The Scissors Model of Microcrack Detection in Bone: Work in Progress

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ABSTRACT

We have proposed a new model for microcrack detection by osteocytes in bone. According to this model, cell signalling is initiated by the cutting of cellular processes which span the crack. We show that shear displacements of the crack faces are needed to rupture these processes, in an action similar to that of a pair of scissors. Current work involves a combination of cell biology experiments, theoretical and experimental fracture mechanics and system modelling using control theory approaches. The approach will be useful for understanding effects of extreme loading, aging, disease states and drug treatments on bone damage and repair; the present paper presents recent results from experiments and simulations as part of current, ongoing research.

INTRODUCTION

The idea for the so-called “scissors model” first came about in 2001: the concept has now appeared in a number of publications [1-5] and currently is being actively researched by a team which includes materials scientists, cell biologists, microscopists and experts in engineering control theory. In this paper we take the opportunity to present for the first time some of our most recent results, from experimental activities which are still ongoing. Because this paper was written to form part of a special symposium devoted to body tissues under extreme loading and disease, we have pointed out those areas where our approach has contributed, or may contribute in the future, to the understanding of bone mechanics under these particular conditions. However it should be emphasised that we have still a lot to do in the future in considering these particular aspects in more detail.

REPAIR OF BONE MICRODAMAGE: DOES IT MATTER?

Before considering the detection and repair of small cracks in bone – which is the subject of this paper – it is worth asking: “Does it matter?”. Do these cracks threaten the structural integrity of bone in normal use? This is a question which we can answer using a different model which we have developed over the last 12 years, a phenomenological model which describes fatigue damage and repair in bone using the available data, which are now quite extensive, and including statistical scatter via the Weibull approach. This model has been described in a number of previous publications [6-9]; it does not consider at all the underlying physical mechanisms, but it is able to predict experimental data from different animals and different test protocols.
rather well, it gives reasonable predictions of phenomena such as the effect of exercise on stress fracture risk and as a result has recently been employed in the field of sports science [10].

Here we use the model to consider the risk of fatigue failure in a typical bone, loaded with physiologically normal stresses, comparing bone from a typical individual capable of repairing small cracks as they initiate, and an individual suffering from a disease state in which this repair process does not occur. Figure 1 below presents the results in terms of a stress reduction factor, defined as the ratio between the stresses in the normal person and in the diseased person which give the same fatigue behaviour. The behaviour chosen was a 1% risk of fatigue failure per bone during the lifetime of the person, as this is typical of the actual rates of failure of bones in primates in the wild [11]. The stress factor is thus the same as the factor by which the compromised individual would have to reduce the forces on their bones in order to prevent failure from occurring.

Obviously the magnitude of this factor depends on the activity level of the individual: more active individuals will be more at risk, especially those engaging in extreme loadings such as professional athletes and soldiers. We considered six different activity levels as defined by Whalen et al [12]. The results show that repair has a very strong influence: for normal or active individuals it’s necessary to reduce loadings by a factor of the order of 2 to 3 if repair is not available. In practice this would mean that any activity more strenuous than slow walking would be dangerous.

![Figure 1: Estimated stress factor for similar fatigue performance of bones, comparing normal individuals and those having no ability to repair microcracks. Six different activity levels are considered, after Whalen et al [12].](image)

**MICROCRACK DETECTION: THE SCISSORS MODEL**

Many workers have contributed to what is now quite an extensive research activity on the initiation, growth and repair of microcracks in bone. A recent review provides a summary of this work [13]. In brief, cyclic loading *in vivo* causes cracks to initiate in cortical and cancellous bone. These cracks are typically elliptical in shape, being elongated in the direction approximately parallel to the bone’s longitudinal axis due to the material’s anisotropy. On
average the length (from tip to tip) of such a crack is about 100µm on its minor axis and about 400µm on the major axis. We will refer to the minor axis length in what follows because this is the length which appears on transverse sections of bones and so is the value most often reported as the crack length. The number density of cracks varies greatly but can be in excess of 1 crack per cubic millimeter. In vivo crack lengths rarely exceed 200µm, but in cyclic loading tests conducted on bones ex vivo, a proportion (10-20%) of cracks are found to propagate by fatigue mechanisms, one crack eventually causing failure. Based on our experimental experience and some simple fracture mechanics calculations, one can form a picture of the risk which these cracks pose. Very small cracks (say less than 30µm) pose a negligible risk; cracks of the typical length of 100µm can be tolerated but would pose a risk in individuals engaging in strenuous activities; longer cracks (greater than about 300µm) will grow quite quickly under normal in vivo stresses and so must be detected and removed to avoid failure. Thus we have a kind of “specification” for the maintenance system which operates in our bones.

The repair system is quite well described: it consists of two types of cells working together: osteoclasts remove old, damaged bone and osteoblasts fill in the gap by making new bone. Currently though, we don’t understand how microcracks are detected in the first place. There are a number of theoretical models under investigation (see [13]) most of which work from the idea that the crack affects the flow of fluid which is constantly seeping through the porous bone matrix. Our model is quite different: we considered whether the crack would damage any of the cells (osteocytes) which live in bone. We concluded that the cells themselves were unlikely to be fractured by the locally elevated strains near the crack, but that fracture could occur in the long, thin extensions known as cellular processes, which link cell to cell in a network known as the “syncytium”. Each cell has about 100 such processes (see fig.2). We proposed that where a process passes across a crack, the displacements of the crack faces could cause rupture (fig.2). Theoretical analysis using fracture mechanics found that this was most likely to occur due to shear loadings, especially in combination with compressive forces across the crack faces. In fact most cracks in bone experience this combination of shear and compression, due to the typical loadings and crack orientations which occur.

EXPERIMENTAL WORK: MICROSCOPY

The photograph in fig.2 represents the highest level of resolution which we were able to achieve using laser confocal microscopy. To obtain more precise information we examined cracks at higher magnification using scanning electron microscopy. We found fibrous features spanning crack faces: by staining the samples with Phalloidin to reveal the cellular material, we demonstrated that the great majority of these were cellular processes. This is interesting in itself because other workers have proposed that these features are collagen fibrils and have argued that they have a significant toughening effect. For the bone samples which we examined, at least, there were very few collagen fibrils spanning cracks: this finding could be important because cellular processes would be too weak to contribute any significant toughening effect.

We first examined ovine bones which had been taken from animals used in other experiments, which were known to have had only normal loading activities during life. We saw both intact and broken processes across crack faces; we counted the intact ones because it was difficult to count the broken ones reliably.
Figure 2: Schematic illustrating the Scissors Model. Laser scanning confocal image of a bone sample (width of picture = 50µm). Three osteocyte cells can be seen (arrowed) with many processes coming from them. A crack (arrowed, A) crosses the picture: some cellular processes cross the crack and remain intact (circled) whilst others have ruptured.

Figure 3: Experimental data (points) recording the number of intact processes per mm of crack length, as seen on transverse sections of ovine bone after normal \textit{in vivo} loading. Predictions of our theoretical model (line). Recent data from samples subjected to loadings greater than \textit{in vivo} levels (grey circle).

Figure 3 shows the results, plotting the number of broken processes per millimeter of crack length, as a function of crack length. The figure also shows the predictions of our theoretical model, using a typical osteocyte density (20,000/mm$^3$) and assuming normal \textit{in vivo} loading activities. There is a lot of scatter in the data points from individual cracks, but it’s clear that
longer cracks have relatively fewer unbroken processes and that this behaviour is well predicted by the model. Figure 3 also indicates some more recent data obtained from ovine bone samples which had been subjected to cyclic loading in the laboratory at stress levels greater than experienced in vivo. This data, currently incomplete, is indicated by the grey circle, showing that for crack lengths around 100µm there are many more broken processes. All of this is nicely in accordance with our “specification” above: for in vivo loading there is a threshold at somewhat less than 100µm, below which no processes break and so the crack would be undetectable. This threshold shifts downwards if higher stresses are applied which would make the 100µm cracks potentially much more dangerous.

EXPERIMENTAL WORK: IN VITRO CELL STUDIES

In parallel with the above work we have been conducting studies on living cells cultured in vitro. We used MLO-Y4 cells, a line of cells developed for laboratory study, which have been shown to have many properties similar to those of natural osteocytes, but which are easier to culture and maintain for experimental work. We cultured these cells in 3D gels in pots of diameter 10mm, depth 10mm. A crack-like planar defect was introduced, width 5mm, depth 10mm, of varying thickness, using wires of different diameters. We made various measurements to detect the production of different proteins which are known to play roles in cell signalling. As figure 4 shows, we found that after introducing the cracks, there was a significant increase in the production of RANKL, a protein which is known to stimulate the production of osteoclasts. This is highly significant because osteoclast production is the first step in the repair process.

Figure 4: Production of RANKL by osteocyte-like cells in vitro at various times following introduction of a crack-like defect 5mm x 10mm, various widths (160-400µm), compared to undamaged controls.
These MLO-Y4 cells are similar to real osteocytes in bone, but there are some important differences, which we investigated experimentally. We found that real osteocytes have an average separation of 37 µm and have typically 100 processes per cell. Our MLO-Y4 cells, on the other hand, had a separation of 100 µm and only 13 processes per cell on average. This means that the “crack” created in our cell experiments is (in terms of the number of processes which cross it) equivalent to a much smaller crack in bone: in fact it is equivalent to a crack of length 500 µm, somewhat larger than the typical 100 µm, but more comparable than first appears. Given that our experimental cells are 100 µm apart, the 160 µm wire is probably cutting through processes but not rupturing cells (though obviously this is something we need to check) whilst the thicker wires have a greater effect because they probably are also breaking cell bodies, something which we believe does not happen in bone.

THEORETICAL MODEL

A theoretical model, written in the form of a MATLAB simulation, was first developed some years ago [14; 15], based around a fracture mechanics description of fatigue crack propagation in short and long cracks. This model is currently being improved to make several aspects more realistic. One feature which required attention was the simulation of crack initiation, which is a common problem in fracture mechanics models because we have no real theoretical framework for this stage of a crack’s life. To circumvent this problem we examined the data on microcrack lengths in bone samples taken from our own work and other sources. We found that the distribution of crack lengths can be described as a two parameter Weibull distribution, whose constants change in a predictable manner with stress level, number of cycles, animal type etc. Figure 5 shows an example of one of these distributions. We intend to incorporate this information into our model, to provide a realistic description of the lengths and numbers of cracks which initiate; these cracks can then be allowed to grow using standard short-crack growth formulae.

![Figure 5: Example of our analysis of microcrack data, showing a two-parameter Weibull fit to the crack length distribution in bovine bone samples subjected to normal in vivo loadings.](image-url)
DISCUSSION

As mentioned above, this paper is very much a description of work in progress rather than of a finished project. Results to date are very encouraging, suggesting that several aspects of our scissors model can be demonstrated in experiments and predicted theoretically. For example, we have now shown conclusively that cellular processes can span cracks, remaining intact in small cracks under in vivo loadings and breaking in increasingly large numbers if the crack length or the stress increases. Through our cell experiments we have demonstrated that planar defects of approximately the same size as microcracks (when differences in the cell types are accounted for) cause cells to increase production of a substance which stimulates bone-resorbing cells. We aim to incorporate all of these effects into our theoretical model, thus building a simulation of the entire feedback loop whereby microdamage is detected and repaired. One important and difficult step in the development of that model is the simulation of crack initiation: this problem has recently been overcome by studying and modeling crack length data.

What consequences does this work have for tissues under extreme loading and disease? High levels of loading, equivalent to very strenuous or active lifestyles, are known to give rise to stress fractures. This is a major problem, for example, in military recruits, football players and racehorses. Stress fractures occur because fatigue cracks are growing so quickly that the time between detection and repair is no longer sufficient to prevent the development of a macrocrack. Stress fractures can also happen as a result of diseases such as osteoporosis, which may affect the detection and repair system and also the mechanical properties of the bone itself. Drug treatments for osteoporosis work in different ways, targeting different parts of the feedback system. In all of these cases an understanding of the mechanisms of crack detection and repair is clearly vital.

Further work is required to confirm the scissors mechanism and to investigate it in more detail; this work is ongoing at present.

CONCLUSIONS

1. The repair of microdamage by living bone is vital to maintain the necessary level of structural integrity which allows us to go about everyday life without suffering fatigue failures.
2. The scissors model, in which cracks are detected because they cause rupture of cellular processes, is demonstrated to be a plausible mechanism.
3. In particular, the numbers of broken processes rise significantly when crack lengths increase and when stresses increase, within and just above normal physiological levels.
4. In addition, crack-like defects introduced into cell cultures cause increased production of RANKL, which is known to stimulate osteoclast production.
5. Microcracks in bone have a length distribution which can be described by the two-parameter Weibull equation: this finding is useful for the simulation of fatigue crack initiation in our theoretical model.
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REFERENCES


