

Comparative study of the quantitative accuracy of oncological PET imaging based on deep learning methods

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Background: [¹⁸F] Fluorodeoxyglucose (FDG) positron emission tomography/computed tomography (PET/CT) is an important tool for tumor assessment. Shortening scanning time and reducing the amount of radioactive tracer remain the most difficult challenges. Deep learning methods have provided powerful solutions, thus making it important to choose an appropriate neural network architecture.

Methods: A total of 311 tumor patients who underwent ¹⁸F-FDG PET/CT were retrospectively collected. The PET collection time was 3 min/bed. The first 15 and 30 s of each bed collection time were selected to simulate low-dose collection, and the pre-90s was used as the clinical standard protocol. Low-dose PET was used as input, convolutional neural network (CNN, 3D Unet as representative) and generative adversarial network (GAN, P2P as representative) were used to predict the full-dose images. The image visual scores, noise levels and quantitative parameters of tumor tissue were compared.

Results: There was high consistency in image quality scores among all groups [Kappa =0.719, 95% confidence interval (CI): 0.697–0.741, P<0.001]. There were 264 cases (3D Unet-15s), 311 cases (3D Unet-30s), 89 cases (P2P-15s) and 247 cases (P2P-30s) with image quality score \geq 3, respectively. There was significant difference in the score composition among all groups (χ^2 =1,325.46, P<0.001). Both deep learning models reduced the standard deviation (SD) of background, and increased the signal-to-noise ratio (SNR). When 8%PET images were used as input, P2P and 3D Unet had similar enhancement effect on SNR of tumor lesions, but 3D Unet could significantly improve the contrast-noise ratio (CNR) (P<0.05). There was no significant difference in SUVmean of tumor lesions compared with s-PET group (P>0.05). When 17%PET image was used as input, SNR, CNR and SUVmax of tumor lesion of 3D Unet group had no statistical difference with those of s-PET group (P>0.05).

Conclusions: Both GAN and CNN can suppress image noise to varying degrees and improve image quality. However, when 3D Unet reduces the noise of tumor lesions, it can improve the CNR of tumor lesions. Moreover, quantitative parameters of tumor tissue are similar to those under the standard acquisition protocol, which can meet the needs of clinical diagnosis.

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Introduction

In terms of the detection, treatment, and follow-up of cancer, positron emission tomography (PET) is a sensitive, noninvasive imaging method that provides molecular-level metabolic information and fine anatomical morphological information of lesions (1-4). In clinical practice, the activity and acquisition time of injected radiopharmaceuticals are often limited by safety, patient tolerance, or compliance. Reducing acquisition time may positively impact patient comfort and increase patient throughput in the Division of Nuclear Medicine. Children, healthy volunteers, and cancer patients should receive lower doses of the tracer to reduce radiation exposure, as well as when multiple followup scans are performed and different tracers are used to monitor the progress of treatment. However, reducing the injection dose/acquisition time can increase image noise, reduce the signal-to-noise ratio (SNR), and increase potentially unnecessary artifacts, thus affecting diagnostic and quantitative accuracy.

In recent years, large datasets have been made available, optimization algorithms have advanced, and efficient network structures have emerged, which have been widely used to segment images (5), detect lesions (6), and restore image super-resolution in computer vision tasks (7). Medical imaging has recently benefited from deep learning-based reconstruction methods (8-11), such as using convolutional neural networks (CNNs) or generative adversarial networks (GANs) to denoise and reduce artifacts in images and improve the SNR of low-dose PET (LDPET) images. Deep learning has proven remarkably effective in predicting full-dose PET (FDPET) images from either LDPET or ultra-low-dose PET (ULDPET) images. Kaplan and Zhu suggested reconstructing FDPET images from 1/10thdose PET images using residual CNN preserved image edges and metabolic parameters (12). By using LDPET sinograms as 3D Unet inputs, Sanaat reduced the bias of quantitative parameters in 83 brain regions using fulldose sinograms (13). Using 3D conditional generative adversarial networks (3D C-GANs), Wang et al. designed

an end-to-end approach to predict high-dose PET images from the corresponding LDPET images (14). Ouyang *et al.* used GAN to generate PET images of amyloid plaques of high quality and simultaneously applied contrary learning, analyzing features, and perceptual impairments specific to particular tasks to considerably improve high-quality PET images (15). According to our research, PET image noise can be reduced, and image quality can be improved using deep learning. Different network structure frameworks have different absolute quantitative accuracy and visual image quality (16), but the impact on clinical diagnosis of tumors and quantitative accuracy of lesions remains to be explored.

According to our literature investigation, we found that the improved model based on CNN network is more used in PET image recovery for tumor lesions at present (17). Since PET images are gray images with relatively simple semantic meaning and fixed structure, high-level semantic information and low-level features are very important. CNN represented by Unet contains Skip connection and U-shaped structure, which can well retain low-level features and high-level features. The multi-layer continuous image input can provide enough information to distinguish the noise from the organization structure. In the literature of Lu et al. (16), it is confirmed that 3D Unet has the best quantitative performance compared with 2D and 2.5D Unet, so we choose 3D Unet to represent CNN. GAN network is an unsupervised deep learning Model. It generates satisfactory input results through the game learning of two types of Generative Model (generative model and Discriminative model) in the framework. In medical image reconstruction, segmentation, registration and other work, most work is based on Pixel2Pixel (P2P) GAN frame and Cycle GAN frame, P2P GAN has been applied in CT noise reduction, MR Reconstruction and PET noise reduction (18). Therefore, in this study, P2P was chosen to represent GAN.

In this study, we systematically compared two deep learning reconstruction models based on 3D Unet and Pix2Pix GAN to predict FDPET tumor images from LDPET images. The original LDPET (1/6) and (1/12) 3762

Fotal = 311

Figure 1 Proportion of tumor types in the included cases.

ULDPET images were used as input data to the 3D Unet and Pix2Pix GAN networks, respectively, to compare the effectiveness of noise reduction and accuracy in the quantitative measurements of the two deep learning models and explore their clinical applicability.

Methods

Patient clinical data collection

In total, 311 tumor patients (205 male and 106 female) who underwent PET/CT scanning in the Department of Nuclear Medicine at the First Hospital of Shanxi Medical University from March 2020 to December 2021 were included in this study. The tumor types of the included cases are shown in *Figure 1*. Their ages ranged between 20–90 years, with the mean patient age of 61 ± 13 years. This study was conducted in accordance with the principles of the Declaration of Helsinki (as revised in 2013). The study protocol was approved by the Ethics Committee of First Hospital of Shanxi Medical University (No. K-GK001) and informed consent was taken from all individual participants.

PET/CT scanning protocol and reconstruction methods

The patients were scanned using the Discovery MI PET/ CT apparatus (GE Healthcare, USA), and ¹⁸F-FDG was produced by the HM-10 cyclotron and ¹⁸F-FDG synthesis module of the Sumitomo Corporation of Japan (radiochemical purity >99%). The patients fasted for 4 to 6 hours before the examination, and their blood sugar levels were within the normal range. They were intravenously injected with ¹⁸F-FDG 3.7 MBq/kg according to their body weight, and then rested for 50–70 minutes after the injection. Scanning was performed from the base of the skull to the middle of the femur. The CT scanning parameters were as follows: tube voltage 120 kV, tube current 60-150 mA (automatic adjustment in x-y direction/z direction), noise index 18, pitch 0.984:1, slice thickness 2.75 mm, rotation time 0.5 s; PET scan in list mode, 3 min/bed, 5-7 beds. The patients were breathing with normal tidal airflow. Select PET list mode data for image reconstruction. The reconstruction algorithm was TOF (time of flight)-PSF (point spread function)-BSREM (block sequential regularized expectation maximization) with penalty factor β =350 (19). Images collected for 3 min per bed were regarded as full dose PET (FD-PET). Low-dose imaging was selected for the first 15 and 30 s of each bed collection (92% reduction and 87% reduction), which were denoted as 8% PET and 17% PET, respectively. At the same time, the pre-90s of each bed collection time was selected as the clinical Standard protocol collection, which was recorded as standard PET (s-PET).

Deep learning reconstruction methods

3D Unet model

- (I) Preprocessing: perform a normalization operation with a mean value of 0 and a variance of 1 on the PET image taken at a low dose and uniformly crop the image to a size of 256×256×64.
- (II) Model structure: the entire model structure is shown in *Figure 2*. The 3D Unet was proposed based on the improvement in the 3D Unet (20). It includes an encoding structure for capturing semantics and a coding structure for an accurately positioned decoding structure. The entire model uses an LDPET image with a size of 256×256×64 as input, extracts image semantic features through an "encoding structure" constructed by three downsampling layers, uses ReLU as an activation function, and then constructs through three upsampling layers. The 3D Unet model is an improved version of the traditional 3D Unet model in the following three aspects:
 - (i) In the encoding structure, the traditional 3D Unet model applies a max pooling layer to downsample the image, whereas the 3D Unet model uses a convolution operation with a 3×3×3 convolution kernel and stride of 2 to replace the max pooling layer.
 - (ii) In the decoding structure, the 3D Unet model applies transposed convolution to upsample



Figure 2 Process diagram of the 3D Unet model. Conv, convolution; BN, batch normalization; ReLU, rectified linear unit; 3D Unet, a deep network model based on CNN; CNN, convolutional neural network.

the image, whereas the 3D Unet uses bilinear interpolation instead of transposed convolution to remove potential checkerboard artifacts.

- (iii) In the traditional 3D Unet, the downsampled feature image and upsampled feature image are spliced through concatenation. In the 3D Unet, two feature images are directly added.
- (III) Loss function: losses are calculated using the mean squared error (MSE), which is calculated as follows:

$$MSE(X,Y) = \frac{1}{M} \sum_{m=1}^{M} (X_m - Y_m)^2$$
[1]

where M represents the number of pixels, m represents the mth pixel, and X and Y represent the reconstructed and labeled images of the model, respectively.

(IV) Post-processing: the image predicted by the model was stitched to restore the original image size, and the normalized pixel values were mapped back to the data range of the original image, according to the mean and variance.

P2P model

- Model structure: the overall model structure design is based on the P2P framework (21):
 - (i) The PET image generation network adopts an adjusted Unet network. The Unet encoder is downsampled three times, for a total of eight downsamplings. Furthermore, the decoder part is upsampled three times in accordance with the

high-level semantic feature map of the encoder, which comprises more low-level and multidimensional picture features for multi-scale prediction, and skip connection is employed at the same stage.

- (ii) Obtain the actual collected image (3 min) scanned for a long time and the image discriminant model reconstructed by the model. Further, input the real image and generated image into the discrimination network in a different sequence, such as [real image generated image], the corresponding discrimination label is (1, 0). A discriminant network was constructed by stacking multilayer CNN convolution layers. After multiple convolution downsampling, the obtained features were straightened, and the discriminative probability of the real 3 min image and the generated 3 min image were predicted through two fully connected layers (Figure 3).
- (II) Loss function.
 - (i) The loss function in the training stage of model generation is based on the L1 loss (regression loss function), as shown in the following formula:

$$L_G = \frac{1}{N} \sum_{i=1}^{n} \alpha_i * \left| y_{i_{predict}} - y_i \right|$$
[2]

where $y_{i_predict}$ represents the ith layer image



Figure 3 Model flow diagram Pixel2Pixel reconstruction. G, Generator model; D, Discriminator model.

reconstructed by the generative model; y_i represents the real voxel value, corresponding to layer i, after normalization of the image; α_i represents the weight of the layer-i image; n represents the number of layers in the image.

(ii) Bidirectional cross-entropy loss functions were used in the training stage of the discriminant models, as shown in the formula:

$$L_{D} = P * \log Q - (1 - P) * \log (1 - Q)$$
[3]

where P indicates the label (P=0 indicates that the image is generated by the model, P=1 indicates that the image is derived from the real image), and Q indicates the probability predicted by the discriminant model.

(iii) Final loss function obtained from the generation and discriminant models:

$$L_{GAN} = \operatorname{argmin}_{G} \max_{D} E\left(\log\left(L_{D}(g, y)\right)\right) + E\left(\log\left(1 - L_{D}(g, y)\right)\right) + E\left(\log\left(L_{G}(x)\right)\right)$$
[4]

When the image output from the generated model is closer to the real image, the loss of

the generated model G is increasingly smaller, whereas the loss of the discriminant model D is greater, thus realizing the optimization process of training.

(III) Image post-processing stage: all the slices predicted by the network are spliced together in order, and the resultant output in the range of 0–1 is mapped back to the data range of the PET image according to the maximum and minimum values of the original input data.

Training set and test set

In this study, 311 patients were randomly divided into training and test groups in the ratio of 7:3, with 218 patients used for training and 93 for testing; 8% PET and 17% PET raw data, respectively, were used as inputs to train the two models.

Visual assessment of PET image quality

The quality of PET images in the deep learning, 8% PET,

17% PET, and s-PET groups was evaluated by two experienced PET/CT physicians (with 5 and 8 years of experience in PET/CT diagnosis, respectively) without knowledge of image acquisition, reconstruction methods, and clinical information. Visual scoring was performed on a 1–5 scale (22): 1, unacceptable; 2, acceptable (subdiagnostic); 3, average; 4, good; and 5, excellent. When there was a difference between the two physicians, an additional score was assigned by a third physician.

Semi-quantitative analysis of PET images

AK software (Artificial Intelligent Kit, GE Healthcare, China) was used to delineate the spherical volume of interest (VOI) and measure the quantitative parameters of the liver background and tumor lesions in each group of images. On the CT tomographic images, measurements were conducted in the right liver lobe using a 3 cm VOI. The software automatically measured the maximum standardized uptake value (SUVmax), mean standardized uptake value (SUVmean), and standard deviation (SD) in the VOI and calculated the SNR in the liver [SNR = liver SUVmean/SD] (23). Similarly, an automated software program measured the SUVmax and SUVmean of ¹⁸F-FDG high-uptake tumor lesions by manually defining the VOIs on the CT images. The median diameter of the tumor lesions was 2.14 cm (1.65-3.94 cm), and the liver was used as the background to calculate the SNR in tumor lesions (SNR = SUVmax/liver SD) and the contrast-noise ratio (CNR) in tumor lesions [CNR = (lesion SUVmean - liver SUVmean)/liver SD] (24).

Statistical analysis

Data analysis was performed using IBM SPSS 22.0, and measurements with non-normal distributions were expressed as medians (P25, P75). The reconstruction data with 17% PET raw data as input were defined as 3D Unet-30s and P2P-30s groups, and the reconstruction data with 8% PET raw data as input were also defined as 3D Unet-15s and P2P-15s groups. The background of s-PET group images and quantitative parameters of tumor lesions were used as reference standards to evaluate the quality of the reconstructed image and its quantitative accuracy. The kappa test was used to analyze the agreement between the visual scores of the two physicians. The chi-square test was used to analyze the differences in the proportion of cases in the visual score of PET images among the groups. Multiple group comparisons were conducted using the Kruskal Wallis H test with Bonfferoni correction, whereas pairwise comparisons were conducted using the Wilcoxon signed-rank test. Statistical significance set at P<0.05 was regarded as consistent.

Results

Visual assessment of PET image quality

The visual scores of the two physicians for the 3D Unet-15s/3D Unet-30s and P2P-15s/-30s groups and between groups (8% PET, 17% PET, and s-PET) were highly consistent, with a kappa value of 0.719 (95% CI: 0.697– 0.741), P<0.001). Image visual scores of 236 subjects in the 3D Unet-15s group were \geq 3 points, compared with 303 subjects in the 3D Unet-30s group. A total of 280 patients in the P2P-15s and P2P-30s groups had visual scores \geq 3, while 1,130,303 patients in the 8% PET, 16% PET and s-PET groups had visual scores \geq 3, respectively. Statistically significant differences were found in the composition ratio of visual scores between groups (χ^2 =1,252.81 and 1,286.98, respectively, P<0.001 for both groups) (*Figure 4*), and the images of typical cases are shown in *Figure 5* and *Figure 6*.

3D Unet deep learning model based on CNN

The ability to suppress background tissue noise

In this study, we selected liver, lung, aorta and lumbar spine as background tissues, and measured the noise levels of four background tissues in 8%PET, 17%PET and two deep learning model reconstruction image groups.

With 8%PET as the input of the model, 3D Unet can reduce the SD of the background tissue and significantly increase the SNR. Compared with standard dose images, SD and SNR in liver and aorta of the 3D Unet-15s group were lower than those of the s-PET group.

With 17% PET as the input of the model, 3D Unet also reduced the SD of background tissue and increased the SNR. Noise levels in lung and lumbar tissues were not statistically different from those in the s-PET group (*Tables 1-4*).

Effect on noise and SNR of tumor tissue

Regardless of whether 8% PET or 17% PET was used as input to the model, SNRs of tumor lesions in 3D Unet-15s group were higher than those in the original image, but there was no statistical difference in SNR compared with



Figure 4 Distribution of image quality assessments. X-axis represents image grouping obtained by multiple image reconstruction methods. Taking the 15 s PET image as input, the reconstructed image is recorded as 3D Unet-15s/P2P-15s. Similar to the above recording method, another set of images is recorded as 3D Unet-30s/P2P-30s; Y-axis represents the percentage of each score in each group; the legend in the upper right corner represents scores of 1–5. PET, positron emission tomography; s-PET, standard positron emission tomography (per bed time: 90 s); 3D Unet, a deep network model based on CNN; P2P, Pixel2Pixel deep network model based on GAN; CNN, convolutional neural network; GAN, generative adversarial network.

the standard dose group.

NR is an important indicator to measure image contrast. The larger CNR, the larger the difference between the focal signal and the background area, and correspondingly, the better the image contrast. Therefore, we further compared the influence of the deep learning model on the CNR of tumor lesions, when 8%PET or 17%PET was used as the model input, 3D Unet could increase the CNR of tumor lesions to a level similar to that of the standard dose group (*Tables 5, 6, Figures 7, 8*).

Effect on semi-quantitative parameters of tumor tissue

The metabolic parameters of tumor have strong correlation with the evaluation of benign and malignant tumor, the evaluation of tumor efficacy, and the evaluation of tumor prognosis. Therefore, we further observed the influence of deep learning model on semi-quantitative parameters of tumor lesions. When 8%PET was used as model input, SUVmean and SUVmax of tumor lesions in the 3D Unet-15s group were lower than those in the original image group. Compared with standard dose images, SUVmax of tumor lesions in the 3D Unet-15s group was significantly decreased, while SUVmean had no significant difference.



Figure 5 Comparison of reconstructed images using 3D Unet and P2P models with 8%PET images as input. SUVbw, standardized uptake value body weight; 3D Unet, a deep network model based on CNN; P2P, Pixel2Pixel deep network model based on GAN; PET, positron emission tomography; FD-PET, full dose-positron emission tomography; CNN, convolutional neural network; GAN, generative adversarial network.



Figure 6 Comparison of reconstructed images using 3D Unet and P2P models with 17% PET images as input. SUVbw, standardized uptake value body weight; 3D Unet, a deep network model based on CNN; P2P, Pixel2Pixel deep network model based on GAN; PET, positron emission tomography; FD-PET, full dose-positron emission tomography (per bed time: 180 s); CNN, convolutional neural network; GAN, generative adversarial network.

Table 1 Comparison of SE	value of background tissue	between reconstructed images and 1	eal images using 8%	PET images as input
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Group	Liver	Lung	Aorta	Lumbar spine
3D Unet-15s	0.20 (0.14, 0.28)*	0.06 (0.05, 0.10)	0.12 (0.09, 0.15)*	0.27 (0.16, 0.37)
P2P-15s	0.31 (0.23, 0.37)*	0.11 (0.09, 0.13)*	0.16 (0.15, 0.23)	0.27 (0.24, 0.34)
8%PET	0.86 (0.64, 1.06)	0.22 (0.16, 0.25)	0.55 (0.46, 0.77)	1.20 (0.75, 1.57)
s-PET	0.25 (0.19, 0.29)	0.08 (0.06, 0.09)	0.19 (0.16, 0.20)	0.27 (0.23, 0.41)
Н	32.13	35.22	35.11	28.65
Р	<0.0001	<0.0001	<0.0001	<0.0001

The italic font represents a statistical difference between this value and that of the 8%PET group. And * indicates that the value is statistically significant compared with the s-PET group. SD, standard deviation; PET, positron emission tomography; s-PET, standard positron emission tomography (per bed time: 90 s); 3D Unet, a deep network model based on CNN; P2P, Pixel2Pixel deep network model based on GAN; H, H value for the Kruskal-Wallis method; CNN, convolutional neural network; GAN, generative adversarial network.

When 17% PET was used as model input, SUVmean and SUVmax of tumor lesions in the 3D Unet-30s group were lower than those in the original image group, and only SUVmax was not significantly different in the 3D Unet-30s group compared with the standard dose group (*Tables 5,6, Figures 9,10*).

P2P deep learning model based on GAN

The ability to suppress background tissue noise

When 8%PET was used as model input, P2P significantly

decreased SD and increased SNR in four background tissues. Compared with standard dose images, SD and SNR in liver and lungs were higher and lower in the P2P-15s group than in the s-PET group, and SD and SNR in the aorta and lumbar spine were not different from those in the s-PET group.

When 17%PET was used as model input, the SD of background tissue was decreased and SNR was increased in P2P-30s group. Compared with standard dose images, there were statistical differences in SD and SNR in lung and liver in the P2P-30s group, but no statistical differences in aortic

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Group	Liver	Lung	Aorta	Lumbar spine	
3D Unet-15s	13.42 (10.04, 19.71)*	7.83 (6.10, 8.84)*	15.73 (12.15, 22.36)*	7.69 (6.15, 10.82)	
P2P-15s	7.49 (6.66, 10.34)*	3.58 (2.73, 4.36)*	8.14 (7.30, 10.48)	6.29 (5.73, 7.31)	
8%PET	2.69 (1.95, 3.28)	2.00 (1.18, 2.14)	2.34 (1.55, 3.01)	1.51 (1.20, 2.26)	
s-PET	8.29 (7.77, 10.02)	5.00 (4.23, 5.64)	7.95 (6.22, 9.32)	5.95 (4.77, 8.28)	
Н	36.86	40.84	37.06	31.10	
Р	<0.0001	<0.0001	<0.0001	<0.0001	

Table 2 Comparison of SNR value of background tissue between reconstructed images and real images using 8% PET images as input

The italic font represents a statistical difference between this value and that of the 8%PET group. And * indicates that the value is statistically significant compared with the s-PET group. SNR, signal-to-noise ratio; PET, positron emission tomography; s-PET, standard positron emission tomography (per bed time: 90 s); 3D Unet, a deep network model based on CNN; P2P, Pixel2Pixel deep network model based on GAN; H, H value for the Kruskal-Wallis method; CNN, convolutional neural network; GAN, generative adversarial network.

Table 3 Comparison of SD value of background tissue between reconstructed images and real images using 17% PET images as input

Group	Liver	Lung	Aorta	Lumbar spine
3D Unet-30s	0.19 (0.15, 0.24)*	0.06 (0.05, 0.09)	0.12 (0.10, 0.14)*	0.26 (0.20, 0.43)
P2P-30s	0.28 (0.25, 0.33)*	0.10 (0.07, 0.13)*	0.16 (0.13, 0.20)	0.29 (0.22, 0.37)
17%PET	0.45 (0.38, 0.67)	0.12 (0.11, 0.16)	0.35 (0.25, 0.42)	0.66 (0.45, 0.90)
s-PET	0.25 (0.19, 0.29)	0.08 (0.06, 0.09)	0.19 (0.16, 0.20)	0.27 (0.23, 0.41)
Н	29.46	25.18	32.96	19.93
Р	<0.0001	<0.0001	<0.0001	<0.0001

The italic font represents a statistical difference between this value and that of the 17%PET group. And * indicates that the value is statistically significant compared with the s-PET group. SD, standard deviation; PET, positron emission tomography; s-PET, standard positron emission tomography (per bed time: 90 s); 3D Unet, a deep network model based on CNN; P2P, Pixel2Pixel deep network model based on GAN; H, H value for the Kruskal-Wallis method; CNN, convolutional neural network; GAN, generative adversarial network.

Table 4 Con	parison of SNR	value of background	tissue between	reconstructed	images and r	eal images us	ing 17%PET	🛾 images as inp	ut
	1				0	0	0	0 1	

Group	Liver	Lung	Aorta	Lumbar spine
3D Unet-30s	11.29 (10.11, 12.95)*	7.40 (5.00, 9.01)*	13.85 (9.15, 17.47)*	7.19 (5.34, 11.07)*
P2P-30s	11.32 (9.46, 14.59)*	5.88 (4.43, 7.75)*	13.31 (10.05, 15.87)*	8.55 (6.78, 10.74)*
17%PET	4.70 (3.49, 5.19)	2.64 (2.03, 3.17)	3.93 (2.84, 5.30)	3.15 (2.05, 3.81)
s-PET	8.29 (7.77, 10.02)	5.00 (4.23, 5.64)	7.95 (6.22, 9.32)	5.95 (4.77, 8.28)
Н	31.06	27.54	35.80	27.63
Р	<0.0001	<0.0001	<0.0001	<0.0001

The italic font represents a statistical difference between this value and that of the 17%PET group. And * indicates that the value is statistically significant compared with the s-PET group. SNR, signal-to-noise ratio; PET, positron emission tomography; s-PET, standard positron emission tomography (per bed time: 90 s); 3D Unet, a deep network model based on CNN; P2P, Pixel2Pixel deep network model based on GAN; H, H value for the Kruskal-Wallis method; CNN, convolutional neural network; GAN, generative adversarial network.

1	0 1	1 1		0		
Group –	PET quantitative parameters of tumor lesions					
	SNR	CNR	SUVmean	SUVmax		
3D Unet-15s	33.81 (24.79, 45.52)	9.94 (5.63, 13.72)	3.78 (3.06, 4.86)	6.57 (4.54, 9.61)*		
P2P-15s	20.45 (15.08, 29.24)*	3.97 (1.06, 6.91)*	3.03 (2.62, 3.99)*	4.72 (3.70, 6.27)*		
8%PET	16.38 (10.78, 19.90)	3.19 (2.18, 4.82)	4.25 (3.62, 5.34)	10.81 (7.44, 16.41)		
s-PET	34.82 (24.92, 51.82)	9.68 (5.90, 13.95)	3.93 (3.23, 4.86)	7.29(5.78, 10.59)		
Н	54.90	27.25	16.81	37.28		
Р	<0.0001	<0.0001	0.001	<0.0001		

Table 5 Comparison of multi-group model results for PET quantitative parameters of tumor lesions based on 8% PET images

The italic font represents a statistical difference between this value and that of the 8%PET group. And * indicates that the value is statistically significant compared with the s-PET group. SNR, signal-to-noise ratio; CNR, contrast-noise ratio; SUVmean, mean standardized uptake values for all voxels in the ROI; SUVmax, maximum standardized uptake values for all voxels in the ROI; PET, positron emission tomography; s-PET, standard positron emission tomography (per bed time: 90 s); 3D Unet, a deep network model based on CNN; P2P, Pixel2Pixel deep network model based on GAN; H, H value for the Kruskal-Wallis method; ROI, region of interest; CNN, convolutional neural network; GAN, generative adversarial network.

Table 6 Comparison of multi-group model results for PET quantitative parameters of tumor lesions based on 17% PET images

Groups	PET quantitative parameters of tumor lesions				
	SNR	CNR	SUVmean	SUVmax	
3D Unet-30s	39.29 (29.10, 50.56)	9.40 (4.66, 12.77)	3.55 (3.27, 4.60)*	7.45 (5.42, 9.73)	
P2P-30s	25.31 (19.83, 39.49)*	4.88 (1.18, 8.96)*	3.78 (3.32, 5.09)	5.47 (4.47, 7.51)*	
17%PET	17.61 (13.26, 24.68)	4.39 (3.12, 7.10)	4.17 (3.47, 5.30)	8.37 (6.38, 11.29)	
s-PET	34.82 (24.92, 51.82)	9.68 (5.90, 13.95)	3.93 (3.23, 4.86)	7.29 (5.78, 10.59)	
Н	38.19	17.67	2.63	12.20	
Р	<0.0001	0.001	0.452	0.007	

The italic font represents a statistical difference between this value and that of the 17%PET group. And * indicates that the value is statistically significant compared with the s-PET group. SNR, signal-to-noise ratio; CNR, contrast-noise ratio; SUVmean, mean standardized uptake values for all voxels in the ROI; SUVmax, maximum standardized uptake values for all voxels in the ROI; PET, positron emission tomography; s-PET, standard positron emission tomography (per bed time: 90 s); 3D Unet, a deep network model based on CNN; P2P, Pixel2Pixel deep network model based on GAN; H, H value for the Kruskal-Wallis method; ROI, region of interest; CNN, convolutional neural network; GAN, generative adversarial network.

and lumbar values (Tables 1-4).

Effect on noise and SNR of tumor tissue

When 8%PET or 17%PET was used as model input, SNR of P2P-15s group was higher than that of original image group, but still significantly lower than that of standard dose group (s-PET group).

Then, the influence of P2P deep learning model on CNR of tumor lesions was observed. When 8%PET was used as model input, CNR of P2P-15s group was higher than that of original image group, but significantly lower than that of s-PET group. When 17%PET was used as model input, CNR of tumor lesions in the P2P-30s group showed no statistical difference compared with the original image group, and was significantly lower than that in the s-PET group (*Tables 5,6, Figures 7,8*).

Effect on semi-quantitative parameters of tumor tissue

When 8%PET was used as model input, the SUVmean and SUVmax of tumor lesions in the P2P-15s group were lower than those in the original image group and the standard dose group. When 17%PET was used as model input,

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Figure 7 Comparison of SNRs in tumor lesions between multiple groups. SNRs, signal-to-noise ratios; 3D Unet, a deep network model based on CNN; P2P, Pixel2Pixel deep network model based on GAN; PET, positron emission tomography; s-PET, standard positron emission tomography (per bed time: 90 s); CNN, convolutional neural network; GAN, generative adversarial network.



Figure 8 Comparison of CNRs in tumor lesions between multiple groups. CNRs, convolutional neural networks; 3D Unet, a deep network model based on CNN; P2P, Pixel2Pixel deep network model based on GAN; PET, positron emission tomography; s-PET, standard positron emission tomography (per bed time: 90 s); CNN, convolutional neural network; GAN, generative adversarial network.



Figure 9 Comparison of SUVmean in tumor lesions between multiple groups. SUVmean, mean standardized uptake values for all voxels in the ROI; 3D Unet, a deep network model based on CNN; P2P, Pixel2Pixel deep network model based on GAN; PET, positron emission tomography; s-PET, standard positron emission tomography (per bed time: 90 s); SUVmean, mean standardized uptake value; ROI, region of interest; CNN, convolutional neural network; GAN, generative adversarial network.

SUVmax of tumor lesions in P2P-30s group was significantly lower than that in standard dose group, while SUVmean had no statistical difference (*Tables 5,6, Figures 9,10*).

Discussion

A systematic comparison of the effectiveness of deep learning reconstruction models, based on CNN and GAN, for improving LDPET images was conducted in this study to determine their impact on tumor clinical diagnosis. The results showed that both the 3D Unet and P2P models improved the quality of the LDPET tumor image and improved the objective quantitative parameter SNR of image quality, both in the background tissue and tumor lesions. However, the 3D Unet model was significantly better than the P2P model in improving the quality of the clinical images of the PET tumor; it could meet the accuracy of tumor lesion quantification and reach the level of tumor diagnosis.



Figure 10 Comparison of SUVmax in tumor lesions between multiple groups. SUVmax, maximum standardized uptake values for all voxels in the ROI; 3D Unet, a deep network model based on CNN; P2P, Pixel2Pixel deep network model based on GAN; PET, positron emission tomography; s-PET, standard positron emission tomography (per bed time: 90 s); SUVmax, maximum standardized uptake value; ROI, region of interest; CNN, convolutional neural network; GAN, generative adversarial network.

Deep learning methods using GANs and CNNs have shown promise in medical image denoising (25). Wang et al. (14), Islam and Zhang (26), and Sajjad et al. (27) used a GAN synthesis model to generate PET images of the brain at different stages of progression of Alzheimer's disease. Ouvang et al. also used a GAN model to generate images of a PET scan with a standard dose of amyloid from intensively low-dosage (1%) images (15). Xie et al. achieved a tradeoff between lesion contrast recovery and background noise using an improved GAN model (28). A GAN-based deep learning model was used to restore LDPET images of the brain and body (29-32). The results show that the GANbased model can retain more details in the image while reducing SUVmean and SUVmax errors. However, for the recovery of LDPET images of tumor lesions, an improved reconstruction model based on a CNN was adopted. Using CNN, Kaplan and Zhu obtained better image edge, structure, and texture details and refined the quantitative parameters using 1/10 LDPET images as inputs (12). Lu et al. used a 3D Unet deep learning reconstruction model to significantly reduce the dose of ¹⁸F-FDG in patients with tumors and ensure diagnostic accuracy (16). Furthermore, 2D and 2.5D Unets were found to have inferior quantitative performance when compared to 3D Unet, and when using 10% low counts to predict standard-dose PET images.

3D Unet was able to reduce image noise and SUVmean deviation even for subcentimeter lung nodules, consistent with the results of this study. There are also studies (13,33-35) using fewer doses of PET images, and CNN also shows an advantage in tumor quantification. A 3D Unet model was developed by Sanaat *et al.* to predict FDPET images from 5% LDPET images (13), thereby improving both image quality and SUV accuracy. Wang *et al.* used 6.25% ¹⁸F-FDG ULDPET images and MRI images of the same machine from 33 children and young adults and input them into the CNN network for image quality recovery (35). The results showed a significant improvement in SUVmax values of tumor lesions and reference tissues and achieved similar parameter values as the standard PET clinical scan.

We found that the 3D Unet model was significantly better than the P2P model for denoising the PET images. 3D Unet builds the entire model based on a 3D convolution. With 3D convolution, we can better acquire the three-dimensional spatial information of an image, and the network model can perform more reliably and robustly if the data contain more dimensions; therefore, reconstructed images are of better quality. Simultaneously, compared with the workflow of P2P stitching to obtain the final image layer by layer, the 3D Unet stitching has fewer layers and uses bilinear interpolation to replace the upsampling layer, which can effectively eliminate the checkerboard artifacts in the reconstructed image. The P2P model takes 2D data as the training target; therefore, the processing speed is high and hardware requirement is low. In this study, high-count reconstructions were used as gold standard labels to represent the unknown PET images to be reconstructed. In addition, instead of feeding noisy images directly into the CNN, a feasible set of PET images was determined using the CNN, thus the entire network was more data-driven. The most desirable feature of this model is that it can include more extensive prior information in the image representation, such as interpatient scanning information. In addition, when the prior information comes from multiple resources, such as time and anatomical information, we can use multiple input channels to aggregate the information and let the network decide the best combination during the training phase. In this study, 3D Unet and P2P models had essentially the same ability to reduce noise, restore image quality, and improve SNR in (1/6 and 1/12) LDPET, both of which can meet the image quality requirements of clinical tumor diagnosis. However, the two models had different recovery effects on the accuracy of the quantitative tumor parameters.

A significant reduction was observed in the quantitative parameters SUVmean and SUVmax for cancerous lesions for both models. 3D Unet-30s significantly reduced SUVmean (P=0.016) compared with 90 s, whereas SUVmax did not differ significantly (P=0.313). In the 3D Unet-15s group, SUVmax was significantly reduced (P=0.034), whereas SUVmean was not significantly different (P=0.315). SUVmean may have been affected by differences in the metabolic volume delineated by the tumor. Among P2P-15s/-30s participants, SUVmax was significantly lower than that of the 90 s participants (all P<0.0001). Thus, with the 3D Unet model, the quantitative parameters of the tumors were maintained more accurately.

In addition, we found that the 3D Unet model had obvious advantages in improving the CNR of foci with high FDG uptake. By constraining the measured data, 3D Unet can help recover some small features that have been removed by image denoising methods, facilitating the recovery of CNR (36). This study confirmed the recovery effect of the 3D Unet model on the CNR index of LDPET images, which could allow the CNR to reach the level of 90s-PET images with lower dose (1/12 dose) PET data input and clearly display and effectively detect small lesions (37). However, the size of lesions in this study ranged from 1.65 to 3.94 cm, and stratified analysis was not performed according to the size of lesions. As a result, it is uncertain whether this model would perform well in other tasks (such as detecting small lesions). Moreover, the robustness of the 3D Unet model should be determined by further testing via quantification of other tasks.

For the optimization of model performance, this study also used 1/6 and 1/12 noise level data to train the model. da Costa-Luis and Reader suggested that a 3D Unet network design is robust when only a limited number of training data are available while accepting multiple inputs (including competitive denoising methods) (38). When the test data dose levels exceed the training dose levels, a new training session is required to achieve an optimal neural network performance. In this study, data with two noise levels were used to train the model separately, instead of superimposing the two types of data. At the same time, some studies have proposed that training models with anatomic morphological imaging information and LDPET images can further improve the image details and quantitative accuracy of lesions. In addition, a combination of different PET reconstruction algorithms, such as the Bayesian algorithm (BSREM) with a CNN network, has also been used to optimize model performance (39).

The applicability of this model is limited, mainly because of the SiPM digital PET/CT equipment and data obtained based on the BSREM reconstruction algorithm. However, the equipment used in most clinical trials is not uniform and PET data with the same acquisition reconstruction parameters cannot be used. Moreover, the study was limited in sample size, which made it impossible to apply fair judgment to the performance of the deep learning model. To make the model more robust, the training dataset must be increased. In addition to cross-validating the generalizability of the model, we hope to conduct more multicenter studies. Furthermore, a variety of tumors were included in this study, and deep-learning model research focusing on a single tumor may be more conducive to the clinical application of the model, such as the application of LDPET in the screening of pulmonary nodules.

Conclusions

In this study, the advantages of two different models in the clinical tumor diagnosis environment were compared. Both GAN and CNN can suppress image noise to varying degrees and improve the overall quality of the image. However, when CNN represented by 3D Unet reduces the noise of tumor lesions, it can improve the CNR of tumor lesions and correspondingly improve the salience of tumor lesions. Moreover, the SUVmean and SUVmax of tumor lesions are similar to those under the collection conditions of clinical standard protocols, which can meet the needs of clinical diagnosis.

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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. This study was conducted in accordance with the Declaration of Helsinki (as revised in 2013). The study protocol was approved by the Ethics Committee of First Hospital of Shanxi Medical University (No. K-GK001) and informed consent was taken from all individual participants.

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