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A Randomized Trial Comparing 2 Techniques of Balloon Kyphoplasty and Curette Use for Obtaining Vertebral Body Height Restoration and Angular-Deformity Correction in Vertebral Compression Fractures due to Osteoporosis

L. Bastian, F. Schils, J.B. Tillman, and G. Fueredi, on behalf of the SCORE Investigators

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ABSTRACT

BACKGROUND AND PURPOSE: Vertebral compression fractures often result in pain and vertebral deformity. We compared 2 different balloon kyphoplasty techniques both using intraoperative curettage.

MATERIALS AND METHODS: Adults 50 years of age or older with osteoporosis and 1 acute VCF were randomized to undergo bilateral BKP in which the curette was used first (n = 57) followed by inflatable bone tamps or in which IBTs were used first, followed by curettage and a second IBT inflation (n = 55).

RESULTS: Mean procedure duration was 33.5 and 36.8 minutes, and fluoroscopy duration was 3.8 and 3.7 minutes for the CF and IBTF groups, respectively. Two-thirds of VCFs were wedge-shaped, and one-half had dynamic mobility. Anterior height restored postoperatively was 2.28 mm (95% CI, 1.49–3.08 mm; P < .001) and 2.78 mm (95% CI, 1.89–3.66 mm; P < .001) for CF and IBTF groups, representing ~35% and 39% of lost height restored, but group differences were not significant (P = .4). Intraoperative anterior height gain attributed to dynamic mobility was 2.96 mm (95% CI, 1.92–4.00 mm; P < .001) and 3.05 mm (95% CI, 2.10–4.00 mm; P < .001); additional height attributed to IBT inflation was 1.09 mm (95% CI, 0.77–1.41 mm; P < .001) and 1.25 mm (95% CI, 0.68–1.82 mm; P < .001), representing a 37% and 41% increase. There was no significant height loss on IBT removal and cementation. Both groups had improved pain and ambulation. Asymptomatic leakage occurred in 15% of VCFs. There was 1 nonserious device-related hematoma (IBTF group). One new clinical VCF occurred in each group, but they were not device-related.

CONCLUSIONS: Both techniques resulted in significant vertebral body height and pain improvement. Procedure and adverse event data demonstrated safe curette use in conjunction with balloon kyphoplasty procedures.

 $\label{eq:BBREVIATIONS: BKP = balloon kyphoplasty; CF = curette-first; CI = confidence interval; IBT = inflatable bone tamp; IBTF = IBT-first; IQR = interquartile range; VCF = vertebral compression fracture$

Vertebral compression fractures are associated with vertebral deformity and pain and a decrease in a patient's mobility, function, and quality of life.¹ Approximately 25% of women older than 50 years of age and 40% of women older than 80 years have a

VCF each year¹; approximately two-thirds of vertebral fractures do not come to clinical attention,² though the effects of the associated deformity are not inconsequential.¹

Balloon kyphoplasty is a minimally invasive treatment option designed to stabilize VCFs and correct angular deformity. During kyphoplasty, 2 inflatable bone tamps are placed into the vertebral body under fluoroscopic guidance. Inflation of the IBTs compacts the cancellous bone, creating a cavity within the vertebral body,

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and pushes the endplates apart, often restoring lost height and correcting angular deformity.³⁻⁶ After the balloons are removed, the void is filled with bone cement. In 2 randomized trials, BKP was shown to provide better pain, disability, and quality-of-life outcomes than nonsurgical management for patients with osteo-porosis and cancer.^{5,7,8}

The ability to restore vertebral anatomy is affected by many factors, including patient positioning, IBT inflation and position within the vertebral body, and the presence of sclerotic bone. A curette may be used as an adjunct tool to scrape and score cancellous bone either before or after the IBTs are inflated in an attempt to disrupt sclerotic bone that may hinder vertebral body height restoration.

Currently there is limited clinical evidence regarding the use of the curette in balloon kyphoplasty procedures. The overall objective of this study was to document the relative roles of postural reduction and systematic curette use, in combination with the IBT, in obtaining vertebral body anatomy height restoration during the balloon kyphoplasty procedure.

MATERIALS AND METHODS

Patients/Subjects

The SCORE trial was a randomized nonblinded trial comparing 2 surgical techniques for performing balloon kyphoplasty. Patients were randomized from February 2009 through August 2010. All participants had 1 acute (with edema on MR imaging), painful, thoracic or lumbar (T5-L5) vertebral fracture due to primary or secondary osteoporosis; the acute fracture also had to have caused at least 15% height loss. In the absence of cancer or trauma (see exclusions below), the presence of a VCF is a hallmark of osteoporosis9; therefore, the decision to perform diagnostic bone mineral attenuation examinations was determined by treating physicians and was not a study requirement. Patients were allowed to have chronic fractures as long as at least 1 adjacent segment was not fractured; chronic fractures were not treated. Patients were excluded from participating if they could not undergo MR imaging or standing lateral x-rays; had >1 acute fracture; had fractures due to cancer or high-energy trauma; had a body mass index >35 kg/m²; required procedures other than BKP for fracture stabilization; or had contraindications to BKP such as irreversible coagulopathy, known allergies to bone cement or contrast, or evidence of systemic infection or infection within the spine. Participants gave written informed consent before enrollment. The protocol and consent forms were approved by local ethics committees.

Because this study was descriptive in nature, a sample size of 120 patients was deemed adequate to obtain results for primary and secondary end points.

Research Design

Patients were randomly assigned in a 1:1 ratio to have kyphoplasty by using the IBT-first followed by curette usage and a second IBT inflation (IBT-first) or a kyphoplasty procedure where the curette was used first followed by IBT inflation (curette-first). We investigated these 2 approaches within the study design because they are standard methods used clinically. Study randomization took place in the operating room and was stratified by the presence or absence of dynamic fracture mobility when the patient was placed prone on the operating table; this assessment was made by treating physicians by using intraoperative fluoroscopy. The study was performed at 11 investigational centers in 3 countries (Belgium, Germany, and the United States).

Balloon kyphoplasty was performed by using introducer tools, inflatable bone tamps, curettes and polymethylmethacrylate bone cement, and delivery devices (all devices were manufactured by Medtronic Spine, Sunnyvale, California), by using a percutaneous, bilateral, transpedicular, or extrapedicular approach.¹⁰ The curette tip can be ratcheted 0°, 30°, 60°, and 90° and locked in place for each setting. In the curette-first arm, after vertebral body access and appropriate channel creation with standard bone access systems, the curette was inserted first on 1 side to scrape the cancellous bone to establish a specific and directed path of least resistance for balloon insertion. The 30° position was used first with scraping from anterior to posterior throughout the vertebral body; articulations at 60°-90° were used when possible subsequently; similar use of the curette occurred on the contralateral side. In the IBT-first group, balloons were inserted into the working channel first; on inflation, the operator may visualize irregular or undesired balloon inflation profiles or limited balloon inflation volume due to hard bone. In these cases, the curette was appropriately oriented to scrape cancellous bone in the areas hindering balloon inflation. In all cases of curettage in both treatment arms, care was taken so as not to breach the cortical bone. Most procedures were performed with the patient under general anesthesia; some patients had conscious or deep sedation with local anesthesia.

The primary end point of the study was the difference in the percentage of vertebral body height restored (anterior, medial, and posterior vertebral body measurements¹¹) in each study arm as measured at discharge by using standing lateral radiographs. Secondary radiographic end points included vertebral body height restored in millimeters; kyphosis angle, defined as the angle formed by lines drawn parallel to the caudal and cranial fractured vertebral body endplates; and the local Cobb angle, defined as the angle formed by lines drawn parallel to the superior endplate of the vertebral body above and the inferior endplate of the vertebral body below.6 Clinical secondary end points included postoperative back pain relief by using a numeric rating scale that assessed current pain (scale 0-10),¹² ambulatory status (subjective patient questionnaire), and all adverse events. Radiographic outcomes were assessed at baseline/screening and intra- and postoperatively. Back pain and ambulation were assessed at baseline/screening and postoperatively (discharge), while adverse events were collected throughout the study through 30 days postoperatively.

Adverse events and serious adverse events were coded according to the Medical Dictionary for Regulatory Activities (http:// www.meddramsso.com).¹³ All adverse events and serious adverse events within 30 days were reported, as randomized. An adverse event was considered serious if it resulted in death, life-threatening injury, or permanent impairment; required intervention to prevent impairment; or required prolonged hospitalization. Site investigators reviewed and signed all case report forms, which were source-verified for 100% of the patients.

Radiographic imaging guidelines were established before study start for consistency; for pre- and postoperative images,

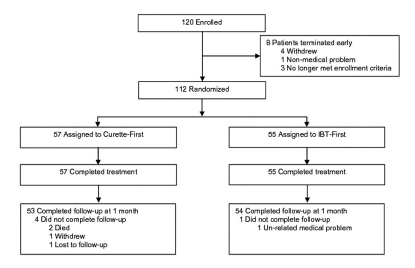


FIG 1. Patient accountability.

Table 1: Baseline p	atient demo	graphics
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Characteristic	CF	IBTF
No. of patients evaluable ^a	57	55
Age (yr) (mean)	73.6 ± 8.4	76.0 ± 9.4
Weight in kg (mean)	71.0 ± 16.2	71.8 ± 16.3
Height in cm (mean)	162.1 ± 8.3	162.2 ± 8.8
Female (No.) (%)	47 (82.5)	41 (74.5)
Ethnicity (No.) (%)		
Caucasian	55 (96.5)	55 (100.0)
Hispanic	1 (1.8)	0 (0)
Middle Eastern	1 (1.8)	0 (0)
Smoking history (No.) (%)		
Never	36 (63.2)	36 (65.5)
Former	14 (24.6)	14 (25.5)
Current	7 (12.3)	5 (9.1)
Index fracture age (wk) (median) (IQR)	4.0 (2.0, 9.0)	5.0 (3.0, 8.0)
Index vertebral body location (No.) (%)		
Т5-Т9	7 (12.3)	6 (10.9)
T10-L2	42 (73.7)	37 (67.3)
L3-L5	8 (14.0)	12 (21.8)
Index fracture morphometry (No.) (%) ^b		
Wedge	38 (66.7)	34 (61.8)
Crush	4 (7.0)	4 (7.3)
Concave	15 (26.3)	17 (30.9)
Prior symptomatic VCF (No.) (%) ^b		
0	44 (77.2)	43 (78.2)
1	5 (8.8)	11 (20.0)
2	2 (3.5)	1 (1.8)
3	6 (10.5)	0 (0)

^a A total of 120 patients enrolled. Eight patients withdrew early, did not receive surgery, and, therefore, did not receive randomization assignment.

^b As determined by treating physicians.

patients were in a standing position with legs straight and the shoulder against the Bucky with shoulders and hips perpendicular to it. Arms were either at a right angle to the body holding a support bar or across the chest if no bar was available. The beam and Bucky were centered at the index level to minimize parallax. A specific calibration marker, attached to a lanyard, was draped over the shoulder and positioned near the treated level. Intraoperatively, patients were placed prone with bolsters under the hips and shoulders (some patients had images obtained pre- and postbolstering) and imaged to assess postural reduction. Images were also taken after each IBT inflation step and cement placement. All spine radiographs and images were assessed centrally (Medical Metrics, Houston, Texas); an independent radiologist, blinded to treatment assignment, used quantitative morphometry¹¹ to determine anterior, mid-, and posterior vertebral body locations by using a computer-based marking system. Vertebral body height restoration and kyphotic angulation measurements were based on the quantitative morphometry.

Statistical Methods

Results were analyzed by modified intention to treat, including all data available at baseline and follow-up from all (112) patients randomized and treated. Eight additional patients were enrolled in the study but were terminated early before the procedure; because randomization was conducted in the operating room due to stratification based on dynamic fracture mobility, these 8 pa-

tients were not randomized to treatment. We used analysis of variance to analyze the continuous primary and secondary radiographic and pain end points between treatment groups; for categoric variables, between-group comparisons were assessed by using χ^2 , and, when appropriate, the Fisher exact test. A *P* value of .05 was considered statistically significant.

Role of the Funding Source

This study was sponsored and funded by Medtronic Spine LLC, which contributed to the study design, data monitoring, statistical analysis, and reporting of results and paid for core laboratory services (Medical Metrics, Houston, Texas). All authors had complete access to data and were provided all analyses requested.

RESULTS

Patient Disposition and Demographics

Patients were enrolled between February 2009 through August 2010 and were randomized to curette-first (n = 57) or IBT-first (n = 55) groups. Patient disposition is shown in Fig 1. Eight patients were discontinued before the planned procedure and were not randomized to treatment. Of the 112 patients who were randomized, 5 patients discontinued before 1 month.

Patients were, on average, 75 years of age, and 79% were women (Table 1). In the overall cohort, 78% of patients did not have any prior VCF. At baseline, wedge fractures were the most common type of VCF identified by investigators (64%). Median fracture age was 4.0 and 5.0 weeks in the CF and IBTF groups, respectively. In the overall cohort, 73 of 112 (65%) patients randomized had bone mineral attenuation examinations; in this subset of patients, the average *t*-score was -1.9 ± 1.0 .

Procedure and Fracture Characteristics

Randomization was stratified by fracture mobility and was conducted in the operating room; one-half of treated fractures had dynamic mobility observed by the treating physician when patients were placed prone with bolsters on the operating table (Table 2). Most patients were treated by using general anesthesia (83%) and a bipedicular transpedicular approach (94%). The

Tabl	e 2:	Proced	lure c	haract	eristics
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Procedure ^a	CF	IBTF
No. of patients treated	57	55
Patients with mobile fracture (No.) (%)	29 (50.9)	28 (50.9)
Procedure duration (min) (mean)	33.5 ± 10.8	36.8 ± 12.5
Fluoroscopy duration (min) (mean)	3.8 ± 2.0	3.7 ± 3.3
Anesthesia		
General	49 (86.0)	44 (80.0)
Local	8 (14.0)	11 (20.0)
Approach		
Extrapedicular	3 (5.3)	3 (5.5)
Transpedicular	54 (94.7)	51 (92.7)
Initial IBT inflation volume (mL)		
Left	2.76 ± 1.21	2.52 ± 1.38
Right	2.86 ± 1.23	2.52 ± 1.45
Initial IBT inflation pressure (PSI)		
Left	167.3 ± 73.0	168.5 ± 79.5
Right	169.9 ± 84.8	159.5 ± 75.9
Final IBT inflation volume (mL)		
Left	2.79 ± 1.29	3.00 ± 1.48
Right	2.85 ± 1.35	3.03 ± 1.51
Final IBT inflation pressure (PSI)		
Left	164.1 ± 87.3	151.8 ± 86.2
Right	157.6 ± 89.0	138.6 ± 77.5
Final cement volume (mL)		
Left	2.78 ± 1.20	2.94 ± 1.16
Right	2.89 ± 1.25	
Length of stay (hours)	96.1 ± 153.9	77.5 ± 108.2

^a Thirteen patients in the curette-first group had an additional curettage and IBT inflation; 7 patients in the IBT-first group had a third IBT inflation.

mean kyphoplasty procedure duration was 33.5 and 36.8 minutes, with mean fluoroscopy durations of 3.8 and 3.7 minutes for the CF and IBTF groups, respectively. On average, final bilateral IBT inflation volumes for the CF and IBTF groups were 5.6 \pm 2.6 and 6.0 mL \pm 2.8 mL, which were consistent with cement volumes of 5.7 \pm 2.3 and 5.9 \pm 2.2 mL. For the CF and IBTF groups, hospitalization duration was an average of 96.1 and 77.5 hours, respectively.

As assessed by the radiographic core laboratory, most fractures were wedge-shaped and were considered moderate or severe. Nineteen percent had a vacuum cleft present (Table 3).

Primary End Point

The primary end point of the study was height restoration assessed postoperatively by using standing lateral films. The mean baseline anterior height was 60.2% (95% CI, 54.6%–65.8%) and 58.6% (95% CI, 53.8%–63.4%) of adjacent normal vertebrae for the CF and IBTF groups, which was improved to 70.5% (95% CI, 66.2%–74.8%; P < .001) and 71.3% (95% CI, 67.2%–75.3%; P < .001), respectively (Fig 2). The improvement observed was not statistically significantly different between the 2 treatment groups (P = .4). Similar results were found at the midvertebral measurements (Fig 2).

Secondary Radiographic End Points

At baseline, compared with normal adjacent vertebrae, the average anterior height loss of index fractures measured was -6.49 mm (95% CI, -7.51 to -5.47 mm) and -7.13 mm (95% CI, -8.21 to -6.06 mm) for the CF and IBTF groups, respectively. Compared with the preoperative standing assessment, the intraoperative anterior height gain, attributed to unloading the spine

Table 3: Fracture characteristics^a

able 5. Fracture characteristics					
Preoperative Fracture Evaluation	CF	IBTF			
No. of patients treated	57	55			
No. of fractures evaluable	56	54			
Fracture type					
Wedge	47 (83.9)	44 (81.5)			
Biconcave	9 (16.1)	10 (18.5)			
Crush	0 (0)	0 (0)			
Severity of fracture					
Slight	13 (23.2)	10 (18.5)			
Moderate	21 (37.5)	23 (42.6)			
Severe	22 (39.3)	21 (38.9)			
Vacuum cleft					
Absent	43 (76.8)	46 (85.2)			
Present	13 (23.2)	8 (14.8)			
Intraoperative evaluation					
No. of fractures evaluable	55	51			
Cement leakage absent	46 (83.6)	44 (86.3)			
Cement leakage present	9 (16.4)	7 (13.7)			
Disk	8	6			
Epidural/foraminal	1	0			
Paraspinal	2	1			
Vascular	2	1			
>15 mm	1	0			

^a Parameters assessed by radiographic core laboratory.

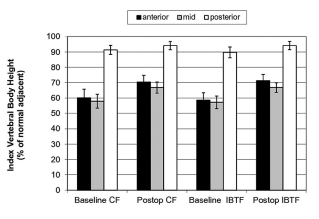


FIG 2. Primary end point of absolute height restored at index vertebrae as a percentage of adjacent normal vertebrae. Group means and 95% confidence intervals are shown for measurements at the posterior, anterior, and midpoint of index vertebral bodies for each group at baseline and at 48 hours postoperatively by using standing lateral x-ray films.

and providing postural reduction, was 2.96 mm (95% CI, 1.92-4.00 mm; P < .001) and 3.05 mm (95% CI, 2.10–4.00 mm; P <.001) for the CF and IBTF groups, respectively (Fig 3); however, the improvement observed between groups was not statistically significantly different (P = .9). The additional benefit in intraoperative height gain attributed to the first IBT inflation was 1.09 mm (95% CI, 0.77–1.41 mm; P < .001) in the CF group and 1.25 mm (95% CI, 0.68–1.82 mm; P < .001) in the IBTF group, representing a 37% and 41% increase in the anterior measurement over postural reduction; there was no significant height loss on IBT removal and cement placement (Fig 3). The final preoperative-to-postoperative anterior height improvement was 2.28 mm (95% CI, 1.49–3.08 mm; *P* < .001) and 2.78 mm (95% CI, 1.89– 3.66 mm; P < .001) for the CF and IBTF groups (Fig 3), representing ~35% and 39% of lost height restored. Differences were not statistically different between groups in the anterior improve-

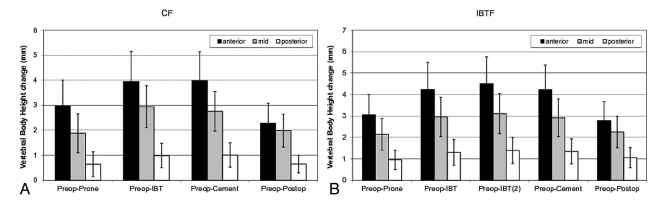


FIG 3. Index vertebral body height gain in millimeters. CF (A) and IBTF (B) group means and 95% confidence intervals are shown for vertebral body height improvement (compared with preoperative) in millimeters, measured at the posterior, anterior, and midpoint of index vertebral bodies after prone positioning (postural reduction), IBT inflation, cementation, and postoperative steps. Preoperative and postoperative assessments used standing lateral x-rays, and intraoperative assessments used lateral fluoroscopic images.

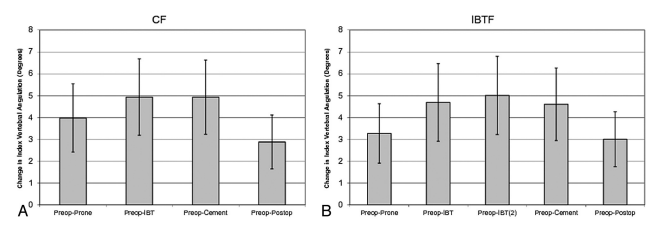


FIG 4. Index vertebral body kyphosis angle. CF (A) and IBTF (B) group means and 95% confidence intervals are shown for kyphotic angulation improvement (compared with preoperative) of index vertebral bodies measured as the angle between the inferior and superior endplates of index vertebrae after prone positioning (postural reduction), IBT inflation, cementation, and postoperative steps. Preoperative and postoperative assessments used standing lateral x-rays, and intraoperative assessments used lateral fluoroscopic images.

ment (P = .4); similar results were found in the midvertebral measurements (Fig 3).

We also measured the vertebral body kyphosis angle for treated fractures (Fig 4). Compared with preoperative measurements, intraoperative postural reduction resulted, on average, in 3.98° (95% CI, 2.42°–5.54°; P < .001) and 3.27°(95% CI, 1.90°– 4.63°; P < .001) of correction for the CF and IBTF groups (Fig 4); the additional benefit in amount of angulation attributed to the first IBT inflation was an additional 1.10° (95% CI, 0.51°-1.68°; P < .001) and 1.51° (95% CI, 0.70°–2.33°; P < .001) of correction, respectively. The IBT contribution represented a 27% and 46% increase from the postural reduction measurements for CF and IBTF groups, but the results were not statistically different between groups (P = .4). The final pre- to postoperative kyphosis correction was 2.88° (95% CI, 1.64°-4.11°; P < .001) and 3.00° (95% CI, 1.74°-4.25°; P < .001) for the CF and IBTF groups, but differences were not statistically different between groups (P = .9).

The local Cobb angle was measured for treated vertebrae. The postoperative correction of the local Cobb angle was 1.36° (95% CI, 0.33°–2.38°; P = .010) and 1.87°(95% CI, 0.81°–2.92°; P < .001) for the CF and IBTF groups; the groups were not statistically significantly different (P = .5).

Mobile Fractures

In evaluating the intraoperative contribution of postural reduction, IBT/curette use, and overall height restoration, improvement was better in each category for mobile fractures. Average anterior height gain attributed to postural reduction was 4.39 mm (95% CI, 3.38-5.39 mm) for mobile fractures versus 1.40 mm (95% CI, 0.68–2.11 mm) for nonmobile ones (P < .001 for difference), and the additional gain attributed to IBT inflation was 1.57 mm (95% CI, 1.08-2.06 mm) versus 0.91 mm (95% CI, 0.50-1.31 mm), respectively; the difference between mobile and nonmobile groups for IBT contribution was also statistically significant (P = .043) and represents an approximate 35.8% and 65.0% increase over postural reduction, respectively. The final pre-/postoperative height change was better in the mobile fracture group (3.46 mm; 95% CI, 2.58–4.34 mm; P < .001) compared with nonmobile fractures (1.52 mm; 95% CI, 0.84-2.21 mm; P < .001); this difference was statistically significant between the groups (P < .001).

Figure 5 is an illustrative case example depicting angular-deformity correction for the treated fracture at all stages of the kyphoplasty procedure. The case example is from a 72-year-old female patient with a mobile L1 fracture. Placing the patient prone on the operating table resulted in an 8.9° correction without bol-

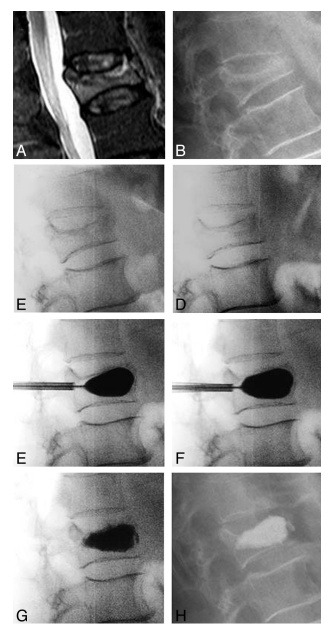


FIG 5. Case illustration of kyphosis correction for each step in the kyphoplasty procedure. *A*, Preoperative T2 short- τ inversion recovery MR imaging. *B*, Preoperative standing x-ray. *C*, Intraoperative postural reduction without bolsters. *D*, Intraoperative postural reduction with a bolster. *E*, Intraoperative first balloon inflation. *F*, Intraoperative curette and second balloon inflation. *G*, Intraoperative cement placement. *H*, Postoperative standing x-ray.

sters and an additional degree of correction after placing chest and hip bolsters. The patient was randomized to the IBTF group, and the first balloon inflation was uniform, providing an additional 7° of angular-deformity correction; the IBT inflation pressures were 120 PSI (bilateral) and were 4.0 (right) and 4.2 (left) mL in volume. After curette use and a second balloon inflation, there was a better inflation pattern and volume, with an additional degree of angular-deformity correction; for the second inflation, pressures were 106 (right) and 120 (left) PSI with an inflation volume of 5.0 mL for each side. The overall postoperative kyphosis correction of the fracture was 12.4°, and the local Cobb angle correction was 9.1°. In the overall cohort, 57 of 112 patients (51%) had fractures

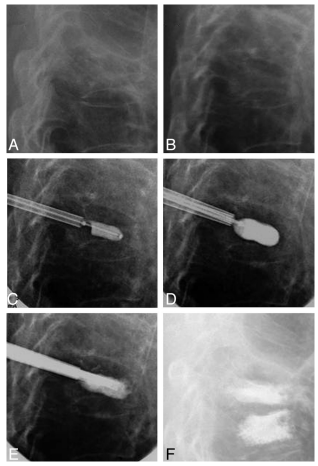


FIG 6. Case illustration of a nonmobile fracture. *A*, Preoperative standing x-ray. *B*, Intraoperative postural reduction with a bolster. *C*, Intraoperative first balloon inflation. *D*, Intraoperative curette and second balloon inflation. *E*, Intraoperative cement placement. *F*, Postoperative standing x-ray.

with dynamic mobility (Table 2); we performed analyses comparing mobile versus nonmobile fractures. The median age for mobile fractures was statistically significantly lower (3.0 weeks; IQR, 2.0-6.0 weeks) than nonmobile fractures (6.0 weeks; IQR, 3.0-10.0 weeks; P = .042).

Figure 6 is an illustrative case in which there was no postural reduction at T7; the 73-year-old female patient was randomized to the IBTF group. The first balloon inflation was poor, had an irregular inflation pattern (119 [right] and 75 [left] PSI and 2.0 [right] and 1.0 [left] mL volume), and provided no angular-deformity correction. After curette use and a second balloon inflation, the result was a more uniform fill (119 [right] and 96 [left] PSI and 2.0 mL, bilateral) with deformity correction. There was 2.4° of correction overall at T7 in the standing postoperative measurement (the inferior T8 was treated within 30 days of initial treatment).

Figure 7 is an illustrative case in curette and balloon use; the 71year-old female patient had a nonmobile L1 fracture and was randomized to the IBTF group. The initial balloon inflation was poor and had an irregular inflation pattern (1.0 mL and 230 PSI, bilateral). After bilateral curette use and a second balloon inflation, the result was a more uniform fill (230 PSI, bilateral; and 2.5 [right] and 3.0 mL

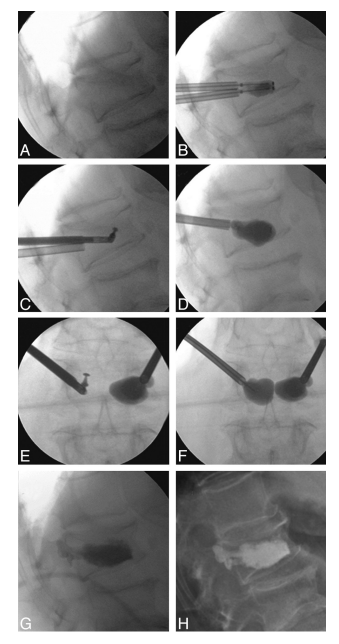


FIG 7. Case illustration of nonmobile fracture and use of curette. *A*, Intraoperative postural reduction with a bolster. *B*, Intraoperative first balloon inflation. *C*, Intraoperative curette usage (used bilaterally but image shows use only on 1 side). *D*, Intraoperative second balloon inflation. *E*, Anteroposterior (AP) film of *C*. *F*, AP film of *D*. *G*, Intraoperative cement placement. *H*, Postoperative standing x-ray.

[left]); however, the degree of vertebral body angulation correction was minimal overall (0.2°).

Anesthesia Type

We analyzed some parameters by general and local anesthesia groups. Fewer patients were treated with local anesthesia (Table 2); within the local anesthesia cohort, hospital duration was shorter (24.0 hours; IQR, 20.0–29.0 hours) compared with the general anesthesia group (72.0 hours; IQR, 24.0–96.0 hours; P < .001). More patients with local anesthesia (16/19, 84%) had nonmobile fractures compared with those with general anesthesia (39/93, 42%; P < .001) and had less fracture reduction in general (data not shown).

Pain and Mobility

Assessed by the 10-point numeric rating scale, the CF kyphoplasty group had a baseline back pain score of 7.7 points (95% CI, 7.4–8.1 points), which was similar to that in the IBTF group (7.5 points; 95% CI, 7.1–7.9 points). Both groups had statistically and clinically significant postoperative back pain relief (Fig 8). On average, the CF group had postoperative improvement to 3.0 points (95% CI, 2.5–3.5 points; P < .001) and the IBTF group improved to 3.3 points (95% CI, 2.8–3.8 points; P < .001). This represented 61% and 56% improvement for the CF and IBTF groups, which was not statistically significantly different (P = .14); patients with mobile fractures had similar pain relief compared with patients with nonmobile fractures (data not shown). Concomitant with the pain reduction observed, more patients were able to freely ambulate postoperatively in both kyphoplasty groups (Fig 8).

Safety

No subject experienced an adverse event during surgery. Asymptomatic cement leakage occurred in 16% and 14% of vertebral bodies treated in the CF and IBTF groups (Table 3). The most common adverse events within 30 days were back pain (1 CF, 4 IBTF) and headache (2 CF, 1 IBTF). There was 1 nonserious device-related hematoma that occurred in the IBTF group (On-line Table 1). There were 5 nonserious adverse events that were possibly related to anesthesia (On-line Table 1), which included 2 cases of postoperative confusion and 1 case each of vomiting and arthralgia in the CF group and 1 case of headache in the IBTF group.

New vertebral fracture serious adverse events occurred in 2 patients, 1 in each group (On-line Table 2), both of which resulted in a subsequent surgical procedure. Three patients had serious adverse events that resulted in death (On-line Table 2). One patient with a lung infection did not respond to antibiotic therapy and died 35 days postoperatively. Another patient had pneumonia 4 days after surgery, which resulted in death; this patient had predisposing risk factors of cardiac insufficiency and previous cardiac valvuloplasty. A third patient died 6 days after surgery from septic shock; this patient presented with abdominal pain 4 days postkyphoplasty and was hospitalized with total colectomy and ileostomy. None of the serious adverse events, including those that resulted in death, were related to the device or procedure (On-line Table 2).

DISCUSSION

This study is the first that systematically used a curette in balloon kyphoplasty procedures documenting multiple outcomes. The results of this randomized trial comparing 2 different kyphoplasty techniques demonstrate statistically significant vertebral body height restoration and kyphosis correction in both treatment arms (CF and IBTF); the results also demonstrate that fracture mobility, fracture age, patient positioning, and IBT combined with curette use are important parameters that influence fracture reduction. Anesthesia type may be a factor influencing fracture mobility and vertebral body height restoration, but interpretation is limited by the small cohort and nonrandomized nature of this comparison. Both randomized treatment groups showed similar

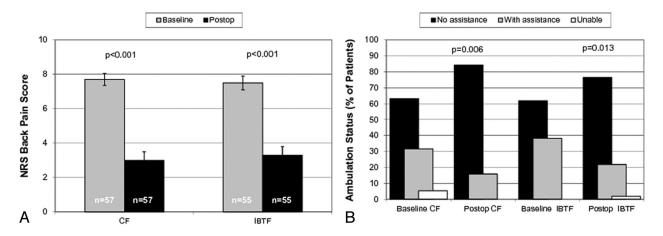


FIG 8. Postoperative back pain and ambulation. *A*, Group means and 95% confidence intervals are shown for the back pain numeric rating scale (0–10) measured at baseline and discharge. Paired *t* test *P* values for comparison with baseline are shown for each group. *B*, Percentage of patients in each ambulation category is shown for baseline and at discharge for each group. Stuart-Maxwell *P* values comparing postoperative status with baseline are shown for each kyphoplasty group.

vertebral body height restoration, pain and mobility, and safety outcomes. By qualitatively comparing the adverse event profile, cement extravasation rate, and the procedure and fluoroscopy duration that were observed in this trial with others,^{4,5,7,8,14,15} we found that the routine use of the curette should be considered safe.

Mounting evidence demonstrates that kyphoplasty and vertebroplasty provide better outcomes to nonsurgical management in randomized controlled studies.7,8,16-18 Although 2 randomized trials concluded that vertebroplasty was no different from a "sham" intervention for back pain, disability, and quality of life outcomes,^{19,20} these trials have been criticized due to several important limitations that make it hard to draw definitive conclusions.^{21,22} Kyphoplasty has not yet been compared with a sham intervention; this comparison would be a reasonable and definitive next step. However, kyphoplasty had better pain and EuroQol 5 dimensions quality-of-life outcomes compared with nonsurgical management throughout 2 years,⁷ an outcome not consistent with placebo effects. Currently, there is 1 small randomized trial that compared kyphoplasty with vertebroplasty; pain outcomes were similar but kyphoplasty provided statistically significantly better height-restoration outcomes.²³ A few studies have demonstrated a possible connection between angulation and pulmonary function,^{24,25} but no definitive study has demonstrated a clinical benefit to height restoration.

Dynamic mobility of vertebral fractures has been investigated and described in the published literature.^{26,27} McKiernan et al²⁷ in a prospective case series of patients with vertebroplasty estimated that 35% of fractures had dynamic mobility, and these authors reported improvement of kyphosis in the subset of patients with mobile fractures. In our series, 50% of patients (and fractures treated) had mobile fractures; overall kyphosis correction improved from -15.2° to -10.7° , an approximate 30% improvement for mobile fractures treated in this study with BKP. Improvement in nonmobile fracture was approximately half as much at 14% improvement. In our series, mobile fractures, not surprisingly, were younger fractures.

Postural reduction and additional IBT contribution have also been documented in several studies.^{6,28} In our study, for both treatment groups, the intraoperative contribution of postural reduction was statistically significant and substantial (Figs 3 and 4); the intraoperative IBT contribution was also statistically significant and represented a 27%-46% increase over postural reduction. This additional correction of angulation (and vertebral body height) by the IBT/curette was maintained after IBT removal and subsequent cement placement. It is difficult to translate intraoperative contributions to the final results because the spine is unloaded intraoperatively but loaded in baseline and postoperative assessments. Overall, the final kyphosis correction/vertebral body height restoration in our study was lesser to some extent compared with other results such as those by Voggenreiter⁶; this finding may be due to the multicenter nature of our study, larger sample size, and the independent radiographic core lab assessments, which minimize reader bias. It will be of interest in future studies to understand deformity-correction outcomes with new inflatable bone tamp technologies that are made of less compliant materials.

Kyphoplasty with curette usage was found to be safe for the treatment of VCFs in patients with osteoporosis. Both treatment groups had a similar safety profile; as shown in On-line Tables 1 and 2, medical adverse events were similar between the 2 groups during the 1-month period, as randomized. There was 1 nonserious device-related hematoma that occurred in the IBTF group. Several other nonserious adverse events were attributed to the effects of anesthesia. No serious adverse event was device-related. Asymptomatic cement extravasation occurred in 15% of treated vertebral bodies overall. The safety profile and cement leakage are similar to those in several other studies in which the curette was not routinely used, suggesting that routine use of the curette does not alter the safety profile of kyphoplasty.^{4,7,8}

A limitation of this study is that it was not blinded; though ours was primarily a radiographic study, the core lab was blinded to treatment assignment. Although some may consider the short follow-up a limitation, the study was purposefully designed as such to focus on intraoperative assessments for determining height restoration. A limitation may be perceived by the difficulty in standardizing the exact use of the curette due to variations in anatomy and the bone environment; on the contrary, most importantly, the variability of use adds to the generalizability of the study because it represents the reality of curette use in clinical practice. Furthermore, the contribution of the curette itself is difficult to ascertain because both treatment arms used the curette. Nonetheless, the strengths are the randomized multicenter trial design and the multiple independent radiographic assessments that included several measurements within the intraoperative time frame.

Both treatment groups (CF and IBTF) showed statistically significant improvements in anterior and mid-vertebral body height, kyphosis correction, pain improvement, and ambulation status, consistent with several other kyphoplasty studies.⁴⁻⁷ However, we were not able to demonstrate statistically significant differences between treatments groups, though there may have been a trend for better radiographic results in the IBTF group. It is also difficult to ascertain the contribution of the curette itself because both treatment groups used the curette; however, when we compared the 2 groups at the initial balloon inflation step (preoperative IBT in Figs 3 and 4), there was little difference between groups. This finding provides some information because in the CF group, the curette was used just before IBT inflation and in the IBTF group, the curette was not yet used in the first IBT step. Therefore, in general, we do not think it is necessary to use the curette in every kyphoplasty case. Physicians may expect increased fluoroscopy time or procedure duration with additional tool use; however, we have demonstrated that curette use can be performed safely and, in comparison with other BKP studies, does not seem to increase overall operation time or fluoroscopy time.^{14,15} We have demonstrated, in several individual cases, that after initial IBT placement, the use of the curette can help better position the IBT and provide better inflation patterns on the second or third IBT inflations; this outcome is likely in sclerotic or harder bone environments and older fractures. In our experience, preoperative films with increased attenuation in the fractured vertebral body or resistance in introducing the needle may be indicative of hard/sclerotic bone; in these cases, the use of the curette may be warranted before balloon inflation to help direct the balloon tamps. However, most often the initial balloon inflation is most telling. Small balloon volumes at high pressures or irregular balloon shapes indicate the presence of hard/sclerotic bone; therefore, decisions regarding curette use are made intraoperatively. We, therefore, recommend the use of the IBT-first approach and then determining the need for curette use in kyphoplasty procedures. Anatomy correction is an important consideration in fracture management, and the treating physician should consider all parameters to achieve this end point in vertebral augmentation procedures.

CONCLUSIONS

Two different techniques with BKP and curette use resulted in significant vertebral body height restoration, kyphotic deformity correction, and pain improvements. Procedure and adverse event data demonstrated safe curette use in conjunction with balloon

kyphoplasty procedures. Fracture mobility, fracture age, patient positioning (bolstering), IBT and curette use, and possibly anesthesia type are important parameters that may influence fracture reduction. Treating physicians should consider all of these parameters in achieving maximal anatomy correction when performing vertebral augmentation procedures.

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APPENDIX

SCORE Investigators

- Belgium (35 enrolled patients)
 - F. Schils, Liège
 - K. De Smedt, Wilrijk
- Germany (50 enrolled patients)
 - G. Voggenreiter, Eichstätt
 - M. Markmiller, Kempten
 - L. Bastian, Leverkusen
 - H.C. Harzmann, München
 - B. Böhm, Nürnberg
 - M. Bierschneider, Regensburg
 - U. Laupichler, Schwandorf
- United States (35 enrolled patients)
- G. Fueredi, Burlington, Wisconsin J. Small, Temple Terrace, Florida

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