

Introducing the Chartis® Precision XL Catheter

The Next Generation in Collateral Ventilation Assessment



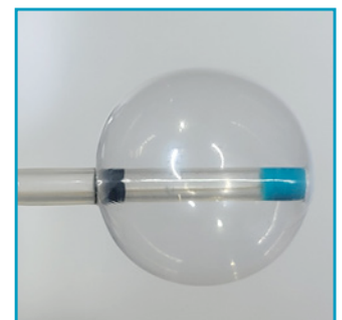
- 1 Updated Balloon Profile**
 - Designed to facilitate positioning during procedures. The design aims to help clinicians navigate the distal carina and may help address the challenge of clogging.
- 2 Expanded Range**
 - Designed to enable a broader range of targets, including shorter lobar airways.
- 3 Modified Insertion Force**
 - A thinner shaft diameter is designed to improve movement through the bronchoscope.
- 4 Streamlined Setup**
 - Comes pre-packaged with all necessary accessories and pre-attached stopcocks to simplify preparation.
- 5 Scope Compatibility**
 - Incorporates a built-in strain relief to help prevent kinking through any 2.8 mm bronchoscope working channel.

Updated Balloon Profile:

- The maximum diameter of the balloon is reached toward the distal tip designed to help in the assessment of shorter airways.
- Forward inflating balloon is designed to reduce contact with distal carina and minimize mucus clogging.
- The balloon can reach a maximum diameter of 13mm to enable large airway assessments.



Chartis Precision Catheter Balloon



Chartis G3 Balloon

Ease of Use:



- Pre-attached stopcocks reduce assembly steps.



- A thinner shaft diameter is designed to improve movement through the bronchoscope.

Ordering Information

- SKU Number: CHR-CA-1.0

International Important Safety Information: The Chartis® System is indicated for use by bronchoscopists during a diagnostic bronchoscopy in adult patients with Chronic Obstructive Pulmonary Disease (COPD) and emphysema in a bronchoscopy suite. The system, composed of the Chartis Catheter and Chartis Console, is designed to measure pressure and flow in order to calculate resistance to airflow and quantify collateral ventilation in isolated lung compartments. The Chartis Catheter is used through the working channel of a bronchoscope and connects to the Chartis Console. The Chartis Console is capital equipment that is reusable and displays the patient information. The Chartis System is contraindicated in the presence of active infection or major bleeding diathesis. There are no known interfering substances. Use is restricted to a trained physician. Prior to use, please reference the Chartis System Instructions for Use/ User Manual for more information on indications, contraindications, warnings, all precautions, and adverse events.

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