

Imaging Knee MRI Segmentation and 3D Reconstruction Based on Deep Learning

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Abstract. Osteoarthritis is a chronic degenerative disease that causes knee pain and movement disorder in patients. There is still no drug to treat the disease, so scientists are working on the drugs. To test new drugs, medical images have played a vital role in medical treatment and diagnosis, and their analysis has become the top priority in clinical treatment. In medical image analysis, accurate segmentation of medical images plays a crucial role in its analysis. In recent years, with the continuous development of artificial intelligence, people have tried to use deep learning and convolutional neural networks to segment images to achieve the effect of reducing the workload of doctors. In this study, we do image segmentation using UNet network from three directions, and we use three 2D segmentation model from different directions to rebuild a 3D segmentation model to have an accurate and reproducible volumetric quantification of articular cartilage in the knee.

Keywords: Osteoarthritis, U-Net, Segmentation, Deep Learning, Knee, Artificial Intelligence, MRI Images, 3D Reconstruction

1. Introduction

Osteoarthritis is a chronic degenerative joint disease that causes knee pain and movement disorders in patients. According to the World Health Organization, about 355 million people suffer from osteoarthritis worldwide. The prevalence of osteoarthritis is 10% - 17% in people over 40 years old, 50% over 60, and 80% in those over 75. An early sign of OA is knee cartilage degeneration. Recent studies have shown that knee osteoarthritis is reversible if found in early stages of the biochemical changes in knee articular cartilage. In order to diagnose OA, doctors must confirm symptoms such as cartilage loss just as mentioned before. As a result, it is very important to monitor the volume of the cartilage. The traditional method for image-based knee cartilage assessment is having MRI images annotated by humans' slice by slice. This process is very time-consuming and labor-intensive. Meanwhile, manual annotation is also prone to errors caused by subjective factors of

personnel. Regardless, manual segmentation results is still the gold standard reference for deep learning segmentation algorithms [1].

At present, With the continuous development of artificial intelligence, people have tried to use deep learning and convolutional neural networks to segment images to achieve the effect of reducing the workload of doctors. The use of deep learning for image processing has become the current mainstream, and medical image segmentation has also become a successful example of deep learning application in the field of image processing with its unique application scenarios [2]. However, there are only 2D models after image segmentation, so the doctors cannot have an accurate and reproducible volumetric quantification of articular cartilage in the knee, and they cannot diagnose the condition of the knee joint precisely.

To solve this problem, we propose a 3D model combining three individual 2D networks to perform segmentation efficiently and accurately; Also, doctors can have an accurate volumetric quantification of the knee joint through the 3D model.

2. Review work

There have been many researches on image segmentation. This section elaborates on some image segmentation frameworks developed from the past to the present. TABLE 1 shows some common frameworks and describe their features and pros and cons.

Table 1. Deep learning segmentation network frameworks

Frame works	Features	Notes
FCN	Dense prediction, pixel-level segmentation	Duplicate convolutional calculations caused by overlapping image blocks are avoided
DeconvNet	Multiple deconvolutional layers and depooling layers are designed	Solved the size problem in the original FCN network, making the detailed information of the object more detailed
Deep Lab	The convolutional neural network is fused with the traditional probability graph model and uses empty convolution	Enlargement of the receptive field without evolutionary pooling prevents the loss of local information features of the image
SegNet	A maximum value pooled index method is proposed	The information lost during pooling can be obtained during the decoding phase by the maximum value index
PSPNet	A pyramid model is proposed to extract multi-scale information features of images	The fusion of detailed features and global features greatly enriches the semantic information of the image
U-Net	The codec network structure of hop splicing is used for feature fusion	The impact is far-reaching, and the U-Net network is widely used
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3. Material and methods

3.1. Network structure

In recent years, U-Net, as a fully convolutional network, has been the most common approach for medical image segmentation. However, existing 2D U-Net models can only analyze images from a single orientation, making it challenging to capture information from different anatomical planes. To address this limitation, we extended the U-Net to 3D by combining three orientation-based 2D U-

Nets in the axial, coronal, and sagittal orientations. Using this approach, we obtained predictions from three different orientations, and then weighted and combined these predictions to derive the optimal weights. This allowed us to gather information from multiple anatomical planes and improve the accuracy of our segmentation. The flow diagram for our algorithm is illustrated in Fig 1.

The U-Net architecture used in this study is a convolutional neural network composed of a Contracting Path for feature extraction and an Expansive Path for segmentation. The Contracting Path involves repeated 3x3 convolutions with ReLU activations and 2x2 max-pooling for down-sampling. The Expansive Path performs up-sampling, and each step includes up-convolution, feature concatenation, and additional convolutions. A 1x1 convolution at the final layer maps feature vectors to the desired output classes. This network utilizes VGG16 as the main feature extraction backbone, facilitating the use of pre-trained weights. All three networks share the same structure and parameter value. They use Adam optimizer with maximum learning rate of $1e - 4$ and trained for 100 epochs. The parameters and their values used for are listed in TABLE 2.

Table 2. Parameters

Parameter	Value
Backbone	VGG16 + Unet
Batch size	2
Optimizer	Adam
η	1E-04
β_1	0.9
β_2	0.999
ϵ	1E-08
Weight initialization	Pre-train
Activation	Softmax
Loss	Soft Dice

3.2. Dataset and dataset processing

In this study, the dataset consisting of three-dimensional knee MRI from 88 retrospective patients at two time points (baseline and 1-year follow-up) with ground truth articular cartilage segmentations was standardized. The original data, stored with a DICOM format, are of 384*384*160 pixels each image.

The original data is stored in a DICOM format. Since the U-Net model we choose to use can only be trained in 2 dimensions, we must transform three-dimensional MRI DICOM files into two-dimensional images for training as shown in figure 1. Each knee has 160 slices in coronal direction and 192 slices in axial and sagittal direction. So, the number of the slices is 5120 in coronal direct, and 6144 in axial and sagittal direct. This dataset contains labeled images with corresponding ground truth segmentation masks. The MRI image is transformed from DICOM format into a PNG format image, which is of 384*384 (or 384*160) pixels per slice.

3.3. Training

The ultimate goal for this research is to create a model that takes three multi-channel 2D image slices of x size $w \times h$ and outputs a 3D probabilistic segmentation map of size $x \times w \times h$ for predicting K class.

To begin the training process, a pre-trained U-Net model is employed as the backbone architecture. This model has been previously trained on a large dataset to capture general image features. And then we replace the weight of the model with new weight to accommodate the cartilage segmentation task during the training.

To optimize training, this study employed a dynamic learning rate scheduler. The learning rate is adjusted during training epochs, enabling the network to efficiently navigate the optimization landscape.

To improve efficiency, this study trained three directions on three computers at the same time and sent the results of the three directions to one computer after the training [3].

3.4. Combination and evaluation

We apply the models to their corresponding testing dataset and obtain the cartilage probability of each pixel. For example, if the model is trained to predict dataset sliced along coronal. The probability data is stored as two-dimensional matrix, represented as C_n . Then these 2D matrixes are piled up along coronal to form a 3D matrix that store the cartilage distribution in 3D space, represented as C . Repeat the same process three times, we can get three 3D matrixes of same size $384 \times 384 \times 160$, represented as C, A, S .

Next, we combine the three matrixes through linear blending at the weight of (α, β, γ) to obtain the final prediction probability matrix $F = \alpha C + \beta A + \gamma S$.

To further analyze and compare the performance of different models, we conducted dice coefficient analysis on the predicted probability matrix Y to the ground truth matrix X to see if probability linear blending improved the accuracy of the prediction. The four outcomes can be formulated in a 2×2 confusion matrix (TP, FN, FP, and TN) where each row of the matrix represents the instances in an actual class while each column represents the instances in a predicted class. The Mean Intersection-over-Union (MioU) and Dice similarity coefficient (DSC) are defined as follows:

$$MIoU = \frac{1}{k+1} \sum_{i=0}^k \frac{TP}{TP + FN + FP}$$

$$Dice(X, Y) = \frac{2|X \cap Y|}{|X| + |Y|} = \frac{2TP}{2TP + FN + FP}$$

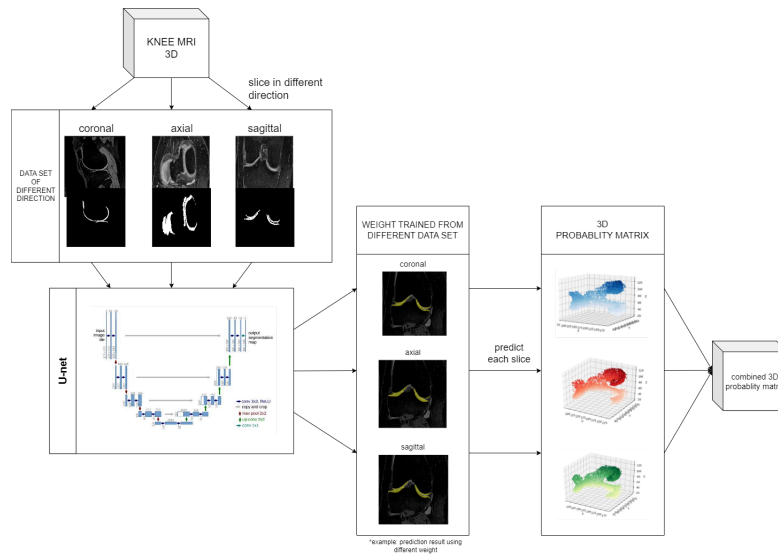


Figure 1. Flow diagram for 3D cartilage distribution prediction

4. Results

The three deep learning models are trained to their best performance on three different datasets, with their MIOU being 83.54%, 77.93% and 72.85% respectively. After piling up each slice, the cartilage distribution in 3D space is shown in Figure 2.

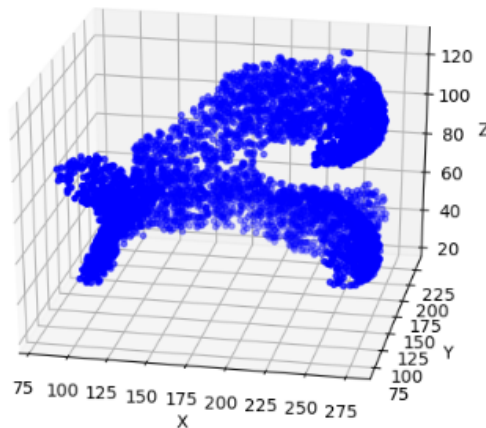


Figure 2. Cartilage distribution in 3D space

The final fusion model is a linear blending of the prediction values using the weights of (0.5077, 0.4012, 0.0911). This linear blending weight should be slightly altered to fit different models. After applying each model to 3D MRI images, we conducted further analysis using evaluation metrics including confusion Matrix and dice coefficient. The discrepancy between predicted probability and ground truth matrix of the four model can be shown using confusion matrixes. Dice coefficient is introduced for a clearer accuracy measurement in 3D space. The results are shown in Fig. 5.

We can tell from the table that the fusion model predicts the highest number of accurate cartilage pixels with 254,158 and the lowest number of wrong cartilage pixels with 18045. Fusion model achieves the highest dice coefficient at 0.9129, slightly surpassing the coronal plane model at 0.9103, overwhelming the sagittal plane model at 0.8759 and axial plane model at 0.8429. This

shows the fusion model has less miss-typed pixel and achieves higher accuracy than all the individual plane models [4].

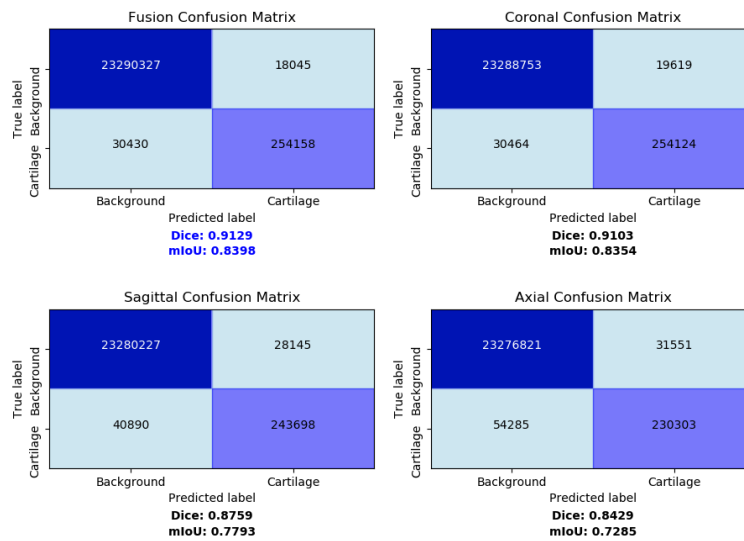


Figure 4. Confusion matrixes and dice coefficient of different models

5. Discussion

All three individual plane models are expected to achieve the same accuracy. However, we can't find good explanation for the bad performance of axial plane model. Why the dice coefficient of axial plane model is so much lower than the other two needs further discussion. The problem may lie in the original data set, or the structure of the cartilage itself. That is the assumption that certain planes of the cartilage just don't give good features for neural network to learn. Therefore, when doing linear blend, weight of these planes should be reduced modestly.

In this paper, our 3D segmentation model is actually a combination of three 2D segmentation models. The original intention is to make up the connection between different planes that the individual 2D model is likely to ignore. If the models are from multiple axes than three, the accuracy of the final fusion model is expected to improved. Moreover, there are now 3D U-net model that does 3D convolution to the MRI image. This allows the network to study the connection between plane while training, but requires a large more effort to train. Further research can compare the pros and cons of different segmentation method [5].

6. Conclusion

In conclusion, this study demonstrates the potential of deep learning algorithms in accurately performing segmentation on knee MRI images for the early detection of knee osteoarthritis. By utilizing a U-Net model and combining three individual 2D networks, the researchers were able to achieve efficient and accurate segmentation of medical images, reducing the workload of doctors and providing an accurate volumetric quantification of articular cartilage in the knee. The fusion model, in particular, outperformed the individual plane models, with a dice coefficient of 0.9129, indicating higher accuracy. However, it is important to acknowledge the limitations of this study, such as the need for a larger dataset and further research to understand the discrepancies in model

performance. Overall, the findings highlight the potential of deep learning and segmentation algorithms in aiding the diagnosis of knee osteoarthritis and suggest further exploration in this field.

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Bowen Luan, Keran Xu, Liyuan Zhou, Yaxing Zhang and Jiayu Jin contributed equally to this work and should be considered co-first authors. The authors would like to thank all colleagues and students who contributed to this study.

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