

copernic^{rc}

venous remodeling
balloon



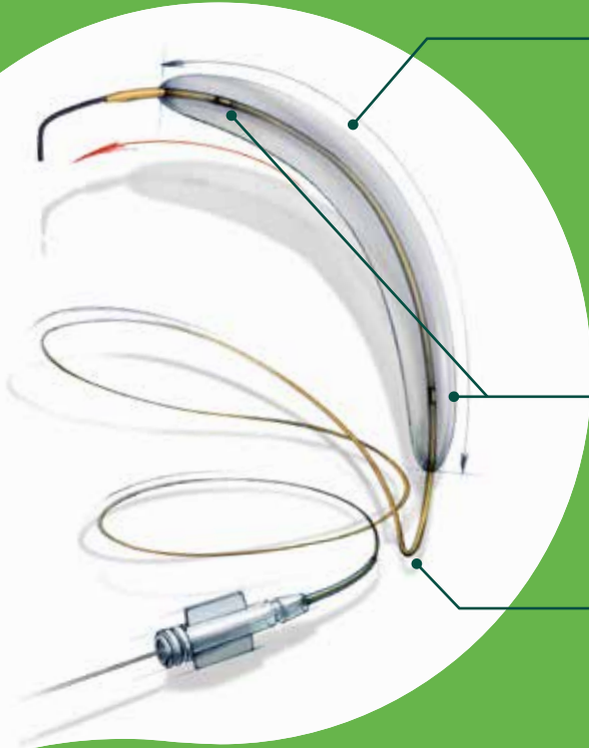
avm* treatment



copernic^{rc}

Balloon catheter

Designed for use in the neurovasculature and peripheral system to temporarily stop or control blood flow.

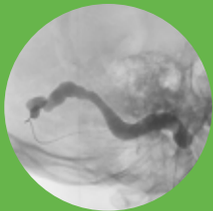


● Preservation of the sinus
Large and long balloon
for a complete occlusion of the sinus
to preserve it during embolization

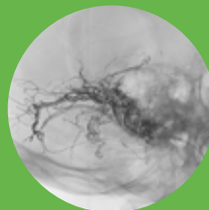
Ease of inflation & deflation
designed for large volume balloon

● Adequate positioning
Visibility of the balloon
thanks to 2 radiopaque platinum rings
at its distal and proximal sides

● Smooth navigation
Thanks to its hydrophilic coating surface



Proper balloon
conformability



DMSO** compatible

With courtesy of Pr. Chapot

ordering information

Unique features on the market

Reference	Balloon diameter (mm)	Balloon length (mm)	Catheter length (cm)	Max. outer diameter	Compatible guidewire
COPERNIC8x80RC	8	80	160	4,35F (1,45mm)	.014" guidewire
COPERNIC10x80RC	10	80		5F (1,6mm)	

*Arteriovenous Malformation. **Dimethyl sulfoxide

COPERNIC: 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, 95160 Montmorency, FRANCE. Carefully read the instructions for use before use. First CE marking: 2001.

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