Embolisation Device help in delivering the device smoothly and releasing it simply, as they create less friction during the delivery through the microcatheter. Additionally these ends are less traumatic, even if the implant is oversized in the distal part of the vessel.

Flared Ends

The DERIVO® Embolisation Device has flared ends for a secure wall apposition immediately after the initial distal opening, while the foreshortening on the proximal end is reduced.





Flow Diversion

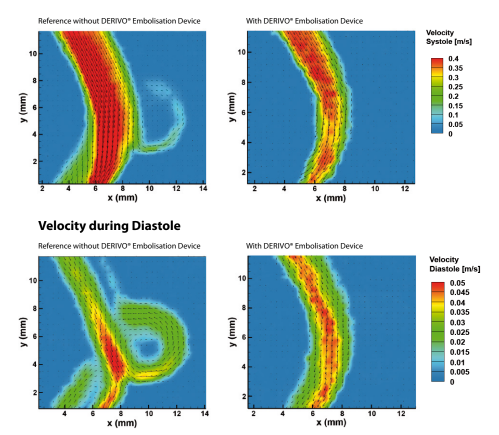
The mesh density enables flow diversion away from the aneurysm while maintaining the flow into the side branches.

Particle Image Velocimetry (PIV) proves the effectiveness of the DERIVO® Embolisation Device flow diversion properties.

Vessel Wall Conformability

The braiding design ensures a good vessel wall conformability, even in highly variable vessel diameters and in tortuous anatomies.

Velocity during Systole



Particle Image Velocimetry (PIV) by courtesy of: Dept. of Cardiovascular Engineering RWTH Aachen (CVE/AME)

PROCEDURE - RELIABLE AND EFFECTIVE

s.e.c.u.r.e. GP Technology

The DERIVO® Embolisation Device is equipped with a Nitinol transport wire using the s.e.c.u.r.e. GP Technology engineered to meet the demands of a reliable and effective procedure.

S-safe

E- enhanced

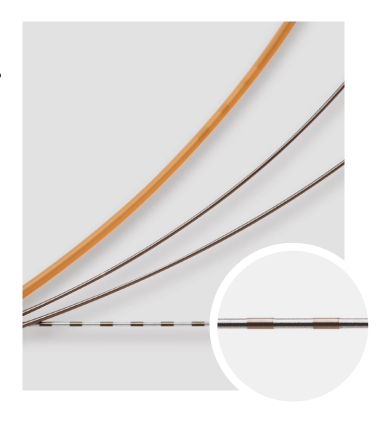
C- controlled

U- unique

R- reliable

E- effective

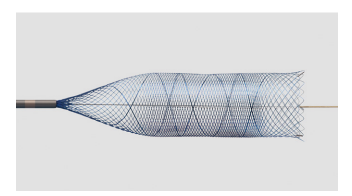
The sleek surface of the transport wire changes into a unique – optically and tactile perceptible – checkered surface at the fluoroscopy marker point, to enhance the grip and push for a controlled and safe placement of the DERIVO® Embolisation Device.



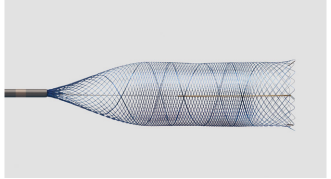
Resheathability

The device can be safely recaptured and repositioned if an adjustment and superior placement is needed.

Tip Design



With tip – for additional distal support and retention of device access after release.



Without tip (only applicable for 40 mm and 50 mm device lengths) – for more flexibility and tip control in the treatment of long lesions.

SIZING SUPPORT CHART – DERIVO® EMBOLISATION DEVICE

Labelled DERIVO® Dimensions (mm)	Reference Number		Unconstrained DERIVO® Dimensions (mm)	Inte	DERIVO® Lengths in corresponding nded Use Diameters ((mm)
		Ø	3.7	3.5	3.0	2.5
3.5 × 15	01-000408	_	10	15	20	25
3.5 × 20	01-000409	Device Length	13	20	27	32
3.5 × 25	01-000410	e Le	16	25	35	41
3.5 × 30	01-000411	evic	19	30	41	48
3.5 × 40	01-000415		25	40	53	66
		Ø	4.2	4.0	3.5	3.0
4.0 × 15	01-000381	Device Length	11	15	20	25
4.0 × 20	01-000330		14	20	27	32
4.0 × 25	01-000335	e Le	17	25	35	41
4.0 × 30	01-000340	Devi	20	30	41	48
4.0 × 40	01-000360	_	26	40	53	66
		Ø	4.7	4.5	4.0	3.5
4.5 × 15	01-000382	ے	11	15	20	25
4.5 × 20	01-000331	engt	14	20	27	32
4.5 × 25	01-000336	e Le	17	25	35	41
4.5 × 30	01-000341	Device Length	20	30	41	48
4.5 × 40	01-000361		26	40 53 66	66	
		Ø	5.2	5.0	4.5	4.0
5.0 × 15	01-000383		11	15	20	23
5.0 × 20	01-000332	gth	14			
5.0 × 25	01-000337	Device Length	17		41	
5.0 × 30	01-000342	evice	20	30	41	48
5.0 × 40	01-000362	۵	26	40	53	62
5.0 × 50	01-000363		34	50	68	82
		Ø	5.7	5.5	5.0	4.5
5.5 × 15	01-000384	Device Length	11	15	20	23
5.5 × 20	01-000333		14	20	27	32
5.5 × 25	01-000338		17	25	35	41
5.5 × 30	01-000343	evic	20	30	41	48
5.5 × 40	01-000364	٥	26	40	53	62
5.5 × 50	01-000365		34	50	68	82
		Ø	6.2	6.0	5.5	5.0
6.0 × 15	01-000385		11	15	20	23
6.0 × 20	01-000334	Device Length	14	20	27	32
6.0 × 25	01-000339	e Lei	17	25	35	41
6.0 × 30	01-000344	evic	20	30	41	48
6.0 × 40	01-000366	Δ	26	40	53	62
6.0 × 50	01-000367		34	50	68	82

Note: all indicated lengths can vary within a tolerance range of +/- 1mm $\,$

For optimal case preparation, Acandis also offers software-based 3D Sizing Support.

For further information please contact the Clinical Support Team: clinical-support@acandis.com

Labelled DERIVO® Diameter (mm)	Labelled DERIVO® Length (mm)	Reference Number	Recommended Vessel Diameter (mm)	Required Microcatheter for Delivery ** (inch)	
3.5	15	01-000408			
	20	01-000409			
	25	01-000410	2.5 – 3.5		
	30	01-000411			
	40	01-000415*			
	15	01-000381			
	20	01-000330		0.027	
4.0	25	01-000335	3.0 – 4.0		
	30	01-000340			
	40	01-000360*			
	15	01-000382			
	20	01-000331			
4.5	25	01-000336	3.5 – 4.5		
	30	01-000341			
	40	01-000361*			
	15	01-000383			
	20	01-000332			
5.0	25	01-000337	40.50		
5.0	30	01-000342	4.0 – 5.0		
	40	01-000362*			
	50	01-000363*			
	15	01-000384			
	20	01-000333			
5.5	25	01-000338	4.5 – 5.5		
	30	01-000343	4.5 – 5.5		
	40	01-000364*			
	50	01-000365*			
6.0	15	01-000385			
	20	01-000334			
	25	01-000339	5.0 – 6.0		
	30	01-000344	3.0 - 6.0		
	40	01-000366*			

 $All \ changes \ or \ modifications, \ may \ they \ be \ technical \ or \ other, \ or \ changes \ in \ the \ availability \ of \ products \ are \ expressively \ reserved.$

01-000367*

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^{*} Indicated on package as "without Tip" as the tip always stays inside the stent for the 40 mm and 50 mm length

^{**} Please contact your local Acandis® representative for information on compatible microcatheters