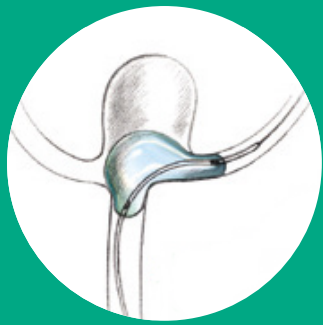
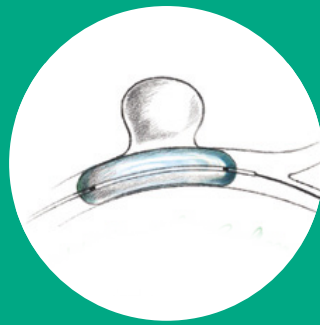


Eclipse (hyper compliant):



Copernic (compliant):



ordering information

Eclipse and Copernic are single lumen balloon catheters packaged with the corresponding guidewire.

Eclipse

Reference	Total length (cm)	Balloon length (mm)	Balloon diameter (mm)	Maximum volume (ml)	RX markers	Guidewire recommended
SECLIPSE7	160	7	4 to 6	0.2	3 markers	Compatible with .012" guidewire (packaging includes a HYBRID1214D)
SECLIPSE9		9		0.25		
SECLIPSE12		12		0.3		
SECLIPSE15		15		0.4		
SECLIPSE20		20		0.5		

Copernic

Reference	Total length (cm)	Balloon length (mm)	Balloon diameter (mm)	Maximum volume (ml)	RX markers	Guidewire recommended
SCOPERNIC10	160	10	3 to 5	0.25	2 markers	Compatible with .012" guidewire (packaging includes a HYBRID1214D)
SCOPERNIC15		15		0.35		
SCOPERNIC20		20		0.4		
SCOPERNIC30		30		0.5		

*Internal data

COPERNIC and ECLIPSE are occlusion catheters indicated for use in the neurovasculature and peripheral system to temporarily stop or control blood flow, to treat vasospasms and embolization of aneurysms with balloons. Class III CE0297 in compliance with Medical Device Directive (MDD 93/42/EEC amended by 2007/47/EC). Manufactured by BALT EXTRUSION SAS, 10 rue de la Croix Vigneron, 96160 Montmorency, France. Carefully read the instructions for use before use. First CE marking: 2001 (COPERNIC), 2006 (ECLIPSE). HYBRID guidewires are designed to facilitate the insertion of catheters into intracranial vascular branches for diagnostic or therapeutic use. Class III CE0297 in compliance with Medical Device Directive (MDD 93/42/EEC amended by 2007/47/EC). Manufactured by BALT EXTRUSION SAS. Carefully read the instructions for use before use. First CE marking: 2010. The guiding catheter FARGO is intended to facilitate the introduction of micro-catheters for therapeutic and diagnostic use. Class III CE0297 in compliance with Medical Device Directive (MDD 93/42/EEC amended by 2007/47/EC). Manufactured by BALT EXTRUSION SAS. Carefully read the instructions for use before use. First CE marking: 2009.

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Balt

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95160 Montmorency
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eclipse & copernic

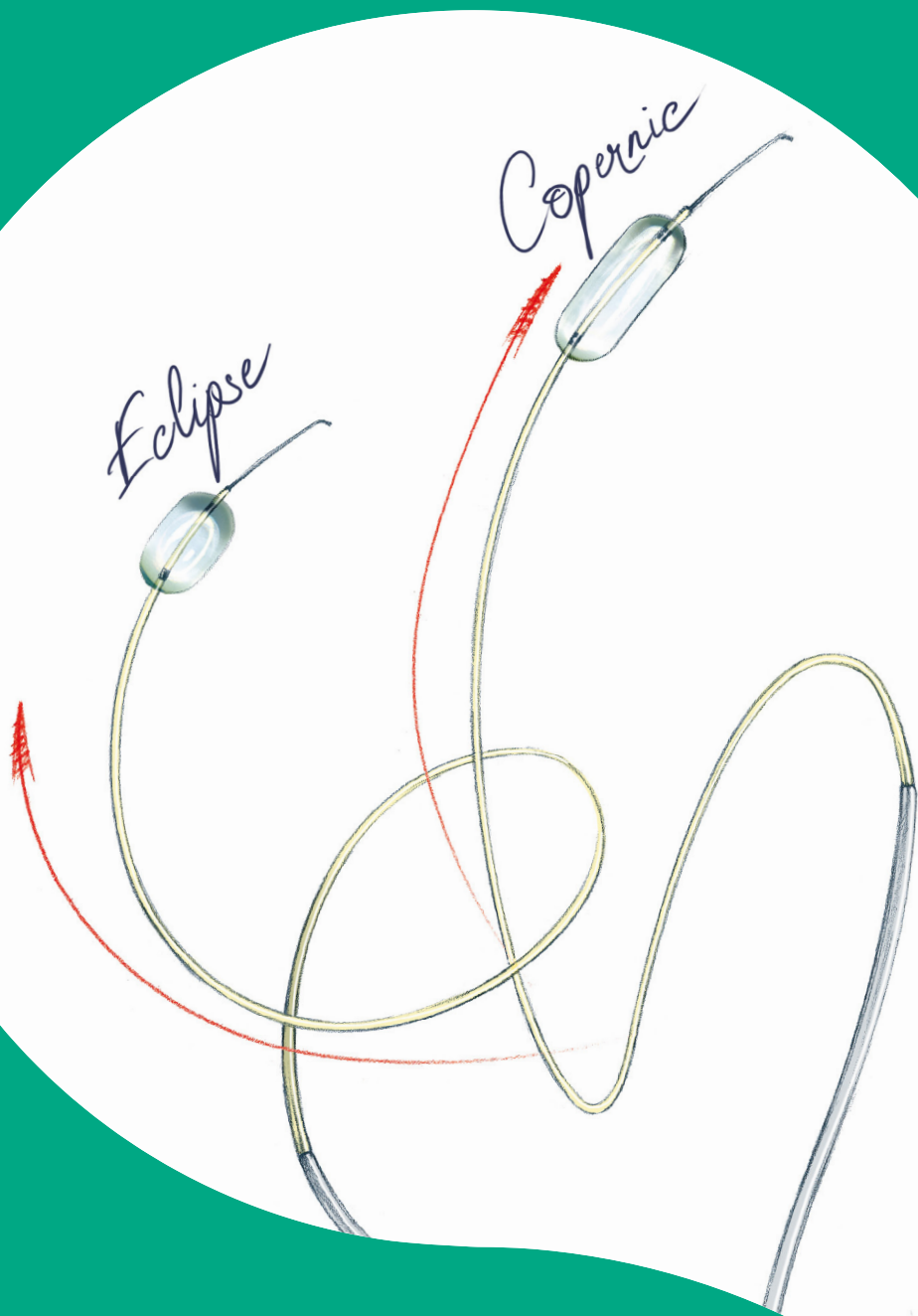
remodeling balloon catheters



eclipse & copernic

Compliant and hypercompliant balloon catheters

Designed to temporarily stop or control blood flow, to treat vasospasms, and for embolization of aneurysms with balloons.



balloon
size from 4 to 6
& 3 to 5mm

Navigation

Hydrophilic coating
on both the catheter & the balloon
for a smooth progression

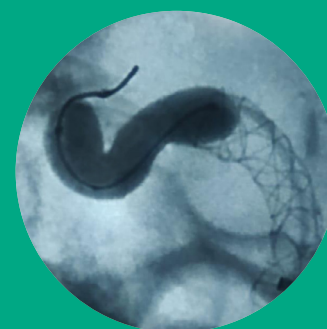
Compatibility with
hybrid1214D guidewire
which ensures support & trackability

Compliant & hypercompliant balloons

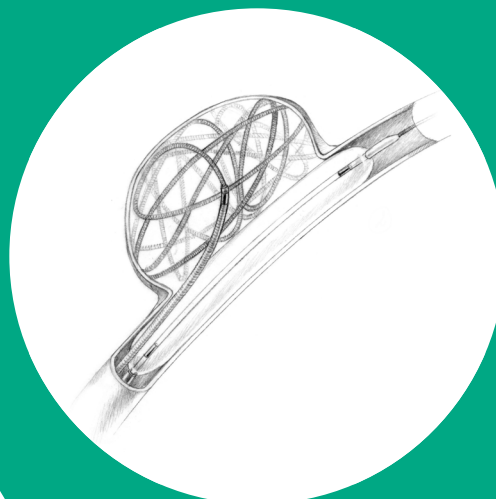
Made from a very supple elastomer*
to easily conform to the anatomy

Rapid inflation deflation
with 1/2 contrast*

Good visibility
thanks to 2RX markers providing the
balloon length



With courtesy of
Dr Sukalyan Purkayastha, India



Remodeling

Both Eclipse and Copernic
can be used in parallel with
a coiling microcatheter in a
single 6F guiding catheter
(FARGOMAX6F)

aneurysm treatment