










Cost Benefit of Implementation of Risk Stratification Models for Adult Spinal Deformity Surgery

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Peter G. Passias, MD¹ , Tyler K. Williamson, DO² , Nicholas A. Kummer, BS¹ , Ferran Pellisé, MD PhD³, Virginie Lafage, PhD⁴ , Renaud Lafage, MSc⁵ , Miguel Serra-Burriel, PhD⁶, Justin S. Smith, MD PhD⁷, Breton Line, BS⁸ , Shaleen Vira, MD⁹, Jeffrey L. Gum, MD¹⁰ , Sleiman Haddad, MD³, Francisco Javier Sánchez Pérez-Grueso, MD¹¹, Andrew J. Schoenfeld, MD MSc¹², Alan H. Daniels, MD¹³, Dean Chou, MD¹⁴, Eric O. Klineberg, MD¹⁵, Munish C. Gupta, MD¹⁶ , Khaled M. Kebaish, MD¹⁷ , Michael P. Kelly, MD¹⁸, Robert A. Hart, MD¹⁹, Douglas C. Burton, MD²⁰, Frank Kleinstück, MD²¹, Ibrahim Obeid, MD²², Christopher I. Shaffrey, MD²³, Ahmet Alanay, MD²⁴, Christopher P. Ames, MD¹⁴, Frank J. Schwab, MD⁴, Richard A. Hostin Jr., MD²⁵, Shay Bess, MD⁸, and International Spine Study Group²⁶

¹ Departments of Orthopedic and Neurological Surgery, NYU Langone Orthopedic Hospital, New York Spine Institute, NY, NY, USA

² Department of Orthopaedic Surgery, University of Texas Health San Antonio, San Antonio, TX, USA

³ Spine Surgery Unit, Vall d'Hebron Hospital, Barcelona, Spain

⁴ Department of Orthopaedics, Lenox Hill Hospital, New York, NY, USA

⁵ Department of Orthopaedics, Hospital for Special Surgery, New York, NY, USA

⁶ Center for Research in Health and Economics, Universitat Pompeu Fabra, Barcelona, Spain

⁷ Department of Neurosurgery, University of Virginia Medical Center, Charlottesville, VA, USA

⁸ Denver International Spine Center, Presbyterian St. Luke's/Rocky Mountain Hospital for Children, Denver, CO, USA

⁹ Department of Orthopedic Surgery, UT Southwestern Medical Center, Dallas, TX, USA

¹⁰ Norton Leatherman Spine Center, Louisville, KY, USA

¹¹ Spine Surgery Unit, Hospital Universitario La Paz, Madrid, Spain

¹² Department of Orthopaedic Surgery, Brigham and Women's Hospital, Harvard Medical School, Boston, MA, USA

¹³ Department of Orthopaedic Surgery, Warren Alpert School of Medicine, Brown University, Providence, RI, USA

¹⁴ Department of Neurosurgery, University of California, San Francisco, CA, USA

¹⁵ Department of Orthopedic Surgery, University of California Davis, Sacramento, CA, USA

¹⁶ Department of Orthopaedic Surgery, Washington University in St. Louis, Missouri, USA

¹⁷ Department of Orthopaedic Surgery, The Johns Hopkins Medical Institutions, Baltimore, MD, USA

¹⁸ Department of Orthopaedic Surgery, Rady Children's Hospital, San Diego, CA, USA

¹⁹ Department of Orthopaedic Surgery, Swedish Neuroscience Institute, Seattle, WA, USA

²⁰ Department of Orthopaedic Surgery, University of Kansas Medical Center, Kansas City, KS, USA

²¹ Spine Center Division, Department of Orthopedics and Neurosurgery, Schulthess Klinik, Zürich, Switzerland

²² Spine Surgery Unit, Bordeaux University Hospital, Bordeaux, France

²³ Spine Division, Departments of Neurosurgery and Orthopaedic Surgery, Duke University School of Medicine, Durham, NC, USA

²⁴ Department of Orthopedics and Traumatology, Acibadem University, Istanbul, Turkey

²⁵ Department of Orthopaedic Surgery, Baylor Scoliosis Center, Dallas, TX, USA

²⁶ Rocky Mountain Scoliosis and Spine, Denver, CO, USA

Corresponding Author:

Peter G. Passias, MD, New York Spine Institute, Division of Spinal Surgery, Departments of Orthopedic and Neurological Surgery, NYU Langone Medical Center, Orthopedic Hospital – NYU School of Medicine, 301 East 17th St, New York, NY 10003, USA.

Email: Peter.Passias@nyumc.org



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Abstract

Study Design/Setting: Retrospective cohort study.

Objective: Assess the extent to which defined risk factors of adverse events are drivers of cost-utility in spinal deformity (ASD) surgery.

Methods: ASD patients with 2-year (2Y) data were included. Tertiles were used to define high degrees of frailty, sagittal deformity, blood loss, and surgical time. Cost was calculated using the Pearl Diver registry and cost-utility at 2Y was compared between cohorts based on the number of risk factors present. Statistically significant differences in cost-utility by number of baseline risk factors were determined using ANOVA, followed by a generalized linear model, adjusting for clinical site and surgeon, to assess the effects of increasing risk score on overall cost-utility.

Results: By 2 years, 31% experienced a major complication and 23% underwent reoperation. Patients with ≤ 2 risk factors had significantly less major complications. Patients with 2 risk factors improved the most from baseline to 2Y in ODI. Average cost increased by \$8234 per risk factor ($R^2 = .981$). Cost-per-QALY at 2Y increased by \$122,650 per risk factor ($R^2 = .794$). Adjusted generalized linear model demonstrated a significant trend between increasing risk score and increasing cost-utility ($r^2 = .408, P < .001$).

Conclusions: The number of defined patient-specific and surgical risk factors, especially those with greater than two, were associated with increased index surgical costs and diminished cost-utility. Efforts to optimize patient physiology and minimize surgical risk would likely reduce healthcare expenditures and improve the overall cost-utility profile for ASD interventions.

Level of evidence: III

Keywords

adult spinal deformity, spine, cost-utility, risk stratification, major complications, mechanical failure, surgical factors

Introduction

The prevalence of surgical intervention for adult spinal deformity has increased over the last two decades.^{1,2} These procedures are associated with an increased risk of complications, mechanical failure, readmissions and revisions. Such adverse events also impact patient-reported outcomes and long-term healthcare expenditures.³⁻⁶ The combination of higher cost and decreased clinical improvement in the setting of a complicated postoperative course can significantly impact the cost-effectiveness profile of these procedures.

Numerous risk factors for complications, mechanical failure and revisions following ASD surgery have been described previously, as well as their impact on clinical outcomes.⁷⁻¹⁷ The most representative and parsimonious cohort, as contextualized in the work of *Pellise et al*, include Lowest Instrumented Vertebrae (LIV) at pelvis, frailty, sagittal deformity (global sagittal alignment [SVA], lordosis gap [PILL], and T1 sagittal tilt), blood loss, and surgical time.⁷ The impact of these individual parameters on cost-utility of interventions for ASD, however, is not well characterized.

In this context, we sought to better understand the extent to which these defined risk factors of adverse events following ASD surgery are drivers of poor cost-utility. In the current investigation we examined the complication rates, clinical outcomes, initial and overall cost of patients who underwent ASD surgery based on their number of baseline risk factors. Each of these components play an integral role in the cost-utility of surgical intervention for spinal deformity. Therefore, ultimately, we used

these outcomes to evaluate the influence of risk factors on cost-utility following surgery. We hypothesized that an increase in the number of risk factors at baseline would be associated with reduced cost-utility for ASD surgeries at 2 years.

Materials and Methods

Study Design and Inclusion Criteria

We queried an adult spinal deformity database to identify patient records eligible for inclusion in this analysis. As described in previous publications, this database consists of 14 distinct centers across the United States that contribute consecutively enrolled, consented patient data with Institutional Review Board approval.¹⁸⁻²⁰ Patients that were included in the present study also had complete 2-year (2Y) Health Related Quality of Life (HRQL) data and radiographic measurements.

Data Collection and Radiographic Parameters

We abstracted demographic (age, body mass index [BMI], biological sex, Charlson Comorbidity Index [CCI], frailty [as measured by the *Passias et al* ASD-mFI), surgical (levels fused, operative time, length of stay, surgical approach, performance of decompressions and osteotomies), and clinical (complications, reoperations, DRG codes) data.^{21,43} Complication assessments were made based on review of imaging, patient reports, and clinical follow-up data. These forms were submitted by individual surgeons and centers and imparted into the database by a

standardized committee of study coordinators. Patient-reported outcome measures (Oswestry Disability Index [ODI], Short Form-36 [SF-36]) were collected at baseline and follow-up intervals (6 weeks, 6 months, 1 year, and 2 years).

Full length free-standing lateral spine radiographs (EOS or 36-inch cassette if unavailable) were collected and assessed at baseline and follow-up. Radiographic images were analyzed using SpineView® (ENSAM, Laboratory of Biomechanics, Paris, France) software.²²⁻²⁴ Spinopelvic radiographic parameters measured were Pelvic Tilt (PT), Pelvic Incidence (PI), Sagittal Vertical Axis (SVA), Thoracic Kyphosis (TK, T4-12), Lumbar Lordosis (LL, L1-S1), T1 Sagittal Tilt (T1-SPi) and mismatch between Pelvic Incidence and Lumbar Lordosis (PI-LL).

Complication Assessment

Mechanical complications were defined as any complication related to the implant, including implant prominence, implant malposition, painful implant, implant failure, interbody dislocation, screw nerve impingement, screw fracture, rod dislocation, and rod fracture. Mechanical and radiographic complications were classified as major if involving invasive intervention or causing prolonged or permanent morbidity or mortality. Proximal junctional kyphosis (PJK) was defined by a PJK angle of $<-10^\circ$ and a PJK angle difference of $<-10^\circ$ from baseline at any time point up to two years. Proximal junctional failure (PJF) was defined using the criteria of Lafage et al.: a PJK angle of $<-28^\circ$ and a difference in PJK angle of $<-22^\circ$ from baseline at any follow-up time point up to 2 years.³⁹

Previous Risk Stratification Criteria

Risk factors used in this study were defined based on the prior work of Pellise et al. and included: Lowest Instrumented Vertebrae (LIV) at pelvis, frailty, sagittal deformity (global sagittal alignment [SVA], lordosis gap [in this study, PI-LL], and T1 sagittal tilt [T1-SPi]), blood loss, and surgical time.^{7,25} Patients within the highest tertiles, sagittal deformity measures, blood loss, and surgical time, as well as Frail by the ASD-mFI, were defined as possessing the risk factor of interest for the purposes of this work. An incremental risk score adding the number of risk factors (range from 0 to 5) was generated for each patient.

Utility Calculation

Utility data was calculated converting ODI to SF-6D based on a previously published conversion methodology.²⁶⁻²⁸ The utilities were then transformed into Quality Adjusted Life Years (QALYs) and characterized as QALYs gained by comparing baseline point estimates to 2-year results. Quality adjusted life years were discounted at an annual 3% rate as recommended by the World Health Organization to account for decline in function associated with aging.

Table 1. Cohort Breakdown by Number of Risk Factors.

# Risk Factors	0	1	2	3	4	5	Total
Total	81	129	180	165	128	41	724
Pelvis	0	85	157	164	128	41	575
High frailty	0	18	55	76	101	41	291
High deformity	0	11	71	114	115	41	352
High blood loss	0	3	32	73	92	41	241
High surgical time	0	12	45	68	76	41	242

Cost Calculation

PearlDiver data was utilized to calculate national average Medicare costs using job order cost accounting (“charge analysis”). PearlDiver is one of the most comprehensive datasets with access to Medicare claims. We used mean costs associated with procedures based on 2018 adult spinal deformity diagnosis-related groups. We also accounted for costs associated with the occurrence of complications and comorbidities (CC), major complications and comorbidities (MCC), and revisions according to CMS.gov manual definitions, per previously published work.³⁰ Two-year reimbursement consisted of a standardized estimate using regression analysis of Medicare pay-scales for all services rendered within a 30-day window, including estimates regarding costs of postoperative complications, outpatient healthcare encounters, revisions and medical related readmissions. We used a multivariable approach, accounting for surgical approach, CC, MCC, length of stay (LOS), revisions, and death, to calculate cost per Quality Adjusted Life Year (QALY) at 2-year follow-up.^{29,30}

Statistical Analysis

We compared demographics, surgical details, complication rates, clinical improvement, and cost-utility (cost per QALY) at two years between cohorts based on the number of risk factors present at baseline. Number of risk factors were treated as a count variable. Linear regression analysis demonstrated correlation with increasing risk factors and clinical outcomes, complication rates, initial and overall cost. Statistically significant differences in cost-utility by number of baseline risk factors were determined using ANCOVA, controlling for baseline disability. A generalized linear model, adjusting for clinical site, surgeon, and age of patient, was performed using logit link and gamma distribution to assess the effects of increasing risk score on overall cost-utility.

Statistical Sub-Analysis

The cohort was further divided into four sub-cohorts by the High Deformity and Frailty variables. A risk score was generated out of the three remaining surgical factors and assessed against outcomes in each frailty/deformity sub-cohort. Multivariable logistic regression analyses and log linear models accounting for frailty and baseline deformity examined the effect of increasing

Table 2. Baseline Characteristics by Number of Risk Factors.

# Risk Factors	0	1	2	3	4	5	P-value
Baseline Characteristics							
Age	42.1 ± 17.8	59.6 ± 13.9	63.1 ± 10.8	64.4 ± 9.7	64.5 ± 9.0	63.7 ± 7.8	<.001
Gender	78%	85%	82%	73%	71%	68%	.016
Body Mass index (kg/m ²)	23.8 ± 3.7	27.0 ± 6.4	27.5 ± 5.4	28.1 ± 5.7	30.3 ± 5.7	31.4 ± 6.5	<.001
Charlson comorbidity index	.6 ± 1.1	1.4 ± 1.3	1.9 ± 1.7	1.9 ± 1.7	2.6 ± 1.9	2.7 ± 1.8	<.001
ASD frailty index	.15 ± 0.1	.28 ± 0.1	.31 ± 0.1	.36 ± 0.1	.45 ± 0.1	.49 ± 0.1	<.001
Osteoporosis	2%	12%	19%	17%	22%	20%	.003
Baseline radiographic parameters							
SVA (mm)	-1.6 ± 34.0	22.9 ± 46.7	53.3 ± 58.9	90.0 ± 68.1	115.4 ± 66.8	137.6 ± 61.3	<.001
PI-LL (°)	-3.7 ± 14.1	6.1 ± 14.4	13.3 ± 18.5	23.9 ± 19.1	29.0 ± 18.5	36.3 ± 17.7	<.001
T1 sagittal tilt (°)	-5.4 ± 2.5	-4.3 ± 3.8	-2.5 ± 5.4	.5 ± 6.3	3.0 ± 6.2	4.1 ± 6.3	<.001
Surgical details							
Number of levels fused	8.6 ± 4.1	9.0 ± 4.2	10.6 ± 4.8	11.8 ± 4.0	12.4 ± 3.8	13.8 ± 3.8	<.001
Operative time (min)	281 ± 97	362 ± 132	417 ± 154	489 ± 178	533 ± 181	563 ± 108	<.001
Estimated blood loss (mL)	748 ± 464	852 ± 601	1337 ± 1031	1963 ± 1126	2880 ± 1816	3765 ± 2178	<.001
Length of stay (days)	5.9 ± 2.4	6.7 ± 3.9	7.6 ± 4.4	8.8 ± 4.9	9.6 ± 4.6	10.5 ± 7.1	<.001
Osteotomy	62%	65%	63%	79%	78%	83%	.001
Major osteotomy	4%	6%	13%	32%	45%	55%	<.001
Decompression	16%	51%	58%	74%	67%	83%	<.001
Prior thoracolumbar fusion	6%	14%	21%	33%	41%	46%	<.001
Complication rates							
Major complication	6%	24%	23%	38%	55%	41%	<.001
Reoperation	12%	19%	21%	28%	27%	41%	.004
Admission to SICU	57%	58%	62%	72%	85%	88%	<.001
Proximal junctional kyphosis	36%	50%	58%	61%	50%	74%	.001
Proximal junctional failure	2%	5%	9%	14%	11%	20%	.004
Pseudarthrosis	2.5%	2.3%	1.7%	3.6%	7.0%	7.3%	.116
Surgical implant failure	1%	10%	16%	22%	27%	27%	<.001
Rod breaks	1.2%	5.4%	8.9%	15.8%	18.8%	24.4%	<.001
Screw breaks	.0%	.8%	2.8%	1.8%	5.5%	4.9%	.086
Cost-utility analysis							
Baseline ODI	25.9	40.0	43.5	48.1	52.5	56.0	<.001
Two-year ODI	17.2	22.9	23.5	30.5	34.6	40.7	<.001
Total cost 2Y	\$70,678.78	\$83,979.59	\$86,370.35	\$98,217.21	\$104,035.88	\$113,915.99	<.001
Utility gained	.084	.098	.104	.083	.075	.054	.015
QALYs gained 2Y	.163	.190	.202	.161	.146	.105	.015
Cost per QALY 2Y	\$434,452.91	\$438,425.43	\$430,936.83	\$611,173.02	\$721,936.73	\$1,086,846.81	<.001

risk score on the likelihood of developing certain complications and outcomes of overall cost-utility. Significance was set at $P < .05$. All statistical tests were performed using SPSS software (v25.0, Armonk, NY, USA).

Results

Patient Demographics

Overall, there were 724 patients that met inclusion criteria for this study. The mean baseline characteristics for this cohort were as follows: average age of 60.7 ± 13.6 , 77% were female,

and BMI 27.8 ± 5.9 kg/m², CCI of 1.8 ± 1.7 , and frailty index of 3.4 ± 1.6 .

Surgical Characteristics

The cohort had a mean operative time of 437 minutes (median: 375, standard deviation: 174.7) and a mean estimated blood loss of 1736 ± 1498 mL. Overall, 59% of patients received a decompression and 71% had an osteotomy. By approach, 67% received a posterior-only approach, 33% received a combined approach. Of those included in the cohort, 25.7% were undergoing a revision. There were 79.4% of patients fused to the pelvis. The mean length of stay for the cohort was 8.0 ± 4.7 days.

Table 3. Odds of Experiencing Complication by Increasing Number of Risk Factors.

Complication	Odds Ratio [Confidence Interval]	P-value
Major complication	1.60 [1.41-1.81]	<.001
Reoperation	1.28 [1.12-1.45]	<.001
Admission to SICU	1.43 [1.27-1.61]	<.001
Proximal junctional kyphosis	1.18 [1.06-1.32]	.003
Proximal junctional failure	1.42 [1.17-1.71]	<.001
Pseudarthrosis	1.43 [1.07-1.92]	.017
Surgical implant failure	1.53 [1.32-1.78]	<.001
Rod breaks	1.62 [1.36-1.94]	<.001
Screw breaks	1.63 [1.13-2.35]	.009

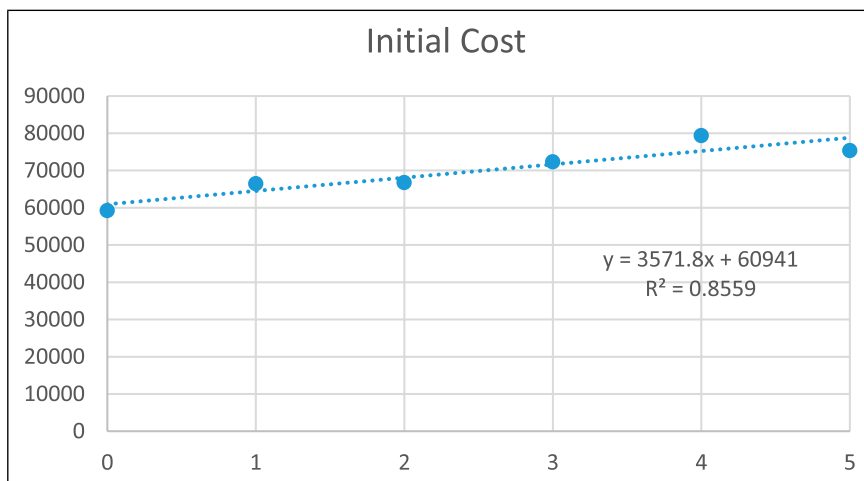


Figure 1. Correlation between risk score and initial cost.

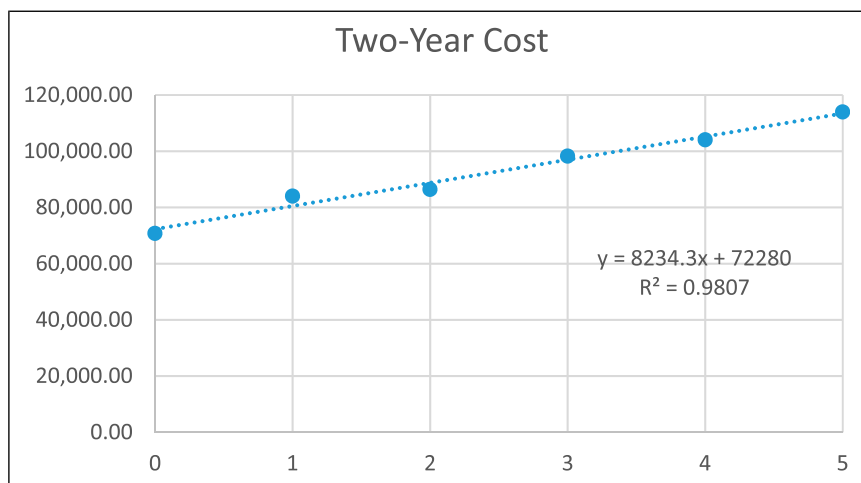


Figure 2. Correlation between risk score and two-year cost.

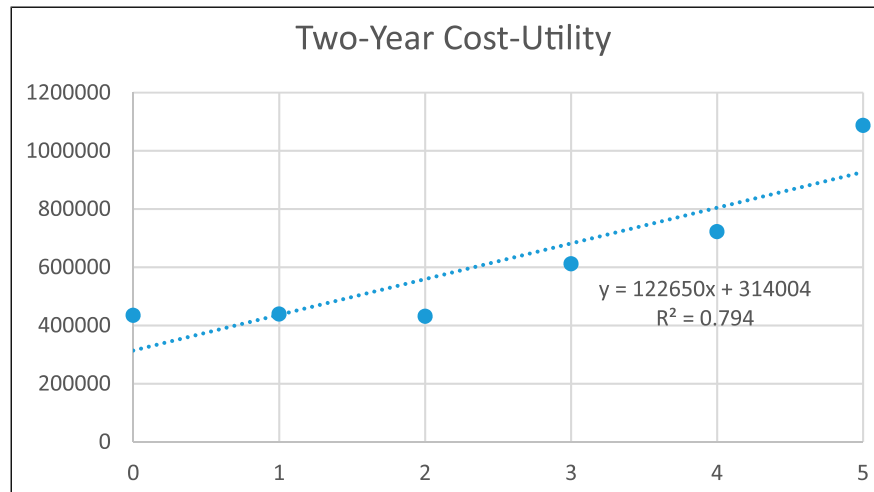


Figure 3. Correlation between risk score and two-year cost-utility.

Radiographic Characteristics

Patients presented radiographically at baseline with an average PI-LL of $16.6 \pm 20.8^\circ$, SVA 65.3 ± 71.7 mm, and a T1 Sagittal Tilt of $-1.2 \pm 6.1^\circ$. Addressing high deformity as per *Pellise* et al, 239 patients had high deformity in SVA (33.3%), 241 in PI-LL (33.4%), and 243 in T1-slope (33.4%).

Cohort Radiographic and Complication Outcomes

At 2 years, patients demonstrated radiographic parameters as follows: an average PI-LL of $3.9 \pm 14.8^\circ$, SVA 30.8 ± 53.0 mm, and a T1 Sagittal Tilt of $-3.9 \pm 4.9^\circ$. By 2 years, 74% of the cohort experienced any type of complication, 31% experienced a major complication, and 23% underwent a reoperation.

Overall Risk Stratification Model Quantification

Of the patients included in the study, 81 met 0 risk factors, 129 met 1, 180 met 2, 165 met 3, 128 met 4, and 41 met 5. The intersections between risk factors and the quantity of factors met can be visualized in [Table 1](#). When stratifying by number of risk factors, [Table 2](#) demonstrate the differences in demographics and surgical details, respectively.

Overall Complication and Reoperation Outcomes by Risk Stratification

With increasing number of risk factors, [Table 2](#) indicates the significant differences in between rates of complications. Patients who met criteria for 3, 4 or 5 risk factors had significantly higher major complication rates compared to those who met 0, 1, or 2 (all $P \leq .01$). Similar results were found for the association for major complications (OR: 1.6, 95% CI: [1.41-1.81], $P < .001$) and reoperation (OR: 1.3, 95% CI: [1.12-1.45], $P < .001$), along with admissions to SICU, development of proximal junctional

failure, surgical implant failure, rod breaks, and screw breaks by two years (all $P < .01$), as indicated in [Table 3](#).

Overall Correlation to Oswestry Disability Index

The amount of risk factors met had a positive correlation to baseline ODI scores ($r = .442$, $P < .001$) and 2-year ODI scores ($r = .311$, $P < .001$). When controlling for baseline disability, the group with 2 risk factors during ASD surgery gained the most utility (change in ODI from baseline to two years) out of all groups, but did not significantly outgain the group with 0 risk factors (-20.2 vs -16.2 , $P = .240$), depicted in [Table 2](#).

Overall Cost Evaluation

When analyzing risk factor quantity, there was a \$3572 increase in initial DRG cost-per-risk factor met ($R^2 = .856$; [Figure 1](#)), a \$8234 increase in initial national average cost-per-risk factor ($R^2 = .981$; [Figure 2](#)), and an increase by \$122,650 per risk factor in costs-per-QALY at two years ($R^2 = .794$; [Figure 3](#)). These findings can be visualized in [Table 2](#). Patients with 0, 1, or 2 risk factors had the lowest costs-per-QALY at two years, and these three groups each had significantly lower costs-per-QALY when compared to groups with 3, 4, or 5 risk factors (all $P < .001$). Adjusted generalized linear model accounting for surgery site, surgeon, and age of patient, demonstrated a significant trend between increasing risk score and increasing cost-utility ($r^2 = .408$, $P < .001$) with no significant interaction seen by site, surgeon, or age (all $P > .05$).

Frailty/Deformity Sub-Analysis

The overall cohort was divided into groups by deformity severity and frailty. After categorization, the following proportions were defined as: 30.4% were Not Frail/Low Deformity, 18.8% Not Frail/High Deformity, 21.0% were Frail/Low

Table 4. The Impact of Number of Risk Factors on Cost-Utility in Each Frailty/Deformity Cohort.

# Risk Factors	0	1	2	3	P-value
Not Frail Low Deformity					
Baseline ODI	25.8	39.6	33.0	33.4	<.001
2-Year ODI	15.5	21.0	16.4	18.6	.189
Total cost 2Y	\$68,478.08	\$86,105.06	\$94,728.08	\$109,610.96	<.001
Utility gained	.073	.077	.086	.075	.839
QALYs gained 2Y	.142	.150	.167	.146	.839
Cost per QALY 2Y	\$484,267.24	\$576,123.20	\$572,654.96	\$752,956.89	.002
Not frail high deformity					
Baseline ODI	43.3	40.5	38.0	37.4	.651
2-Year ODI	26.9	22.3	21.1	28.5	.407
Total cost 2Y	\$81,812.99	\$77,603.32	\$94,110.67	\$107,189.11	.039
Utility gained	.077	.092	.090	.050	.225
QALYs gained 2Y	.150	.179	.175	.097	.225
Cost per QALY 2Y	\$547,405.19	\$434,580.12	\$542,538.85	\$1,104,480.49	<.001
Frail low deformity					
Baseline ODI	45.5	49.1	52.1	52.7	.271
2-Year ODI	31.1	30.0	33.0	42.4	.085
Total cost 2Y	\$76,929.57	\$91,444.47	\$94,930.07	\$94,138.41	.400
Utility gained	.086	.102	.093	.043	.106
QALYs gained 2Y	.167	.198	.181	.084	.106
Cost per QALY 2Y	\$460,863.41	\$463,725.64	\$525,894.02	\$1,152,667.41	<.001
Frail high deformity					
Baseline ODI	59.4	56.8	54.0	53.4	.339
2-Year ODI	35.8	36.1	35.2	35.6	.995
Total cost 2Y	\$83,034.12	\$95,483.12	\$103,232.97	\$111,592.37	.171
Utility gained	.115	.104	.100	.095	.924
QALYs gained 2Y	.223	.202	.194	.184	.924
Cost per QALY 2Y	\$371,994.19	\$473,010.38	\$535,976.00	\$605,185.20	.006

Deformity, and 29.8% were Frail/High Deformity. Multi-variable regression analysis, controlling for baseline severity of deformity and frailty, assessed the odds of experiencing certain complications in each sub-cohort. Not Frail/Low Deformity patients with a risk score of 0 were less likely to any major complication (OR: .1, 95% CI: [.03-.3]; $P < .001$), major mechanical complication (no patients with risk score of zero suffered a major mechanical complication), and reoperation (OR: .4, 95% CI: [.2-.9]; $P = .041$) compared to those with a risk score of 1, when accounting for baseline deformity. In the Not Frail/High Deformity cohorts, patients with a risk score of 0 demonstrated higher rates of major complications (36%) and major radiographic complications (18%) than those with a risk score of 1 or 2 (both $P < .05$). In the Frail/Low Deformity cohort, patients with a risk score of zero were less likely to experience a major radiographic complication, specifically PJF (0%). While there were no differences between those with a risk score of 0 or 1 in the Frail/High Deformity population, patients with a risk score of 0 or 1 were less likely to develop a major complication (OR: .4, 95% CI: [.2-.8]; $P = .005$) or implant failure (OR: .5, 95% CI: [.3-.9]; $P = .037$) compared to those with a risk score of 2 or 3.

Frailty/Deformity Cost-Utility Analysis

When examining cost-utility by increasing risk score in [Table 4](#), there was a positive correlation with increasing risk score within each frailty/deformity cohort (all $P < .001$). Utility gained was highest in risk scores 1 or 2 compared to 0 or 3 in all frailty/deformity groups, except for Frail/High Deformity, where there was a decrease in utility gained with increasing risk score. When dividing total cost by QALYs gained over two years, there were no differences in the cost-utility between risk scores of 0, 1, and 2 for the first three cohorts (Not Frail/Low Deformity, Not Frail/High Deformity, Frail/Low Deformity), and all three scores had significantly favorable cost-utility compared to patients with a risk score of 3 (all $P < .05$). Patients with a risk score of 1 in the Not Frail/High Deformity cohort had the most favorable cost-utility, even compared to those with a risk score of 0 (\$434,580.12 vs \$547,405.19). However, patients in the Frail/High Deformity cohort generated the most favorable cost-utility with a risk score of 0, as increasing cost-utility demonstrated correlation with increasing risk score ($P < .001$).

Discussion

Risk factors for major complications, readmission, and reoperation were analyzed and their level of importance were determined in order to provide further insight for spine surgeons in patient selection and risk stratification in a study by *Pellise et al.* The study analyzed non-modifiable, modifiable, radiographic, surgical, and surgical site characteristics and found that certain factors had significant influence in providing time-risk predictions according to adverse events. Among these, blood loss, Sagittal Vertical Axis measurement, variables associated with frailty measures such as age and disability, T1 Sagittal Tilt, operative time, lowest instrumented vertebrae extension to pelvis, and lordosis gap were relevant in predicting major complications in the preoperative and postoperative models.⁷ Given that major complications have an influence on cost, the present study sought to utilize these stratification variables to determine the impact on cost-utility.^{15,27,30}

Among the patients who met 4 risk criteria, a vast majority in this group due to fusion to pelvis, frailty, deformity, or blood loss. Given that this group also had the most reduced cost utility, this validates a stronger weight with regard to these risk factors in comparison to fusion to pelvis, which has varying discussion regarding its involvement in patient outcomes.³¹⁻³⁴ We note that there was a fairly steady decrease in cost utility alongside an increase in risk factors. Given cost utility is calculated via the ratio of cost per QALY, this is most likely a compounded effect of the increasing cost associated with risk factors and the increased likelihood of complications as noted by the original *Pellise et al.* study. The same trend was also seen in regards to radiographic alignment, as an increase in number of risk factors was also correlated to a higher degree of deformity in all three parameters at baseline. Ultimately, because the amount of risk factors was also significantly associated with baseline ODI and follow-up ODI at 2 years, our cost-utility results represent a conglomerate of factors significantly impacting both complication rates and, most importantly, patient-reported outcomes.

Improvement in disability was significantly hindered by an accumulating number of risk factors. However, more interestingly, the patient group with 2 risk factors gained the most utility over the span over two years, even compared to patients with no risk factors at all. Likewise, because their total costs estimated over two years did not significantly differ from patients with a lower amount of risk factors ($P > .05$), they were able to achieve the lowest cost-utility profile overall, along with patients experiencing zero or one risk factor. Patients with one and two risk factors, however, amassed higher rates of complications more often compared to their counterparts with no risk factors. Yet, we did not see these significantly impact the total cost estimated when taking into account major and minor complications, as well as reoperation.

Therefore, the arising question is: “Are we overestimating the impact of these major and minor complications on clinical

outcomes to a degree?” While there were trends between increasing number of factors and increasing complication rates, overall cost, and decreased utility gained, there seemed to be a delineation of successful outcomes when parsing out the risk factors individually. For instance, there was a difference between patients with no risk factors and one risk factor in terms of utility gained and cost-utility, as well as between those with two and three risk factors. While the latter observation could be explained by the trend analysis, the former is perplexing. A reason for this could be that, in the correct patients with a lower number of risk factors, concerted efforts to achieve durable, optimal outcomes may also incur a slightly greater rate of complications. However, these minor blemishes do not preclude these patients from achieving better clinical outcomes. In fact, in our study, patients acquiring one to two risk factors actually performed slightly better clinically than those with no risk factors. Future research is needed to determine if these patients with a minor amount of risk for complications ultimately fare better long term in relation to those with higher or no risk at all.

Patients were then categorized by specific factors, severity of deformity and frailty, upon presentation that are often deemed unmodifiable prior to surgical intervention in order to examine the effect of modifying other factors to decrease complication rates and increase utility gained to ultimately produced a favorable economic outcome. When doing so for Not Frail patients with Low Deformity, patients with a risk score of zero (no presence of high EBL, high operative time, or fixation to the pelvis) generated the lowest cost-utility. As patients in this cohort lack the deformity severity compared to other cohorts in an analysis, it is intuitive lower amounts of invasiveness are required to generate a good outcome. However, when examining Not Frail patients with High Deformity, a risk score of 1 demonstrated the lowest cost-utility when accounting for baseline deformity. It would also make sense that these patient necessitate a more significant degree of correction, inquiring either higher blood loss, operative time, or fixation to the pelvis, in order to realign their deformity to optimal target goals. Therefore, a mild amount of risk in these patients, given their low frailty status, may result in significant clinical and economic benefit within this cohort.

However, this trend was the opposite for frail patients presenting for correction of adult spinal deformity. While the Frail patients with Low Deformity had similar cost-utility with risk score of 0, 1, or 2, Frail patients with High Deformity gained the greatest utility, the lowest cost, and most favorable cost-utility outcome out of any group when encountering a risk score of 0. Frailty has proven throughout the literature to be a better metric for assessing risk prior to surgery when compared to other standard demographic measures like age, BMI, and comorbidities. Therefore, this cohort of patients is already placed at higher risk for complications regardless of presenting deformity. With the addition of high baseline deformity, preventative measures against complication development should be emphasized and have significant effects.

Similarly, our study showed the highest rates of mechanical complications and reoperations among this group of patients. It is for this reason there has been a recent trend in modifying current realignment classifications to adjust for age, frailty, and even osteoporosis.³⁹⁻⁴¹ These patients often present with higher severity of deformity and disability, while also requiring less correction to regain significant functionality and the ability to perform activities of daily living.^{39,42} Frail patients have shown capability to surpass their Not Frail peers in clinical improvement from ASD surgery and achieve overall equivocal cost-utility from intervention. Therefore, the findings of this study are further testament to the utilization of adjusted realignment goals to tailor invasiveness and correction to both the patient-specific factors and needs in order to modify their inherent risk for higher rates of complications and maximize their utility gained.

Like frail patients, patients with one to two risk factors also started with a higher degree of deformity and disability than those with zero risk factors, and it is important to note they had more potential to improve at baseline. Patients with zero risk factors also began, on average, with lower deformity in each radiographic parameter when assessing them by either their SRS-Schwab or Ames-ISSG classification.^{35,36} Therefore, as baseline frailty has been assumed to have a parabolic effect on clinical improvement in ASD surgery, so may baseline deformity or disability, as both the lowest and highest degrees of each at baseline did not fare as well as those within the middle range when examining utility gained at two years.^{37,38}

Furthermore, knowing that adverse events also have a direct economic impact, this study potentially extended these associations to cost. This was evidenced by the present study showing a drop-off in cost-utility for patients with 3-5 risk factors, as well as the correlation between the risk factors and subsequently developing complications. Alongside major complications, our study demonstrated the risk factors tested correlated with a broad array of both early and late complications, from likelihood of being admitted to SICU to the development of either junctional or implant failure by two years. Each of these observed complications play their own role in the accrual of cost and, due to their predictability based on risk criteria, become further evidence for the utilization of a risk stratification system to identify the patient destined for complications that will significantly impact their clinical success and make appropriate decisions to prevent or minimize the likelihood of those risk factors when possible.

However, despite the significant findings, our study is not without limitations. The population under study may be prone to selection, indication and expertise bias. We also cannot identify, or control for, errors that may occur during the data entry process and are limited to consider factors routinely collected by the participating centers. The use of generalized cost estimates derived from PearlDiver may also mean the findings may not be translatable to all clinical contexts and healthcare environments. Further, more granular, testing in larger and more diverse samples remains to be performed.

While the results suggest that an increase in quantity of risk factors, regardless of which factor, increases the potential cost associated with medical intervention, it is important to state there is a potential confounding component as the makeup of each factor quantity grouping (0,1,2,3,4,5) by risk factor is not entirely uniform. However, the initial study by *Pellise et al* shed light on this question, as the study ranked the influence of these factors in determining the risk of major complication or revision. Concerning our study specifically, we recognize cost is not limited to the factors we assessed and, as we identified, there are other complications that may play a significant role in the differences in cost and overall cost-utility. Yet, it is worth noting many of the complications we reported in our study also correlated with the risk factors tested, aligning with our overall message. Lastly, as a cost-utility analysis, we caution against use of this work to deny patients surgery or define who may, or may not, undergo adult spinal deformity correction. There may still be patients in a higher risk stratification category who benefit substantially from surgical correction of adult spinal deformity, even if the procedure is not as cost-effective as it might be for individuals in lower risk stratification categories.

Conclusion

As advances in spine surgery enable the physician to take on more challenging cases, it has become even more imperative to recognize which pre-existing and surgical factors play a role in development of complications. This study shows an increase in risk factors involving frailty, baseline deformity, and operative factors adversely affect cost-utility by increasing total cost and reducing QALYs at 2 years. The clinical influence of the study may be multiple-fold, but predominantly may aid with the prioritisation and planning of risk-factor amelioration when planning and executing surgical intervention for ASD wherever possible. Preoperative or intraoperative measures should be taken to reduce a patient's amount of applicable risk factors in order to concurrently reduce higher rates of complications, increase cost-utility, and potentially maximize optimal clinical outcomes.

Authors Note

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Ethical Statement

Ethical Approval

Institutional Review Board approval was obtained before enrolling patients in the prospective database. Informed consent was obtained from each patient prior to enrollment.

ORCID iDs

Peter G. Passias [ID](https://orcid.org/0000-0002-1479-4070) <https://orcid.org/0000-0002-1479-4070>
 Tyler K. Williamson [ID](https://orcid.org/0000-0003-2584-291X) <https://orcid.org/0000-0003-2584-291X>
 Nicholas A. Kummer [ID](https://orcid.org/0000-0003-0334-4434) <https://orcid.org/0000-0003-0334-4434>
 Virginie Lafage [ID](https://orcid.org/0000-0002-0119-7111) <https://orcid.org/0000-0002-0119-7111>
 Renaud Lafage [ID](https://orcid.org/0000-0002-4820-1835) <https://orcid.org/0000-0002-4820-1835>
 Breton Line [ID](https://orcid.org/0000-0003-0395-1066) <https://orcid.org/0000-0003-0395-1066>
 Jeffrey L. Gum [ID](https://orcid.org/0000-0003-0471-9437) <https://orcid.org/0000-0003-0471-9437>
 Munish C. Gupta [ID](https://orcid.org/0000-0002-4711-4377) <https://orcid.org/0000-0002-4711-4377>
 Khaled M. Kebaish [ID](https://orcid.org/0000-0002-1235-6202) <https://orcid.org/0000-0002-1235-6202>

References

- Kim YC, Lee KH, Kim GL, et al. Improvements in lower-extremity patient-reported outcomes after lumbar interbody fusion. *J Neurosurg Spine*. 36, 2021 Sep 3:1-8. doi: [10.3171/2021.2.SPINE201494](https://doi.org/10.3171/2021.2.SPINE201494). Epub ahead of print.
- Lomelí-Rivas A, Larrinúa-Betancourt JE. Biomecánica de la columna lumbar: un enfoque clínico [Biomechanics of the lumbar spine: A clinical approach]. *Acta Ortop Mex*. 2019; 33(3):185-191. Spanish.
- Worley N, Buza J, Jalai CM, et al. Diabetes as an independent predictor for extended length of hospital stay and increased adverse post-operative events in patients treated surgically for cervical spondylotic myelopathy. *Int J Spine Surg*. 2017;11(2): 10. doi:[10.14444/4010](https://doi.org/10.14444/4010)
- Pierce KE, Kapadia BH, Bortz C, et al. Operative fusion of patients with metabolic syndrome increases risk for perioperative complications. *J Clin Neurosci*. 2020;72:142-145. Epub 2019 Dec 30. doi:[10.1016/j.jocn.2019.12.043](https://doi.org/10.1016/j.jocn.2019.12.043)
- Pierce KE, Passias PG, Daniels AH, et al. Baseline frailty status influences recovery patterns and outcomes following alignment correction of cervical deformity. *Neurosurgery*. 2021;88(6): 1121-1127. doi:[10.1093/neuros/nyab039](https://doi.org/10.1093/neuros/nyab039)
- Tabatabai S, Do Q, Min J, et al. Obesity and perioperative outcomes in older surgical patients undergoing elective spine and major arthroplasty surgery. *J Clin Anesth*. 2021;75:110475. Epub ahead of print. doi:[10.1016/j.jclinane.2021.110475](https://doi.org/10.1016/j.jclinane.2021.110475)
- Pellisé F, Serra-Burriel M, Smith JS, et al. Development and validation of risk stratification models for adult spinal deformity surgery. *J Neurosurg Spine*. 2019;31:1-13. doi:[10.3171/2019.3.SPINE181452](https://doi.org/10.3171/2019.3.SPINE181452)
- Pizones J, Moreno-Manzanaro L, Sánchez Pérez-Grueso FJ, et al. Restoring the ideal Roussouly sagittal profile in adult scoliosis surgery decreases the risk of mechanical complications. *Eur Spine J*. 2020;29(1):54-62. doi:[10.1007/s00586-019-06176-x](https://doi.org/10.1007/s00586-019-06176-x)
- Agarwal N, Goldschmidt E, Taylor T, et al. Impact of frailty on outcomes following spine surgery: A prospective cohort

- analysis of 668 patients. *Neurosurgery*. 2021;88(3):552-557. doi:10.1093/neuros/nyaa468
10. Passias PG, Bortz CA, Segreto FA, et al. Development of a modified cervical deformity frailty index: A streamlined clinical tool for preoperative risk stratification. *Spine (Phila Pa 1976)*. 2019;44(3):169-176. doi:10.1097/BRS.0000000000002778
 11. Horn SR, Passias PG, Bortz CA, et al. Predicting extended operative time and length of inpatient stay in cervical deformity corrective surgery. *J Clin Neurosci*. 2019;69:206-213. doi:10.1016/j.jocn.2019.07.064
 12. Horn SR, Pierce KE, Oh C, et al. Predictors of hospital-acquired conditions are predominately similar for spine surgery and other common elective surgical procedures, with some key exceptions. *Global Spine J*. 2019;9(7):717-723. doi:10.1177/2192568219826083
 13. Bortz C, Alas H, Segreto F, et al. Complication risk in primary and revision minimally invasive lumbar interbody fusion: A comparable alternative to conventional open techniques? *Global Spine J*. 2020;10(5):619-626. doi:10.1177/2192568219867289
 14. Ahmed SI, Bastrom TP, Yaszay B, Newton PO, Harms Study Group. 5-Year reoperation risk and causes for revision after idiopathic scoliosis surgery. *Spine (Phila Pa 1976)*. 2017;42(13):999-1005. doi:10.1097/BRS.0000000000001968
 15. Passias PG, Poorman GW, Bortz CA, et al. Predictors of adverse discharge disposition in adult spinal deformity and associated costs. *Spine J*. 2018;18(10):1845-1852. doi:10.1016/j.spinee.2018.03.022
 16. Daniels AH, Reid DBC, Durand WM, et al. Assessment of patient outcomes and proximal junctional failure rate of patients with adult spinal deformity undergoing caudal extension of previous spinal fusion. *World Neurosurg*. 2020;139:e449-e454. doi:10.1016/j.wneu.2020.04.024
 17. Gerling MC, Radcliff K, Isaacs R, et al. Two-year results of the prospective spine treatment outcomes study: An analysis of complication rates, predictors of their development, and effect on patient derived outcomes at 2 Years for surgical management of cervical spondylotic myelopathy. *World Neurosurg*. 2017;106:247-253. doi:10.1016/j.wneu.2017.06.147
 18. Blondel B, Schwab F, Ungar B, et al. Impact of magnitude and percentage of global sagittal plane correction on health-related quality of life at 2-years follow-up. *Neurosurgery*. 2012 Aug;71(2):341. doi:10.1227/NEU.0b013e31825d20c0
 19. Liu S, Schwab F, Smith JS, et al. Likelihood of reaching minimal clinically important difference in adult spinal deformity: A comparison of operative and nonoperative treatment. *Ochsner J*. 2014;14(1):67-77.
 20. Gum JL, Line B, Carreon LY, et al. Reaching the medicare allowable threshold in adult spinal deformity surgery: multi-center cost analysis comparing actual direct hospital costs versus what the government will pay. *Spine Deform*. 2022;10:425-431. Epub ahead of print. doi:10.1007/s43390-021-00405-4
 21. Passias PG, Bortz CA, Pierce KE, et al. A simpler, modified frailty index weighted by complication occurrence correlates to pain and disability for adult spinal deformity patients. *Int J Spine Surg*. 2020;14(6):1031-1036. doi:10.14444/7154
 22. Champain S, Benchikh K, Nogier A, Mazel C, Guise JD, Skalli W. Validation of new clinical quantitative analysis software applicable in spine orthopaedic studies. *Eur Spine J*. 2006;15:982-991. DOI: 10.1007/s00586-005-0927-1
 23. O'Brien MF, Kuklo TR, Blanke KM, Lenke LG. *Spinal Deformity Study Group Radiographic Measurement Manual*. Memphis, TN, USA: Medtronic Sofamor Danek; 2005.
 24. Rillardon L, Levassor N, Guigui P, et al. Validation of a tool to measure pelvic and spinal parameters of sagittal balance. *Rev Chir Orthop Reparatrice Appar Mot*. 2003;89:218-227.
 25. Elshamly M, Windhager R, Toegel S, Grohs JG. Long-term impact of sagittal malalignment on hardware after posterior fixation of the thoracolumbar spine: a retrospective study. *BMC Musculoskel Disord*. 2020 Jun 16;21(1):387. doi:10.1186/s12891-020-03405-z
 26. Carreon LY, Glassman SD, McDonough CM, Rampersaud R, Berven S, Shainline M. Predicting SF-6D utility scores from the Oswestry disability index and numeric rating scales for back and leg pain. *Spine (Phila Pa 1976)*. 2009;34:2085-2089. doi:10.1097/BRS.0b013e3181a93ea6
 27. Poorman GW, Passias PG, Qureshi R, et al. Cost-utility analysis of cervical deformity surgeries using 1-year outcome. *Spine J*. 2018:1552-1557. doi:10.1016/j.spinee.2018.01.016.LK. http://elinks.library.upenn.edu/sfx_local?sid=EMBASE&issn=18781632&id=doi:10.1016%2Fj.spinee.2018.01.016&atitle=Cost-utility+analysis+of+cervical+deformity+surgeries+using+1-year+outcome&stitle=Spine+J.&title=Spine+Journal&volume=&issue=&spage=&epage=&aulast=Poorman&aufirst=Gregory+W.&aunit=G.W.&afulfill=Poorman+G.W.&coden=SJPOA&isbn=&pages=-&date=2018&aunit1=G&aunitm=W
 28. Carreon LY, Bratcher KR, Das N, Nienhuis JB, Glassman SD. Estimating EQ-5D values from the Oswestry Disability Index and numeric rating scales for back and leg pain. *Spine (Phila Pa 1976)*. 2014;39(8):678-682. doi:10.1097/BRS.0000000000000220.
 29. Gum JL, Hostin R, Robinson C, et al. Impact of cost valuation on cost-effectiveness in adult spine deformity surgery. *Spine J*. 2017;17(1):96-101. doi:10.1016/j.spinee.2016.08.020
 30. Brown AE, Lebovic J, Alas H, et al. A cost utility analysis of treating different adult spinal deformity frailty states. *J Clin Neurosci*. 2020;80:223-228. doi:10.1016/j.jocn.2020.07.047
 31. Martin CT, Holton KJ, Jones KE, Sembrano JN, Polly DW. Bilateral open sacroiliac joint fusion during adult spinal deformity surgery using triangular titanium implants: Technique description and presentation of 21 cases. *J Neurosurg Spine*. 2021;36:1-7. Epub ahead of print. doi:10.3171/2021.3.SPINE202218
 32. Chen S, Luo M, Wang Y, Liu H. Stopping at sacrum versus nonsacral vertebra in long fusion surgery for adult spinal deformity: Meta-analysis of revision with minimum 2-year follow-up. *World Neurosurg*. 2018;S1878-8750(18):32925. Epub ahead of print. PMID: 30605759. doi:10.1016/j.wneu.2018.12.102
 33. Bridwell KH, Edwards CC 2nd, Lenke LG. The pros and cons to saving the L5-S1 motion segment in a long scoliosis fusion construct. *Spine (Phila Pa 1976)*. 2003;28(20):S234. doi:10.1097/01.BRS.0000092462.45111.27

34. Kim YJ, Bridwell KH, Lenke LG, Cho KJ, Edwards CC 2nd, Rinella AS. Pseudarthrosis in adult spinal deformity following multisegmental instrumentation and arthrodesis. *J Bone Joint Surg Am.* 2006;88(4):721. doi:[10.2106/JBJS.E.00550](https://doi.org/10.2106/JBJS.E.00550)
35. Schwab F, Ungar B, Blondel B, et al. Scoliosis Research Society-Schwab adult spinal deformity classification: A validation study. *Spine (Phila Pa 1976).* 2012;37(12):1077-1082. doi:[10.1097/BRS.0b013e31823e15e2](https://doi.org/10.1097/BRS.0b013e31823e15e2)
36. Ames CP, Smith JS, Eastlack R, et al. Reliability assessment of a novel cervical spine deformity classification system. *J Neurosurg Spine.* 2015;23(6):673-683. Epub 2015 Aug 14. doi:[10.3171/2014.12.SPINE14780](https://doi.org/10.3171/2014.12.SPINE14780)
37. Reid DBC, Daniels AH, Ailon T, et al. Frailty and health-related quality of life improvement following adult spinal deformity surgery. *World Neurosurg.* 2018;112:e548-e554. Epub 2018 Jan 31. doi:[10.1016/j.wneu.2018.01.079](https://doi.org/10.1016/j.wneu.2018.01.079)
38. Than KD, Park P, Fu KM, et al. Clinical and radiographic parameters associated with best versus worst clinical outcomes in minimally invasive spinal deformity surgery. *J Neurosurg Spine.* 2016;25(1):21-25. Epub. doi:[10.3171/2015.12.SPINE15999](https://doi.org/10.3171/2015.12.SPINE15999)
39. Lafage R, Schwab F, Glassman S, et al. Age-adjusted alignment goals have the potential to reduce PJK. *Spine (Phila Pa 1976).* 2017;42:1275-1282.
40. Noh SH, Ha Y, Park JY, et al. Modified global alignment and proportion scoring with body mass index and bone mineral density analysis in global alignment and proportion score of each 3 categories for predicting mechanical complications after adult spinal deformity surgery. *Neurospine.* 2021;18(3):484-491. doi:[10.14245/ns.2142470.235](https://doi.org/10.14245/ns.2142470.235)
41. Hong YG, Kim HC, Jeon H, et al. Association of frailty with regional sagittal spinal alignment in the elderly. *J Clin Neurosci.* 2022;96:172-179. Epub 2021 Nov 24. doi:[10.1016/j.jocn.2021.10.008](https://doi.org/10.1016/j.jocn.2021.10.008)
42. Passias PG, Ahmad W, Kummer N, et al. Examination of the economic burden of frailty in patients with adult spinal deformity undergoing surgical intervention. *Neurosurgery.* 2022;90(1):148-153. doi:[10.1227/NEU.0000000000001756](https://doi.org/10.1227/NEU.0000000000001756)
43. Passias PG, Moattari K, Pierce KE, et al. Performance of the modified adult spinal deformity frailty index in preoperative risk assessment. *Spine (Phila Pa 1976).* 2022;47:1463-1469. Epub ahead of print. doi:[10.1097/BRS.0000000000004342](https://doi.org/10.1097/BRS.0000000000004342)