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Diagnosis after Zooming in: A Multi-label Classification Model by Imitating Doctor Reading Habits to Diagnose Brain Diseases

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15 Abstract

Purpose: Computed tomography (CT) has the advantages of being low cost and noninvasive and is a primary diagnostic method for brain diseases. However, it is a challenge for junior radiologists to diagnose CT images accurately and comprehensively. It is necessary to build a system that can help doctors diagnose and provide an explanation of the predictions. Despite the success of deep learning algorithms in the field of medical image analysis, the task of brain disease classification still faces challenges: researchers lack attention to complex manual labeling requirements and the incompleteness of prediction explanations. More importantly, most studies only measure the performance of the algorithm, but do not measure the effectiveness of the algorithm in the actual diagnosis of doctors.

Methods: In this paper, we propose a model called DrCT2 that can detect brain diseases without using image-level labels and provide a more comprehensive explanation at both the slice and sequence levels. This model achieves reliable performance by imitating human expert reading habits: targeted scaling of primary images from the full slice scans and observation of suspicious lesions for diagnosis. We evaluated our model on two open-access datasets: CQ500 and the RSNA Intracranial Hemorrhage Detection Challenge. In addition, we defined three tasks to comprehensively evaluate model interpretability by measuring whether the algorithm can select key images with lesions. To verify the algorithm from the perspective of practical application, three junior radiologists were invited to participate in the experiments, comparing the effects before and after human-computer cooperation in different aspects.

Results: The method achieved F1-scores of 0.9370 on CQ500 and 0.8700 on the RSNA dataset. The results show that our model has good interpretability under the premise

of good performance. Human radiologist evaluation experiments have proven that our model can effectively improve the accuracy of the diagnosis and improve efficiency.

Conclusions: We proposed a model that can simultaneously detect multiple brain diseases. The report generated by the model can assist doctors in avoiding missed diagnoses, and it has good clinical application value.

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Keywords: Medical image classification, Interpretability, Attention mechanism, Human-AI Interaction.

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₅ I. Introduction

Brain disease is one of the ailments that threaten health and damages the life of humans¹. Due to the increase in patients, professional doctors are insufficient². Training an experienced doctor usually takes more than seven years. Therefore, it is necessary to build a computeraided medical diagnosis system, which can help doctors diagnose effectively and accurately. 79 In recent years, deep learning algorithms have been applied in brain disease classification tasks, e.g. ^{3,4,5,6}. Although these algorithms have achieved good performance, this domain 81 still faces challenges. Most studies label sequence CT images at the image level, which is time-consuming. More important, separately labeling images or diagnosing by algorithm 83 violates common medical knowledge. Diagnosing sequenced medical images requires the observation of adjacent images. Doctors usually browse adjacent slices and capture the changes between the slices to make a judgment on the disease. There are subtle differences between adjacent brain CT images, but they play a decisive role in disease judgment. The 87 key images and points almost determine all the judgments of the disease. How to focus on 88 them to obtain more information about the disease is essential.

By considering all the challenges we analyzed before, we proposed a model called DrCT2. The method is inspired by the diagnostic habits of radiologists. As shown in Figure 1, in clinical diagnosis, radiologists browse a complete set of brain CT scans and focus on the key images that may reflect diseases, observing closer to get more information about 93 the lesion. Our model imitates the doctor's reading habit by selecting key images and zooming in to obtain detailed information at different scales. It consists of three parts: primary network, attention proposal slices network (APS), and knowledge fusion network. The primary network includes feature attention, dependencies learning, and slice attention 97 module. These two attention modules focus on key features and images that may reflect the lesion, respectively. Dependencies learning module can learn the relationship between sequence slices. The APS zooms in on key images proposed by the slice attention module 100 and obtains more details from different scales. The knowledge fusion network merges the 101 knowledge learned from two neural networks to make a final judgment. Our model inputs 102 sequence CT, and only requires the label at the sequence level. It is not necessary to label 103 each CT image as in most studies, which reduces the workload of data labeling and is more 104 reasonable. 105

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The medical computer vision (CV) model is different from the normal CV model. The normal CV model focuses on the improvement of algorithm performance, while the medical CV model needs to focus more on whether the model can truly assist doctors in diagnosis under the premise of ensuring performance. In this paper, we put forward a new perspective: for medical models, it is not only to maximize the prediction accuracy of the algorithm itself but to measure the benefits that practitioners obtained after interaction with the algorithm. We defined 3 tasks to evaluate whether the model can select key images. The evaluation of key slices selection proves the reliability and interpretability of our model. Selecting slices accurately is also very important for prompting radiologists to avoid misdiagnosis. We further invited junior radiologists to simulate real diagnosis scenarios for evaluation. The results show that our model can effectively improve diagnostic efficiency and accuracy. Compared with previous work⁷, we improve the performance of the algorithm and propose interpretive evaluation tasks. In the evaluation experiments, the effect of the model to assist doctors in decision-making is verified. We not only evaluated the test set like most studies but also focused on doctors to evaluate the potential of our model to assist clinical decision-making.

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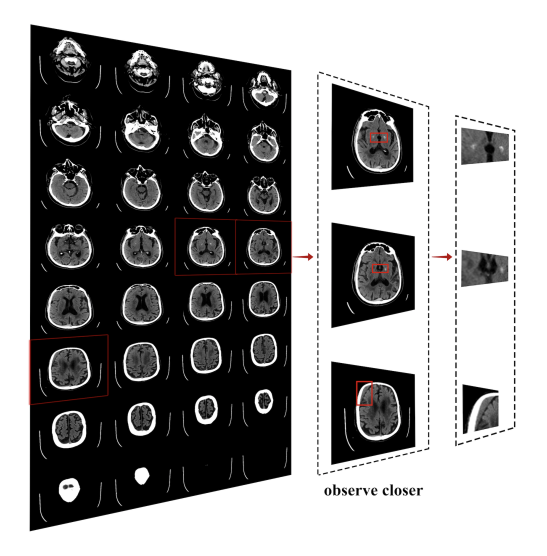


Figure 1: Some diseases are straightforward to misdiagnose, which requires observing and considering adjacent images. In this CT scan, we need to observe adjacent images to distinguish between a bleeding point or calcification. It is easy to ignore subdural hemorrhage without zooming in on the image.

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122 II. Related Works

In this paper, we propose a model inspired by radiologists' diagnostic habits by zooming in on key slices, analyzing at the sequence level, and giving multi-label classification results. Our model can provide the interpretability of the model, and we evaluate this through a sequence-level interpretative evaluation task. In Section II.A., we first analyze some sequence-level image classification algorithms from two types of research directions, 2D and 3D. Our research pays special attention to interpretability, so we also analyze some interpretable medical image analysis algorithms in Section II.B..

30 II.A. Sequence-level Image Classification

In reference ^{8,9,10,11}, the temporal attention mechanism which focuses on important frames has been widely used. The algorithm proposed by Yang et al. ¹² can adaptively capture the regions of interest in each frame and learn the key features based on these areas. Yu et al. ¹³ proposed a method for generating sentences to describe a video and Tu et al. ¹⁴ proposed the spatial-temporal attention (STAT) method for the video description task. These two studies ^{13,14} takes into account both the spatial and temporal information in a video.

Although these studies have been successful, they cannot solve the problem of sequence-level brain disease classification. Many studies ^{15,16,17} have tried to use 3D convolutional neural networks to solve the problem of sequence-level brain disease classification. Nie et al. ¹⁸ proposed a novel 3D convolutional neural network architecture for learning supervised features. Gao et al. ¹⁹ integrated 2D and 3D CNN networks to classify brain diseases. Nawali et al. ¹⁶ and Ker et al. ²⁰ proved the performance of 3D CNN in the classification task of a cerebral hemorrhage.

These studies prove the effectiveness of 3D convolutional neural networks in medical image processing tasks. However, the 3D architecture requires a large amount of calculation, low-resolution images are selected as input to preserve the complete brain structure during training. If the resolution of the image is reduced, the need for judging subtle bleeding points and lesions cannot be met. Therefore, if the model can fuse the input multi-resolution images, such as using high-resolution images for key images, it can provide more sufficient information from different aspects which may achieve better performance. Multi-resolution

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image fusion analysis has many advantages²¹. For sequence-level medical images, key images 151 and areas play a decisive role in disease judgment. Fu et al.⁷ proved its reliability and 152 reasonableness. Jiang et al.²² proposed a model called MFI-Net that can avoid the loss of 153 coarse-grained feature information in the shallow layer by extracting local and global feature 154 information at different resolutions. Liu et al.²³ proposed a multi-resolution medical image 155 fusion network with iterative back-projection (IBPNet). Experimental results show that it 156 has better performance in visual perception and objective evaluation. Li et al. 24 proposed a 157 model for lung nodule detection that employed patch-based multi-resolution CNNs to extract 158 the features and employed four different fusion methods for classification. In this work, we 159 focus on key slices and areas at the same time and try to obtain information about lesions 160 on slices from multiple resolutions. 161

Dependencies between slices should be considered, and some research has recently learned slice dependencies through 3D networks. Zhuang et al.²⁵ proposed a new self-supervised learning model that contains a Rubik's cube recovery task. It can pre-train 3D neural networks from raw 3D medical data. Compared with the training strategy from scratch, it can achieve better performance on various tasks. Zhu et al.²⁶ further developed this method, enriched the pre-training tasks, and achieved better performance in the downstream tasks. Zhu et al.²⁷ proposed a novel SSL approach for 3D medical image classification. It embeds task knowledge into training 3D neural networks. The experimental results demonstrate the effectiveness of embedding lesion-related prior knowledge into neural networks for 3D medical image classification. The above research proves that modeling and learning dependencies between slices are effective. In our model, we learn about the dependencies between slices through the cooperation of the primary and auxiliary (APS) networks.

II.B. Interpretive Medical Image Analysis Model

In the medical image analysis domain, the interpretability of the model is significant. Only giving the prediction result is not credible; the basis for the model's decision also needs to be explained. Pranav et al. ²⁸ proposed a convolutional neural network, CheXNet, that can output the probability of disease and use the class activation mapping method to indicate the lesion areas of the lung disease. Zoom-in-Net ²⁹ can generate four bounding boxes based

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on the attention maps of the highlighted suspicious area, and four bounding boxes can cover 80% of the lesions. Zhang et al. 30 proposed a solution for pathological diagnosis and interpretation. The model proposed by this team can generate an interpretive report for the pathologist's reference. The report not only displays the selected regions of interest but also generates explanatory text for different regions. Wang et al. 31 proposed an algorithm for the classification of intracranial hemorrhage diseases, which can increase the interpretability of the model by outputting a prediction basis and image-level attention maps.

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These studies all indicated suspicious areas on a single image. However, in our task, the doctor is concerned not only about the suspicious part of the image but also about which slice in the sequence CT can reflect the lesion. It is significant to automatically select key images, which can improve the efficiency of doctors' reading and provide an explanation for the prediction. In addition, the key images selected by the model can be used as a reference for the doctor's diagnosis, prompting the doctor to avoid missed diagnosis or misdiagnosis. As shown in Figure 2, our model can generate reports to assist doctors in diagnosis.

Human-computer Interaction (HCI) in Medical Image Anal-II.C. ysis Domain

The focus of the medical image analysis model is to assist doctors in their work, so it should be human-centered³². The combination of AI-based frameworks with complementary human intervention could result in synergistic effects inpatient management, interpretation, and diagnosis³³. Sayres et al.³⁴ invited 10 ophthalmologists to diagnose the diabetic retinopathy disease in three conditions (unassisted, grades only, or grades plus heatmap) based on retinal fundus images to evaluate their model. The results found that algorithm-assisted diagnosis 202 improved the diagnostic accuracy and confidence of ophthalmologists, and the diagnosis time 203 was reduced after providing model explanations (the third condition). Zhao et al. 35 developed a deep learning model to help doctors diagnose musculoskeletal tumors. The expert evaluation experiments showed that their models improved the sensitivities of six of seven doctors and accuracy in three of seven doctors, while they did not significantly reduce the specificities of any. Choi et al. ³⁶ compared the diagnostic performance of radiologists on two datasets (breast ultrasound images alone or with computer-aided diagnosis) and showed that the computer-aided diagnosis method could improve radiologists' diagnostic performance by 210

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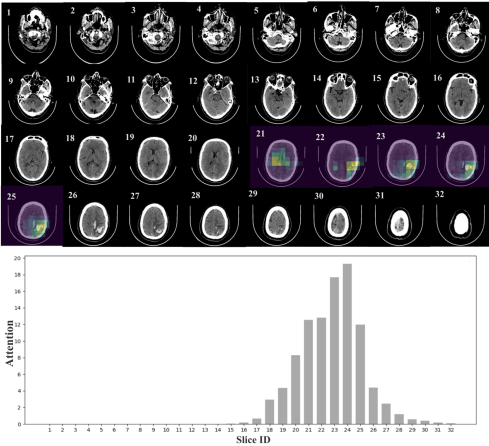
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increasing specificity, accuracy, and positive predictive value. Ding et al.³⁷ invited 20 gastroenterologists to conduct experiments, and the results showed that an algorithm-assisted method can identify abnormalities more sensitively and rapidly than conventional diagnosis by gastroenterologists. Esmaeili et al.³⁸ trained some explainable deep learning models based on the Grad-CAM mechanism and evaluated whether the trained model could localize tumor regions. The results show that deep learning models may classify some tumor brains based on other non-relevant features. We believe that to evaluate model performance should not only consider under experimental data but more importantly, compare the effects of assisting doctors.



This patient may have the following diseases: ICH, IPH, Mass Effect

Figure 2: An example of the diagnostic report generated by the DrCT2 model. The model can identify the 5 slices most likely to reflect the lesion and highlight the suspicious area automatically. The histogram is the model's attention to each image. It can be seen from the figure that our algorithm accurately selects the key images and key areas.

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III. Task Analysis

Difficulty in labeling data Labeling medical data is a costly and time-consuming task.
To avoid the waste of expert resources, the algorithm proposed in this paper is based on
the sequence level: can be trained without labeling on a single image. Labeling data at
the sequence level is easier to implement, and expert diagnosis based on full slice CT obeys
medical commonsense.

Adjacent slice dependence In clinical practice, a patient's condition cannot be inferred from a single slice, and doctors usually browse adjacent slices and catch the changes between the slices to diagnose. As shown in Figure 3, observing changes in continuous CT images is conducive to the diagnosis of the disease and reduces the occurrence of misdiagnosis and missed diagnosis. The sequence-level model we proposed is more in line with the doctor's diagnostic habits.

Dependency between diseases There are dependency relationships between brain diseases. For example, skull fractures are often accompanied by epidural hemorrhage. Hypertensive cerebral parenchymal hemorrhage may break into the ventricle and cause intraventricular hemorrhage and form a mass effect. For our model, the dependencies learning module learns the sequence images from positive and negative directions and considers the relationship between different diseases at the same time.

Easy to miss diagnosis Some diseases are easily missed in the judgment of brain diseases for the doctor, such as subtle bleeding points and subarachnoid hemorrhage³⁹. As shown in Figure 1, if the doctor does not observe the brain CT carefully, it is difficult to find the bleeding point, because some small bleeding points are not significantly different from calcification. An algorithm with interpretability can provide references for doctors' diagnoses. For our task, the algorithm should be able to pick out the key images and highlight the suspicious areas in the images.

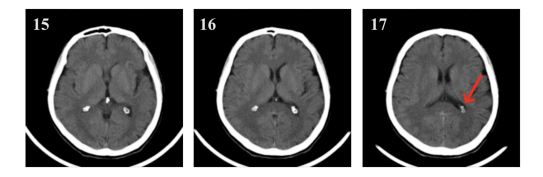


Figure 3: The three slices in this figure are adjacent. There is a small white dot where the arrow points. It is difficult to judge whether there is intraventricular hemorrhage only by observing the slice with ID 17. However, by observing the changes between the three slices, we can see that the white spot is calcification, not intraventricular hemorrhage.

IV. Sequence-level Interpretive Evaluation Tasks

Whether the key images can be selected correctly is crucial to assist in diagnosis. It is also an important indicator for evaluating model performance. To evaluate the performance under different disease conditions, we define the following tasks from different levels.

Easy condition task We define in a set of CT (28 images/set), having five or more images that can reflect the disease, which is the simple condition for our algorithm. We evaluated how many of the top five images with high attention selected by the algorithm reflected the lesion.

Difficult condition task We defined in a set of CT images (28 images/set) as having five or fewer images that can reflect the disease as the difficult condition for our algorithm. We evaluated whether the top five images with high attention selected by the algorithm could reflect lesions. This task can evaluate whether the algorithm can achieve good performance in the case of an easily missed diagnosis. If the algorithm can achieve good performance under this task, it proves that it has the potential to assist doctors in discovering difficult-to-detect diseases.

Comprehensive condition task There are h images in a set of CT that can reflect the disease, to evaluate whether the top h images selected by our algorithm with high attention

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can cover. This task can comprehensively evaluate the explanatory performance at the sequence level.

The pseudo-code for our three tasks can be seen in algorithm 1. U is a dataset containing 264 n_{scan} scans, and the composition of the dataset is different under different disease conditions. 265 S_i represents the i-th set of scans. The function Model is the trained model, which inputs 266 a set of slices and outputs the attention value of each slice. The function Sort sorts the 267 attention value in descending order. The function Select select the slices (T_i) corresponding to the top m_{select} values of the sort result. s_j represents the slice in T_i . For easy task and 269 difficult task, m_{select} is equal to 5; for the comprehensive task, m_{select} is the number of slices that can reflect the lesion in the set of slices. D_i is the number of images selected correctly. 271 $Score_i$ is the evaluation result of the model on CT scan S_i . Sum is a variable used to record scan evaluation results. We use the average accuracy for evaluation. 273

Please note that the easy and difficult tasks defined here are only defined from the number of diseased slices, not all from the prediction mechanism of the algorithm or from the perspective of radiologists.

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Algorithm 1: Evaluation algorithm for key image picking

```
Input: U = \{S_i\}, i \in [1, n_{scan}]
   Output: Accuracy: evaluation results
 1 Initialize: Accuracy \leftarrow 0; Sum \leftarrow 0;
 2 for each S_i \in U do
       D_i \leftarrow 0
 3
       Score_i \leftarrow 0
 4
       A_i \leftarrow Model(S_i)
 5
       V_i \leftarrow Sort(A_i)
 6
       T_i \leftarrow Select(V_i, S_i)
 7
       if easy condition task then
 8
           for each s_i \in T_i do
 9
               if slice s_i is the key slice then
10
                   D_i = D_i + 1
11
               end
12
           end
13
           Score_i = D_i/5
14
           Sum = Sum + Score_i
15
       end
16
       if difficult condition task then
17
           if T_i contains key slices then
18
               Sum = Sum + 1
19
           end
20
       end
\mathbf{21}
       if comprehensive condition task then
\mathbf{22}
           for each s_i \in T_i do
23
               if slice s_i is the key slice then
24
                   D_i = D_i + 1
25
               end
26
27
           Score_i = D_i/m_{select} Sum = Sum + Score_i
28
       end
29
30 end
31 Accuracy = (Sum/n_{scan}) * 100\%
```

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V. Methods

As shown in Figure 4, our proposed model DrCT2 contains three modules: primary network, attention proposal slices network (APS), and knowledge fusion network. The primary network work learns the dependencies between slices and diseases. Two-step attention mechanisms keep the algorithm focused on key images and suspicious parts. The APS network zooms in key images proposed by the primary network and learns more details on different scales. The knowledge fusion network merges all the information learned from two modules and gives a final prediction. The cooperation of the three modules makes the model have good performance and explanatory.

56 V.A. Primary Network

The primary network is an encoder-decoder network, which we call DrCT1⁷. It inputs a set of brain CT images that after sampling (we sampled fix-length images from a set of CT images with different numbers of slices) to the encoder and outputs a feature matrix (each image is represented as a feature vector, and we combine them into a feature matrix). A set of brain CT images after sampling contains m slices.

$$S = (s_1, s_2, ..., s_m) (1)$$

We used the VGG16⁴⁰ model pre-trained on ImageNet⁴¹ to extract features from each slice. We froze the convolutional layers and discarded the fully-connected layers during feature extraction. The adaptive max-pooling method is used to compress the feature maps (512 channels, 7×7 size) into 512-dimensional vectors (512 channel, 1×1 size feature map). Each slice is represented by a 512-dimensional feature vector:

$$s_t = (z_1^{< t>}, z_2^{< t>}, ..., z_{512}^{< t>}), t \in [1, m]$$
 (2)

The decoder consists of three parts: the feature attention part, the dependencies learning part, and the slice attention part. The feature vectors passed through these three modules to learn the probability of each disease.

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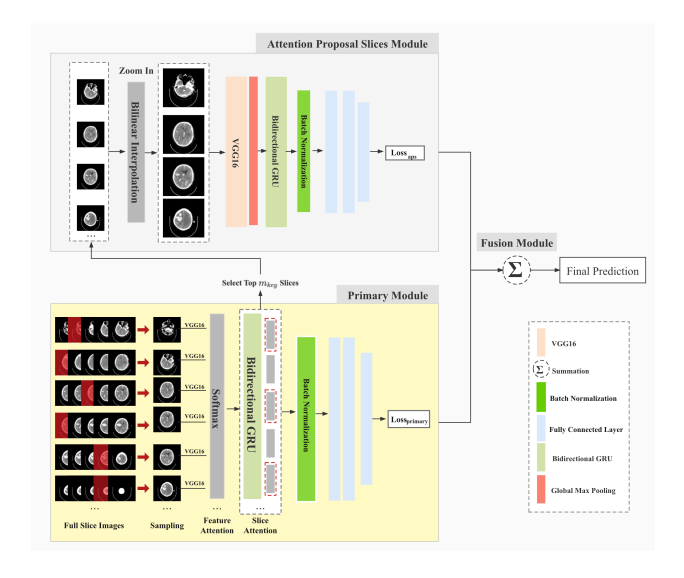


Figure 4: An overview of our proposed model consists of three parts: i) The primary network is encoder-decoder-based and assigns more weight to areas or slices that may contain lesions through the attention mechanism. ii) The APS network selects the m_{key} slices most likely reflecting the lesions and zooms in them as the input of the APS network to learn more detailed information. iii) The fusion network considers the knowledge learned by the two networks to make judgments about the disease.

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V.A.1. Feature Attention

The feature vector learned by CNN is the high-dimensional feature representation of each image. Feature attention can assign different weights to each node. Nodes with high attention can be visualized to explain model prediction. It can be defined as:

$$w_i^{\langle t \rangle} = \frac{exp(z_i^{\langle t \rangle})}{\sum_{j=1}^{512} exp(z_j^{\langle t \rangle})}$$
 (3)

where $z_i^{< t>}$ is the value of the *i*-th dimension vector in a 512-dimensional vector, and $w_i^{< t>}$ is the weight assigned to the *i*-th dimension vector. The feature vectors $(z_1^{< t>}, ..., z_{512}^{< t>})$ of the original slice (s_t) merge with the assigned weights $(w_1^{< t>}, ..., w_{512}^{< t>})$ to obtain a new slice feature representation x_t .

V.A.2. Dependencies Learning

This module can learn the dependencies between slices through a bidirectional recurrent neural network (BRNN)⁴². It can capture information from both positive and negative directions. Gated recurrent unit (GRU⁴³) as the base unit of BRNN. GRU solves the long-term dependency problem in RNN networks. The new feature representation $(x_1, ..., x_t, ...x_m)$ is put into the BGRU for dependencies learning. The input of base units at each time step is the new feature representation weighted by the feature attention module. We use element-wise multiplication to merge these two vectors in this step. The output formula of each time step of BGRU is:

$$\hat{y}^{\langle t \rangle} = g(W_y[\overrightarrow{a}^{\langle t \rangle}, \overleftarrow{a}^{\langle t \rangle}] + b_y) \tag{4}$$

where $\hat{y}^{< t>}$ is the output at the t-th time step, $\overrightarrow{a}^{< t>}$ and $\overleftarrow{a}^{< t>}$ represent the information in the positive and the negative directions respectively, W_y and b_y are weight and bias respectively, and function q represents the activation function.

This module learns the dependencies between sequence-weighted features learned by feature attention.

6 V.A.3. Slice Attention

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This module can assign weights to each vector as shown below:

$$a_t = Mapping(\hat{y}^{< t>}) \tag{5}$$

$$w_t = \frac{exp(a_t)}{\sum_{j=1}^{m} exp(a_j)}$$
 (6)

where $\hat{y}^{< t>}$ is the output vector of at the t-th time step in the dependencies learning part. The Mapping function represents a fully connected layer, which maps the output vector to a value of length 1. m represents the number of slices after sampling. w_t represents the importance of the t-th slice in disease judgment. $(\hat{y}^{<1>},...,\hat{y}^{<m>})$ and $(w_1,...w_m)$ are fused (using the element-wise multiplication method) to obtain the result after slice attention.

Highly attended feature vectors are proposed for the APS network. Weighted feature vectors pass through three fully connected layers and output the probability of each disease, and use the dropout ⁴⁴ layer to prevent over-fitting, the dropout rate is 0.5. The activation function of the first two fully-connected layers is ReLU, and for the output layer, we used sigmoid function.

V.B. Attention Proposal Slices Network

The APS network can learn more detailed information from key images. The model can improve performance with the help of this network. The APS network zooms in on m_{key} key images proposed by the slice attention module and learns more details from different scales.

$$key \ images = (k_1, k_2, ..., k_{m_{key}}) \tag{7}$$

We used bilinear interpolation to zoom-in on m_{key} key images from 224×224 to 512×512 . In large-resolution images, key details are easier to obtain. The formula is as follows:

$$K_i = BilinearInterpolation(k_i) \tag{8}$$

$$F_i = P(K_i) \tag{9}$$

$$F_l = merge(F_1, ..., F_{m_{key}})$$
 (10)

where the function BilinearInterpolation represents the bilinear interpolation algorithm; k_i is the i-th slice in the m_{key} key slices, and K_i is the i-th slice after zoom-in operation. The function P is the VGG16 model pre-trained on ImageNet. F_i is the extracted feature vector, and F_l is the result of combining m_{key} feature vectors. This feature matrix passed through

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a fully connected layer after using batch normalization ⁴⁵. Then, the dependencies learning module and three fully connected layers were passed through to obtain the probabilities of diseases.

360 V.C. Knowledge Fusion

In clinical diagnosis, doctors make the final diagnosis by considering overall and details. The knowledge fusion module merges the knowledge from the primary network and the APS network. We summed the two feature matrices and after three fully connected layers, we obtained the final prediction.

365 V.D. Loss Functions

We optimized three losses during the overall training:

$$Loss = Loss_{primary} + Loss_{aps} + Loss_{fusion}$$
 (11)

Loss_{primary} and Loss_{aps} is the mean square error (MSE):

$$MSE = \frac{1}{n} \sum_{i=1}^{n} (y_i - y_i')^2$$
(12)

Loss_{fusion} is the binary cross entropy error (BCE):

$$BCE = -\frac{1}{n} \sum_{i=1}^{n} (y_i * log(y_i') + (1 - y_i) * log(1 - y_i'))$$
(13)

y' is the prediction of our model and y is the true label. n is the number of samples.

DrCT2 contains three modules: the primary network and the APS network use the MSE loss function for supervision and the BSE loss for the fusion module.

$_{\scriptscriptstyle{5}}$ VI. Evaluation

We evaluated the performance of our model from two perspectives. In the first part (Section VI.A.), we evaluate from the perspective of algorithm performance, conduct ablation experiments, and compare experiments. However, the medical task-oriented model is different from

the open domain computer vision model. The evaluation of computer vision models is more inclined to improve the performance of the algorithm, and the medical task is more focused on measuring whether it could assist doctors with diagnosis in the actual applications. More emphasis is placed on the cooperation between humans and machines. In the second part (Section VI.B.), from the perspective of human-computer interaction, we invited three junior radiologists to measure the impact of utilizing our model on the human diagnosis.

385 VI.A. Algorithm Performance Evaluation

6 VI.A.1. Dataset and Training Details

Our experiment was evaluated using two datasets: CQ500⁴⁶ and the dataset provided by the RSNA Intracranial Hemorrhage Challenge (represented as RSNA)⁴⁷. Table 1 shows 388 the data distribution of these two datasets. We use a fixed-length sampling method to 389 sample scans with different numbers of slices to the same number (32 for CQ500, 28 for 390 RSNA). The dataset after sampling is called the sampled dataset. For example, in the 391 0.625 mm scan data, 256 brain CTs will be generated, which will increase the computational 392 complexity of the algorithm. The unsampled data are used to generate a new CT set for data 393 enhancement. We use k-fold cross-validation (k equal to 10 for the CQ500 dataset, and 5 394 for the RSNA dataset). We keep the same data distribution as the first place solution in the 395 RSNA challenge for comparison reasons. We use PyTorch ⁴⁸ as a framework for implementing 396 our deep learning models. The experiments are performed on a workstation equipped with 397 a 3.90 GHz CPU and 2070 GPU with 8 GB memory. The batch size is 32 for the CQ500 398 dataset and 256 for the RSNA dataset. We trained for 100 epochs and used the Adam 49 399 optimizer for optimization. In DrCT2, we used different learning rates for the three modules: 400 0.0003 for APS and 0.0005 for the primary network and fusion network. In the following 401 experiments, we used the Macro-F1 score as our evaluation metric. Macro-F1 can treat each 402 category equally, and it will be more sensitive about rare categories ⁵⁰. In this task, we hope 403 that the model can consider the impact of fewer diseases on the performance of the model; 404 while Micro-F1 is more easily affected by common categories. Therefore, we choose Macro 405 F1 as our evaluation metrics. 406

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Table 1: Data distribution of CQ500 and RSNA.

Dataset	CQ500	RSNA
Patients	490	18923
Scans	1181	21540
Diseases	9	6
Number of scans	32-396	21-60
After Sampling	32	28
Disease label statistics (a	at sequen	ce level)
Any(ICH)	701	7855
Intraparenchymal(IPH)	356	4688
Intraventricular(IVH)	72	3236
Subdural(SDH)	102	3388
Epidural(EDH)	20	309
Subarachnoid(SAH)	128	3468
Calvarial fracture	181	
Mass Effect	248	
Midline shift	173	

VI.A.2. Supervised with Different Loss Functions

We chose the MSE loss as the loss function of the primary network and the auxiliary network (APS). However, in principle, both modules can use the BCE as the loss function. We construct experiments on two datasets (CQ500 and RSNA) to compare the performance of models supervised by different loss functions. The result can be seen in Table 2. It can be seen from the experimental results that better performance can be obtained by using the MSE as the loss function. Therefore in the following experiments, we use the MSE as the loss function for these two modules.

Table 2: Comparative experiment of using different loss functions for $Loss_{Primary}$ and $Loss_{Aps}$.

Model	Dataset	Loss Function	Precision	Recall	F1 Score
	CQ500 RSNA	BCE	94.23%	93.94%	0.9360
DrCT2		MSE	94.29%	94.10%	0.9370
DICIZ		BCE	87.93%	88.10%	0.8682
		MSE	88.05%	88.23%	0.8700

VI.A.3. Effect of APS Module

The APS module zooms in on m_{key} slices and puts them into the network. The selection of m_{key} slices depends on the slice attention mechanism, and they are the key images selected by the algorithm. We construct the following experiments to verify the role of this module from 3 aspects.

- With or without APS Network;
- Under a different number of slices, select through attention mechanism or random;
- Input under different resolutions.

With or without APS Network Under Different Datasets To evaluate the effect of the APS network, we evaluated the performance of the DrCT1 (without APS module) and DrCT2 (with APS module) model on two datasets are shown in Table 3.

Table 3: Experiment between the DrCT1 and DrCT2 models on the two datasets.

Dataset	Model	Precision	Recall	F1 Score
CQ500	DrCT1	93.10%	93.14%	0.9262
CQ300	DrCT2	94.29%	94.10%	0.9370
RSNA	DrCT1	87.51%	87.83%	0.8650
IGNA	DrCT2	88.05%	88.23%	0.8700

The precision-recall curve of DrCT2 in the training process is drawn in Figure 5.

From the results, we can see that with the help of the APS network, the DrCT2 model has better performance (+1.08% F1 for CQ500; +0.5% F1 for RSNA) than DrCT1. This proves the importance of the targeted fusion of multi-scale information.

Key Slices Selection We design experiments to compare the performance of random selection or selection through attention mechanisms. The architecture of our network is as described above, and the resolution of the image input to the APS network is 512×512 . The two methods select m_{key} slices from m brain CT images (m is the number of slices after sampling). We selected 4 to 28 slices in the following experiments, and the results are shown in Figure 6. We can see that it is effective to input after selection through the attention

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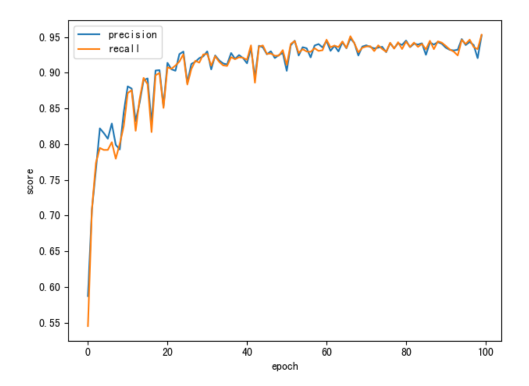


Figure 5: Recall and precision curve when training the DrCT2 model on dataset CQ500.

mechanism under a different number of slices. In the DrCT2 model, the best results were achieved when 16 images were selected. Therefore, in the following experiments, we set $m_{key} = 16$ as the default setting.

To further explore the potential rules of model selection of slices, we statistically analyzed the 5 slices in which the model gave the highest attention weight. On the CQ500 dataset, there are 744 sets of CT scans that reflect the disease. The statistical graph is shown in Figure 7. We found that the key slices are often concentrated from the 16th to the 22nd slices.

Under Different Resolutions To evaluate the effect of the zoom-in operation, we design the following 5 experiments. For the DrCT2 model, we input images of different resolutions (224×224 vs. 512×512) into the APS network to evaluate its performance. For the DrCT1 model, we construct experiments to evaluate the performance under original resolution (224×224) or high resolution (512×512). After statistical analysis, as shown in Figure 7, we found that the key slices were often concentrated from the 16th to the 22nd slices. So we

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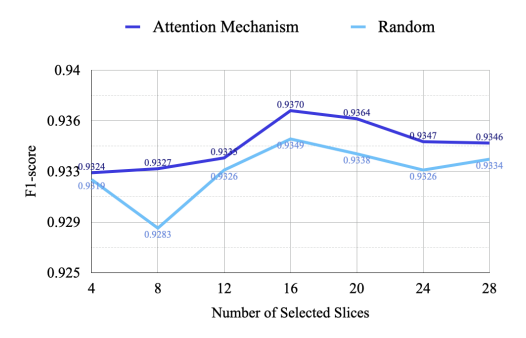


Figure 6: Evaluation results when m_{key} slices are input to the APS network by random selection and selection by our attention mechanism.

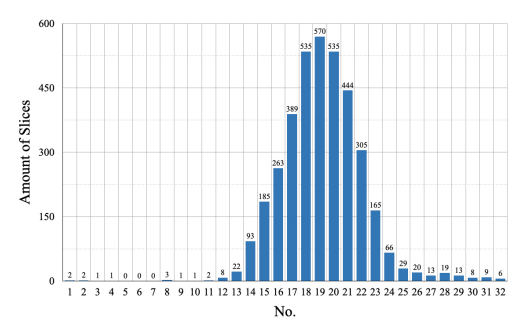


Figure 7: Statistics of the top five slice positions with high attention selected by our model in 744 sets of diseased brain CT (CQ500's total diseased CTs).

simulate the mechanism of DrCT2 and replace these slices (from 16th to 22nd) with highresolution images, while the rest still maintain low-resolution images. We call this strategy page 22 Ruigian Wang

the model fusion of DrCT1. Note that all the experiments are evaluated based on the CQ500 dataset. The experiment result can be seen in Table 4.

Table 4: Comparative experiment on different strategies for DrCT1 and DrCT2.

Model	Dataset	Strategy	Input image resolution	Precision	Recall	F1 Score
		Original	Primary: 224×224	93.10%	93.14%	0.9262
DrCT1		High-resolution Input	Primary: 512×512	91.70%	91.39%	0.9105
	CQ500	Model Fusion	Primary 1: 224×224	92.57%	92.27%	0.9179
		Woder rusion	Primary 2: 512×512	92.01/0	92.2170	
		With APS	Primary: 224×224;	93.96%	93.62%	0.9327
DrCT2	Γ2	WIGHTALS	APS: 224×224	93.9070	95.02/0	0.9321
		With APS	Primary: 224×224;	94.29%	94.10%	0.9370
		With Al 5	APS: 512×512	34.29/0	94.1070	0.9310

VI.A.4. Performance on Interpretive Evaluation Tasks

Model performance is certainly important in computer-assisted diagnosis algorithms, but our research not only focuses on performance but also model interpretability. This model can jointly select the key slices that the model focuses on through the attention mechanism and the auxiliary network (APS network). We elaborated in detail in Section IV.. The experiment is based on the dataset RSNA because it has annotations for each slice. We determined the number of slices for the three tasks we defined, and evaluated the performance of DrCT1 and DrCT2 on these three tasks, as shown in Table 5.

Table 5: The results of the two models in the interpretive evaluation task.

Task	Scans Number	DrCT1	DrCT2
Easy Task	6,529	81.59%	83.92%
Difficult Task	1,759	58.76%	62.67%
Comprehensive Task	7,855	65.72%	67.15%

To evaluate the performance of the key images selection algorithm on different diseases, we calculated the accuracy for each disease: for all slices that reflect a certain disease, how many slices our algorithm can correctly pick out. The results are shown in Table 6. The results show that our model can be selected more accurately than DrCT1.

Disease	Amount	DrCT1 Accuracy	DrCT2 Accuracy
EDH	2376	77.90%	79.46%
IPH	28270	76.70%	79.11%
IVH	20473	80.71%	81.96%
SAH	27784	80.74%	80.96%
SDH	36672	84.82%	84.84%
Any	83923	77.41%	78.23%

Table 6: The accuracy of the key slice selection for each disease.

66 VI.A.5. Comparison with Image-level Algorithm

The first place in the RSNA Intracranial Hemorrhage Challenge ⁵¹ is an image-level algorithm (represented as RSNA-1). The RSNA-1 utilizes total labels for each slice in a scan, and we only use one label in a scan. In the case of the RSNA dataset, for a set of sequences with 28 slices, RSNA-1 has 28 slice-level labels, while we only have 1 sequence-level label. From this perspective, it is unfair to compare our model trained with weak labels with a model trained with full labels such as RSNA-1. However, to compare the differences between the two models, we did the following experiment.

We maintained the same distribution as them based on the results they submitted ⁵¹.

We convert the image-level prediction of the RSNA-1 algorithm to the sequence-level (if a slice reflects a certain disease, the whole CT set will also reflect it), and compare it with the results of DrCT1 and DrCT2. The precision, recall, and F1 score were used to evaluate the performance of the three models, and the results as shown in Table 7.

Table 7: Comparative experiment between sequence-level (DrCT1 and DrCT2) and image-level models (RSNA-1) on RSNA dataset.

Dataset	Model	Precision	Recall	F1 Score
	RSNA-1	93.59%	93.03%	0.9265
RSNA	DrCT1	87.51%	87.83%	0.8650
	DrCT2	88.05%	88.23%	0.8700

The results show that our proposed algorithm has great potential. Compared to image-level algorithms, we can obtain relatively good performance (-5.65% F1 score) using only weakly supervised labels (without using the image-level label). To further analyze the performance in predicting each disease, we calculated the accuracy for the three models. The prediction results are shown in Figure 8. We can see that our model performs better than the

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DrCT1 model. For EDH, the performance of our model is even better than that of RSNA-1 (+0.04% accuracy).

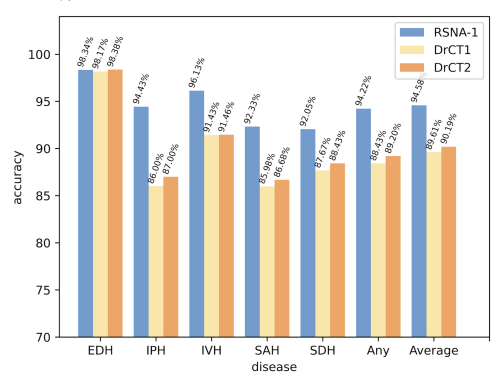


Figure 8: The accuracy of the three models on each disease.

486 VI.B. Human Radiologists Evaluation

Although these evaluation metrics in the above experiments can reflect the performance of the algorithm, how much help the artificial intelligence algorithm can bring to the radiologists in the real scene needs to be evaluated. For this reason, we designed experiments to simulate clinical scenarios to verify whether junior radiologists can be helped with our algorithms. We invited three junior radiologists (junior radiologists A, B, and C) with more than 2 years of experience to perform these experiments. Note that, we did not notify junior radiologists of the performance of the model in advance to prevent humans from having some prior judgments on the results of our model. We measured it from three aspects: diagnostic accuracy, diagnostic time, and diagnostic confidence. We use precision, recall, and F1-score to evaluate diagnostic accuracy. The diagnostic confidence is divided into four levels from 0 to 3: 0 represents a completely uncertain diagnosis; 1 represents a slightly certain diagnosis;

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⁴⁹⁸ 2 represents a high probability diagnosis, and 3 represents full confidence in the diagnosis.

The experimental data are 148 sets of brain CT scans in the RSNA dataset. And they are

randomly selected from the test set to ensure that the model has never seen the data during

the training process. The labels given by the RSNA dataset are used as the gold standard.

The resolution of the CT slice is 224 × 224. The details of the experiments can be seen

below:

- Experiment 1 (radiologist only): The radiologists diagnose the selected brain CT scan, and record the radiologist's diagnosis time and diagnosis conclusion for each set of data. The three radiologists need to score their diagnosis with four diagnostic confidence levels (the scoring time was not included in the diagnosis time) and record the degree of confidence.
- Experiment 2 (radiologist + AI): After an interval of one week from the start of experiment 1, maintain the radiologists are in the same state and randomly shuffle the data sequence to ensure that they forget the impression of the data in experiment 1. The radiologists combine the report generated by the DrCT2 algorithm to diagnose the selected brain CT scan. The experimental report contains three parts, as shown in Figure 2, including all 28 CT slices, their attention weight histograms; the top 5 images with high attention, and their highlighted areas generated by slice attention and feature attention respectively; and the model prediction results.

Table 8: Human Radiologists evaluation task for Experiment 1 (radiologist only) and Experiment 2 (radiologist + AI model).

	I	Experiment	1	E	Experiment 2	2	Experiment
	JR A	JR B	JR C	JR A	JR B	JR C	DrCT2
Date (2021)	11.03	11.04	11.06	11.11	11.10	11.13	×
Precision	0.7411	0.6154	0.7127	0.7558	0.7522	0.8081	0.8551
Recall	0.7364	0.5715	0.7045	0.7474	0.7457	0.8048	0.8454
F1-score	0.7295	0.5780	0.6992	0.7423	0.7348	0.7977	0.8337
Confidence	2.48	2.05	2.10	2.41	2.65	2.48	×
Total time	3h9m27s	2h52m2s	2h39m44s	2h52m28s	2h25m43s	2h22m4s	2 m1s
Time/scan(s)	76.80s	68.88s	64.76s	69.92s	62.29s	57.59s	0.82s

The results of our experiment can be seen in Table 8. From the experimental results, we can find that our model can be more accurate than junior radiologists. With the help of

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our model, radiologists have effectively improved the diagnostic accuracy (+0.16 F1-score), efficiency (-10s average diagnosis time), and more confidence. For radiologist B, we found that even with the help of our model, it still cannot surpass the diagnostic accuracy of our model. The reason for this problem is insufficient trust in our model, although the diagnostic confidence has been improved. We analyzed this issue in detail in the Discussion section.

VII. Discussion

From the perspective of algorithm performance: Our model combines normal resolution and high-resolution input by the primary network and the APS network. It can 526 achieve better performance evaluated by the above experiments. Under the same primary 527 network, DrCT2 effectively improves the performance compared with DrCT1. To compare 528 the performance of DrCT2 and DrCT1 on the CQ500 and RSNA datasets, we reported the 529 precision, recall, and F1 score, and determined statistical significance by using a t-test with 530 a threshold of 0.05 (P<0.05). The P-values for precision, recall, and F1 scores are 0.01940, 0.04971, and 0.02634, respectively. This improvement is because of the APS network. The 532 cooperation of the APS network and the primary network can help the model to improve 533 performance and also help guide the attention mechanism to better select the key slices. As 534 shown in Tables 5 and 6, DrCT2 achieved better performance than DrCT1 in the interpre-535 tive evaluation task. Selecting slices more accurately not only helps improve performance 536 but also enables the model to generate more accurate reports to better assist doctors. It is 537 worth noting that improving interpretability is more important than improving performance 538 because interpretability is a prerequisite for the clinical application of medical models. 539

We did not use some mechanism to force the model to focus on adjacent slices, but the model automatically focuses on sequence slices from 15 to 21 under the supervision of the APS network. This proves that our auxiliary (APS) network is more inclined to make judgments in conjunction with adjacent slices rather than making decisions based on scattered slices. This is consistent with the doctor's diagnosis habit, which is to diagnose by adjacent slices. We explained its necessity in Section III.. These slices are indeed more likely to reflect cerebral hemorrhage from a medical point of view. Both the performance improvement and the medical point of view have proven that the main and auxiliary network (APS network) mechanism we proposed is effective.

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From Table 4, we found that it is meaningless to learn only on high-resolution slices. For example, DrCT1 achieved the lowest performance under 512 × 512 resolution (-1.57% F1 score than original image resolution). Even under the model fusion strategy, it is less effective than the original 224 × 224 resolution (-0.83% F1 score). Our strategy of learning the key slices through the auxiliary network (APS network) again can achieve better results, regardless of whether the original resolution or high-resolution images are input in the auxiliary network (APS network). If we input high-resolution slices in the APS network, the best performance will be achieved. This is also in line with the characteristics of doctors' diagnosis, that is, to pay more attention to the key slices. In addition, key slices were zoomed in on to find detailed clues for diagnosis.

In Table 7, we can see that although we still have a gap with the image-level supervision algorithm. However, we have achieved better performance (+0.04% accuracy) in the EDH category with the least data, even if we only use relatively weak labels. This reflects the potential of our model on small datasets.

There are some limitations to the deep learning method applied in the medical domain. The main factors could include: enough high-quality annotated data, temporality (the diseases are always progressing and changing over time in a non-deterministic way), domain complexity (e.g., different imaging protocols, different types of data) and interpretability ⁵². In our research, we attempt to address data limitations with pre-training on ImageNet, but natural images are different from medical images. We provide explanations at the sequence and slice level, but in clinical applications, doctors expect the AI model to provide more explanation for its predictions. How to learn more information from unlabeled medical data through self-supervised learning, and provide more explanatory is our future direction.

From the perspective of human-computer interaction: In medical applications, more attention should be given to how the algorithm assists the doctor in improving the diagnostic accuracy (rather than maximizing the prediction accuracy of the algorithm itself). In addition, one may also focus on how an algorithm could minimize clinician time or maximize confidence in the diagnosis, for example.

From this point of view, we evaluated the algorithm using two experiments. The first one focused on explainability (whether the model pays attention to suspicious slices) and the page 28 Ruiqian Wang

second on the prediction gain from the human-computer interaction. Through the experimental results as shown in Table 5 and Table 6 we have proven that our model can better select the key slices than DrCT1. We measure the performance changes after AI model assistance from three perspectives: diagnostic accuracy, diagnostic time, diagnostic confidence. From Table 8, we can summary that:

- Our model has better performance than three junior radiologists;
 - Radiologists have improved the diagnostic efficiency and accuracy with the help of AI;
 - Diagnosed with the aid of AI will be more confident usually, but there are also situations that confidence has declined, such as junior radiologist A. Decline in confidence causes low accuracy of improvement.

Both DrCT1⁷ and this study considers the important role of doctors in medical AI. The differences are as follows:

- Experiment participants: in DrCT1, the participant in the evaluation is a medical expert; in this paper, the participants are three junior radiologists. Doctors at two professional levels can evaluate and experience from different perspectives.
- Evaluation angle and the role of AI: In DrCT1, the reliability of the model was evaluated by whether an expert can make a diagnosis based on the selected key images. There was no AI involved in decision-making. In this study, the algorithm generates reports to assist doctors in diagnosis, and experiments can demonstrate the impact of our AI model on doctors' diagnostic decisions.

Our innovation not only includes the improvement of algorithm performance, but also takes into account the role of doctors, and is committed to building models that are more likely to be applied to the real world.

In the clinic, doctors not only give a diagnosis through medical imaging but also need to comprehend the clinical case history and body examination. Therefore, we will comprehensively consider multi-modal data for more comprehensive prediction in our future research.

We found that our model has the potential to learn abnormal brain structures. As shown in Figure 9, our model predicts correctly: there is no intracranial hemorrhage in

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this set of CT scans. However, if the doctor pays attention to the proposed slices with high attention, the focus of encephalomalacia will be found. This may help doctors to avoid missed diagnoses.

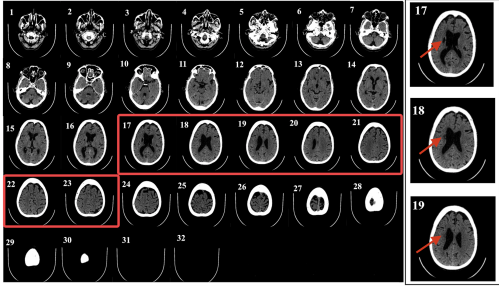


Figure 9: The red box circles the slices selected by the slice attention mechanism. The slices with ID 17, 18, and 19 contain cerebral infarction lesions.

Insufficient trust in the AI model & how to gain the trust of humans: We observed 610 (from Table 8 that the use of artificial intelligence models could improve the diagnostic ac-611 curacy of junior radiologists. However, the combined (human-computer) accuracy remained 612 lower than that achieved using AI only. One reason for this may be the lack of trust in the 613 AI model, as seen for JR A in Table 8. This emphasizes the need for explainability of the 614 AI systems, which can convince the practitioners of their pertinence. From our interviews 615 with radiologists, they emphasized that prompting suspicious slices and highlighting key 616 areas will help AI assist practitioners to clarify their diagnosis. Future directions include 617 improving human-computer interaction performance, assisting practitioners, and improving 618 patient care. 619

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。VIII. Conclusion

Our contributions are as follows: first, we proposed a model DrCT2 that can simultaneously 621 detect multiple brain diseases, which imitates the reading habits of human experts: observing 622 closer at key images from a set of slice scans and observing suspicious lesions for diagnosis. 623 The performance of the model was evaluated on two open-access datasets, with the F1 scores 624 of 0.9370 and 0.8700. Second, we proposed three tasks to evaluate the performance of the algorithm for selecting key images. The accuracy of our model on these three tasks was 626 81.59%, 58.76%, and 65.72%, and it achieved better performance than DrCT1. This proves 627 that our primary and auxiliary network coordination mechanism can achieve better results. 628 The three tasks are of great significance for evaluating the interpretability of the model. 629 The key slices selected by the algorithm can provide a reference for the doctor's diagnosis 630 and reduce the occurrence of misdiagnosis and missed diagnosis. From the perspective of 631 human-computer interaction, we invited three junior radiologists to verify the diagnostic 632 effect after using the model. Experimental results prove that our model can effectively 633 improve the accuracy of diagnosis and help doctors improve efficiency. The algorithm avoids 634 complex annotations, is easy to implement and explanatory, and has good application value. 635 In future work, we will explore the potential of the algorithm in small sample data and 636 continue to increase the application potential of the algorithms.

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