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1 2	Efficacy, Safety and Economics of Bracing after Spine Surgery: A Systematic Review of the Literature
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10 11	Corresponding author: Jefferson R. Wilson, jeffersonwilson7@gmail.com, 30 Bond Street, Toronto, Ontario, Canada M5W 1W8
12	Abstract
13 14 15	Background Context: Bracing is often used after spinal surgery to immobilize the spine, improve fusion, and relieve pain. However, controversy exists regarding the efficacy, necessity and safety of various bracing techniques in the post-surgical setting.
16 17 18	Purpose: In this systematic review, we aimed to compare the effectiveness, safety and cost-effectiveness of postoperative bracing versus no postoperative bracing following spinal surgery in patients with several common operative spinal pathologies.
19	Study Design/Setting: Systematic Review
20	Patient Sample: N/A
21	Outcome Measures: N/A
22	Methods: A systematic search was conducted of MEDLINE. Embase and the Cochrane
23	Collaboration Library from 1970 to May 2017, supplemented by manual searching of the
24	reference list of relevant studies and previously published reviews. Studies were
25	included if they compared disability, quality of life, functional impairment, radiographic
26	outcomes, cost-effectiveness and/or complications between patients treated with
27	postoperative bracing versus those not receiving any postoperative bracing. Each article
28	was critically appraised independently by 2 reviewers, and the overall body of evidence

- was rated using guidelines outlined by the Grading of Recommendation Assessment, Development and Evaluation (GRADE) Working Group. 29
- 30

- 1 Results: Of the 858 retrieved citations, 5 studies met inclusion criteria and were
- 2 included in this review, consisting of 4 randomized controlled trials and 1 prospective
- 3 cohort study. Low to moderate evidence suggests that there are no significant
- 4 differences in most measures of disability, pain, quality of life, functional impairment,
- 5 radiographic outcomes, and safety between groups. Isolated studies reported
- 6 statistically significant and inconsistent differences between groups with respect to Neck
- 7 Disability Index at 6 weeks postoperatively and/or Short Form-36 Physical Component
- 8 Score at 1.5, 3, 6, and 12 months postoperatively.
- 9 **Conclusions:** Based on limited evidence, postoperative bracing does not result in
- 10 improved outcomes following spinal surgery. Future high quality randomized trials will
- 11 be required to confirm these findings.
- 12 Keywords: outcomes; postoperative bracing; surgery; spinal pathology; complications.

13 Introduction

- 14 Bracing is routinely used after surgery for a number of spinal pathologies, including degenerative
- 15 disease of the lumbar and cervical spine, thoracolumbar fractures and scoliosis.[1] The intended
- 16 goal of this practice is to immobilize the spine, relieve pain, improve fusion rates, and remind
- 17 patients to avoid certain activities that may compromise their recovery.[1, 2] However, a number
- 18 of important complications can arise from bracing, including dysphagia, nerve palsies, pressure
- 19 ulcers, and skin rashes.[1] Furthermore, braces can be uncomfortable for some patients as well as
- 20 costly.[2]
- 21 Given the paucity of high-quality comparative studies, it is unclear whether postoperative
- 22 bracing can effectively limit and restrict spinal movements, reduce rates of pseudoarthrosis and
- 23 optimize patient recovery. Certain advances in spinal surgery have allowed for rigid internal
- stabilization of the spine and, arguably, have decreased the requirement for external
- 25 immobilization. Although these techniques may be sufficient to achieve successful fusion, there
- 26 may still be a role for postoperative bracing in higher risk patients, including those who smoke,
- 27 suffer from osteoporosis or require an extensive multilevel surgery.
- As a result of the limited body of evidence available, spine surgeons often base their decision to
- use postoperative bracing on their own clinical experience and training.[1] This finding is
- 30 supported by a survey that highlighted substantial disagreement among spinal surgeons with
- respect to the appropriate type, duration and indication for use of postoperative bracing after
- anterior cervical spine surgery.[3] Given the heterogeneity in management strategies, there is a
- need to synthesize results from high quality studies and establish recommendations surrounding
- 34 care following spinal surgery.
- 35 This systematic review addresses 4 key questions (KQs). KQ1: What is the efficacy and
- 36 effectiveness of postoperative bracing compared with no bracing based on disability, pain,

- 1 quality of life and functional outcomes? KQ2: What is the impact of postoperative bracing
- 2 compared with no bracing on radiographic outcomes? KQ3: What is the safety profile of
- 3 postoperative bracing compared with no bracing? KQ4: What is the cost-effectiveness of
- 4 postoperative bracing? Importantly, this systematic review will assess the overall strength of
- 5 the evidence using methodology developed by the Grading of Recommendation
- 6 Assessment, Development and Evaluation (GRADE) working group.

7 Methods

8 Eligibility Criteria

- 9 Population
- 10 Our review targeted studies including patients undergoing surgery for any spinal pathology,
- 11 including cervical and lumbar degenerative disease, trauma, oncology and adolescent idiopathic

X

- scoliosis. Studies were excluded if patients under study were treated non-operatively (Table 1).
- 13 Intervention and Comparison
- 14 This review focused on studies that had an intervention group who received postoperative
- bracing, and a control group who received standard of care and no postoperative bracing (Table
- 16 1).

17 *Outcomes*

- 18 For KQ1, we sought studies that considered the clinical efficacy of postoperative bracing by
- 19 measuring patient disability, pain, quality of life, and/or functional outcomes. For KQ2, we
- 20 focused on studies that assessed radiographic outcomes, including fusion rate, sagittal alignment,
- and range of motion. For KQ3, we sought studies that compared complication rates and adverse
- events between the intervention and control groups. For KQ4, we focused on studies that
- 23 examined various measures of cost-effectiveness including incremental cost-effectiveness ratio
- 24 and cost per unit of outcome (Table 1).

25 Study Characteristics

- For KQ1, 2 and 3, we sought comparative studies (i.e. randomized controlled trials, cohort
- studies) designed to evaluate differences between a postoperative bracing group and a control
- group. To be included, studies needed to have at least 10 patients per group. Case reports,
- 29 nonclinical studies, and animal studies were excluded. For KQ4, we focused on full economic
- 30 studies. For all KQs, abstracts, editorials, letters, narrative and systematic reviews were
- 31 excluded. Duplicate publications of the same study that did not report on different outcomes
- 32 were also excluded.

33 Information Sources

- 1 A systematic search of MEDLINE, Embase and Cochrane Collaboration Library was conducted
- 2 to identify relevant studies. Manual searching of the reference lists of included studies and
- 3 previously published reviews was also conducted to ensure all relevant studies were located.

4 Search Strategy

- 5 The search strategy was first developed in MEDLINE and then appropriately modified for the
- 6 other databases. We used the following search terms to search all databases: Orthotic Devices
- 7 AND Spinal Diseases AND Postoperative Complications/Care AND Treatment Outcome or
- 8 Outcome Assessment. Only studies involving humans, written in English and published in peer-
- 9 review journals between 1970 and May 2017 were considered for inclusion, with no other limits
- 10 applied. A detailed search strategy is provided in the supplemental digital material.

11 Study Selection

- 12 All abstracts and titles were reviewed and sorted by our predefined inclusion criteria. Studies
- 13 were classified as relevant, possibly relevant, or irrelevant. Full text investigation of all relevant
- 14 and possibly relevant studies was done for further clarification.

15 Data Extraction and Synthesis

- 16 The following data were extracted from each included article: study design; patient sample and
- 17 characteristics, including diagnosis, surgical summary and type of bracing; outcome assessment
- tools; follow-up schedule; drop-out rate; and results of association, including standard deviation,
- 19 odds ratio, confidence intervals, and p-values.

20 Risk of Bias in Individual Studies

- 21 The class of evidence for each article was rated (Class I, II, III, IV) independently by 2 reviewers
- 22 using criteria outlined by the Journal of Bone and Joint Surgery for therapeutic studies and
- 23 modified to encompass both methodological quality and risk of bias. Randomized controlled
- trials were rated based on patient allocation, intention to treat analysis, independent or blinded
- assessment, whether co-interventions were applied equally, rates of follow-up, statistical power,
- and control for possible confounding. Prospective cohort studies were rated based on
- 27 independent or blinded assessment, whether co-interventions were applied equally, rates of
- follow-up, statistical power, and control for possible confounding. Due to the nature of the
- 29 intervention, studies were rated as having independent or blinded assessment if surgeons were
- 30 blinded to the randomization group until after surgery, patients were blinded to the
- 31 randomization group until day of admission of surgery or after surgery, and/or radiologists
- 32 reviewing radiographs were blinded to the randomization group.

33 Risk of Bias Across Studies

1 The overall body of evidence was assessed using a scoring system developed by the **<u>GRADE</u>**

2 working group with recommendations from the Agency for Healthcare Research and Quality

- 3 (AHRQ). This methodology allows for an assessment of the overall strength of the evidence
- 4 and is particularly valuable for highlighting critical knowledge gaps.

5 The initial strength of the overall body of evidence was graded as "high" if half or more of the

- 6 studies were randomized controlled trials and "low" if the majority of studies were observational
- 7 studies. The body of evidence was downgraded 1, 2, or 3 levels if there was risk of bias, results
- 8 were inconsistent or consistency was unknown, the evidence was indirect, the effect estimates
- 9 were imprecise (e.g. wide confidence intervals), or if there was publication bias. If no
- 10 downgrades were made, the body of evidence was upgraded 1, 2 or 3 levels based on large
- 11 magnitude of effect, dose-response gradient or if all plausible biases would decrease the
- 12 magnitude of an apparent effect.
- 13 The final rating of the body of evidence expresses our confidence in the estimate of effect and
- 14 the impact of further research on this topic. An overall strength of "high" means we have high

15 confidence that the evidence reflects the true effect. Further research is very unlikely to change

- 16 our confidence in the estimate of effect. The overall strength of "moderate" means we have
- 17 moderate confidence that the evidence reflects the true effect. Further research may change our
- confidence in the estimate of effect and may change the estimate. A grade of "low" means we
- 19 have low confidence that the evidence reflects the true effect. Further research is likely to change
- 20 the confidence in the estimate of effect and likely to change the estimate. A grade of
- 21 "insufficient" means that evidence is either unavailable or does not permit a conclusion.

22 **Results**

23 Study Selection

- 24 The initial electronic search yielded a total of 853 citations. Five additional citations were
- 25 identified through reference scanning. After initial review of abstracts and titles, 841 studies did
- not meet our inclusion criteria. Following full text investigation, an additional 12 studies were
- excluded because 1) they were not comparative studies; 2) patients were not treated surgically; 3)
- there was no postoperative comparison of intervention and control groups; 4) they had a different
- 29 outcome of interest; 5) they had no control group; 6) they were a duplicate publication with no
- new results; and/or 7) they were not in English. A total of 5 studies were deemed relevant
- 31 following this review process.

32 Study Characteristics

- 33 For KQ1, we identified 4 studies (3 randomized controlled trials, 1 prospective cohort)
- discussing the effect of postoperative bracing on disability, pain, quality of life and functional
- outcomes.[4-7] Sample sizes ranged from 33 to 257 surgical patients with mean ages between
- 43.9 and 72.7 years. All patients were diagnosed with degenerative cervical myelopathy or

- 1 radiculopathy, or degenerative disease of the lumbar spine. Bracing included Philadelphia
- 2 collars, cervical collars, and lumbar corsets for differing lengths of time. Various outcome
- 3 measures were used across the studies, with the Short Form-36 (SF-36) Physical Component
- 4 Score (PCS) reported the most frequently (n = 4), [4-7] followed by the SF-36 Mental
- 5 Component Score (MCS) (n = 3), [4, 6, 7] SF-36 subscales (n = 3), [4, 6, 7] Neck Disability Index
- 6 (NDI) (n = 2), [4, 5] neck pain (n = 2), [4, 5] and arm pain (n = 2). [4, 5]
- 7 For KQ2, a total of 5 studies (4 randomized controlled trials, 1 prospective cohort) met our
- 8 inclusion criteria.[4-8] These studies were designed to assess the impact of postoperative bracing
- 9 compared with no bracing on radiographic outcomes. Sample sizes ranged from 33 to 257
- 10 surgical patients with mean ages between 14.3 and 72.7 years. Patients were diagnosed with
- 11 degenerative cervical myelopathy or radiculopathy, degenerative disease of the lumbar spine, or
- 12 adolescent idiopathic scoliosis. Bracing included Philadelphia collars, cervical collars, body
- 13 casts, and lumbar corsets for various lengths of time. Fusion rate was the most frequently
- 14 reported outcome measure (n = 3).[4, 5, 7]
- 15 For KQ3, we identified 3 studies (2 randomized controlled trials, 1 prospective cohort)
- 16 examining the safety profile of postoperative bracing compared with no bracing.[5-7] Sample
- sizes ranged from 50 to 257 surgical patients with mean ages between 43.9 and 72.7 years.
- 18 Patients were diagnosed with degenerative cervical myelopathy or radiculopathy, or degenerative
- 19 disease of the lumbar spine. Bracing included Philadelphia collars, cervical collars, and lumbar
- 20 corsets for various lengths of time. Outcome measures included complications (n = 3)[5-7] and
- revision surgery or second procedure (n = 2).[5, 7]
- 22 No studies met our inclusion criteria for KQ4 on the cost-effectiveness of postoperative bracing.

23 **Risk of Bias**

- 24 We critically appraised the 5 studies included in our review. The inter-rater reliability was
- 25 80%; disagreement on the fifth study, by Yee et al., surrounding the "intention to treat"
- 26 <u>analysis was resolved through discussion. It was noted that although 90 patients were</u>
- 27 randomized, only 72 were included in their analysis, and therefore the study did not use
- 28 **intention to treat analysis.**[7] Of **the studies included**, 4 were considered Class II and 1 was
- rated Class III. The 4 Class II studies were randomized controlled trials, and were downgraded
- 30 from Class I because they did not include intent-to-treat analysis, independent or blind
- assessment, adequate sample size, random sequence generation and/or statement of concealed
- allocation, had unreported follow-up rates or follow-up rates < 80%, and/or did not control for
 possible confounders. The Class III study was a prospective cohort study that was downgraded
- from Class II because co-interventions were not applied equally and follow-up was not reported.

35 **Results of Individual Studies**

- 36 *KQ1*: What is the efficacy and effectiveness of postoperative bracing compared with no bracing
- 37 based on disability, pain, quality of life and functional outcomes?

1 <u>Degenerative cervical myelopathy or radiculopathy</u>

- 2 Three studies compared disability, pain, quality of life and/or functional outcomes between the
- postoperative bracing and non-bracing groups in patients with degenerative cervical myelopathy
- 4 or radiculopathy.[4-6]
- 5 According to Abbott et al., patients who received postoperative bracing had better NDI scores at
- 6 6 weeks after surgery compared to patients who did not receive postoperative bracing (mean
- 7 difference between groups -4.4, 95% CI -8.6 to -0.2, p = 0.042).[4] However, Campbell et al.,
- 8 reported that patients in the non-braced group had better NDI scores at 6 weeks after surgery
- 9 compared to patients in the braced group (p = 0.008).[5]
- 10 Abbott et al. also found that patients in the control group had better SF-36 PCS scores at 6 weeks
- 11 (mean difference between groups 5.8, 95% CI 0.8-10.7, p = 0.025), 3 (mean difference between
- 12 groups 6.8, 95% CI 0.4-13.1, p = 0.038), 6 (mean difference between groups 7.4, 95% CI 1.4-
- 13 13.4, p = 0.017), and 12 months (mean difference between groups 7.5, 95% CI 0.3-14.6, p =
- 14 0.041) after surgery compared to patients in the postoperative bracing group.[4] In contrast,
- 15 Campbell et al. and Hida et al. found no significant differences in SF-36 PCS scores between the
- 16 two groups at all time points assessed (Campbell et al. 6 months p = 0.481, 12 months p = 0.260,
- 17 24 months p = 0.279; Hida et al. p = 0.537).[5, 6]
- 18 Two studies evaluated differences in various subscales of the SF-36 between a postoperative
- bracing and a control group.[4, 6] A single study by Abbott et al. reported significantly better
- 20 SF-36 Bodily Pain (BP) scores in the postoperative bracing group at 6 (mean difference between
- 21 groups 21.4, 95% CI 4.4-38.5, p=0.016) and 12 (mean difference between groups 17.5, 95% CI
- 22 1.7-33.2, p=0.031) months, as well as better SF-36 Social Functioning (SF) scores at 12-months
- 23 (mean difference between groups 16.5, 95% CI 0.1-32.9, p=0.049) than in the non-bracing
- 24 group.[4] The other six subscales (Physical Functioning (PF), General Health (GH), Role
- 25 Limitations Physical (RP), Vitality (VT), Role Limitations Emotional (RE) and Mental Health
- 26 (MH)) were not significantly different between treatment groups at all time points assessed (1.5,
- 27 3, 6, 12 and 24 months).[4] A second study by Hida et al., identified no differences between the
- collar-fixation group and the control group with respect to SF-36 BP subscale (p=0.848).[6]
- 29 Other measures of disability, pain, quality of life, and functional impairment, including SF-36
- 30 MCS, Visual Analog Scale (VAS), Japanese Orthopedic Association (JOA) recovery rate, Falls
- Efficacy Scale (FES), unipedal balance standing test, and neck and arm pain were not
- 32 significantly different between the postoperative bracing and non-bracing groups (Table 3).[4-6]
- 33 <u>Degenerative disease of the lumbar spine</u>
- A single study by Yee at al. examined pain and quality of life outcomes for patients with a
- degenerative disease of the lumbar spine.[7] There were no significant differences in the Dallas
- Pain Questionnaire (DPQ, daily activity category p = 0.34, work/leisure category p = 0.67,

- anxiety-depression category p = 0.17, social category p = 0.40), SF-36 PCS (p = 0.30), SF-36
- 2 MCS (p = 0.57) and SF-36 subscales (PF p = 0.38, BP p = 0.28, GH p = 0.23, RP p = 0.41, VT p
- 3 = 0.25, SF p = 0.79, RE p = 0.86, MH p = 0.30) between the braced and non-braced groups at all
- 4 time points assessed.[7]

5 *KQ2:* What is the impact of postoperative bracing compared with no bracing on radiographic6 outcomes?

- 7 <u>Degenerative cervical myelopathy or radiculopathy</u>
- 8 Three studies assessed the radiographic outcomes between the postoperative bracing and non-
- 9 bracing groups in patients with degenerative cervical myelopathy or radiculopathy.[4-6]
- 10 Abbott et al. reported no significant differences in cervical range of motion (ROM) (p > 0.05),
- fusion rate (p = NR), and sagittal alignment (p = NR) between the two groups.[4] Campbell et al.
- 12 also found no significant differences in fusion success between the bracing and non-bracing
- 13 groups (p = NR).[5] Similarly, Hida et al. concluded that there were no significant differences in
- 14 ROM (p = 0.61) or decrease in lordotic angle C2-7 (p = 0.82) between groups.[6]
- 15 Degenerative disease of the lumbar spine
- 16 Yee at al. found no significant differences in fusion rate at 12 months (p = 0.8) or 24 months (p =
- 17 0.9) postoperatively between patients who wore a lumbar corset and those who did not.[7]
- 18 Adolescent idiopathic scoliosis
- 19 Based on a single study, there was no significant difference in mean loss of spinal curve
- 20 correction between the braced and non-braced groups at all time points assessed in patients with
- 21 adolescent idiopathic scoliosis (p = NR).[8]
- 22 KQ3: What is the safety profile of postoperative bracing compared with no bracing?
- 23 <u>Degenerative cervical myelopathy or radiculopathy</u>
- 24 Campbell et al. reported rates of instrumentation failure, graft extrusion, and second procedures

including revisions, removals, reoperations, supplemental fixations, and external bone growth

stimulators,[5] while Hida et al. considered all perioperative complications including surgical

- site infection, epidural hematoma, and C5 palsy.[6]
- 28 Both Campbell et al. and Hida et al. identified no significant differences in the incidence of
- 29 complications between the bracing and non-bracing groups (Campbell et al. no events of
- instrumentation failure or graft extrusion in either group, p = NC; Hida et al. p = 0.53).[5, 6] In
- addition, Campbell et al. found no significant differences between groups in the rate of revision
- surgery (p=0.653), removals (p=0.724), reoperations (p=1.000), supplemental fixations
- 33 (p=0.286) or any second operation (p = 0.184).[5]

1 <u>Degenerative disease of the lumbar spine</u>

- 2 Based on a single study, there were no significant differences in rates of revision surgery, a
- 3 second procedure or complications between the braced and non-braced groups at all time points
- 4 assessed (p = 0.8).[7] Revision surgery included later-stage revision surgery due to symptomatic
- 5 nonunion or later-stage hardware removal due to prominence/bursitis, and complications
- 6 included intraoperative incidental durotomy, intraoperative pedicle-screw-placement issues, new
- 7 postoperative radiculopathy, early postoperative pulmonary embolism, wound
- 8 seroma/hematoma, deep wound infection or persistent lumbar radiculopathy postoperatively.
- 9 KQ4: What is the cost-effectiveness of postoperative bracing?
- 10 No studies were identified that evaluated the cost-effectiveness of postoperative bracing.

11 Summary of Evidence

- 12 The overall quality of evidence ranged from "insufficient" to "moderate". Evidence was
- 13 downgraded due to risk of bias, unknown or inconsistent consistency of results, and/or imprecise
- 14 effect estimates (e.g. wide confidence intervals).
- 15 Based on low evidence, postoperative bracing in patients with degenerative cervical myelopathy
- 16 or radiculopathy does not result in improved SF-36 MCS, VAS, JOA recovery rate, sagittal
- alignment, ROM, decrease in lordotic angle C2-7, and rate of revision surgery or second
- 18 procedure outcomes. Based on moderate evidence, postoperative bracing does not result in
- 19 improved neck or arm pain, fusion rate, and incidence of complications in this patient
- 20 population.
- 21 Low evidence suggests no improvement in DPQ, SF-36 PCS, SF-36 MCS, SF-36 subscales,
- fusion rate, incidence of complications, and rate of revision surgery or second procedure
- following postoperative bracing in patients with a degenerative disease of the lumbar spine.
- 24 Based on low evidence, postoperative bracing is not associated with improved loss of spinal
- 25 curve correction in adolescents with idiopathic scoliosis.

26 **Discussion**

- 27 The use of bracing after surgery for a variety of spinal pathologies remains controversial with
- 28 limited evidence available to the surgeon to make an informed decision. Historically cited
- 29 reasons to use bracing include to limit mobility and stabilize the spine, improve fusion rates,
- 30 prevent graft dislodgement or subsidence, reduce postoperative pain and optimize outcomes.
- 31 Bracing, however, can be uncomfortable, lead to social isolation and be associated with
- 32 complications such as dysphagia, nerve palsies and pressure ulcers. This review summarizes the
- current literature on the efficacy, safety and cost-effectiveness of bracing after spinal surgery
- 34 <u>using rigorous methodology and is the first to synthesize results using methodology</u>

1 **proposed by the GRADE working group**. This knowledge is valuable in a clinical setting, and

- 2 can be used by clinicians to determine the most appropriate postoperative management
- 3 strategies. Clinical judgement, however, is still required to determine whether a patient may
- 4 benefit from additional external immobilization.

5 Degenerative cervical myelopathy or radiculopathy

- 6 Based on this review, postoperative bracing of the cervical spine has no impact on 1) most
- 7 measures of pain, disability, functional impairment, and quality of life; 2) radiographic outcomes
- 8 such as fusion rate and range of motion; and 3) rates of complications and reoperations. In the
- 9 study by Abbott et al., patients who received postoperative bracing exhibited superior
- 10 improvements in neck disability (NDI at 6 weeks) and various metrics of quality of life (SF-36
- PCS at 6 weeks to 12 months, SF-36 SF at 12-months and SF-36 BP at 6 to 12 months) than
- 12 those who did not.[4] These findings can be partly explained by psychological factors, including
- a sense of security provided by the brace, increased coping mechanisms, improved functional
- self-efficacy, and less fear avoidance.[4] In contrast, Campbell et al identified that patients with
- 15 postoperative bracing had worse NDI scores, likely due to the discomfort and disability
- 16 associated with wearing a brace.[5]
- 17 Previous studies in healthy subjects have demonstrated that cervical bracing reduces velocity of
- eye movements and causes deterioration in the anterior to posterior body sway induced by
- vibration of the calf muscles.[9, 10] Given these findings, it is hypothesized that restricting
- 20 cervical motion through external immobilization may significantly impair static postural control
- and disturb balance during dynamic movement. In the study by Abbott et al., however, there
- 22 were no differences between a bracing group and a control group with respect to the unipedal
- 23 balance standing test.[4]
- 24 Biomechanical studies have indicated that cervical collars help to restrict motion during routine
- 25 activities and stabilize the spine.[11-13] However, early mobilization exercises can prevent spine
- contracture and improve range of motion after surgery.[6] In studies by Hida et al. and Abbott et
- al., there were no significant differences in cervical range of motion between the postoperative
- bracing group and the control group.[4, 6] Although range of motion often decreases following
- surgery, this is more likely due to fusion and fixation techniques, damage to the cervical flexors
- 30 and extensors and injury to the facet joints. Postoperative bracing may also help to decrease the
- risk of graft or cage migration, maintain spinal alignment and improve fusion rates.
- 32 Advancements in surgical procedures, however, have allowed for internal stabilization of the
- 33 spine and may have decreased the requirement for external immobilization. For example, the use
- of anterior plates has shown to increase fusion and decrease subsidence rates by limiting motion
- between the graft and vertebral bodies.[14, 15] This finding was confirmed by Campbell et al.
- 36 who reported no significant difference in rates of fusion between a bracing and non-bracing
- 37 group;[5] these results question the need for postoperative bracing, especially in patients
- undergoing internal stabilization. There may still be a role for postoperative bracing in patients at

- 1 a higher risk of pseudoarthrosis and disease progression, including those who smoke, have had a
- 2 previous spine operation, and/or are treated without rigid internal fixation.[16-21] Furthermore,
- 3 surgeons are more likely to use postoperative bracing following a multilevel anterior cervical
- 4 discectomy and fusion (ACDF) (76%) compared to a single level ACDF (55%).[3]

5 Degenerative Disease of the Lumbar Spine

- 6 Based on this review, the use of a lumbar corset following surgery for degenerative lumbar
- 7 disease has no impact on pain, disability, functional impairment, quality of life, radiographic
- 8 outcomes, incidence of complications, and rate of reoperations. Postoperative bracing is often
- 9 used in this population to relieve pain, limit mobility, improve fusion rates and optimize
- 10 outcomes; however, the study by Yee et al. indicated no advantage or disadvantage to the use of
- a lumbar corset.[7] There may still be a role for bracing in patients at a higher risk of nonunion
- 12 or pseudoarthrosis, such as those who smoke or require a multilevel fusion.

13 Adolescent Idiopathic Scoliosis

- 14 Historically, molded plaster braces were used to correct the curve following posterior fusions
- 15 without instrumentation and maintain this correction until solid bony fusion. Techniques
- 16 proposed in the study by Christodoulou et al., including the use of Harrington distraction rods,
- 17 however, has decreased the need for postoperative external bracing due to more rigid internal
- 18 fixation.[8] In this review, a single study examined postoperative bracing in an adolescent
- 19 idiopathic scoliosis population and found no differences in radiographic outcomes between
- 20 patients who received bracing versus those who did not. Further investigation, however, is
- 21 needed to determine the effectiveness of bracing in this population based on other outcome
- 22 measures.
- 23 Our finding that bracing may not confer additional benefits following spine surgery will have
- 24 relevant applications in a clinical setting. First, complications such as skin reactions, dysphagia,
- 25 pressure ulcers and nerve palsies, as well as costs associated with bracing can be eliminated.
- 26 Second, if postoperative bracing is not required, there may be less of an impact on activities of
- 27 daily living, decreased social isolation, body anxiety, self-perception and body image issues and
- an improved ability to return to work or school following surgery.

29 Strengths and Limitations

- 30 To our knowledge, no other reviews have evaluated the merits of bracing in the post-surgical
- 31 setting for patients with various spinal pathologies <u>using the GRADE approach</u>. <u>This</u>

32 methodology allows for rigorous evaluation of the overall strength of the evidence and

- 33 **helps to identify critical knowledge gaps in the literature (e.g** a lack of high-quality
- 34 comparative studies and limited data on the cost-effectiveness of postoperative bracing).
- Furthermore, the majority of current studies published on this topic have moderately high to high

- 1 risk of bias and imprecise estimates of effect (or estimates with unknown precision). Consistency
- 2 of results is also largely unknown as results are based on single studies.
- 3 Our review also has its limitations. First, our search was restricted to studies published in English
- 4 and, as a result, some articles with relevant titles or abstracts were excluded. Second, although
- 5 results were separated based on patient population, the type and length of bracing, as well as
- 6 surgical technique varied substantially among studies, preventing pooling of data and meta-
- 7 analysis.

8 Conclusions

- 9 Based on the results of this review, postoperative bracing does not result in improved outcomes
- 10 after spine surgery in patients with various spinal pathologies. Although some outcomes were
- significantly different between bracing and non-bracing groups, firm conclusions cannot be made
- 12 due to small sample sizes, risk of bias and low quality of evidence. Finally, given the paucity of
- 13 studies available, no conclusions can be made regarding the cost-effectiveness of bracing after
- 14 surgery.

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31		
32		
	T ¹	

- 33 Figure 1. Flow diagram of study selection
- 34

Characteristic	Inclusion	Exclusion				
Population	Patients w/ any spinal pathology treated	Patients treated non-				
	surgically and followed postoperatively	operatively				
	(e.g. scoliosis, spinal trauma, cervical					
	myelopathy)					
Intervention	Postoperative bracing	·. 0				
Comparison	No postoperative bracing					
	Standard of care					
Outcome	KQ1: Efficacy and effectiveness	KQ1: Subjective neurological				
	• Disability (e.g. NDI)	status, patient satisfaction,				
	• Pain (e.g. VAS)	improvement of symptoms				
	• Quality of life (e.g. SF-36)					
	• Functional outcomes (e.g. JOA,					
	mJOA, Nurick)					
	KQ2: Radiographic Outcomes					
	Fusion rate					
	Sagittal alignment					
	• ROM					
	KQ3: Safety					
	• Complications or adverse events					
	(e.g. dysphagia, hardware, skin					
	complications, reoperation, revision	l				

1 Table 1. Inclusion and exclusion criteria for studies reviewed

	surgery, infection, hematoma)	
	KQ4: Cost-effectiveness	
	• Incremental cost-effectiveness ratio	
	(or similar)	
	• Cost per unit of outcome	
Study design	KQ1, 2, 3: Comparative studies (e.g. RCT,	Case reports
	prospective cohort, case-control studies)	Nonclinical studies
	designed to compare a postoperative	Animal studies
	bracing group with a control group; $n \ge 10$	
	per group.	
	KQ4: Full economic studies	
Publication	Studies published in peer-review journals	Abstracts, editorials, letters
	and in English	Duplicate publications of the
	×Õ	same study that do not report
		on different outcomes
		Narrative or systematic
	20	reviews

1 JOA = Japanese Orthopedic Association; mJOA = modified Japanese Orthopedic Association;

- 2 NDI = neck disability index; ROM = range of motion; SF-36 = short form-36; VAS = visual
- 3 analog scale

1 Table 2. Characteristics of included studies

		Mean				Outcome		Dron
Populatio	Sex,	Age		~		Sucome		ыор
n, n	%	(Range	Diagnosis	Surgery	Bracing	Factors	Follow-up	-out,
		SD)*			.0	Assessed		n†
		, 5D)						
Brace	М,	53.4	Cervical	ACDF with	Philadelphi	KQ1: Borg	1.5, 3, 6,	55%
(n = 17)	53%	(NR,	spondylosis (n =	interbody	a Collar	CR-10 (neck	12, and 24	(n =
		13)	8), Cervical disk	cage	daytime	and arm	months	18)
			herniation $(n = 4)$,		only for 6	pain), NDI,	postoperati	
			Cervical		weeks	SF-36 (PCS	ve	
			degenerative disc			and MCS),		
			disease $(n = 5)$			FES, unipedal		
Control	М,	47.3	Cervical	-		balance		
(n = 16)	69%	(NR,	spondylosis (n =			standing test		
		11)	5), Cervical disk			KQ2: CROM,		
			herniation $(n = 4)$,			fusion rate,		
	Populatio n, n Brace (n = 17) Control (n = 16)	Populatio Sex, n, n % Brace M, (n = 17) 53% Control M, (n = 16) 69%	Mean Populatio Sex, Age n, n % (Range , SD)* , SD)* Brace M, 53.4 (n = 17) 53% (NR, 13) 13) Control M, 47.3 (n = 16) 69% (NR, 11) 11 11	PopulatioSex,Age (Range , SD)*Diagnosisn, n%(Range , SD)*DiagnosisBraceM,53.4Cervical $(n = 17)$ 53%(NR,spondylosis (n = 13)8), Cervical disk herniation (n = 4), Cervical degenerative disc disease (n = 5)ControlM,47.3Cervical (NR,spondylosis (n = 11)11)5), Cervical disk herniation (n = 4),	MeanPopulatioSex,Age (Range , SD)*DiagnosisSurgeryBraceM, 53.4 CervicalACDF with(n = 17) 53% (NR,spondylosis (n =interbody13)8), Cervical diskcageherniation (n = 4),Cervicaldegenerative discdisease (n = 5)ControlM, 47.3 Cervical(n = 16)69%(NR,spondylosis (n =11)5), Cervical diskherniation (n = 4),	MeanPopulatioSex,Age (Range , SD)*DiagnosisSurgeryBracing n, n %(Range , SD)*SD)*BracingMarcingBracingBraceM,53.4CervicalACDF withPhiladelphi $(n = 17)$ 53%(NR,spondylosis (n =interbodya Collar13)8), Cervical diskcagedaytimeherniation $(n = 4)$,only for 6Cervicalweeks(n = 16)69%(NR,spondylosis $(n =$ interbody11)5), Cervical diskherniation $(n = 4)$,interbodyinterbody	MeanMeanOutcomePopulaioSex,AgeBracingBracingBracingn, n%(RangeSurgeryBracingBracingAssessed, SD)* \cdot \cdot AssessedAssessedBraceM,53.4CervicalACDF withPhiladelpiKQ1: Borg(n = 17)53%(NR,spondylosis (n =interbodya CollarCR-10 (neck)1308), Cervical diskcagedaytimeand armherniation (n = 4),LervicalweeksSF-36 (PCS)ControlM,47.3CervicalweeksSF-36 (PCS)(n = 16)G9%(NR,spondylosis (n =LervicalweeksSF-36 (PCS)(n = 16)69%(NR,spondylosis (n =Lervicalstanding teststanding test(n = 16)69%(NR,spondylosis (n =LervicalLervicalstanding test(n = 16)11)5), Cervical diskLervicalLervicalLervicalKQ2: CROM(n = 16)11)5), Cervical diskLervicalLervicalLervicalLervical(n = 16)11)5), Cervical diskLervicalLervicalLervicalLervical(n = 16)11)5), Cervical diskLervicalLervicalLervicalLervical(n = 16)11)5), Cervical diskLervicalLervicalLervicalLervical(n = 16)11)5), Cervical diskLervicalLervicalLervical	MeanPopulatioSex,Age (Range , SD)*DiagnosisSurgery SurgeryBracing BracingFactors FactorsFollow-up Follow-upBraceM,53.4CervicalACDF withPhiladelphiKQ1: Borg1.5, 3, 6,(n = 17)53%(NR,spondylosis (n =interbodya CollarCR-10 (neck12, and 2413)8), Cervical diskcagedaytimeand armmonthsherniation (n = 4),Cervicalcagedaytimeand armpostoperatiControlM,47.3CervicalweeksSF-36 (PCS)ve(n = 16)69%(NR,spondylosis (n =standing testtrue11)5), Cervical diskcagestanding testtrue(n = 16)69%(NR,spondylosis (n =standing testKQ2: CROM,(n = 16)69%(NR,spondylosis (n =standing testkcy2: CROM,

		G	Mean				Outcome		Drop
Authors &	Populatio	Sex,	Age	Diagnosis	Surgery	Bracing	Factors	Follow-up	-out,
Year	n, n	%	(Range	C		C	Assessed	Ĩ	
			, SD)*			×	Assessed		nţ
				Cervical			sagittal		
				degenerative disc			alignment		
				disease $(n = 7)$			KQ3: None		
							KQ4: None		
Campbell et	Brace	М,	44.3	Single-level	Decompressio	Cervical	KQ1: NDI,	1.5, 3, 6,	NR
al., 2008	(n =	43.6	(NR,	radiculopathy or	n and	collar	neck and arm	12, and 24	
(Prospectiv	149)	%	8.8)	myelopathy	arthrodesis		pain scales,	months	
e cohort)‡	Control	М,	43.3	COX	using		SF-36	postoperati	
	(n =	49.1	(NR,	G	allograft and		KQ2: Fusion	ve	
	108)	%	9.0)	~	anterior		success		
					cervical plate		KQ3: Second		
					or		procedure§,		

			Mean				0.4		D
Authors &	Populatio	Sex,	Age				Outcome		Drop
Year	n, n	%	(Range	Diagnosis	Surgery	Bracing	Factors	Follow-up	-out,
		70	(100.80				Assessed		n†
			, SD)*			X			
					arthoplasty	X	instrumentatio		
					C	5	n failure, graft		
					S		extrusion		
					0		KQ4: None		
Christodoulo	Brace	NA	NR	Adolescent	Posterior	Plaster	KQI: None	3, 6, 12, and	NR
u et al.,	(n = 25)			idiopathic	decompressi	body cast	KQ2: Spinal	24 months	
1987 (RCT)	Control	NA	NR	scoliosis with	on and fusion	for 6	curve	postoperati	
	(n = 25)			thoracic curves \geq	with	months	KQ3: None	ve	
				35°	Harrington		KQ4: None		
					instrumentati				
					on				
					augmented				

		a	Mean				Outcome		Drop
Authors &	Populatio	Sex,	Age	Diagnosis	Surgery	Bracing	Factors	Follow-up	-out,
Year	n, n	%	(Range				Assessed		n†
			, SD)*			X			I
					by a Cotrel				
					bar or by				
					sublaminal				
					Luque wires				
Hida et al.,	Brace	М,	72.0	Cervical	Double-door	Philadelphi	KQ1: VAS,	0.5, 3, 6,	18%
2017 (RCT)	(n = 45)	73%	(NR,	myelopathy	cervical	a collar	JOA, SF-36	and 12	(n =
			8.7)	secondary to	laminoplasty	for 2	(PCS and	months	16)
	Control	М,	71.6	multisegmental	without	weeks	MCS)	postoperati	
	(n = 45)	62%	(NR,	cervical	instrumentati		KQ2: ROM	ve	
			9.6)	spondylotic	on		(total,		
				stenosis			extension and		
							flexion),		

			Mean				Outcome		Drop
Authors &	Populatio	Sex,	Age	Diagnosis	Surgary	Procina	Factors	Follow up	out
Year	n, n	%	(Range	Diagnosis	Surgery	Dracing	Tactors	Follow-up	-001,
			, SD)*			×	Assessed		nŢ
						2	lordotic angle		
					C	G	C2-7		
					J.		KQ3:		
							Perioperative		
							complications		
				0			ſ		
				Č.			KQ4: None		
Yee et al.,	Brace	М,	52	Spondylosis/stenos	Posterior	Canvas	KQ1: DPQ,	12 and 24	20%
2008 (RCT)	(n = 46)	43%	(NR,	is (n = 11),	lumbar spinal	lumbar	SF-36	months	(n =
		**	15.2)	degenerative	arthrodesis	corset	KQ2: Fusion	postoperati	18)
			**	spondylolisthesis		full-time	rate	ve	
				(n = 13), isthmic		for 8	KQ3: Revision		

			Mean				Outcome		Drop
Authors &	Populatio	Sex,	Age	Diagnosis	Surgamy	Draging	Eastors	Follow up	out
Year	n, n	%	(Range	Diagnosis	Surgery	Dracing	Factors	ronow-up	-001,
			, SD)*			×	Assessed		nŢ
				spondylolisthesis		weeks	surgery††,		
				(n = 8), junctional			complications		
				syndrome $(n = 1)$,			* * * *		
				pseudarthrosis (n			KQ4: None		
				= 2),					
				iatrogenic/post-op					
				instability $(n = 2)$					
				††					
	Control	M.	53	Spondylosis/stenos					
	(n-44)	54%	(NR	is $(n - 12)$					
	(11 – ++)	**	(INK,	13(11-12),					
			15.4)	degenerative					
			**	spondylolisthesis					

			Mean				Outcome		Dron
Authors &	Populatio	Sex,	Age	Diagnosis	Surgary	Proving	Factors	Follow up	out
Year	n, n	%	(Range	Diagnosis	Surgery	Bracing	Factors	ronow-up	-out,
			, SD)*			×	Assessed		n†
				(n = 12), isthmic		~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~			
				an on dy lo listhosis					
				spondylolistnesis					
				(n = 5), junctional					
				syndrome $(n = 2)$,					
				pseudarthrosis (n					
				= 2),					
				iatrogenic/post-op					
				instability $(n = 1)$,					
				congenital					
				stenosis (n = 1)					
				* †					

1 ACDF = anterior cervical discectomy and fusion; CROM = cervical range of motion; DPQ = Dallas pain questionnaire; FES = falls

2 efficacy scale; JOA = Japanese Orthopedic Association; MCS = mental component score; NDI= neck disability index; NR = not

- 1 reported; PCS = physical component score; ROM = range of motion; SD = standard deviation; SF-36 = short form-36; VAS = visual
- 2 analog scale
- 3 * Age in years
- 4 † Drop-out before end of follow-up
- 5 ‡ Postoperative care, including immobilization techniques and activity restrictions, was left to the discretion of the attending surgeon
- 6 § Included revisions, removals, reoperations, supplemental fixations, and external bone growth stimulators
- 7 ¶ Included surgical site infection, epidural hematoma, and C5 palsy
- 8 ** Based on patients who completed follow-up
- 9 *†*[†] Included later-stage revision surgery due to symptomatic nonunion or later-stage hardware removal due to prominence/bursitis
- 10 *‡* Included intraoperative incidental durotomy, intraoperative pedicle-screw-placement issues, new postoperative radiculopathy, early
- 11 postoperative pulmonary embolism, wound seroma/hematoma, deep wound infection or persistent lumbar radiculopathy
- 12 postoperatively

1 Table 3. Results of statistical analysis

Authors & Year	Statistical Analysis	Differences at Baseline	Postoperative Differences
			<u> </u>
Abbott et al.,	ANCOVA with	There were no significant differences	ANCOVA:
2013	adjustment for	between the cervical collar group and	NDI Scores at 1.5 months $(p = 0.042)^*$
	covariates,	the control group with respect to	Mean difference between groups: -4.4 (95% CI: -8.6 to
	repeated measure	gender, age, diagnosis, level of	-0.2); Cohen's effect size: -0.77
	analysis of	operation, cervical range of motion,	SF-36 BP scores
	covariance	unipedal balance and baseline NDI,	6 months $(p = 0.016)^*$
		SF-36 (subscales, PCS and MCS),	Mean difference between groups: 21.4 (95% CI: 4.4 to
		FES and Borg CR-10 (neck and arm pain) scores (p = NR)	38.5); Cohen's effect size: 0.73
			12 months $(p = 0.031)^*$
			Mean difference between groups: 17.5 (95% CI: 1.7 to
			33.2); Cohen's effect size: 0.57
			SF-36 SF scores at 12 months $(p = 0.049)^*$

Mean difference between groups: 16.5 (95% CI: 0.1 to
32.9); Cohen's effect size: 0.45
SF-36 PCS
$1.5 months (p = 0.025)^*$
Mean difference between groups: 5.8 (95% CI: 0.8 to
10.7); Cohen's effect size: 0.84
3 months (p = 0.038)*
Mean difference between groups: 6.8 (95% CI: 0.4 to
13.1); Cohen's effect size: 0.63
6 months (p = 0.017)*
Mean difference between groups: 7.4 (95% CI: 1.4 to
13.4); Cohen's effect size: 0.80
12 months $(p = 0.041)^*$
Mean difference between groups: 7.5 (95% CI: 0.3 to
14.6); Cohen's effect size: 0.66

There were no significant differences between the cervical collar group and the control group with respect to 1) neck pain, arm pain, FES, SF-36 PF, SF-36 RP, SF-36 GH, SF-36 VT, SF-36 RE, SF-36 MH, SF-36 MCS at all time points assessed (1.5, 3, 6, 12, and 24 months) (p > 0.05) 2) all components of the unipedal balance test (right/ foot, right/left foot soft surface, right/left foot eyes closed) at all time points assessed (1.5, 3, 6, 12, and 24 months) (p > 0.05) 3) all components of CROM (right/left lateral flexion, flexion, extension, right/left rotation) at all time points assessed (1.5, 3, 6, 12, and 24 months) (p > 0.05)

4) fusion rates and sagittal alignment (p = NR)

			Repeated measures analysis of covariance showed that controlling for the combined effects of all prospective
			measures gave significantly better outcome for the
			cervical collar group in neck pain ($p = 0.038$), SF-36
			PCS (p = 0.010) and SF-36 BP (p = 0.029).
Campbell et	ANOVA,	SF-36 PCS was higher in the control	Mean improvement in NDI Scores at 1.5 months ($p =$
al., 2008	ANCOVA, Fisher	group (31.1 ± 7.2) than in the braced	0.008)
	exact test, Student	group (33.4 \pm 7.8) (p = 0.019)	Braced: 21.6±18.4
	t test	6	Not Braced: 28.4±19.0
		There were no significant differences	
		between the braced group and the	There were no significant differences between the
		control group with respect to age (p =	braced group and the control group with respect to NDI
		0.367), gender (p = 0.447), worker's	scores at 3-months ($p = 0.468$), 6-months ($p = 0.169$),
		compensation ($p = 0.458$), litigation (p	12-months (p = 0.415) and 24-months (p = 0.693), SF-
		= 1.000), smoking status (p = 1.000)	36 PCS at 6-months (p = 0.481), 12-months (p = 0.260)

and occupational status ($p = 0.695$),	and 24-months ($p = 0.279$), average neck pain scores at
and baseline NDI ($p = 0.141$), neck	24-months ($p = 0.622$), and average arm pain scores at
pain (p = 0.523) and arm pain (p =	24-months ($p = 0.260$).
0.710) scores.	There were no significant differences in fusion success at any time period between groups, though higher rates of fusion were reported in the non-braced group ($p =$ 0.552 at 24-months).
cedeo .	There were no significant differences in rates of secondary surgeries or procedures between the braced group and the control group: revisions ($p = 0.653$),
PC0	removals (p = 0.724), reoperations (p = 1.000), supplemental fixations (p = 0.286), any surgery (p =
V	0.184).
<i>Mean curves</i> $(p = NR)$	<i>Postoperative mean curves</i> $(p = NR)$

Christodoulou NR

et al., 1987		Braced: 58.0°	Braced: 23.0°
		Not Braced: 54.0°	Not Braced: 22.8°
			There was no significant difference between the braced
			group and the control group with respect to the mean
			loss of correction at 24 months (7.0° for braced group,
			6.3° for control group, p = NR).
Hida et al.,	Two-way	There were no significant differences	There were no significant differences between the
2017	repeated	between the collar-fixation group and	collar-fixation group and the control group with respect
	ANOVA, Fisher	the control group with respect to age	to
	exact test, Student	(p = 0.73), gender (p = 0.26), height (p	1) VAS ($p = 0.487$), JOA recovery rates ($p = 0.80$),
	t test, x^2 test,	= 0.59), weight (p = 0.66), operation	SF-36 PCS (p = 0.537), SF-36 MCS (p = 0.504),
		time (p = 0.57), intraoperative blood	and SF-36 BP subscores ($p = 0.848$) at 12
		loss ($p = 0.69$), number of operated	months follow-up.
		levels (p = 0.67), VAS (p = 0.33),	2) the decrease in the C2-7 lordotic angle ($p =$
		JOA ($p = 0.67$), lordotic angle ($p =$	0.82) and ROM (p = 0.61).

0.84), ROM (p = 0.88) and SF-36 PCS	3) incidence of complications ($p = 0.53$).
(p = 0.68), MCS $(p = 0.80)$ and BP $(p = 0.68)$	
= 0.57).	

Yee et al.,	Mann-Whitney U	There were no significant differences	There were no significant differences between the brace
2008	test, chi-square	between the brace group and the	group and the control group with respect to
	test, Fisher exact	control group with respect to age (p =	1) the distribution of surgical complications or
	test, two-way	0.97), gender (p = 0.35), CCI (p =	subsequent revision rates $(p = 0.8)$
	ANOVA	0.6), number of levels included in the	2) postoperative DPQ category scores ($p = 0.34$ for
		arthrodesis ($p = 0.42$), smoking status	the daily activity category, $p = 0.67$ for the
		(p = 0.89), worker's compensation or	work/leisure category, $p = 0.17$ for the anxiety-
		litigation ($p = 0.48$), revision surgery	depression category and $p = 0.40$ for the social
		(p = 0.88), BMI (p = 0.74), diagnosis	category)
		(p = 0.8), and preoperative SF-36	3) postoperative SF-36 domain and component
		MCS (p = 0.9), SF-36 PCS (p = 0.19),	scores (p = 0.38 for PF, p = 0.28 for BP, p =
		SF-36 domain scores ($p > 0.05$) and	0.23 for GH, p = 0.41 for RP, p = 0.25 for VT, p
		DPQ category scores ($p > 0.05$)	= 0.79 for SF, p = 0.86 for RE, p = 0.30 for MH,

p = 0.30 for PCS and p = 0.57 for MCS)

4) rates of fusion seen radiographically at 12

months (p = 0.8) or 24 months (p = 0.9)

postoperatively.

5) Rates of revision surgery due to symptomatic

non-union (p = 0.43)

1 ANCOVA = analysis of covariance; ANOVA = analysis of variance; BMI = body mass index; BP = bodily pain; CCI = Charlson

- 2 comorbidity index; CI = confidence interval; CROM = cervical range of motion; DPQ = Dallas pain questionnaire; FES = falls
- 3 efficacy scale; GH = general health; JOA = Japanese Orthopedic Association; MCS = mental component score; MH = mental health;
- 4 NDI = neck disability index; NR = not reported; OR = odds ratio; PCS = physical component score; PF = physical functioning; RE =
- 5 role limitations emotional; ROM = range of motion; RP = role limitations physical; SF = social functioning; SF-36 = short form-36;
- 6 VAS = visual analog scale; VT = vitality
- 7 * No significant differences at all other time points assessed (1.5, 3, 6, 12, and 24 months postoperative)

1 Table 4. Summary and Strength of Evidence

		~	
		Strength	
Patient Population and	Studies	of	Overall Effect and Conclusions
Outcome of Interest		Evidence	
KQ1: What is the efficacy and effe	ctiveness of postoperative bracing of	compared with	n no bracing based on disability, pain, quality of life and
functional outcomes?			
Degenerative cervical			Results were inconsistent across studies: 1) a prospective
myelopathy or radiculopathy			cohort study reported that patients in the non-braced group
NDI	n = 2(Campbell, Carreon et al.	Insufficient	had better NDI scores at 6 weeks after surgery compared to
SF-36 PCS	2009, Abbott, Halvorsen et al.	Insufficient	patients in the braced group,(Campbell, Carreon et al. 2009)
SF-36 MCS	2013)	Low	whereas 2) a pilot randomized controlled trial indicated that
SF-36 BP subscale	n = 3(Campbell, Carreon et al.	Insufficient	patients in the braced group had better NDI scores at 6 weeks
SF-36 SF subscale	2009, Abbott, Halvorsen et al.	Insufficient	after surgery compared to patients in the non-braced
SF-36 other subscales	2013, Hida, Sakai et al. 2017)	Insufficient	group.(Abbott, Halvorsen et al. 2013)
VAS	n = 2(Abbott, Halvorsen et al.	Low	
JOA recovery rate	2013, Hida, Sakai et al. 2017)	Low	Results were inconsistent across studies: 1) a pilot
FES	n = 2(Abbott, Halvorsen et al.	Insufficient	randomized controlled trial indicated that patients in the non-

	Strength	
Studies	of	Overall Effect and Conclusions
	Evidence	
2013, Hida, Sakai et al. 2017)	Insufficient	braced group had better SF-36 PCS scores at 6 weeks, 3, 6,
n = 1(Abbott, Halvorsen et al.	Moderate	and 12 months after surgery compared to patients in the
2013)	Moderate	braced group,(Abbott, Halvorsen et al. 2013) whereas 2) two
n = 1(Abbott, Halvorsen et al.		studies (a prospective cohort study and a randomized
2013)	C C	controlled trial) reported no significant differences in SF-36
n = 1(Hida, Sakai et al. 2017)	NO.	PCS scores between the braced and non-braced groups at all
n = 1(Hida, Sakai et al. 2017)	6.	time points assessed.(Campbell, Carreon et al. 2009, Hida,
n = 1(Abbott, Halvorsen et al.		Sakai et al. 2017)
2013)		
n = 1(Abbott, Halvorsen et al.		Results were inconsistent across studies: 1) a pilot
2013)		randomized controlled trial indicated that patients in the
n = 2(Campbell, Carreon et al.		braced group had better SF-36 BP subscale scores at 6 and
2009, Abbott, Halvorsen et al.		12 months after surgery compared to patients in the non-
2013)		braced group,(Abbott, Halvorsen et al. 2013) whereas 2) a
n = 2(Campbell, Carreon et al.		randomized controlled trial reported no significant
	Studies 2013, Hida, Sakai et al. 2017) n = 1(Abbott, Halvorsen et al. 2013) n = 1(Abbott, Halvorsen et al. 2013) n = 1(Abbott, Halvorsen et al. 2013) n = 1(Hida, Sakai et al. 2017) n = 1(Hida, Sakai et al. 2017) n = 1(Hida, Sakai et al. 2017) n = 1(Abbott, Halvorsen et al. 2013) n = 1(Abbott, Halvorsen et al. 2013) n = 1(Abbott, Halvorsen et al. 2013) n = 2(Campbell, Carreon et al. 2013) n = 2(Campbell, Carreon et al. 2013) n = 2(Campbell, Carreon et al.	Studies Strength 2013, Hida, Sakai et al. 2017) Insufficient n = 1(Abbott, Halvorsen et al. Moderate 2013) Moderate n = 1(Abbott, Halvorsen et al. Moderate 2013) Moderate n = 1(Abbott, Halvorsen et al. Moderate 2013) n = 1(Hida, Sakai et al. 2017) n = 1(Hida, Sakai et al. 2017) Networken et al. 2013) n = 1(Abbott, Halvorsen et al. 2013) n = 2(Campbell, Carreon et al. 2013) In = 2(Campbell, Carreon et al.

		Strength	
Patient Population and	Studies	of	Overall Effect and Conclusions
Outcome of interest		Evidence	
	2009, Abbott, Halvorsen et al.		differences in SF-36 BP subscale scores between the braced
	2013)		and non-braced groups at all time points assessed.(Hida,
			Sakai et al. 2017)
		Ċ	Patients in the braced group had better SF-36 SF subscale
		0.	scores at 12 months after surgery compared to patients in the
		6.	non-braced group.(Abbott, Halvorsen et al. 2013)
	×°		There were no significant differences in SF-36 MCS and
			other subscales, VAS, JOA recovery rate, FES, unipedal
	G		balance standing test, neck pain, and arm pain between the
			braced and non-braced groups at all time points assessed.
Degenerative disease of the			
lumbar spine			
DPQ	n = 1(Yee, Yoo et al. 2008)	Low	There were no significant differences in DPQ, SF-36 PCS,

		Strength	
Patient Population and	Studies	of	Overall Effect and Conclusions
Outcome of Interest			
		Evidence	
SF-36 PCS	n = 1(Yee, Yoo et al. 2008)	Low	MCS and subscales between the braced and non-braced
SF-36 MCS	n = 1(Yee, Yoo et al. 2008)	Low	groups at all time points assessed.
SF-36 subscales	n = 1(Yee, Yoo et al. 2008)	Low	
KQ2: What is the impact of postope	erative bracing compared with no b	racing on radi	ographic outcomes?
Degenerative cervical			There were no significant differences in ROM, fusion rate,
myelopathy or radiculopathy			sagittal alignment, and decrease in lordotic angle C2-7
ROM	n = 2(Abbott, Halvorsen et al.	Low	between the braced and non-braced groups at all time points
Fusion rate	2013, Hida, Sakai et al. 2017)	Moderate	assessed.
Sagittal alignment	n = 2(Campbell, Carreon et al.	Low	
Lordotic angle C2-7	2009, Abbott, Halvorsen et al.	Low	
	2013)		
	n = 1(Abbott, Halvorsen et al.		
	2013)		
	n = 1(Hida, Sakai et al. 2017)		
Degenerative disease of the			There was no significant difference in fusion rate between

		Strength	
Patient Population and			
Outcome of Interest	Studies	of	Overall Effect and Conclusions
Outcome of Interest		Evidence	
lumbar spine			the braced and non-braced groups at all time points assessed.
Fusion rate	n = 1(Yee, Yoo et al. 2008)	Low	
Adolescent idiopathic scoliosis			There was no significant difference in mean loss of spinal
with thoracic curves $\geq 35^{\circ}$			curve correction between the braced and non-braced groups
Spinal curve	n = 1(Christodoulou, Prince et	Low	at all time points assessed.
	al. 1987)	NO.	
KQ3: What is the safety profile of	postoperative bracing compared wi	th no bracing?	
Degenerative cervical	.0	<i>y</i>	There were no significant differences in rate of revision
myelopathy or radiculopathy	Ô		surgery, second procedure or complications between the
Revision surgery or second	n = 1(Campbell, Carreon et al.	Low	braced and non-braced groups at all time points assessed.
procedure	2009)		
Complications		Moderate	
	n = 2 (Campbell, Carreon et al.		
	2009, Hida, Sakai et al. 2017)		
Degenerative disease of the			There were no significant differences in rate of revision

Detions Deputation and		1	
Outcome of Interest	Studies	of Evidence	Overall Effect and Conclusions
lumbar spine			surgery, second procedure or complications between the
Revision surgery or second	n = 1(Yee, Yoo et al. 2008)	Low	braced and non-braced groups at all time points assessed.
procedure			
Complications	n = 1(Yee, Yoo et al. 2008)	Low	S
KQ4: What is the cost-effective	ness of postoperative bracing?		
None	$\mathbf{n} = 0$	NA	None
CROM = cervical range of moti	on; DPQ = Dallas pain questionnair	re; FES = falls	efficacy scale; JOA = Japanese Orthopedic Association; MCS
mental component score; NDI =	neck disability index; NA = not ap	plicable; PCS :	= physical component score; ROM = range of motion; SF-36 =
short form-36; $VAS = visual and$	alog scale		



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