

# Certificate of Registration

QUALITY MANAGEMENT SYSTEM - ISO 13485:2016

This is to certify that:

**VitalPath**  
9409 Science Center Drive  
New Hope  
Minnesota  
55428  
USA

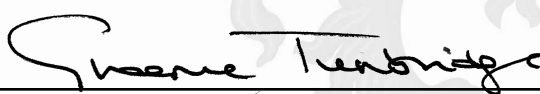
Holds Certificate Number: MD 823386

and operates a Quality Management System which complies with the requirements of ISO 13485:2016 for the following scope:

Vital Path is a Catheter Manufacture, including Design, Development, and Manufacture for OEM Medical Device Industries.

Transfer from SARA Registrar  
Certificate number: SARA-2009-CA-0097-D

For and on behalf of BSI:

  
Graeme Tunbridge, Senior Vice President Global Regulatory & Quality

Original Registration Date: 2025-06-20

Latest Revision Date: 2025-06-20

Effective Date: 2025-06-20

Expiry Date: 2027-06-25

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...making excellence a habit.™

Certificate No: MD 823386

| Location  | Registered Activities   |
|---|---|
| VitalPath<br>9409 Science Center Drive<br>New Hope<br>Minnesota<br>55428<br>USA | Vital Path is a Catheter Manufacture, including Design, Development, and Manufacture for OEM Medical Device Industries. |
| VitalPath<br>1758 Terrance Drive<br>Roseville<br>Minnesota<br>55113<br>USA      | Vital Path is a Catheter Manufacture, including Design, Development, and Manufacture for OEM Medical Device Industries. |



Original Registration Date: 2025-06-20  
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This certificate was issued electronically and remains the property of BSI and is bound by the conditions of contract.  
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Printed copies can be validated at [www.bsigroup.com/ClientDirectory](http://www.bsigroup.com/ClientDirectory)

Information and Contact:  
BSI Group The Netherlands B.V., John M. Keynesplein 9, 1066 EP Amsterdam, The Netherlands | Tel: +31 20 3460 780  
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Americas Headquarters: 12950 Worldgate Drive, Suite 800, Herndon, VA 20170-6007 USA