The available evidence on demineralised bone matrix in trauma and orthopaedic surgery

Citation for published version (APA):

Document license:
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DOI:
10.1302/2046-3758.67.BJR-2017-0027.R1

Document status and date:
Published: 01/07/2017

Document Version:
Publisher’s PDF, also known as Version of Record (includes final page, issue and volume numbers)

Please check the document version of this publication:

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The available evidence on demineralised bone matrix in trauma and orthopaedic surgery

A SYSTEMATIC REVIEW

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Maastricht University Medical Centre, Maastricht, The Netherlands

Objectives
The aim of this systematic literature review was to assess the clinical level of evidence of commercially available demineralised bone matrix (DBM) products for their use in trauma and orthopaedic related surgery.

Methods
A total of 17 DBM products were used as search terms in two available databases: Embase and PubMed according to the Preferred Reporting Items for Systematic Reviews and Meta Analyses statement. All articles that reported the clinical use of a DBM-product in trauma and orthopaedic related surgery were included.

Results
The literature search resulted in 823 manuscripts of which 64 manuscripts met the final inclusion criteria. The included manuscripts consisted of four randomised controlled trials (level I), eight cohort studies (level III) and 49 case-series (level IV). No clinical studies were found for ten DBM products, and most DBM products were only used in combination with other grafting materials. DBM products were most extensively investigated in spinal surgery, showing limited level I evidence that supports the use of Grafton DBM (Osteotech, Eatontown, New Jersey) as a bone graft extender in posterolateral lumbar fusion surgery. DBM products are not thoroughly investigated in trauma surgery, showing mainly level IV evidence that supports the use of Allomatrix (Wright Medical, London, United Kingdom), DBX (DePuy Synthes, Zuchwil, Switzerland), Grafton DBM, or OrthoBlast (Citagenix Laval, Canada) as bone graft extenders.

Conclusions
The clinical level of evidence that supports the use of DBM in trauma and orthopaedic surgery is limited and consists mainly of poor quality and retrospective case-series. More prospective, randomised controlled trials are needed to understand the clinical effect and impact of DBM in trauma and orthopaedic surgery.

Cite this article: Bone Joint Res 2017;6:423–432.

Keywords: Bone defects, Bone grafts, Demineralised bone matrix

Article focus
What is the available clinical level of evidence that supports the use of demineralised bone matrix (DBM) in trauma and orthopaedic surgery?

Key messages
The majority of the available clinical level evidence supporting the use of DBM products in trauma and orthopaedic surgery consist of case series (level IV evidence).

The use of DBM products has been most extensively investigated in spinal surgery, with level I evidence that supports the use of Grafton DBM (Osteotech, Eatontown, New Jersey) as a bone graft extender in posterolateral lumbar fusion surgery. The use of DBM products is not well investigated in trauma surgery, with only case series that mainly describe the use of Allomatrix (Wright Medical, London,
United Kingdom), DBX (DePuy Synthes, Zuchwil, Switzerland), Grafton DBM, and OrthoBlast (Citagenix Laval, Canada) as bone graft extenders in trauma surgery (level III/IV evidence).

**Strengths and limitations**
- Product-specific overview of DBM used for specific trauma- and orthopaedic-related indications.
- Inclusion of a large number of papers describing the clinical outcomes of DBM products used in trauma and orthopaedic surgery.
- Not all DBM products that are commercially available worldwide could be included.

**Introduction**
Bone grafting is a common procedure in trauma and orthopaedic surgery, with more than two million procedures being performed worldwide each year. Autologous bone is the gold standard grafting material and it is mostly harvested from the iliac crest. However, this harvesting procedure has a considerable morbidity rate (8% to 39%), and the amount of bone that can be harvested is not always sufficient. These issues resulted in the development of alternative bone graft materials. These alternative bone graft materials have been used as a bone graft extender (reinforcing autologous bone) or as a bone graft substitute (replacing autologous bone).

Demineralised bone matrix (DBM) is processed allogeneic bone that has been demineralised by extensive decalcification procedures. These procedures include chemical and radiation steps to minimise immunogenic response and the risk of infection. The resulting material consists of matrix proteins containing certain quantities of osteo-inductive growth factors (e.g. bone morphogenetic proteins). In 1965, Urist showed that DBM maintains its osteo-inductive potential, since subcutaneous implantation led to de novo bone formation in rabbits. This was later confirmed by Geesink et al in a small human series, where five out of six critical fibular bone defects were successfully treated with DBM. However, whether these osteo-inductive properties can be consistently reproduced in a DBM product remains heavily debated. The preserved amount of bone morphogenetic protein (BMP) after demineralisation merely lies within the nanogram range and, moreover, the absolute amount of BMP per DBM product varies up to fourfold between various batches of the same DBM product.

Nevertheless, many trauma and orthopaedic surgeons consider DBM to be a useful bone graft substitute for a wide range of clinical indications in trauma and orthopaedic surgery. Subsequently, the number of commercially available DBM products is constantly increasing, which is possible due to favourable regulatory pathways that allow quick access of new products onto the clinical market (i.e. DBMs are not regulated under 510(k) regulation, but are considered minimally manipulated tissue for transplantation). Nevertheless, there is no evidence-based guideline available that assists trauma and orthopaedic surgeons in making evidence-based decisions regarding the use of DBM products in different clinical indications. Therefore, the aim of this systematic review was to provide the clinical level of evidence that supports the use of DBM products in trauma- and orthopaedic-related surgery.

**Materials and Methods**

**Literature search.** Product names of commercially available DBM products in the Netherlands (Table I) were used as search terms in two available online databases: PubMed and Embase. The search key used for PubMed was: “(product name) AND bone[Title/Abstract]” and for Embase, it was: “product name AND bone:ab,”. The databases were searched from the earliest date available until 01 January 2017. Independently, two researchers (JvDS and KAH) performed the search. Manuscripts describing original studies on the use of DBM products written in English, German and Dutch were considered eligible. In order to select manuscripts primarily concerning the clinical use of DBM products for trauma- and orthopaedic-related indications, manuscripts were excluded if they only contained in vitro data or animal experiments or reported on the use of DBM products in non-orthopaedic-related indications (e.g. dental or maxillofacial surgery). References in the selected manuscripts were reviewed in order to ensure that no papers were missed with the chosen search strategy. All included manuscripts were assigned a level of evidence by two agreeing reviewers (JvDS and KAH) for their use as a bone graft extender or bone graft substitute, as described by Wright et al (Table II).

**Results**

**Literature search.** The initial literature search in PubMed and Embase resulted in a total of 823 papers. After screening all titles and abstracts, 101 manuscripts from PubMed and 140 manuscripts from Embase were considered eligible. Exclusion of duplicate articles resulted in a total of 149 eligible manuscripts. Of these 149 eligible manuscripts, 93 were excluded based on the exclusion criteria described, and eight manuscripts were added based upon the reference list. This resulted in a final number of 64 manuscripts that fulfil all selection criteria (Fig. 1). A detailed overview specified per product is provided in Table III.

**Orthopaedic-related indications.** DBM products have been used as a bone graft extender or as a bone graft substitute for a wide range of trauma- and orthopaedic-related indications (Table IV). Most studies were case series (n = 49); there were only four randomised control trials (RCTs) and eight cohort studies. The remaining three
Table I. Product specifications of commercially available demineralised bone matrix (DBM)

<table>
<thead>
<tr>
<th>Product</th>
<th>Manufacturer</th>
<th>DBM %</th>
<th>Carrier</th>
<th>Form</th>
<th>FDA</th>
<th>Indication</th>
</tr>
</thead>
<tbody>
<tr>
<td>Accell Connexus</td>
<td>Citagenix, Laval, Canada</td>
<td>70</td>
<td>Reverse phase medium</td>
<td>Paste</td>
<td>510(k)</td>
<td>Bone void filler/bone graft extender</td>
</tr>
<tr>
<td>Accell TBM</td>
<td>Integra, Irvine, California</td>
<td>100</td>
<td>-</td>
<td>Strips</td>
<td>510(k)</td>
<td>Bone void filler/bone graft extender</td>
</tr>
<tr>
<td>AlloCraft</td>
<td>Stryker, Mahwah, New Jersey</td>
<td>80</td>
<td>Acellular matrix</td>
<td>Paste</td>
<td>510(k)</td>
<td>Bone void filler</td>
</tr>
<tr>
<td>AlloMatrix</td>
<td>Wright Medical, London, United Kingdom</td>
<td>40 to 86</td>
<td>Calcium sulphate</td>
<td>Paste</td>
<td>510(k)</td>
<td>Bone void filler</td>
</tr>
<tr>
<td>AlphaGRAFT</td>
<td>Alphatech, Carlsbad, California</td>
<td>80</td>
<td>Acellular matrix</td>
<td>Paste</td>
<td>510(k)</td>
<td>Bone void filler</td>
</tr>
<tr>
<td>Altiva</td>
<td>Exactech, Gainesville, Florida</td>
<td>ND</td>
<td>Gelatin</td>
<td>Paste</td>
<td>510(k)</td>
<td>Bone void filler</td>
</tr>
<tr>
<td>BioSet</td>
<td>Penta Biomedical, Verona, Italy</td>
<td>24</td>
<td>Gelatin</td>
<td>Paste/strips</td>
<td>510(k)</td>
<td>Bone void filler</td>
</tr>
<tr>
<td>DBX</td>
<td>DePuy Synthes, Zuchwil, Switzerland</td>
<td>32</td>
<td>Hyaluronic acid</td>
<td>Paste/strips</td>
<td>510(k)</td>
<td>Bone graft extender/bone void filler</td>
</tr>
<tr>
<td>Grafton DBM</td>
<td>Osteotech, Eatontown, New Jersey</td>
<td>17 to 31</td>
<td>Glycerol</td>
<td>Paste/strips</td>
<td>510(k)</td>
<td>Bone graft substitute/bone graft extender/bone void filler</td>
</tr>
<tr>
<td>InterGro</td>
<td>Zimmer Biomet., Westminster, California</td>
<td>40</td>
<td>Lecithin</td>
<td>Paste</td>
<td>510(k)</td>
<td>Bone graft extender/bone void filler</td>
</tr>
<tr>
<td>Optefil</td>
<td>Exactech</td>
<td>24</td>
<td>Gelatin</td>
<td>Paste</td>
<td>510(k)</td>
<td>Bone void filler</td>
</tr>
<tr>
<td>Opteform</td>
<td>Exactech</td>
<td>ND</td>
<td>Cortical and cancellous bone chips suspended in collagen-gelatin</td>
<td>Paste</td>
<td>510(k)</td>
<td>Bone void filler</td>
</tr>
<tr>
<td>Optium DBM</td>
<td>DePuy, Warsaw, Indiana</td>
<td>ND</td>
<td>Glycerol</td>
<td>Paste</td>
<td>510(k)</td>
<td>Bone void filler</td>
</tr>
<tr>
<td>OrthoBlast</td>
<td>Citagenix</td>
<td>ND</td>
<td>Reverse phase medium</td>
<td>Paste</td>
<td>510(k)</td>
<td>Bone void filler</td>
</tr>
<tr>
<td>OrthoBlast II</td>
<td>Citagenix</td>
<td>20</td>
<td>Reverse phase medium</td>
<td>Paste</td>
<td>510(k)</td>
<td>Bone void filler</td>
</tr>
<tr>
<td>Osteofil</td>
<td>Medtronic, Minneapolis, Minnesota</td>
<td>24</td>
<td>Collagen</td>
<td>Paste/strips</td>
<td>510(k)</td>
<td>Bone void filler</td>
</tr>
<tr>
<td>VIAGRAF</td>
<td>Smith &amp; Nephew, London, United Kingdom</td>
<td>ND</td>
<td>Glycerol</td>
<td>Paste/strips</td>
<td>510(k)</td>
<td>Bone void filler</td>
</tr>
</tbody>
</table>

FDA, Food and Drug Administration; ND, no data available

Table II. Level of evidence as described by Wright et al.

<table>
<thead>
<tr>
<th>Level of evidence</th>
<th>Type of study</th>
</tr>
</thead>
<tbody>
<tr>
<td>I</td>
<td>1. RCT</td>
</tr>
<tr>
<td></td>
<td>2. Systematic review of level I RCT</td>
</tr>
<tr>
<td>II</td>
<td>1. Prospective cohort study</td>
</tr>
<tr>
<td></td>
<td>2. Poor-quality RCT</td>
</tr>
<tr>
<td></td>
<td>3. Systematic review level II studies</td>
</tr>
<tr>
<td>III</td>
<td>1. Case-control study</td>
</tr>
<tr>
<td></td>
<td>2. Retrospective cohort study</td>
</tr>
<tr>
<td></td>
<td>3. Systematic review of level III studies</td>
</tr>
<tr>
<td>IV</td>
<td>Case series (no, or historical, control group)</td>
</tr>
<tr>
<td>V</td>
<td>Expert opinion</td>
</tr>
</tbody>
</table>

RCT, randomised control trial

Studies reported the random use of DBM products among different study groups without reporting outcomes related to the use of DBM.11-13 The number of clinical studies per DBM product varies extensively. The most frequently examined DBM products, Grafton DBM (Osteotech, Eatontown, New Jersey) and AlloMatrix (Wright Medical, London, United Kingdom), were described in 33 and 18 studies, respectively. No clinical studies were found for ten of the 15 remaining DBM products.

**Spine surgery.** Four DBM products have been used in spinal surgery: Accell Connexus (Citagenix, Laval, Canada); AlloMatrix; Grafton DBM; and Osteofil (Medtronic, Minneapolis, Minnesota). These DBM products were used as bone graft extenders in most studies; only in two studies44,46 was Grafton DBM used as a bone graft substitute.

**Lumbar spinal fusion.** Accell Connexus was used as a bone graft extender in instrumented posterolateral lumbar fusions in 33 patients.44 In this randomised study, iliac crest bone was augmented with a mixture of Accell Connexus and autologous bone marrow aspirate, and compared with the use of iliac crest bone alone. After one year, grafting had resulted in similar fusion rates (70% versus 76%). There was no difference...
in pain or duration of surgery on the visual analogue scale (VAS). This study provides level II evidence that Accell Connexus may be used as a bone graft extender for lumbar spinal fusion.

Allomatrix has been used in posterolateral lumbar fusions. Fu et al.\textsuperscript{15} showed in a case-control study that Allomatrix or autologous bone resulted in comparable fusion rates when used with hydroxyapatite/tricalcium phosphate granules: 81% fusion with Allomatrix and 86% fusion with autologous bone respectively. Girardi and Cammisa\textsuperscript{16} described a retrospective case series of 65 patients where Allomatrix was mixed (1:1) with iliac crest bone. Radiological follow-up showed an improvement in the Lenke scores, 3.7 after one month to 1.6 after 12 months. In another retrospective case series\textsuperscript{13} following 32 patients who underwent posterior lumbar interbody fusion for 36 months (18 to 42), clinical and radiological scores improved significantly. At the latest follow-up, the mean Oswestry Disability Index improved from 52% to 22%. The mean Roland-Morris Disability Questionnaire improved from 52% to 29%, while > 90% of the operated levels were fused. These studies provide level III evidence that Allomatrix may be used as a bone graft extender for lumbar spinal fusion.

Grafton DBM was studied as a bone graft extender for posterolateral spinal fusion in a RCT by Cammisa et al in 2004.\textsuperscript{35} In 120 patients, posterolateral lumbar fusions were carried out with pedicle screw fixation and one side of the spine was grafted with autograft (17.2 standard deviation (sd) 9.7 ml), while the contralateral side was grafted with autograft and Grafton DBM (17.2 sd 9.7 ml, mixed 1:2). After two years, autograft with Grafton DBM resulted in fusion in 42 cases (52%) and autograft alone resulted in fusion of 44 cases (54%). In another prospective cohort study, 36 patients undergoing instrumented posterolateral lumbosacral spinal fusion were grafted with Grafton DBM and aspirated bone marrow (19 cases), Grafton DBM and autologous bone (27 cases), or autologous bone alone (27 cases). All groups showed similar fusion rates after two years' follow-up (63%, 70% and 67%, respectively). Use of Grafton DBM as a bone graft extender in posterolateral lumbar fusions is further described in three other studies\textsuperscript{37-39} encompassing 138 patients in total, where fusion rates after more than two years' follow-up ranged from 86% (Kang et al\textsuperscript{37}) to 93% (Thalgott et al\textsuperscript{38}). These studies include level I evidence that Grafton DBM can be used as a bone graft extender for lumbar spinal fusion.

<table>
<thead>
<tr>
<th>Products</th>
<th>Inclusion</th>
<th>Exclusion</th>
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<tbody>
<tr>
<td><strong>PubMed papers (n)</strong></td>
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<td></td>
<td></td>
</tr>
<tr>
<td><strong>Eligible papers (n)</strong></td>
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<td></td>
<td></td>
</tr>
<tr>
<td><strong>Duplicates</strong></td>
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<td></td>
<td></td>
</tr>
<tr>
<td><strong>Total</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>In vitro experiments</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>Animal experiments</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>Non-trauma- or orthopaedic-related indication</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>Additional references</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

**Table III.** Search results of systematic literature search

- Grafton DBM was studied as a bone graft extender for posterolateral spinal fusion in a RCT by Cammisa et al in 2004.\textsuperscript{35} In 120 patients, posterolateral lumbar fusions were carried out with pedicle screw fixation and one side of the spine was grafted with autograft (17.2 standard deviation (sd) 9.7 ml), while the contralateral side was grafted with autograft and Grafton DBM (17.2 sd 9.7 ml, mixed 1:2). After two years, autograft with Grafton DBM resulted in fusion in 42 cases (52%) and autograft alone resulted in fusion of 44 cases (54%). In another prospective cohort study, 36 patients undergoing instrumented posterolateral lumbosacral spinal fusion were grafted with Grafton DBM and aspirated bone marrow (19 cases), Grafton DBM and autologous bone (27 cases), or autologous bone alone (27 cases). All groups showed similar fusion rates after two years' follow-up (63%, 70% and 67%, respectively). Use of Grafton DBM as a bone graft extender in posterolateral lumbar fusions is further described in three other studies\textsuperscript{37-39} encompassing 138 patients in total, where fusion rates after more than two years' follow-up ranged from 86% (Kang et al\textsuperscript{37}) to 93% (Thalgott et al\textsuperscript{38}). These studies include level I evidence that Grafton DBM can be used as a bone graft extender for lumbar spinal fusion.
Osteofil was used as a bone graft extender for one-level (95 cases) and two-level (45 cases) posterolateral spinal fusions in a study from Epstein and Epstein.63 In this study, Osteofil was mixed (1:1) with autologous bone, and patient outcome was assessed by the sF-36 Questionnaire. One year post-operatively, clinical improvement was observed on six out of eight health scales of the SF-36 Questionnaire. In one-level fusion, 2D-cT showed fusion rates of 93% after an average of 5.2 (sd)1.8 months. Two patients required secondary surgery to treat nonunion or instability. In two-level fusion, 2D-cT showed fusion rates of 92% after an average of 6.1 to 1.9 months. Again, two patients required secondary surgery due to nonunion or instability.63 These studies provide level IV evidence that Osteofil may be used as a bone graft extender for lumbar spinal fusion.

Cervical spinal fusion. DBM products used in anterior cervical fusions include Allomatrix, Grafton DBM, and Osteofil. Allomatrix was used as a bone graft substitute in a study that included 29 patients suffering from craniocervical spine instability requiring occipitocervical fusion (OCF) due to trauma, rheumatoid arthritis or neoplasms.63 Two instrumentation techniques were compared (screw-rod instrumentation versus hook-and-screw-rod instrumentation), although the authors mention that Allomatrix was used in most cases. In fact, neither graft-specific outcome, nor graft-related complications were mentioned. This study provides no graft-specific outcomes on the use of Allomatrix as a bone graft substitute for cervical spinal fusions.

Grafton DBM was used in a RCT by An et al41 which included 77 patients undergoing anterior cervical fusion. Grafton DBM was combined with tricortical allografts and this was compared with tricortical autografts alone. Nonunion developed in 46% of patients who were grafted with Grafton DBM and allografts, compared with only 26% of patients who received a tricortical autograft (p = 0.11), suggesting that the combination of Grafton DBM and allograft results in a higher rate of nonunion. Grafton DBM was also used to fill, partially42 or fully,45 polyetheretherketone (PEEK) cages used for cervical fusion. Park et al42 used PEEK cages containing autologous bone chips and Grafton DBM for cervical fusion of 42 levels in 31 patients. After one year, a fusion rate of
97% was observed. Both the VAS score for neck and arm pain and the modified Japanese Orthopaedic Association (JOA) scoring system for myelopathy were significantly improved. Elsayaf et al describe complete filling of the PEEK cage with Grafton DBM in anterior cervical discectomy and fusion in a case series of 20 patients; the mean Cobb angle improved (3.4° pre-operatively vs 14.5° post-operatively) and JOA myelopathy scores and neck disability index also subsequently improved. These studies include level I evidence that Grafton DBM is not useful as a bone graft extender for cervical spinal fusion.

Osteofil was mixed with autologous bone (n = 11) in a study by Epstein, which also included 24 patients in which Vitoss (β-tricalcium phosphate) was mixed with autologous bone. Radiological follow-up showed that all levels were fused after an average of 5.2 months. Less than 50% of the original fusion mass remained visible on 2D-CT scans after six months in 64% of fusions grafted with Osteofil, compared with 21% of fusions grafted with Vitoss (Stryker, Mahwah, New Jersey), which suggested a quicker resorption rate of Vitoss. This study provides level IV evidence that Osteofil may be used as a bone graft extender with autologous bone for cervical spinal fusion. Thoracic spinal fusion. Grafton DBM was the only product also used for thoracic fusions. In a retrospective cohort, Weinzapfel et al used Grafton DBM in patients who underwent anterior thoracic discectomies and compared their results with using morsellised cancellous allografts. On the final radiographs, 82% of the allograft group and 92% of the Grafton DBM group were rated as fused. There was no significant difference between the two groups. This study provides level III evidence that Grafton DBM may be used as a bone graft substitute for thoracic spinal fusion.

Trauma surgery. Four DBM products have been used to treat bone defects that accompany fractures or were the result of nonunion: Allomatrix, DBX (DePuy Synthes, Zuchwil, Switzerland), Grafton DBM and OrthoBlast (Citagenix). They are used as a graft extender in most cases, however, they may also be used as a graft substitute. Fractures. Allomatrix was used to treat distal radial fractures. In a RCT, unstable distal radial fractures were treated by operative fixation with Kirschner-wires, with (n = 24) or without (n = 26) augmentation of the fracture site with Allomatrix. The physical and radiological outcomes did not show any significant difference between wrist function, speed to recovery, union rate, or complication rate during a one-year follow-up. Allomatrix has also been used for primary treatment of fresh bone defects caused by small-calibre gunshot wounds in the hand. In a retrospective case series of 12 patients, 11 bone defects healed without further intervention and one defect required a second bone grafting procedure. In another case report, Allomatrix was used to fill metaphyseal bone defects after elevation of the impressed osteochondral fragments in the treatment of reverse Hill-Sachs lesions of the humeral head. These studies include level I evidence that Allomatrix is not a useful bone graft substitute to treat unstable distal radial fractures already fixed with Kirschner-wires.

DBX has been used to treat sternal segment dislocations. Sternal segment dislocations in eight patients were treated with titanium screws and DBX. Titanium screws and DBX reduced the length of hospitalisation, and led to rapid functional recovery and excellent aesthetic results according to the authors. This study provides level IV evidence that DBX may be used as a bone graft substitute to treat fractures of the sternum.

Grafton DBM and OrthoBlast were applied as a bone graft extender by Cheung et al. Grafton DBM or OrthoBlast were mixed with cancellous allografts and used to graft bone defects encountered in periarticular fractures of the tibia, fibula, femur, humerus, forearm and acetabulum. Fracture healing occurred in 69% of the patients who received Grafton DBM (n = 13) compared with 100% in patients who received OrthoBlast (n = 15). Grafton DBM mixed with cancellous allograft was also used to graft bone defects to stimulate bone healing in smokers and non-smokers. After an average follow-up of 32 months, treatment was successful in 68% of the smokers compared with 88% of the non-smokers. Four patients with foot and ankle fractures (tibial/fibular fracture, fifth metatarsal fracture) were grafted with Grafton DBM or autografts. No complications were described. Grafton DBM was also used to enhance cancellous allografts in two tibial stress fractures treated by drilling and bone grafting, and for reconstructing large segmental bone defects of the tibia (n = 2) and humerus (n = 1) using a titanium mesh cage filled with Grafton DBM and cancellous allograft chips. These studies include level III evidence that both Grafton DBM and OrthoBlast may be used in combination with allograft as a bone graft substitute to treat bone defects during fracture surgery.

Nonunion. Allomatrix, mixed with cancellous allograft chips, was used to treat 41 atrophic or avascular nonunions. These nonunions were located in the femur, radius, tibia, and humerus. In 13 cases the nonunion recurred (32%), and revision surgery was necessary in 19 cases. Additionally, Allomatrix has been used to graft bone defects that resulted after nonunion (n = 35). Allomatrix was mixed (1:3) with calcium sulphate pellets, and after seven months, 85% of the grafted nonunions were healed. These studies provide level IV evidence that Allomatrix may be used as a bone graft substitute to treat nonunion.

DBX has been described in a case report, showing the successful treatment of a subtrochanteric nonunion of an 11-year-old patient with an adult proximal humeral locking plate and additional grafting with DBX. This study provides level IV evidence that DBX may be used as a bone graft substitute to treat nonunion.
Grafton DBM was used in a cohort study by Hierholzer et al.\textsuperscript{53} Treatment of 33 humeral nonunions with Grafton DBM were compared with 45 nonunions that received autologous bone grafts. Autologous bone grafts resulted in union in all cases, whereas Grafton DBM resulted in union in 97\% in a mean time to union of 4.2 and 4.5 months, respectively. Ziran et al.\textsuperscript{21,49} described the use of Grafton DBM to treat nonunion in two studies. First, Grafton DBM was mixed with cancellous allograft to stimulate bone healing in smokers and non-smokers.\textsuperscript{66} After an average follow-up of 32 months, treatment was successful in 68\% of the smokers versus 88\% of the non-smokers. Subsequently, treatment of nonunion in smokers with Grafton DBM (n = 25) was compared with OrthoBlast (n = 13).\textsuperscript{49} Treatment with Grafton DBM was only successful in 52\%, whereas treatment with OrthoBlast was successful in 85\%, which was not a statistically significant difference. In a case report,\textsuperscript{54} Grafton DBM was used to treat a non-displaced coracoid fracture. After screw fixation, the nonunion site was debrided and successfully grafted with Grafton DBM. These studies include level III evidence that Grafton DBM may be used as a bone graft substitute to treat nonunion and level IV evidence that Grafton DBM or OrthoBlast may be used as a bone graft extender to treat nonunion.

**Orthopaedic surgery.** Four DBM products, Allomatrix, DBX, Grafton DBM, and OrthoBlast, have been described for other clinical indications that included the treatment of arthrodesis, avascular necrosis, and tumour treatment. Allomatrix was added to a calcium sulphate bone substitute (OsteoSet) in a small case series by Deheshi, Allen and Kim,\textsuperscript{12} to be used as a graft to restore retroacetabular bone stock in seven patients who showed retroacetabular osteolysis after primary total hip arthroplasty with cementless acetabular sockets. Specific graft-related outcomes were not reported.\textsuperscript{12} These studies provide level IV evidence that Allomatrix can be used as a bone graft extender to treat acetabular bone defects.

Allomatrix has been used to treat benign bone tumours in the tibia (n = 17), humerus (n = 11), fibula (n = 3), and radius (n = 2). Defects were filled with a mixture of Allomatrix and a calcium sulphate bone substitute (OsteoSet, 1:3). After seven months, 93\% of the bone defects were healed, tumour recurrence was seen in three cases, and one wound infection required antibiotic treatment.\textsuperscript{23} In addition, in a study that investigated the treatment of 98 benign bone tumours located in the tibia, humerus, femur and pelvis with various bone grafts, Allomatrix was used in 34 of these grafting procedures but no Allomatrix-specific outcomes were reported.\textsuperscript{28} These studies provide level IV evidence that Allomatrix may be used as a bone graft substitute with or without a calcium sulphate bone substitute to treat benign bone tumours.

DBX was used as a graft in the treatment of enchondromas of the hand. Kwon and Wong,\textsuperscript{34} and Dietz, Kachar and Nagle,\textsuperscript{33} reported small case series of five and two patients, respectively. No recurrence or pathological fractures were reported. These studies provide level IV evidence that DBX may be used as a bone graft substitute to treat hand enchondromas.

Grafton DBM has been described in ankle and foot surgery. In a case series by Kado, Gambetta and Perlman, Grafton\textsuperscript{56} DBM was combined with autologous or allogenic bone for arthrodesis (n = 18) or osteotomies (n = 6). In addition, Grafton DBM (n = 37) and OrthoBlast (n = 26) were also used for complex ankle or hindfoot arthrodesis.\textsuperscript{52} Nonunions developed in five patients who had Grafton DBM (14\%) and in two patients who had OrthoBlast (8\%). Furthermore, Grafton DBM was used as a bone graft after core decompression of asymptomatic avascular necrosis of the hip in 37 patients,\textsuperscript{57} but it did not reduce the need for total hip arthroplasty compared with treatment of symptomatic avascular necrosis during the two years of follow-up. Finally, Grafton DBM was used in combination with autologous bone marrow for percutaneous injection of unicameral bone cysts in 33 patients. After six weeks, patients returned to everyday activity, and bone healing was confirmed radiologically after six months.\textsuperscript{59} Furthermore, solitary bone cysts in children could be treated with Grafton DBM.\textsuperscript{60,62} After filling the defects with Grafton DBM in seven cases, a continuous decrease in radiographic bone transparency was observed over a period of two years.\textsuperscript{60} These studies include level IV evidence that Grafton DBM can be used as a bone graft substitute with or without autologous bone marrow to treat benign bone tumours.

**Discussion**

The aim of this systematic review was to provide the clinical level of evidence for using DBM products in trauma and orthopaedic surgery. This review reveals that the clinical level of evidence is mainly limited to level IV studies: case series in which DBM products are described for numerous indications with or without additional autografts, allografts or calcium-based bone substitutes. Combined with the fact that the composition and amount of osteo-inductive growth factors may vary extensively between and within DBM products,\textsuperscript{8,67} it is not possible to point out specific indications where DBM products have added value in trauma and orthopaedic surgery. Formulating an evidence-based guideline to assist trauma and orthopaedic surgeons requires more well designed clinical studies that prove a beneficial effect of DBM compared with treatment with a bone graft substitute (negative control) or treatment including the current gold standard autografts (positive control).

The clinical use of DBM is most thoroughly described for cervical and lumbar spinal fusion surgery.\textsuperscript{68} In a systematic review by Zadegan et al.\textsuperscript{69} which included 12 studies, the fusion rate of DBM in cervical fusion surgery
was comparable with that of other graft materials. However, most studies did not specify which DBM product was used, which may possibly lead to false claims of the effect of specific DBM products. This review shows that there is only one level I study available that determines the effect of Grafton DBM in cervical fusion surgery. However, this study shows a higher rate of nonunion compared with that found in autologous bone grafting when it is used in cervical spine fusions. In another RCT, Grafton DBM was used in lumbar spinal fusion, showing that similar fusion rates were achieved when Grafton DBM was added to autologous bone versus autologous bone alone. Except for Allomatrix, there is hardly any evidence for using other DBM products in spinal fusion surgery.

The clinical use of DBM products in trauma surgery is not well supported. For example, the use of DBX is only described in a small case series of sternum fractures and a single case report of nonunion. Furthermore, the case-control study that used Grafton DBM or OrthoBlast in fracture treatment included only 13 and 15 cases and did not have a control group. The RCT by Hierholzer et al. which concludes that grafting humeral nonunions with Grafton DBM is as effective as using autologous bone reported union rates of 97% and 100%, respectively, after four months. These high union rates in both groups raise the question as to whether the re-osteosynthesis per se in these studies is effective or if other factors are responsible for such high union rates.

In conclusion, preclinical studies have shown that the in vitro and in vivo efficacy of DBM depends upon the preparation methods and carrier materials used. Due to these differences, it is important for trauma and orthopaedic surgeons to make evidence-based decisions based on available literature that specifies the DBM product used for each indication. This review provides an up-to-date overview of the currently available clinical evidence for 17 different DBM products. Overall, the level of evidence of the studies available to date is generally low, mainly including retrospective case series. The highest clinical level of evidence is available for using DBM products in spinal surgery where its performance is comparable with that of autologous bone grafting. The clinical level of evidence for using DBM products in trauma surgery is minimal (mainly level IV). Thus, no evident benefit is proven for treatment of fractures or nonunion. The same holds for orthopaedic indications such as arthrodesis, avascular necrosis, and tumour surgery. Therefore, there is a need for further studies.
for more level I (RCTs or prospective case-cohort) studies in order to understand clearly the clinical effects of DBM in trauma and orthopaedic surgery, and to stimulate the discussion regarding the effectivity of using DBM in trauma and orthopaedic surgery.

References
