A NEW COMPREHENSIVE APPROACH
FOR BONE REMODELING UNDER MEDIUM AND HIGH
MECHANICAL LOAD BASED ON CELLULAR ACTIVITY
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Daniel George, Rachele Allena, Céline Bourzac, Stéphane Pallu, Morad Bensidhoum, Hugues Portier and Yves Rémond

Most of the last century, bone remodeling models have been proposed based on the observation that bone density is dependent on the intensity of the applied mechanical loads. Most of these cortical or trabecular bone remodeling models are related to the osteocyte mechanosensitivity, and they all have a direct correlation between the bone mineral density and the mechanical strain energy. However, experiments on human athletes show that high-intensity sport activity tends not to increase bone mineral density but rather has a negative impact. Therefore, it appears that the optimum bone mineral density would develop for “medium”-intensity activity (or medium mechanical loads) and not for the highest-intensity one.

In this work, we propose a new continuum approach based on bone cell activity being either positive or negative as a function of the intensity of the applied mechanical load. At standard earth gravity without exercise, bone homeostasis is observed with cell activity being at equilibrium. When “medium loads” such as “low-intensity” or “optimized” sport activity are applied, cells are activated and an increase of bone density occurs. On the other hand, “high-intensity loads” such as over-training lead to bone density decrease or bone degradation. Our results are in agreement with the literature and enable us to foresee applications such as optimal sport training for best physical conditions.

1. Introduction

The last hundred years or so have seen many bone remodeling models being developed under the hypothesis that the mechanical energy is the main driving parameter of this complex phenomenon. According to the first law of bone remodeling defined in [Wolff 1986] and reprinted many times, bone mineral density is directly

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dependent on the intensity of the applied mechanical loads. Many strain-related models for bone remodeling have followed since. To name a few, see for example [Carter 1984], [Frost 1987] and its “mechanostat” proposal, [Cowin 1986], [Beaupré et al. 1990], [Turner 1998], or more recently [Lekszycki 2002] where the highly heterogeneous bone microstructure is seen as an optimum structural response to given external mechanical boundary conditions. Many theoretical frameworks of bone remodeling were proposed for cortical [Pivonka et al. 2008] and trabecular [Ruimerman et al. 2005] bone accounting for different bone cells at the origin of this remodeling [Pivonka and Komarova 2010; Klein-Nulend et al. 2013]. More recently, further theoretical models were proposed [Madeo et al. 2011; 2012; Lekszycki and dell’Isola 2012; Scala et al. 2017], some of them also taking into account the complex viscous mechanical behavior of the bone [Andrears et al. 2014b; Giorgio et al. 2016; 2017].

Nowadays, it is generally accepted that without considering the specific effects of the bone cells, whatever the theoretical model, the prediction of bone remodeling remains at best phenomenologically driven. Although at the continuous level (scale of the bone) the continuum mechanics is “manageable”, the integration of continuum biology is highly risky since, for the time being, there are no experimental measurements available in the literature able to link the local cell phenomena to the bone continuum. The full understanding of the bone mechanobiology is still unknown, but some literature exists on its basic principles (see for example [Burr and Allen 2014, pp. 85–86]). It is therefore possible to start developing more precise mechanobiological models, but at the local scale (scale of the cells), and to try understanding what the main biological parameters driving this evolution are [Lemaire et al. 2011]. Nonetheless, bridging the local and the global scales through multiscale or homogenization models remains a challenging task [Lemaire et al. 2005; 2010; 2015]. Multiscale theories on biological materials have been developed recently (see, e.g., [Rémond et al. 2016; George et al. 2017; Spingarn et al. 2017]), but the uncertainties remain about the multiscale models themselves [Sansalone et al. 2015], the scale growth response [Louna et al. 2016], or even the influence of the microstructure on the overall behavior [Sheidaei et al. 2019].

The obvious next step in bone remodeling is to try integrating more biological actions that are at the origin of the bone tissue evolution such as the capillary growth [Bednarczyk and Lekszycki 2016], the nutrient supply [Lu and Lekszycki 2016], or the cell migration [Allena and Maini 2014; Schmitt et al. 2015; Frame et al. 2019; 2018]. However, the homogenization of such local effects at more macroscopic scales [George et al. 2018a; 2018b; 2019] is complex to transpose and interpret. This is even without accounting for thermodynamically consistent models [Martin et al. 2017] or the influence of the bone microstructure distribution over its macroscopic behavior [Bagherian et al. 2019].
At this stage, it is clear that bone mechanobiology is a highly complex phenomenon and, even with the major progress made in the past fifty years, we still know very little to properly describe the bone remodeling scenario. In this work, we want to address the bone remodeling phenomenon at a macroscopic scale based on the direct relationship between the mechanical strain and the bone cell responses (see, e.g., [Ehrlich and Lanyon 2002]). Our approach is evidently dependent on the activation of osteoblasts for the bone creation and of the osteoclasts for the bone resorption triggered by specific signals that are transmitted by the osteocytes, as proposed in [Ignatius et al. 2005; Andreaus et al. 2014a; Rochefort et al. 2010]. In agreement with the experimental literature [Herman et al. 2010; Hao et al. 2017], we find that bone mineral density does not continually increase with the developed mechanical strain, contrarily to the proposed “mechanostat” model by Frost [1987], and that reverse effects can be observed.

2. Model development

2.1. Model construction. We propose a strain energy density (SED) based approach accounting for external mechanical loads driving the cell activity and leading to a macroscopic bone mineral density change at the continuous scale.

It has been observed that practicing a regular physical activity is healthy not only for the heart, but also to reinforce bone structure and stiffness. However, it was also acknowledged that over-training could lead to a worse health condition than the moderate training scenario [Forwood and Parker 1989; Grimston et al. 1991; Herman et al. 2010; Hao et al. 2017]. Thus, for “medium” mechanical loads (i.e., intermittent sport activity under a certain threshold), the bone mineral density increases, whereas for “high” mechanical loads (i.e., critical above the given threshold), the bone mineral density decreases.

We want to keep a simplified approach in order to be able to identify the model’s parameters and to validate it. Our main assumption is that the cell activity is directly proportional to the intensity of the mechanical load, but with a predefined scenario. We define such a cell activity as the quantity of bone formation/resorption (or bone mineral density change) as a function of time and mechanical energy. Hence, for a given mechanical energy level, the cell activity will change the bone mineral density in a given time. The cell activity increases with the mechanical load up to a certain value that is proportional to the amount of available space within the structure (related to the porosity and defined in Figure 1 by the maximum osteoblast activity per unit volume). The cell activity cannot exceed this value and a maximum cellular density exists. The maximum cellular activity (of osteoblasts or osteoclasts) is dependent on the cell density at a given location and at a given time. It is assumed that this activity linearly increases (with the mechanical load)
Figure 1. Schematic of the mechanoregulatory model of cellular activity as a function of the developed mechanical energy $W$ within the structure. Blue dashed line: osteoblast activity $A_{ob}$. Green dashed line: osteoclast activity $A_{oc}$. Red line: bone density $\rho_{bone}$ obtained by the combined blue and green cellular activities.

up to the maximum value (defined by the porosity). Finally, as we are working at the continuous macroscopic scale, we define a continuous representative volume element (RVE) that is large enough so that both osteoblasts and osteoclasts can be active within it at the same time (as a function of the given bone microstructure distribution and the given mechanical strain field) depending on the signals transmitted by the osteocytes. It is therefore assumed that the total bone mineral density change is the sum of the positive (through osteoblasts) and the negative (through osteoclasts) effects within this given RVE. Therefore, bone mineral density is directly proportional to the cell activity and defined by the corresponding units (i.e., the cellular activity is given in $\text{kg} \cdot \text{m}^{-3} \cdot \text{unit time}^{-1}$ of fabricated (or degraded) bone per cell density).

A schematic of the cell activities as a function of the intensity of the developed mechanical energy $W$ within the structure is presented in Figure 1. The homeostasis condition ($W_0$) corresponds to the equilibrium state where it is acknowledged that bone remodeling and cellular activity are nonzero, but they are not integrated within the model. The modeled variations (i.e., bone density change) are dependent on cell activity differences away from the equilibrium state.
The osteoblast activity is shown as positive whereas the osteoclast activity is shown as negative. Both activities increase linearly up to a maximum level as the mechanical energy increases. Two conditions are required here:

1. for a bone density increase for “medium low” mechanical energy \((W_0 < W < W_2)\), the osteoblast activity must be higher than the osteoclast one, and
2. for a bone density decrease with “higher” mechanical energy \((W > W_2)\), the osteoclast activity must be higher than the osteoblast one.

On the graph in Figure 1, the initial osteoblast activity is triggered by osteoclasts (osteoblasts will start being active only after osteoclasts have “cleaned up” the bone surface so bone remodeling can initiate). At \(W_0\), we are at homeostasis state where neither bone formation nor resorption occurs at a constant living load condition. Only continuous bone remodeling of living is present. This means that Figure 1 is correct at a given time \(t\) and corresponds to an equilibrium state once bone formation or degradation has finished. In the current work, this condition corresponds to the start of the analysis (initial zero condition). From \(W_0\) to \(W_1\), both osteoblast and osteoclast activity increase at the same time leading to an increase of the bone mineral density since the sum of both cell activities is positive. When reaching the maximum osteoblast activity, i.e., \(W \geq W_1\), the bone mineral density increase is impacted by the osteoclast activity taking over the osteoblast one. Above a given energy threshold, from \(W_2\) to \(W_3\), the bone mineral density decreases as the combined cell effect is negative. For a mechanical energy between \(W_0\) and \(W_2\), we have a bone density increase, while for a mechanical energy above \(W_2\), we have a bone density decrease. \(W_2\) corresponds to the energy threshold not to overtake to keep a good bone health. The following equations interpret the scheme in Figure 1:

\[
\begin{align*}
\text{if } W_0 < W < W_1, & \quad A_{ob} = k_1 \cdot W + \rho_{\text{Bone}}^{\text{ini}} \quad \text{and} \quad A_{oc} = -k_2 \cdot W + \rho_{\text{Bone}}^{\text{ini}}, \\
\text{if } W_1 < W < W_2, & \quad A_{ob} = A_1 + \rho_{\text{Bone}}^{\text{ini}} \quad \text{and} \quad A_{oc} = -k_2 \cdot W + \rho_{\text{Bone}}^{\text{ini}}, \\
\text{if } W_2 < W < W_3, & \quad A_{ob} = A_1 + \rho_{\text{Bone}}^{\text{ini}} \quad \text{and} \quad A_{oc} = -k_2 \cdot W + \rho_{\text{Bone}}^{\text{ini}}, \\
\text{if } W > W_3, & \quad A_{ob} = A_1 + \rho_{\text{Bone}}^{\text{ini}} \quad \text{and} \quad A_{oc} = -A_2 + \rho_{\text{Bone}}^{\text{ini}},
\end{align*}
\]

where \(k_1\) and \(k_2\) are the coefficients of osteoblast and osteoclast activity increase, \(A_1\) and \(A_2\) are their corresponding maximum values, and \(\rho_{\text{Bone}}^{\text{ini}}\) is the initial bone density, constant over time, in \(\text{kg} \cdot \text{m}^{-3} \cdot \text{unit time}^{-1}\). Thus, we obtain a model with only four parameters \((k_1, k_2, A_1, \text{and } A_2)\) defined as

\[
\begin{align*}
A_{ob} &= k_1 \cdot W + \rho_{\text{Bone}}^{\text{ini}} \quad \text{for } W < W_1, \quad A_{oc} = -k_2 \cdot W + \rho_{\text{Bone}}^{\text{ini}} \quad \text{for } W < W_3, \\
A_{ob}^{\text{max}} &= A_1 + \rho_{\text{Bone}}^{\text{ini}} \quad \text{for } W > W_1, \quad A_{oc}^{\text{max}} = -A_2 + \rho_{\text{Bone}}^{\text{ini}} \quad \text{for } W > W_3.
\end{align*}
\]

The units of these four parameters are given in \(\text{kg} \cdot \text{m}^{-3} \cdot \text{unit time}^{-1}\) for \(A_1\) and \(A_2\), and in \(\text{kg} \cdot \text{mJ}^{-1} \cdot \text{m}^{-3} \cdot \text{unit time}^{-1}\) for \(k_1\) and \(k_2\). Such parameters can be quantified.
Figure 2. Schematic of the mechanoregulatory model or cellular activity as a function of time $t$ for different levels of applied mechanical load (energy) within the structure. Blue dashed line: osteoblast activity $A_{ob}$. Green dashed line: osteoclasts activity $A_{oc}$. Red line: bone density $\rho_{bone}$.

experimentally by histochemistry through the measurement of the different cell density activities as a function of the applied mechanical load at any given time and point of the structure. For instance, osteocyte viability and osteoclast activity can be determined through the ratio between empty (i.e., without cell) and full (i.e., with cell) lacunae. More specifically, osteocyte viability can be determined by cell apoptosis through cleaved caspase-3 activation [Lavrik 2005; Nicholson et al. 1995; Maurel et al. 2013], and osteoclast activity through the measure of the resorption area surrounding the cells (TRAP activity) [Kodama et al. 2009].

Once the cell activity has been defined and quantified as a function of the mechanical energy $W$, we can describe it as a function of time for different levels of mechanical loads (as presented in Figure 2).
Four different cases are considered:

1. When forces correspond to everyday life conditions, we are at equilibrium state ($W_{\text{homeostasis}}$) where the bone density is in the homeostasis condition. In this case, the “average” osteoblastic and osteoclastic cell activities are equal and opposite. Their sum is equal to zero (corresponding to homeostatic state) and bone mineral density remains constant over time.

2. When forces increase ($W_{\text{intermediate}}$), corresponding to a “sport training”-like activity at a moderate level, osteoblastic cells become predominant and the sum of these effects is positive leading to an increase in the bone mineral density.

3. If the “exercise” level increases (over-training-like) leading to an increase of the averaged developed mechanical energy, the energy level reaches a threshold ($W_{\text{max}} = W_2$) corresponding to the maximum energy under which bone density evolution remains at best positive or null. Here again, and similarly to the homeostasis case, but not for the same reason, the combined effect of osteoblasts and osteoclasts provides equilibrium. However, contrary to the homeostasis case, this equilibrium is unstable, depending on the biology that is patient-dependent.

4. Finally, when the average mechanical energy goes beyond the threshold ($W = W_2$), bone density decreases due to an over-expression of the osteoclastic cells reacting to a solicitation that needs to be biologically quantified.

In essence, an average person’s life is the result of the combined effect of all these different cases (being dependent on both the mechanical energy level and time) and bone mineral density evolves accordingly in space and time.

An interpretation of the schematic bone mineral density evolution in time of Figure 2 is given by the four energy intervals

$$
\begin{align*}
\text{if } W_0 < W < W_1, & \quad \frac{\partial \rho_{\text{bone}}}{\partial t} > 0 \quad \text{and} \quad \rho_{\text{bone}} > 0, \\
\text{if } W_1 < W < W_2, & \quad \frac{\partial \rho_{\text{bone}}}{\partial t} < 0 \quad \text{and} \quad \rho_{\text{bone}} > 0, \\
\text{if } W_2 < W < W_3, & \quad \frac{\partial \rho_{\text{bone}}}{\partial t} < 0 \quad \text{and} \quad \rho_{\text{bone}} < 0, \\
\text{if } W > W_3, & \quad \frac{\partial \rho_{\text{bone}}}{\partial t} < 0 \quad \text{and} \quad \rho_{\text{bone}} < 0,
\end{align*}
$$

(3)

where $\rho_{\text{bone}}$ is the bone density.

When combining the two schemes of Figures 1 and 2 on the same graph, we obtain the corresponding Figure 3 where the bone mineral density is a function
of time $t$ and the elastic strain energy $W$ (when we suppose that the mechanical behavior of bone is linear elastic).

As the cell activity is defined as an amount of fabricated (or degraded) bone as a function of its density, the amount of mechanical energy developed, and the time, the variation of bone mineral density can be computed directly. It remains only to calculate the corresponding Young’s modulus (as a function of the bone density), given by the relation $E = E_0 \cdot \rho_{\text{bone}}^2$ [Rho et al. 1995], that is classically accepted in the literature, where $E_0$ is the cortical bone Young’s modulus.

2.2. Model application. A numerical application of the above proposed model is made on a simplified geometry accounting for simple load conditions. We consider a femur diaphysis loaded under compression (i.e., body weight) as presented in Figure 4, left. Since we assume a constant distribution of the stresses through the thickness of the femur diaphysis, we propose to study a simplified 2D rectangular beam of length $L = 50$ mm and height $H = 20$ mm (Figure 4, right).

Of course, it is acknowledged that the exact quantification of the model parameters will depend on the given geometry and structure of the exact experimental model to represent. In this case, we will quantify them only for validation purposes.
Figure 4. Geometry for the numerical application. Left: real geometry of a femoral diaphysis under simple tension. Right: simplified geometry of a rectangular beam under simple tension.

The other material parameters are defined by the Young’s modulus of the cortical bone $E_0 = 20.3$ GPa and Poisson ratio $\nu = 0.3$ [Bernard et al. 2013]. The applied mechanical load is equivalent to a human body weight $F = 400$ N on one leg and an initial normalized bone density $\rho_{\text{bone}}^{\text{mi}} = 0.5$ is taken for validation purposes. We suppose a closed system with no external input.

The model parameters were identified without a priori knowledge of the corresponding biological quantifications of the in vivo conditions and serve here only for validation. The homeostatic energy $W_0 = 1 \times 10^{-5}$ mJ was determined based on an average femur cross-sectional area and its corresponding bone density and body weight load conditions. The other two energies $W_1 = 1.456 \times 10^{-5}$ mJ and $W_3 = 3 \times 10^{-5}$ mJ were defined based on identification of the four parameters of the model by $k_1 = 1 \times 10^5$ kg·mJ$^{-1}$·m$^{-3}$·unit time$^{-1}$, $k_2 = 0.7 \times 10^5$ kg·mJ$^{-1}$·m$^{-3}$·unit time$^{-1}$, $A_1 = 1.456$ kg·m$^{-3}$·unit time$^{-1}$, and $A_2 = 2$ kg·mJ$^{-1}$·m$^{-3}$·unit time$^{-1}$, where $k_1 > k_2$ and $A_2 > A_1$. The intermediate energy $W_2$ is the linear interpolation between $W_1$ and $W_3$. The model was implemented within the software COMSOL Multiphysics, and the results are presented in the following section.

3. Results and discussion

3.1. Validation. The model was subjected to a constant force $F = 400$ N, corresponding to the body weight, for an arbitrary length of time, leading to a constant energy distribution throughout the structure and hence a bone density evolution as defined by Figure 3. An initial numerical validation of the model was made by testing three simplified cases: (i) osteoblasts and osteoclasts have the same activity (i.e., the models parameters are equal and opposite: $k_1 = -k_2$ and $A_1 = -A_2$), (ii) only osteoblasts are active (i.e., $k_2 = A_2 = 0$), and (iii) only osteoclasts are active (i.e., $k_1 = A_1 = 0$). The results are presented in Figure 5.

As expected, an equal intensity of osteoblast and osteoclast activity (Figure 5, left) leads to a constant bone density as a function of time. This is similar to homeostasis conditions when homogenized to long periods of time (everyday life activity). In Figure 5, center, when only osteoblasts are considered (i.e., $k_2 = A_2 = 0$), there is no osteoclast activity; therefore, if the energy level is high enough to trigger
osteoblasts, bone density increases and since the bone stiffness increases correspondingly, then the developed mechanical energy (for a constant applied force) decreases, and so does the osteoblast activity. Finally, when only osteoclasts are considered (i.e., $k_1 = A_1 = 0$, in Figure 5, right), there is no osteoblast activity, bone density decreases and so does its stiffness, and hence the developed mechanical energy increases (for a constant applied force) together with the osteoclast activity.

### 3.2. Results and sensitivity study for different load cases.

Different intensities of $F$ were applied from 360 N to 480 N. The results are presented in Figures 6, 7, and 8. For constant-load cases below 478 N, we observe a convergence towards an increase of bone density (Figure 7) and a corresponding decrease of mechanical energy (Figure 6). For load cases above 478 N, the opposite occurs with a decrease
Bone mineral density $\rho_{\text{bone}}$ variation as function of time for different values of the applied force $F$.

of bone density and an increase of energy. Here, the energy threshold $W_2$ shows up around 478.1 N with the given values of the model parameters.

The threshold (i.e., the switch from positive to negative bone density) is very sensitive to the intensity of the applied force and it is located within a very thin force range around 478 N, deregulating the system very quickly. It is not possible at this stage to know if this is a limitation of the proposed model (due to the simplified assumptions made or the lack of more detailed information on the existing couplings at the mechanobiological level) or if this is really occurring in reality. Such an aspect needs to be investigated further experimentally at a later stage. For the case of the control force (400 N), the mechanical energy $W$ and the bone density $\rho_{\text{bone}}$ reach final values of $9.019 \times 10^{-6}$ mJ and 0.64.

Cell activities are presented in Figure 8. For the control case, their final values are equal to $9.019 \times 10^{-1}$ kg $\cdot$ m$^{-3}$ $\cdot$ unit time$^{-1}$ and $-6.313 \times 10^{-1}$ kg $\cdot$ m$^{-3}$ $\cdot$ unit time$^{-1}$ for $A_{ob}$ and $A_{oc}$, respectively. Thus, we can conclude that for this specific case, a fast stabilization is observed to reach the mechanobiological equilibrium. Since we observed a fast switch in the variable trends around 478 N, we decided to perform more simulations around this threshold in order to highlight such a transition.

Similarly to the developed mechanical energy and bone mineral density variations, when the applied mechanical force increases above 478 N, osteoblast activity reaches its maximum value, and we observe an inversion of the osteoclast activity.
Figure 8. Osteoblast activity $A_{ob}$ and osteoclast activity $A_{oc}$ variations as functions of time for different values of the applied force $F$. Osteoblast activity is shown positive in continuous lines up to a force of 478.15 N above which it remains maximal at $1.46 \text{ kg} \cdot \text{m}^{-3} \cdot \text{unit time}^{-1}$. Osteoclast activity is shown negative in dashed lines up to a force of 480 N showing the transient behavior through 478 N towards bone degradation.

(decreasing to increasing). As the osteoblast activity is at its maximum of $1.46 \text{ kg} \cdot \text{m}^{-3} \cdot \text{unit time}^{-1}$, when increasing the applied force, the osteoclast activity will pass from “ineffective” (below the osteoblast activity of $-1.5 \text{ kg} \cdot \text{m}^{-3} \cdot \text{unit time}^{-1}$) to “effective” as it reaches a final value of $-2 \text{ kg} \cdot \text{m}^{-3} \cdot \text{unit time}^{-1}$. Once the new equilibrium reached, where both osteoblast and osteoclast activities have reached their maximum value, a fast bone degradation is observed (Figure 7). A return to normal physical conditions depends upon a decrease of the applied mechanical force and a new stabilization of the system under study.

The proposed model only relies on a few parameters: $\rho_{bone}^{ini}, A_1, A_2, k_1, k_2, W_0, W_1, \text{ and } W_3$. A sensitivity study was performed in order to evaluate their influence on the overall results. We vary one parameter at a time in the range $\pm 10\%$ and we recorded the obtained results for the energy $W$, the osteoblast ($A_{ob}$) and osteoclast ($A_{oc}$) activities, and the bone density $\rho_{bone}$. The obtained results show that $\rho_{bone}^{ini}, A_1, A_2, W_0, W_1, \text{ and } W_3, \text{ within the } 10\% \text{ range variation, have}
Results of the sensitivity study performed for $k_1$ and $k_2$, where “ut” stands for “unit time”.

<table>
<thead>
<tr>
<th>$k_1$ ($\text{kg} \cdot \text{mJ}^{-1} \cdot \text{m}^{-3} \cdot \text{ut}^{-1}$)</th>
<th>$9 \times 10^4$</th>
<th>$9.5 \times 10^4$</th>
<th>$10 \times 10^4$</th>
<th>$10.5 \times 10^4$</th>
<th>$11 \times 10^4$</th>
</tr>
</thead>
<tbody>
<tr>
<td>( W ) (mJ)</td>
<td>$+11.76%$</td>
<td>$+5.39%$</td>
<td>$9.019 \times 10^{-6}$</td>
<td>$-4.63%$</td>
<td>$-8.67%$</td>
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<tr>
<td>$\rho_{\text{bone}}$ ($\times 100%$)</td>
<td>$-5.42%$</td>
<td>$-2.58%$</td>
<td>$0.6352$</td>
<td>$+2.41%$</td>
<td>$+4.64%$</td>
</tr>
<tr>
<td>$A_{\text{ob}}$ ($\text{kg} \cdot \text{m}^{-3} \cdot \text{ut}^{-1}$)</td>
<td>$+0.61%$</td>
<td>$+0.12%$</td>
<td>$0.9019$</td>
<td>$+0.13%$</td>
<td>$+0.47%$</td>
</tr>
<tr>
<td>$A_{\text{oc}}$ ($\text{kg} \cdot \text{m}^{-3} \cdot \text{ut}^{-1}$)</td>
<td>$-11.80%$</td>
<td>$-5.39%$</td>
<td>$-0.6313$</td>
<td>$+4.63%$</td>
<td>$+8.66%$</td>
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</table>

<table>
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<tr>
<th>$k_2$ ($\text{kg} \cdot \text{mJ}^{-1} \cdot \text{m}^{-3} \cdot \text{ut}^{-1}$)</th>
<th>$6.3 \times 10^4$</th>
<th>$6.65 \times 10^4$</th>
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<th>$7.35 \times 10^4$</th>
<th>$7.7 \times 10^4$</th>
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</thead>
<tbody>
<tr>
<td>( W ) (mJ)</td>
<td>$-6.31%$</td>
<td>$-3.32%$</td>
<td>$9.019 \times 10^{-6}$</td>
<td>$+3.68%$</td>
<td>$+7.82%$</td>
</tr>
<tr>
<td>$\rho_{\text{bone}}$ ($\times 100%$)</td>
<td>$+3.32%$</td>
<td>$+1.7%$</td>
<td>$0.6352$</td>
<td>$-1.78%$</td>
<td>$-3.68%$</td>
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<tr>
<td>$A_{\text{ob}}$ ($\text{kg} \cdot \text{m}^{-3} \cdot \text{ut}^{-1}$)</td>
<td>$-6.31%$</td>
<td>$-3.32%$</td>
<td>$0.9019$</td>
<td>$+3.68%$</td>
<td>$+7.82%$</td>
</tr>
<tr>
<td>$A_{\text{oc}}$ ($\text{kg} \cdot \text{m}^{-3} \cdot \text{ut}^{-1}$)</td>
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<td>$+8.14%$</td>
<td>$-0.6313$</td>
<td>$-8.87%$</td>
<td>$-18.60%$</td>
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Table 1. Results of the sensitivity study performed for $k_1$ and $k_2$.

little effect. The initial bone mineral density value has no effect since we apply a constant force on the structure, so whatever the initial state equilibrium, it will lead to the same final results with a different kinematic evolution. Since most of the evolution occurs during the osteoblast ($A_{\text{ob}}$) and osteoclast ($A_{\text{oc}}$) activities through $k_1$ and $k_2$ parameters, the other parameters do not impact either of the obtained results in this study.

However, it appeared that $k_1$ and $k_2$ have a higher impact on the final results as reported in Table 1. As $k_1$ decreases, $\rho_{\text{bone}}$ decreases too ($-11.8\%$), whereas as $k_2$ decreases, $\rho_{\text{bone}}$ increases ($+15.68\%$). For $W$ and $A_{\text{ob}}$, inverse trends can be noticed when varying $k_1$ and $k_2$. For $k_1$, $W$ and $A_{\text{ob}}$ fluctuate between $+11.76\%$ and $-8.67\%$ and between $-5.42\%$ and $+4.64\%$, respectively. For $k_2$, $W$ and $A_{\text{ob}}$ fluctuate between $-6.31\%$ and $+7.82\%$ and between $+3.32\%$ and $-3.68\%$, respectively. Finally, $A_{\text{oc}}$ varies between $+0.61\%$ and $+0.47\%$ for $k_1$ and between $+6.31\%$ and $+7.82\%$ for $k_2$.

Most of the bone mineral density variation occurs within the $k_1$ and $k_2$ phases. Hence, modifying only one parameter ($k_1$ or $k_2$) at a time has a direct impact on the final results. This effect would be compensated for if both parameters would change oppositely in the same proportions, which would lead to the same final results.

After identification and validation of the mechanobiological phenomena at play in this model (developed mechanical energy, osteoblast and osteoclast activities, and bone mineral density variations) as a function of time for given intensities of mechanical forces, the results were plotted not as a function of time, but as a
Figure 9. Numerical results (in comparison to the initial model hypotheses of Figure 1).

function of the intensity of mechanical energy for all the simulations. The results are presented in Figure 9.

Figure 9 shows the results obtained with the newly proposed mechanoregulatory model. These results should be compared with Figure 1 showing the theoretical hypotheses defined initially. An observation is made that a very strong correlation is obtained between the proposed model and the obtained numerical results. We have an initial increase of the osteoblast and osteoclast activities up to their maximum values that remain constant afterwards. The only difference is that in the numerical model, the initial osteoblast activity being triggered by the osteoclasts was not included, which explains why it is not visible in Figure 9, but starts directly increasing at the origin. We also observe that the bone mineral density evolution follows the sum of both cell activities (function of the applied mechanical force) being positive initially for lower mechanical energy and negative later on for higher energy from an original normalized bone density of 0.5. The maximum osteoblast activity is reached first (energy level $W_1$) followed by the maximum of the osteoclast activity (energy level $W_3$). We note that the equilibrium between osteoblast and osteoclast activity does not correspond to the bone mineral density variation equaling zero (i.e., a return to the initial bone density of 0.5) as there appears to be a time delay of the structure response to the mechanical load. The passage of the bone mineral density from positive to negative variation occurs later. Thereafter, there is a continuing decrease of the bone mineral density with stabilization later on, dependent on the intensity of the applied force.

3.3. Application to variable load conditions. The numerical results were validated on constant load and showed good correlation with the proposed model hypotheses.
Hence, to extract bone density variations in a more “real application”-like environment, variable mechanical forces were considered. A constant force is applied on the model until equilibrium is reached (constant bone density that occurs in normal living conditions after several weeks). Then the intensity of the applied force is changed to reach a new equilibrium. This is done for two intensities of the mechanical force, one below and one above the overload degradation threshold. Results are presented in Figure 10.

At initial mechanical load (300 N), bone density quickly reaches equilibrium (≈ 0.55) with decreasing cellular activity. Next, the force is increased above the threshold (520 N). The cell activity is changed to reach its maximum with immediate response from osteoblasts and delayed response from osteoclasts. This is dependent on the evolution of the elastic strain energy within the model that takes time to develop as a function of the bone density that is present at a given point of the structure and at a given time. Once the energy has developed above the threshold, bone density starts to decrease to reach a new equilibrium that is lower than the previous one. When cycling this effect by applying in turn these two mechanical forces, we observe the corresponding bone density variations and new equilibrium being formed. Decreasing the overloading force will lead to a reincrease of the
bone density coming back to its “normal” healthy working conditions. However, it is anticipated that keeping an overload condition on the bone will probably lead to a complete degradation of the structure. This was not modeled here as it certainly requires accounting for extra mechanobiological actions that were not integrated within this simplified mechanoregulatory model.

4. Conclusion

A new comprehensive approach based on cell activity to describe bone remodeling is proposed to assess the possible bone degradation kinetics when under high-intensity mechanical loads. Despite the complexity of the mechanobiological process, only four experimentally measurable parameters are required to tune this model for specific cases. These are the variations of the bone density kinetics with the intensity of the applied mechanical loads up to their maximum value, and the two maximum values for osteoblasts and osteoclasts activities. The results show the respective contributions of each process on the bone mineral density evolution and are in agreement with the experimental data provided in the literature. In fact, the model is able to depict both the harmful and the favorable effects of high and medium mechanical loads, respectively. With this approach, when experimentally measuring the four model parameters for different load scenario (force intensity, test specimen, aging, diseased, etc.), potential differences are expected between the cases and possible foreseen applications for the optimization process for sport activities.

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