# TUBING KIT

Use with SOFIA® Flow Plus Aspiration Catheter and Gomco® 405 Aspiration Pump



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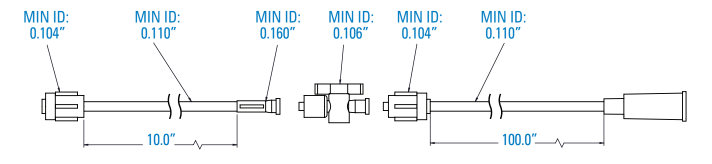
Use with SOFIA® Flow Plus Aspiration Catheter and Gomco® 405 Aspiration Pump

### TUBING KIT

### LARGE ID | HIGH FLOW | OPTIMIZED ASPIRATION

| PRODUCT | PROXIMAL TUBING | DISTAL TUBING | TOTAL TUBING | MINIMUM    | MINIMUM 1-WAY | MINIMUM    |
|---------|-----------------|---------------|--------------|------------|---------------|------------|
| CODE    | LENGTH          | LENGTH        | LENGTH       | TUBING ID  | STOPCOCK ID   | LUER ID    |
| MVTK110 | 100 in /        | 10 in /       | 110 in /     | 0.110 in / | 0.106 in /    | 0.104 in / |
|         | 254 cm          | 25 cm         | 279 cm       | 2.8 mm     | 2.7 mm        | 2.6 mm     |

One kit per box, includes proximal tubing, 1-way stopcock, and distal tubing





#### LARGE BORE 0.106" ID STOPCOCK

controls flow without limiting aspiration power<sup>1</sup>

## LARGE 0.110" ID BRAIDED KINK RESISTANT TUBING

provides consistent high flow aspiration<sup>1</sup>

#### STANDARD PROXIMAL CONNECTOR

offers secure connection to vacuum pump<sup>1</sup>

<sup>1</sup>TR17-226, TR18-226



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MicroVention Worldwide Innovation Center 35 Enterprise Aliso Viejo, CA 92656 PH 714.247.8000 PH 1.800.990.8368 For professional use. CAUTION: Federal (United States) law restricts this device to sale by or on the order of a physician. See instructions for use for full prescribing information, including indications, contraindications, warnings, precautions and adverse events.

The SOFIA® Flow Plus Aspiration Catheter with the Gomco® 405 Aspiration Pump and MicroVention® Tubing Kit is intended for use in the revascularization of patients with acute ischemic stroke secondary to intracranial large vessel occlusive disease (within the internal carotid, middle cerebral – M1 and M2 segments, basilar, and vertebral arteries) within 8 hours of symptom onset. Patients who are ineligible for intravenous tissue plasminogen activator (IV t-PA) or who fail IV t-PA therapy are candidates for treatment.