Revision by S2-alar-iliac instrumentation reduces caudal screw loosening while improving sacroiliac joint pain—a group comparison study

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Received: 19 June 2020 / Revised: 10 August 2020 / Accepted: 20 August 2020
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Abstract
Lumbosacral instrumentation continues to be challenging due to complex biomechanical force distributions and poor sacral bone quality. Various techniques have therefore been established. The aim of this study was to investigate the outcome of patients treated with S2-alar-iliac (S2AI), S2-alar (S2A), and iliac (I) instrumentation as the most caudal level. Sixty patients underwent one of the 3 techniques between January 2012 and June 2017 (S2AI 18 patients, S2A 20 patients, I 22 patients). Mean age was 70.4 ± 8.5 years. Screw loosening (SL) and sacroiliac joint (SIJ) pain were evaluated during the course at 3-month and maximum follow-up (FU). All patients completed 3-month FU, the mean FU period was 2.5 ± 1.5 years (p = 0.38), and a median of 5 segments was operated on (p = 0.26), respectively. Bone mineral density (BMD), derived opportunistically from computed tomography (CT), did not significantly differ between the groups (p = 0.66), but cages were more frequently implanted in patients of the S2A group (p = 0.04). SL of sacral or iliac screws was more common in patients of the S2A and I groups compared with the S2AI group (S2AI 16.7%, S2A 55.0%, I 27.3% of patients; p = 0.03). SIJ pain was more often improved in the S2AI group not only after 3 months but also at maximum FU (S2AI 61.1%, S2A 25.0%, I 22.7% of patients showing improvement; p = 0.02). Even in shorter or mid-length lumbar or thoracolumbar constructs, S2AI might be considered superior to S2A and I instrumentation due to showing lower incidences of caudal SL and SIJ pain.

Keywords S2AI • Instrumentation • Sacrum • Degenerative spine • Screw loosening

Introduction
Instrumentation of the lumbosacral spine continues to be a challenging area in spine surgery, particularly due to complex local anatomy, unique biomechanical force distributions, and comparatively poor sacral bone quality. Various concepts for construct improvement have therefore been developed. However, there is a high rate of screw loosening (SL), instrumentation failure, pseudarthrosis, and sacroiliac joint (SIJ) pain (up to 83%) [1–3].

The Galveston technique and iliac screws were the first approaches trying to improve caudal instrumentation by extending the construct down to the pelvis [4, 5]. However, although iliac screws were proven to be superior to the Galveston technique in terms of construct strength, iliac screws were shown to cause pain due their prominence, ending in revision surgery in about 22% of cases [5, 6]. Moreover, instrumentation down to the sacrum puts a long cranial cantilever on the SIJ, not only reducing the durability of the construct but also causing a biomechanical overload and therefore
severe pain of the SIJ, which was reported in a systematic review to occur in $37 \pm 28.48\%$ (range 6–75%) of patients [7].

The search for alternative techniques to this point ended in the description and clinical establishment of the S2-alar-iliac (S2AI) screw trajectory [8]. By crossing 3 cortical bone structures (sacral bone plus SIJ) and allowing for long screw pinching between the compact bone of the pelvis, while also allowing for small incisions due to diverging trajectories and in-line rod bending, this technique offers a variety of advantages over iliac screw placements [9]. Likewise, due to the rigid trans-SIJ trajectory, this technique inhibits any SIJ motion, in contrast to iliac screws and is therefore assumed to better avoid postoperative SIJ pain. On the other hand, some colleagues argue that this rigid SIJ fixation causes reduced but necessary motion inside the SIJ, thus potentially reducing the patients’ quality of life [10]. Moreover, the trajectory is more demanding, thus causing potentially more vascular complications inside the small pelvis or even requiring spinal navigation to ease the approach [11].

We therefore hypothesize that instrumentations including S2AI screws are less prone to SL and SIJ pain compared with other established techniques. Thus, the aim of this study was to investigate differences in outcome between patients treated with S2AI, S2-alar (S2A), and iliac (I) instrumentation as the most caudal level.

**Materials and methods**

**Study design**

We reviewed all medical records and specifically checked the pre- and postoperative examinations including imaging studies, as well as postoperative follow-up (FU). The study enrolled 60 consecutive patients (35 male and 25 female) who underwent instrumentation by one of the 3 techniques between January 2012 and June 2017 as revision surgery after initial instrumentation (S2AI 18 patients [Fig. 1], 50% females, 72.1 ± 7.4 years; S2A 20 patients, 35% females, 69.8 ± 8.6 years; I 22 patients, 41% females, 69.5 ± 9.0 years). Outcome (SL and gluteal pain due to SIJ pain) was compared between the 3 groups considering preoperative, 3-month FU, and maximum FU examinations. Pain was routinely evaluated over the whole observation period during treatment and FU using a 10-point visual analogue scale.

Bone mineral density (BMD) was opportunistically assessed in preoperative imaging by computed tomography (CT) [12]. The 3 groups were comparable regarding the surgical approach and strategy of the treatment, especially in terms of the degree to which their deformities were corrected. The decision to perform one procedure versus another was based on the surgeons’ general preference. No patient criteria were taken into account in making this decision. SL was determined via CT images in all 3 planes by a conference of neurosurgeons and neuroradiologists.

**Ethics**

This study was approved by our local ethics committee (registration number: 159/16S) and was conducted in accordance with the Declaration of Helsinki. Informed consent was not required due to the retrospective character of the study and the conditions of our local ethics committee.

**Statistics**

All analyses were performed with the statistical software R (version 3.6.1; https://www.r-project.org/). $p$ values < 0.05 were considered statistically significant.

General characteristics of the investigated cohort were presented stratified in 3 groups: S2AI, S2A, and I, depending on the surgical technique. Means and standard deviation (SD), median and ranges, or absolute or relative numbers were
calculated. Depending on data distribution, analyses of variance (ANOVA), Kruskal-Wallis tests, or Chi-squared tests were used to assess differences in the analyzed parameters between the 3 groups.

The proportions and absolute numbers of patients with SIJ pain according to preoperative assessment, as well as at 3-month and maximum FU, were calculated stratified by group, respectively. Furthermore, proportions and absolute numbers of patients with improvement, deterioration, and unchanged status in SIJ pain at maximum FU compared with the preoperative status were calculated. Differences in these proportions between the 3 groups were assessed using Chi-squared or Fisher’s exact tests.

Potential predictors for improvement in SIJ pain were assessed, again stratified by group. Specifically, mean and SD, median and ranges, or absolute or relative numbers were calculated in patients with improvement and in patients with deterioration or unchanged status regarding SIJ pain. Associations with this status were tested using \( t \) tests, Mann-Whitney \( U \) tests, or Chi-squared tests. Improvement at maximum FU compared with the preoperative status was considered. The Benjamini-Hochberg procedure was used for correction of multiple testing regarding these prediction analyses, assuming a false discovery rate of 25%.

**Results**

**General results and group differences**

Participants’ mean age was 70.4 ± 8.5 years. The general demographics of all 3 groups are outlined in Table 1. Overall, the 3 groups were highly comparable and showed no statistically significant differences regarding baseline characteristics at the preoperative state. All patients received a 2-rod titanium construct.

All patients completed 3-month FU; maximum FU times were 2.3 ± 0.9 (S2AI), 3.0 ± 1.7 (S2A), and 2.4 ± 1.6 (I) years \( (p = 0.38) \). A median of 4.5 to 5 segments (S2AI, S2A) and 3 segments (I) was operated on \( (p = 0.26) \), extending to S2 or the os ilium, respectively. BMD as assessed opportunistically by CT did not significantly differ between the groups \( (p = 0.66) \). Cages were more frequently implanted in patients of the S2A group \( (p = 0.04) \).

**Caudal screw loosening**

SL of sacral or iliac screws was more common in patients of the S2A and I groups when compared with the S2AI group (S2AI 16.7%, S2A 55.0%, I 27.3% of patients; \( p = 0.03 \)).

**Improvement in sacroiliac joint pain**

Gluteal pain as a clinical sign for SIJ pain was more often improved in the S2AI group not only after 3 months but also at maximum FU (S2AI 61.1%, S2A 25.0%, I 22.7% of patients showing improvement; \( p = 0.02 \); Table 2). While the S2A group already had a low rate of preoperative SIJ pain, the pain was comparable in the S2AI and I groups but significantly better in the S2AI group at 3-month and maximum FU. While 61.1% of patients in the S2AI group improved from their preoperatively existing SIJ pain, only 25.0% in the S2A and 18.2% in the I group improved at 3-month FU.

Specifically, predictors of SIJ pain at maximum FU were the number of fused segments and the number of screws per side from previous surgery among the S2AI group (Table 3), but these predictors did not survive correction for multiple comparisons. However, patients with SIJ pain at FU also required revision surgery significantly more often, in most cases due to and as a sign of SL.

**Complications**

There were no perioperative screw-related vascular or visceral surgical complications due to sacral or iliac screws; however, there was one screw-related aortic dissection due to an L2 screw in the I group. In the S2AI group, cerebrospinal fluid (CSF) leakage due to surgery was observed in 3 patients (16.7%), whereas hematoma occurred in 1 patient (5.6%). Furthermore, 3 patients (15.0%) of the S2A group showed CSF leakage, and 2 (10.0%) showed hematoma, whereas another 2 patients (10.0%) showed screw dislocation with cement leakage in 1 case. Among patients of the I group, 1 patient (4.5%) showed screw dislocation.

Regarding postoperative medical complications, urinary tract infection was most common (S2AI 4 patients, S2A 2 patients), followed by deep vein thrombosis (S2AI 2 patients, S2A 1 patient, I 2 patients), partially with related pulmonary artery embolism (S2AI 1 patient, I 2 patients). Two patients of the S2A group developed pneumonia, 1 patient of the S2AI group had a postoperative non-ST-elevation myocardial infarction, and 1 patient of the I group had postoperative endocarditis.

**Discussion**

Our main findings were that S2AI showed superiority compared with S2A and I techniques in terms of reduced caudal SL and a lower rate of SIJ pain. Additionally, none of the differences between groups significantly promoted these effects.

Considering the results, any of the 3 techniques is better than using S1 instrumentation as the caudal part at all [3]. While spinopelvic fixation is nowadays achieved via a range of approaches, iliac and S2AI screws are presently the most commonly used ones. Caudal instrumentation down to the pelvic ring offers reinforced biomechanical strength, especially if long fusions extend to the sacrum or in cases of sacrectomy, osteoporosis, and deformity surgery necessitating
osteotomies or general revisions [13]. Clinically, additional iliac screws have been shown to be superior in neuromuscular spinal deformities and overall pediatric patients [14]. The same is true in adult revision cases after failed lumbosacropelvic fixation. Even in those cases, lumbosacral fusion could be promoted [15].

Table 1 General characteristics

|                          | S2AI (n = 18) | S2A (n = 20) | I (n = 22) | p  |
|--------------------------|--------------|--------------|------------|----|
| Age (in years; mean (SD))| 72.1 (7.4)   | 69.8 (8.6)   | 69.5 (9.0) | 0.364|
| Sex (% of patients (N))  | Female       | Female       | Female     | 0.646|
| Osteopenia / osteoporosis (% of patients (N)) | None | Osteopenia | Osteoporosis | 0.607|
| Spondylolisthesis (% of patients (N)) | 27.8 (5) | 20.0 (4) | 36.4 (8) | 0.551|
| Previous surgery         |              |              |            |     |
| No. of decompressed vertebrae (median (range)) | 3.0 (0–6) | 2.0 (0–4) | 2.5 (0–5) | 0.300|
| No. of screws one side (median (range)) | 3.0 (0–7) | 3.0 (0–6) | 3.0 (0–8) | 0.664|
| Cement augmentation (% of patients (N)) | 6.3 (1) | 12.5 (2) | 25.0 (5) | 0.311|
| No. of cages (median (range)) | 0.5 (0–4) | 0.5 (0–3) | 1.0 (0–8) | 0.731|
| Vertebral body replacement (% of patients (N)) | 6.3 (1) | 6.3 (1) | 20.0 (4) | 0.351|
| Anterior fusion (% of patients (N)) | 12.5 (2) | 0.0 (0) | 10.0 (2) | 0.537|
| Surgery                  |              |              |            |     |
| Time since last spine surgery (in years; mean (SD)) | 2.0 (2.7) | 1.7 (2.7) | 1.4 (1.1) | 0.458|
| Screw length (in mm; mean (SD)) | 82.8 (9.0) | 49.3 (7.5) | 97.7 (13.1) | 0.014|
| No. of decompressed vertebrae new (median (range)) | 0.5 (0–5) | 1.0 (0–7) | 0.0 (0–6) | 0.039|
| No. of fused segments overall (median (range)) | 5.0 (2–9) | 4.5 (1–9) | 3.0 (2–11) | 0.258|
| No. of screws one side overall (median (range)) | 6.0 (3–11) | 6.5 (3–11) | 5.0 (3–13) | 0.294|
| Cement augmentation new (% of patients (N)) | 27.8 (5) | 35.0 (7) | 22.7 (5) | 0.675|
| No. of levels with newly implanted cages (median (range)) | 0.0 (0–4) | 1.5 (0–5) | 0.0 (0–3) | 0.039|
| Newly implanted vertebral body replacement (% of patients (N)) | 5.6 (1) | 15.0 (3) | 4.6 (1) | 0.507|
| Additional anterior fusion (% of patients (N)) | 0.0 (0) | 5.0 (1) | 4.6 (1) | 1.000|
| Intraoperative blood loss (in ml; mean (SD)) | 1911.0 (2523.0) | 1120.0 (593.3) | 1545.0 (1436.0) | 0.899|
| Duration of surgery (in min; mean (SD)) | 289.3 (98.3) | 247.4 (73.6) | 248.0 (88.3) | 0.245|

General characteristics of the investigated cohort divided into three groups: S2-alar-iliac (S2AI), S2-alar (S2A), and iliac (I) instrumentation. Furthermore, information on the preoperative status regarding previous surgery as well as info on characteristics of the present surgery are given. Italicized p values indicate statistical significance (p < 0.05)

Table 2 Improvement in sacroiliac joint (SIJ) pain

|                          | S2AI (n = 18) | S2A (n = 20) | I (n = 22) | p  |
|--------------------------|--------------|--------------|------------|----|
| SIJ pain—preoperative (% of patients (N)) | 61.1 (11) | 30.0 (6) | 63.6 (14) | 0.059|
| SIJ pain—3-month follow-up (% of patients (N)) | 27.8 (5) | 10.0 (2) | 40.9 (9) | 0.078|
| SIJ pain—maximum follow-up (% of patients (N)) | 11.1 (2) | 10.0 (2) | 40.9 (9) | 0.030|
| SIJ pain—3-month follow-up compared with preoperative status (% of patients (N)) | Worse | Better | 5.6 (1) | 61.1 (11) | 6.6 (11) | 25.0 (5) | 18.2 (4) | 0.007|
| SIJ pain—maximum follow-up compared with preoperative status | Unchanged | 33.3 (6) | 75.0 (15) | 81.8 (18) | 0.015|
| SIJ pain—maximum follow-up compared with preoperative status | Worst | Better | 5.6 (1) | 61.1 (11) | 6.6 (11) | 25.0 (5) | 22.7 (5) | 0.015|
| SIJ pain—maximum follow-up compared with preoperative status | Unchanged | 33.3 (6) | 75.0 (15) | 77.3 (17) | 0.015|

Overview on SIJ pain during the clinical course considering the preoperative status as well as assessments at 3-month and maximum follow-up (FU). Italicized p values indicate statistical significance (p < 0.05)
Table 3  Sacroiliac joint (SIJ) pain at maximum follow-up (FU)

|                          | S2A1 (n = 18) | S2A (n = 20) | I (n = 22) |
|--------------------------|---------------|--------------|------------|
|                          | Unchanged or worse (N = 7) | Improved (N = 11) | p       | Unchanged or worse (N = 15) | Improved (N = 5) | p       | Unchanged or worse (N = 17) | Improved (N = 5) | p       |
| Age (in years; mean (SD))| 72.1 (6.7)    | 72.1 (8.4)   | 0.992      | 69.7 (9.1)   | 70.1 (8.9)   | 0.931   | 70.7 (10.0)      | 65.6 (4.5)   | 0.125   |
| Sex (% of patients (N))  | Female 71.4 (5) | 36.4 (4) | 0.335 | Male 60.0 (9)   | 80.0 (4)   | 0.613   | 58.8 (10) | 60.0 (3)   | 1.000   |
| Osteopenia/osteoporosis (% of patients (N)) | None 0.0 (0) | 0.0 (0) | 0.593 | Osteopenia 14.3 (2) | 2.5 (0-5) | 0.530 | 2.0 (0-8) | 4.0 (2-6) | 0.132 |
| Spondylolisthesis (% of patients (N)) | 42.9 (3) | 8.0 (8) | 0.326 | 20.0 (3) | 20.0 (1) | 1.000 | 29.4 (5) | 60.0 (3) | 0.309 |
| Previous surgery | No. of decompressed vertebrae (median (range)) | 3.0 (0-4) | 3.0 (0-6) | 1.000 | 2.0 (0-4) | 2.5 (0-3) | 1.000 | 2.0 (0-5) | 3.0 (2-4) | 0.392 |
|                        | No. of fused segments (median (range)) | 2.0 (0-4) | 4.0 (0-7) | 0.055* | 2.5 (0-5) | 3.0 (0-6) | 0.530 | 2.0 (0-8) | 4.0 (2-6) | 0.132 |
|                        | No. of screws one side (median (range)) | 3.0 (0-4) | 5.0 (0-6) | 0.020* | 4.0 (2-5) | 4.0 (0-7) | 0.935 | 3.0 (0-9) | 4.0 (3-6) | 0.127 |
|                        | Cement augmentation (% of patients (N)) | 14.3 (1) | 0.0 (0) | 0.438 | 8.3 (1) | 25.0 (1) | 0.450 | 20.0 (3) | 40.0 (2) | 0.560 |
|                        | No. of cages (median (range)) | 0.0 (0) | 1.0 (0-4) | 0.394 | 0.5 (0-3) | 1.0 (0-3) | 0.948 | 1.0 (0-8) | 2.0 (0-4) | 0.114 |
|                        | Vertebral body replacement (% of patients (N)) | 0.0 (0) | 11.1 (1) | 1.000 | 8.3 (1) | 0.0 (0) | 1.000 | 13.3 (2) | 40.0 (2) | 0.249 |
| Surgery | Time since last spine surgery (in years; mean (SD)) | 1.7 (2.6) | 2.2 (2.9) | 0.725 | 2.1 (3) | 0.6 (0.4) | 0.120 | 1.2 (0.8) | 2.1 (1.7) | 0.292 |
|                        | Screw length (in mm; mean (SD)) | 80.0 (8.2) | 84.6 (9.3) | 0.295 | 48.3 (8.2) | 52.0 (4.5) | 0.229 | 96.5 (14.1) | 102.0 (8.4) | 0.298 |
|                        | No. of decompressed vertebrae new (median (range)) | 0.0 (0-2) | 1.0 (0-5) | 0.697 | 2.0 (0-7) | 1.0 (0-2) | 0.301 | 0.0 (0-6) | 0.0 (0-0) | 0.146 |
|                        | No. of fused segments overall (median (range)) | 5.0 (2-9) | 5.0 (2-8) | 0.520 | 4.0 (1-9) | 5.0 (2-8) | 1.000 | 3.0 (2-11) | 6.0 (2-8) | 0.375 |
|                        | No. of screws one side overall (median (range)) | 5.0 (3-11) | 6.0 (3-9) | 0.583 | 6.0 (3-11) | 7.0 (4-10) | 0.755 | 5.0 (3-13) | 7.0 (5-10) | 0.096 |
|                        | Cement augmentation new (% of patients (N)) | 0.0 (0) | 45.5 (5) | 0.101 | 40.0 (6) | 20.0 (1) | 0.613 | 29.4 (5) | 0.0 (0) | 0.290 |
|                        | No. of levels with newly implanted cages (median (range)) | 0.0 (0-3) | 0.0 (0-4) | 0.909 | 1.0 (0-5) | 2.0 (0-4) | 0.853 | 0.0 (0-3) | 0.0 (0-1) | 0.831 |
|                        | Newly implanted vertebral body replacement (% of patients (N)) | 14.3 (1) | 0.0 (0) | 0.389 | 20.0 (3) | 0.0 (0) | 0.539 | 5.9 (1) | 0.0 (0) | 1.000 |

Potential predictors for an improvement in SIJ pain are shown in this table, differentiating between patients with improvement and patients with deterioration or unchanged status regarding SIJ pain. Italicized p values indicate statistical significance (p < 0.05); p values with asterisks did not survive correction for multiple comparisons using the Benjamini-Hochberg procedure with a false discovery rate of 25%.

Although literature reports a considerably high rate of lumbo-sacral non-fusion of up to 83%, the rate was considerably lower in our series, despite the analyzed groups and despite all cases being revision cases [1, 2, 16–18]. Although iliac screws became standard over the last decade in many centers due to superior construct endurance, screw prominence can induce pain, sitting difficulty, and can even require screw removal [1, 19, 20]. However, this was not the case in our iliac screw cohort. Despite new technical reports on reducing iliac screw prominence, S2AI screws do not harbor this issue at all [21]. Typically, and as our series does, such surgeries harbor not only an increased risk of perioperative surgical but also medical complications [22], despite the outcome usually being affected [23].

The current best level of evidence originates from a meta-analysis of 5 retrospective studies reporting a significantly lower rate of revision surgery, wound infection, and screw-related pain for S2AI versus iliac screws [22]. However, this study reported heterogeneous data of different centers with therefore even reported selection bias. Furthermore, although our study, as does the reported one, also reflects level III evidence, it not only reports on homogeneous single-center data but also investigates the potential of such instrumentation to treat SIJ pain originating from overstressed SIJ due to long cranial cantilever transmitted by the sacral instrumentation, which is a sparsely investigated issue. While another study showed that S2AI screws cause less postoperative SIJ pain compared with S1, S2, or L5 screws as the caudal end of the construct, our study actually shows that in this...
matter, S2AI screws are even superior to iliac screws [24]. With these results, our study is the first to prove not only that S2AI screws improve already existing SIJ pain, but also the superiority of this approach to iliac screws. Although both techniques bridge the SIJ, there seems to be some remaining minor but still sufficient movement within the SIJ in case that iliac screws are used. S2AI screws, in contrast, go directly through the SIJ and thereby cease even minor movement within the SIJ. This not only is relevant surgically considering SIJ pain as a disabling sequelae of sacral instrumentation but also shows us, impressively, how little movement within the SIJ is able to elicit this pain and how large the forces onto the SIJ after instrumentation down to the sacrum need to be. This is relevant especially when considering trans-SIJ plating as a sufficient treatment of SIJ pain.

In previous studies, such as the already mentioned meta-analysis, there was a difference in indications for spinopelvic instrumentation and in the rate of patients receiving anterior cage support [22], which was not the case in our series. Furthermore, the lower rate of SL of the S2AI screws compared with sacral and iliac screws is well in accordance with previous data, which seems mostly due to omission of connectors as a potential source of failure and the stronger cortical purchase by crossing 3 cortical bone structures. Crossing 3 cortical bone structures seems to be the main issue defining the persistence of the S2AI screws. A cadaveric study proved that 65-mm S2AI screws were as strong as 80-mm S2AI and 90-mm iliac screws, showing that the tricortical purchase and not the overall length seems decisive [25]. Considering additional anterior column support via the anterior, oblique, and lateral of posterior cages is regarded as essential to promote fusion and relieve stress from posterior elements [13, 26]. Despite our cohort reporting a direct comparison of 3 very homogeneous groups of one center, mean FU was long enough to be within the commonly reported time prone to SL. Nonetheless, our cohort does not report standardized questionnaires and can therefore only report the subjectively experienced gluteal pain plus the objectively detected SL.

Conclusion

In conclusion, even in shorter or mid-length lumbar or thoracolumbar constructs, S2AI might be considered superior to S2A and I instrumentation due to lower incidences of caudal SL and SIJ pain. However, the superiority not only is in the outcome, but the surgical technique also provides some advantages by allowing for small incisions due to diverging trajectories, less dissection, and in-line rod bending without the need for additional connectors. Because the trajectory is more demanding, thus potentially causing more anterior complications, we recommend spinal navigation to ease the approach, as done in all cases of this study. Although our data might be partially biased, the results and clinical experience are clear. We therefore advocate for the S2AI technique as the caudal end of longer instrumentations. Future prospective studies enrolling larger series should be conducted to confirm these initial results.

Authors’ contributions

Conceptualization: Sandro M. Krieg, Nico Sollmann, Bernhard Meyer
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Supervision: Bernhard Meyer

Funding information
Open Access funding provided by Projekt DEAL.
The study was financed by institutional grants.

Data availability
The data that support the findings of this study are available from the corresponding author, upon reasonable request.

Compliance with ethical standards

Conflict of interest
SK is consultant for Nexstim Plc (Helsinki, Finland) and Spineart Deutschland GmbH (Frankfurt, Germany) and received honoraria from Medtronic (Meerbusch, Germany) and Carl Zeiss Meditec (Oberkochen, Germany). SK and BM received research grants and are consultants for Brainlab AG (Munich, Germany). BM received honoraria, consulting fees, and research grants from Medtronic (Meerbusch, Germany), icotec ag (Altstätten, Switzerland), and Relievant Medsystemy Inc. (Sunnyvale, CA, USA), honoraria and research grants from Ulrich Medical (Ulm, Germany), honoraria and consulting fees from Spineart Deutschland GmbH (Frankfurt, Germany) and DePuy Synthes (West Chester, PA, USA), and royalties from Spineart Deutschland GmbH (Frankfurt, Germany). NS received honoraria from Nexstim Plc (Helsinki, Finland). SI is consultant for Brainlab AG (Munich, Germany). However, all authors declare that they have no conflict of interest regarding the materials used or the results presented in this study.

Ethics approval
All procedures performed in studies involving human participants were in accordance with the ethical standards of the institutional research committee (registration number: 159/16S) and with the 1964 Helsinki declaration and its later amendments or comparable ethical standards.

Consent to participate
Informed consent was not required due to the retrospective character of the study and conditions of our local ethics committee.

Consent for publication
Not applicable.

Code availability
Not applicable.

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