A single center retrospective clinical evaluation of anterior cervical discectomy and fusion comparing allograft spacers to silicon nitride cages

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Background: Iliac crest autograft or allograft spacers have been traditionally utilized in anterior cervical discectomy and fusion (ACDF) to provide vertebral stabilization and enhanced osteogenesis. However, abiotic cages have largely replaced these allogenic sources due to host-site morbidities and disease transmission risks, respectively. Although devices made of polyetheretherketone (PEEK) or titanium-alloys (Ti) have gained wide popularity, they lack osteoinductive or conductive capabilities. In contrast, silicon nitride (Si₃N₄) is a relatively new implant material that also provides structural stability and yet purportedly offers osteopromotive and antimicrobial behavior. This study compared radiographic outcomes at ≥12 months of follow-up for osseous integration, fusion rate, time to fusion, and subsidence in ACDF patients with differing intervertebral spacers.

Methods: Fifty-eight ACDF patients (108 segments) implanted with Si₃N₄ cages were compared to thirty-four similar ACDF patients (61 segments) implanted with fibular allograft spacers. Lateral radiographs (normal, flexion, and extension) were obtained at 3, 6, 12, and 24 months to assess osseous integration, the presence of bridging bone, the absence of peri-implant radiolucencies, subsidence, and fusion using both interspinous distance (ISD) and Cobb angle methods.

Results: In patients with ≥12 months of follow-up, fusion for the allograft spacers and Si₃N₄ cages was 86.84% and 96.83%, respectively (ISD method, P=0.10), and 67.65% and 84.13%, respectively (Cobb angle method P=0.07), while osseointegration was 76.32% and 93.65%, respectively (P=0.02). The time to fusion significantly favored the Si₃N₄ cages (4.08 vs. 8.64 months (ISD method, P=0.01), and 6.76 vs. 11.74 months (Cobb angle method, P=0.04). The assessed time for full osseointegration was 7.83 and 19.24 months for Si₃N₄ and allograft, respectively (P=0.00). Average subsidence at 1-year follow-up was 0.51 and 2.71 mm for the Si₃N₄ and allograft cohorts, respectively (P=0.00).

Conclusions: In comparison to fibular allograft spacers, Si₃N₄ cages showed earlier osseointegration and fusion, higher fusion rates, and less subsidence.

Keywords: Silicon nitride (Si₃N₄); allograft; clinical trial; anterior cervical discectomy and fusion (ACDF); osseointegration

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Introduction

Anterior cervical discectomy and fusion (ACDF) is a common corrective procedure for various degenerative spinal disorders including stenosis, disc herniation, spondylolysis, osteophyte formation, and spondylolisthesis (1,2). Autologous bone harvested from the iliac crest has been traditionally utilized as a replacement spacer within the operative cervical segment (3). While the use of autograft is considered to be the “gold standard” in ACDF (1,4), it has also been associated with a long-history of chronic harvest-site maladies including pain, seromas, hematomas, infections, and fractures (5). Persistent patient complaints about these comorbidities have led practitioners to substitute allograft spacers or abiotic biomaterial cages (e.g., Ti-alloys, PEEK) for autograft (6). Indeed, there has been a remarkable shift from the use of autogenous and allograft spacers to abiotic cages over the past 20 years. In 1998, 86% of all ACDF cases used autograft. Allograft comprised just 14% of total implantations, whereas cage use was still in its infancy. By 2008, autograft had declined to 10%, while allograft and interbody cages increased to 59% and 31%, respectively (7). Today, interbody cages dominate with ~60% of the devices produced from PEEK and 10% from Ti-alloys, with the balance being primarily a mixture of autograft and allograft (8).

While there have been numerous clinical studies comparing the safety and efficacy of autograft, allograft, and abiotic cages through the years, there is still a paucity of strong evidence supporting any one material. Most studies collectively suggest the equivalence of autograft and allograft, but generally favor autograft because of earlier arthrodesis (9-12). Clinical evaluations or reviews have compared autograft to Ti-cages (13-16), autograft to PEEK-cages (17), allograft to Ti-cages (18), allograft to PEEK-cages (19), Ti- to PEEK-cages (20-22), or multiple interbody materials (1,23,24) and have found little or no differences in reported outcomes, fusion rates, or subsidence. While these studies generally conclude that iliac crest grafts are still the “gold standard” due to earlier and more robust fusions, they concede that abiotic cages containing autologous bone are acceptable lower-morbidity alternatives. This conclusion is not particularly surprising given the greater osteogenic character of autograft compared with denatured allograft or the abiological cages. In fact, the initial requirements for allograft and the abiotic cages excluded biological activity. They were accepted solely based on sterility, biocompatibility, and mechanical properties (25-27). It has only been during the past 5 years that alterations of these devices (e.g., composites and coatings) have been engineered to upregulate their osteogenic capabilities (28,29).

Silicon nitride (Si₃N₄) is a relatively new spinal arthrodesis cage that has favorable osteopromotive and bacteriostatic properties. It was selected for this comparative study because its characteristics appear ideal for an interbody device including biocompatibility (30-36), enhanced strength and fracture toughness (37,38), moderate radiolucrency (39,40), osteoconductivity (41-48), and bacterial resistance (41,47,49-51). Si₃N₄ was first successfully used as an intervertebral device in a small clinical trial in Australia in 1986 (52), and a number of recent case and clinical studies have reported its favorable interbody characteristics (53-57).

The purpose of this study was to retrospectively assess ACDF outcomes for 58 patients (108 segments) who received a Si₃N₄ cage compared to 34 patients (61 segments) who were implanted with fibular allograft spacers all by the same surgeon (MWS). Fusion was assessed using lateral radiographs (normal, flexion, and extension) at 3, 6, 12, and 24 months post-operatively using ISD and Cobb angle methods. Assessments were also made for the extent of osseous integration, the presence of bridging bone, an absence of peri-implant radiolucrencies, and subsidence. The null hypotheses assumed that there would be no significant differences for any of these outcomes.

Methods

This study was designed as a retrospective chart review of patients on whom a single fellowship-trained spine surgeon (MWS) performed ACDF with either a Si₃N₄ cage (Valeo™C-II, Amedica Corporation, Salt Lake City, UT) or a structural fibular allograft spacer between August 1, 2013 and December 31, 2016. Inclusion and exclusion criteria are listed in Table 1. The medical procedures reported herein were consistent with the standard-of-care for these types of cases and did not involve more than minimal risk to the patients. Institutional review board (IRB) approval was obtained; but due to the retrospective nature of the study, informed consent by the IRB was not required. Nevertheless, all patient information and data remain completely anonymous and in compliance with IRB standards.
Surgical procedure

After radiographically localizing the affected level, a standard Smith-Robinson approach was made on the left-side (58). Complete discectomy, release of the posterior longitudinal ligament (PLL), and decompression of the uncinate process were then performed. A high-speed burr was used to prepare the endplates in a box-like fashion without violating their structural integrity. For all surgeries, burr shavings were harvested from local bone. The resulting autograft and DBM putty were pressed into either the endplates of a fibular allograft spacer or the center of a cage. After sizing, the bone spacers and Si$_3$N$_4$ cages, all lordotic, were placed at the appropriate levels. A standard anterior plate and screw construct was introduced for stabilization. Post-operatively, patients were mobilized as soon as possible. No cervical orthoses were used following surgery. The patients were instructed to restrict lifting to less than ~4.5 kg (<10 lbs.) during the first 6 weeks and no more than ~11.3 kg (25 lbs.) between weeks 6 and 12, as well as to avoid repetitive bending or twisting of the neck for at least 3 months.

Radiological assessment

Lateral radiographs were acquired at the following nominal post-operative intervals (within a margin of error): 2 weeks (<4 weeks); 6 weeks (≥4 and <8 weeks); 3 months (≥8 and <18 weeks); 6 months (≥18 and <40 weeks); 12 months (≥40 and <78 weeks); and 24 months (≥78 weeks) to assess fusion via visible osseous integration—defined as the presence of bridging bone and the absence of peri-implant radiolucency (55). These lateral radiographs were also used to evaluate subsidence—defined as the difference in segmental height (i.e., fused vertebrae plus disc space) between 2-week postoperative and subsequent follow-up radiographs (59). Additionally, flexion and extension radiographs at 3, 6, 12, and 24 months were used to assess fusion via ISD and Cobb angle methods (60,61). In the former method, nonunion at a given time-point was defined as a difference in ISD (i.e., between the spinous process tips of adjacent, fused vertebrae) of >2 mm between the patient’s flexion and extension radiographs, indicating excessive sagittal motion. Nonunion was similarly defined in the latter method as a difference in Cobb angle (i.e., between the line perpendicular to the top surface of the upper fused vertebral body and the line perpendicular to the bottom surface of the lower fused vertebral body) of >2° between the patient’s flexion and extension radiographs. Examples of these methods, applied to a Si$_3$N$_4$ cage, are provided in Figure 1. All radiographs were subsequently evaluated for fusion by the senior author and independently checked by the second author.

Outcome measures

The time to fusion using the ISD and Cobb methods was selected as the primary outcome measure. Secondary measures included fusion rate, osseous integration, and subsidence.

Statistical analyses

The Wilcoxon rank-sum test for ordinal data and Fisher's exact test for categorical data were used for statistical
analyses with $\alpha=0.05$. All statistical analyses were run using commercially available statistical software (Minitab 18, Minitab Inc., State College, PA, USA).

### Results

Between August of 2013 and December of 2016, fifty-eight patients underwent ACDF with $\text{Si}_3\text{N}_4$ cages (108 total levels) and thirty-four patients (61 total levels) underwent ACDF with allograft spacers. A patient accountability flowchart is given in Figure 2.

### Demographics and patient characteristics

Demographics, comorbidities, and operative details are presented in Table 2. It should be noted that not all patients appeared for all of the follow-up periods; therefore, each evaluation time-point represents a slightly different patient population (cf., Figure 2). Data on a total of 52 patients and 101 operative levels were available at $\geq12$-month follow-up (i.e., $31\text{ Si}_3\text{N}_4$ patients, 63 levels; and $21\text{ allograft patients,}$ 38 levels). There were no significant differences in patient demographics, comorbidities, or operative details between the two cohorts with the possible exception of a higher proportion of females in the $\text{Si}_3\text{N}_4$ group at $\geq12$-month follow-up (cf., Table 2). The majority of patients in both cohorts were overweight or obese with a primary diagnosis of radiculopathy or a combination of radiculopathy and myelopathy. About 50% were smokers and a significant percentage was also diabetic (i.e., between $\sim22\%$ and $41\%$).

### Complications and reoperations

Complication rates for the two cohorts are provided in Table 3. The surgical procedure and hospital stay were uneventful for most of the patients. Seven patients from the $\text{Si}_3\text{N}_4$ group and four patients from the allograft group suffered from recurrent dysphagia or dysphonia for more than six weeks. All were resolved within three months of follow-up using Medrol dospaks (Pfizer, NY, USA). Five $\text{Si}_3\text{N}_4$ patients and one allograft patient underwent subsequent ACDF operations for adjacent level disorders (i.e., stenosis with radiculopathy) that were not symptomatic at the time of their index procedures. Additionally, a recurrence of symptoms (e.g., persistent pain or paresthesia) was the sole reason for three revision surgeries in the $\text{Si}_3\text{N}_4$ cohort, whereas nonunion (five patients) and a post-operative infection (one patient) were reasons for revisions in the allograft cohort. In the latter case, Staphylococcus aureus was identified as the causative pathogen. All revisions resulted in solid arthrodesis and resolution of pain.

### Clinical outcomes

Primary and secondary outcomes are presented in Table 4 and in Figures 3-6. For the primary measure of time to fusion, both the ISD and Cobb angle methods demonstrated that the $\text{Si}_3\text{N}_4$ cages provided earlier arthrodesis, averaging $\sim4$ and $\sim7$ months, respectively. This contrasted with the fusion assessments for the allograft group of $\sim9$ and $\sim12$ months, respectively. The difference in time to fusion between the two cohorts for both measurement methods was statistically significant ($P=0.01$ and $0.04$, cf., Table 4). All of the secondary outcomes also favored $\text{Si}_3\text{N}_4$ over allograft. At $\geq12$-month follow-up, $\sim97\%$ and $\sim87\%$ of levels were fused in the $\text{Si}_3\text{N}_4$ and allograft groups using the ISD method, respectively ($P=0.10$, cf., Table 4). This compares to $\sim84\%$ and $\sim68\%$ using the
Table 2 Demographics, comorbidities, and operative details

<table>
<thead>
<tr>
<th>Patient characteristics</th>
<th>All patients</th>
<th>P value</th>
<th>Patients at ≥1 year follow-up</th>
<th>P value</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Si$_3$N$_4$</td>
<td>Allograft</td>
<td></td>
<td>Si$_3$N$_4$</td>
</tr>
<tr>
<td>Demographics</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Average age (years)</td>
<td>52.1</td>
<td>50.2</td>
<td>0.29</td>
<td>54.3</td>
</tr>
<tr>
<td>% Female</td>
<td>58.6</td>
<td>44.1</td>
<td>0.20</td>
<td>67.7</td>
</tr>
<tr>
<td>Average BMI (kg/m$^2$)</td>
<td>31.5</td>
<td>30.7</td>
<td>0.45</td>
<td>31.0</td>
</tr>
<tr>
<td>% Underweight</td>
<td>1.72</td>
<td>0.00</td>
<td>1.00</td>
<td>3.23</td>
</tr>
<tr>
<td>% Normal weight</td>
<td>15.51</td>
<td>23.53</td>
<td>0.41</td>
<td>16.13</td>
</tr>
<tr>
<td>% Overweight</td>
<td>29.31</td>
<td>23.53</td>
<td>0.63</td>
<td>25.81</td>
</tr>
<tr>
<td>% Obese</td>
<td>53.45</td>
<td>52.94</td>
<td>1.00</td>
<td>54.84</td>
</tr>
<tr>
<td>% Radiculopathy</td>
<td>82.76</td>
<td>67.65</td>
<td>0.12</td>
<td>77.42</td>
</tr>
<tr>
<td>% Myelopathy</td>
<td>0.00</td>
<td>2.94</td>
<td>0.37</td>
<td>0.00</td>
</tr>
<tr>
<td>% Radiculopathy + myelopathy</td>
<td>17.24</td>
<td>29.41</td>
<td>0.20</td>
<td>22.58</td>
</tr>
<tr>
<td>Comorbidities</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>% Tobacco use</td>
<td>44.83</td>
<td>47.06</td>
<td>1.00</td>
<td>51.61</td>
</tr>
<tr>
<td>% Diabetic</td>
<td>22.41</td>
<td>41.18</td>
<td>0.06</td>
<td>32.26</td>
</tr>
<tr>
<td>% Osteoporotic</td>
<td>1.72</td>
<td>0.00</td>
<td>1.00</td>
<td>3.23</td>
</tr>
<tr>
<td>% Osteopenic</td>
<td>1.72</td>
<td>2.94</td>
<td>1.00</td>
<td>0.00</td>
</tr>
<tr>
<td>Operative details</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>% 1 level</td>
<td>36.201</td>
<td>38.24</td>
<td>1.00</td>
<td>29.03</td>
</tr>
<tr>
<td>% 2 levels</td>
<td>43.10</td>
<td>44.11</td>
<td>1.00</td>
<td>41.94</td>
</tr>
<tr>
<td>% 3 levels</td>
<td>18.97</td>
<td>17.65</td>
<td>1.00</td>
<td>25.81</td>
</tr>
<tr>
<td>% 4 levels</td>
<td>1.72</td>
<td>0.00</td>
<td>1.00</td>
<td>3.23</td>
</tr>
<tr>
<td>% C1/C2</td>
<td>0.00</td>
<td>0.00</td>
<td>1.00</td>
<td>0.00</td>
</tr>
<tr>
<td>% C2/C3</td>
<td>0.00</td>
<td>0.00</td>
<td>1.00</td>
<td>0.00</td>
</tr>
<tr>
<td>% C3/C4</td>
<td>8.33</td>
<td>3.28</td>
<td>0.33</td>
<td>9.52</td>
</tr>
<tr>
<td>% C4/C5</td>
<td>18.52</td>
<td>19.67</td>
<td>0.84</td>
<td>19.05</td>
</tr>
<tr>
<td>% C5/C6</td>
<td>41.67</td>
<td>44.26</td>
<td>0.75</td>
<td>39.68</td>
</tr>
<tr>
<td>% C6/C7</td>
<td>31.48</td>
<td>32.79</td>
<td>0.87</td>
<td>31.75</td>
</tr>
<tr>
<td>% C7/T1</td>
<td>0.00</td>
<td>0.00</td>
<td>1.00</td>
<td>0.00</td>
</tr>
</tbody>
</table>

BMI, body mass index.

Cobb angle method, respectively (P=0.07, cf., Table 4). The percent of osseous integration for the Si$_3$N$_4$ group was ~94% compared to ~76% for allograft (P=0.02, cf., Table 4) and the time for osseous integration was ~8 months for the Si$_3$N$_4$ cohort compared to ~19 months for the allograft group (P=0.00, cf., Table 4). Average subsidence after at least 1-year follow-up was 0.51 and 2.71 mm for the Si$_3$N$_4$ cages and allograft spacers, respectively, (P=0.00, cf., Table 4). The time-series results at the earlier follow-up periods are shown in Figures 3 through 6 for fusion rate using ISD and
Cobb angle methods, osseous integration, and subsidence, respectively. At each time-point, these data showed that the Si₃N₄ cohort had earlier and more effective fusion (cf., Figures 3, 4), greater osseous integration (cf., Figure 5), and lower subsidence (cf., Figure 6). With the exception of the Cobb angle measurements for fusion rate, all of the data for fusion rate and % osseous integration at 3, 6, and 12 months were significant. The subsidence results also favored the Si₃N₄ group at every follow-up time-point (cf., Figure 6). At 24-month follow-up, 100% of the patients in the Si₃N₄ group were fused compared to ~96% of the allograft group using the ISD method (P=1.00, cf., Figure 3), whereas for the Cobb method, ~95% and ~81% fusion results were observed, respectively (P=0.35, cf., Figure 4). A representative radiographic example of the progressive fusion of a single level ACDF using a Si₃N₄ cage is shown in Figure 7.

Table 3 Complication rate for the patient population

<table>
<thead>
<tr>
<th>Complication</th>
<th>Si₃N₄</th>
<th>Allograft</th>
<th>P value</th>
</tr>
</thead>
<tbody>
<tr>
<td>% Dysphagia/dysphonia</td>
<td>12.07</td>
<td>11.78</td>
<td>1.00</td>
</tr>
<tr>
<td>% Adjacent level surgery</td>
<td>8.62</td>
<td>2.94</td>
<td>0.41</td>
</tr>
<tr>
<td>% Recurrence</td>
<td>5.17</td>
<td>0.00</td>
<td>0.29</td>
</tr>
<tr>
<td>% Non-union</td>
<td>0.00</td>
<td>14.71</td>
<td>0.01</td>
</tr>
<tr>
<td>% Post-operative infection</td>
<td>0.00</td>
<td>2.94</td>
<td>0.37</td>
</tr>
<tr>
<td>% Revision surgery</td>
<td>5.17</td>
<td>17.65</td>
<td>0.07</td>
</tr>
</tbody>
</table>

Table 4 Primary and secondary outcome measures at ≥12 months follow-up

<table>
<thead>
<tr>
<th>Outcome</th>
<th>Si₃N₄</th>
<th>Allograft</th>
<th>P value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Time to fusion using ISD method (months)*</td>
<td>4.08</td>
<td>8.64</td>
<td>0.01</td>
</tr>
<tr>
<td>Time to fusion using Cobb angle method (months)*</td>
<td>6.76</td>
<td>11.74</td>
<td>0.04</td>
</tr>
<tr>
<td>% Fusion using ISD method</td>
<td>96.83</td>
<td>86.84</td>
<td>0.10</td>
</tr>
<tr>
<td>% Fusion using Cobb angle method**</td>
<td>84.13</td>
<td>67.65</td>
<td>0.07</td>
</tr>
<tr>
<td>% Osseous integration</td>
<td>93.65</td>
<td>76.32</td>
<td>0.02</td>
</tr>
<tr>
<td>Time to osseous integration (months)</td>
<td>7.83</td>
<td>19.24</td>
<td>0.00</td>
</tr>
<tr>
<td>Average subsidence (mm)**</td>
<td>0.51</td>
<td>2.71</td>
<td>0.00</td>
</tr>
</tbody>
</table>

*, of those levels that were fused by ≥12 months follow-up; **, could not assess four levels in the allograft group; ***, could not assess three levels in the allograft group. ISD, interspinous distance.

Figure 3 Time series comparison of % fusion between allograft spacers and Si₃N₄ cages using the ISD method. P values indicate significance at each time point. n-values are the number of operated levels. Error bars represent binomial standard deviations.

Figure 4 Time series comparison of % fusion between allograft spacers and Si₃N₄ cages using the Cobb angle method. P values indicate significance at each time point. n-values are the number of operated levels. Error bars represent binomial standard deviations.

Discussion

For nearly 60 years, ACDF has been the standard of care for patients presenting intractable radiculopathy or myelopathy due to cervical disc herniation, stenosis, spondylosis, and spondylolisthesis (62,63). Historically, autologous bone harvested from the patient’s iliac crest has been utilized to stabilize the operative segment and prevent
kyphotic collapse during the fusion process. However, pain associated with the donor site—sometimes lasting up to 12 months post-operatively—has been the major complaint and significant challenge to satisfactory patient outcomes (6,64). To circumvent this persistent morbidity, processed and sterilized cadaver bone began to be used in the 1980s after passage of the National Organ Transplantation Act (65); and thereafter, allograft use rapidly increased while autograft procedures commensurately declined (7). Reviews by Malloy and Hilibrand in 2002 (66) and by Tuchman et al. in 2017 (12) summarized the usage of these two allogenic sources for approximately the past 50 years. Overall, higher arthrodesis rates have been observed with autograft, particularly for earlier fusion and in multi-level procedures. However, for single-level ACDF, patient outcomes appear to be nearly equivalent, and both sources are considered to be effective arthrodesis materials.

Today, the use of both autograft and allograft has...
been largely supplanted by abiotic cages mostly made from either Ti-alloys or PEEK. Ti-alloys have been around since just after World War II and have been actively used as implants since the 1970’s (26). However, biomedical titanium is essentially bioinert because of a thin passivation layer of titanium dioxide (TiO$_2$) which prevents significant biochemical interactions. Because of its inert nature, surface functionalization of Ti-alloys has been conducted since the 1990s in an effort to enhance the material’s osteoconductivity (67-70). Biomedical PEEK was introduced in the 1990s and rapidly gained acceptance as an intervertebral cage because of its lower cost, favorable modulus, and ease of use (27). Its rise in popularity was accelerated because of subsidence concerns associated with Ti-alloys. However, subsequent studies have shown that the initial and long-term mechanical stability of a spinal implant may be more dependent upon its overall size and geometry rather than its hardness or elastic modulus (54,71,72). Unfortunately, PEEK does not integrate into adjacent host bone and is not visible on plain X-rays (73). In vivo, PEEK spacers heal by the formation of fibrous tissue. This observation has been referred to as the PEEK “halo effect” (74,75). In reality, the hydrophobic nature of PEEK discourages osseointegration by inhibiting protein absorption and cell adhesion on the implant’s surface (41,49,76). Clearly, autograft, allograft, Ti-alloys, and PEEK have their limitations. This has led a number of authors to conclude that the ideal spinal fusion material has yet to be determined (77-79).

In contrast, Si$_3$N$_4$ exhibits many of the important elements of an ideal interbody device. Its inherent chemistry, hydrophilicity, surface charge, and topography are likely responsible for the accelerated fusion observed in this study. Its underlying osteopromotive mechanism involves the release of minute amounts of silicic acid (H$_4$SiO$_4$) and ammonia (NH$_3$). These compounds are known to play active roles in the bone healing process (48,80-83). Eluted ammonia from the ceramic’s surface is also enzymatically converted to nitric oxide (NO) and peroxynitrite (ONOO$^{-}$) which eventually results in the lysis of adherent bacteria (50,84). Additionally, the anisotropic needle-like Si$_3$N$_4$ grains (<1.0 μm in cross-section and ≤10 μm in length) at the ceramic’s surface have also been shown to play important roles in promoting attachment of proteins and eukaryotic cells while resisting adhesion of prokaryotic cells (47,49,51). In summary, spinal implants produced from Si$_3$N$_4$ appear to have distinct advantages over allograft spacers and other abiotic materials. These benefits include enhanced osteoconductivity (41-48), earlier fusion (56), anti-infective behavior (41,47,49-51,57), and improved radiolucency (39,40).

There are a number of limitations associated with this study. Foremost among them is its retrospective nature. Although the data provide clear statistical evidence of the superiority of Si$_3$N$_4$ cages over allograft spacers, the results presented herein were not compiled using a prospective, randomized, and blinded protocol. The study is also limited from the perspective that the clinical work was conducted only by one surgeon at a single institution.

Conclusions

This retrospective 92-patient ACDF clinical evaluation compared outcomes for traditional allograft spacers and Si$_3$N$_4$ cages over a 24-month follow-up period. The results demonstrated that both materials were effective in achieving acceptable levels of fusion and osseous integration at ≥12 months. However, earlier periods significantly favored the use of the Si$_3$N$_4$ cages. At 3-, 6-, and 12-month follow-ups, the Si$_3$N$_4$ cohort showed faster fusion by both the ISD and Cobb angle methods and greater osseous integration. The Si$_3$N$_4$ group also showed smaller amounts of subsidence at all follow-up periods. At a minimum, the results demonstrate that Si$_3$N$_4$ cages are at least as effective as traditional allograft in ACDF patient outcomes.

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Footnote

Conflicts of Interest: Dr. MW Smith is a consulting surgeon to Amedica Corporation. Drs. BJ McEntire and BS Bal are principals and employees of Amedica Corporation which was the manufacturer of the Si$_3$N$_4$ devices used in this study. Dr. DR Romano has no conflicts of interest to declare.
Ethical Statement: Institutional review board (IRB) approval was obtained; but due to the retrospective nature of the study, informed consent by the IRB was not required.

References


