Prophylactic vertebroplasty procedure applied with a resorbable bone cement can decrease the fracture risk of sandwich vertebrae: long-term evaluation of clinical outcomes

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Abstract
A sandwich vertebra is formed after multiple osteoporotic vertebral fractures treated by percutaneous vertebroplasty, which has a risk of developing new fractures. The purpose of our study was to (i) investigate the occurrence of new fractures in sandwich vertebra after cement augmentation procedures and to (ii) evaluate the clinical outcomes after prophylactic vertebral reinforcement applied with resorbable bone cement.

From June 2011 to 2014, we analysed 55 patients with at least one sandwich vertebrae and treated with percutaneous vertebroplasty. Eighteen patients were treated by prophylactic vertebroplasty with a resorbable bone cement to strengthen the sandwich vertebrae as the prevention group. The others were the non-prevention group. All patients were examined by spinal radiographs within 1 day, 6 months, 12 months, 24 months and thereafter.

The incidence of sandwich vertebra is 8.25% (55/667) in our study. Most sandwich vertebrae (69.01%, 49/71) are distributed in the thoracic–lumbar junction. There are 24 sandwich vertebrae (18 patients) and 47 sandwich vertebrae (37 patients) in either prevention group or non-prevention group, respectively. No significant difference is found between age, sex, body mass index, bone mineral density, cement disk leakage, sandwich vertebrae distribution or Cobb angle in the two groups. In the follow-up, 8 out of 37 (21.6%) patients (with eight sandwich vertebrae) developed new fractures in non-prevention group, whereas no new fractures were detected in the prevention group. Neither Cobb angle nor vertebral compression rate showed significant change in the prevention group during the follow-up. However, in the non-prevention group, we found that Cobb angle increased and vertebral height lost significantly ($P<0.05$).

Prophylactic vertebroplasty procedure applied with resorbable bone cement could decrease the rate of new fractures of sandwich vertebrae.

Keywords: vertebroplasty; resorbable bone cement; sandwich vertebrae; spinal fracture

Introduction
Osteoporotic vertebral compression fractures (OVCFs) are a common complication of osteoporosis, which are also a major cause of morbidity and health costs among patients [1–3]. Percutaneous vertebroplasty (PVP) is a radiological procedure, which consists of percutaneous puncture and injection of polymethylmethacrylate
Committee of our hospital. prior to the study. All the investigations were approved by Ethics

Surgical procedures
All procedures have been performed fluoroscopically guided by unilateral pedicular approach under local anesthesia.

PVP procedure: local anesthesia (2% lidocaine and 1% rapivacaine, 1:1) was administered. Patients were positioned prone radio-lucent table. The orientation of puncture was located in the anterior 3/4 of the vertebral body under the guidance of C-arm X-ray machine. Subsequently, 3–5 ml of PMMA (Osteopal V, Heraeus Medical, Germany) was injected until adequate filling of the vertebral body with lateral fluoroscopic guidance.

The prophylactic vertebroplasty: the protocol was the same with PVP and the two operations could be done at the same time. But, the cement injected was a resorbable calcium sulfate/calcium phosphate composite bone graft (GeneX, Biocomposites Ltd, UK).

Evaluation of sandwich vertebrae
Postoperational examination including spinal radiographs was taken within 1 day, 6 months, 12 months, 24 months and thereafter. If a patient complaint a back pain or the spinal radiographs showed missing height of sandwich vertebral body, further magnetic resonance imaging or 99mTc bone scan was performed to confirm the presence of a new vertebral fracture.

Data collection
Data of patients within sandwich vertebra between two groups was analysed. A demographic database was created including age, sex, height, weight, bone mineral density (BMD) and follow-up period. The information was collected, including location of sandwich vertebra level, cement leakage into the intervertebral disk of sandwich vertebra. BMD of lumbar vertebrae (L1–4) and hip was measured by dual-energy X-ray absorptiometry (Hologic Inc., USA). The cobb angle of each sandwich vertebra between two groups was also measured. The height compression ratio of sandwich vertebra (anterior height/posterior height) was calculated to prevent errors caused by X-ray magnification (Fig. 1).

Statistical analysis
All statistical analysis was conducted in SPSS 20.0. The age, body mass index (BMI), BMD and vertebral height compression rate were shown as average ± standard deviation. Between groups, independent sample T test was performed. Within the group, paired sample T test was performed. A chi square test was performed for data comparison. \( P < 0.05 \) was taken as statistically significant.

Results
We performed PVP on 667 cases of osteoporotic vertebral fractures in the past 3 years. Fifty-five cases (10 males, 45 females; mean age 75.31 ± 7.91 years) with 71 sandwich vertebrae were involved in the study. The mean follow-up was 18.5 ± 7.3 months (range, 9–36 months). The incidence of sandwich vertebra was 8.25% (55/667). The mean BMI was 22.66 ± 2.77 kg/m². The BMD of lumbar and total hip was \( 0.688 \pm 0.123 \) g/cm² and \( 0.627 \pm 0.226 \) g/cm², respectively.

Overall, we observed 47 vertebrae (37 cases) in the non-prevention group and 24 vertebrae (18 cases) in the prevention group. There was no significant difference in age, sex, height, body weight, BMI or follow-up between the two groups (Table 1). In the non-prevention group, there were seven cases had cement leakage into the disc of the sandwich vertebra. However, in the prevention group, three patients suffered from cement leakage into the disc of the sandwich vertebra. No significant difference of occurrence rates of cement leakage had been observed (Table 1). Most of the sandwich vertebrae were located in the thoracic–lumbar junction (Figs 2 and 3). The distribution of sandwich vertebra was similar between the two groups (Table 1).

In the non-prevention group, eight patients (21.6%) had experienced eight sandwich vertebrae (17%) fractures. We described pre- and postoperation images of a case in the non-prevention group (Fig. 4). However, there was no patients who suffered new fracture
of sandwich vertebra in the prevention group. We also presented pre- and postoperation images of a prophylactic case (Fig. 5). Significant difference of the new fracture occurrence was observed between the two groups (Table 2). However, rates of new fracture of sandwich vertebra were 21.9% in thoracic–lumbar junction (T11-L1 segment), compared with 6.7% in the other segments ($P = 0.381$, Table 3). Furthermore, we showed that the cobb angle increased and vertebral anterior height decreased significantly in the non-prevention group ($P < 0.05$, Table 4). No significant change of cobb angle or vertebral height was observed in the prevention group.

**Discussion**

PVP is gaining popularity in the treatment of OVCFs, for its minimal invasion and rapid pain relief [14–16]. However, the vertebral bodies adjacent to the vertebrae previously treated by PMMA are more likely to develop new fractures. Most studies showed that the rates of new fractures in adjacent vertebra were relatively higher than non-adjacent level [10, 17–20]. In this study, we found that the rates of new fractures in sandwich vertebrae fractures were 21.60%, which was similar with the previous report (21.43%) [10]. It seemed that the sandwich vertebrae were easier to fracture than the adjacent vertebral fractures.

It was altered biomechanics in the treated vertebrae that developed an adjacent segment fracture [8, 9]. But, there were no reports about altered biomechanics in sandwich vertebrae. It is generally recognized that sandwich vertebra sustained double load shift and suffered strain concentration produced by the two adjacent augmented vertebrae. The risk factors of new fractures of adjacent vertebra were investigated in many studies. The amount of PMMA injected per vertebral body [8], intradiskal cement leakage [21–24], location of the adjacent vertebra [25], local kyphotic angulation [10] and lower BMD [17, 24] were all proved to increase risk of developing new fractures in adjacent vertebra. However, no significant difference of the risk factors was observed between the two groups. Interestingly, most of the sandwich vertebrae in our study were located in the thoracic–lumbar junction, especially T11-L1 segment. But, there was no statistically significant difference. The sample size may be not enough or others reasons.

Prophylactic vertebral reinforcement was limited in vitro studies so far. The injected PMMA was shown to increase the loading of on-axis [26, 27] or off-axis [28]. Regardless of beneficial results in the research, the spinal column was far more complicated than the single sample. There were many limits for PMMA in prophylactic vertebral reinforcement, for its biomechanical properties such as too high stiffness, strength and inabsorable. So it remained controversial in prophylactic vertebral reinforcement with PMMA [29, 30]. In Uebelhart’s report [31], they found refractures in the prophylactic

![Figure 1. Lateral radiograph showed that the sandwich vertebral body located between two cement treated vertebrae. Cobb angle was the angle between a and b. The ratio between anterior height and posterior height was c/d](image)

![Figure 2. Locations of total 25 sandwich vertebrae in prevention group, whereas most of them were T11 and L1](image)
vertebral reinforcement with PMMA. The reasons might be the increase of bone fragility caused by long history of glucocorticoid intake. Meanwhile, it was confirmed that only the endplate-to-endplate filling of PMMA would increase the vertebral intensity [32, 33]. It seemed that sandwich vertebrae was more easily to refraction when the PMMA was not adequately filled, especially in patients with increased bone fragility. Moreover, during PMMA polymerization, the induced heat and MMA toxicity damage has been proved in some studies [34, 35], resulting in aseptic inflammation and bone necrosis, thereby developing new fractures [36].

Nowadays, some absorbable bone graft substitutes were studied in vitro and in vivo trials [11, 12, 37–40], including calcium sulfate, calcium phosphate and the mixture of both materials. In our research, the bioactive and resorbable cement was GeneX, which was an injectable bone substitute consisting of calcium sulfate and beta-tricalcium phosphate with a weight ratio of 1/1. The calcium sulfate/calcium phosphate composite bone cement powder had a reaction with curing liquid with 10–15 min of coagulation, which was similar to PMMA. Once mixed, powder and liquid composed a viscous paste able to be easily injected. During the complete hardening period, the composite bone cement became solid providing mechanical strength. The calcium sulfate will be gradually resorbed allowing the implant to be remodeled through bone ingrowth [37]. However, the strength of calcium sulfate alone is too weak, and its rate of resorption is too high. The calcium phosphate could slow down the absorption speed of calcium sulfate and at the same time act as an osteoconductive template for new bone ingrowth. The calcium phosphate could embedded inside new bone tissue during the new bone ingrowth [11, 12, 38–40]. However, there were few reports about prophylactic vertebral reinforcement and the localized treatment of osteoporosis applied with resorbable composite bone cement. In our opinion, the prophylactic vertebroplasty with the resorbable composite bone cement is not only used for the prevention of the new fracture but also a novel localized treatment for osteoporosis. Because the resorbable calcium sulfate/calcium phosphate composite bone cement could help the induction of new bone and increase the bone strength in patients with osteoporosis. As the strength of sandwich vertebra, or others with a risk of fractures, could be increased by the calcium sulfate/calcium phosphate composite bone cement, the possibility of the new fracture would decrease.
There were some limitations in our study. First of all, the level of osteoporosis was one of the risk factors of developing new fractures. Our study didn’t involve the effect of antiosteoporosis drugs on new fractures. Besides, the scale of the group was relatively small, especially for the prevention group. Therefore, we needed to expand the size of sample to further investigate the risk factors of new sandwich vertebral fractures. Finally, we couldn’t get histological sections of the sandwich vertebral during the follow-up, which can confirm osteogenesis effect of the absorbable bone graft.

Table 2. Incidence of new fractures in the two groups

| Patient, n | New F | No F | (%) | Vertebra, n | New F | No F | (%) |
|------------|-------|------|-----|------------|-------|------|-----|
| Prevention group | 0     | 18   | 0   | 0          | 24    | 0    | 0   |
| Non-prevention group | 8     | 29   | 21.6| 8          | 39    | 17.0 |      |
| Chi square test |       |      |     | P = 0.033* |       |      |     |

New F, new fracture; No F, no fracture. The difference is statistically significant *P < 0.05.

Table 3. Decrease of new fractures in non-prevention group

| Vertebr, n | New F | No F | (%) |
|------------|-------|------|-----|
| Thoracic-lumbar junction | 7     | 25   | 21.9|
| Non-thoracic-lumbar junction | 1     | 14   | 6.7 |
| Chi square test |       |      | P = 0.381 |

New F, new fracture; No F, no fracture. The difference is no statistically significant.
Table 4. Cobb angle and vertebral compression ratio between two groups

|                      | Prevention group | Non-prevention group |
|----------------------|------------------|----------------------|
| **Cobb angle**       |                  |                      |
| Post op              | 5.28 ± 3.62      | 3.92 ± 3.17          |
| L-FLU                | 5.85 ± 3.91      | 5.87 ± 4.27          |
| **P**                | 0.285            | 0.000*               |

L-FLU, last follow-up. In the non-prevention group, the Cobb angle increased and the vertebral compression rate decreased at follow-up (*P < 0.05).

### Conclusion

PVP, a safe and effective procedure, is used to treat multiple thoracic or/and lumbar fractures caused by osteoporosis, which could form the sandwich vertebra. It has a risk of developing new fractures in the sandwich vertebra. However, prophylactic vertebroplasty applied with calcium sulfate/calcium phosphate composite bone cement can decrease the rate of new fractures in sandwich vertebra.

### Conflict of interest statement

None declared.

### References

1. Delmas PD, van de Langerijt L, Watts NB et al. Underdiagnosis of vertebral fractures is a worldwide problem: the IMPACT study. *J Bone Miner Res* 2005;20:557–63.
2. Riggs BL, Melton LJ III. The worldwide problem of osteoporosis: insights afforded by epidemiology. *Bone* 1995;17:505–11S.
3. Silverman SL. The clinical consequences of vertebral compression fractures. *Bone* 1992;13(Suppl 2):S27–31.
4. Mathis JM, Barr JD, Belkoff SM et al. Percutaneous vertebroplasty: a developing standard of care for vertebral compression fractures. *AJNR Am J Neuroradiol* 2001;22:373–81.
5. Wanger M, Markwalder TM. Re: percutaneous vertebroplasty for pain relief and spinal stabilization (Spine 2000; 25: 923–8). *Spine* 2000;25:2968–9.
6. Diamond TH, Hartwell T, Clarke W et al. Percutaneous vertebroplasty for acute vertebral body fracture and deformity in multiple myeloma: a short report. *Br J Haematol* 2004;124:485–7.
7. McGirt MJ, Parker SL, Wolinsky JP et al. Vertebroplasty and kyphoplasty for the treatment of vertebral compression fractures: an evidenced-based review of the literature. *Spine* 2009;34:501–8.
8. Berlemann U, Ferguson SJ, Nolte LP et al. Adjacent vertebral failure after vertebroplasty. A biomechanical investigation. *J Bone Joint Surg Br* 2002;84:748–52.
9. Polkeik A, Nolte LP, Ferguson SJ. The effect of cement augmentation on the load transfer in an osteoporotic functional spinal unit: finite-element analysis. *Spine* 2003;28:991–6.
10. Wang L, Yang H, Shi Y et al. Sandwich vertebral fracture in the study of adjacent-level fracture after vertebral cement augmentation. *Orthopedics* 2012;35:e1225–30.
11. Smeets R, Kolk A, Gerressem M et al. A new biphasic osteoconductive calcium composite material with a negative Zeta potential for bone augmentation. *Head Face Med* 2009;5:13.
12. Nilsson M, Wang JS, Wielanek L et al. Biodegradation and biocompatibility of a calcium sulphate-hydroxyapatite bone substitute. *J Bone Joint Surg Br* 2004;86:120–5.
13. Masala S, Anselmetti GC, Marcia S et al. Percutaneous vertebroplasty in multiple myeloma vertebral involvement. *J Spinal Disord Tech* 2008;21:344–8.
14. Clarencou F, Fahed R, Gabrieli J et al. Safety and clinical effectiveness of percutaneous vertebroplasty in the elderly (≥80 years). *Eur Radiol* 2016;26:2352–8.
15. Diamond TH, Bryant C, Browne L et al. Clinical outcomes after acute osteoporotic vertebral fractures: a 2-year non-randomised trial comparing percutaneous vertebroplasty with conservative therapy. *Med J Aust* 2006;184:113–7.
16. Santos RF, Ribeiro JC, de Moraes FB et al. Evaluation of the quality of life after vertebroplasty to treat compressive osteoporotic fractures. *Rev Bras Ortop* 2014;49:477–81.
17. Uppin AA, Hirsch JA, Centenera LV et al. Occurrence of new vertebral body fracture after percutaneous vertebroplasty in patients with osteoporosis. *Radiology* 2003;226:119–24.
18. Kim YY, Rhyu KW. Recompression of vertebral body after balloon kyphoplasty for osteoporotic vertebral compression fracture. *Eur Spine J* 2010;19:1907–12.
19. Trout AT, Kallmes DF, Kaufmann TJ. New fractures after vertebroplasty: adjacent fractures occur significantly sooner. *AJNR Am J Neuroradiol* 2006;27:217–23.
20. Tanigawa N, Kariya S, Komemushi A et al. Percutaneous vertebroplasty for osteoporotic compression fractures: long-term evaluation of the technical and clinical outcomes. *AJR Am J Roentgenol* 2011;196:1415–8.
21. Komemushi A, Tanigawa N, Kariya S et al. Percutaneous vertebroplasty for osteoporotic compression fracture: multivariate study of predictors of new vertebral body fracture. *Cardiovasc Intervent Radiol* 2006;29:580–5.
22. Lin EP, Ekholm S, Hiwatashi A et al. Vertebroplasty: cement leakage into the disc increases the risk of new fracture of adjacent vertebral body. *AJNR Am J Neuroradiol* 2004;25:175–80.
23. Chen WJ, Kao YH, Yang SC et al. Impact of cement leakage into disks on the development of adjacent vertebral compression fractures. *J Spinal Disord Tech* 2010;23:35–9.
24. Kim MH, Lee AS, Min SH et al. Risk factors of new compression fractures in adjacent vertebrae after percutaneous vertebroplasty. *Asian Spine J* 2011;5:180–7.
25. Kim SH, Kang HS, Choi JA et al. Risk factors of new compression fractures in adjacent vertebrae after percutaneous vertebroplasty. *Acta Radiol* 2004;45:440–5.
26. Furtado N, Oakland RJ, Wilcox RK et al. A biomechanical investigation of vertebroplasty in osteoporotic compression fractures and in prophylactic vertebral reinforcement. *Spine* 2007;32:E480–7.
27. Huggins KB, Harten RD, Langrana NA et al. Biomechanical effects of unipedicular vertebroplasty on intact vertebrae. *Spine* 2003;28:1540–7; discussion 8.
28. Aquarius R, Homminga J, Hosman AJ et al. Prophylactic vertebroplasty can decrease the fracture risk of adjacent vertebrae: an in vitro cadaveric study. *Med Eng Phys* 2014;36:944–8.
29. Becker S, Garosch M, Meissner J et al. Is there an indication for prophylactic balloon kyphoplasty? A pilot study. *Clin Orthop Relat Res* 2007;458:83–9.
30. Kobayashi N, Numaguchi Y, Fuwa S et al. Prophylactic vertebroplasty: cement injection into non-fractured vertebral bodies during percutaneous vertebroplasty. *Acad Radiol* 2009;16:136–43.
31. Uebelhart B, Casez P, Rizzoli R et al. Prophylactic injection of methylenbis(tacrylate in vertebrae located between two previously cemented levels does not prevent a subsequent compression fracture in a patient with bone fragility. *Joint Bone Spine* 2008;75:322–4.
32. Steens J, Verdoescht N, Aalsma AM et al. The influence of endplate-to-endplate cement augmentation on vertebral strength and stiffness in vertebroplasty. *Spine* 2007;32:E419–22.
33. Chevalier Y, Pahr D, Charlebois M et al. Cement distribution, volume, and compliance in vertebroplasty: some answers from an anatomy-based nonlinear finite element study. *Spine* 2008; 33: 1722–30.

34. Anselmetti GC, Manca A, Kanika K et al. Temperature measurement during polymerization of bone cement in percutaneous vertebroplasty: an in vivo study in humans. *Cardiovasc Intervent Radiol* 2009; 32:491–8.

35. Lieberman IH, Togawa D, Kayanja MM. Vertebroplasty and kyphoplasty: filler materials. *Spine* 2005; 30:555–68.

36. Heo DH, Chin DK, Yoon YS et al. Recollapse of previous vertebral compression fracture after percutaneous vertebroplasty. *Osteoporos Int* 2009;20:473–80.

37. Stubbs D, Deakin M, Chapman-Sheath P et al. In vivo evaluation of resorbable bone graft substitutes in a rabbit tibial defect model. *Biomaterials* 2004;25:5037–44.

38. Masala S, Nano G, Marcia S et al. Osteoporotic vertebral compression fracture augmentation by injectable partly resorbable ceramic bone substitute (Cerament®SPINESUPPORT): a prospective nonrandomized study. *Neuroradiology* 2012; 54:1245–51.

39. Yang HL, Zha XS, Chen L et al. Bone healing response to a synthetic calcium sulfate/beta-tricalcium phosphate graft material in a sheep vertebral body defect model. *J Biomed Mater Res B Appl Biomater* 2012;100:1911–21.

40. Abramo A, Geijer M, Kopylov P et al. Osteotomy of distal radius fracture malunion using a fast remodeling bone substitute consisting of calcium sulphate and calcium phosphate. *J Biomed Mater Res B Appl Biomater* 2010;92:281–6.