CHANGE IN 3D PERIARTICULAR BONE DENSITY AFTER KNEE JOINT DISTRACTION OR HIGH TIBIAL OSTEOTOMY IN THE TREATMENT OF OSTEOARTHRITIS

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Abstract

Background: Osteoarthritis (OA) is a degenerative joint disease, associated with both cartilage and periarticular bone change. High Tibial Osteotomy (HTO) is a generally considered method for prolonging the time before a total knee replacement is necessary and to reduce the pain in patients suffering from OA. A relatively new technique is Knee Joint Distraction (KJD). Although there is evidence for an improvement of cartilage after KJD, changes in periarticular bone have not yet been investigated.

Objective: The main goal of this research is to determine the difference in quality of periarticular bone of the tibia before and two years after treatment with KJD or HTO using 3D Computed Tomography (CT).

Methods: Coronal CT images were obtained from two previous conducted studies, a total of 23 patients (mean age 51 ± 7 years; 15 males, 10 KJD) were included. Changes in bone density are related to changes in intensity, measured in Hounsfield Units (HU). In the assessment of the periarticular bone quality, a distinction was made between subchondral and trabecular bone, by calculating intensities in five different layers to a depth of 5 mm beneath the joint surface of the tibia. Bone quality was expressed in mean absolute deviation (MAD) and mean intensity.

Results: Mean intensities seem to be decreased at two year follow up compared to baseline, but these differences were statistically insignificant in both HTO and KJD. Interestingly, in the case of KJD, the MAD of the intensities in all layers of the lateral compartment and some layers of the medial and other compartments, were significantly decreased.

Conclusions: The results suggest that periarticular bone density neutralizes. This was statistically indistinguishable for HTO, but MAD decreased significantly for KJD. This indicates that joint distraction has a positive effect on the quality of periarticular bone.
Preface

Before you lies the thesis that we made as a completion of the bachelor and premaster in Technical Medicine at the University of Twente. It is a reflection of the competences and an opportunity to show the knowledge we acquired during the past years.

We have gained more experience in using software like LaTeX, SPSS and especially Matlab. We cheered and raised our hands in the air when Matlab showed us what we wanted it to show, but we execrated it when the red lines popped up for the umpteenth time.

To us, performing this research was an interesting challenge, in which we have felt delighted, as well as disheartened. But: “If the job was easy, there should be no need of the technical medical profession.” We would like to thank Cees Slump for this insight and many more he gave us during the process. Another thanks to Mattiënne van der Kamp for keeping an eye on the process and guiding our personal and professional development. Furthermore we would like to thank our supervisors from the UMC Utrecht for their supervision and contribution in this project. Special thanks to Nick Besselink, who was always available and willing to guide and support us during these ten weeks.

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1 Introduction

1.1 Osteoarthritis

Osteoarthritis (OA) is one of the most common disabling diseases in developed countries. Worldwide, the prevalence of OA is estimated to be 9.6% for men and 18.0% for women aged over 60 years.\(^1\) OA can have effect on various different joints, but knee OA is most commonly seen.\(^2\)

OA is a degenerative joint disease, which is characterized by progressive degeneration of cartilage, subchondral sclerosis, osteophyte formation, changes in periarticular structures and joint inflammation, as shown in figure 1.\(^3\) These changes often lead to pain, stiffness and reduced mobility of the joint. The main cause of OA is still unclear, but there is evidence that biomechanical changes lead to damaged cartilage and bone, deteriorating the joint and inducing OA.\(^4\) It is believed that this process is counteracted by the formation of osteophytes which try to restore mechanical load within the joint.\(^5\)

Periarticular bone is composed of subchondral and trabecular bone. In OA, the formation of osteophytes and cysts can be found in the subchondral bone and trabecular bone. Subchondral bone thickness in the knee joint can vary from approximately 0.1 – 1.5 millimeter, where maxima can be seen at the central zones of the tibia and minima at the peripheral zones.\(^6\)

![Figure 1: Schematic illustration of changes in osteoarthritis.\(^7\)](image)

1.2 Treatments

Because OA still is an incurable disease, treatment focuses on reducing the symptoms. This is mainly done by reducing the load on the knee, physiotherapy and painkilling.\(^8\) If these options do not improve the circumstances sufficiently, surgery could be another possibility. High Tibial Osteotomy (HTO) relieves the pressure on the affected side of the knee by correcting the angle between the femur and tibia. This procedure is usually done if the medial side of the knee is affected by OA. An HTO can be done by the medial opening wedge or the lateral closing wedge technique. With both approaches the load on the medial side decreases while the load increases on the lateral side of the knee joint. In this way, the stress on the affected side is reduced.\(^9\)

A relatively new treatment is Knee Joint Distraction (KJD). In this surgical procedure the knee is externally fixated and distracted by a few millimeters.\(^10\) This distraction changes the pressure in the knee and takes the load of the cartilage and the periarticular bone tissue. These factors have shown to be critical for recovering of cartilage and bone tissue.\(^11\) However, it is not
well understood how these principles exactly work.\textsuperscript{[12]} The aim of both HTO and KJD, especially in relatively young patients, is to reduce the pain and stiffness related to OA and postpone the placement of a prostatic knee, which is the final option to treat OA.\textsuperscript{[10]}

\begin{figure}[h]
\centering
\includegraphics[width=\textwidth]{figure2.png}
\caption{Two possible treatments for OA.\label{fig:figure2}}
\end{figure}

1.3 Study goal

In OA, periarticular bone density is increased due to sclerosis. However, formation of cysts causes decreased densities at certain locations. Previous research showed a normalization of the subchondral bone after distraction of the ankle joint. The mean subchondral bone density was decreased, while abnormally low densities due to cysts increased to more normal densities.\textsuperscript{[12]} KJD also seems to have good results, but there has never been a qualitative assessment of bone tissue after KJD. Therefore, the main goal of this study is to determine the quality of the periarticular bone by composing 3D images of the knee joint using Computed Tomography (CT). In this research, the quality of periarticular bone was assessed by the bone density of subchondral bone and trabecular bone for both treatments. The main question of this research is: What is the difference in quality of the periarticular bone before and two years after treatment with KJD or HTO, determined by 3D-CT?
2 Methodology

2.1 Computed Tomography

For this research, CT images were used to compose a 3D reconstruction of the tibia, so the intensity displayed by the voxels represent the bone density at a specific anatomical position. The use of CT is an advantage over the conventional bone density measurements, so called DEXA scans, because it assesses the volumetric density (mg/cm$^3$) rather than the 2D area density (g/cm$^2$) generated from a DEXA scan.\textsuperscript{[15]}

Because the DEXA scan is only two dimensional, it calculates the mean intensity along the direction of the scan axis. For this reason, a DEXA scan would be insufficient to assess changes in bone density because the increased and decreased intensities at different locations that can be found in OA will give a mean value that is comparable to the mean intensity in healthy bone tissue. Therefore, the displayed values by use of a DEXA scan are not truly representative for the bone density. The 3D CT reconstruction can display the intensity for every different voxel in the 3D space, making it more suitable to assess changes in bone density.\textsuperscript{[16]}

2.2 Subjects

Subjects were selected from two trials reviewed by the Medical Ethical Committee (MEC #11-072 and MEC #10-359). Patients were all diagnosed with severe OA of the knee and indicated for a HTO by an orthopaedic surgeon. The inclusion criteria for both trials were set as followed:\textsuperscript{[17]}

- Patients with medial or lateral tibio-femoral compartmental OA considered for HTO according to regular clinical practice;
- Age < 65 years;
- Radiological joint damage: Kellgren and Lawrence score > 2;
- Intact knee ligaments;
- Normal range-of-motion (min. of 120° flexion);
- Normal stability;
- Body Mass Index < 35.

Exclusion criteria for this research were:

- Mechanic axis-deviation (varus-valgus) < 10 degrees;
- Psychological inabilities or difficult to instruct;
- Not able to undergo MRI examination (standard daily clinical practice protocol);
- Inflammatory or rheumatoid arthritis present or in history;
- Post traumatic fibrosis due to fracture of the tibial plateau;
- Bone-to-bone contact in the joint (absence of any joint space on X-ray);
- Surgical treatment of the involved knee < 6 months ago;
- Contra-lateral knee OA that needs treatment;
- Primary patello-femoral OA.
2.3 Data Collection

CT images of the subjects were provided by the department of Rheumatology and Clinical Immunology, University Medical Center Utrecht, and made by a Phillips Brilliance ‘64 CT scanner. These CT images were taken prior to treatment (baseline) and at two years of follow up and were obtained in the period from July 2012 to September 2015. To process the data, as described in the following section, scans containing a coronal view of the knee joint were used. Therefore, CT data of 23 patients (mean age 51 ± 7 years; 15 males, 10 KJD) were included.

2.4 Data Processing

In order to obtain 3D results that can be analysed to assess the quality of the periarticular bone, the data was processed through the actions written below. All processing was done with the use of Matlab R2015b. An overview of the processing steps can be found in a flowchart in appendix I. An explanatory list of certain Matlab commands can be found in appendix II.

2.4.1 Load DICOM files

The script that was used to read the DICOM files in Matlab was based on a script retrieved from Matlab Central/File Exchange. Some adjustments were made to meet specific requirements. The output of the script is a volume image matrix of the loaded DICOM files.

2.4.2 Segmentation

Medical image segmentation is an important part of the data processing. Using segmentation, a region of interest, in this case bone tissue, can be extracted from surrounding tissues in medical images. Many segmentation methods exist, both automatic and semi-automatic.

(a) Segmentation based on a global threshold of 306 Hounsfield Units.

(b) Segmentation based on a local threshold obtained with the \textit{imrect} function in Matlab.

(c) Segmentation based on a global threshold obtained with Otsu’s method.

(d) Segmentation based on a global threshold by visual assessment.

Figure 3: Different segmentation methods that were considered to segment bone from surrounding tissue.
To extract bone from the surrounding tissue, techniques like global, local and Otsu’s thresholding and region growing were considered. A comparison of different segmentation methods, as can be seen in figure 3, showed minimal differences in accuracy. Global thresholding by use of one certain threshold value appeared to be the fastest and least subjective method to segment the bone tissue from the surrounding soft tissue. Therefore, segmentation was done based on one threshold value of 306 Hounsfield Units (HU). This value can separate high intensity voxels (e.g. cortical bone) from low intensity voxels (e.g. soft tissue).[19]

2.4.3 Splitting tibia and femur

In order to look at the tibial plateau, the femur had to be removed from the 3D reconstruction. Based on the assumption that both tibia and the femur are two different connected structures, the tibia and femur were labelled by the function `bwconncomp` after segmentation. After the labelling of both bones, the femur was removed from the 3D image, based on the height. In order for this method to work the tibial and femur bone cannot be connected at any location in the scan. By using the function `imerode`, all the edges of both bones were reduced by one pixel to minimize the chance that the tibia and femur would still be connected. This created small gaps in the images which were partially repaired using the `imclose` function. Scans that had the tibia and femur still connected after this adjustment were excluded from this research, which was the case in the scans of eight subjects. Apart from splitting the bones, this section of the script was also used to filter the scan. Smaller groups of pixels that were not connected to either the tibia or femur were extracted from the data.

2.4.4 Intensity and location of tibial plateau

Bone changes were tracked by five layers of each 1 mm following the bone contours, to a total depth of 5 mm from the joint surface as shown in figure 4. Layers three up to and including five represent trabecular bone, while layers one and two can contain subchondral bone as well as trabecular bone, depending on anatomical variations. To calculate the number of pixels that was needed to create layers that represented 1 mm the pixelspacing was extracted from the DICOM-info.

To create matrices for different depths in the segmented and splitted tibia that includes data of location and intensity of the tibial plateau, a script was written to extract these values. These matrices were used for visual as well as statistical analysis. The intensity value of each layer was determined by taking the mean of the intensities of the number of pixels that form 1 mm.

![Figure 4: Grayscale images of the tibial plateau at different depths from the surface.](image)
2.4.5 Reconstruction

To check the result of the segmentation visually, a script was retrieved from Matlab Central/File Exchange. This script requires input data of a 3D image volume of the type double, single, (u)int8, (u)int 16 or (u)int32. In this research, an input of the type uint16 was used. The 3D reconstructions were created with Stradwin 5.1.

Because most changes in intensity and surface were expected to be found in the tibial plateau, an intensity based colour map was plotted over a surface based reconstruction. This was done by plotting two different matrices: one which contained information about the surface, and another with information about the intensity. The tibial plateau was reconstructed by plotting the intensity based colour map on top of the matrix with surface information. In this way, an indication of bone density was given at each location. This was solely used to check if the tibial plateau was created correctly. This was the case in all but one patients, which was removed from the research.

2.5 Measurements and Calculations

In order to show changes in quality of bone tissue, three different areas at the tibial plateau were defined based on the amount of weight bearing. The defined areas were: the tibial plateau underneath the medial and lateral condyle of the femur, and all other parts of the tibial plateau, as shown in figure 5. The two condyles were created by forming a matrix which contained the information of the femur location viewed from bottom up. This matrix was reduced to the point where only the medial and lateral condyles remained. Using this matrix, the weight bearing areas of the tibial plateau were selected.

Changes in bone density were analysed by comparing the mean intensity of each compartment at baseline and two year follow up. Besides the mean value, the Mean Absolute Deviation (MAD) was also calculated, because it was expected that there would be an increase as well as a decrease in the bone density which would be unnoticeable using the mean intensity. The MAD represents the dispersion around the mean intensity of the data and can give more information about the different densities that can be found in the bone. In this way, the quality of the bone can be calculated more reliable. The MAD is expected to be decreased after the treatment. The lateral and medial side of the knee joint were considered separately to see the change in MAD at each side of the knee. Since it is assumed there will be an increase in pressure at one side of the knee and a decrease at the other after HTO.

![Image](image.png)

Figure 5: Selection of the medial (blue) and lateral (red) compartments on the tibial plateau was based on the location of the medial and lateral condyle of the femur. The remainder of the tibial plateau is defined as other compartment.
2.6 Statistics

All data was analysed using the statistical analysis software package IBM SPSS Statistics 23. The Shapiro-Wilk Test was used in order to detect if the data showed a normal distribution. For normally distributed data, statistical significance was determined by a paired samples T-test. The variables at baseline and at two year follow-up served as a pair in the samples T-test. When data showed no normal distribution, the nonparametric test Related-Samples Wilcoxon Signed Rank Test was applied. For both parametric and nonparametric tests, significance level was set at $p = 0.05$. 
3 Results

3.1 Mean intensity outcome

In total, 23 patients could be included to examine bone quality after treatment (13 HTO, 10 KJD). Mean intensities in all three compartments (medial, lateral and other) followed a normal distribution in both KJD and HTO. Although mean intensities at two year follow-up seem to show a decrease relative to baseline, these differences were statistically indistinguishable.

3.2 Mean Absolute Deviation outcome

In case of HTO, a nonparametric test showed no significant differences in the MAD in all three compartments. In KJD, not all variables showed a normal distribution in case of MAD so different statistical tests were used. Table 1 shows the results of the parametric and nonparametric tests that were used to determine if KJD had significant change in MAD. Especially in the lateral compartment significant change in MAD can be found.

<table>
<thead>
<tr>
<th>Compartment</th>
<th>Test used</th>
<th>Significance per millimeter from joint surface</th>
</tr>
</thead>
</table>
| Medial      | Related-Samples Wilcoxon Signed Rank Test (nonparametric) | 1: \( p = 0.028 \) (sig.)  
2: \( p = 0.059 \)  
3: \( p = 0.053 \)  
4: \( p = 0.041 \) (sig.)  
5: \( p = 0.059 \) |
| Lateral     | Paired-Samples T Test (parametric) | 1: \( p = 0.029 \) (sig.)  
2: \( p = 0.025 \) (sig.)  
3: \( p = 0.026 \) (sig.)  
4: \( p = 0.027 \) (sig.)  
5: \( p = 0.029 \) (sig.) |
| Other       | Related-Samples Wilcoxon Signed Rank Test (nonparametric) | 1: \( p = 0.047 \) (sig.)  
2: \( p = 0.059 \)  
3: \( p = 0.059 \)  
4: \( p = 0.059 \)  
5: \( p = 0.059 \) |

Table 1: Results statistical significance per millimeter from joint surface for KJD treatment. Baseline and two year follow-up were compared.
In figure 6 can be seen that MAD has decreased for both HTO and KJD treatment in all compartments in 1-5 mm from the joint surface. MAD decreased more after KJD in comparison to HTO at baseline and two year follow-up. MAD also decreased with each millimeter in depth. It should be noticed that at baseline MAD at 1 mm from the joint surface was higher than at 2-5 mm for both the HTO and KJD treatment.
After treatment with HTO, changes in bone density were visible in especially the lateral compartment. In figure 7 the lateral compartment shows an increase in intensity, and therefore bone density, at the two year follow-up image. After treatment with KJD a change in intensity was also visible. In figure 8 the medial compartment shows a decrease in intensity.

Figure 7: Intensity based colour map of the lateral and medial compartment was reconstructed on tibial plateau. Images of one subject at baseline and two year follow-up after treatment with HTO are shown in HU.

Figure 8: Intensity based colour map of the lateral and medial compartment was reconstructed on tibial plateau. Images of one subject at baseline and two year follow-up after treatment with KJD are shown in HU.
4 Discussion

The aim of this study is to determine what the influence is of KJD and HTO on the quality of the periarticular bone. This was done by assessment of different variables in different depths of the tibial bone that represent bone density. The mean intensity and the MAD were reviewed for the weight bearing areas underneath the medial and lateral condyles of the femur in particular. The results show that there is no statistical significant difference in mean intensity for all the different areas after both HTO and KJD. However, the results show a slight decrease in intensity after both treatments. The fact that there is no significant change in the mean intensity could be explained by cyst formation in OA. The cysts that are formed in bone tissue have low intensity values on CT images. The bone tissue also contains parts with scleroses that have high values on CT images. Because these values neutralize each other in a mean intensity value, the mean intensity at two year follow-up may not be that much different from the mean intensity calculated at baseline. As a result, change in bone density is harder to statistically prove.

To overcome this effect, this research focuses on the MAD because it displays absolute changes in the bone density. The MAD after KJD significantly decreases in the medial, lateral and other compartments of the tibial plateau. This shows that after the treatment the bone density of the periarticular bone of the tibia neutralizes, meaning the quality shifts towards the quality of healthy bone. This indicates that KJD is a suitable method to treat patients suffering from OA. However, the results of the patients treated with HTO do not show significant changes in the MAD. This can be explained by the fact that HTO is a more invasive procedure which results in more damage to the bone. Because the shift in weight bearing areas caused by HTO, which is the main principle of this procedure, quality improvement of the subchondral and trabecular bone does not necessarily have to occur. As a result of this shift in weight bearing, pressure areas also change. In some scans there was an increase in intensity visible in the lateral compartment due to the increased pressure on this side of the tibial plateau. Since patients treated with HTO have a main affected side, the treatment could help in reducing pain, because HTO showed little decrease in bone density on the main affected side.

4.1 Computed Tomography settings

The displayed HU for a certain structure might not be the same for every scan, since the HU scale is normalized to the brightness and contrast of distilled water at standard pressure and temperature. The Hounsfield scale is linear between two points:

\[
1000 \times \frac{\mu(x,y) - \mu_{\text{water}}}{\mu_{\text{water}}}
\]  

(1)

Because the polyenergetic output for each CT scanner is different, but also for the same scanner with different settings, the attenuating properties may vary. This means the x-ray beam spectrum can vary and therefore the number of HU can not be compared directly. To overcome this problem there are two valid solutions, the first one is using dual energy radiography. This is a method using a high and low energy acquisition, this way the energy dependant factors can be mathematically excluded. The other solution is by use of a phantom with known density. In this case, the density of the phantom is known and a HU value will be given, which makes it possible to compare the other structures in the scan to this density.\[21\]
Since the CT images were made before the start of this research, the use of a phantom nor dual energy radiography was possible. For the CT images used in this research the same CT scanner was used and all settings were equal except for the Peak KiloVoltage (kVp) for one patient. The kVp was set at 120 kVp for all patients but one, a value of 100 kVp was chosen for this CT scan. for the scans at baseline and at two year follow-up of that patient that were to be compared with unequal settings, the x-ray spectra were found within a similar window as shown in figure 9. Because of the similar x-ray spectrum it was decided to include these CT scans in the research.

![Figure 9: X-Ray spectrum of the CT-images. In both spectra can be seen that the values focus around the same point, indicating that there is no shift in values due to the difference in kVp.](image)

### 4.2 Future recommendations

#### 4.2.1 Partial volume effect

An aspect that has to be taken into account when using CT data is the partial volume effect. Because voxels have a certain size, different types of tissue can be captured within the same voxel, resulting in a mean voxel intensity of those tissues. This effect could have had a significant influence on the outcome of this research, since it mainly looked at bone surfaces.

For future research, this effect could be reduced to a minimum by selecting scans with the lowest possible slice thickness.

#### 4.2.2 Ultra thin slices

Just after the script was nearly finished, ultra thin slices were available. These ultra thin slices were made in axial view, instead of the coronal view the script was originally written for. Because ultra thin slices consist of more data, they were expected to be more accurate. Also, the partial volume effect would be reduced when using these ultra thin slices. Therefore, this data was loaded into the script after using the `permute` function to reconstruct the coronal slices. As a result, the proportions of the images were incorrect and there was more noise after segmentation than in the original coronal slices. To fix these problems, a large part of the script should be rewritten and keeping the time to the deadline in mind, the choice was made to keep using the original, less thin, coronal slices.

For the future it would be recommended to use ultra thin slices, since it can highly improve the accuracy of the calculations.
4.2.3 Segmentation

Several threshold methods for segmentation were considered, but eventually a value based on literature study was chosen. This was done because the difference in accuracy for the different methods seemed minimal, the results would be less influenced because the threshold value is the same in every scan and the script was much faster this way. Nevertheless, a more advanced threshold method could improve the segmentation of the bone tissue, especially because the written script is not capable of splitting the bones when the segmented tibia and femur where connected with several voxels. During the research this lead to the exclusion of eight subjects. Since the excluded subjects were of both the KJD and HTO groups, it is assumed that the exclusion did not affect the outcome of this research. For future researches, it is recommended to include a larger population which would increase the validity of the study.

Also, when looking at the results this threshold method seems not fully capable of filtering all soft tissue. The first layer, surface until 1 mm depth, had a very large range of intensity values, which is not expected in bone tissue. When zooming in at the image of this layer, a small layer of one pixel of surrounding soft tissue could be seen. This layer was filtered away with an `erode` function in Matlab, but in some cases the segmentation and the erode function created gaps in the tibial bone. Smaller gaps could be closed by using the function `imclose` in Matlab but this was insufficient for larger gaps. This resulted in a loss of data that could have influenced the outcome of this research. In this research the limited time resulted in imperfect segmented bone.

Since the segmentation of the tibial bone plays such a crucial role in the reconstruction of the tibial plateau, it is essential that this process works as good as possible. For future research the segmented bone should be optimized before continuing the calculations. When the bone contains less gaps and noise the measurements will be more reliable and significant changes will be easier to prove. Perhaps different segmentation methods should be considered apart from thresholding. Altogether, a more advanced segmentation method could improve the segmentation step and therefore all following steps in the script.

4.3 Conclusion

The results of this research indicate that periarticular bone density changes in response to KJD and HTO. The mean densities showed a slight decrease after two years but could not be significantly proven for either procedure. The changes of MAD in patients treated with HTO were also statistically indistinguishable. However, the MAD did significantly change with patients undergoing KJD. This suggests that periarticular bone density neutralizes and that joint distraction has a positive effect on the quality of the bone.
References


Appendices

Appendix I  Flowchart Methodology

Data of 35 patients were provided by UMC Utrecht

Data of 32 patients were suited for matlab

DICOM files were read into matlab

Images were segmented based on thresholding

Data of 24 patients were used to extract results

The femur was extracted from the image matrix

Axial images of tibial plateau were created

Measurements and calculations were based on these images of 23 patients

3 patients were excluded due to missing BL or 2 year data

8 patients were excluded due to insufficient segmentation

1 patient was excluded because the tibial plateau was created incorrectly
Appendix II  List of Matlab Commands

*Bwconncomp*: finds the connecting pixels in 3D space. A value of connectivity can be specified.

*Imclose*: closes gaps in an image with a specified shape and size.

*Imerode*: removes specified number of pixels from the boundaries of connected pixels.

*Imrect*: creates an interactive tool in which the user can select a certain rectangle area using the mouse. Different values can be extracted from the selected area, for example position of the edges and values of the pixels within the rectangle.

*Permute*: rearranges dimensions of 3D matrix.
%% read dicom files
for g = 1:2
clearvars -except g;
close all;

default_dicom_fields = {...
    'Filename', ...
    'Height', ...
    'Width', ...
    'Rows', ...
    'Columns', ...
    'PixelSpacing', ...
    'SliceThickness', ...
    'SliceLocation', ...
    'ImagePositionPatient', ...
    'ImageOrientationPatient', ...
    'FrameOfReferenceUID', ...
};

% We need these checks because to calculate the "extra_fields", we
% need to have the PixelSpacing and SliceThickness data. If not, we
% leave out the extra_fields.
no_pixel_spacing = false;
no_slice_thickness = false;

extra_fields = {...
    'PhysicalHeight', ... % Height (cols) of slice in mm
    'PhysicalWidth', ... % Width (rows) of slice in mm
    'PixelSliceLocation', ... % Slice z-location in pixels
    'PixelSliceThickness', ... % Slice thickness in pixels
    'SliceData' ... % The slice image data
};

dicom_directory = uigetdir();
all_fields = [default_dicom_fields, extra_fields];

% Get directory listing
listing = dir(dicom_directory);
% number of files
N = numel(listing); % How many entries in the directory listing
if (N<3)
    error('Empty folder');
    return
end

slice_data(N) = cell2struct(cell(size(all_fields)), all_fields, 2);
h = waitbar(0,'Reading DICOM Files...', 'WindowStyle', 'modal');
true_index = 0; % a sequential index of dicom files, that is ignoring
% files of other types.

for i = 3:length(listing) % loop through directory listing, but skip '.' and '..'
    filename = listing(i).name;
    [dummy_path, just_the_name, extension] = fileparts(filename);
    full_path = fullfile(dicom_directory, filename);

    goodfile = false;

    % Check for good dicom file
    if isdicom(full_path)
        true_index = true_index + 1;
        header = dicominfo(full_path);
        slice_image = dicomread(header);

        % Save selected header data into the structure slice_data
        for j = 1:numel(default_dicom_fields) % loop through dicom field names
            current_field = default_dicom_fields{j};
            % Deal with requested fields not found in header
            if isfield(header, current_field)
                slice_data(true_index).(current_field) = header.(current_field);
            else
                ['header did not contain the field ' current_field]
            end %if

        end % loop through dicom field names

        % Save slice data
        slice_data(true_index).SliceData = slice_image;

        % Save extra fields
        needed_header_tags = [...
            isfield(header, 'PixelSpacing'), ...
            isfield(header, 'SliceThickness'), ...
            isfield(header, 'SliceLocation')... ];

        if all(needed_header_tags)
            pixel_spacing = header.PixelSpacing;
            slice_data(true_index).PhysicalHeight = ...
            double(pixel_spacing(1)*header.Columns);
            slice_data(true_index).PhysicalWidth = ...
            double(pixel_spacing(2)*header.Rows);
            % need to double check which aspect ratio goes with cols/rows
            slice_data(true_index).PixelSliceLocation = ...
            header.SliceLocation / mean(pixel_spacing);
            slice_data(true_index).PixelSliceThickness = ...
            header.SliceThickness / mean(pixel_spacing);
        else
            no_pixel_spacing = true;
        end % if pixel spacing
end % if isdicom

waitbar(i/N,h);
end % loop through directory listing

% Eliminate empty structs at end.
slice_data = slice_data(1:true_index);

waitbar(1,h);
close(h);
warning on;

% Check that some dicom slice was found
if true_index < 1
    'No dicom slices found...returning empty'
    volume_image = [];
slice_data = [];
image_meta_data = [];
return
end

% If SliceLocation is known, sort by that. This is deemed more
% accurate than going by filename order (or file number).
if isfield(slice_data(1), 'SliceLocation')
    [S,I] = sort([slice_data.SliceLocation]);
slice_data = slice_data(I);
end

pixelspacing = round(slice_data(true_index).PixelSliceThickness);
% Pre-allocate volume image array
[rows, cols] = size(slice_data(1).SliceData);
volume_image = ...
zeros(rows, cols, (length(slice_data)*pixelspacing));

% Build volume image array
h = waitbar(0,'Writing slice images to volume image array...','WindowStyle','modal');
for i = 1:length(slice_data)
    waitbar(i/N,h);
    volume_image(:,:,i*pixelspacing) = slice_data(i).SliceData;
    for j = 1:pixelspacing-1
        volume_image(:,:,i*pixelspacing)-j) = slice_data(i).SliceData;
    end
end
close(h);
a = size(volume_image);
num_slices = a(3);

% Threshold 3d-volume based on value from literature
middleslice = round(num_slices/2);
HU = 306;% Thresholdvalue in HU
Threshold = HU + 1024;% Thresholdvalue in grayscale. 1024 is from the dicominfo
T = volume_image > Threshold;
VI_T1 = volume_image .* T;
Threshold2 = Threshold;

% filling and smoothing bone surface
se = strel('disk',1);
SE = strel('line',3,0);
for i = 1:num_slices
    test = VI_T(:,:,i);
    test = imclose(test,SE);
    IR = imerode(test,se);
    VI_T(:,:,i) = IR;
end

%% Filtering & Splitting based on position of labels
figure(1), imshow(VI_T(:,:,middleslice),[])
title('Click inbetween tibia and femur')
[xtb, ytb, tbvalue] = impixel
close(1)

Label = bwconncomp(VI_T,18);
umlabels = max(size(Label.PixelIdxList));
L = zeros([rows,cols,num_slices]);
n = 1;
for i = 1:numlabels
    freq = max(size(Label.PixelIdxList{1,i}));
    if freq > 10000
        for j = 1:freq
            p = Label.PixelIdxList{1,i}(j);
            L(p) = n;
        end
        n = n + 1;
    end
end
numlabels = max(L(:));
FFL = L;
for i = 1:numlabels
    [Y1,X1] = find(L==i); % finding coordinates of areas
    MY1 = mean(Y1); % mean Y coordinate
    if MY1 < (ytb)% deleting areas below mean frequency
        L(L==i) = 0;
    end
end
% finding and deleting unwanted stuff
figure(1), imshow(L(:,:,middleslice))
title('Finding and deleting unwanted stuff, if none: click black')
pixel_values = impixel;
size_pixel_values = size(pixel_values);
close(1)
if pixel_values(1,1) ~= 0
    for i = 1:size_pixel_values(1)
        P = pixel_values(i,1);
        L(L==P) = 0;
    end
end
L = logical(L);
result = VI_T .* L;
213 % creating femur location
214 figure(1), imshow(FFL(:,:,middleslice))
215 title('Click femur')
216 pixel_values = impixel;
217 pixel_values = mean(pixel_values);
218 close(1)
219 h = waitbar(0, 'Creating femur location', 'WindowStyle', 'modal');
220 FL = zeros([10,10]);
221 for i = 1:num_slices
222    for j = 1:512
223       for k = 200:512
224          if FFL(513-k,j,i) == pixel_values
225             FL(j,i) = 513-k;
226             break
227          else
228             FL(j,i) = 0;
229          end
230       end
231    end
232    waitbar(i/num_slices,h)
233 end
234 FL(FL==0) = NaN;
235 close(h);
236 % finding place where condyles are separted
237 FL1 = uint16(FL);
238 b = bwlabel(FL1);
239 for i = 1:1000
240    flmin = min(FL1(FL1>0));
241    FL1(FL1==flmin) = 0;
242    b = bwlabel(FL1);
243    if max(b(:)) > 1
244       for j = 1:max(b(:))
245          if max(size(find(b==j))) < 100
246             [x,y] = find(b==j);
247             b(b==j) = 0;
248             FL1(x,y) = 0;
249          end
250       end
251    end
252    b = bwlabel(b);
253 end
254 if max(b(:)) > 1
255    figure(1), imshow(b,[])  
256    bweg = impixel;
257    bweg = mean(bweg);
258    if bweg == 0
259       close(1)
260       cut_off = min(FL1(FL1>0));
261       flmin = min(FL1(FL1>0));
262       FL1(FL1==flmin) = 0;
263       flmin = min(FL1(FL1>0));
264       FL1(FL1==flmin) = 0;
265       break
266    else
267       [x,y] = find(b==bweg);
268    end
269 end
for k = 1:max(size(x))
    FL1(x(k),y(k)) = 0;
end
end
end
condyles = logical(FL1);
condyles = imfill(condyles,'holes');

%% crop image based on cut off value
[X,Y] = find(L==1);
cut_off = cut_off - 15;
result_cropped = result(cut_off:end,:,:);
VI_cropped = volume_image(cut_off:end,:,:);

%% creating tibia plateaus for different layers.
info = dicominfo(full_path);
difY = (512-cut_off);
difX = 512;
num_pixels = 10;
pixpermm = round((slice_data(true_index).PixelSliceThickness)/(slice_data(true_index).SliceThickness));
h = waitbar(0,'Creating tibial plateau grayscale-map','WindowStyle','modal');
for i = 1:num_slices
    for j = 1:(difX)
        if result_cropped(k,j,i) == 0
            for l = 1:pixpermm
                pixels1(l) = VI_cropped((k+(l-1)),j,i);
pixels2(l) = VI_cropped((k+(l-1)+pixpermm),j,i);
pixels3(l) = VI_cropped((k+(l-1)+(2*pixpermm)),j,i);
pixels4(l) = VI_cropped((k+(l-1)+(3*pixpermm)),j,i);
pixels5(l) = VI_cropped((k+(l-1)+(4*pixpermm)),j,i);
            end
            t_plat_gray(j,i,1) = mean(pixels1);
t_plat_gray(j,i,2) = mean(pixels2);
t_plat_gray(j,i,3) = mean(pixels3);
t_plat_gray(j,i,4) = mean(pixels4);
t_plat_gray(j,i,5) = mean(pixels5);
t_plat_loc(j,i) = k;
            break
        else
            t_plat_gray(j,i,:) = NaN;
t_plat_loc(j,i) = NaN;
        end
    end
    waitbar(i/num_slices,h)
end
close(h)
H1=t_plat_gray(:,:,1);H2=t_plat_gray(:,:,2);H3=t_plat_gray(:,:,3);H4=t_plat_gray(:,:,4);
H5=t_plat_gray(:,:,5);
montage1 = [H1 H2 H3 H4 H5];
imshow(montage1,[])
%% deleting holes from tibia plateau location
TPL = 512 - t_plat_loc;

depth = 10;
cut_off1 = mode(TPL(:))-(depth*pixpermm);
TPL(TPL<cut_off1) = 0;
imshow(TPL,[])

%creating logical of tibia plateau
TPL(isnan(TPL))=0;
TPlogical = bwlabel(TPL,8);
um_labels = max(TPlogical(:));
for i = 1:num_labels
    freq(i) = max(size(find(TPlogical(TPlogical==i))));
end
maxfreq = find(freq==max(freq(:)));
TPlogical = logical(TPlogical);

%creating logical of condyles (lateral and medial)
TPG = t_plat_gray(:,:,1);
num_layers = 5;
b = bwlabel(condyles);
figure(1)
imshow(TPG,[])
figure(2)
imshow(b)
title('Select lateral condyle')
pixlatcon = impixel;
close([1 2])
pixlatcon = pixlatcon(1,1);
bm = b;
bm(bm==pixlatcon) = 0;
bm = logical(bm);
bl = b;
bl(bl==pixlatcon) = 0;
bl = logical(bl);
rest = ~condyles;
rest = logical(rest);

for i = 1:5
    TPG = t_plat_gray(:,:,i);
    TPG = TPG.*TPlogical;
    plateaus(:,:,i) = TPG;
end

%% Save data
fileloc = char(dicom_directory);
filename = [ fileloc '.mat' ];
save(filename, 'num_slices',...
    'volume_image',...
    'info',...
    'result_cropped',...
    'TPL',...
    'condyles',...
    'plateaus',...
% open saved data
clear all
voor = load(['VOOR.mat']);
a = load(['NA.mat']);

%% selecting which condyles you want to use
montage = [voor.condyles na.condyles];
figure(1)
imshow(montage,[])
title('Voor Na')
prompt = 'Which condyles do you want to use?? '
str = input(prompt,'s');
close(1)
if strcmp(str,'voor') == 1
    welke_is_wat = 'voor is fixed';
    condyles = voor.condyles;
    bm = voor.bm;
    bl = voor.bl;
    rest = voor.rest;
    Fixed = voor.plateaus;
    Moving = na.plateaus;
else
    welke_is_wat = 'na is fixed';
    condyles = na.condyles;
    bm = na.bm;
    bl = na.bl;
    rest = na.rest;
    Fixed = na.plateaus;
    Moving = voor.plateaus;
end

%% imregister before and after
for k = 1:5
    fixed = uint16(Fixed(:,:,k));
    moving = uint16(Moving(:,:,k));
    [optimizer, metric] = imregconfig('monomodal');
    MR(:,:,k) = imregister(moving,fixed,'affine',optimizer,metric);
end

%% calculating different values for fixed
for i = 1:5
    TPG = Fixed(:,:,i);
    A = TPG;
    TPGM = A.*bm;
    TPGL = A.*bl;
Rest = A.*rest;
TPGC = A.*condyles;

meantotal = round(mean(A(A>0)));
modustotal = mode(A(A>0));
MADtotal = round(nanmean(mad(A)));
skewtotal = skewness(A(:));
kurttotal = kurtosis(A(:));

meanmedial = round(mean(TPGM(TPGM>0)));
modusmedial = mode(TPGM(TPGM>0));
MADmedial = round(nanmean(mad(TPGM)));
skewmedial = skewness(TPGM(:));
kurtmedial = kurtosis(TPGM(:));

meanlateral = round(mean(TPGL(TPGL>0))); 
moduslateral = mode(TPGL(TPGL>0));
MADlateral = round(nanmean(mad(TPGL)));
skewlateral = skewness(TPGL(:));
kurtlateral = kurtosis(TPGL(:));

meanrest = round(mean(Rest(Rest>0))); 
modusrest = mode(Rest(Rest>0));
MADrest = round(nanmean(mad(Rest)));
skewrest = skewness(Rest(:));
kurtrest = kurtosis(Rest(:));

meancondyles = round(mean(TPGC(TPGC>0))); 
moduscondyles = mode(TPGC(TPGC>0));
MADcondyles = round(nanmean(mad(TPGC)));
skewcondyles = skewness(TPGC(:));
kurtcondyles = kurtosis(TPGC(:));

mean_values_fixed(i) = struct('meantotal',meantotal,'modustotal',modustotal,'MADtotal',MADtotal,
'skewtotal',skewtotal,'kurttotal',kurttotal,...
'meanmedial',meanmedial,'modusmedial',modusmedial,'MADmedial',MADmedial,'skewmedial',
skewmedial,'kurtmedial',kurtmedial,...
'meanlateral',meanlateral,'moduslateral',moduslateral,'MADlateral',MADlateral,'skewlateral',
skewlateral,'kurtlateral',kurtlateral,...
'meancondyles',meancondyles,'moduscondyles',moduscondyles,'MADcondyles',MADcondyles,
'skewcondyles',skewcondyles,'kurtcondyles',kurtcondyles,...
'meanrest',meanrest,'modusrest',modusrest,'MADrest',MADrest,'skewrest',skewrest,'kurtrest',
kurtrest);
end

%% calculating different values for movingregistered
MR = double(MR);
for i = 1:5
    TPG = MR(:,:,i);
    A = TPG;
    TPGM = A.*bm;
    TPGL = A.*bl;
    Rest = A.*rest;
    TPGC = A.*condyles;
meantotal = round(mean(A(A>0)));  
modustotal = mode(A(A>0));  
MADtotal = round(nanmean(mad(A)));  
skewtotal = skewness(A(:));  
kurttotal = kurtosis(A(:));  

meanmedial = round(mean(TPGM(TPGM>0)));  
modusmedial = mode(TPGM(TPGM>0));  
MADmedial = round(nanmean(mad(TPGM)));  
skewmedial = skewness(TPGM(:));  
kurtmedial = kurtosis(TPGM(:));  

meanlateral = round(mean(TPGL(TPGL>0)));  
moduslateral = mode(TPGL(TPGL>0));  
MADlateral = round(nanmean(mad(TPGL)));  
skewlateral = skewness(TPGL(:));  
kurtlateral = kurtosis(TPGL(:));  

meanrest = round(mean(Rest(Rest>0)));  
modusrest = mode(Rest(Rest>0));  
MADrest = round(nanmean(mad(Rest)));  
skewrest = skewness(Rest(:));  
kurtrest = kurtosis(Rest(:));  

meancondyles = round(mean(TPGC(TPGC>0)));  
moduscondyles = mode(TPGC(TPGC>0));  
MADcondyles = round(nanmean(mad(TPGC)));  
skewcondyles = skewness(TPGC(:));  
kurtcondyles = kurtosis(TPGC(:));  

%save calculated data to struct  
meanvalues_MR(i) = struct('meantotal',meantotal,'modustotal',modustotal,'MADtotal',MADtotal,  
'skewtotal',skewtotal,'kurttotal',kurttotal,...  
'meanmedial',meanmedial,'modusmedial',modusmedial,'MADmedial',MADmedial,'skewmedial',  
skewmedial,'kurtmedial',kurtmedial,...  
'meanlateral',meanlateral,'moduslateral',moduslateral,'MADlateral',MADlateral,'skewlateral',  
skewlateral,'kurtlateral',kurtlateral,...  
'meancondyles',meancondyles,'moduscondyles',moduscondyles,'MADcondyles',MADcondyles,  
skewcondyles',skewcondyles,'kurtcondyles',kurtcondyles,...  
'meanrest',meanrest,'modusrest',modusrest,'MADrest',MADrest,'skewrest',skewrest,'kurtrest',  

% renaming fixed and MR  
if strcmp(str,'voor') == 1  
MVvoor = meanvalues_fixed;  
MVna = meanvalues_MR;  
else  
MVvoor = meanvalues_MR;  
MVna = meanvalues_fixed;  
end  

%  
MR = double(MR);  
for i = 1:5  
for j = 1:size(Fixed,1)  

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for k = 1:size(Fixed,2)
    if Fixed(j,k,i) == 0
        diff(j,k,i) = 0;
    elseif MR(j,k,i) == 0
        diff(j,k,i) = 0;
    else
        if strcmp(str,'voor') == 1
            diff(j,k,i) = abs(Fixed(j,k,i) - MR(j,k,i));
        else
            diff(j,k,i) = abs(MR(j,k,i) - Fixed(j,k,i));
        end
    end
end
end

diff(isnan(diff)) = 0;
se = strel('disk',1);
for i = 1:5
    test = diff(:,:,i);
    IR = imerode(test,se);
    diff(:,:,i) = IR;
    diffc(:,:,i) = diff(:,:,i).*condyles;
end