



# CERTIFICATE



This is to certify that the company

## Precision Spine, Inc.

2050 Executive Drive  
Pearl, MS 39208  
United States of America

with the organizational units/sites as listed in the annex

has implemented and maintains a **Quality Management System**.

Scope of certification:

Design and develop, manufacture, and distribution of sterile and non-sterile spinal implants and surgical instruments. Service and repair of surgical instruments.

- AUS (a), BRA, USA (a,b,c,d)

Through an audit, documented in a report, performed by DQS Medizinprodukte GmbH, it was verified that the management system fulfills the requirements of the following standard:

## ISO 13485 : 2016

including applicable country-specific regulatory requirements, as indicated below the scope (full references of abbreviations are listed in the annex)

|                              |                  |
|------------------------------|------------------|
| Certificate registration no. | 31622620 MDSAP16 |
| Certificate unique ID        | 1000262466       |
| Effective date               | 2026-05-11       |
| Expiry date                  | 2028-06-16       |
| Frankfurt am Main            | 2026-05-11       |



## DQS Medizinprodukte GmbH

Heinrich von Mettenheim  
Managing Director



August-Schanz-Straße 21, 60433 Frankfurt am Main,  
Tel. +49 (0) 69 95427-300, [info-med@dqs.de](mailto:info-med@dqs.de)

**DQS Medizinprodukte GmbH is recognized under the Medical Devices Single Audit Program.**

Visit <https://www.dqs.de/en/customer-database/> to validate this certificate.

The validity of the certification can only be verified by the QR-code.



**Annex to certificate**  
**Certificate registration No.: 31622620 MDSAP16**  
**Certificate unique ID: 1000262466**  
**Effective date: 2026-05-11**

## **Precision Spine, Inc.**

2050 Executive Drive  
Pearl, MS 39208  
United States of America

### **Audited site**

**31622546**  
**Precision Spine, Inc.**  
2050 Executive Drive  
Pearl, MS 39208  
United States of America

### **REPs FEI No.: site scope and country-specific requirements**

Design and develop, manufacture, and  
distribution of sterile and non-sterile spinal  
implants and surgical instruments. Service and  
repair of surgical instruments.  
**- AUS (a), BRA, USA (a,b,c,d)**  
**REPs FEI No.: F007699**

**31622621**  
**Precision Spine, Inc.**  
300 Interpace Parkway Building C, 2nd Floor,  
Parsippany, NJ 07054  
United States of America

Design and develop, manufacture, and  
distribution of sterile and non-sterile spinal  
implants and surgical instruments. Service and  
repair of surgical instruments.  
**- AUS (a), BRA, USA (a,b,c,d)**  
**REPs FEI No.: F007702**



**Annex to certificate**  
**Certificate registration No.: 31622620 MDSAP16**  
**Certificate unique ID: 1000262466**  
**Effective date: 2026-05-11**

## **Precision Spine, Inc.**

2050 Executive Drive  
Pearl, MS 39208  
United States of America

### **Full references of country-specific requirements of MDSAP participating Regulatory Authorities**

| <b>Abbreviation</b> | <b>Jurisdiction</b> | <b>Reference</b>                                                                                                                                                                                                                         |
|---------------------|---------------------|------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| AUS                 | Australia           | (a) Therapeutic Goods (Medical Devices) Regulations 2002, Schedule 3, Part 1 – Full Quality Assurance Procedure<br>(b) Therapeutic Goods (Medical Devices) Regulations 2002, Schedule 3, Part 4 – Production Quality Assurance Procedure |
| BRA                 | Brazil              | RDC ANVISA n. 665/2022<br>RDC ANVISA n. 551/2021<br>RDC ANVISA n. 67/2009                                                                                                                                                                |
| CND                 | Canada              | Medical Devices Regulations – Part 1- SOR 98/282<br>Medical Devices Regulations – Part 1.1 – SOR 98/282 (as applicable)                                                                                                                  |
| JPN                 | Japan               | MHLW Ministerial Ordinance 169, Article 4 to Article 68<br>Japan PMD Act (as applicable)                                                                                                                                                 |
| USA                 | United States       | (a) 21 CFR Part 803<br>(b) 21 CFR Part 806<br>(c) 21 CFR Part 807<br>(d) 21 CFR Part 820<br>(e) 21 CFR Part 821                                                                                                                          |



# CERTIFICATE

This is to certify that the company

## Precision Spine, Inc.

2050 Executive Drive  
Pearl, MS 39208  
United States of America

with the organizational units/sites as listed in the annex

has implemented and maintains a **Quality Management System**.

Scope of certification:

Design and develop, manufacture, and distribution of sterile and non-sterile spinal implants and surgical instruments. Service and repair of surgical instruments.

- AUS (a), BRA, USA (a,b,c,d)

Through an audit, documented in a report, performed by DQS Medizinprodukte GmbH, it was verified that the management system fulfills the requirements of the following standard:

## ISO 13485 : 2016

including applicable country-specific regulatory requirements, as indicated below the scope (full references of abbreviations are listed in the annex)

|                              |                  |
|------------------------------|------------------|
| Certificate registration no. | 31622620 MDSAP16 |
| Certificate unique ID        | 1000262466       |
| Effective date               | 2026-05-11       |
| Expiry date                  | 2028-06-16       |
| Frankfurt am Main            | 2026-05-11       |



## DQS Medizinprodukte GmbH

Heinrich von Mettenheim  
Managing Director



August-Schanz-Straße 21, 60433 Frankfurt am Main,  
Tel. +49 (0) 69 95427-300, [info-med@dqs.de](mailto:info-med@dqs.de)

**DQS Medizinprodukte GmbH is recognized under the Medical Devices Single Audit Program.**

Visit <https://www.dqs.de/en/customer-database/> to validate this certificate.

The validity of the certification can only be verified by the QR-code.



**Annex to certificate**  
**Certificate registration No.: 31622620 MDSAP16**  
**Certificate unique ID: 1000262466**  
**Effective date: 2026-05-11**

## **Precision Spine, Inc.**

2050 Executive Drive  
Pearl, MS 39208  
United States of America

### **Audited site**

**31622546**  
**Precision Spine, Inc.**  
2050 Executive Drive  
Pearl, MS 39208  
United States of America

### **REPs FEI No.: site scope and country-specific requirements**

Design and develop, manufacture, and  
distribution of sterile and non-sterile spinal  
implants and surgical instruments. Service and  
repair of surgical instruments.

**- AUS (a), BRA, USA (a,b,c,d)**

**REPs FEI No.: F007699**

**31622621**  
**Precision Spine, Inc.**  
300 Interpace Parkway Building C, 2nd Floor,  
Parsippany, NJ 07054  
United States of America

Design and develop, manufacture, and  
distribution of sterile and non-sterile spinal  
implants and surgical instruments. Service and  
repair of surgical instruments.

**- AUS (a), BRA, USA (a,b,c,d)**

**REPs FEI No.: F007702**



**Annex to certificate**  
**Certificate registration No.: 31622620 MDSAP16**  
**Certificate unique ID: 1000262466**  
**Effective date: 2026-05-11**

## **Precision Spine, Inc.**

2050 Executive Drive  
Pearl, MS 39208  
United States of America

### **Full references of country-specific requirements of MDSAP participating Regulatory Authorities**

| <b>Abbreviation</b> | <b>Jurisdiction</b> | <b>Reference</b>                                                                                                                                                                                                                         |
|---------------------|---------------------|------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| AUS                 | Australia           | (a) Therapeutic Goods (Medical Devices) Regulations 2002, Schedule 3, Part 1 – Full Quality Assurance Procedure<br>(b) Therapeutic Goods (Medical Devices) Regulations 2002, Schedule 3, Part 4 – Production Quality Assurance Procedure |
| BRA                 | Brazil              | RDC ANVISA n. 665/2022<br>RDC ANVISA n. 551/2021<br>RDC ANVISA n. 67/2009                                                                                                                                                                |
| CND                 | Canada              | Medical Devices Regulations – Part 1- SOR 98/282<br>Medical Devices Regulations – Part 1.1 – SOR 98/282 (as applicable)                                                                                                                  |
| JPN                 | Japan               | MHLW Ministerial Ordinance 169, Article 4 to Article 68<br>Japan PMD Act (as applicable)                                                                                                                                                 |
| USA                 | United States       | (a) 21 CFR Part 803<br>(b) 21 CFR Part 806<br>(c) 21 CFR Part 807<br>(d) 21 CFR Part 820<br>(e) 21 CFR Part 821                                                                                                                          |