Introduction

Bone adapts to its mechanical environment and it has been presumed that this adaptation is stimulated by either strain or some strain-related quantity. Mechanical force applied to bone produces two localized mechanical signals on the cell: deformation of the extracellular matrix (substrate strain) and extracellular fluid flow-induced shear stresses and streaming potentials. Cowin et al. (1991) proposed that osteocytes are mechanically activated by interstitial fluid flow through the lacunocanalicular porosity rather than by the bulk strains resulting from loading the whole bone. This proposal was strengthened by findings that in vitro strains of the order of 10,000 to 30,000 μstrain were required to obtain an increased cellular production of osteopontin mRNA and nitric oxide comparable to response to fluid flow.

It has been assumed that the maximum physiological levels of strain experienced by bones are those measured experimentally by strain gauges in vivo on the external bone surfaces up to 2,500 μstrain in humans. However, local strain can reach levels several times greater than the macroscopic fracture strain and possibly this stimulus is significant for bone adaptation. Experimentally it has been found that a nominal continuum strain level of 2,000 μstrain induces local, osteocyte level strains as high as 12,000 to 15,000 μstrain. Trabecular tissue stress and strain estimates from global loading of whole bones have been found to be three or three hundred times that predicted by analyzing trabecular bone as a continuum.

In a work of this group the loads applied to a whole femur during the stance phase of the gait cycle were related to the strain at the level of a single lacuna and of canaliculi. Cell deformations were predicted from detailed linear FE analysis of the microstructure, consisting of an arrangement of cells embedded in bone matrix material. The local macroscopic stress assigned to each point was used to calculate the macroscopic and consequently the microscopic nodal forces. The actual tissue principal strain along the femur deviated considerably from an average tissue value. The predicted bone matrix strains around osteocyte lacunae and canaliculi, were nonuniform and differed significantly from the macroscopically measured strains. Peak stresses and strains in the walls of the lacuna were up to six times those in the bulk extracellular matrix. Significant strain concentrations were observed at sites where the process meets the cell body. The dimensions of the lacunar axes influenced slightly the local strain.

Keywords: Finite Element Model, Microstructure, Cell Local Strain, Lacunar Axis

Abstract

A multilevel finite element approach is applied to predict local cell deformations in bone tissue. The loads applied to a whole femur during the stance phase of the gait cycle were related to the strain at the level of a single lacuna and of canaliculi. Cell deformations were predicted from detailed linear FE analysis of the microstructure, consisting of an arrangement of cells embedded in bone matrix material. The local macroscopic stress assigned to each point was used to calculate the macroscopic and consequently the microscopic nodal forces. The actual tissue principal strain along the femur deviated considerably from an average tissue value. The predicted bone matrix strains around osteocyte lacunae and canaliculi, were nonuniform and differed significantly from the macroscopically measured strains. Peak stresses and strains in the walls of the lacuna were up to six times those in the bulk extracellular matrix. Significant strain concentrations were observed at sites where the process meets the cell body. The dimensions of the lacunar axes influenced slightly the local strain.
Previously, multilevel approaches have been successfully applied to describe heterogeneous technical materials\textsuperscript{10}. Multi-scale finite element models were also developed to predict local cell deformations in the extracellular matrix of a cartilage explant\textsuperscript{11} and in a skeletal muscle tissue construct\textsuperscript{12}.

In this work, a multilevel FE approach was applied to predict local stress and strain field at the cellular level in bone tissue using a loading configuration system of the femur which simulates optimally the \textit{in vivo} loading conditions during walking. Strain was predicted from linear FE analysis of the microstructure, consisting of an arrangement of cells embedded in bone matrix material and the interconnecting canaliculi. The influence of the dimensions of the lacunar axes on the tissue strain in the cell microenvironment was examined.

**Materials and methods**

The microscopic length scale of a lacuna is orders of magnitude smaller than the macroscopic dimensions of a whole femur. In order to overcome the difficulties from the enormous difference in length scales involved, the analysis was divided into three separate problems. A FE analysis of the macroscale problem was performed to determine the strain

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**Figure 1.** Multi-level analysis for the prediction of the strain field in the vicinity of an osteocyte. Macroscale level: a whole femur. Mesoscale level: a quadrant of the cylindrical shell of femur diaphysis. Microscale level: a periodic array of lacunae interconnected through canaliculi.

**Figure 2.** Geometry and dimensions of the model of the bone tissue surrounding a lacuna and the connected to it canaliculi.

<table>
<thead>
<tr>
<th>Geometry</th>
<th>Major axis</th>
<th>Minor axis</th>
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</thead>
<tbody>
<tr>
<td>G1</td>
<td>10 µm</td>
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<tr>
<td>G2</td>
<td>15 µm</td>
<td>7.5 µm</td>
</tr>
<tr>
<td>G3</td>
<td>20 µm</td>
<td>10 µm</td>
</tr>
</tbody>
</table>
distribution of the femur during one-legged stance. The results of the macroscale FE analysis were used to define the applied boundary conditions of a separate mesoscale FE model, representing a quadrant of the cylindrical shell of femur diaphysis with length of 10 mm. In turn, the results of the mesoscale FE analysis were used to determine boundary conditions for a microscale FE model, representing one osteocyte lacuna, surrounded by bone tissue, with the emerging canaliculi (Figure 1).

Cortical bone was assumed to be linearly elastic, homogeneous and transversely isotropic with a Young’s modulus of 17 GPa and Poisson ratio of 0.41 in the axial direction and 11.5 GPa and 0.38 in the transverse directions, respectively\(^{13,14}\). The same material properties values were used in the macroscopic and mesoscopic levels of analysis. Cancellous bone was assumed to be isotropic material with \(E=206\) MPa and Poisson ratio of 0.33\(^{15}\). The hypothesis of transversely isotropic material is very close to the real mechanical behavior of cortical bone material. The values of Young’s modulus of cortical bone tissue in the transverse directions of the femoral shaft are nearly equal (less than 10% differences) and about 30% lower than those in the longitudinal direction\(^{13,16}\).

The cortical and cancellous bone geometry was reconstructed three-dimensionally with data provided by the Internet site http://www.tecno.ior.it/VRLAB. The mesh of the femur geometry consisted of 5469 tetrahedral 10-noded SOLID 92 elements of ANSYS® (v. 8.1) FE code.

The used musculoskeletal system of the hip, developed by Heller et al. (2005), represented the simplest model of the hip muscles resulting in physiological-like hip joint loading throughout the entire gait cycles. The single instant of maximum in vivo hip contact force in the walking cycle was taken to define the load profile\(^{17}\). The hip contact force acts at P0 (Figure 3). The attachments or wrapping points of the muscles involved in walking are labelled P1 and P2. The values of muscle and joint reaction forces are displayed in Table 1.

The boundary condition, applied to the model of the femur, was full constraint of all degrees of freedom at the nodes (gray scale, Figure 4) that lie at the distal end of the femur (knee joint). The mesoscale FE model was discretized using the same element type as in the macroscale model (SOLID 92). A total of 10,432 elements were used. To couple the mesoscale FE analysis to the macroscale FE problem, the local macroscopic stress assigned to a certain macroscopic point (Figure 1) was used to calculate the mesoscopic nodal forces, which were used as external loads at the mesoscale analysis. A linear interpolation of these values was applied to the nodal points on the faces of the mesoscale model. These nodal forces are depicted in Figure 5 and their values are presented in Table 2.

The microstructural FE model has been described in a previous study\(^8\). Briefly, a three-dimensional model of a periodic array of lacunae and the interconnecting them canaliculi was formed as a part of the bone tissue. Lacunae shapes were assumed to be ellipsoid inclusions within the extracellular matrix, interconnected with cylindrical channels representing...
groups of canaliculi. The long axis of the ellipsoid lacuna in the microscopic model was directed to the circumferential direction of the cross-section of the femur. The microstructure representative volume with the details of its geometry is displayed in Figure 2. The dimensions of the lacunar axes are also included in Figure 2. The geometry of the model has been highly idealized. Confocal microscopy images of the lacunae suggest that an ellipsoidal model of the lacuna is a reasonable estimate. McCreadie et al. (2004) have measured lacunar major axis from about 7 to 14 μm in not fractured female indi-
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In our measurements from a male iliac crest, lacunar long axis was found as long as 15 μm, and anisotropy was almost constant throughout the examined region. Based on this result, the anisotropy ratio was kept constant and the size of the long axis of the lacuna varied from 10 to 20 μm. The dimensions of the other microstructural elements (canaliculi, canals, etc.) were obtained from measurements in two-dimensional images of stained transverse sections of a human femur diaphysis. The radius of the Haversian canal was taken to be 25 μm, the distance between two adjacent Haversian canals 200 μm and the width of the canalicular channel was taken to be 5 μm. This latter canalicular channel represents a group of 10 canaliculi, taken together as a single canaliculus. It was assumed that cells do not contribute to the mechanical properties of the tissue. For the discretization of the microscale models, the same element type (SOLID 92) was used as for the macroscale model. An axisymmetric model was utilized and only one-eighth of the microstructure representative volume was used by virtue of symmetry (Figure 6). A total of 11,561 elements were created. The assessed from the mesoscopic scale analysis local stress, assigned to a certain point (Figure 1), which was used to calculate the microscopic nodal forces, is presented.

<table>
<thead>
<tr>
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<th>2</th>
<th>3</th>
<th>4</th>
<th>5</th>
</tr>
</thead>
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<td>-0.031</td>
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<td>-2.584</td>
<td>5.282</td>
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</table>

Table 2. The resultants of forces and moments and the surfaces of the mesoscale model on which they are applied are presented (Figure 5). These forces are calculated from the local macroscopic stress assigned to a certain macroscopic point of analysis (Figure 1) and were used as external loads at the mesoscale analysis.

<table>
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<tr>
<th>Stress component</th>
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<tr>
<td>Y</td>
<td>-0.52</td>
</tr>
<tr>
<td>Z</td>
<td>-35.89</td>
</tr>
</tbody>
</table>

Table 3. The assessed from the mesoscopic scale analysis local stress, assigned to a certain point (Figure 1), which was used to calculate the microscopic nodal forces, is presented.

The dimensions of the other microstructural elements (canaliculi, canals, etc.) were obtained from measurements in two-dimensional images of stained transverse sections of a human femur diaphysis. The radius of the Haversian canal was taken to be 25 μm, the distance between two adjacent Haversian canals 200 μm and the width of the canalicular channel was taken to be 5 μm. This latter canalicular channel represents a group of 10 canaliculi, taken together as a single canaliculus. It was assumed that cells do not contribute to the mechanical properties of the tissue. For the discretization of the microscale models, the same element type (SOLID 92) was used as for the macroscale model. An axisymmetric model was utilized and only one-eighth of the microstructure representative volume was used by virtue of symmetry (Figure 6). A total of 11,561 elements were created. The assessed from the mesoscopic scale analysis local stress, assigned to a certain point, was used to calculate the microscopic nodal forces. A linear interpolation of the values of stress was applied to the nodal points on the faces of the microscale model. These stress values are presented in Table 3 and the nodal forces, used as suitable boundary conditions for the solution of the symmetric problem, are depicted in Figure 6. This technique assumes that the changes in stress and strain fields within the microscale mesh caused by the presence of the lacuna and the canaliculi are not apparent at the edges of the mesh. The microstructure can be thought of as repeating itself near a mesoscopic point. However, the microstructure may differ from one mesoscopic point to another.

The solid bone material at the microscopic level was also assumed to be transversely isotropic. For the determination of the values of elastic moduli at this level, an iteration procedure was followed: The strain field was determined from the solution of the microscopic FE model and the imposed at the nodes forces calculated from the transferred mesoscopic stresses. If the calculated displacements at the microscopic nodes do not coincide with those calculated from the mesoscopic analysis, the modulus of elasticity is gradually increased.

![Figure 8](image_url)
and the procedure is continued to arrive at an updated estimation of the microscopic displacement field, until convergence is succeeded. The calculated in this way elastic modulus was found 10% increased relatively to the macroscopic one. This can be attributed to lack of microporosity at this level.

The computational engine itself at the microscale was validated by computing the stress and strain field for the simplified case of an included spherical hole subjected to a uniform stress and comparing this result with the analytical solution.

**Results**

The used loading configuration, representing one legged-stance and taking into consideration the abductors, the tensor fascia latae and the vastus lateralis muscle groups, produced almost uniform strain distribution along the diaphysis and the longitudinal strain ranged between 2,400 (laterally) and -2,800 μstrain (medially) (Figure 7). The maximum tensile longitudinal strain was observed at the lateral diaphysis and the maximum compressive longitudinal strain at the medial diaphysis and did not exceed 2,800 μstrain. The actual tissue principal strain along the femur deviated considerably from an average tissue value.

Figures 8-10 display the prediction of the principal local tissue stresses and strains on a cellular level at the diaphysis of the femur, where the maximum macroscopic compressive longitudinal strain was observed, for geometry 2. The values of stresses are in MPa.

![Figure 9](image1.png)

**Figure 9.** Prediction of the principal strain and stress II in the environment of an osteocyte within the extracellular matrix, following application of a loading configuration, corresponding to one-legged stance, at the diaphysis of the femur, where the maximum macroscopic compressive longitudinal strain was observed, for geometry 2. The values of stresses are in MPa.

![Figure 10](image2.png)

**Figure 10.** Prediction of the principal strain and stress III in the environment of an osteocyte within the extracellular matrix, following application of a loading configuration, corresponding to one-legged stance, at the diaphysis of the femur, where the maximum macroscopic compressive longitudinal strain was observed, for geometry 2. The values of stresses are in MPa.
the lacunae was considerably larger in comparison to literature values obtained from in vivo strain-gauge measurements in human bone (Lanyon et al., 1975). Peak stresses and strains in the walls of the lacuna were up to six times those in the bulk extracellular matrix. The maximum compressive strain, of the order of 10,000 μstrain, was observed along canaliculi and particularly at sites where the process meets the cell body.

Figures 11 and 12 display the influence of lacunar major and minor axes dimensions on local tissue stress and strain at the nodes 1, 4 and 13. The lacunar size increased or decreased the tissue strain, depending on the location and the direction. The maximum compressive strain was observed at node 1 and its value increased at 3.5% with an increase of the major lacunar axis from 10 μm to 20 μm.

Figure 11. Influence of lacunar axes dimensions (G1, G2 and G3) on local tissue principal strain (I, II and III) at the periphery of the lacuna at nodes 1, 4 and 13.

Figure 12. Influence of lacunar axes dimensions (G1, G2 and G3) on local tissue principal stresses (I, II and III) at the periphery of the lacuna at nodes 1, 4 and 13.
Cells are constantly subjected to mechanical stimuli. Though many studies have been performed in vitro to investigate the mechanisms of detection and adaptation to mechanical strains, the cellular strains remain largely unknown. Most researchers regard as physiological osteocyte deformation the measured bone surface strains. However, the stress-strain fields in the vicinity of the cell cannot be established a priori from macroscopic tissue strains, as cells experience a complex stress and strain state caused by the heterogeneous microstructural architecture and the physical behaviour of both cells and extracellular matrix. Since cellular deformation is likely involved in mechanical adaptation and remodeling, further understanding of the biomechanical origins of alterations in cellular responses can be aided by the use of computational models. 

In this work, a multilevel biomechanical finite element model has been developed to determine the strain in the area of an individual cell and the canalicular processes during the stance phase of gait cycle. The multilevel approach has been validated and proven to be successful for describing technical materials. The effective macroscopic tissue behavior is derived from the microstructural calculations by a computational homogenization procedure. Using this homogenization procedure, the effective material properties can be computed for arbitrary complex microstructural material behavior and cell geometries, even under large strain conditions. In our model, an opposite procedure is followed: the local cell deformations are determined by detailed linear finite element analysis of the heterogeneous microstructure, from the external loading of the macrostructure. A similar procedure was followed by Guilak et al. (2000) in a macro-micro finite element model for describing the local mechanical environment of chondrocytes in articular cartilage.

This model incorporates a detailed analysis of the complex microstructure of cortical bone, consisting of an arrangement of cells embedded in matrix material. This has the advantage that influences of porosity, cell distribution and cell geometry on local strain can be studied. The geometry of the model has been highly idealized. Lacunar ellipsoid shape and perpendicular emanating of canaliculi from a lacuna are reasonable representations of bone microstructure as it has been illustrated by confocal microscopy images and microphotographs under reflected polarized light.

Local periodicity has been assumed in most micromechanical models of bone microstructure. An advantage of this model is that it is 3-dimensional in order to take into account the known local anisotropy behaviour of the bone tissue.

Overall good agreement was observed between the present numerical results for the whole femur and those of previous solutions. The strain magnitudes determined from this finite element analysis appear comparable to those determined from in vivo measurements by strain gauges on tibial shaft surface (up to 2,500 strain in humans), by in vitro measurements on embalmed or synthetic femur models or by FE modeling. The maximum principal strain of the current study was about 7% higher than that found by Speirs et al. (2007), when physiological boundary conditions were applied. The difference can be attributed to the anisotropic elastic modulus adopted in the current study.

The strain at the osteocytes level during a usual daily loading may be up to six times higher than the nominal values predicted for the extracellular matrix and this strain does not depend strongly on lacunar size. An increase of the long lacunar axis from 10 μm to 20 μm increased the maximum compressive strain at 3.50%. The canalicular size does not influence strongly perilacunar strain. An increase of 2.31% in the predicted maximum strain has been observed with an increase in canalicular diameter from 0.362 to 0.421 μm. The microscopic model was indirectly validated by comparison of its results to literature measurements. The strain values, calculated by the FE model at the microscopic level, were comparable to those found by direct in situ measurements of the deformation behavior of cortical bone specimens. A nominal continuum strain level on bone specimens

### Table 4.

<table>
<thead>
<tr>
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<th>G3</th>
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### Table 5.

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approximately equal to that measured in humans in vivo, was found that produced local, osteocyte level, strains as high as 12,000 to 15,000 μstrain. A point of criticism might concern the maximum strain that bone can withstand without failure. Tensile microcracks were found that first appear at approximately 4,000 μstrain, and considerable growth in microcrack density appeared once the tensile strain has passed 8,000 μstrain. Compressive microcracks were first formed at higher macroscopic strains 8,000-10,000 μstrain. These are macroscopic strains calculated from load-deformation curves of the specimens, and they are average over the measured area. Local regions in bone, like lacunae, erosion areas, microcracks, can develop much higher local strain. According to these findings, even the estimated by our model maximum strain of 10,000 μstrain may not be catastrophic to bone. Microcracks have been observed to initiate frequently at osteocyte lacunae providing evidence that strain in the matrix around osteocytes is considerably higher than in the rest of the tissue. The findings of the current work are consistent with the above experimental results.

Cell responses are found to depend on the magnitude and form of the applied strains. In most studies the levels of strain magnitude applied in vitro to cells to elicit responses are those measured experimentally by strain gauges in vivo on the external bone surfaces (2,500 μstrain) and these strains are considered as physiological. According to the results of this study as well of other studies much higher strains, considered as supraphysiological, can be developed in bone at the osteocyte level during a usual activity like gait. Thus, the very high deformation found in vitro necessary to activate a direct mechanotransduction might indeed be observed in physiological conditions at the cell level, because of the local strain amplification.

The current model can be refined in the future by including additional microstructural characteristics (various sizes of lacunae and a larger number of canaliculi with smaller diameters) and extended to include influence of pericellular non-mineralized matrix, fluid flow, or even the influence of the cell properties on the stress-strain fields within the extra-cellular matrix.

References


