



UMRFormer-net: a three-dimensional U-shaped pancreas segmentation method based on a double-layer bridged transformer network

Kun Fang^{1,2}, Baochun He², Libo Liu², Haoyu Hu³, Chihua Fang^{3,4}, Xuguang Huang¹, Fucang Jia^{2,4,5}

¹School for Information and Optoelectronic Science and Engineering, South China Normal University, Guangzhou, China; ²Research Center for Medical Artificial Intelligence, Shenzhen Institute of Advanced Technology, Chinese Academy of Sciences, Shenzhen, China; ³Department of Hepatobiliary Surgery (I), Zhujiang Hospital of Southern Medical University, Guangzhou, China; ⁴Pazhou Lab, Guangzhou, China; ⁵Shenzhen Key Laboratory of Minimally Invasive Surgical Robotics and System, Shenzhen Institute of Advanced Technology, Chinese Academy of Sciences, Shenzhen, China

Contributions: (I) Conception and design: K Fang, F Jia; (II) Administrative support: X Huang, F Jia; (III) Provision of study materials or patients: H Wang, C Fang; (IV) Collection and assembly of data: H Wang, B He, F Jia; (V) Data analysis and interpretation: K Fang, B He, L Liu; (VI) Manuscript writing: All authors; (VII) Final approval of manuscript: All authors.

Correspondence to: Fucang Jia. Shenzhen Institute of Advanced Technology, Chinese Academy of Sciences, 1068 Xueyuan Avenue, Xili University Town, Nanshan District, Shenzhen 518055, Guangdong, China. Email: fc.jia@siat.ac.cn; Xuguang Huang. School for Information and Optoelectronic Science and Engineering, South China Normal University, 55 West of Zhongshan Avenue, Tianhe District, Guangzhou 510631, Guangdong, China. Email: huangxg@sncu.edu.cn.

Background: Methods based on the combination of transformer and convolutional neural networks (CNNs) have achieved impressive results in the field of medical image segmentation. However, most of the recently proposed combination segmentation approaches simply treat transformers as auxiliary modules which help to extract long-range information and encode global context into convolutional representations, and there is a lack of investigation on how to optimally combine self-attention with convolution.

Methods: We designed a novel transformer block (MRFormer) that combines a multi-head self-attention layer and a residual depthwise convolutional block as the basic unit to deeply integrate both long-range and local spatial information. The MRFormer block was embedded between the encoder and decoder in U-Net at the last two layers. This framework (UMRFormer-Net) was applied to the segmentation of three-dimensional (3D) pancreas, and its ability to effectively capture the characteristic contextual information of the pancreas and surrounding tissues was investigated.

Results: Experimental results show that the proposed UMRFormer-Net achieved accuracy in pancreas segmentation that was comparable or superior to that of existing state-of-the-art 3D methods in both the Clinical Proteomic Tumor Analysis Consortium Pancreatic Ductal Adenocarcinoma (CPTAC-PDA) dataset and the public Medical Segmentation Decathlon dataset (self-division). UMRFormer-Net statistically significantly outperformed existing transformer-related methods and state-of-the-art 3D methods ($P < 0.05$, $P < 0.01$, or $P < 0.001$), with a higher Dice coefficient (85.54% and 77.36%, respectively) or a lower 95% Hausdorff distance (4.05 and 8.34 mm, respectively).

Conclusions: UMRFormer-Net can obtain more matched and accurate segmentation boundary and region information in pancreas segmentation, thus improving the accuracy of pancreas segmentation. The code is available at <https://github.com/supersunshinefk/UMRFormer-Net>.

Keywords: Pancreas; image segmentation; transformer; deep learning; U-Net

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Introduction

Computer vision and artificial intelligence applications are becoming increasingly important, especially in the field of medical imaging technology (1). In medical imaging, accurate organ segmentation is a prerequisite and an important factor in many subsequent image computing-based analyses. With the rapid development of deep neural networks, automatic segmentation has achieved good results in many organs and tissues, including the brain, lung, liver, and heart (2). For the pancreas, however, the segmentation accuracy is still relatively low, despite considerable improvements in results during the pre-deep learning era (3).

In recent years, deep learning methods based on convolutional neural networks (CNNs) have achieved remarkable success in medical image segmentation. Based on the original two-dimensional (2D) U-Net and three-dimensional (3D) U-Net architectures, Isensee *et al.* (4) proposed nnU-Net, a robust and adaptive deep learning framework. This method focuses on the configuration of preprocessing (resampling and normalization), training strategy, inference, and post-processing, which allows the network to achieve a good improvement in segmentation accuracy. The network provides an efficient and accurate solution for solving any new task in the biomedical field, including automatic segmentation of the pancreas and pancreatic tumors in clinical medicine, and has also been applied in the pancreatic segmentation competition of the Medical Segmentation Decathlon (MSD) (5). Further, Chen *et al.* (6) proposed a model-driven deep learning segmentation method for the pancreas based on helical transformation. They investigated a uniformly sampled helical transformation algorithm that maps a 3D image onto a 2D plane and preserves the spatial relationship between textures. The method effectively deals with 3D context information in 2D models. Salanitri *et al.* (7) proposed a novel 3D CNN for automatic segmentation of the pancreas from magnetic resonance and computed tomography (CT) images which learns to extract and decode volumetric features at different scales in the encoder-decoder structure. Most of the models in the abovementioned pancreatic segmentation-related studies were conducted from the

perspective of spatial locality features.

Recently, transformer-based models (8,9) have attracted much attention in computer vision after achieving state-of-the-art benchmarks in natural language and text processing areas. They have a strong ability to model long-range dependencies and capture global context. However, unlike the local formulation of convolutions, the transformer focuses more on global semantic information as it encodes images as a sequence of one-dimensional (1D) patch embeddings and utilizes self-attention modules to learn a weighted sum of values calculated from hidden layers (10). For example, vision transformer architecture achieved outstanding performances in image classification via pre-training on large-scale datasets (11,12). Subsequently, many transformer-based architectures (13,14) have been explored for dense prediction tasks. Peiris *et al.* (15) proposed the transformer architecture VT-Unet for volumetric medical image segmentation. The encoder has two consecutive self-attention layers to simultaneously encode local and global cues, and the decoder has novel parallel shifted window-based self- and cross-attention blocks to capture fine details. Although the aforementioned methods provide reliable technical guidance, pancreas segmentation remains a challenging task. The difficulty is mainly due to the following aspects: (I) the shape, size, and position of the pancreas in the abdomen varies greatly among patients, which makes it difficult to use reliable prior to improve the delineation procedure; (II) the contrast between the pancreas and surrounding tissue is weak; (III) the pancreas occupies only a very small portion of the CT image; and (IV) the pancreas is relatively soft and can easily be pushed by surrounding tissues with similar intensities, which could result in significant deformation. Although CNN-based methods for automatic pancreas segmentation have excellent representational capabilities, it is difficult to build an explicit long-distance dependence due to the limited receptive fields of convolution kernels. This limitation of convolution operation poses challenges to the learning of global semantic information, which is critical for dense prediction tasks such as segmentation. Existing models utilize convolution to focus more on local features of the pancreatic region, ignoring the global features associated with the pancreas and surrounding tissues; this obfuscates

the border between the pancreas and surrounding tissues.

To overcome these challenges, learning from the transformer idea, we contemplated whether it is possible to design a module that pays deep attention to global long-range dependencies and local feature information to be embedded in the CNN network. Therefore, we proposed a model-driven deep learning method based on UMRFormer-Net for pancreas segmentation. A double transformer-like module named MRFormer was designed to help extract the deep feature information generated by the encoder. We connected two layers of residual units, composed of a 3×3 depthwise convolutional block and a post-layer normalization block after the multi-head self-attention (MHSA) layer inside the module, to achieve the advantage of endogenously integrating the local features of the convolution and the global long-range dependence features of the transformer. This method models the local and global dependencies between features of different scales in the bridging encoder. Furthermore, an MHSA mechanism and a residual depthwise convolution combination block was introduced on the path from the encoder's deep feature input to the skip connection to facilitate feature aggregation at different semantic scales on the decoder sub-network and subsequently, highly flexible feature fusion schemes for differences in coarseness and fine grains.

The main contributions of this paper are as follows:

- (I) For the 3D deep learning segmentation model, we propose a transformer-like module that extracts feature information at different scales on the last two deep layer features of U-Net, which can obtain more effective deep feature context information in the 3D model.
- (II) We propose an MRFormer module that aims to extract spatially fine-grained local information from long sequence feature outputs. It has the advantage of maintaining long-range dependencies, as well as feature recognition and local context capabilities.
- (III) Compared with existing transformer-related methods, our method achieved good segmentation results in CT pancreas datasets from the Clinical Proteomic Tumor Analysis Consortium Pancreatic Ductal Adenocarcinoma (CPTAC-PDA) Pancreas dataset (16) and the public MSD dataset (self-division) (5), which showed it to be superior to the existing methods.

Methods

The study was conducted in accordance with the Declaration of Helsinki (as revised in 2013). In this section, we first describe the overall pipeline and specific structure of UMRFormer-Net, and then show the details of the basic unit (MRFormer) of UMRFormer-Net.

Architecture

As shown in *Figure 1*, the proposed UMRFormer-Net has a U-shaped encoder-decoder architecture with a hybrid CNN and transformer, and adopts a double MRFormer context bridge module at the bottom of the whole network. Specifically, we generally assume that input a 3D image $X \in \mathbb{R}^{C \times H \times W \times D}$ with a spatial resolution of $H \times W$, depth dimension of D (# of slices), and C channels (# of modalities). UMRFormer-Net first utilizes a five-layer 3D CNN to generate feature maps at different scales to capture spatial and deep features, and then utilizes a double transformer-like module to encode the long-range dependencies semantic information of the fourth- and fifth-layer feature maps in a global space. Next, the upsampling layer and the convolution layer are repeatedly stacked to gradually obtain high-resolution segmentation results. Next, we describe the components of UMRFormer-Net in detail.

Network encoder

Feature embedding of the double MRFormer encoder

Given the feature maps M_1 and M_2 are output by the fourth and fifth layers of the network encoder. We used linear projection (3×3×3 convolution layers) to increase the channel size of the fourth and fifth layers from $K_1=128$ and $K_2=256$ to $d_1=512$ and $d_2=1024$. Since the double MRFormer module requires a sequence as input separately, we uniformly compressed the spatial and depth dimensions of the input 3D feature map into 1D, and obtained $d_1 \times Q_1$ ($Q_1 = \frac{H}{8} \times \frac{W}{8} \times \frac{D}{8}$) and $d_2 \times Q_2$ ($Q_2 = \frac{H}{16} \times \frac{W}{16} \times \frac{D}{16}$) feature maps (f_1 and f_2 , respectively), which can be regarded as $Q_1 d_1$ -dimensional and $Q_2 d_2$ -dimensional labels, respectively. The input 3D feature map was compressed into 1D and then added to a learnable positional encoding, creating feature embedding, as follows:

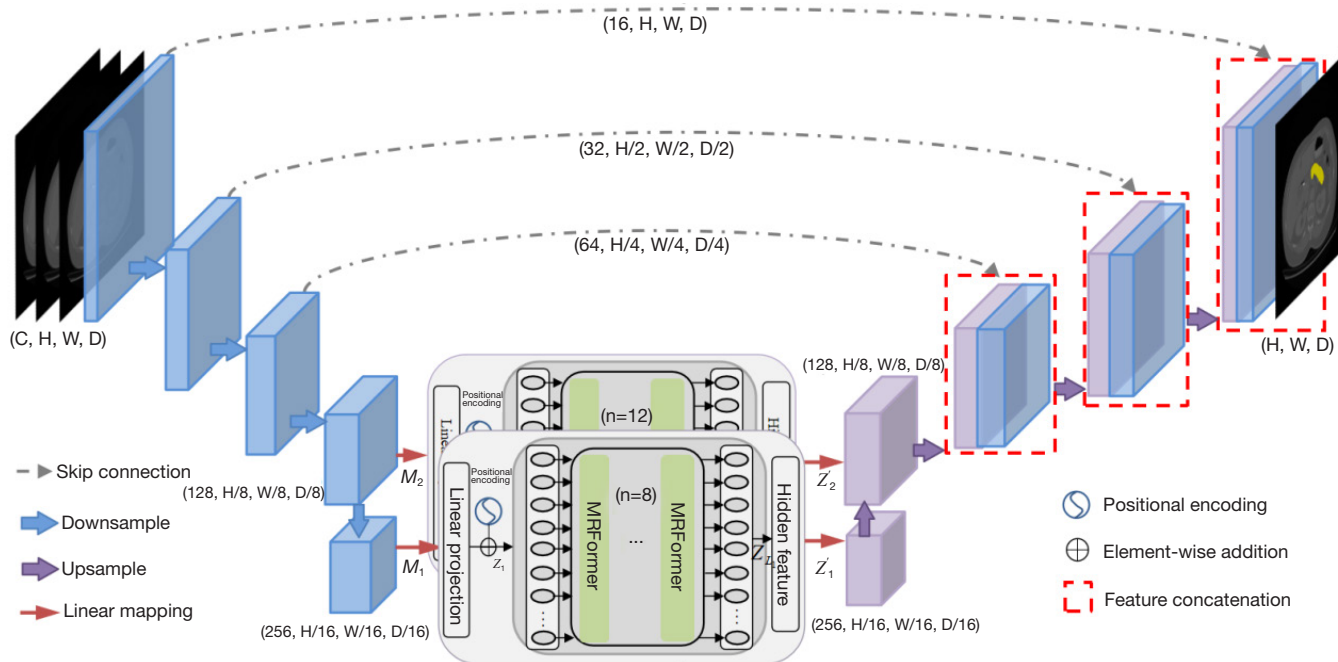


Figure 1 Overall architecture of the proposed UMRFormer-Net, which presents details of the double MRFormer layers.

$$z_1 = f_1 + PE_1 = LP_1 \times M_1 + PE_1 \tag{1}$$

$$z'_\ell = LN(MHA(z_{\ell-1})) + z_{\ell-1} \tag{3}$$

$$z_2 = f_2 + PE_2 = LP_2 \times M_2 + PE_2 \tag{2}$$

$$z_\ell = LN_2(DConv_2(LN_1(DConv_1(z'_\ell)))) + z'_\ell \tag{4}$$

where LP_1 and LP_2 are the linear projection operation, $PE_1 \in R^{d_1 \times Q_1}$ and $PE_2 \in R^{d_2 \times Q_2}$ denote the positional encoding, and $z_1 \in R^{d_1 \times Q_1}$ and $z_2 \in R^{d_2 \times Q_2}$ refer to the feature embedding.

MRFormer block

As shown in *Figure 2*, the MRFormer block consists of MHSA (8,17) and RDCB modules. The RDCB module mainly consists of two depthwise convolutions (18,19) with layer normalization after each layer. We also added a dense residual connection between each layer. Compared with the original transformer, which used only a feed-forward network, we proposed to use an additional two depthwise convolution layers. This was conducted with the aim of focusing on local receptive fields and to attract 2D or 3D spatial features which cannot be captured by 1D vectors. The layer normalization operations produced milder activation value at the network layers, facilitating more stable model training. The output of the ℓ -th ($\ell \in [1,2,\dots,L]$) transformer layer can be calculated by:

where $LN_{(\cdot)}$ denotes the layer normalization, z_ℓ is the output of the ℓ -th MRFormer layer, and $DConv_{(\cdot)}$ denotes the depthwise convolution. Similar to previous works (20,21), the calculation formula of self-attention in MHSA (8,17) is as follows:

$$head_i = Attention(Q_i, K_i, V_i) = SoftMax\left(\frac{Q_i K_i^T}{\sqrt{d_i}}\right) V_i, \quad i = 1, 2, \dots \tag{5}$$

$$MultiHead(Q, K, V) = Concat(head_1, head_2, \dots) W^O \tag{6}$$

where $Q_{(\cdot)}, K_{(\cdot)}, V_{(\cdot)} \in R^{M^2 \times d_{(\cdot)}}$ denote the query, key, and value matrices, respectively; M^2 and $d_{(\cdot)}$ represent the number of patches in a window and the dimension of the query or key, respectively; and $W^O \in R^{d_{(\cdot)} \times d_{(\cdot)}}$ represents the weight matrix of the dot product.

Network decoder

In order to match the input size of the decoder, it was

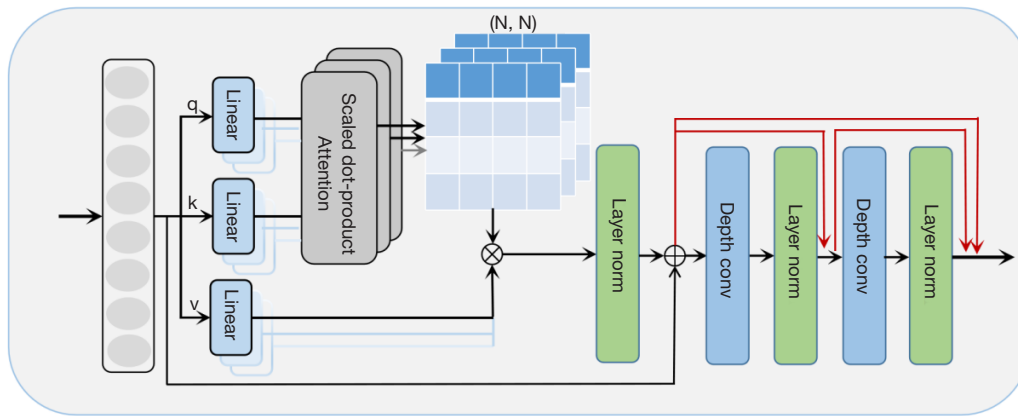


Figure 2 The MRFormer module for deep fusion of global and local features. It consists of MHSA and RDCB modules. MHSA, multi-head self-attention; RDCB, residual depthwise convolutions block.

necessary to map the feature sequence output by the double MRFormer module into a 3D feature map. Specifically, the double MRFormer $Z_{L_1} \in R^{d_1 \times Q_1}$ and $Z_{L_2} \in R^{d_2 \times Q_2}$ were reshaped to $d_1 \times \frac{H}{8} \times \frac{W}{8} \times \frac{D}{8}$ and $d_2 \times \frac{H}{16} \times \frac{W}{16} \times \frac{D}{16}$, respectively. After feature mapping, upsampling and convolution were performed on the outputs of Z_1' and Z_2' , respectively, and then fused with encoder features via long-range skip connections in order to capture finer semantic and fine-grained information as well as richer spatial details.

Results

MSD Pancreas and CPTAC-PDA Pancreas Datasets

The model was validated in pancreas segmentation experiments. For a fair comparison with previous transformer-based architectures, we evaluated our methods on the public MSD dataset (self-division) and the CPTAC-PDA Pancreas dataset.

For the MSD dataset, the original training datasets with open ground truth segmentation consist of 281 3D CT volumes, divided into 169, 56, and 56 at a 6:2:2 ratio, and is used for training, validation, and testing of pancreas segmentation. The number of slices ranges from 36 to 174, and the CT images are 512x512 in size. The labels contain two categories: background (label 0) and pancreas (label 1).

CPTAC-PDA Pancreas dataset consists of reference segmentations for 91 abdominal CT images delineating

the pancreas and pancreas adenocarcinoma (PDA). The 91 CT images are divided into 55, 18, and 18 for training, validation, and testing, respectively. The labels contain two categories: background (label 0) and pancreas (label 1).

Evaluation metrics

To compare the performance between different models in our experiments, the evaluation metrics used to measure the segmentation effect of medical images included the Dice coefficient (22) and 95% Hausdorff distance (23). Mean and standard deviation (SD) indicate the dispersion of the results of the sample data, respectively. With I_{seg} and I_{GT} representing the segmentation image and the ground truth segmentation, respectively, the Dice formula can be written as follows:

$$Dice = \frac{2(I_{seg} \cap I_{GT})}{I_{seg} + I_{GT}} \tag{7}$$

Regarding set X and set Y, the maximum Hausdorff distance represents the maximum distance of one point set to the nearest point of another point set, defined as:

$$d_H(X, Y) = \max\{d_{XY}, d_{YX}\} \tag{8}$$

$$d_{XY} = \max_{x \in X} \left\{ \min_{y \in Y} \|x - y\| \right\} \tag{9}$$

$$d_{YX} = \max_{y \in Y} \left\{ \min_{x \in X} \|y - x\| \right\} \tag{10}$$

Table 1 Pre-input data preprocessing configuration according to the characteristics of the datasets

Data processing mode	UMRFormer-Net
Target spacing	0.73 mm × 0.73 mm × 3.0 mm
Median image size	145×512×512
Crop size	64×128×128
Batch size	2
Stride of downsampling	[2 2 2], [2 2 1], [2 2 2], [2 2 2], [2 2 2]

Table 2 Comparison of segmentation results in Dice score and 95% HD in the MSD dataset

Method	Pancreas	
	Dice score ↑	95% HD ↓
3D U-Net	72.69±0.08***	9.45±8.68**
Trans UNet	62.84±0.13**	16.22±11.75
Swin UNet	63.72±0.32	14.89±10.24*
nnFormer	65.86±0.18**	13.92±9.05**
VT-UNet-B	70.04±0.26	10.13±5.24***
nnU-Net	76.88±0.09*	8.62±7.26**
UMRFormer-Net (ours)	77.36±0.11	8.34±6.69

Data were presented as mean ± SD. Paired t-test was used to test for the difference between our method and other methods in Dice score, 95% HD individually, *P<0.05, **P<0.01, ***P<0.001. MSD, Medical Segmentation Decathlon; SD, standard deviation; HD, Hausdorff distance.

The 95% Hausdorff distance is the 95th percentile of the distances between the points of sets X and Y.

Implementation details

We performed all experiments using Python 3.8 (Guido van Rossum, Netherlands) and Pytorch 1.8.0 (FAIR, Menlo Park, USA) on Ubuntu 18.04 (Canonical, Mann, UK). All training procedures were performed on a single NVIDIA RTX 3090 GPU (Nvidia, Santa Clara, CA, USA) with 24 GB memory for 1000 epochs using a batch size of 2. The initial learning rate was set to 0.002 with a poly learning rate strategy, in which the initial rate decayed by each iteration with power 0.9. To train the model, the SGD optimizer was adopted (24). The weight decay was set to 1e-5. We utilized the sum of the Dice loss and cross-entropy loss.

Pre-processing and augmentation strategies

All CT images were resampled to the same spacing (0.73 mm × 0.73 mm × 3.0 mm). During the training process, the augmentation strategy was implemented in the following order: rotation, scaling, Gaussian blurring, brightness and contrast adjustment, simulation of low resolution, gamma augmentation, and mirroring.

Network configurations

In *Table 1*, we show experimental network configurations on the CPTAC-PDA Pancreas dataset and the Public MSD dataset.

Experiments on the MSD Pancreas dataset

To investigate the segmentation performance of our model, we conducted experiments on the MSD Pancreas dataset, comparing our model's performance with those of the 3D U-Net (25), Swin-UNet (26), Trans-UNet (27), nnFormer (28), and VT-UNet (15) models. The best-performing existing engineering method, nnU-Net (4), was also used as a reference for segmentation accuracy. We directly used the results obtained using these methods for comparison. For a fair comparison, we divided the datasets according to VT-UNet-B (15). At the time of inference, we generated multi-class predictions using a sliding window technique used in previous papers (29,30). As shown in *Table 2*, the quantitative experimental results of this model suggest that our approach obtained a better segmentation performance than all the baselines, as measured by either the Dice coefficient or 95% Hausdorff distance. The differences were statistically significant. Additionally, UMRFormer-Net obtained dominance over the 95% Hausdorff distance of the pancreas with our redesigned MRFormer block.

For qualitative analysis, *Figure 3* displays a visual comparison of the pancreas segmentation results of various methods. UMRFormer-Net obtained more matched and accurate segmentation boundary and region information, which improved the accuracy of pancreas segmentation.

Experiments on CPTAC-PDA Pancreas dataset

To better demonstrate the generalization ability and efficiency of our model, we further evaluated its performance in another public dataset, CPTAC-PDA Pancreas. We also compared our method with

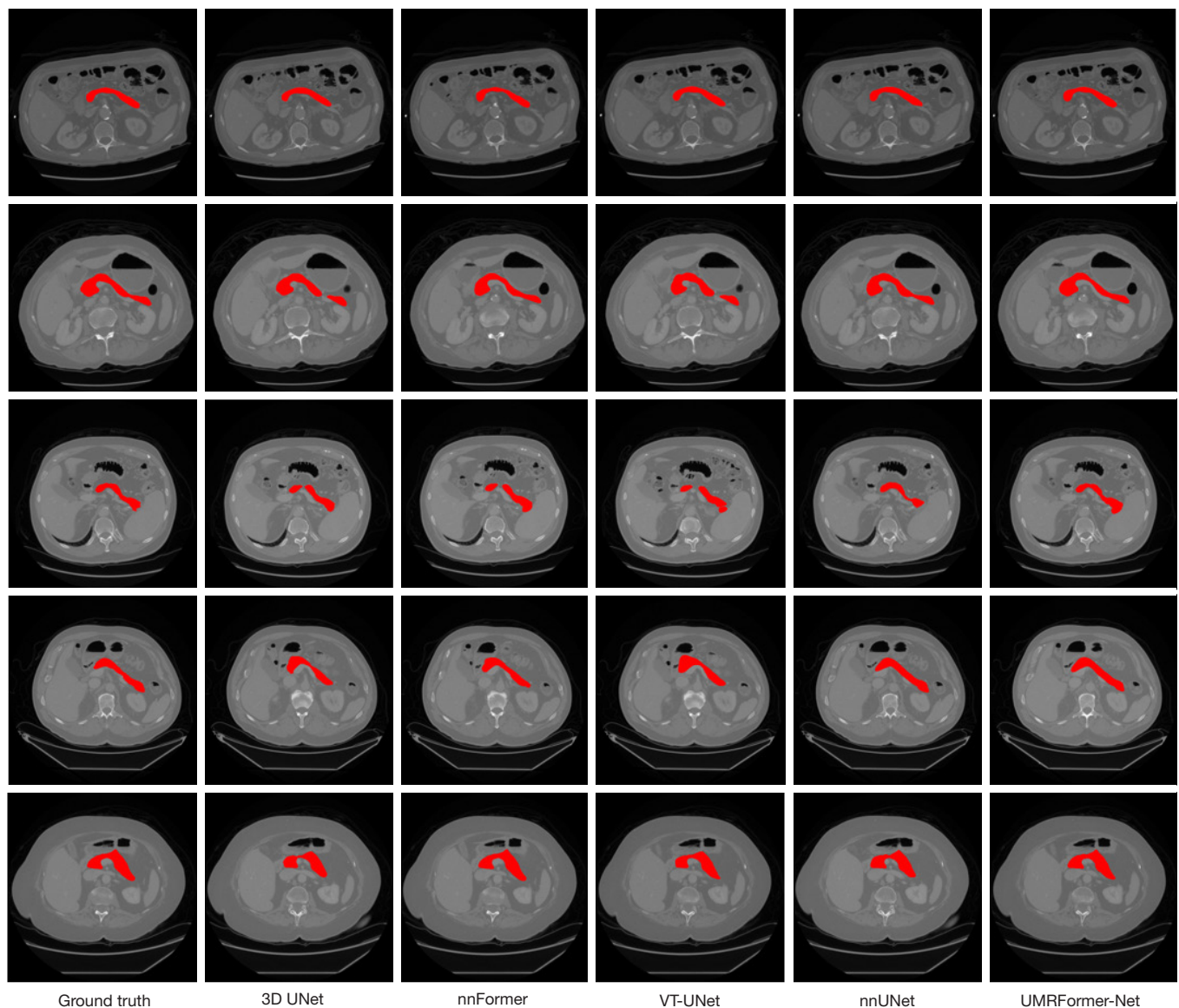


Figure 3 Visual comparison of CT pancreas segmentation results between different methods. CT, computed tomography.

other existing methods including 3D U-Net (25), Swin-Unet (26), Trans-Unet (27), nnFormer (28), VT-UNet (15), and nnU-Net (4), as shown in *Table 3*. The results showed that our proposed UMRFormer-Net achieved state-of-the-art accuracy compared with the classical 3D U-Net and other transformer-based methods.

Ablation study

We conducted extensive ablation experiments to validate the effectiveness of the proposed transformer-like decoder

layers and the necessity of using transformer-like at the skip connections of the last two layers of the encoder and decoder. The experimental results were yielded by five cross-validation evaluations on the CPTAC-PDA Pancreas dataset. First, we analyzed the impact of different layers of MRFormer decoders. Further, we studied the effect of the original transformer and the improved MRFormer module on the segmentation performance. Different from the native transformer module, the MRFormer module is composed of a multi-head attention module and two convolution layers. Finally, we studied and compared the impact of the addition

of MRFormer on the segmentation performance of the base 3D CNN network.

Depth of the double MRFormer (n)

As shown in *Figure 1*, we conducted ablation experiments to

Table 3 Comparison of segmentation results in Dice score and 95% HD in the CPTAC-PDA Pancreas dataset

Method	Pancreas	
	Dice score ↑	95% HD ↓
3D U-Net	81.43±0.03**	7.31±6.52*
Trans UNet	79.95±0.07*	9.40±10.36
Swin UNet	80.92±0.08	8.24±9.65**
nnFormer	82.46±0.05**	6.35±8.34
VT-UNet-B	83.21±0.06**	5.78±6.42***
nnU-Net	85.06±0.02**	4.63±5.84*
UMRFormer-Net (ours)	85.54±0.04	4.05±5.15

Data were presented as mean ± SD. Paired *t*-test was used to test for differences between our method and other methods in Dice score and 95% HD individually, **P*<0.05, ***P*<0.01, ****P*<0.001. CPTAC-PDA, Clinical Proteomic Tumor Analysis Consortium Pancreatic Ductal Adenocarcinoma; SD, standard deviation; HD, Hausdorff distance.

Table 4 Ablation study on the number of double MRFormer (n)

Depth of double MRFormer (n)	Pancreas	
	Dice score ↑	95% HD ↓
{{fourth,14}, {fifth,10}}	81.86±0.05	7.28±8.34
{{fourth,12}, {fifth,8}}	85.54±0.04	4.05±5.15
{{fourth,10}, {fifth,6}}	84.88±0.03	6.15±6.92
{{fourth,8}, {fifth,4}}	83.01±0.04	7.13±7.96

Data were presented as mean ± SD. HD, Hausdorff distance; SD, standard deviation.

Table 5 Ablation study on compare native and improved transformers

Native and improved transformer	Configuration of double-layer embedded	Pancreas	
		Dice score ↑	95% HD ↓
Transformer	{{fourth,512}, {fifth, 1024}}	84.31±0.06	5.65±6.74
MRFormer		85.54±0.04	4.05±5.15

Data were presented as mean ± SD. HD, Hausdorff distance; SD, standard deviation.

verify the effect of the depth of the double-layer MRFormer (n) on the segmentation performance.

As shown in *Table 4*, the optimal depth for the MRFormer module was 12 and 8 for the fourth and fifth layers, respectively. A greater depth did not necessarily improve the performance.

Additionally, we investigated the effects of the original transformer and the improved MRFormer module on the segmentation performance. Different from the native transformer module, the MRFormer module is composed of a multi-head self-attention module and two convolution layers. We set the convolution layers in MRFormer to two layers, considering the memory constraints and computational efficiency.

Comparison of the native and improved MRFormer module

We compared the impact of the double native transformer and the double improved transformer in performing network segmentation. The embedded dimensions input to the double transformer-like module in a given network were as follows: the fourth and fifth layers of the network were set to 512 and 1024, respectively. Of note, replacing the feed-forward network layer with two residual units combined with depthwise convolution and layer normalization made the network pay more attention to its local feature information while ensuring global semantic information of long feature sequences, which significantly improved the model's performance. The specific results are shown in *Table 5*. Compared with the native transformer module, our MRFormer module had better segmentation accuracy in the CPTAC-PDA Pancreas dataset.

Transformer-like decoder layers

Our method is based on a five-layer symmetric 3D CNN encoder-decoder architecture. *Table 6* presents a comparison of transformer-like ablation experiments for different layers. It counts from the bottom layer of the network and

Table 6 Ablation study on transformer-like decoder layers

Transformer-like decoder types	Position of transformer-like module	Pancreas	
		Dice score ↑	95% HD ↓
Zero	–	81.43±0.03	7.31±6.52
One	Fifth layer	83.17±0.05	5.17±6.68
Two	Fourth and fifth layers	85.54±0.04	4.05±5.15
Three	Fourth, fifth, and third layers	84.08±0.02	4.79±5.92

Data were presented as mean ± SD. HD, Hausdorff distance; SD, standard deviation.

compares the segmentation performance with one and two decoding layers connected to transformer-like layers. As the results show, after the network decoding layer was connected to a double-layer transformer, the segmentation performance showed an improvement compared to that with a single-layer transformer. We also set up an experiment which aimed to simultaneously access the transformer-like layer to the feature sequences output by more than three decoding layers. The result showed that as the computational efficiency decreased, the time complexity increased, and the computational memory footprint became too large during training. The final segmentation accuracy also decreased compared to that with the two-layer transformer-like method.

Discussion

Accurate segmentation of the pancreas is crucial in the quantitative analysis of radiology and surgery, as it can assist radiologists and surgeons in achieving the best diagnosis and treatment strategies for patients. To this end, we proposed a model-driven deep learning method based on UMRFormer-Net for pancreas segmentation. Our segmentation experiments demonstrated that the model has high segmentation accuracy and outperforms some existing methods in segmentation.

Our proposed model has the following two characteristics. First, a novel double MRFormer was proposed to enable a 3D network to efficiently apply 3D contextual information for 3D CT image segmentation. The UMRFormer-Net method embedded in the double MRFormer module can be used to more effectively obtain deep feature contextual information in 3D models. It forms a spatially local and global unified framework to directly optimize 3D segmentation results, thereby improving

model performance; Second, the proposed MRFormer module aims to focus on spatial local information in long sequence features. It has the advantage of maintaining long-range dependencies, as well as feature recognition and local context capabilities. It helps to solve the problem of unclear boundary texture segmentation of targets such as the pancreas. As shown in *Table 2* and *Table 3*, we can clearly see that the UMRFormer-Net using MRFormer is superior to other methods (4,15,25,26,27,28) in terms of the Dice score and 95% Hausdorff distance.

However, our proposed model still has a few limitations. First, the performance of the model depends on the accuracy of manual annotation during training, and manual annotation by doctors is based on their existing clinical knowledge. The size of pancreatic organ tissue varies with age (e.g., between children and adults), and the subjective standards of doctors for pancreas delineation differ slightly, which affects the performance of the model for pancreas segmentation. Therefore, it is necessary to fine-tune the model parameters according to the data characteristics of the case.

Image quality and annotation quality are critical in artificial intelligence application research. Suman *et al.* (31) studied the MSD dataset and found that there were numerous biliary stents and post-chemotherapy cases in the training datasets, which may impact the overall segmentation performance. It is therefore essential to further optimize the model structure or adopt a more effective way of enhancing data to improve the model's segmentation performance.

Conclusions

This paper has presented a novel medical segmentation framework named UMRFormer-Net and applied it

in pancreatic segmentation of CT images. The model effectively integrates the transformer-like structure into the CNN, which not only retains the advantages of the CNN in modeling local receptive field information, but also fully utilizes and innovates the advantages of the transformer module's learnable global dependence semantic information. Our experimental results from two datasets (the CPTAC-PDA Pancreas dataset and the MSD Pancreas dataset) fully demonstrate the effectiveness of the proposed UMRFormer-Net. Based on this endogenous hybrid architecture, UMRFormer-Net achieves better segmentation accuracy in pancreas than previous transformer-based segmentation models. In the future, we will test UMRFormer-Net in other medical segmentation tasks.

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Footnote

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Ethical Statement: The authors are accountable for all aspects of the work in ensuring that questions related

to the accuracy or integrity of any part of the work are appropriately investigated and resolved. The study was conducted in accordance with the Declaration of Helsinki (as revised in 2013).

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