

Certificate of Registration

QUALITY MANAGEMENT SYSTEM - ISO 13485:2016

This is to certify that:

AxoGen Corporation
13631 Progress Blvd, Suite 400
Alachua
Florida
32615
USA


Holds Certificate No:

FS 608440

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

Receipt, Handling, Storage and Distribution of Medical Devices related to nerve repair - AxoGuard Nerve Connector and AxoGuard Nerve Protector.

For and on behalf of BSI:


Chief Operating Officer Assurance - Americas

Original Registration Date: 2014-06-03

Latest Revision Date: 2018-07-05

Effective Date: 2017-06-03

Expiry Date: 2020-06-02

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Certificate No: **FS 608440**

Location	Registered Activities
AxoGen Corporation 13631 Progress Blvd, Suite 400 Alachua Florida 32615 USA	Receipt, Handling, Storage and Distribution of of Medical Devices related to nerve repair- AxoGuard Nerve Connector and AxoGuard Nerve Protector.
AxoGen Corporation 300 Boone Road Burlison Texas 76028 USA	Receipt, Handling, Storage and Distribution of of Medical Devices related to nerve repair- AxoGuard Nerve Connector and AxoGuard Nerve Protector.



Original Registration Date: 2014-06-03

Effective Date: 2017-06-03

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This certificate remains the property of BSI and shall be returned immediately upon request.
An electronic certificate can be authenticated [online](#). Printed copies can be validated at www.bsigroup.com/ClientDirectory
To be read in conjunction with the scope above or the attached appendix.

Americas Headquarters: BSI Group America Inc., 12950 Worldgate Drive, Suite 800, Herndon, VA 20170-6007 USA
A Member of the BSI Group of Companies.