Meta-analyses comparing spine simulators with cadavers and finite element models by analysing range-of-motion data before and after lumbar total disc replacement

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Highlights
- Data from two different in vitro test methods for the same evaluation subjects were compared. It was investigated whether spine simulators with real human cadavers (SSCs) and finite element models (FEMs) provide the same data exemplarily for range of motion (ROM) before and after insertion of motion-retaining devices.
- Only fifty-nine percent of SSC meta-analyses show restored ROM after insertion of the device compared to the intact spinal segment. In FEM meta-analyses, ROM is restored in ninety percent.
- Ten percent of ROM analyses show significantly different data between SSCs and FEMs.
- With regard to the included studies, data generated by SSCs and FEMs cannot be used unrestricted as alternative and complementary data.
- Our analysis provides a new approach to compare data from associated test methods.

Abstract

Background: Range-of-motion (ROM) data generated by the in vitro test methods of spine simulators with cadavers (SSCs) and finite element models (FEMs) are used alternatively and complementarily for in vitro evaluations.
Introduction

In the approval process for motion-retaining devices to be applied to the human spine, animal studies are not suitable. The reason is the different mechanics of quadrupedal animals compared to bipedal humans during locomotion. Thus, approvals of devices for the spine depend on in vitro test results, although even mandatory tests do not provide data reflecting the complete function of spinal segments and the entire spine.

In order to acquire as much information as possible about a motion-retaining device, it is necessary to have detailed knowledge about the function of the device. One of the main functions of a spinal segment is motion, measured as range of motion (ROM), e.g., by the in vitro test methods of spine simulators with real human cadavers (SSCs) and finite element models (FEMs). FEMs are developed on the basis of SSC results; thus, SSC and FEM ROM data are alternatively and complementarily used for further evaluations. However, it is unclear whether SSC and FEM provide the same results for the same evaluation subjects, whether generated data on spinal function motion are reliable. We evaluated exemplary ROM data from SSCs and FEMs before (intact) and after and-ball-and-socket total disc replacement (bsTDR) to determine whether the same data are provided or to what extent the results differ.

BsTDR was developed as spinal disc substitute to retain motion within the intervertebral space if degenerative disc disease would lead to fusion surgery. Whereas an intervertebral disc is made of fibrocartilage for the performance of motion and further functions, bsTDR consists of at least two articulating prosthetic components to enable motion.

Why this review is important

In our extensive literature research, we identified no analysis comparing ROM data generated by SSCs and FEMs from intact and bsTDR-treated spinal segments to determine whether both in vitro test methods provide the same data for the same test subjects and consequently whether the two methods could be used alternatively and complementarily for further evaluations of the spine.

Material and methods

Study protocol

Our protocol, with a detailed description of our evaluation procedure, is available at https://spinefoundation.info/en/meta-analyses-database [1]: https://spinefoundation.info/medaba/studyprotocols?u=8.

Aim of Review: Our purpose is to compare exemplary segmental ROM data from SSCs and FEMs before and after ball-and-socket total disc replacement (bsTDR) to determine whether the two test methods provide the same data for the same evaluation subjects.

Key Scientific Concepts of Review: We performed 70 meta-analyses (MAs) and 20 additional comparative analyses based on data from 21 SSC studies used for 39 MAs and 16 FEM studies used for 31 MAs. Only fifty-nine percent (n = 23/39) of SSC MAs show a restored ROM after bsTDR, whereas in FEM MAs, the ROM is restored in ninety percent (n = 28/31). Among the analyses comparing data from the same spinal segments, motion directions and bsTDR, SSC and FEM data are significantly different in ten percent (n = 2/20). According to our results, data generated by SSCs and FEMs cannot be used as alternative and complementary data without restriction. The quality of the evaluation methods itself as well as potential technical reasons for the discrepant results were not our evaluation target. Further SSC and FEM data should be compared using the same approach.

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(including L4(6)/S1); single-level bsTDR interventions; and studies for which the full text of the publication was available.

Studies/publications were excluded if any of the following criteria were met:

1. Experimental research studies using the following evaluation techniques:
   - spine simulator without real human cadavers
   - human in vivo testing
   - animal study
2. Measurement of the cervical spine
3. No segmental measurement
4. Total disc replacement of any type other than ball and socket.

Search resources

For our search, we used the databases MEDLINE, Embase, Cochrane Library, Global Health Database (formerly CAB abstracts), World Health Organization regional databases, Google and Google Scholar, as well as doctoral theses and books addressing the topic.

Search terms

We searched for studies with the following terms in the title or abstract: “lumbar”, “spine”, “spinal”, “segment”, “vertebral”, “vertebra”, “biomechanics”, “biomechanical”, “measurements”, “measuring”, “numerical”, “movement”, “motion”, “total disc replacement”, “TDR”, “bsTDR”, “ball and socket”, “ball and cup”, “artificial disc”, “total disc”, “discectomy”, “spinal implant(s)”, “device”, “prosthesis”, “human”, “simulator”, “cadaveric”, “cadaver”, “test”, “testing”, “finite element”, “FE”, “analysis”, “in vitro”, “range of motion”, “ROM”, and “kinematic”.

Selection of studies

Our literature search for potentially eligible studies included title and abstract screening for preselection and full-text retrieval for a second round of selection.

Data extraction, management and assessment for risk of bias

An online system (OS) was created that allows a structured collection of data taken from complete original publications, including quality checks of the studies, evaluations of heterogeneity, and performance of MAs [1]. Basic information on the publications (e.g., title, author(s), date of publication and digital object identifier number) is stored within the OS, whereas duplicates are not stored. The OS provides tools to assess the quality of the studies using a short checklist (eight items) and long checklists (five additional items). Both checklists contain, among others, a question about bias. The short checklist includes a numeric system. When a predetermined score is achieved, the long checklist can be used for a continued quality assessment. Studies passing the long checklist are available for meta-analytic calculations, forest and funnel plots.

Response to missing data

We contacted authors (Cunningham [3], Chen [4], Chung [5], Wang [6]) to provide further information when the data reported in the studies were insufficient. The authors responded and sent the required data in all cases.

Assessment of statistical heterogeneity

Heterogeneity among studies was assessed using forest plots and the chi-squared test, with a statistical significance threshold of \( p < 0.01 \); \( I^2 \) statistics (ranging from 0 to 100%) were used to classify the degree of heterogeneity, with 0% indicating absence of heterogeneity, >0–25% indicating low heterogeneity, >25–50% indicating moderate heterogeneity, >50–75% indicating considerable heterogeneity, and >75–100% indicating high heterogeneity.

Data synthesis

A fixed-effects model was used for the MAs. SSC results were weighted according to the inverse of the variance. Depending on the data, FEM results were weighted equally. All MAs for ROM data from SSCs and FEMs and analyses for the comparison of the two test methods were performed in the OS.

Results

Studies were found for L3/4, L4/5, and L5/S1, with a total number of 21 SSC studies and 16 FEM studies (Table 1). The most evaluated level was L4/5 (n = 18/37). The most evaluated devices were the CAD (n = 12/37), the Maverick (n = 11/37) and the Prodisc-L (n = 17/37). The data of all analyses are given in Table 2 (39 MAs for SSCs), Table 3 (31 MAs for FEMs), and Table 4 (20 analyses for comparison of SSC and FEM). Due to the large number of MAs (n = 70), only one forest plot for each test method is shown (Fig. 1; further forest plots: https://spinefoundation.info/medaba/results?u=8). Table 5 presents an overview of the maintenance of ROM after bsTDR devices with regard to the CAD, Maverick, Prodisc-L and AbsTDR. A significant change in ROM (CROM) (\( p < 0.01 \)) after bsTDR compared to the intact spinal segment is interpreted as non-restored ROM. Table 6 shows an overview of significant and non-significant differences between SSCs and FEMs with regard to each spinal segment, motion direction and device. As an additional result of our analyses, Table 7 shows the best-performing devices for each test method, spinal segment and direction of motion (best performance defined as the p-value closest to 1 compared to the intact spinal segment).

SSC results

Table 2 shows the ROM results for the different spinal segments, directions of motion and devices. Furthermore, spinal segments and directions of motion with insufficient numbers of studies to perform MAs are shown. Hereafter, only results with restored ROM after bsTDR are presented, and they are always compared to intact spinal segments.

CAD

The CAD restores motion at L3/4 in lateral bending (CROM = 1.47\(^{\circ}\)). At L4/5, ROM is restored in overall sagittal motion (CROM = 0.82\(^{\circ}\)). At L5/S1, ROM is restored in overall sagittal motion (CROM = 1.05\(^{\circ}\)) and axial rotation (CROM = 0.75\(^{\circ}\)). In all MAs, the CAD increases ROM after insertion, up to 5.27\(^{\circ}\) in overall sagittal motion at L3/4. None of the eight MAs with CAD shows a decrease in ROM after insertion.

Maverick

At L4/5, ROM is restored in overall sagittal motion (CROM = 2.06\(^{\circ}\)), flexion (CROM = −0.38\(^{\circ}\)) and axial rotation (CROM = −0.03\(^{\circ}\)). At L5/S1, ROM is restored in overall sagittal motion (CROM = −1.07) and lateral bending (CROM = −0.22\(^{\circ}\)). The Maverick increases ROM after insertion in 4 out of 10 MAs, up to 2.7\(^{\circ}\) at L5/S1 in extension. In 6 out of 10 MAs, the Maverick decreases ROM after insertion, down to −3.86\(^{\circ}\) at L5/S1 in flexion.
Table 1

Studies included in the meta-analyses. SSCs spine simulators with cadavers, FEMs finite element models, segment spinal segment, CAD Charité artificial disc, e extension, f flexion, l lateral bending, a axial rotation, N/A not available, *prototype of an artificial disc in development, displ. contr. displacement controlled ROM with a required moment.

| Study          | Segment | Device(s) | Number of specimen | Applied moment (Nm) | Compressive load (N) |
|----------------|---------|-----------|--------------------|---------------------|---------------------|
| SSCs           |         |           |                    |                     |                     |
| Cunningham et al. [3] | L4/5 | CAD       | 8                  | 8                   | N/A                 |
| Demetropoulos et al. [16] | L4/5 | ProDisc-L  | 10                 | 10                  | 200                 |
| DiAngelo et al. [17] | L5/S1 | CAD, Maverick, ProDisc-L | 20 | 8 | 100 |
| Erkan et al. [18] | L5/S1 | Maverick  | 6                  | 7.5                 | 100                 |
| Gaffey et al. [19] | L4/5 | ProDisc-L  | 7                  | (f), (e)            | 400                 |
| Ha et al. [20] | L4/5 | activL    | 5                  | 8                   | 400                 |
| Hitchon et al. [21] | L4/5 | Maverick  | 7                  | 6                   | N/A                 |
| Ingalhalikar et al. [22] | L4/5 | Maverick  | 10                 | displ. contr.       | N/A                 |
| Kikawa et al. [23] | L4/5 | Triumph   | 7                  |                     | N/A                 |
| Kim et al. [24] | L4/5 | CAS       | 5                  | 8                   | 400                 |
| Le Huez et al. [10] | L4/5 | Maverick  | 6                  | 7                   | 400                 |
| Meyers et al. [15] | L5/S1 | ProDisc-L  | 12                 | 10                  | 600/1200            |
| Moldavsky et al. [25] | L4/5 | InOrbit   | 7                  | 8                   | N/A                 |
| O’Leary et al. [13] | L5/S1 | CAD       | 5                  | (f), (e)            | 400                 |
| Panjabi et al. [15] | L5/S1 | ProDisc-L  | 6                  | 10 (c.f), 8 (l,a)   | 400                 |
| Panjabi et al. [27] | L5/S1 | CAD       | 5                  | 10                  | 400                 |
| Takigawa et al. [28] | L4/5 | Maverick  | 7                  | 7.5                 | 400                 |
| Tsitsopulos et al. [29] | L5/S1 | ProDisc-L  | 12                 | (f), 6 (e)          | 400                 |
| Voronov et al. [30] | L3/4 | CAD       | 6                  | (f), (e), 5 (a)     | 400                 |
| Wilke et al. [27] | L3/4 | CAD, ProDisc-L, Prototype*  | 6 | 7.5 | 0 |
| Wong [31] | L5/S1 | Maverick, ProDisc-L | 7 | 7.6 | 100 |
| FEMS           |         |           |                    |                     |                     |
| Chen et al. [4] | L3/4 | ProDisc-L  | –                  | 10                  | 150                 |
| Chen et al. [32] | L3/4 | ProDisc-L  | –                  | 10                  | 400                 |
| Choi et al. [33] | L3/4 | ProDisc-L  | –                  | 10                  | 280                 |
| Choi et al. [34] | L3/4 | ProDisc-L, NewPro* | – | 10 | 280 |
| Chung et al. [5] | L4/5 | CAD, ProDisc-L | – | 6 | 400 |
| Di Mascio et al. [35] | L4/5 | Maverick | –                  | 10                  | N/A                 |
| Dooris et al. [36] | L3/4 | Maverick  | –                  | 6                   | 400                 |
| Goel et al. [37] | L5/S1 | CAD       | –                  | 10.6                | 400                 |
| Kim et al. [38] | L4/5 | CAD, Maverick, ProDisc-L | – | 5 | 400 |
| Knupfel et al. [39] | L5/S1 | ProDisc-L  | –                  | N/A                 | 734*                |
| Le Huez et al. [10] | L4/5 | Maverick  | –                  | 7                   | 400                 |
| Rundell et al. [40] | L3/4 | ProDisc-L  | –                  | 7.5                 | 500                 |
| Schmidt et al. [41] | L4/5 | CAD, Slide-Disc (mobile/immobile core) | – | 7.5 | 1000 |
| Wang et al. [6] | L4/5 | Triumph   | –                  | 6                   | 400                 |
| Zander et al. [42] | L4/5 | CAD, ProDisc-L, activL | 10(e,f), 7.5(l,a) | 500 |
| Zhong et al. [43] | L3/4 | ProDisc-L  | –                  | 10                  | 150                 |

**Prodisc-L**

At L4/5, ROM is restored in overall sagittal motion (−0.8°) and axial rotation (0.17°). At L5/S1, ROM is restored in overall sagittal motion (CROM = −0.64°), extension (CROM = 1.37°), lateral bending (CROM = 0.07°) and axial rotation (CROM = 0.08°). Prodisc-L increases ROM after insertion in 4 out of 8 MAs, up to 1.32° at L5/S1 in extension. In 4 out of 8 MAs, Prodisc-L decreases ROM, down to −2.52° at L5/S1 in flexion.

**AbsTDR**

ROM is restored at L3/4 in lateral bending (CROM = 1.01°), at L4/5 in overall sagittal motion (CROM = −0.8°), in extension (CROM = −0.24°), in flexion (CROM = 0.69°), and in lateral bending (CROM = −0.35°), and at L5/S1 in overall sagittal motion (CROM = −0.76°), lateral bending (CROM = −0.23°), and axial rotation (CROM = 0.2°). ROM after AbsTDR insertion is increased in 7 out of 13 MAs, up to 5.28° at L3/4 in overall sagittal motion. ROM decreases in 6 out of 13 MAs, down to −3.61° at L5/S1 in flexion.

**FEM results**

The results for the different spinal segments, motion directions and devices are presented in Table 3, including the results for segments and directions of motion with insufficient numbers of studies to perform MAs. Hereafter, we present only results with restored ROM after bsTDR, which are always compared to intact spinal segments.

**CAD**

ROM is restored at L4/5 in overall sagittal motion (CROM = 2.27°), extension (CROM = 3.32°), flexion (CROM = −1.0°), lateral bending (CROM = 1.16°) and axial rotation (CROM = −0.05°). The CAD increases ROM after insertion in 4 out of 5 MAs, up to 3.32° at L4/5 in extension. In 1 out of 5 MAs, ROM after CAD is decreased, down to −1.03° at L4/5 in flexion.

**Maverick**

ROM is restored at L4/5 in overall sagittal motion (CROM = 1.32°), extension (CROM = 3.32°), flexion (CROM = 0°), lateral bending (CROM = 0.29°) and axial rotation (CROM = −0.05°). The Maverick increases ROM after insertion in 4 out of 5 MAs, up to 1.32° at L4/5 in overall sagittal motion. In 1 out of 5 MAs, ROM after Maverick is decreased, down to −0.05° at L4/5 in axial rotation.

**Prodisc-L**

ROM is restored at L3/4 in overall sagittal motion (CROM = 3.84°), extension (CROM = 2.96°), flexion (CROM = 0.88°), lateral bending (CROM = 1.45°) and axial rotation (CROM = 1.59°), as well as at L4/5 in extension (CROM = 3.32°), flexion (CROM = −0.97°), lateral bending (CROM = 2.8°) and axial rotation (1.79°). The Prodisc-L increases ROM after insertion in 9 out of 10 MAs, up to 3.84° at L3/4 in overall sagittal motion. In 1 out of 10 MAs, ROM after Prodisc-L is decreased, down to −0.97° at L4/5 in flexion.
Table 2
Results of meta-analyses calculating the difference in range of motion of intact spinal segments and after ball-and-socket total disc replacement of each spinal segment, motion direction and device using spine simulators with cadavers (SSCs). CAD Charité artificial disc, bsTDR ball-and-socket total disc replacement, AbsTDR all bsTDR (activL, CAD, InOrbit, Maverick, NewPro, Prodisc-L, prototype of an artificial disc in development, Slide-Disc, Triumph). CROM change in range of motion in degree, CI confidence interval (l lower and u upper limit), n number of studies, reference no. reference number.

| SSCs          | L3/4 overall sagittal motion | L4/5 overall sagittal motion | L5/S1 overall sagittal motion |
|---------------|-----------------------------|-----------------------------|-------------------------------|
|               | extension flexion lateral bending axial rotation | extension flexion lateral bending axial rotation | extension flexion lateral bending axial rotation |
|               | CROM 5.27 | 1.47 | 2.01 | 0.82 | 2.19 | 1.21 | 1.05 | 0.75 |
|               | CI 95% I 3.22 | 0.32 | 1.49 | –2.04 | 1.2 | 0.7 | –1.43 | –0.06 |
|               | CI 95% u 7.32 | 2.61 | 2.54 | 3.69 | 3.18 | 1.72 | 3.54 | 1.56 |
|               | p-value <0.0001 | 0.012 | <0.0001 | 0.57 | 0.000014 | <0.00001 | 0.41 | 0.0069 |
|               | Chi² 2.22 | 1.58 | 0.26 | 3.57 | 17.63 | 6.54 | 1.37 | 0.71 |
|               | I² 55% | 36.8% | 0% | 72% | 94.3% | 84.7% | 0% | 0% |
|               | n 2 | 2 | 2 | 2 | 2 | 2 | 3 | 2 |
|               | reference no. | [2,30] | [2,30] | [3,24] | [3,24] | [13,17,27] | [17,27] |
|               | Maverick CROM | 2.06 | 1.26 | –0.38 | –1.31 | –0.03 | –1.07 | 2.7 | –3.86 | –0.22 | 0.36 |
|               | CI 95% I 0.09 | 0.42 | –1.81 | –2.29 | –2.29 | –0.59 | –2.27 | 2.04 | –4.41 | –0.61 | 0.1 |
|               | CI 95% u 4.04 | 2.11 | 1.06 | –0.32 | 0.54 | 0.14 | 3.35 | –3.31 | 0.16 | 0.63 |
|               | p-value 0.04 | 0.0033 | 0.6 | 0.0095 | 0.93 | 0.083 | <0.00001 | <0.00001 | 0.25 | 0.0072 |
|               | Chi² 2.41 | 4.9 | 8.6 | 14.22 | 1.01 | 4.9 | 0.94 | 14.26 | 23.13 | 4.54 |
|               | I² 0% | 59.1% | 76.7% | 78.9% | 0% | 59.2% | 0% | 86.0% | 91.4% | 56% |
|               | n 4 | 3 | 4 | 3 | 4 | 3 | 3 | 3 |
|               | reference no. | [10,21,22,28] | [21,22,28] | [21,22,28] | [10,21,22,28] | [10,21,28] | [18,17,31] | [17,18,31] | [17,18,31] | [17,18,31] |
|               | Prodisc-L CROM | –0.8 | | –0.32 | | –0.32 | | –0.32 | | 0.07 | 0.08 |
|               | CI 95% I –1.88 | | –2.14 | | –2.14 | | –2.14 | | 0.18 | –0.43 |
|               | CI 95% u 0.28 | | –1.02 | | –0.49 | | –0.81 | | 2.55 | –1.33 | 0.57 | 0.48 |
|               | p-value 0.15 | | <0.00001 | | 0.31 | | 0.39 | | 0.024 | 0.000032 | 0.79 | 0.7 |
|               | Chi² 0.72 | | 6.43 | | 12.82 | | 0.59 | | 0.61 | 0.07 | 10.48 | 5.84 |
|               | I² 68.9% | | 84.4% | | 0% | | 0% | | 0% | 71.4% | 48.6% |
|               | n 3 | | 4 | | 4 | | 4 | | 4 | 4 |
|               | reference no. | [16,19,29] | [16,19,29] | [16,19,29] | [15,17,26,31] | [17,31] | [17,31] | [15,17,26,31] | [15,17,26,31] |
|               | AbsTDR CROM | 5.28 | | 1.01 | | 1.62 | | 0.8 | | 2.49 | –3.61 | –0.23 | 0.2 |
|               | CI 95% I 3.41 | | 0 | | 1.18 | | –1.54 | | 0.67 | –0.19 | –0.72 | 0.14 | –1.57 | –4.12 | –0.53 | –0.02 |
|               | CI 95% u 7.16 | | 2.02 | | 2.05 | | –0.06 | | 0.18 | 1.56 | 0.03 | 0.57 | 0.06 | 3.08 | –3.09 | 0.06 | 0.43 |
|               | p-value <0.0001 | | 0.05 | | <0.0001 | | 0.034 | | 0.26 | 0.12 | 0.07 | 0.0012 | 0.068 | <0.00001 | <0.00001 | 0.12 | 0.078 |
|               | Chi² 2.07 | | 0.47 | | 51.84 | | 27.27 | | 14.95 | 83.94 | 29.98 | 7.37 | 1.2 | 19.22 | 55.3 | 13.69 |
|               | I² 51.8% | | 0% | | 0% | | 80.7% | | 81.7% | 66.5% | 88.1% | 70% | 5% | 0% | 89.6% | 91% | 56.2% |
|               | n 2 | | 2 | | 11 | | 6 | | 6 | 11 | 10 | 8 | 3 | 3 | 6 |
|               | reference no. | [2,30] | [2,30] | [2,30] | [3,10,16,19,20,21,22,23,24,25,28] | [22,24,28] | [22,24,28] | [22,23,24,25,28] | [23,24,25,28] | [13,15,17,18] | [17,18,31] | [17,18,31] | [15,17,18,26] | [15,17,18,26] | [17,18,31] | [17,18,31] | [27,29,31] | [27,29,31] |

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### Table 3
Results of meta-analyses calculating the difference in range of motion of intact spinal segments and after ball-and-socket total disc replacement of each spinal segment, motion direction and device using finite element models (FEMs).

| FEMs | L3/4 | L4/5 | L5/S1 |
|------|------|------|------|
|      | overall sagittal motion | extension | flexion | lateral bending | axial rotation | overall sagittal motion | extension | flexion | lateral bending | axial rotation | overall sagittal motion | extension | flexion | lateral bending | axial rotation |
| CAD  | CROM  | 2.27  | 3.32  | −1.03 | 2.85  | 1.16 |
|      | CI 95% l | 0.16  | −0.33 | −3.1 | 1.73  | −1.03 |
|      | CI 95% u | 4.38  | 6.96  | 1.05 | 3.97  | 3.34 |
|      | p-value | 0.033 | 0.074 | 0.33 | <0.00001 | 0.3 |
|      | I² | 1.95  | 6.22  | 2.17 | 3.95  | 1.87 |
|      | n | 4 | 4 | 4 | 3 | 3 |
|      | reference no. | [5,38,41, 42] | [5,38,41,42] | [5,38,41,42] | [5,41,42] | [5,41,42] |
| Maverick | CROM  | 1.32  | 1.32  | 0 | 0.29  | −0.05 |
|      | CI 95% l | −1.41 | −0.9 | −1.58 | −1.08 | −0.36 |
|      | CI 95% u | 4.05 | 3.54 | 1.58 | 1.66 | 0.25 |
|      | p-value | 0.34 | 0.24 | >0.99999 | 0.68 | 0.72 |
|      | I² | 0% | 51.8% | 0% | 49.4% | 0% |
|      | n | 3 | 3 | 2 | 2 |
|      | reference no. | [10,35,38] | [10,35,38] | [10,35,38] | [10,35] | [10,35] |
| Prodisc | CROM  | 3.84  | 2.96  | 0.88 | 1.45  | 1.59 |
|      | CI 95% l | −4.34 | −0.2 | −4.17 | 0.29 | −0.11 |
|      | CI 95% u | 12.01 | 6.12 | 5.92 | 2.61 | 3.28 |
|      | p-value | 0.36 | 0.067 | 0.73 | 0.014 | 0.067 |
|      | I² | 0.000016 | 0.058 | 0.42 | 0.098 | 0.15 |
|      | n | 5 | 5 | 5 | 3 |
|      | reference no. | [4,32,33, 34,36,40,43] | [4,32,33,40,43] | [4,32,33,40,43] | [5,38,42] | [5,38,42] |
|      | p-value | 0.13 | 0.85 | 0.25 | 0.41 | 0.34 |
|      | I² | 0% | 0% | 0% | 0% | 0% |
|      | n | 7 | 7 | 6 | 6 |
|      | reference no. | [37,39] | [34,36,40,43] | [36,40,43] | [40,43] | [40,43] |
| AbsTDR | CROM  | 2.88  | 2.42  | 0.46 | 1.09  | 1.1 |
|      | CI 95% l | −4.79 | −0.74 | −4.19 | −0.79 | −1.52 |
|      | CI 95% u | 10.55 | 5.58 | 5.1 | 2.97 | 3.72 |
|      | p-value | 0.46 | 0.13 | 0.85 | 0.25 | 0.41 |
|      | I² | 11.08 | 8.66 | 6.63 | 2.62 | 4.8 |
|      | n | 7 | 7 | 6 | 6 |
|      | reference no. | [4,32,33, 40,43] | [4,32,33,40,43] | [4,32,33,40,43] | [5,6,10,35,38,41,42] | [5,6,10,35,38,41,42] |

**CAD**: Charité artificial disc, **bsTDR**: ball-and-socket total disc replacement, **AbsTDR**: all bsTDR (activL, CAD, InOrbit, Maverick, NewPro, Prodisc-L, prototype of an artificial disc in development, Slide-Disc, Triumph). **CROM**: change in range of motion in degree, **CI**: confidence interval ([l <i>lower</i> and <i>u</i> upper limit]), **n**: number of studies, **reference no.**: reference number.
Table 4
Comparison of results of meta-analyses for each spinal segment, motion direction and device, using spine simulators with cadavers (SSCs) and finite element models (FEMs). CAD Charité artificial disc, bsTDR ball-and-socket total disc replacement, AbsTDR all bsTDR (activL, CAD, InOrbit, Maverick, NewPro, Prodisc-L, prototype of an artificial disc in development, Slide-Disc, Triumph), D ROM difference between SSC and FEM, CROM change in range of motion in degree, CI confidence interval, n number of studies, reference no. reference number, italic not available.

| CAD | L3/4 | Overall sagittal motion | Extension flexion lateral bending axial rotation | L4/5 | Overall sagittal motion | Extension flexion lateral bending axial rotation | L5/S1 | Overall sagittal motion | Extension flexion lateral bending axial rotation |
|-----|------|------------------------|-----------------------------------------------|------|------------------------|-----------------------------------------------|------|------------------------|-----------------------------------------------|
|     | CAD  | D in ROM                | CROM                                          |      | D in ROM                | CROM                                          |      | D in ROM                | CROM                                          |
|     |      |                        | lower CI 95%                                  |      |                        | lower CI 95%                                  |      |                        | lower CI 95%                                  |
|     |      |                        | upper CI 95%                                  |      |                        | upper CI 95%                                  |      |                        | upper CI 95%                                  |
|     |      |                        | p-value                                       |      |                        | p-value                                       |      |                        | p-value                                       |
|     |      |                        | Chi²                                           |      |                        | Chi²                                           |      |                        | Chi²                                           |
|     |      |                        | I²                                             |      |                        | I²                                             |      |                        | I²                                             |
|     |      |                        | n                                              |      |                        | n                                              |      |                        | n                                              |
|     |      |                        | reference no.                                  |      |                        | reference no.                                  |      |                        | reference no.                                  |
|     | Maverick | D in ROM                | CROM                                          |      | D in ROM                | CROM                                          |      | D in ROM                | CROM                                          |
|     |      |                        | lower CI 95%                                  |      |                        | lower CI 95%                                  |      |                        | lower CI 95%                                  |
|     |      |                        | upper CI 95%                                  |      |                        | upper CI 95%                                  |      |                        | upper CI 95%                                  |
|     |      |                        | p-value                                       |      |                        | p-value                                       |      |                        | p-value                                       |
|     |      |                        | Chi²                                           |      |                        | Chi²                                           |      |                        | Chi²                                           |
|     |      |                        | I²                                             |      |                        | I²                                             |      |                        | I²                                             |
|     |      |                        | n                                              |      |                        | n                                              |      |                        | n                                              |
|     |      |                        | reference no.                                  |      |                        | reference no.                                  |      |                        | reference no.                                  |
|     | Prodisc-L | D in ROM                | CROM                                          |      | D in ROM                | CROM                                          |      | D in ROM                | CROM                                          |
|     |      |                        | lower CI 95%                                  |      |                        | lower CI 95%                                  |      |                        | lower CI 95%                                  |
|     |      |                        | upper CI 95%                                  |      |                        | upper CI 95%                                  |      |                        | upper CI 95%                                  |
|     |      |                        | p-value                                       |      |                        | p-value                                       |      |                        | p-value                                       |
|     |      |                        | Chi²                                           |      |                        | Chi²                                           |      |                        | Chi²                                           |
|     |      |                        | I²                                             |      |                        | I²                                             |      |                        | I²                                             |
|     |      |                        | n                                              |      |                        | n                                              |      |                        | n                                              |
|     |      |                        | reference no.                                  |      |                        | reference no.                                  |      |                        | reference no.                                  |
|     | AbsTDR | D in ROM                | CROM                                          |      | D in ROM                | CROM                                          |      | D in ROM                | CROM                                          |
|     |      |                        | lower CI 95%                                  |      |                        | lower CI 95%                                  |      |                        | lower CI 95%                                  |
|     |      |                        | upper CI 95%                                  |      |                        | upper CI 95%                                  |      |                        | upper CI 95%                                  |
|     |      |                        | p-value                                       |      |                        | p-value                                       |      |                        | p-value                                       |
|     |      |                        | Chi²                                           |      |                        | Chi²                                           |      |                        | Chi²                                           |
|     |      |                        | I²                                             |      |                        | I²                                             |      |                        | I²                                             |
|     |      |                        | n                                              |      |                        | n                                              |      |                        | n                                              |
|     |      |                        | reference no.                                  |      |                        | reference no.                                  |      |                        | reference no.                                  |
AbsTDR ROM is restored at L3/4 in overall sagittal motion (CROM = 2.88°), extension (CROM = 2.42°), flexion (CROM = 0.46°), lateral bending (CROM = 1.09°) and axial rotation (CROM = 1.1°), as well as at L4/5 in overall sagittal motion (CROM = 1.31°), extension (CROM = 1.84°), flexion (CROM = -0.53°), lateral bending (CROM = 1.36°) and axial rotation (CROM = 0.67). ROM after bsTDR insertion is increased in 10 out of 11 MAs, up to 2.88° at L3/4 in overall sagittal motion. In 1 out of 11 MAs, ROM after insertion is decreased, down to -0.53° at L4/5 in flexion.

Comparison of SSC and FEM results

The results for the different spinal segments, directions of motion and devices are presented in Table 4. Table 6 provides an overview of the statistical significance of the differences between the SSC and FEM results. The detailed results are presented here:

**CAD**

There was no significant difference between SSC and FEM at L4/5 in overall sagittal motion (p = 0.42), lateral bending (p = 0.39) or axial rotation (p = 0.97).
There was no significant difference between SSC and FEM at L4/5 in overall sagittal motion (p = 0.67), extension (p = 0.96), flexion (p = 0.73), lateral bending (p = 0.064) or axial rotation (p = 0.95).

There was a significant difference between SSC and FEM at L4/5 in overall sagittal motion (p = 0.000041). There was no significant difference at L4/5 in lateral bending (p = 0.012) or axial rotation (p = 0.2).
There was a significant difference between SSC and FEM at L5/S1 in overall sagittal motion (p < 0.00001). There was no significant difference at L3/4 in overall sagittal motion (p = 0.55), lateral bending (p = 0.94) or axial rotation (p = 0.7), as well as no significant difference at L4/5 in overall sagittal motion (p = 0.13), extension (p = 0.26), flexion (p = 0.31), lateral bending (p = 0.28), or axial rotation (p = 0.76).

**Best performing bsTDR**

As shown in Table 7, the CAD performs best in SSCs at L4/5 and L5/S1 in overall sagittal motion. The Maverick performs best in SSC at L4/5 in lateral bending and axial rotation and in FEMs at L4/5 in overall sagittal motion, extension, flexion, lateral bending, and axial rotation. The Prodisc-L performs best in SSCs at L5/S1 in extension, flexion, lateral bending, and axial rotation.

**Overall results**

Approximately 59% of all SSC MAs show restored ROM (n = 23/39 MAs), whereas in FEM MAs, ROM is restored in 90% (n = 28/31) (Table 5).

The 20 analyses comparing SSCs and FEMs with available data for the same spinal segments, motion directions and bsTDR devices show a statistically significant difference of 10% (n = 2/20) between SSCs and FEMs. For further details, the performance of the Prodisc-L at L4/5 in overall sagittal motion and the performance of AbsTDR at L5/S1 in overall sagittal motion are shown in Table 4.

**Results by spinal level**

Altogether, the L3/4 segment was tested in 24% of studies (n = 9/37), L4/5 in 49% (n = 18/37), and L5/S1 in 27% (n = 10/37). No suitable studies are available for L1/L2 and L2/L3 MAs (Table 1).

Eighty percent (n = 16/20) of analyses for the comparison of SSCs and FEMs are at L4/5 and L5/S1. The number of studies is sufficient (n = 18/37) to perform MAs for ROM data with inserted CAD, Maverick, Prodisc-L, and AbsTDR devices in all directions of motion, except for extension and flexion for the CAD and Prodisc-L in SSC. Our analyses for the comparison of SSCs and FEMs in this segment show a statistically significant difference of 6% (n = 1/16) (Table 6). For the comparison of SSCs and FEMs with regard to L3/4 and L5/S1 segments, only results for AbsTDR are available.

**Discussion**

BsTDR was developed to retain spinal segmental motion in mainly middle-aged patients with degenerative disc disease and at the same time to reduce pain. However, Johnsen et al. [44] found no correlation between lumbar segmental movement and patient reported outcomes.

Another type of disc replacement than bsTDR provides motion by the features of the material between two metal plates, i.e. without articulating prosthetic components. Depending on the number of available studies with segmental ROM data and to avoid data differences from different types of disc replacement, we included bsTDR study data only to our MAs.

### Table 7

Best performing devices with regard to spinal segments and motion directions. Best performing is defined as the p-value closest to 1 compared to the intact spinal segment. In the case of same p-value, the closest range of motion (ROM) of ball-and-socket total disc replacement to the ROM of the intact spinal segment is defined as best performing. SSCs spine simulators with cadavers, FEMs finite element models, CAD Charité artificial disc, greyed out not enough studies to perform meta-analyses.

| SSCs       | overall sagittal motion | extension | flexion | lateral bending | axial rotation |
|------------|-------------------------|-----------|---------|----------------|---------------|
| L3/4       |                         |           |         |                |               |
| L4/5       | CAD                     |           |         |                | Maverick      |
| L5/S1      | CAD                     | Prodisc-L | Prodisc-L| Prodisc-L      |               |

| FEMs       | overall sagittal motion | extension | flexion | lateral bending | axial rotation |
|------------|-------------------------|-----------|---------|----------------|---------------|
| L3/4       |                         |           |         |                |               |
| L4/5       | Maverick                | Maverick  |         |                | Maverick      |
| L5/S1      |                         | Maverick  |         |                |               |

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AbsTDR

There was a significant difference between SSC and FEM at L5/S1 in overall sagittal motion (p < 0.00001). There was no significant difference at L3/4 in overall sagittal motion (p = 0.55), lateral bending (p = 0.94) or axial rotation (p = 0.7), as well as no significant difference at L4/5 in overall sagittal motion (p = 0.13), extension (p = 0.26), flexion (p = 0.31), lateral bending (p = 0.28), or axial rotation (p = 0.76).

**Results of the bsTDR evaluation**

**CAD**

Comparing the results of the CAD, Maverick and Prodisc-L, the CAD is the best performing bsTDR in 15% (n = 2/13; overall sagittal motion at L4/5 and L5/S1) (Table 7). The CAD restores ROM in 50% (n = 4/8) of SSC MAs and 80% (n = 4/5) of FEM MAs (Table 5). The comparison of SSCs and FEMs shows no significant difference in any of the analyses (n = 3/3) (Table 6).

**Maverick**

The Maverick is the best performing bsTDR in 54% (n = 7/13) (Table 7). The Maverick restores ROM in 50% of SSC MAs (n = 5/10) and in all of the FEM MAs (n = 5/5) (Table 5). The comparison of SSCs and FEMs shows no significant difference in any of the analyses (n = 3/3) (Table 6).

**Prodisc-L**

The Prodisc-L is the best performing bsTDR in 31% (n = 4/13) (Table 7). The Prodisc-L restores ROM in 75% (n = 6/8) of SSC MAs and 90% (n = 9/10) of FEM MAs (Table 5). The comparison of SSCs and FEMs shows a significant difference in 33% (n = 1/3) of analyses (Table 6).

**AbsTDR**

The analyses of AbsTDR allow us to compare evaluations of SSCs and FEMs independently of the specific disc. ROM is restored in 62% (n = 8/13) of SSC MAs and 91% (n = 10/11) of FEM MAs (Table 5). The comparison of SSCs and FEMs shows a significant difference in 11% (n = 1/9) of analyses (Table 6).
Method and evaluation basics

Although more cervical discs than lumbar discs are implanted in humans, our literature search revealed more studies with ROM data for the lumbar spine than for the cervical spine. For this reason, we included single SSC and FEM studies of the lumbar spine in our ROM data MAs.

The cadavers used in the SSC studies of our MAs were confirmed to meet the quality criteria proposed by Wilke et al. for evaluating ROM [7].

According to Jones et al. [8], FEMs were validated by the comparison of data from SSCs and/or from the literature to provide a convincing body of evidence that the calculated results are meaningful.

We evaluated data from SSCs and FEMs to identify whether both evaluation methods provide the same data for the same test subjects, but not the quality of the test methods or issues of the test setups.

For our comparison, we used all available segmental ROM data for extension and flexion as well as overall sagittal motion data. Not all studies provide extension and flexion data separately. A division by two of the overall sagittal motion data is incorrect because the ROM of extension is smaller than the ROM of flexion. Consequently, the provided extension and flexion data were summarized to obtain overall sagittal motion data. In accordance with White and Panjabi [9], lateral bending and axial rotation data were used unilaterally. For this reason, and because some included studies present unilateral data only, provided bilateral motion data were divided by two.

In our analysis, we performed MAs of separately obtained SSC- and FEM-based ROM data of various spinal segments, motion directions and devices to compare SSCs with FEMs. SSC and FEM data from single studies included in our MAs for ROM of intact spinal segments and after bsTDR are independent of each other and are therefore not directly compared. Only one study included in our MAs provides SSC data with dependent FEM data [10], each used as a single dataset for the groups of our meta-analytic calculations.

Results

According to Le Huec et al. [10], SSC and FEM testing are two different but complementary approaches to conduct biomechanical studies of the spine for ROM analyses. Whereas cadaveric tests provide data on motion responses of a spinal segment to varied loading conditions, FEM analyses, using computational models, tend to simulate the biomechanical behaviour of spinal segments.

In order for SSCs and FEMs to be used as complementary test methods, it is necessary to assume that these methods provide the same data for the same evaluated subjects, at least without statistically significant differences.

We compared SSC and FEM ROM data from lumbar spinal segments before and after bsTDR devices to identify whether they are equivalent or to what extent they differ. Our meta-analytic calculations and further analyses with data from the included studies led to unexpected results, demonstrating that SSCs and FEMs do not consistently provide the same results. While 90% (n = 28/31) of FEM MAs show restored ROM, the MAs for SSCs demonstrate restored ROM in only 59% (n = 23/39). A statistically significant difference in the comparison of SSCs and FEMs with data from the same spinal segments, motion directions and bsTDR was found in 10% (n = 2/20).

Recommendations for the standardization of in vitro tests were presented in 1998 [7], i.e., before all studies included in our MAs were published. One reason for the different ROM results in the comparison of SSC and FEM data could be inconsistent validation of the included FEM studies. However, Dreischarf et al. [11] compared and re-validated eight published FEMs and stated that “predicted median segmental intervertebral rotations and disc pressures were in good agreement with measured in vivo values”. Furthermore, technical difficulties in SSC studies and differences in the test setups of included SSC and FEM studies can cause different ROM results. According to Wilke et al. [7], the applied moment should be high enough to achieve normal ROM: 7.5 Nm is suggested for the lumbar spine. In our included studies, the moment varies but remains within the elastic range [7]. The applied moments and compressive loads before (intact) and after bsTDR are consistent in each study. Regarding the compressive load, O’Leary et al. [13] recommend 400 N. Patwardhan et al. [14] describe a decrease in ROM with increasing compressive load. However, according to Meyers et al. [15], the compressive load does not significantly affect ROM in extension, flexion or axial rotation. Volkheimer et al. [12] showed in a review that even with the same testing conditions, the results may vary in some cases.

No further MAs or other analyses with ROM data for the comparison of SSCs and FEMs are available in the literature to discuss our results, which demonstrates the uniqueness of our evaluation.

In summary, SSCs and FEMs do not provide ROM data for unrestricted complementary or alternative evaluations of spinal segments. Otherwise, it is assumed that single SSC and FEM studies can provide high-quality data. Whereas our comparisons of SSC and FEM ROM data are limited to intact spinal segments and inserted bsTDR devices, other motion-retaining devices for the spine could provide ROM results more conveniently.

In order to obtain generally high-quality results from evaluations for, e.g., the approval of motion-retaining devices for the human spine, it will be necessary to further compare data from SSCs and FEMs and to study the origin of the differences for the sake of obtaining more uniform data and thus reliable results. The data and approach of our evaluation can support further evaluations and developments, although the in vivo situation of the spine is complex, not well understood, and variable between individuals [7].

Our evaluation has limitations: The search includes studies in English and German only. For some motion segments and directions of motion, we did not find appropriate studies for further MAs.

Conclusion

Our analyses provide a new approach to compare data from associated in vitro test methods with the same evaluation subjects. Comparing SSC and FEM ROM study data of the same spinal segments, motion directions and devices before and after bsTDR, we detected differences between SSC and FEM data in the endpoint ROM. The conclusion with regard to the included studies is that SSC and FEM ROM data cannot be used unrestricted as alternative and complementary data. We did not evaluate any technical issues as potential reasons for the ROM data differences or whether SSC and FEM test methods can provide high-quality ROM data. Further developments and evaluations should lead to greater consistency of data between SSCs and FEMs.

Compliance with Ethics Requirements

This article does not contain any studies with human or animal subjects.
Declaration of Competing Interest

The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

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