Characterization of the inflammatory response to four commercial bone graft substitutes using a murine biocompatibility model

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Abstract: Bone grafting is utilized in nearly all orthopedic subspecialties and in most anatomic regions. Bone graft substitutes have the potential to offer similar efficacy as autogenous grafts without the morbidity of harvest. Several studies have noted the efficacy of new-generation bone substitute products, but few studies have evaluated their safety. This study characterizes and quantifies the inflammatory reaction to four different commercially available bone graft substitutes, which were examined using the in vivo murine air pouch biocompatibility model. One coralline hydroxyapatite product was chosen as an example of a purely osteoconductive material. Three demineralized bone matrix products were chosen to represent products that are both osteoconductive and osteoinductive. Samples were implanted in a murine air pouch and harvested after 14 days in situ. Pouch fluid was extracted, mRNA isolated, and reverse transcription polymerase chain reactions carried out to detect interleukin-1 gene expression as a marker for inflammation. In addition, multiple histological characteristics were examined to quantify cellular responses to the implanted materials. All bone graft substitutes induced a significant inflammatory response compared with negative controls. Histology and polymerase chain reaction data indicated that the level of inflammatory reaction was elevated in materials with a higher demineralized bone matrix to carrier proportion. The hydroxyapatite product generated a low inflammatory reaction. In conclusion, this study used an in vivo model of biocompatibility to demonstrate that a significant inflammatory reaction occurs when using implanted bone graft substitutes. When choosing a bone grafting method, surgeons should consider both the efficacy and safety of methods and materials used. Further studies are necessary to determine the ideal bone graft material to maximize efficacy while minimizing morbidity.

Keywords: bone graft, biocompatibility, inflammation, animal model

Introduction

Bone grafting is a ubiquitous tool of orthopedic surgery, because it is used in all subspecialties and in most anatomic regions. The bone source may be from the patient (autograft) or a cadaver (allograft). However, problems exist with both of these sources. Harvesting autograft bone prolongs the operative time and may cause significant donor-site complications. The use of allograft bone carries the added risk of disease transmission as well as the potential for inflammatory or immunologic reactions to foreign tissue.

Although autografts remain the gold standard for most bone grafting applications, recent advances have suggested that alternatives may be available that offer less potential morbidity. There have been significant advances in the development of bone graft substitutes over the past decade. If these substitutes can be demonstrated to be
There are several engineered bone graft substitutes available. For example, a fresh autograft possesses structure, necessary signaling molecules, and cells required to produce new bone. The murine air pouch model for biocompatibility testing was ignited by the area in mm² to determine cellular density, as described elsewhere. 

Histologic evaluation and image analysis

Tissue samples were fixed, dehydrated, and embedded in paraffin blocks. Particular care was taken to maintain the original shape and orientation of the sample relative to its position on the mouse. Sections were stained with hematoxylin and eosin to evaluate the cellular infiltrates within the pouch and pouch membrane. A minimum of four separate sections per specimen were examined using ImagePro image analysis software package (Media Cybernetics, Bethesda, MD). Pouch thickness was measured at six points distributed evenly along the circumference of each pouch. The density of nucleated cells was measured within the thickness of the pouch membrane and within the pouch cavity. The total number of cells was measured within at least four representative areas and divided by the area in mm² to determine cellular density, as described previously.

Cytokine gene activation

Activation of inflammatory cytokine genes was measured using reverse transcription-polymerase chain reactions (PCR). Gene expression of the proinflammatory cytokine interleukin-1
(IL-1β) was measured in cells obtained from the pouch fluid aspirate. Total RNA was extracted from this fluid according to the manufacturer’s instructions (Tel-Test, Friendswood, TX). A cDNA library was constructed from 5 µg RNA extract in 50 µL solution containing 5 µL 10X PCR buffer, 1 µL 25 mM MgCl₂, 20 µL deoxynucleotide triphosphates, 50 units RNase inhibitor, 5 µL random hexamers, and 250 units reverse transcriptase (Perkin-Elmer Cetus, Norwalk, CT). The reaction mixture was incubated for 10 minutes at 25°C, 25 minutes at 48°C, and 5 minutes at 95°C. The IL-1 primer sequence was obtained from Clontech Laboratories Inc (Mountain View, CA). A total of 2 µL of cDNA was combined with 25 µL reaction solution containing 0.5 µL primer pair, 2.5 µL PCR buffer, 1.5 µL MgCl₂, 2 µL nucleotide triphosphates (NTPs), and 1.25 U AmpliTaq DNA polymerase (Perkin-Elmer Cetus). PCR was initiated by heating at 94°C for 1 minute to denature RNA-cDNA hybrids, followed by annealing of the primers at 60°C for 1 minute and then extension of the primer sequences at 72°C for 1 minute. This sequence was repeated 35 times using the DNA thermal cycler (Perkin-Elmer Cetus). Following amplification, 10 µL of each reaction solution was mixed with 1 µL loading buffer for agarose gel electrophoresis. Densities of cytokine bands were measured using ultraviolet light and the ISO 2000 Digital Imaging System (Alpha Innotech, San Leandro, CA). All cytokine levels were normalized using the PCR product from the housekeeping gene GAPDH.

Statistics
The analysis of variance (ANOVA) test with post hoc analysis using the least significant difference formula was used to compare means for all histological parameters, including membrane thickness, density of cellular infiltrate within the pouch membrane, and density of cellular infiltrate within the cavity of the pouch. When the overall F-test for differences among treatment means was significant, pairwise comparisons of treatment means were conducted. For the PCR data, single-factor ANOVA was again used to test for differences among means. When the overall F-test was significant, treatment means were compared using Student’s t-test. All results were considered significant for $P < 0.05$.

Results
Air pouch membrane
The histological characteristics of the air pouch membrane were assessed for each of the implanted substances and compared with the saline control, because the inflammatory response to implanted biomaterials can be characterized by increases in both the membrane thickness and the density of cellular infiltrate within the membrane wall. In saline-treated mice, the air pouch membrane was characterized by an outer fibrous layer composed primarily of fibroblastic cells and an inner portion composed of inflammatory cells, predominantly macrophages (Figure 1A). The membranes had a mean thickness of 47.7 µm and a cell number of 83.4 cells/mm². As shown in Figure 2, pairwise comparisons showed a significantly higher membrane thickness for each bone graft substitute when compared with the saline control ($P < 0.05$). Additionally, the ProOsteon and the DBX had significantly thicker membranes than either of the Accell bone graft substitutes ($P < 0.05$). No other mean differences between compounds were significant. The cellular density within the air pouch membrane walls did not differ between the saline control and any of the compounds tested using single-factor ANOVA (data not shown).

Air pouch cellular infiltrate
The density of the cellular infiltrate was examined within the air pouch cavity. The cavities of the test compounds each contained regions of immature bone as well as regions of dense cellular infiltrates (see Figure 1B–E). As shown in Figure 3, using pairwise comparisons, a statistically significant differ-
ence in the density of the cellular infiltrate was observed among all groups ($P < 0.05$). ProOsteon showed significantly less cellular infiltration than all of the DBM products. Among the DBM products, there was an increasing density of the cellular infiltrate as the ratio of DBM to carrier increased. The ratio of macrophage/monocytes to fibroblasts in the infiltrate was determined using image analysis as previously described.20

Significantly elevated numbers of monocytes were observed in pouch tissues associated with Accell Connexus and Accell DBM-100 (Figure 4) when compared with the control tissue ($P < 0.05$), whereas the ratios of monocytes:fibroblasts in pouch tissues associated with ProOsteon and DBX were not significantly different from control pouch tissues.

**Cytokine gene activation in the air pouch**

Reverse transcription-PCR was performed using the mRNA extracted from cells within the pouch fluid. Murine IL-1 primers were used to determine IL-1 expression as a marker to quantify the inflammatory reaction within the cavity of the pouch. Results are shown in Figure 5. Compared with the saline control, all compounds showed significantly higher levels of IL-1 expression ($P < 0.05$). The highest level of IL-1 expression was observed in the Accell DBM-100-containing air pouches; the increased expression was statistically significant when compared with both ProOsteon and Accell Connexus ($P < 0.05$).

**Discussion**

Recent studies have helped to clarify the optimum makeup of DBM products, including the source of allograft, preparation methods, particle size, and different carrier substances.21 Several studies have suggested that differing methods of graft preparation can cause differences in efficacy.14,22 Unfortunately, few studies have examined how differences in bone graft substitute production may cause variations in biocompatibility with adverse effects.

All DBM products are prepared from bone allografts that are first treated with a variety of washes, antibiotic treatments, irradiation, and other treatments to remove potentially infectious agents. In addition to removing bacterial contaminants,
processing has been shown to inactivate multiple infectious viruses, including human immunodeficiency virus, hepatitis B and C viruses, and cytomegalovirus. It is not yet clear whether differences in efficacy result from differences in the carriers or are as a result of other differences in preparation. The tissue is then demineralized using an acid solution, usually hydrochloric acid. Finally, the DBM is then mixed with a carrier to alter the texture, viscosity, and other characteristics of the final packaged product.

There are other factors that can affect the efficacy and safety of DBM products. Animal and human studies have shown that the age of the donor can significantly impact the osteoinductive potential for DBM. There are also significant differences in the type of carrier used and in the concentration of DBM within that carrier. Different carriers include glycerol, porcine collagen, hyaluronic acid, lecithin, calcium sulfate, or no carrier at all. Most DBM products contain approximately 20%–40% DBM by weight. In addition to the carrier substance in the DBM product, some investigators have also looked at various additional delivery systems added at the time of implantation to further alter the physical properties of the DBM. The fact that there are so many different materials and techniques used to manufacture similar products all intended for the same purpose means that the ideal makeup of a bone graft substitute is as yet undefined.

This study was designed to examine the inflammatory characteristics of several commercially available bone graft substitutes. Three demineralized bone graft products and one synthetic hydroxyapatite compound were studied using the murine air pouch. Overall, all products showed a significant inflammatory response within the pouch when compared with the saline control (summarized in Table 1). The hydroxyapatite material showed less inflammation than the DBM products. Within the DBM products tested, there was a trend toward increasing inflammation with increasing DBM concentration. The precise mechanism for this relationship remains to be determined and should be addressed in the clinical setting, given that a wide variety of DBM products with different DBM to carrier ratios are available.

With regard to the efficacy of different DBM products, Peterson et al recently published a study comparing multiple commercially available DBM products using an animal spinal fusion model with three different DBM preparations. They showed significant differences in the fusion rate and osteoinductive activity seen on histological sections. The animal model utilized immunoincompetent, athymic mice, which unfortunately precluded the evaluation of any effects of immune-provoked inflammation on osteoinductivity.

There were two weaknesses to our model of testing for the inflammatory reaction induced by bone graft substitutes. First, an animal model was employed. Any analogies to human biology must be made with caution. Species-specific reactivity to biomaterials should be considered. Specifically, the DBM products were derived from human tissue and theoretically could elicit a xenograft reaction not pertinent to human implantation. Second, this study was designed to elucidate the inflammatory reaction induced by these materials, and the cytokine evaluation was restricted to the measurement of IL-1β, the major mediator elicited in the murine inflammatory air pouch. No assertions were made relative to their efficacy. Therefore, one should use caution when extrapolating from the data presented here to determine the ideal bone graft substitute. Further studies are needed to discover the ideal formulation for a bone graft substitute. A follow-up to this study could include the examination of additional bone graft substitutes. Moreover, bone tissue can be implanted within the air pouch with bone graft substitute materials to quantify bone induction and production. Ultimately, randomized, prospective clinical trials comparing these products should be undertaken. However, studies such as ours help shed some light on the complex biology of bone graft substitutes.

### Table 1 Summary of data from histological analyses and reverse transcription-polymerase chain reaction analysis

| Bone graft substitute | Membrane thickness (mean ± SE, μm) | Pouch density (mean ± SE, cells/mm²) | Macrophages/monocytes: fibroblasts | Interleukin-1 expression (mean ± SE, copies) |
|-----------------------|-----------------------------------|-----------------------------------|----------------------------------|----------------------------------|
| Saline                | 47.7 ± 2.4                        | 1.0 ± 0.2                         | 1:4                              | 50,609 ± 24,083                  |
| ProOsteon®           | 166.7 ± 7.2                       | 30.7 ± 1.2                        | 1:3.3                            | 113,930 ± 23,715                 |
| DBX®                 | 202.3 ± 8.7                       | 63.1 ± 1.0                        | 1:4                              | 167,203 ± 50,079                 |
| Accell Connexus®     | 124.8 ± 6.5                       | 96.0 ± 1.6                        | 1:1.3                            | 146,115 ± 44,487                 |
| Acell DBM-100™       | 134.4 ± 6.3                       | 107.7 ± 2.2                       | 1:2.3                            | 377,137 ± 83,669                 |

**Abbreviation:** SE, standard error.

**Conclusion**

By both histologic and biochemical markers, our results indicate that all the products investigated provoked a significant inflammatory reaction when compared with the saline control. DBM products showed more inflammation than coralline
hydroxyapatite. Within the DBM groups, there was a trend toward increasing inflammation with increasing DBM concentration. Therefore, our preliminary findings suggest that DBM products with varied DBM proportions may influence the level of biocompatibility during clinical use.

**Disclosure**

The authors report no conflicts of interest in this work.

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