We evaluated the effect of low-intensity pulsed ultrasound stimulation (LIPUS) on the remodelling of callus in a rabbit gap-healing model by bone morphometric analyses using three-dimensional quantitative micro-CT. A tibial osteotomy with a 2 mm gap was immobilised by rigid external fixation and LIPUS was applied using active translucent devices. A control group had sham inactive transducers applied. A region of interest of micro-CT was set at the centre of the osteotomy gap with a width of 1 mm. The morphometric parameters used for evaluation were the volume of mineralised callus (BV) and the volumetric bone mineral density of mineralised tissue (mBMD). The whole region of interest was measured and subdivided into three zones as follows: the periosteal callus zone (external), the medullary callus zone (endosteal) and the cortical gap zone (intercortical). The BV and mBMD were measured for each zone.

In the endosteal area, there was a significant increase in the density of newly formed callus which was subsequently diminished by bone resorption that overwhelmed bone formation in this area as the intramedullary canal was restored. In the intercortical area, LIPUS was considered to enhance bone formation throughout the period of observation. These findings indicate that LIPUS could shorten the time required for remodelling and enhance the mineralisation of callus.

The effects of low-intensity pulsed ultrasound stimulation (LIPUS) are reportedly derived from the promotion of cell differentiation, which induces acceleration of fracture healing with earlier restoration of strength.1 Previous studies have investigated which stages of healing are affected by LIPUS. Many have suggested that LIPUS affects the inflammation, angiogenesis and formation stages of soft callus.2-4 However, whether it influences the formation of hard callus or remodelling remains unclear.1,5,6 Most previous studies have investigated the effects of LIPUS by histological evaluation, which is limited to observation in a two-dimensional (2D) plane. Another method adopted has been mechanical testing of harvested specimens, but this is limited since testing is destructive and the strength of the healing site is evaluated in only one of many planes.7

In recent years, several studies have investigated the morphometry of the site of fracture healing using micro-CT.8-10 The advantage of this technique lies in its non-destructive morphological and densitometric assessments in a three-dimensional (3D) plane. However, evaluation by micro-CT of the effects of LIPUS is only possible after healing has progressed to mineralisation of the callus. This period comprises two stages: the modelling period, during which callus is formed, and the remodelling period, during which mineralised tissues are redistributed by the processes of formation and resorption to restore the original structure.11 The effects of LIPUS should be evaluated in both of these stages. In the modelling period, callus is formed externally from the periosteum and endosteally from the endosteum. The subsequent remodelling period partly overlaps the modelling period. If LIPUS enhances fracture healing by shortening either one of these processes, morphometry by means of micro-CT could identify spatial changes in the localisation of callus in 3D planes. However, no quantitative studies have included 3D morphological evaluation using micro-CT.
Our aim was to evaluate quantitatively the effect of LIPUS on bone healing by means of micro-CT throughout both modelling and remodelling using a gap-healing model in the rabbit.

Materials and Methods

Creation of the model. We used 42 skeletally mature male Japanese white rabbits aged between 21 and 23 weeks and weighing between 3.4 kg and 4.0 kg (Kitayama Labs, Nagano, Japan). Under general anaesthesia, the right hind limb was prepared using a routine aseptic technique. Before the creation of an osteotomy four transfixion pins (diameter, 2 mm; length, 50 mm) were inserted at the metaphyseal regions of the tibia in the frontal plane using a custom-made driver. An anteromedial incision 2 cm in length was made, and the tibia exposed using two mini-retractors after elevating the periosteum. A transverse osteotomy was performed using an MDS36-30 T-saw (MANI, Tochigi, Japan) with a blade 0.36 mm thick in the mid-shaft of the tibia to produce a gap of 2 mm at 12 mm distal to the tibiofibular junction. The osteotomy was immobilised using the transfixion pins which were attached to an external fixator with KE double side bars (IMEX Veterinary Inc., Longview, Texas). The gap was confirmed by inserting a 2 mm spacer block. The site was thoroughly irrigated with saline to ensure that no tissue (bone, muscle, or fascia) was left in the gap. The soft tissues were then re-approximated, and the skin was closed. Proper alignment of the bone fragments was confirmed radiologically (Fig. 1). All the procedures were performed in accordance with the guidelines of the Association for Assessment and Accreditation of Laboratory Animal Care.

LIPUS treatment. We used the LIPUS model SAFHS 2000J (Teijin Pharma, Tokyo, Japan) which transmits a burst of 1.5 MHz sine waves of 200 μsec repeated at 1 kHz with a mean intensity of 30 mW/cm². After post-operative day 3, the LIPUS treatment was commenced under general anaesthesia for all animals. The animals were divided into two groups of seven rabbits, comprising a LIPUS and a control group. In the former group, the transducer was placed on the anterior surface of the operated limb using ultrasound coupling gel for 20 minutes, six times per week for four, six or eight weeks. The control group had a sham inactive transducer applied which was used under exactly the same conditions as in the LIPUS group.

Experimental evaluations were conducted in a blinded manner and the animals were assigned to the two groups in a randomised manner using the envelope method. The same method was used to determine the duration of treatment.

Post-operative management. All the animals were allowed to move freely post-operatively in their cages bearing full weight. Subcutaneous buprenorphine (Otsuka Pharmaceutical, Tokyo, Japan) was injected for relief from pain. The environment was provided in accordance with the guidelines of the Institute of Laboratory Animal Resources and in compliance with the Guide for the Care and Use of Laboratory Animals. General conditions were checked for all the animals at least once a day. Food intake, activity and the condition of the scar were checked. Body-weight was measured on the day of operation and every two weeks. For post-operative care of the external fixator, we checked the clamps daily, and tightened the frame as necessary. The fixator was protected using gauze with a shock-absorbing material. In order to prevent infection at the pin site, irrigation of the pin sites was performed daily. There were no cases of fracture, infection or unexpected death.

Radiography. Anteroposterior radiographs of the tibia were taken together with a 2 mm incremental lead gauge under a tube voltage of 48 kV and current of 3.20 mAs immediately post-operatively and every two weeks thereafter to confirm the alignment of the osteotomy and to evaluate the size of the gap. Each radiograph was taken under exactly the same conditions using a standardised protocol. The gap was then measured by counting pixels overlying it using Photoshop CS3 software (Adobe Systems, San Jose, California). Pixel spacing of the digitised radiograph was calibrated by the number of pixels overlying each step of the lead gauge.

Micro-CT. The animals were killed after four, six and eight weeks and the right tibia was removed and scanned by micro-CT (ScanXmate-E090; Comscantecno, Kanagawa, Japan). The scan was performed along the long axis of the diaphysis, with a voltage of 60 kVp and a current of 80 μA. The area of the scan was 5 mm proximal and 5 mm distal to the centre of the gap, with a resolution of voxel size of 28.57 μm³. The region of interest (ROI) was set at the area of callus healing defined by the gap filled with callus in 2D-CR (Fig. 2) and extended 0.5 mm proximally and distally to the centre of the gap with a total of 36 CT axial scans. A 3D rendering was created for each ROI and the mean intensity of 30 mW/cm².
reconstruction of mineralised tissue was performed using TRI-BONE software (Ratoc System Engineering, Tokyo, Japan). A threshold for newly formed mineralised callus was set at 200 mg/cm³. Morphometric parameters used for the evaluation were mineralised callus volume (BV, cm³) and mineralised callus content (BMC, mg) calculated from the contoured ROI in 3D images and volumetric bone mineral density of mineralised tissue comprising the callus (mBMD, mBMD = BMC/BV, mgHA/cm³). The ROI was subdivided into three zones: the periosteal callus zone (external, black zone in right); the medullary callus zone (endosteal, white zone in right); and the remaining area as the cortical gap zone (intercortical, grey zone in right).

The region of interest (ROI) was set at the callus healing area, defined as the centre of the osteotomy gap with a width of 1 mm (left). This area was subdivided into three zones: the periosteal callus zone (external, black zone in right); the medullary callus zone (endosteal, white zone in right); and the remaining area as the cortical gap zone (intercortical, grey zone in right).

The mean BV for each zone is shown in Figure 5. In the LIPUS groups, the mean mBMD was significantly higher at eight weeks than at four weeks for total, external, internal and endosteal zones (total, 558.10 mg/cm³ (SEM 16.32) vs 332.52 mg/cm³ (SEM 57.75); p < 0.001; external, 512.23 mg/cm³ (SEM 12.88) vs 307.65 mg/cm³ (SEM 33.01) (ANOVA, p < 0.001); internal, 508.22 mg/cm³ (SEM 16.54), vs 328.50 mg/cm³ (SEM 40.44) (ANOVA, p < 0.001); endosteal, 441.12 mg/cm³ (SEM 19.18) vs 313.13 mg/cm³ (SEM 45.21) (p = 0.014)). Comparing the results at the same time points, between the two groups the mean mBMD was significantly higher at eight weeks in the LIPUS group than in the control group in both the external and intercortical zones.

The mean BV for each zone is shown in Figure 5. In the control group no significant differences were observed at each interval. However, in the LIPUS group, the mean BV of the bone fragments at the gap by exactly the same procedure as was used to define external callus. Endosteal callus was defined as the callus inside this surface and intercortical callus as the callus existing between these two surfaces models. For each of these three zones, the mBMD and BV were measured.

**Statistical analysis.** The data on body weight, gap size and micro-CT evaluations of the BV and mBMD were analysed by one-way analysis of variance (ANOVA) using SPSS version 17.0 software (SPSS Inc., Chicago, Illinois). Homogeneity of variance was calculated using Levene statistics, according to which multiple pair-wise comparisons using a Tukey post hoc test were conducted to establish homogenous subsets and the Games-Howell post hoc test was used for unequal variance. Data on body weight and gap size were presented as the mean and SD and BV and mBMD data as the mean and the standard error of the mean (SEM). A p-value ≤ 0.05 was considered to be statistically significant.

**Results**

**Body weight.** The mean body-weight of both the LIPUS and control groups on the day of operation and at every two weeks showed no significant differences for all groups at each time point (ANOVA, p = 0.55 on day at operation, p = 0.23 at two weeks, p = 0.17 at four weeks, p = 0.11 at six weeks and p = 0.55 at eight weeks) (Table I).

**Radiological.** The mean size of the gaps in each group is shown in Table II. No significant differences were apparent in any group at each time point (ANOVA, p = 0.74 on day of operation, p = 0.56 at two weeks, p = 0.48 at four weeks, p = 0.30 at six weeks and p = 0.65 at eight weeks).

**Micro-CT.** Typical images of the 3D reconstructed micro-CT scans at the ROI in the gap for both groups are shown in Figure 3. The mineralised areas of callus were larger in the LIPUS groups than in the control groups. The formation of cortex and medullary canal at the gap were more obvious in the LIPUS groups than in the control groups.

The mean volumetric BMD of mineralised tissue for each zone is shown in Figure 4. Within the control groups no significant differences were found at each interval. However, in the LIPUS groups, the mean mBMD was significantly higher at eight weeks than at four weeks for total, external, internal and endosteal zones (total, 558.10 mg/cm³ (SEM 16.32) vs 332.52 mg/cm³ (SEM 57.75); p < 0.001; external, 512.23 mg/cm³ (SEM 12.88) vs 307.65 mg/cm³ (SEM 33.01) (ANOVA, p < 0.001); internal, 508.22 mg/cm³ (SEM 16.54), vs 328.50 mg/cm³ (SEM 40.44) (ANOVA, p < 0.001); endosteal, 441.12 mg/cm³ (SEM 19.18) vs 313.13 mg/cm³ (SEM 45.21) (p = 0.014)). Comparing the results at the same time points, between the two groups the mean mBMD was significantly higher at eight weeks in the LIPUS group than in the control group in both the external and intercortical zones.
for the endosteal zone was significantly lower at eight weeks than at four weeks (0.13, SEM 0.02 × 10⁻² cm³ vs 0.62 SEM 0.14 × 10⁻² cm³, p = 0.02).

Comparing the results at the same time point between the two groups, the mean BV in the intercortical zone at eight weeks was significantly higher in the LIPUS group than in the control group (1.37 SEM 0.13 × 10⁻² cm³ vs 0.37 SEM 0.15 × 10⁻² cm³, ANOVA, p < 0.001).

Discussion
In order to evaluate the effects of LIPUS using an animal model, it is desirable to adopt a method using a fracture with a sufficiently long healing process because osseous healing is usually so rapid that the effects of stimulation can barely be detected even when using sensitive methods of evaluation.² Secondly, since LIPUS is used clinically to encourage healing,¹⁷ it is necessary to ensure that this approach is effective in stimulating retarded healing. In animal models, creating a model with sufficiently long healing is difficult.¹⁸,¹⁹ We created a 2 mm gap at the site of the osteotomy²⁰,²¹ which was stabilised by a relatively stiff construct, as previously reported by mechanical testing.²²

In previous studies, remodelling of callus has been evaluated histologically. In a rat fracture model it has been reported that woven bone is converted to lamellar bone at the fracture site and that bone marrow is created in the remodelling stage.¹¹ Although histological examination is indispensable for observing the restructuring of woven bone from lamellar bone, the interpretation of the 3D callus structure from the 2D histological views is difficult.

One of the advantages of micro-CT is its higher resolution and the very thin slices which are scanned. Many slices can thus be obtained so that spatial differences within areas of the specimen become negligible. In addition, micro-CT allows automated acquisition and non-destructive morphological and densitometric assessments in 3D. In recent years, this technique has been used in many fields of orthopaedic research.⁹,¹⁰ Using micro-CT, data on the geometrical properties of the healing bone can be acquired along with the spatial distribution of BMD by simultaneous scans of a calibration phantom. We selected the BV and the mBMD of the mineralised tissue comprising the callus volume as the denominator rather than tissue volume. The volume of callus represents the volume of mineralised tissue comprising the callus, but tissue volume includes the bone-marrow empty space around the callus. This means that the mBMD reflects the geometrical properties of the callus as well as density.

The availability of micro-CT for the evaluation of healing bones allows precise, 3D reconstruction of the callus.
The mBMD in the endosteal area in the LIPUS group increased from four to eight weeks, whereas BV decreased during the same interval. In the endosteal area, this represented an increase in the density of newly-formed callus which was subsequently reduced by bone resorption that overwhelmed bone formation in this area. This remodelling process contributed to the formation of the medullary cav-