Analyzing the behavior of a porous nano-hydroxyapatite/polyamide 66 (n-HA/PA66) composite for healing of bone defects

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Abstract: The aim of this study was to analyze the behavior of the porous nano-hydroxyapatite/polyamide 66 (n-HA/PA66) composite grafted for bone defect repair through a series of biological safety experiments, animal experiments, and a more than 5-year-long clinical follow-up. The biological safety experiments, carried out in accordance with the Chinese Guo Biao and Tolerancing (GB/T) 16886 and GB/T 16175, revealed that porous n-HA/PA66 composite had no cytotoxicity, no sensitization effect, no pyrogenic reaction, and that its hemolysis rate was 0.59% (less than 5%). Rabbit models of tibia defects with grafted porous n-HA/PA66 composite were established. After 2 weeks, the experiment showed that osteogenesis was detected in the porous n-HA/PA66 composite; the density of new bone formation was similar to the surrounding host bone at 12 weeks. After 26 weeks, the artificial bone rebuilt to lamellar bone completely. In the clinical study, a retrospective review was carried out for 21 patients who underwent serial radiographic assessment after treatment with porous n-HA/PA66 composite grafts following bone tumor resection. All wounds healed to grade A. No postoperative infections, delayed deep infection, nonspecific inflammation, rejection, or fractures were encountered. At a mean follow-up of 5.3 years, the mean Musculoskeletal Tumor Society’s (MSTS) 93 score was 29.3 points (range: 28–30 points) and mean radiopaque density ratio was 0.77±0.10. The radiologic analysis showed that porous n-HA/PA66 composite had been completely incorporated with the host bone about 1.5 years later. In conclusion, this study indicated that the porous n-HA/PA66 composite had biological safety, and good biocompatibility, osteoinduction, and osseointegrations. Thus, the porous n-HA/PA66 composite is an ideal artificial bone substitute and worthy of promotion in the field.

Keywords: biomaterial, artificial bone materials, porous n-HA/PA66 composite (n-HA/PA66), bone defects healing, bone grafting

Introduction

Bone is both an important tissue and a major organ of the human body. Among other things, bones are important for support, protection, loading, hematopoiesis, and calcium storage. Bone defects are quite common clinically. The treatment of large bone defects caused by trauma, infection, and tumor excision represents a great challenge in the clinical practice setting. Bone grafting—which includes autografts, allografts, and synthetic substitutes—is an important treatment for large bone defects. Successful incorporation of grafted material into bone defects depends on the material’s ability to promote new bone formation and provide a scaffold for osteogenesis. As is well known, autogenous bone grafting is the gold standard in the reconstruction of bone defects; it has the advantage of providing both bone stock and osteogenic precursor cells. Due to limited resources, the autograft technique presents difficulties in meeting the demands...
for large bone defects. Because of rejection and the spreading
of infection diseases, allograft use is restricted in clinical
practice. Hence, synthetic bone substitutes are being increas-
ingly used in bone reconstruction techniques. Searching for
the ideal artificial bone substitute for repairing bone defects
is therefore an important subject in orthopedics.

While the use of biological materials has long been stud-
ied, the use of artificial bone materials to repair bone defects emerged in the 1980s and has developed rapidly since. The
basic concept of artificial bone material is that it is used as a
bone tissue substitute, wherein it is incorporated into the bone
defect to reestablish bone clinically. Searching for the ideal
artificial bone material is presently a hot topic in orthopedics.

The properties of the ideal artificial bone materials include
biological safety, fine biocompatibility, good absorbability
and biodegradability, good porous structure (an average pore
diameter 200–400 microns), good mechanical properties,
osteogenesis, osteoinduction, and osteoconduction. Currently, the artificial biomaterials used as bone tissue
substitutes mainly include artificially-synthesized materials
and materials derived from natural biological organisms.
Amongst other materials, artificially-synthesized materi-
als contain calcium phosphate ceramic, calcium phosphate
cement (CPC), bioactive glass, polyamide (PA), polylactic
acid (PLA/PLLA), and polyglycolic acid (PGA). Materials
derived from natural biological organisms include mainly
nano-hydroxyapatite (n-HA), collagen, chitins, coral bone
(CAB), and natural bone. 

Because it is difficult to prepare an ideal bone tissue
substitute with a single material, many studies have being car-
rried out in recent years looking at developing bone substitute
composites for use in repairing bone defects. A novel bio-
medical composite, porous nano-hydroxyapatite/polyamide
66 (n-HA/PA66), has been developed in recent years. This composite has good similarity to human bone; that is,
n-HA has a similar composition and crystal structure to the
mineral of natural bone. n-HA, which has good biocompat-
ability and bone regeneration, can prompt injured bone to
reproduce. PA66, which is a kind of high molecular synthetic
material with sufficient resources, acts in the role of a col-
lagen matrix in the composite, avoiding the immunogenicity
problem of natural polymeric materials. The polar amide
bond, carboxyl group, and carboxyl group of PA66 can
grow, form osteoids, and mineralize tissue cell to promote
healing of bone defects. Moreover, this composite has good mechanical performance and can serve as a load-bearing
bone substitute; for example, its dense composite has been
extensively used in repairing vertebral plates and has proven
to have a good effect on bone reconstruction.

Materials and methods
Biological safety experiments
The porous n-HA/PA66 composite was prepared by n-HA
(molecular weight of 1,000; nanoparticle diameter of
80–100 nm) and PA66 (molecular weight of 18,000–20,000),
which were provided by the Sichuan National Nanotech-
ology Co, Ltd (Chengdu, People’s Republic of China).
The size of the porous n-HA/PA66 composite sample was
4 × 2 × 2 mm³. The average pore size was 500 microns,
with a porosity of 75%–85%, and a compressive strength
of 2–10 mPa (Figure 1). The porous n-HA/PA66 composite
was synthesized using chemical foaming techniques, as
previously described in other studies.

To evaluate the biological safety of the porous n-HA/PA66
composite, a cytotoxicity test, sensitivity test, pyrogen test, and
a hemolysis test were carried out according to the Chinese Guo
Biao and Tolerancing (GB/T)16886 and GB/T16175. The
extract liquid of the porous n-HA/PA66 composite was prepared

Figure 1 The morphology of porous n-HA/PA66 composite.
Notes: (A) A porosity of 75%–85% and an average pore size of 500 microns; (B) A size of 4 × 2 × 2 mm³.
Abbreviation: n-HA/PA66, nano-hydroxyapatite/polyamide 66.
in biological safety experiments. A mouse fibroblast L929 cell strain, which was inoculated into cell culture dish, was prepared for the cytotoxicity test. The experimental group was given porous n-HA/PA66 extract liquid and the control group received sterile physiological saline. All dishes were put into an incubator (37°C, 5% carbon dioxide [CO₂]) for 24 hours to observe the cytolytic ratio. In the sensitization test, the red punctuation and hydrosia of guinea pigs were observed at 24 hours, 48 hours, and 72 hours by subcutaneously injecting porous n-HA/PA66 extract liquid into the back of the guinea pigs. The control group was injected with sterile physiological saline. The extract liquid was injected via rabbit ear vein for the pyrogen test. The temperature changes were measured before and after the injection. The hemolysis test was conducted on anticoagulated fresh rabbit blood. The experimental group received extract liquid and the control group sterile physiological saline.

Animal implantation studies

In the animal implantation study, 16 New Zealand White rabbits (2.0–2.5 kg, randomly male and female, offered by Sichuan University, West China Experimental Animal Center) received tibia bone defects 4 mm in diameter. Performed as a self-contrast in paired design, the right rabbit tibia was grafted with porous n-HA/PA66 composite, while the left tibia (control) of the same rabbit was grafted with autogenous iliac bone. For both groups, we performed macroscopic observations and radiological, histological, and scanning electron microscope analysis at 2, 4, 12, and 26 weeks after surgery.

Clinical studies

From November 2007 to July 2008, there were 21 consecutive patients who fulfilled the following inclusion criteria: 1) benign bone tumor confirmed by pathology; 2) their treatment consisted of one operation and single implantation of the porous n-HA/PA66 composite; 3) there was complete follow-up data; and 4) the patient showed no recurrence. We excluded a woman who suffered from giant cell tumor of the right proximal tibia, where the patient showed no recurrence. We excluded a woman who suffered from giant cell tumor of the right proximal tibia, where the patient showed no recurrence. We excluded a woman who suffered from giant cell tumor of the right proximal tibia, where the patient showed no recurrence. We excluded a woman who suffered from giant cell tumor of the right proximal tibia, where the patient showed no recurrence. We excluded a woman who suffered from giant cell tumor of the right proximal tibia, where the patient showed no recurrence. We excluded a woman who suffered from giant cell tumor of the right proximal tibia, where the patient showed no recurrence. We excluded a woman who suffered from giant cell tumor of the right proximal tibia, where the patient showed no recurrence. We excluded a woman who suffered from giant cell tumor of the right proximal tibia, where the patient showed no recurrence. We excluded a woman who suffered from giant cell tumor of the right proximal tibia, where the patient showed no recurrence. We excluded a woman who suffered from giant cell tumor of the right proximal tibia, where the patient showed no recurrence. We excluded a woman who suffered from giant cell tumor of the right proximal tibia, where the patient showed no recurrence. We excluded a woman who suffered from giant cell tumor of the right proximal tibia, where the patient showed no recurrence.

The clinical studies group included nine male and twelve female patients. Their age at the time of surgery was 18–56 years, with a mean age of 33 years. Eight lesions were located in the diaphysis, twelve in the metaphysis of a long bone, and only one in a short bone. Histological examinations revealed that nine were giant cell tumor, six were bone cysts, and six were fibrous dysplasia. The size of the lesions ranged from 4.0 × 3.0 × 3.0 cm³ to 17.0 × 3.0 × 3.0 cm³, and no case suffered from pathological fracture. All cases underwent curettage, were inactivated, and then the tumor cavities filled with porous n-HA/PA66 composite. While four cases received internal fixation implants, six were implanted with both variant bone plates and internal fixation. The implant amount of porous n-HA/PA66 composite ranged from 10–30 g, with an average of 20 g. One, 3, 6, 9, and 12-month follow-up was observed after treatment, and after we recommended an inspection be carried out each year (Table 1).

The clinical evaluation included objective findings focusing on the operative incision’s healing, blood tests, biochemical assays, and subjective complaints measured with the Musculoskeletal Tumor Society’s (MSTS) 93 limb functional recovery score system (upper limb: pain, adjacent joint function, emotional acceptance, supports, positioning of hand, lifting ability; lower limb: pain, adjacent joint function, emotional acceptance, supports, walking ability, gait; each 0–5 points, total of 30 points). Using standard projections and evaluated independently, the radiographs were analyzed by three investigators, and the data were checked for interobserver agreement. In the event of differences, the patient’s radiograph was reevaluated by all three observers together. The radiographic integration of porous n-HA/PA66 composite was assessed according to the Lane and Sandhu radiological criteria system, to determine the presence of the distinct bridging trabeculae at the recipient bone–implant interface and also the amount of incorporation of porous n-HA/PA66 composite. A software program (Syngo version V35; Siemens Medical Systems, Erlangen, Germany) for quantitative analysis was used to measure the X-ray radiopaque density value. An area of interest was determined over the grafted section (see Figure 2) with number of counts recorded. An identical area of interest was determined over a non-lesion bone area near the operated section, and the grafting-to-non-lesion count ratio (G/N ratio) was then calculated (see Figure 2). The normal value of the G/N ratio was 1.00. After curettage of the bone tumor and implantation of the porous n-HA/PA66 composites, patients were followed-up for between 5–5.5 years (5.3 years on average) by panoramic X-ray and computed tomography (CT) plain scan, and spiral CT 3D reconstruction.

For statistical analysis, SPSS 17.0 (IBM Corporation, Armonk, NY, USA) software was used. The values are presented as mean ± standard deviation (SD). One-way analysis of variance (ANOVA) was used to determine differences between the radiopaque density ratios for dependent variables measured at the normal level (sex, diagnosis, location, bone grafted mass). The correlation was analyzed, with linear correlation used to determine the relationship between radiopaque
density ratios and the age of patients at the time of surgery. A *P*-value of less than 0.05 was considered significant.

**Results**

**Biological safety experiments**

For the cytotoxicity test, the cytolytic ratio of the experimental group was found to be 0 (R=0). There was no significant difference between the experimental group and the control group. In the sensitization test, the erythema and edema failed to be found in the guinea pigs injected by leaching liquor and saline. In the pyrogen test, the temperatures of experimental rabbits rose by 0.6°C, which met the requirements of the pyrogen test. In the hemolysis test, the results showed that the hemolysis rate was 0.59% (less than 5%). Therefore, the porous n-HA/PA66 composite had no cytotoxicity, no sensitization effect, no pyrogenic reactions, and the hemolysis rate was 0.59% (less than 5%), which met biological safety standards according to the Chinese GB/T16886 and GB/T16175.

**Animal implantation studies**

Macroscopic observation showed that there was no postoperative infection or material displacement. Histological assessment indicated that osseous tissue gradually crept into the interconnected porosity and connected with the material directly, without fibrous tissue. The amount of new bone and calcium crystals were increased at 4–8 weeks, and the new bones continuously matured. After 12 weeks, new bone formed in the pores, gradually linked with each other, and connected with materials directly. After 26 weeks, the interconnected porosity was completely filled with new bone tissue, integrated into a whole, and showed good osteoconductivity and osteocompatibility. Marrow cavity was also generated at that time (Figures 3 and 4). Panoramic X-ray analysis showed that by 26 weeks the test side had been repaired completely, the density of which was equivalent to the normal control.

**Table 1** Details of patients after implantation of the porous n-HA/PA66 composite

<table>
<thead>
<tr>
<th>Number</th>
<th>Age</th>
<th>Sex</th>
<th>Diagnosis</th>
<th>Location</th>
<th>Bone grafting</th>
<th>Follow-up</th>
<th>MSTS</th>
<th>Radiographic finding</th>
<th>G/N ratio</th>
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<td>37</td>
<td>F</td>
<td>FD</td>
<td>Right proximal femur</td>
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<td>5.2 years</td>
<td>29</td>
<td>Complete incorporation</td>
<td>0.82</td>
</tr>
<tr>
<td>2</td>
<td>20</td>
<td>F</td>
<td>BC</td>
<td>Left heel bone</td>
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<td>5.5 years</td>
<td>30</td>
<td>Complete incorporation</td>
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<td>5.5 years</td>
<td>29</td>
<td>Complete incorporation</td>
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<tr>
<td>4</td>
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<td>M</td>
<td>GCT</td>
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<td>5.4 years</td>
<td>29</td>
<td>Complete incorporation</td>
<td>0.74</td>
</tr>
<tr>
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<td>F</td>
<td>FD</td>
<td>Right femur</td>
<td>25 g</td>
<td>5.1 years</td>
<td>30</td>
<td>Complete incorporation</td>
<td>0.82</td>
</tr>
<tr>
<td>6</td>
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<td>FD</td>
<td>Right proximal tibia</td>
<td>10 g</td>
<td>5 years</td>
<td>29</td>
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<td>0.75</td>
</tr>
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<td>7</td>
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<td>Complete incorporation</td>
<td>0.79</td>
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<td>Left proximal femur</td>
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<td>5.5 years</td>
<td>30</td>
<td>Complete incorporation</td>
<td>0.78</td>
</tr>
<tr>
<td>9</td>
<td>28</td>
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<td>Right proximal tibia</td>
<td>30 g</td>
<td>5.5 years</td>
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<tr>
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<td>Left proximal femur</td>
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<td>5.3 years</td>
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<td>Complete incorporation</td>
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</tr>
<tr>
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<td>Left distal femur</td>
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<td>Complete incorporation</td>
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<tr>
<td>16</td>
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<td>5.1 years</td>
<td>30</td>
<td>Complete incorporation</td>
<td>0.83</td>
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Abbreviations: n-HA/PA66, nano-hydroxyapatite/polyamide 66; F, female; M, male; FD, fibrous dysplasia of bone; G/N ratio, grafting-to-non-lesion ratio; BC, bone cyst; GCT, giant cell tumor; MSTS, Musculoskeletal Tumor Society score.

Figure 2 Determination of the grafting-to-non-lesion count ratio (G/N ratio).
Clinical studies

All cases were followed-up for 5–5.5 years postoperatively (average of 5.3 years). The results were satisfactory. There was nothing obviously exceptional in the C-reactive protein, routine blood investigations, biochemical tests, and hemagglutination inspections of all patients. All wounds healed to grade A. No postoperative infections, delayed deep infection, nonspecific inflammation, rejection, or fractures were encountered. Limb function was good and so was joint motion near the tumor spots. In addition, no patient complained of pain. The internal steel plates were still rigidly fixed and no joint surface subsidence occurred. The mean MSTS score was 29.3 points (range: 28–30 points).

Radiographs which were obtained immediately after surgery revealed radiolucent zones between the implanted porous n-HA/PA66 composite and the surrounding bone. Periodic assessments showed that 25% of grafting zones were replaced by new bones in 2–3 months; after 6–9 months, new bone formation rose to 50%. The defects were fully occupied by new bone within 1.5 years (Figures 5 and 6). Radiographic observations analyzed the grafting and non-lesion areas for differences. Mean radiopaque density ratio was 0.77±0.10 after grafting with porous n-HA/PA66. This was most obvious in cancellous bone such as the heel bone (G/N = 0.41). No degenerative changes were observed in adjacent joints. After implantation with porous n-HA/PA66, the variables were evaluated; there was no relationship between age, sex, diagnosis, bone graft mass, and radiopaque density ratio (Table 2).

We documented a significant difference in the radiopaque density between metaphyseal and diaphyseal location of grafting (F=9.65, P=0.01). Mean radiopaque density ratio was 0.67±0.10 after implantation with porous n-HA/PA66 in the metaphyseal region, and 0.85±0.03 in the diaphyseal region.

As shown in Table 1, the G/N ratio was <1. Because new bone growth in the grafting zone blocked the X-ray transmission, as time went on, the more new bones grew, the smaller the G/N value, and this value became stationary at about 1.5 years. Due to new bone tissue growing more numerous and faster in the metaphysis, the G/N ratio curve was lower and steeper than the curve for diaphysis (Figure 7).

The excluded female patient previously mentioned suffered from a giant cell tumor of the right proximal tibia and the tumor had shown recurrence after follow-up for 2 years. She underwent a second operation and the resected tumor tissues were sent for pathology inspection. The result showed that the pores of porous n-HA/PA66 composite were filled with new bone and tumor tissues which integrated into a whole (Figure 8).

Discussion

Biological safety

Biomaterials are ones which can be implanted into the body and joined with tissues and organs together. Biological safety
Biocompatibility is one of the most important factors when dealing with biomedical materials. Any new biomedical materials in the early stages of clinical application must be tested according to ISO standards or the appropriate standards of the jurisdiction where it is being used. Laboratory and animal experiments are the main methods for evaluating the biological safety of biomedical materials. In our studies, biological safety experiments showed that the porous n-HA/PA66 composite had no cytotoxicity, no sensitization effect, and no pyrogenic reactions. Further, the hemolysis rate was 0.59%, which met biological safety standards according to the Chinese GB/T16886 and GB/T16175. In animal implantation studies, there was no suppuration or necrosis after grafting the porous n-HA/PA66, and new bone formation rebuilt to lamellar bone completely after 26 weeks. In clinical studies, all wounds healed to grade A. All patients healed well, experiencing no infection, aseptic inflammation, or rejection reactions. None of the patients had serious side effects such as functional damage of the liver and kidney. Therefore, it is evident that the treatment of large bone defects by grafting porous n-HA/PA66 composite is secure.

Figure 5 A 20-year-old man suffering from fibrous dysplasia of the right proximal femur. 
Notes: (A) The original image of the defect site. Periodic radiological assessments showed (B) during the period of 6 months, 50% of the grafting zones were replaced by new bones, and (C) fully occupied at 1.5 years after surgery. (D) With a more than 5 years follow-up, the porous n-HA/PA66 composite had been completely incorporated with the host bone. (E and F) grafting-to-non-lesion count ratio was 0.83, the patient was satisfied with the result, and the MSTS was 30. 
Abbreviations: n-HA/PA66, nano-hydroxyapatite/polyamide 66; MSTS, Musculoskeletal Tumor Society score.

Figure 6 Images of a 47-year-old woman treated by a giant cell tumor resection and fixed with an internal steel plate after the porous n-HA/PA66 composite grafted in the right distal femur. 
Notes: Periodic radiological assessments showed that the new bones developed and the porous n-HA/PA66 composite incorporated with the host bone gradually. The internal steel plate was still rigidly fixed and no joint surface subsidence occurred. (A) Preoperative; (B) a half-year after resection; (C) 1 year; and (D) 5 years later. The grafting-to-non-lesion count ratio was 0.78 and the MSTS score was 29 after 5 years follow-up (E and F). 
Abbreviations: n-HA/PA66, nano-hydroxyapatite/polyamide 66; MSTS, Musculoskeletal Tumor Society score.
Porous nano-hydroxyapatite/polyamide 66 for healing bone

The uniform distribution of n-HA granules in the PA66 matrix enable the scaffolding material to possess good biocompatibility, high bioactivity, and enough mechanical strength. Previous studies have confirmed that an n-HA/PA66 composite with an n-HA:PA66 ratio of 6:4 had more biocompatibility than that of other ratios. An appropriate pore diameter and porosity make a biomaterial more biocompatible. The current research shows that pores of 100–500 microns are suitable for bone tissue growth. Hulbert et al reported 100 microns was the minimum pore diameter to allow bone tissue growth, and 150 microns the ideal. Vaz et al showed that 500 microns was the most appropriate for bone growth. In fact, the human bone is a relatively loose complex structure. The structure of bone consists of dense tissue at the outermost layer, changing gradually through semi-dense to a loose mesh structure in the innermost layer. Therefore, when the porous n-HA/PA66 composite was grafted, the pores of the host bone and the grafted material gradually integrated into the two cross-linked form, which would be able to maintain normal physiological metabolism in the healing of the bone’s defects. In our clinical studies, a significant difference was documented in the radiopaque density between the metaphysis and diaphysis of porous n-HA/PA66 composite filling ($F=9.65, P=0.01$). The pores of metaphyseal bone and materials were better integrated because of the loose mesh structure of the metaphysis. This mesh structure increased the localized supply of nutrition and blood, and as a result new bones grew more numerously and faster; the G/N ratio curve of the metaphysis was therefore lower and steeper (Figure 6). Additionally, porosity is another important indicator of a porous mesh structure. The mutual penetration and the size of the aperture are the most important factors in bone regeneration. The higher porosity materials can provide a better channel structure. Jie and Yubao showed that when porosity of the porous n-HA/PA66 composite was more than 50%, there was a positive effect on cell adherence and there were extracellular matrix deposits on the surface. Higher porosity is also advantageous to the transmission of oxygen and nutrients and to the growth of blood vessels and nerves. Thus, good conditions were provided for new bone formation. However, the increase in pore

<table>
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<tr>
<th>Variable</th>
<th>Total (N=21)</th>
<th>G/N ratio</th>
<th>F</th>
<th>P</th>
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<td>0.36</td>
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<td>0.80±0.06</td>
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<td></td>
</tr>
<tr>
<td>Diagnosisb</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>GCT</td>
<td>9</td>
<td>0.80±0.07</td>
<td></td>
<td></td>
</tr>
<tr>
<td>FD</td>
<td>6</td>
<td>0.77±0.08</td>
<td></td>
<td></td>
</tr>
<tr>
<td>BC</td>
<td>6</td>
<td>0.73±0.16</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Locationb</td>
<td></td>
<td>9.65</td>
<td>0.01</td>
<td></td>
</tr>
<tr>
<td>Metaphysis</td>
<td>12</td>
<td>0.67±0.10</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Diaphysis</td>
<td>8</td>
<td>0.85±0.03</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Bone grafted massb</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>≤20 g</td>
<td>13</td>
<td>0.73±0.16</td>
<td></td>
<td></td>
</tr>
<tr>
<td>&gt;20 g</td>
<td>8</td>
<td>0.78±0.07</td>
<td></td>
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</tr>
</tbody>
</table>

Notes: *Correlation between the two variables was analyzed with linear correlation; *one-way analysis of variance (ANOVA).
Abbreviations: G/N ratio, grafting-to-non-lesion ratio; FD, fibrous dysplasia of bone; BC, bone cyst; GCT, giant cell tumor.

![Figure 7](https://www.dovepress.com/)

Figure 7 Comparing the G/N ratio between the metaphyseal and diaphyseal areas where the porous n-HA/PA66 composites were implanted after bone tumor resection.
Abbreviations: G/N ratio, grafting-to-non-lesion ratio; n-HA/PA66, nano-hydroxyapatite/polyamide 66.
size and porosity decreases the material’s mechanical strength. Therefore, a compromise should be reached between material properties and mechanical strength to satisfy bone conduction requirements. In previous studies, porous n-HA/PA66 composite with an average pore diameter of 500 microns, porosity of 75%–85%, and a compression strength of 2–10 MPa (similar to human cancellous bone) satisfied the requirements to repair bone defects. 

**Osteoinduction and osseointegration**

The beginning of bone formation and bone cell proliferation is a very complex process. Some research has found that three-dimensional pore structures could facilitate cell adhesion, differentiation, and proliferation, and help with fibrovascular and nerve colonization, so that the new bone “crawls” into the material to achieve early osteoinduction. Moreover, the osteoinduction ability of the n-HA/PA66 composite may be associated with a “nanometer-sized effect”. The surface of a nano-material is significantly larger than a micron-material, and the number of atoms are significantly increased on the surface. Thereby, owing to their greater surface area, the nano-sized particles are more biologically active compared with micron-sized particles. The previous studies have shown that micron-HA only had osteoconduction without osteoinduction, yet nano-HA had both. Kim et al reported that nano-HA particles which were compounded into the polypropylene matrix have good biological activity, and that their osteoinduction was superior to ordinary HA particles. In earlier studies where porous n-HA/PA66 composite was implanted into the deep layer of the back muscles of rabbits, newly formed bone tissue was found in pores after 26 weeks. This proved that the porous n-HA/PA66 composite had good osteoinductivity.

Artificial bone substitutes for bone defect repair not only have good osteoinductivity, they can also integrate with host bone so that the implanted materials can play a good physiological function for a long time. The ideal bone substitute material can bond to osseous tissue tightly. Rosen et al have indicated that it was easy to cause loosening (leading to failure) if the fibrous tissue growth wrapped the interface of the bone substitutes after implantation. In our studies, new bone formed in the pores and connected with materials directly. Radiographic observations demonstrated that the grafting and non-lesion area combined as a whole. Owing to good osseointegration of the porous n-HA/PA66 composite, no case showed joint collapse or internal fixation loosening during the average 5.3 years of follow-up. The pathological results of the excluded female patient showed pores were filled with new bones and tumor tissues, which integrated with the porous n-HA/PA66 composite.

**Clinical application**

According to research into biomaterials, there are various artificial bone substitute materials, including biomedical metal materials, polymer materials, and some biological ceramic materials. The results of clinical applications of artificial bone substitute materials are mixed. Demineralized bone matrix (DBM) can impart more osteoinductive potential than standard mineralized allograft. However, variable clinical response is not for structural supports. Synthetic ceramics which are without infectious or immunogenic potential are known to be osteoconductive but lack intrinsic osteogenic potential, as is found with autograft. CPC remains inert for long periods with insubstantial resorption. However, the initial liquid state can lead to difficulties during placement, such as extrusions into intra-articular regions. Collagen/ceramic composites (eg, n-HA/PE, collagen) which have been found to be effective autograft extenders, have osteoconduction and osteoinduction properties. n-HA/PA66 has been found to have biological safety, good biocompatibility, and osteoinduction.
and osseointegration in in vitro and in vivo studies, and has been shown to be a prospective application in the clinical setting.\textsuperscript{33,34}

In our clinical studies, all cases underwent curettage, inactivation, and were filled with porous n-HA/PA66 composite. Solid fusion was obtained in the 6 months after the operation. Thus, the porous n-HA/PA66 composite can be effectively used for repairing bone defects caused by benign bone cysts and tumor-like lesions. Additionally, in clinical practice, the porous n-HA/PA66 composite was used to repair bone defects in limb bone fractures, aseptic necrosis of bone, arthrosis, and orthopedics of grafted bone, with good effects produced. According to our studies, the porous n-HA/PA66 composite implanted into both metaphysal and diaphysic defects and achieved positive results; MSTS scores also revealed favorable functional outcomes (mean: 29.3 points). However, porous n-HA/PA66 is more suitable for filling the diaphysal and short bone defects ($P=0.01$).

After the tumor is resected and the bone grafts adequately, different fixations are used according to the different nature of the lesion: 1) the porous n-HA/PA66 composite is grafted alone with no internal fixation when lesion involves the bone cross-sectional area $<50\%$; 2) the lesion involves a bone cross-sectional area $>50\%$, or the diameter of cortical bone defect is $<50\%$ with risk of pathologic fracture, so a steel plate is internally fixed after the porous n-HA/PA66 composite is grafted; and 3) the lesion involves a bone cross-sectional area $>50\%$, the diameter of cortical bone defect is $>5$ cm, or pathologic fractures have occurred, so an allo- geneic cortex bone plate is tightly embedded in the window of the lesion cortex bone and a parallel plate is fixed at the same time. Owing to the appropriate procedures and repair methods being carried out, no patients saw lesion fractures and were healed satisfactorily in our studies. After surgery, all patients should accept the functional training and regular re-examination schedule which involves blood tests and all methods being carried out, no patients saw lesion fractures and were healed satisfactorily in our studies. After surgery, all patients should accept the functional training and regular re-examination schedule which involves blood tests and imaging examination (panoramic radiography, CT plain scan, and spiral CT 3D reconstruction) to assess any changes in the graft and lesion healing. After the lesion heals well for about 2 years, the internal fixation is considered for removal.

**Conclusion**

In conclusion, the porous n-HA/PA66 composite has biological safety, good biocompatibility, osteoinduction and osseointegration; it is an efficient, safe, and highly cost-effective bone substitute material worthy of promotion in the orthopedic field.

**Acknowledgment**

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**Disclosure**

The authors report no conflicts of interest in this work.

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