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INTRODUCTION TO UNiD™ ASI

UNiD™ Adaptive Spine Intelligence combines service, software, and patient-specific implants providing surgeons with a revolutionary approach to achieving better outcomes.

This clinical brief provides an overview of the clinical rationale, components, and clinical applications of UNiD™ Adaptive Spine Intelligence.

Clinical Issues, Results, and Solutions

This brief explores the clinical issues and related results or solutions with regard to sagittal alignment, surgical precision, O.R. efficiency, and the incidence of rod fracture.

UNiD™ Adaptive Spine Intelligence

Fundamental components of UNiD™ ASI systems-based platform are detailed: LAB, TEK, HUB, and the Iterative Virtuous Cycle.

UNiD™ LAB engineers provide spinopelvic parameters and surgical simulations based on surgeon input and preferences. UNiD™ TEK patient-specific implants are approved by the surgeon via the UNiD™ HUB. Clinical judgment and experience are required to properly use the software.

CLINICAL BENEFITS OF SAGITTAL ALIGNMENT

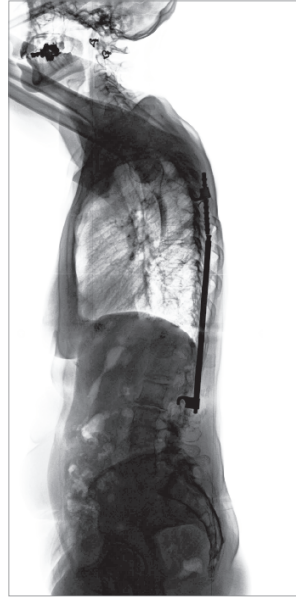
Sagittal alignment is the most dominant radiographic predictor of patient outcomes.^{1,2}

Achieving harmonious alignment of key spinopelvic parameters, such as the sagittal vertical axis (SVA), pelvic incidence/lumbar lordosis mismatch (PI-LL), and pelvic tilt (PT), is a key goal of spinal deformity surgery.^{1,2}

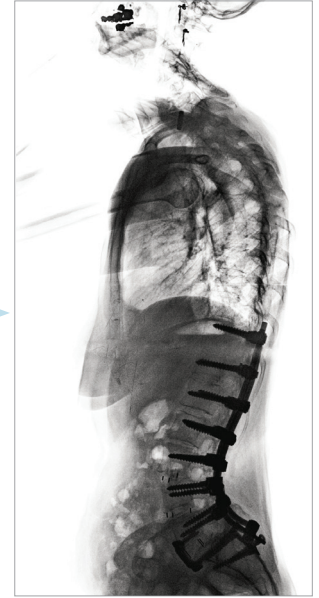
Patients possessing postoperative spinopelvic parameters within normative ranges exhibit improved patient outcomes scores.^{1,2}

One of the risks of not achieving optimal alignment is revision spinal surgery.³

Clinical Transition to Patient-Specific Planning and Alignment



Harrington Rod



Patient-Specific Alignment

Key Clinical Issues



Sagittal re-alignment and clinical outcomes are directly linked.¹

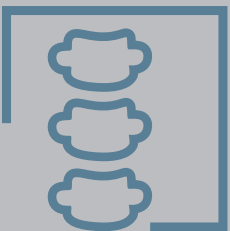
62% of patients remained sagittally malaligned after surgery.⁴

UNiD™ Clinical Results

SVA IMPROVED

81% achieved normative SVA values

Significant improvement for all key parameters postoperatively⁶



10x greater risk of developing adjacent segment disease when postoperative Δ PI-LL $\geq 10^\circ$ for 1 to 3 level degenerative constructs.⁵

PI-LL < 10

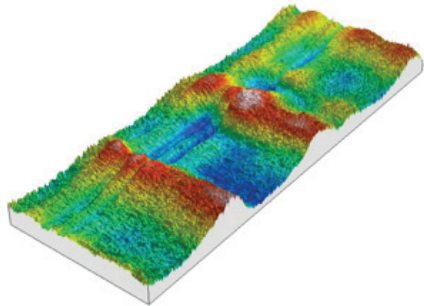
Achieved in all cases

All patients had postoperative PI-LL of less than 10° ⁶

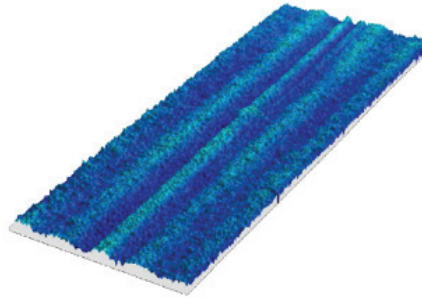
POTENTIAL FOR PRECISION AND EFFICIENCY

UNiD™ Adaptive Spine Intelligence gives surgeons the tools to more precisely achieve their surgical goals, increasing efficiency in both the preoperative planning phase as well as in the operating room.

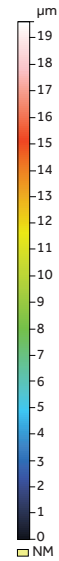
3-D Rod Surface Analysis*



Manually bent



Smoothly contoured UNiD™ Patient-Specific Rod



Strength

Intraoperative rod contouring using a French Bender significantly reduced fatigue strength.⁷

Each UNiD™ Rod is industrially produced in a lab for the highest level of control. The resulting rods are smoothly contoured, aligned with the case plan, and notch-free.

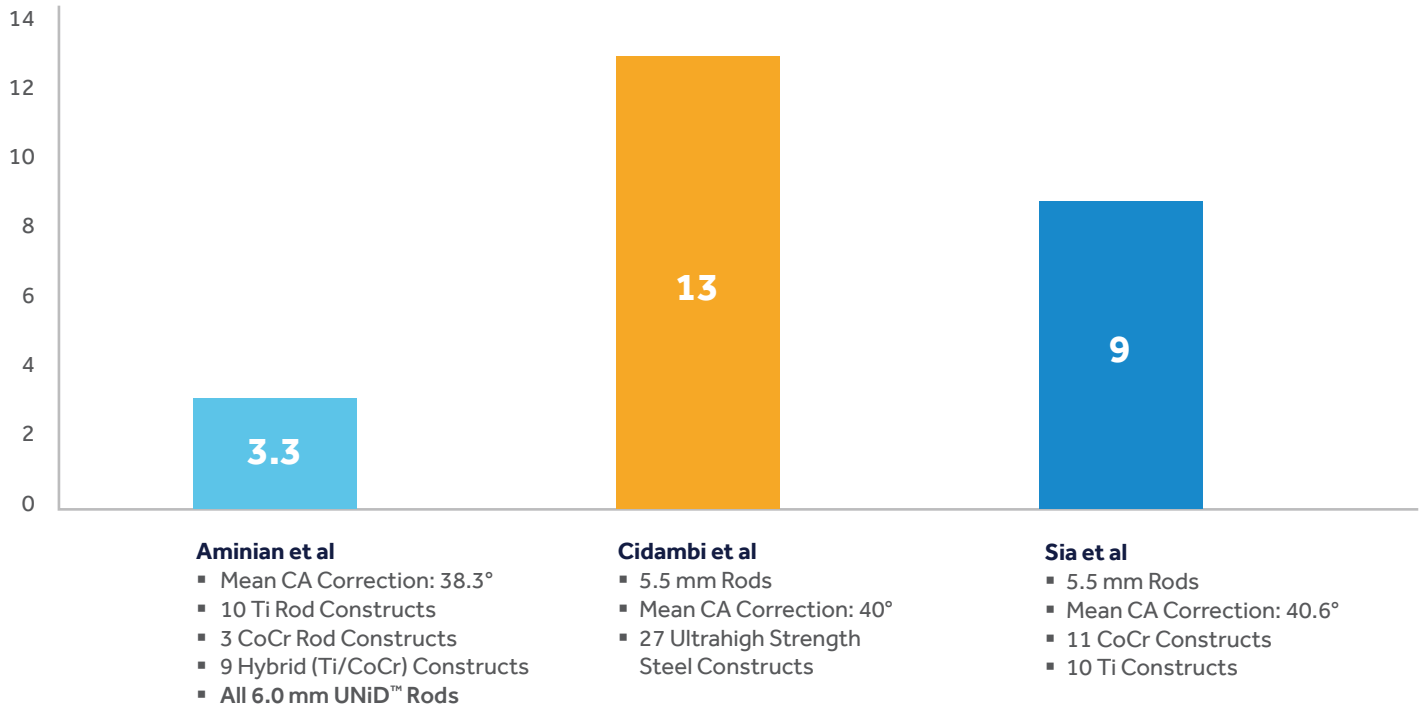
See back cover for the risks of UNiD™ rods.

*3-D Optical Profilometer - Non-Contact Measurement and Analysis

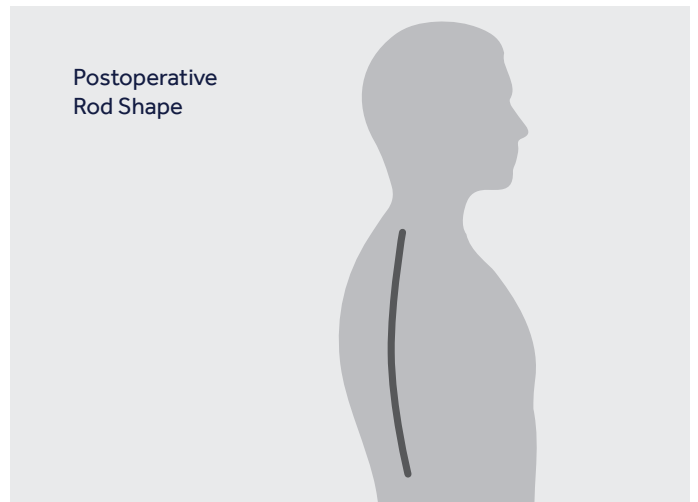
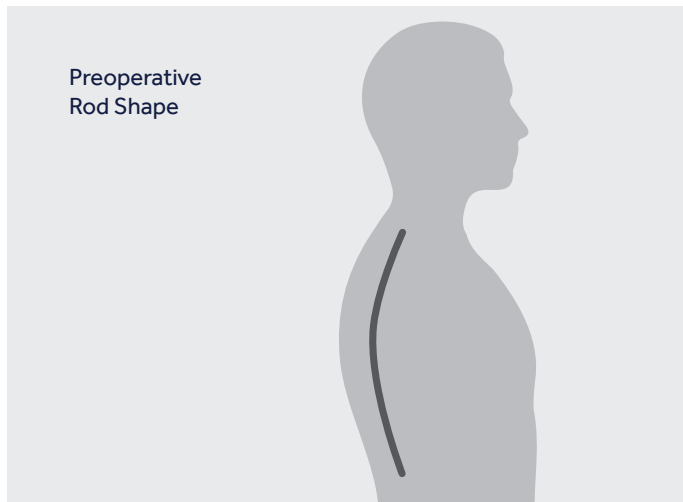
LESS ROD FLATTENING

Four studies assessed the difference in rod contour after implantation, as well as the Cobb Angle (CA) correction they were able to achieve. Change in rod contour (flattening) was assessed by modeling the difference in concave rod deflection between the preoperative and implanted rods.

Mean Concave Rod Deflection (mm) in AIS Patients



Flattening of a Concave Rod after Implantation



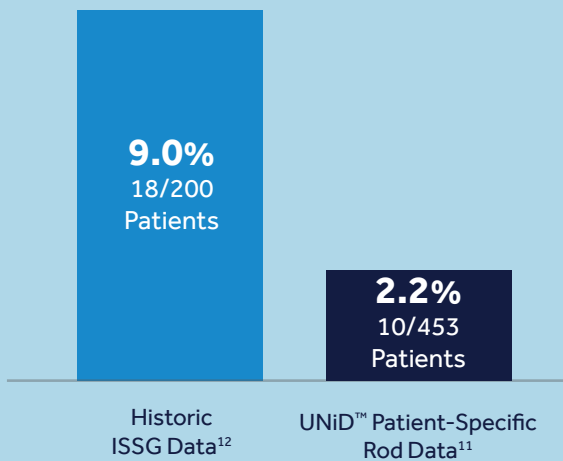
REDUCED INCIDENCE OF ROD FRACTURE

Evaluation of postoperative data indicates a reduction in the rod fracture rate:

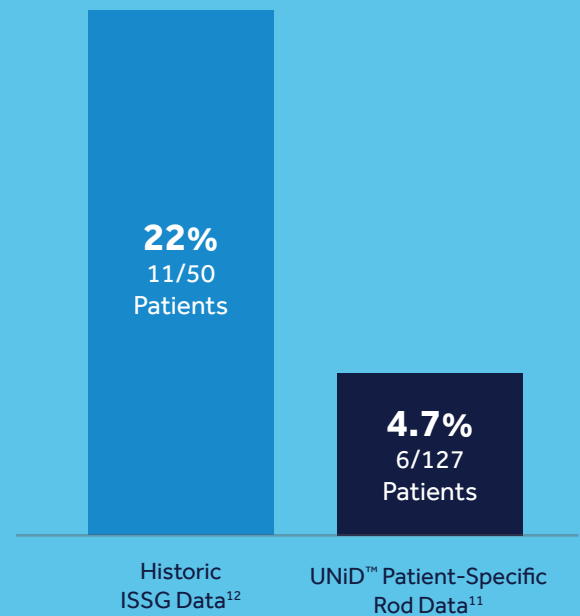
In adult deformity cases (> 5 levels) at least one year after surgery, UNiD Rods had a fracture rate of 10/453 patients, or 2.2%. In a subset of International Spine Study Group (ISSG) data with the same parameters, 18/200 (9.0%) of adult deformity patients experienced rod fractures.^{11,12}

When patients from the same two studies underwent a pedicle subtraction osteotomy (PSO) in the procedure, the rate is **reduced by 79%**, an improvement over the 22.0% rod fracture rate associated with procedures involving a PSO.^{11,12}

Rod Fracture Rates in Adult Deformity Cases



Rod Fracture Rate in Cases involving a PSO

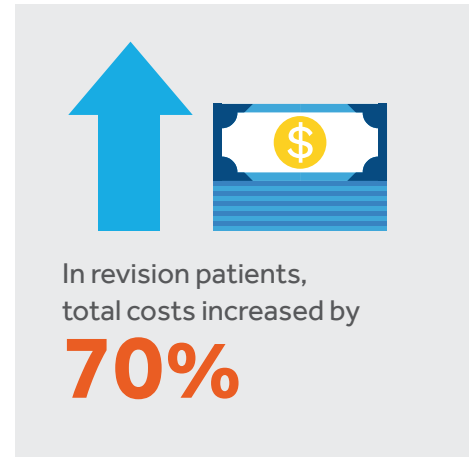
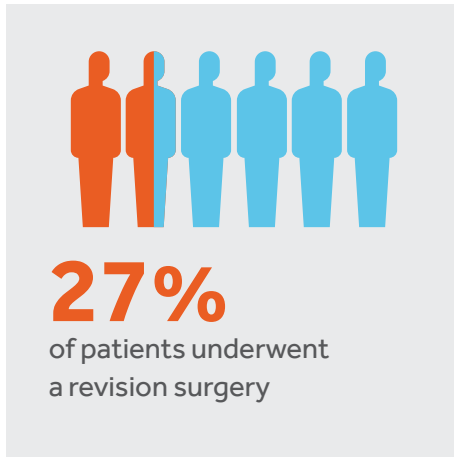


Risks associated with these spinal implants include loosening, disassembly, bending, and/or breakage of components. A successful result is not always achieved in every surgical case. This fact is especially true in spinal surgery where many extenuating circumstances may compromise the results.

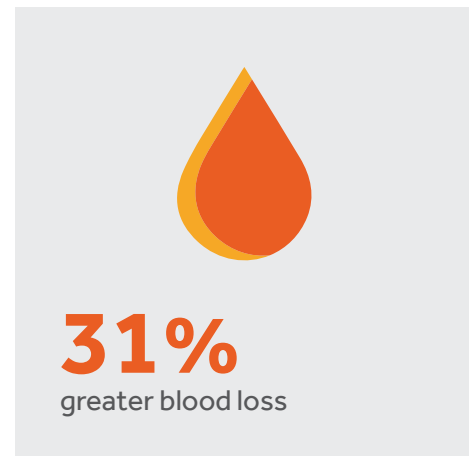
COST TO THE HEALTHCARE SYSTEM

Healthcare providers are increasingly concerned about the cost of complications, readmissions, and reoperations associated with adjacent segment disease and implant failure, such as spinal rod fracture.

In a study¹³ of **484 consecutive adult spinal deformity patients** with an average **follow-up of 4.8 years**:



In a study¹⁴ of consecutive adult scoliosis patients, **126 primary patients** were compared to **124 revision patients**. In this study, revision patients had:





ENABLES SURGEONS

- Plan + Execute: Patient-specific **sagittal alignment**
- Reduce risk of rod breakage¹¹
- Correct deformities with less rod flattening⁸



ENABLES HOSPITALS

- Provide **personalized** spinal solutions
- Reconfirm community leadership in treatment options
- Reduce the risk of revision surgeries due to adjacent segment disease^{4,5} and rod fracture¹¹



UNiD™ ADAPTIVE SPINE INTELLIGENCE

Plan. Execute. Analyze.

The surgeon-centric platform provides a **planning** service staffed by biomedical engineers, precise intraoperative **execution** with personalized solutions, and insightful **analytics** of surgical results with the ultimate goal of improving clinical outcomes.

UNiD™ LAB

Engineering Services

The UNiD™ LAB is our team of biomedical engineers. A dedicated LAB engineer measures patient images, providing the surgeon with detailed analysis, spinopelvic parameters, and normative alignment values.

The LAB engineer then uses sophisticated HUB software to collaborate with the surgeon in developing surgical strategies and simulating surgical case plans. These surgical case plans are based on the latest scientific literature, proprietary algorithms, case strategy, and the surgeon's preferences.

UNiD™ HUB

Interactive Portal

The UNiD™ HUB provides the surgeon with easy access to all components of UNiD™ ASI.

Accessible via desktop or a mobile device, the HIPAA compliant UNiD™ HUB facilitates surgical plan simulations, surgical workflow, outcome analytics, and communications with the UNiD™ LAB. (For more detail, see page 13).

UNiD™ TEK

Personalized Implants

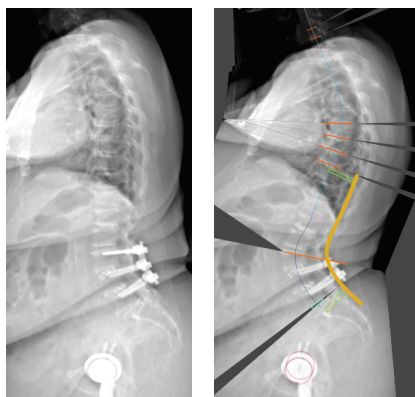
UNiD™ TEK is a suite of technologies enhanced by the UNiD™ HUB platform and UNiD™ LAB Service.

Each TEK implant aligns with the surgical case plan, providing intraoperative plan confirmation.

PLAN PRE-OP PLANNING SERVICES

1 Imaging Analysis

The UNiD™ cycle begins for each patient with the rapid identification of spinopelvic parameters using standard radiographs and, if available, X-Rays (Calibrated X-Rays), EOS, MRI, or CT images. Integration with PACS and communication via the UNiD™ HUB supports the goal of improved patient workflows.

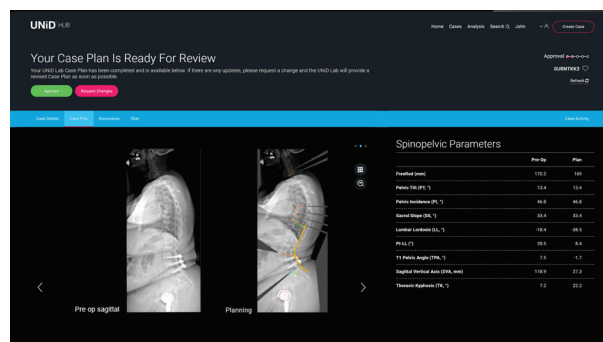


Pre-Op

Plan

2 Case Simulation

The UNiD™ LAB engineer uses sophisticated UNID™ HUB and UNiD™ Spine Analyzer Software to simulate multiple surgical strategies for the surgeon based on the surgical strategy and surgeon preferences. The proprietary algorithms also include the latest scientific literature and proprietary predictive planning models.



EXECUTE INTRA-OP SERVICES

3 Personalized Implants

UNiD™ TEK is a suite of technologies enhanced by the UNiD™ HUB platform and UNiD™ LAB Service. UNiD™ Rods are manufactured following surgical planning performed by a surgeon for a given patient. The UNiD™ Rod implants are fabricated with advanced in-house manufacturing technology. UNiD™ Rods are FDA cleared in the U.S. for compatibility with the CD Horizon™ Solera™ Spinal System.



4 Case Support

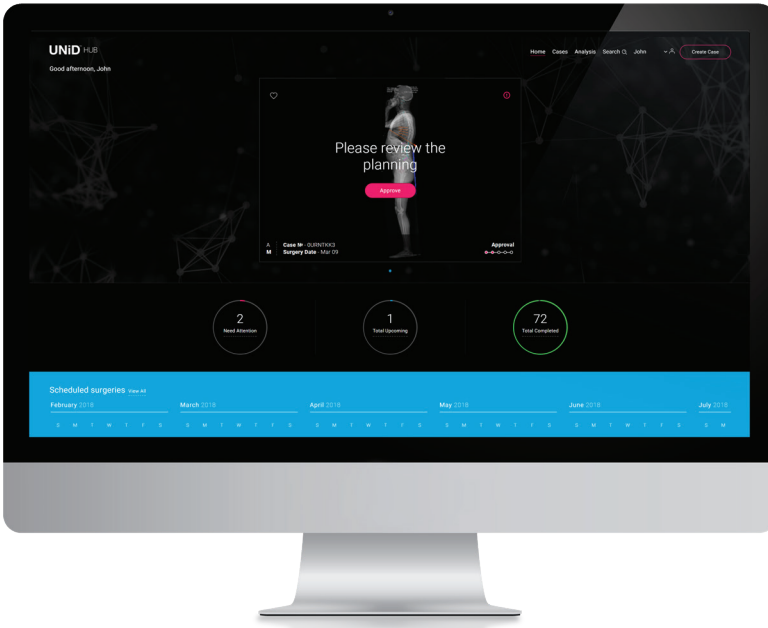
UNiD™ Rods are aligned with the preoperative surgical plan, helping to guide the surgery and provide intraoperative confirmation.



Risks associated with these spinal implants include loosening, disassembly, bending, and/or breakage of components.

UNiD™ HUB INTERACTIVE PORTAL

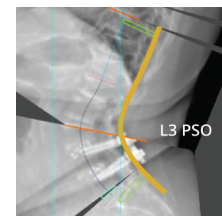
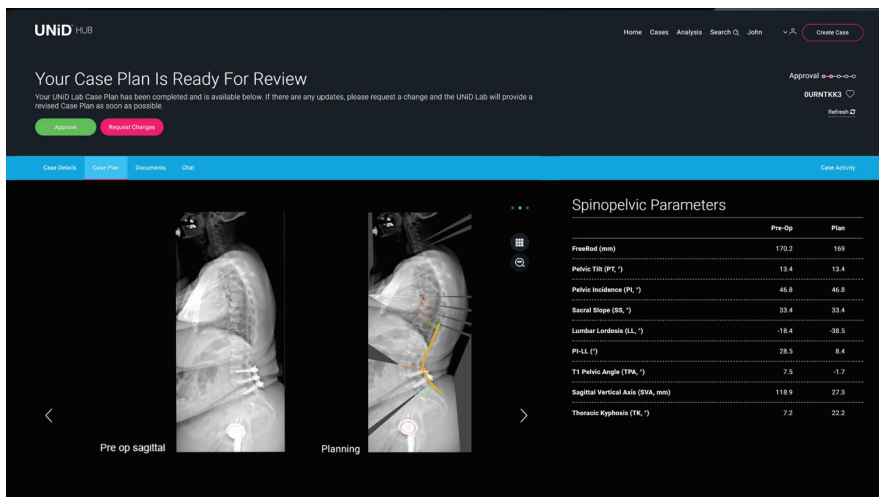
Facilitates communication with the planning service of the UNiD™ LAB and allows the surgeon to leverage the power of Adaptive Spine Intelligence.



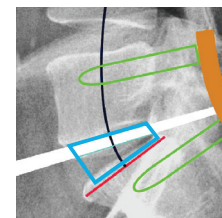
Accessible via desktop or a mobile device, the HIPAA compliant UNiD™ HUB is a centralized location for review and approval of surgical plans as well as a valuable resource for organizing and analyzing surgical results.

PLAN

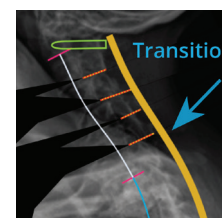
Simulation and Plan Approval



Osteotomies



Cage Planning

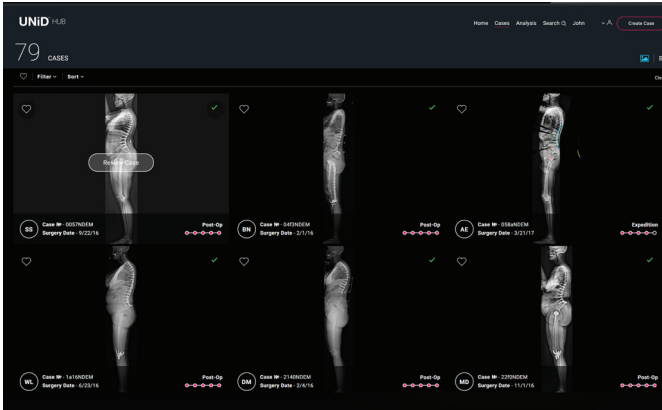


Rod Planning

UNiD™ LAB engineers provide spinopelvic parameters and surgical simulations based on surgeon input and preferences. UNiD™ TEK patient-specific implants are approved by the surgeon via the UNiD™ HUB.

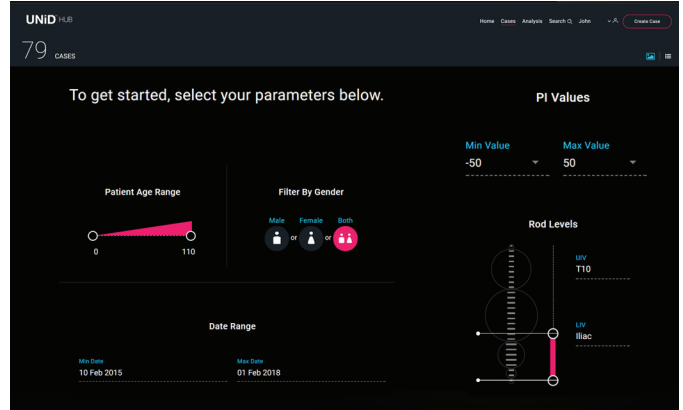
ANALYZE

Database of Surgeries



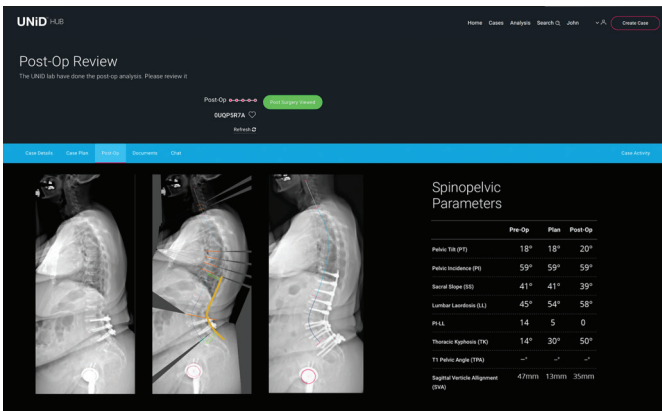
Parameters and images for all surgeries are well organized and easily accessible.

Multiple Search Options



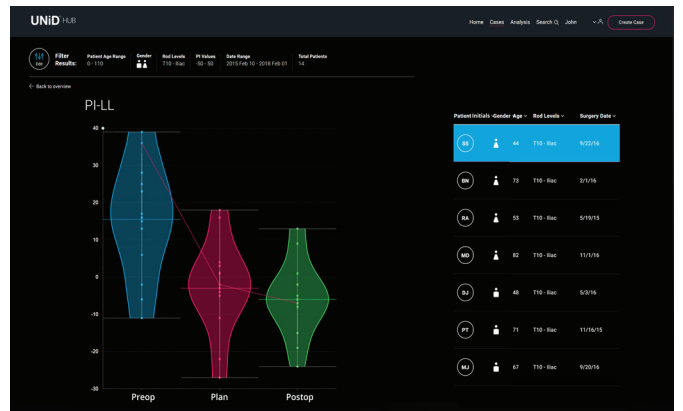
Ability to query along multiple parameters allows focus on a particular subset of cases.

Review Postoperative Results



Compare pre-op, plan, and post-op images and parameters for individual patients or use the Outcome Analysis Toolkit for a broader review of the database.

Outcome Analysis Toolkit



Advanced analytics and visualization of outcome data provides valuable insight and opportunity for improvement. Multiple output options are available for use in presentations, reports, and clinical studies.

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IMPORTANT PRODUCT INFORMATION

CD HORIZON™ SYSTEM INDICATIONS

The CD Horizon™ Spinal System with or without Sextant™ instrumentation is intended for posterior, non-cervical fixation as an adjunct to fusion for the following indications: degenerative disc disease (DDD - defined as back pain of discogenic origin with degeneration of the disc confirmed by history and radiographic studies), spondylolisthesis, trauma (i.e. fracture or dislocation), spinal stenosis, curvatures (i.e. scoliosis, kyphosis, or lordosis), tumor, pseudarthrosis, and/or failed previous fusion. Except for hooks, when used as an anterolateral thoracic/lumbar system, the CD Horizon™ Spinal System titanium, cobalt chrome, and stainless steel implants may also be used for the same indications as an adjunct to fusion.

With the exception of DDD, the CD Horizon™ Legacy™ 3.5mm rods and associated components may be used for the aforementioned indications in skeletally mature patients as an adjunct to fusion. The 3.5mm rods may be used for the specific pediatric indications noted below.

When used for posterior non-cervical pedicle screw fixation in pediatric patients, the CD Horizon™ Spinal System titanium, cobalt chrome, and stainless steel implants are indicated as an adjunct to fusion to treat progressive spinal deformities (i.e. scoliosis, kyphosis, or lordosis) including idiopathic scoliosis, neuromuscular scoliosis, and congenital scoliosis. Additionally, the CD Horizon™ Spinal System is intended to treat pediatric patients diagnosed with the following conditions: spondylolisthesis/spondylolysis, fracture caused by tumor and/or trauma, pseudarthrosis, and/or failed previous fusion. These devices are to be used with autograft and/or allograft. Pediatric pedicle screw fixation is limited to a posterior approach.

The CD Horizon™ PEEK rods are intended to provide posterior supplemental fixation when used with an interbody fusion cage for patients diagnosed with DDD. These DDD patients may also have up to Grade 1 spondylolisthesis or retrolisthesis at the involved level. This device is intended for 1-2 level use in the lumbosacral spine (L2 – S1) in skeletally mature patients. The device is intended for use with an interbody fusion cage at the instrumented level and is not intended for stand-alone use.

The CD Horizon™ Spire™ plate is a posterior, single-level, non-pedicle supplemental fixation device intended for use in the non-cervical spine (T1-S1) as an adjunct to fusion in skeletally mature patients. It is intended for plate fixation/attachment to spinous processes for the purpose of achieving supplemental fixation in the following conditions: DDD (as previously defined), spondylolisthesis, trauma, and/or tumor.

In order to achieve additional levels of fixation, the CD Horizon™ Spinal System rods may be connected to the Vertex™ Reconstruction System with the Vertex™ rod connector. Refer to the Vertex™ Reconstruction System package insert for a list of the Vertex™ indications of use.

RISKS

All of the possible adverse events associated with spinal fusion surgery without instrumentation are possible. With instrumentation, a listing of potential adverse events includes:

- Early or late loosening of any or all of the components.
- Disassembly, bending, or breakage of any or all of the components.
- Post-operative change in spinal curvature, loss of correction, height, or reduction.
- Infection.

PASS LP SYSTEM DESCRIPTION

The internal fixation devices are composed of screws, hooks, rods, plates, cross links, connection and locking devices. The range of different sizes and shapes of the implants allows the surgeon to adapt to the pathology and morphology of each of his patients. The implants are manufactured in titanium alloy Ti-6Al-4V ELI conforming to ISO 5832-3 specifications and ASTM F136 specifications, with the exception of

the rods intended for in situ bending which are manufactured in non-alloyed titanium (CP titanium) conforming to ISO 5832-2 specifications and ASTM F67 specifications and the CoCr rods which are manufactured in cobalt chrome alloy Co-Cr28Mo6 conforming to ISO 5832-12 specifications and ASTM F1537 specifications. The Patient Specific Rod has been designed and manufactured for one specific patient. The Patient Specific Rod should be used during surgery for this patient only and should not be reused (single use only). Refer to the surgical technique brochure for additional information. If this Patient Specific Rod does not perform as intended, use the standard PASS LP rod to complete the surgery. Under no circumstances are the implants reusable.

INDICATIONS

The PASS LP spinal systems include a pedicle system intended to provide immobilization and stabilization of spinal segments in skeletally mature patients as an adjunct to fusion in the treatment of the following acute and chronic instabilities or deformities of thoracic, lumbar, and sacral spine:

- Fractures.
- Dislocation.
- Failed previous fusion (pseudarthrosis).
- Spinal stenosis.
- Degenerative spondylolisthesis with objective evidence of neurological impairment.
- Spinal deformations such as scoliosis or kyphosis.
- Loss of stability due to tumors.

The PASS LP spinal systems are also indicated for pedicle screw fixation for the treatment of severe spondylolisthesis (Grades 3 and 4) of the L5-S1 vertebra in skeletally mature patients receiving fusion by autogenous bone graft having implants attached to the lumbar and sacral spine (L3 to sacrum) with removal of the implants after the attainment of a solid fusion.

The PASS LP also include hooks and rods and sacral/iliac screws indicated for degenerative disc disease (ddd) defined as back pain of discogenic origin with degeneration of the disc confirmed by history and radiographic studies, spondylolisthesis, trauma (i.e., fracture or dislocation), spinal stenosis, deformities or curvatures (i.e., scoliosis, kyphosis, and/or lordosis), tumor, pseudarthrosis and failed previous fusion.

Except for rod plates, when used for posterior non-cervical pedicle screw fixation in pediatric patients, the PASS LP spinal system implants are indicated as an adjunct to fusion to treat adolescent idiopathic scoliosis. The PASS LP spinal system is intended to be used with allograft and/or autograft. Pediatric pedicle screw fixation is limited to a posterior approach.

WARNING: The safety and effectiveness of this device has not been established for use as part of a growing rod construct. This device is only intended to be used when definitive fusion is being performed at all instrumented levels.

RISKS

In addition to the risks associated with surgery of the spine without instrumentation, a number of possible undesirable effects may occur with instrumented surgery (including but not limited to):

- Detachment, deformation, mobilization, slipping, breakage of one or all of the components.
- Pain due to the surgery, the fracture, deformation and or migration of an implant.
- Fracture of the pedicle during insertion of a pedicular screw.
- Postoperative loss of correction and/or reduction of the spine, partial or total loss of the corrections achieved.

Please refer questions on the risks and benefits of UNiD™ ASI to unidsupport@medtronic.com.

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Please see the package insert for the complete list of indications, warnings, precautions, and other important medical information.



Consult instructions for use at this website
www.medtronic.com/manuals.

Note: Manuals can be viewed using a current version of any major internet browser. For best results, use Adobe Acrobat® Reader with the browser.