# A Pseudo-labeling Approach to Semi-supervised Organ Segmentation

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**Abstract.** In this paper, we adopt a "pseudo-labeling" approach to semi-supervised learning based on 50 labeled data and 2000 unlabeled data. This approach yields a model with 0.7496 mean DSC on the validation set, outperforming the 0.6903 mean DSC of the model with only 50 labeled data.

**Keywords:** Abdominal organ segmentation  $\cdot$  Semi-supervised  $\cdot$  Pseudolabeling.

### 1 Introduction

Abdomen organ segmentation has many important clinical applications. Typically, a large number of labeled data is required to train a accurate segmentation model. However, manually annotating organs from CT scans is time-consuming and labor-intensive. This requires us to use appropriate semi-supervised segmentation methods to use unlabeled data, such as disturbance regularization based on data or model [7] [8] and consistency constraint based on multitask [6].

FLARE22 provides 50 labeled images and 2000 unlabeled images to train the segmentation model of 13 organs. There are three main difficulties. First, we need to realize the segmentation of 13 organs. Second, more than 97% of the training data are unlabeled. Third, we need to balance model performance and resource consumption.

Drawing on the idea of developing a classification model using "pseudo-labeling" in [5], we developed a segmentation model using the "pseudo-labeling" approach. Specifically, we first trained a model with labeled data, and then used the model to predict the unlabeled data to give them "pseudo-label". Finally, we fine-tuned the original model using all labeled data and some pseudo-labeled data.

# 2 Method

#### 2.1 Preprocessing

We use several pre-processing strategies as follows.

- Cropping strategy
  - We use the CT scans as the data source to generate the bounding box of foreground, and then crop only the foreground object of the images.
- Resampling method for anisotropic data
  - We resample the original data to unify the voxel spacing into [1.0, 1.0, 1.0].
- Intensity normalization method We change the intensity outside [-300.0, 300.0] to 0.0 and normalize the intensity of [-300.0, 300.0] to [0.0, 1.0].

# 2.2 Proposed Method

Figure 1 illustrates the applied 3D nnU-Net [4], where a 3D U-Net architecture is adopted. We use the "leakyrelu" function with a negative slope of 0.01 as the activation function. Our 3D nnU-Net has 14 out channels, corresponding to the background and 13 organs respectively.

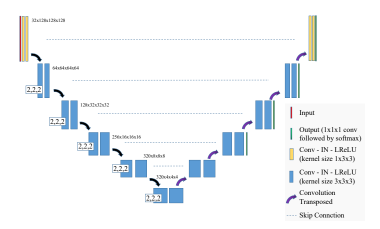


Fig. 1. Network architecture

Strategies to use the unlabelled cases: first, we trained a model with the 50 labeled data. Then we used this model to predict the 2000 unlabeled data to give them "pseudo-label". Finally, we picked out 676 data with at least 2000 voxels for each organ's "pseudo-label", and put them together with the 50 labeled data to fine-tune the original model.

Loss function: we use the sum of Dice loss (after applying a softmax function) and Cross Entropy Loss as the loss function.

Strategies to improve inference speed and reduce resource consumption: when predicting a single image with the trained segmentation model, we first resample it to a voxel spacing of [1.0, 1.0, 1.0], as we did during training, and try to predict. If there is a "CUDA out of memory" error, we resample it to [2.0, 2.0, 2.0] voxel spacing to reduce the size of the resampled image and thus reduce the usage of GPU memory.

# 2.3 Post-processing

During model prediction, we select the label (from 0 to 13) corresponding to the largest of the 14 outputs for each voxel.

# 3 Experiments

#### 3.1 Dataset and evaluation measures

The FLARE2022 dataset is curated from more than 20 medical groups under the license permission, including MSD [10], KiTS [2,3], AbdomenCT-1K [9], and TCIA [1]. The training set includes 50 labelled CT scans with pancreas disease and 2000 unlabelled CT scans with liver, kidney, spleen, or pancreas diseases. The validation set includes 50 CT scans with liver, kidney, spleen, or pancreas diseases. The testing set includes 200 CT scans where 100 cases has liver, kidney, spleen, or pancreas diseases and the other 100 cases has uterine corpus endometrial, urothelial bladder, stomach, sarcomas, or ovarian diseases. All the CT scans only have image information and the center information is not available.

The evaluation measures consist of two accuracy measures: Dice Similarity Coefficient (DSC) and Normalized Surface Dice (NSD), and three running efficiency measures: running time, area under GPU memory-time curve, and area under CPU utilization-time curve. All measures will be used to compute the ranking. Moreover, the GPU memory consumption has a 2 GB tolerance.

# 3.2 Implementation details

**Environment settings** The development environments and requirements are presented in Table 1.

Windows/Ubuntu version Ubuntu 20.04.4 LTS			
CPU	Intel(R) Xeon(R) Gold 5218R CPU @ 2.10GHz		
RAM	128G		
GPU (number and type)	two NVIDIA Tesla T4 16G		
CUDA version	11.6		
Programming language	Python 3.6		
Deep learning framework	Pytorch (Torch 1.10.1, torchvision 0.11.2)		
Specific dependencies	numpy 1.19.5, SimpleITK 2.0.2, monai 0.8.1		

Table 1. Development environments and requirements.

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# Training protocols As described below.

Random flipping strategy (only for initial training stage): each image has a 20% probability of flipping along the x-axis and a 20% probability of flipping along the y-axis.

Random Gaussian smooth (only for initial training stage): each image has a 10% probability of being Gaussian smoothed with sigma in (0.5, 1.15) for every spatial dimension.

Random Gaussian noise (only for initial training stage): each image has a 10% probability of being added with Gaussian noise with mean in (0, 0.5) and standard deviation in (0, 1).

Random intensity change (only for initial training stage): each image has a 10% probability of changing intensity with gamma in (0.5, 2.5).

Random intensity shift (only for initial training stage): each image has a 10% probability of shifting intensity with offsets in (0, 0.3).

Patch sampling strategy: 2 patches of size [128, 128, 128] are randomly cropped from each image. The center of each patch has 50% probability in the foreground and 50% probability in the background.

Optimal model selection criteria: we tried several different training protocols and selected the model with the highest DSC on the validation set.

Some details of the initial training stage and the fine-tuning stage are shown in Table 2 and Table 3 respectively.

Network initialization	"he" normal initialization
Batch size	2
Patch size	128×128×128
Total epochs	1600
Optimizer	Adam
Initial learning rate (lr)	0.0001
Lr decay schedule	Initial learning rate× $(1 - epoch/500)^{0.9}$
Training time	77 hours
Number of model parameters	31.42M

Table 2. Training protocols (initial training stage).

# 4 Results and discussion

#### 4.1 Quantitative results on validation set

Accuracy results on validation set are shown in Table 4. It can be seen from the table that the generalization ability of the model is indeed improved by using unlabeled data through the "pseudo-labeling" method.

Table 3. Training protocols (fine-tuning stage).

Network initialization	model after initial training
Batch size	2
Patch size	128×128×128
Total epochs	40
Optimizer	Adam
Initial learning rate (lr)	0.00005
Lr decay schedule	Initial learning rate× $(1 - epoch/500)^{0.9}$
Training time	39 hours
Number of model parameters	31.42M

 ${\bf Table~4.}~{\bf Results~on~validation~set}.$ 

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	Without using unlabeled data	Using unlabeled data
Mean DSC	0.6903	0.7496
Liver	0.9312	0.9493
RK	0.7151	0.8098
Spleen	0.8180	0.8962
Pancreas	0.6631	0.7506
Aorta	0.7474	0.7953
IVC	0.7003	0.7692
RAG	0.6792	0.6910
LAG	0.5257	0.5400
Gallbladder	0.6235	0.6543
Esophagus	0.6196	0.6641
Stomach	0.7550	0.8219
Duodenum	0.5261	0.5803
LK	0.6703	0.8225

# 4.2 Qualitative results on validation set

Two examples of good segmentation are shown in Figure 2 and two examples of bad segmentation are shown in Figure 3. Visualization is achieved with ITK-SNAP [11] version 3.8.0.

From the perspective of organs, the segmentation results of organs with fewer surrounding organs are better, such as liver and spleen. From the perspective of images, some potential reasons for the bad-segmentation cases are listed below.

- (1) The size of the case is very large, so we have to reduce the size of the case by resampling to avoid GPU memory overflow.
  - (2) The case is not clear, distorted, or skewed.
  - (3) There are rare structures in the case that are not in the training set.

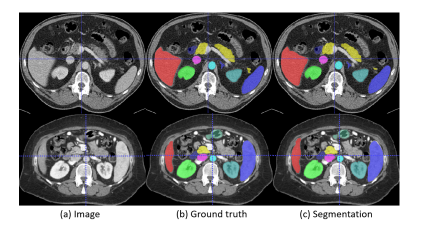


Fig. 2. Good segmentation examples

# 4.3 Segmentation efficiency results on validation set

Segmentation efficiency results for the 5th validation submission are shown in Table 5.

Table 5. Results on validation set.

Running time	1538.14 seconds
Maximal GPU Memory	16327MB
Area under GPU memory-time curve	11050890
Area under CPU utilization-time curve	26722.79

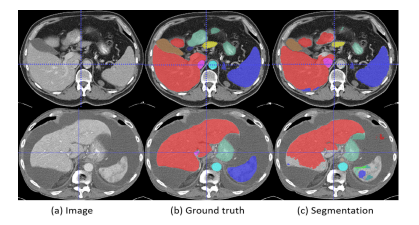


Fig. 3. Bad segmentation examples

The running time is relatively short since we didn't use any cascaded framework. In the testing phase, we used the "sliding\_window\_inference" function of monai to slice the image into several  $128\times128\times128$  patches and predict them separately. This can lead to a large GPU memory consumption when the image size is large.

# 4.4 Results on final testing set

This is a placeholder. We will send you testing results after the challenge.

#### 4.5 Limitation and future work

In terms of model accuracy, we have only tried the "pseudo-labeling" approach at present. In the future, we are going to try some more popular semi-supervised image segmentation methods, such as disturbance regularization based on data or model and consistency constraint based on multitask. In addition, we consider using cascaded method and post-processing method to improve the model, which are not used at present.

In terms of segmentation efficiency, we consider rewriting the slice-and-predict function instead of using the "sliding\_window\_inference" function of monai directly to reduce the GPU memory consumption. In addition, we consider using some optimization methods to improve the running speed of the model in the future.

### 5 Conclusion

Using unlabeled data through "pseudo-labeling" method can improve the performance of the model.

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