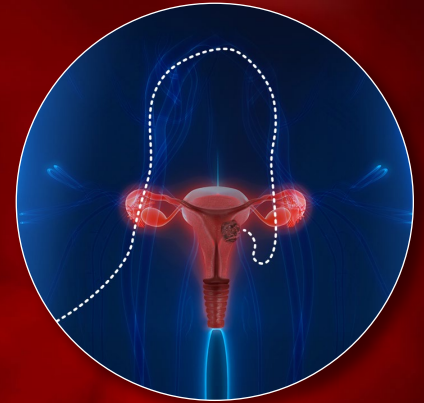


**DraKon™**  
Peripheral microcatheter

**SeQure®**  
Reflux control microcatheter

**3.0 Fr**

**Ultra high flow  
diagnostic microcatheter**



**Deliver up to  
900µm microspheres.**

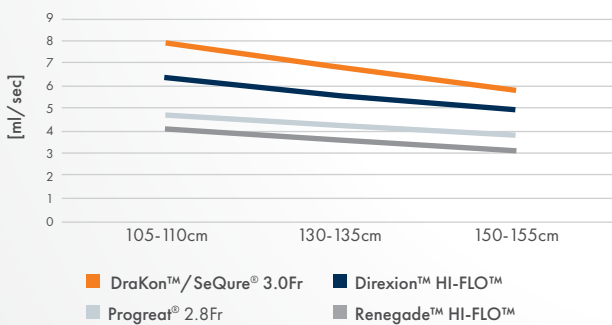
**LARGEST\* INNER DIAMETER (ID)  
MICROCATHETER**

- **0.031" (0.8mm) ID<sup>1</sup>** engineered to:
  - Reduce microcatheters clogging
  - Preserve microspheres integrity

**1200 PSI MAX INJECTION PRESSURE**

- Designed to **optimize flow rate**

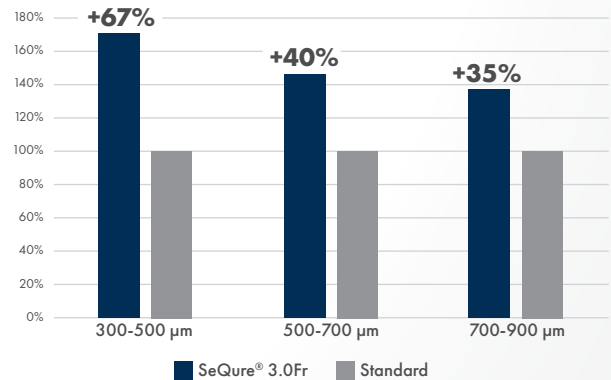
**Flow rates at maximum injection pressure\*\***



**DELIVER MORE<sup>5</sup> WITH CONFIDENCE  
USING SEQURE® 3.0 FR**

- More targeted embolization: injection flow rate **up to +67% higher** without reflux vs. standard microcatheters<sup>6</sup>

**Beads injection flow rate (ratio)**



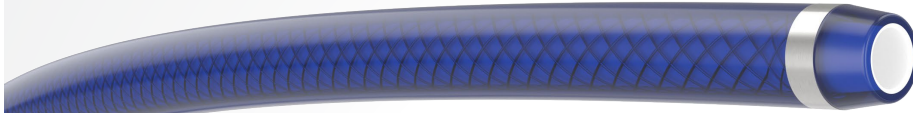
\* Amongst FDA cleared microcatheters in the 2.7Fr – 3.0 Fr class as of May 2021.  
 \*\* DraKon™ & SeQure® 3.0Fr are rated up to 1200PSI. Direxion™ HI-FLO™ is rated up to 1200PSI. Progreat® 2.8Fr is rated up to 900PSI. Renegade™ HI-FLO™ is rated up to 800PSI. Data taken from product IFU. Contrast media could be different between samples.

**DraKon™**  
Peripheral microcatheter

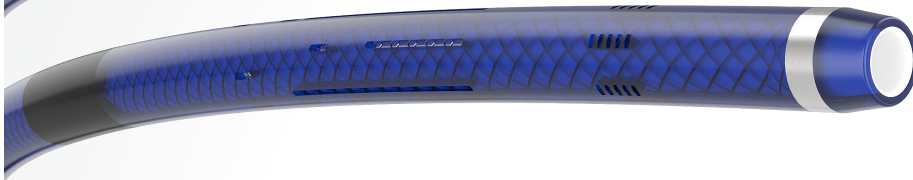
**SeQure®**  
Reflux control microcatheter

**3.0 Fr**

**Ultra high flow  
diagnostic microcatheter**



**DraKon™ 3.0 Fr**



**SeQure® 3.0 Fr**

**Specifications and Compatibilities**

	Length (cm)	Inner Diameter (in/mm)	Max Compatible Guidewire OD (in)	Recommended Guiding Catheter	Max Pressure (PSI)	Dead Space Volume (mL)	Flow Rate (mL/sec)
DraKon™	105	0.031" (0.8 mm)	0.025"	Min. 0.038" (0.97 mm GW compatible) and ID ≥ 1.10 mm	1200	0.74	7.9
	130					0.85	6.8
	155					0.95	5.8
SeQure®	105					0.74	7.9
	130					0.85	6.8
	155					0.95	5.8

Flow Rate with iodinated contrast media (iomeprol 300mg/mL, 37°C).

**Ordering Information**

	Product Reference	Order Number	Length (cm)	RO Markers	Max Embolic Compatibilities (Beads / Coils)	Hydrophilic Coating Length (cm)
DraKon™	DK30_105	238623	105	1	900 µm / Refer to embolic coil IFU	75
	DK30_130	238624	130	1		100
	DK30_155	238625	155	1		125
SeQure®	SQ30_XLB_105	238614	105	2	300 - 900 µm / Refer to embolic coil IFU	75
	SQ30_XLB_130	238615	130	2		100
	SQ30_XLB_155	238616	155	2		125

SeQure® is primarily designed for use with microspheres. SeQure® is not intended for delivery of non-spherical or gelatin sponge.



For more information, please contact us at **877-729-6679**  
or [customer.service-us@guerbet.com](mailto:customer.service-us@guerbet.com)  
[www.guerbet.com/en-us](http://www.guerbet.com/en-us)

1. DraKon™ & SeQure® IFU. 2. Data on File DR-1800178, test done only with SeQure® 2.4 Fr and 2.8 Fr. 3. Data on File TR-057. Bench Tests results may not necessarily be indicative of clinical performance.

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DraKon™ & SeQure® microcatheters are medical devices intended for use by interventional radiologists and interventional oncologists for the infusion of contrast media into all peripheral vessels and for drug infusion in intra-arterial therapy, and infusion of embolic materials. They should not be used in cerebral vessels.

DraKon™ is a trademark of Guerbet Group or its affiliates. SeQure® is a registered trademark of Guerbet Group or its affiliates. Trademarks are the property of their respective owners and are used herein solely for informational purposes.

CAUTION: US Federal Law restricts this device to sale by or on the order of a physician. Illustrations for information purposes – not indicative of actual size or clinical outcome.

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