

Berner Symposium Gefäßmedizin 2025

Donnerstag, 6. November 2025 von 08:30–16:45 Uhr
im Kursaal Bern | Kongresszentrum

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Willkommen

Liebe Kolleginnen, liebe Kollegen

Wir freuen uns, Sie zum Berner Symposium Gefässmedizin 2025 einzuladen.

Für Sie haben wir ein spannendes und praxisnahes Programm zusammengestellt, das neuste Entwicklungen und Behandlungsmethoden in verschiedenen Teilbereichen der Gefässmedizin beleuchtet. Im Plenum referieren nationale und internationale Expertinnen und Experten aus unterschiedlichen Fachrichtungen zu folgenden Themen:

- **Das rupturierte und infizierte Aortenaneurysma**
- **Claudicatio intermittens und kritische Extremitätenischämie**
- **Male health (Prostataembolisation und Behandlung der Varikozele)**
- **Challenging cases: aortal, peripher arteriell und venös**

Wie im vergangenen Jahr wird die Veranstaltung interaktiv gestaltet sein, mit Möglichkeit zum Austausch und zur Diskussion für alle Teilnehmenden. Das Symposium richtet sich an alle, die sich mit der Gefässmedizin beschäftigen. Neben den Ärzt:innen der Angiologie, der interventionellen Radiologie und der Gefässchirurgie laden wir auch herzlich unsere niedergelassenen Kolleginnen und Kollegen ein.

Wir freuen uns sehr, Sie am 6. November 2025 im Kongresszentrum des Kursaals Bern persönlich begrüßen zu dürfen.

Freundliche Grüsse

Prof. Dr. med.
Drosos Kotelis

Prof. Dr. med.
Marc Schindewolf

Dr. med.
Michel Bosiers

Dr. med.
Maria Schaumeier

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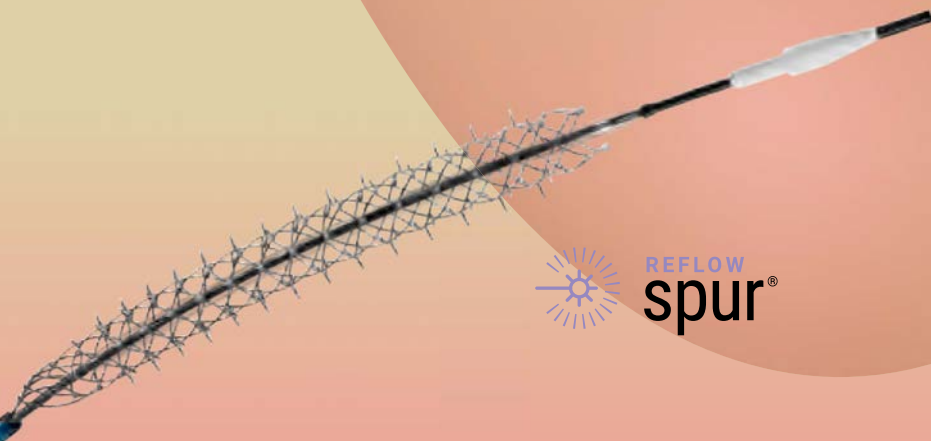
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Anmeldung



Bitte melden Sie sich über den nachfolgenden QR-Code oder Link an:

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Am Veranstaltungstag wird eine Einlasskontrolle stattfinden.
Bei vorhandener Anmeldung erhalten Sie:

- ein Namensschild und ein gedrucktes Programm
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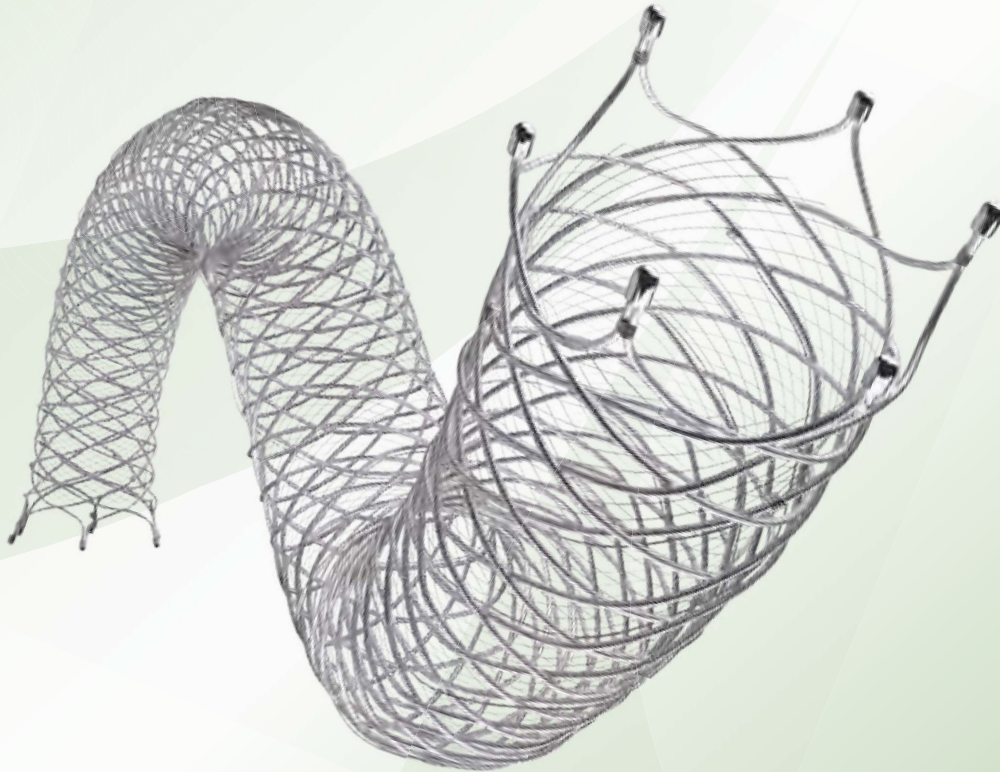
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1. Gore Medical Products Instructions for (IFU). W. L. Gore & Associates, Inc. Accessed May 20, 2025. <https://eifu.goremedical.com/>
2. W. L. Gore & Associates. Post-Market Registry of the GORE® VIABAHN® VBX Balloon Expandable Endoprosthesis Implanted in Peripheral Vessels (EXPAND). VBX 17-04. Published December 18, 2018. Updated February 16, 2024. Accessed May 20, 2025. <https://clinicaltrials.gov/study/NCT03720704>
3. W. L. Gore & Associates. Real-World Data Collection of the GORE® VIABAHN® VBX Balloon Expandable Endoprosthesis When Used as a Bridging Stent with Branched and Fenestrated Endografts in the Treatment of Aortic Aneurysms Involving the Renal-Mesenteric Arteries (EMBRACE). VBX 21-04. Published March 23, 2022. Updated January 1, 2025. Accessed May 20, 2025. <https://clinicaltrials.gov/study/NCT05143138>

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Programm Vormittag

Zeit	Titel	Referent:in / Vorsitz
08.00–08.25	Anmeldung	
08.30–08.35	Begrüßung	Michel Bosiers Drosos Kotelis Maria Schaumeier Marc Schindewolf
08.35–10.00	Aorta	Drosos Kotelis Jürg Schmidli
Das rupturierte Aneurysma	Fallvorstellung	Dimitrios Papazoglou
	How I do it	Edin Mujagic Alexander Zimmermann
	Lösung und Diskussion	Dimitrios Papazoglou
	Update Guidelines & Evidenz	Daniel Becker
Das infizierte Aneurysma	Fallvorstellung	Patrick Do
	How I do it	Florian Dick Thomas Wyss
	Lösung und Diskussion	Patrick Do
	Update Guidelines & Evidenz	Thomas Wyss
10.05–10.25	Kaffee-Pause	
10.30–12.00	pAVK	Michel Bosiers Konstantinos Stavroulakis
Claudicatio intermittens	Fallvorstellung	Aleksandra Tuleja
	How I do it	Lea Attias Pascal Kissling
	Lösung und Diskussion	Aleksandra Tuleja
	Updates Guidelines & Evidenz	Michel Bosiers
Kritische Extremitäten-ischämie	Fallvorstellung BTK	Nicolas Fattinger
	How I do it	Jörn Dopheide Konstantinos Stavroulakis
	Lösung und Diskussion	Nicolas Fattinger
	Updates Guidelines & Evidenz	Konstantinos Stavroulakis
12.05–12.55	Mittagspause	

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Programm Nachmittag

Zeit	Titel	Referent:in / Vorsitz
13.30–14.30	Male health	Maria Schaumeier Marc Schindewolf
Prostata- embolisation	Fallvorstellung	Nicolas Fattinger
	How I do it	Dominik Abt Lukas Hechelhammer
	Lösung und Diskussion	Nicolas Fattinger
	Updates Guidelines & Evidenz	Lukas Hechelhammer
Varikozele	Fallvorstellung	Alois Komarek
	Diskussion	Dominik Abt Lukas Hechelhammer
	Lösung, Updates Guidelines & Evidenz	Alois Komarek
14.35–14.55	Kaffeepause	
15.00–16.30	Challenging cases: artiell und venös	Sébastien Déglise Maani Hakimi
	Aorta	Silvan Jungi
	Periphere Intervention	Michel Bosiers
	Venöse Intervention	Marc Schindewolf
16.30–16.45	Fazit und Schlusswort	Michel Bosiers Drosos Kotelis Maria Schaumeier Marc Schindewolf

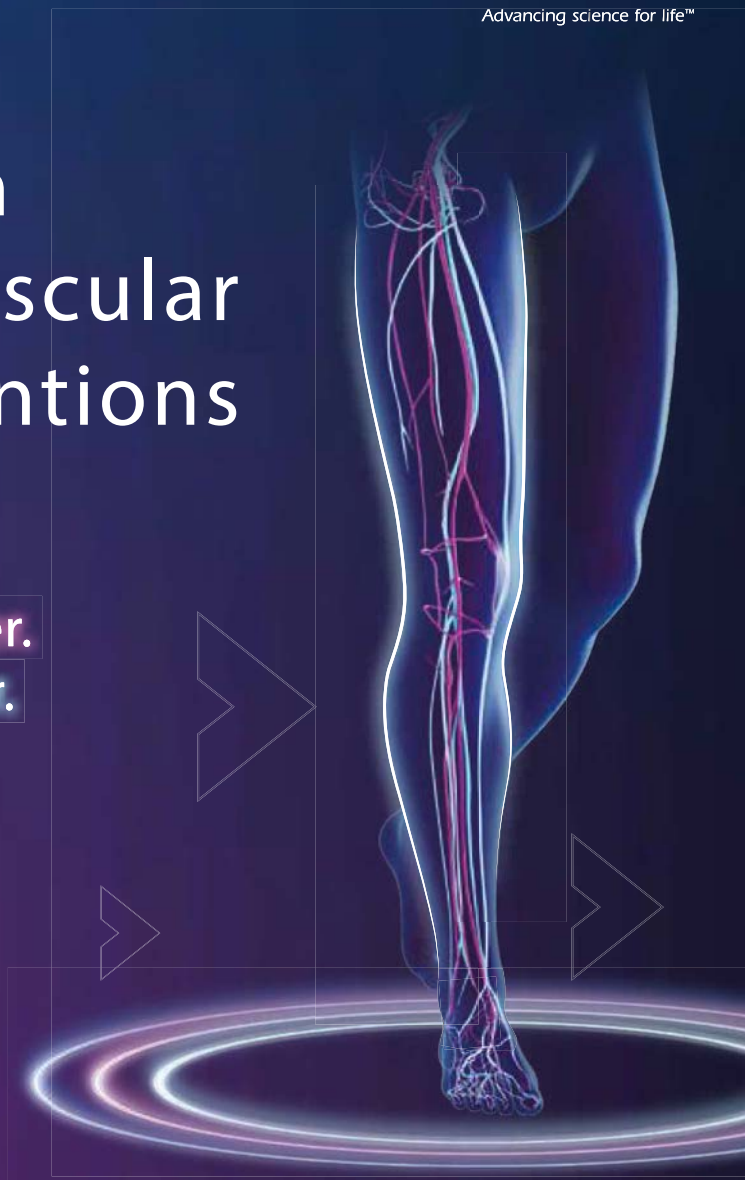
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Das Organisationsteam bedankt sich bei nachfolgenden Personen für Ihren aktiven Beitrag:

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1. Vogel J, Cheng C, Murphy E, et al. Fatigue Test Method to Evaluate the 50 Year Durability of Venous Stents. *Eur J Vasc Endovasc Surg.* 2024;68(4):521-528.

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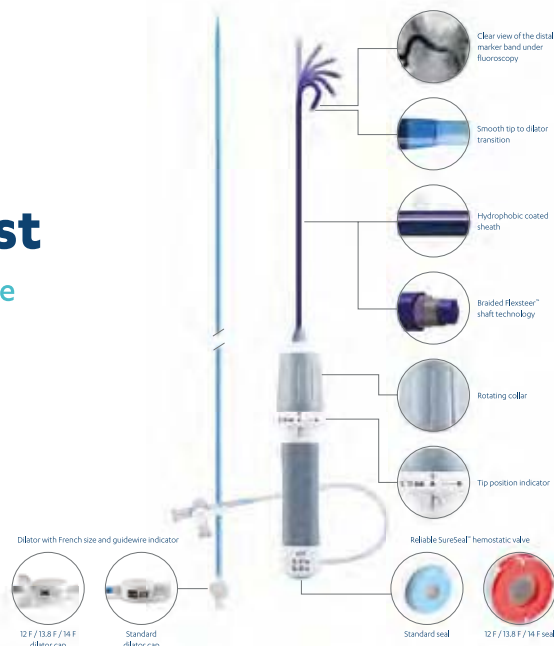
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1. Data on file; 2. De Gregorio M A, et al. Drug-coated balloon for the treatment of long-segment femoropopliteal artery disease: pooled analysis from the BIOLUX P-III SPAIN and BIOLUX P-III all-comers registry long lesion subgroup. J Vasc Interv Radiol. 2023;34:1707-1715. e7; 3. Brodmann M, et al. Real-world experience with a paclitaxel-coated balloon in critical limb ischemia: 24-month subgroup outcomes of BIOLUX P-III. JACC Cardiovasc Interv. 2020;13:2289-2299. 4. Tera G, et al. BIOLUX P-III Paseo-18 Lux all-comers registry: 24-month results in below-the-knee arteries. Cardiovasc Interv Radiol. 2021;44:10-18.

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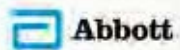
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