



Certificate

SQS herewith certifies that the company named below has a management system which meets the requirements of the standard specified below.



Spineart SA
Chemin du Pré-Fleuri 3
1228 Plan-les-Ouates
Switzerland

Scope of certification

According to appendix

Field of activity

**Design, manufacturing and sales of sterile
and non-sterile spine medical devices**

Normative base

**EN ISO 13485:2016 Medical devices –
Quality Management System**

Validity 03.10.2020 – 02.10.2023
Issue 03.10.2020

Reg. no. H31786


A. Grisard, President SQS


F. Müller, CEO SQS

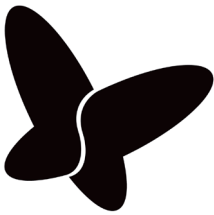


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Swiss Association for Quality
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Bernstrasse 103, 3052 Zollikofen, Switzerland





Spineart SA Chemin du Pré-Fleuri 3 1228 Plan-les-Ouates Switzerland

Central Function	Field of activity	Scope(s)	Norm / Revision	Reg. no.	Validity
Spineart SA Chemin du Pré-Fleuri 3 1228 Plan-les-Ouates Switzerland	Design, manufacturing and sales of sterile and non-sterile spine medical devices	19	EN ISO 13485:2016	H31786	03.10.2020 02.10.2023

Locations	Field of activity	Scope(s)	Norm / Revision	Reg. no.	Validity
Alpes CN SAS 19 route des Marais Z.A.E. de Findrol 74250 FILLINGES France	Machining and manufacturing of Spineart Medical Devices	19	EN ISO 13485:2016	H31786	03.10.2020 02.10.2023
SLI SAS Bâtiment ABC3 Archamps Technopole 80, Rue Douglas Engelbart 74160 ARCHAMPS France	Maintenance activity of Spineart instrumentation, logistic and storage management of Spineart Medical Devices	19	EN ISO 13485:2016	H31786	03.10.2020 02.10.2023
Alpes CN SAS Bâtiment ABC3 Archamps Technopole 80, Rue Douglas Engelbart 74160 ARCHAMPS France	Control, assembly, cleaning and packaging of Spineart Medical Devices in clean room	19	EN ISO 13485:2016	H31786	03.10.2020 02.10.2023

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