Enlarged Training Dataset by Pairwise GANs for Molecular-Based Brain Tumor Classification

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ABSTRACT This paper addresses issues of brain tumor subtype classification using Magnetic Resonance Images (MRIs) from different scanner modalities like T1 weighted, T1 weighted with contrast-enhanced, T2 weighted and FLAIR images. Currently most available glioma datasets are relatively moderate in size, and often accompanied with incomplete MRIs in different modalities. To tackle the commonly encountered problems of insufficiently large brain tumor datasets and incomplete modality of image for deep learning, we propose to add augmented brain MR images to enlarge the training dataset by employing a pairwise Generative Adversarial Network (GAN) model. The pairwise GAN is able to generate synthetic MRIs across different modalities. To achieve the patient-level diagnostic result, we propose a post-processing strategy to combine the slice-level glioma subtype classification results by majority voting. A two-stage course-to-fine training strategy is proposed to learn the glioma feature using GAN-augmented MRIs followed by real MRIs. To evaluate the effectiveness of the proposed scheme, experiments have been conducted on a brain tumor dataset for classifying glioma molecular subtypes: isocitrate dehydrogenase 1 (IDH1) mutation and IDH1 wild-type. Our results on the dataset have shown good performance (with test accuracy 88.82%). Comparisons with several state-of-the-art methods are also included.

INDEX TERMS Molecular-based brain tumor subtype classification, glioma, multi-modal, MRI, data augmentation, generative adversarial networks, deep learning.

I. INTRODUCTION

Gliomas are one of the most common tumors originating from the brain [1]. World Health Organization (WHO) grades gliomas into four classes (grades I-IV) according to their aggressiveness. The diffuse gliomas are classically divided into low-grade gliomas (LGG, WHO grade II) and high-grade gliomas (HGG, WHO grade III and IV). Pre-surgical prediction or classification is important for clinical decision making and planning. Thus, seeking effective prediction/classification methods on Magnetic Resonance Images (MRIs) may provide non-invasive brain tumor diagnostic tools to assist the medical doctors.

According to previous studies, glioma subtype isocitrate dehydrogenase 1 (IDH1) mutations were observed in 12% of glioblastomas [2], and 50% to 80% of LGG [3]. Patients with IDH1 mutated gliomas have a significant increase in overall survival rate than those with IDH1 wild-type gliomas [4]–[6]. Hence, IDH1 mutation information is important for diagnosis, prognosis and guidance in clinical decisions. Since the IDH1 mutation information is at the molecular level which cannot be easily observed from MRIs even to medical experts, the identification of glioma subtype IDH1 mutation from MRIs is challenging, and it usually requires tissue diagnosis from an invasive procedure (e.g. biopsy or resection) that involves some risks to patients. Successful machine learning methods for predicting the glioma subtypes such as IDH1 mutation from MRIs can offer non-invasively alternative diagnostic tools, though many challenges remain before these tools can be put into clinical use.

A. RELATED WORK

Machine learning methods for characterizing gliomas can be roughly divided into 2 classes: those using hand-crafted
features (i.e., features defined by human experts), and those using deep learning methods for automatically learning the features. Kang et al. [7] proposed histogram analysis of apparent diffusion coefficient maps based on the entire tumor volume for grading gliomas. Carrillo et al. [8] used features from MRIs such as tumor size, frontal lobe localization, presence of cysts and satellite lesions to classify glioma patients between IDH1 mutation and wild-type. Qi et al. [9] studied MRI features like the pattern of growth, tumor margins, signal density and contrast enhancement to predict IDH1 mutation. 

Yu et al. [10] extracted features such as location, intensity, shape, texture and wavelet features for grade II glioma classification. Zhang et al. [11] used texture, histogram and Visually Accessible Rembrandt Images (VASARI) features with a SVM classifier to detect IDH1 and TP53 mutations. Shofty et al. [12] extracted features like size, location and texture of gliomas from images in three modalities, and 17 machine learning classifiers were tested for LGG classification with and without 1p/19q codeletion. The above methods used conventional machine learning methods with hand-crafted features from brain MRIs. Since characterizing glioma features related to molecular (e.g. IDH1 mutation) by purely using MRIs is very challenging to clinicians, defining hand-crafted features could be difficult.

Deep learning methods may offer solutions for such a glioma characterization issue by automatically learning such features. Recently, several deep learning methods for glioma classification have been proposed. Li et al. [13] proposed a six-layer CNN to segment tumors. Fisher vector was then applied to encode deep features from the last convolutional layer using image slices of different sizes followed by feature selection and SVM classifier for IDH1 mutation prediction. Chang et al. [14] proposed to predict IDH1 mutation status of gliomas by applying residual CNNs on multi-institutional MRI data with four different modalities T1 weighted, T1 weighted with contrast-enhanced, T2 weighted and FLAIR (abbreviated as T1, T1e, T2, FLAIR in the text below). Dimensional and sequence networks were tested to evaluate the combination of multi-view and multimodal images. Liang et al. [15] applied 3D DenseNets to predict IDH1 mutation status with multi-modal MRIs. The network also showed high generalization to glioma grade classification such as LGG and HGG.

GANs have recently been used for medical data augmentation. Salehinejad et al. [16], [17] proposed to generate synthesized chest X-rays for enlarging the dataset by deep convolutional GANs. Korkinof et al. [18] progressively trained GANs to synthesize mammograms. Gupta et al. [19] used GAN-based data augmentation method for bone lesion pathology. Despite these promising results, MR brain images are very different from the above medical images such as X-Ray chest images in terms of tissues. The rationale of employing GANs for adding synthetic training MRIs for enhancing the classifier’s performance is as follows. Since the design of GAN is based on the criterion that the synthesized MRIs have similar probability density distributions (pdf’s) as that of the real ones [20]. This is equivalent to adding more dense samples to the original pdf, hence synthetic MRIs enrich the tumor statistics in the original distribution. GANs have been widely used in computer vision for augmenting visual data with great success [21]–[23], however, using GANs for MRI brain tumor data augmentation for molecular-level tumor subtype classification is, to the best of our knowledge, the first successful application.

Our work is mainly motivated by the following challenges, to seek robust methods for enlarging the size of the training dataset in order to cover more tumor statistics in the learning. This issue comes from the real scenarios in medical applications where currently most available glioma datasets are relatively moderate in size, and often accompanied by incomplete MRI scans in different modalities. This may lead to overfitting in training and impact the generalization performance of deep learning classifiers on new test data. Since simple approaches for data augmentation, e.g., flipping, shifting and rotation, do not cover new statistics of tumors and surrounding background, seeking more robust augmentation methods is required to tackle this issue.

We propose a novel scheme to improve the performance of gliomas characterization and subtype classification based on the real and pairwise GAN-augmented MR images in multi-modality forms. The main contributions of the paper include:

(a) Propose a novel scheme for improved glioma subtype classification that consists of three main modules.

• Using pairwise GAN model for data augmentation in a bidirectional cross-modal fashion, for augmenting MR images across different modalities.

• Using a post-processing strategy to achieve the patient-level (3D scan-level) diagnosis result, by combining the 2D slice-level classification results.

• Using a two-stage training strategy for deep learning of real and augmented MRIs from pairwise GANs.

(b) Analyze the performance of the proposed scheme by extensive empirical tests on the glioma dataset, including comparison with some state-of-the-art.

It is worth mentioning that although part of the proposed method was presented in [24], this paper differs significantly in terms of: introducing a pairwise GAN model for data augmentation, hence using much larger dataset by mixing real and GAN-augmented MRIs for training; using a two-stage training strategy for improving glioma subtype classification; using post-processing to combine the slice-level classification results for patient-level diagnosis; using four streams of CNNs as well as attention-weighted feature fusion method; and last, including extensive empirical tests and evaluation on a new glioma dataset containing tumor subtypes of IDH1 mutation and wild-type.
II. PROPOSED GAN-ASSISTED MRI AUGMENTATION FOR GLIOMA CLASSIFICATION

A. OVERVIEW OF THE PROPOSED SCHEME

We propose a novel pairwise GAN-assisted training data augmentation strategy for glioma classification, where the training dataset contains both the real and synthetic MRIs by pairwise GAN augmentation. The main idea behind the proposed scheme is to enlarge the training glioma dataset by pairwise GAN for improved performance of glioma classification, which can be further split as: (a) using pairwise GAN-based data augmentation for enlarging the size of the training dataset. The pairwise GAN is used to augment synthetic MRIs across different modalities as well as augmenting synthetic MRIs for fake patients. It also offers more robustness as GAN-augmented MRIs covers more tumor statistics according to their distributions; (b) using post-processing for patient-level prediction on 3D volume images. Post-processing is used to combine the slice-level glioma classification results for each patient based on 3D volume images. This is realized by applying majority voting on the slice classification results of each patient; (c) using a two-stage training strategy. Although GAN-augmented MRIs and real MRIs are rather similar visually, they still have some differences in their distributions [20]. Employing augmented MRIs for pre-training is hence more appropriate to capture the glioma features.

FIGURE 1. The pipeline of the proposed glioma classification scheme.

The pipeline of the proposed scheme is shown in Figure 1. It uses four modalities of MRIs as the inputs (T1, T1e, T2 and FLAIR). In the proposed scheme, 2D image slices are extracted from 3D volume scans in four modalities, and they are partitioned to the training, validation and testing subsets. After that, a pairwise GAN model is employed to generate synthetic MRIs for the training dataset. Real and GAN-augmented MRIs are then utilized to learn the features and the classifier for brain tumor subtypes. Finally, post-processing is conducted for the patient-level diagnosis based on majority voting of slice-level glioma subtype classification results. The main new contributions of this paper are the pairwise GANs and the patient-level-based post-processing parts, as shown in the dashed boxes. In the following, detailed descriptions on these blocks containing the main contributions will be given.

B. PAIRWISE GAN FOR MR IMAGE AUGMENTATION

The pairwise GAN (Generative Adversarial Network) is employed for the augmentation of MR images. A conventional GAN consists of a generator and a discriminator [20]. A generator produces fake images designed to be as similar as possible to the real images in the sense they have similar probability density distribution function. The discriminator is designed to distinguish between the real and fake images. The generator and discriminator are connected and trained iteratively in alternations through iterations to reach an optimal solution. The pairwise GAN uses a pair of inputs in two streams, different from conventional GAN that contains one stream of input.

1) FORMULATION OF THE PAIRWISE GAN

Let the input MR images consist of $M$ modalities ($M = 4$: T1, T1e, T2 and FLAIR in our case). Let us define the image set for the $m$-th modality as $X_m = \{x_{i,m}, i = 1, \ldots, N_m\}$, where $x_{i,m}$ is the $i$-th 2D slice image in the $m$-th MRI modality. Let us consider a pairwise GAN with two input streams, $X_m = \{x_{i,m}\}$ and $X_n = \{x_{i,n}\}$, whose distributions are $x_{i,m} \sim P_{\text{data} m}$ and $x_{i,n} \sim P_{\text{data} n}$, respectively. The aim of the pairwise GAN is to augment images by using a cross-modality generator and discriminator that alternatively generates synthetic (fake) images as close as possible to the real ones according to their probability distributions.

Let the generator and discriminator for the $m$-th modality be denoted by $G_m$ and $D_m$, respectively. In the pairwise GAN, a pair of inputs are being fed into stream-1 and stream-2 of the GAN from $m$-th and $n$-th modality. For stream-1, $G_m(\cdot)$ has the input $x_{i,m}$ and generates the output $\hat{x}_{i,n}$. The discriminator $D_n(\cdot)$ is to distinguish between the real and fake images ($x_{i,n}, \hat{x}_{i,n}$) in the $n$th modality.

FIGURE 2. Example of the pairwise GAN model.

Conversely, for stream-2, $G_n(\cdot)$ has the input $x_{i,n}$ and generates $\hat{x}_{i,m}$ as the output. The aim of the discriminator $D_m(\cdot)$ is to distinguish between the real and fake images ($x_{i,m}, \hat{x}_{i,m}$) in the $m$th modality, similar to the case in stream-1. The pairwise GAN (shown in Figure 2) defines the cost function such that the two generators and discriminators are jointly optimized. These cost functions will be defined in subsequent sections.

It is worth mentioning that the proposed pairwise GAN is designed to handle two scenarios: (a) Augmenting images of fake patients (that is, to create synthetic MRIs for fake patients in order to enlarge the training dataset); (b) Augmenting one missing modality of MR image from another
modality of MRI from a same patient (that is, when some scan modalities from a patient is missing). To deal with the first scenario (a) all existing modalities are used for augmentation, more synthetic MRIs can be generated through this cross-modality manner to enlarge the training dataset. These synthetic MRIs are considered as belonging to fake patients. As the original dataset contains 4 modalities of images, 6 pairs of pairwise GANs are trained for data augmentation (each is used for one pair of two modalities). For the second scenario (b) one pair of modalities is used, to save the computation where augmentation of one modality image is performed by choosing a best suitable modality pair, e.g. if T1 MRI is missing, T1e is used for GAN augmentation; if both T1 and T1e are missing, FLAIR is selected for augmentation; if both T1 and T1e are missing, then FLAIR or T2 is used (and vice versa).

2) ARCHITECTURE OF THE PAIRWISE GAN

The architectures of the generator and discriminator are shown in Figure 3. For the generator, a U-Net architecture is employed to transfer the low-level information through skip connection. The down-sampling path contains four convolutional layers with $4 \times 4$ filter size and stride $= 2$. The number of filters is set to 32, 64, 128, 256, respectively, and each convolutional layer is followed by a leaky rectified linear unit (LeakyReLU) as the activation function, to introduce the non-linear characteristics to the network and generates a small positive gradient when a neuron is not active. After that, an upsampling path is employed, which consists of four convolutional layers with $4 \times 4$ filter size, stride $= 1$ and ReLU as the activation function except Tanh is used in the last layer to generate the output image. The number of filters is set to 128, 64, 32, 1 respectively. It is worth noting that three skip-connections are used by the step concatenation, which combines the low-level features with up-sampled high-level features for obtaining better performance. For the discriminator, a Markov discriminator [21] is used to determine whether the image patches are real or fake. The least squares loss function rather than the conventional negative log likelihood is then used for obtaining stable and better training results. The discriminator consists of five convolutional layers with filter size $4 \times 4$ and the number of filters 64, 128, 256, 512, 1 respectively. The first four convolutional layers use stride $= 2$ and LeakyReLU, while the last one uses only stride $= 1$ and outputs $8 \times 8$ image patches for the discrimination.

3) LOSS FUNCTION OF THE PAIRWISE GAN

The steps for distinguishing real and fake images in the $n$-th modality can be described as follows: first, the generator $G_m$ in stream-1 uses $x_{i,m}$ to generate fake $\hat{x}_{i,n}$, and then the discriminator $D_n$ in stream-2 distinguishes the fake image $\hat{x}_{i,n}$ from the real one $x_{i,n}$. This can be described as firstly seeking the mapping function for $G_m$: $x_{i,m} \rightarrow \hat{x}_{i,n}$.

The discriminator $D_n$ is used to distinguish between the real and fake image, such that $D_n(x_{i,n}) \approx 1$ for the real image, and $D_n(\hat{x}_{i,n}) = G_m(x_{i,m}) \approx 0$, while the aim for the generator is to let $D_n(\hat{x}_{i,n}) = G_m(x_{i,m}) \approx 1$. The adversarial loss for the $n$-th modality can be described as

$$L_n = E_{x_{i,m}} [||D_n(x_{i,n}) - 1||^2_2 + E_{\hat{x}_{i,n}} [||D_n(G_m(x_{i,m}))||^2_2]]$$

(1)

where $E$ is the ensemble average over the dataset of the specified modality.

Similarly one may form the steps and cost function $L_m$ for distinguishing the real and fake images in the $m$-th modality.

$$L_m = E_{\hat{x}_{i,n}} [||D_m(x_{i,m}) - 1||^2_2 + E_{x_{i,m}} [||D_m(G_n(x_{i,n}))||^2_2]]$$

(2)

Since the two streams of GANs are interconnected, the loss function for the pairwise GAN can be described as:

$$L(G_m, G_n, D_m, D_n) = L_n + L_m + \lambda_1 L_1(G_m, G_n)$$

(3)

where $L_1(G_m, G_n)$ is the loss on the generated images to measure the pixel-level difference between the fake and real images:

$$L_1(G_m, G_n) = E_{x_{i,m}, \hat{x}_{i,n}} [||M_{i,m} \odot (G_m(x_{i,m}) - x_{i,n})||_1 + ||M_{i,n} \odot (G_n(x_{i,n}) - \hat{x}_{i,m})||_1]$$

(4)

where $M_{i,m}$ and $M_{i,n}$ are the tumor masks for the images $x_{i,m}$ and $\hat{x}_{i,n}$ respectively, and $\odot$ is the element-wise multiplication. In the mask, the intensity values are set to 1.0 and 1/3 in the tumor and the background regions respectively, in order to focus on learning tumor areas. $\lambda$ in (3) is the regularization parameter. The final generator and discriminator are obtained by the adversarial training based on the full loss function in (3):

$$G_m^* = \arg \max_{G_m, G_n} \min_{D_m, D_n} L(G_m, G_n, D_m, D_n)$$

(5)

where the discriminators $D_m$, $D_n$ and the generators $G_m$, $G_n$ are trained iteratively in alternations. For training the discriminators, the aim is to minimize the loss in (3) while fixing
the parameters of the generators, so that the discriminators can distinguish the synthetic images from the real ones. The generators $G_m$, $G_n$ are trained by maximizing the loss in (3) while fixing the parameters of the discriminators, in order to fool the discriminators with the synthetic images, so that the generator is able to produce synthetic images similar to the real ones.

The pairwise GANs differ from the conventional GANs in terms of its aim and the cost function. It is employed mainly for generating synthetic images across different modalities of MRIs (e.g. from FLAIR to T2, or from T2 to T1) in medical images. This also leads to a different cost function of the pairwise GAN where tumor areas are enhanced for generating synthetic MRIs. Although our choice of the generator and discriminator is similar to [21], it is worth noting that the pairwise GAN is used to train two pairs of generators and discriminators simultaneously with tumor mask added as the prior to focus on the tumor regions. It can be used to augment more training brain images considering the size of brain tumor dataset is usually not so big. Besides, missing MRI scans in some modalities in datasets is a very common and practical issue, the pairwise GAN is thus able to generate synthetic data in these missing modalities.

4) TRAINING PROCESS OF THE PAIRWISE GAN
The training process for the pairwise GAN is shown in the following algorithm where the generators and the discriminators are updated in alternation until the maximum training epoch is reached. After the training process, two generators $G_m^*$ and $G_n^*$ are used for synthesizing MRIs across two modalities.


c. Post-processing for patient-level diagnosis based on 3D volume images
Since the number of 3D brain volume images in a dataset is moderate, 2D brain image slices are used as this leads to more training data. This also leads to reduced dimensionality of the input data (noting that a high dimensional input is subjected to “the curse of dimensionality”), hence it can mitigate the overfitting in the training process. In the proposed scheme, 2D MRI slices are extracted from three different views (axial, coronal, sagittal) in each modality, which increases the diversity of input 2D MRI slices and prevents the network from overfitting to specific image views.

As the 2D image-based classifier outputs the glioma subtype for each image slice, it is necessary to make a patient-level decision based on all slice prediction results for each individual patient. Since the output subtypes from the 2D image-based classifier can be different for different slices from a same patient, due to variations of slices and image view angles, it is hence necessary to introduce a post-processing approach in order to obtain a consistent prediction of the glioma subtype for each patient based on the corresponding 3D scan. We propose a majority voting-based criterion for making the final tumor subtype classification on each patient, as depicted in Figure 4. That is, the final diagnosis of a patient is determined by the majority classification results from all corresponding slices of a patient. Let $s_i$, $(i = 1, \ldots, N)$ be the $i$th slice prediction result of glioma subtype, $N$ is the total number of extracted tumor image slices of a patient, $s_i = 1$ if the slice belongs to IDH1 mutation, and $s_i = 0$ if it belongs to IDH1 wild-type. The final glioma subtype prediction result $s$ for this patient is determined by

$$s = \begin{cases} 1 & \sum_{i=1}^{N} s_i > N/2 \\ 0 & \text{otherwise} \end{cases}$$ (6)
D. IMPLEMENTATION ISSUES

1) REVIEW OF MULTI-STREAM 2D CONVOLUTIONAL NEURAL NETWORKS

For the sake of convenience to the readers, the multi-stream 2D slice-based CNN feature extraction and classification scheme [24] is briefly reviewed. The multi-stream 2D CNN architecture consists of four separate streams for learning glioma features from four modalities of MR images followed by modality-level feature fusion, as shown in Figure 5.

For each stream, MR images from a single modality are used as the input, where multi-stream CNNs form a set of parallel independent CNN networks. In such a way, modality-specific features are learned through each individual CNN stream.

The CNN architecture for each stream consists of seven layers, that is carefully selected after numerous empirical tests. Filter of size $3 \times 3$ is used in each layer, similar to the filter settings used in the VGG net [25]. Four feature maps are then extracted from the final convolutional layer (i.e., 7th layer) in the four streams and fed to the next layer for feature fusion and refinement.

Observing that features from different modalities contribute differently and complement each other, we introduce a different fusion strategy, namely, attention-weighted fusion strategy as compared with that in [24]. As shown schematically in Figure 5, we apply weighted sum on these features, so that the weights can be learned adaptively according to the characteristics of features from different modalities.

For the refinement of fused features, a bilinear layer similar to [26], is then employed that maps the fused features to a high dimensional feature space. The refined feature map is obtained by exploiting the interactions of fused features at different spatial locations as $H = (F)'^T F$, where $F' \in \mathbb{R}^{h \times w \times c}$ is reshaped from the fused feature map $F \in \mathbb{R}^{h \times w \times c}$, and $h, w, c$ are the height, width and the number of channels in $F$, respectively. The bilinear layer leads to a high-dimensional feature representation that contains complementary features from different modalities. After the feature refinement in the bilinear layer, the refined feature map $H$ is then fed to the classifier. The classifier consists of three fully-connected layers where the number of neurons is 256, 256, 2, respectively.

2) A TWO-STAGE TRAINING STRATEGY WITH AN END-TO-END TRAINING

For effectively training the networks, we propose a two-stage training strategy, where the augmented and real images are treated separately instead of mixing them during the training. This is based on the observation that distributions from GAN-augmented MRIs still present some differences from the real ones. Hence, a two-stage training strategy is adopted, aimed at learning generic features and fast convergence by applying initial training from augmented MRIs, followed by refined feature learning from real MRIs. The two-stage training is performed as follows: the augmented images are used initially for pre-training the whole network for glioma subtype feature learning and classification, and then the real MRI data is used for refined-training. This has led to some performance improvement as have been noticed from our empirical tests. Furthermore, this baseline scheme of multi-stream CNN feature extraction and glioma subtype classification is able to be trained end-to-end.

III. EXPERIMENTS AND RESULTS

A. SETUP, DATASETS, AND METRICS

1) SETUP

KERAS library [27] with TensorFlow [28] backend was used for our experiments. All experiments were conducted on a workstation with Intel-i7 3.40GHz CPU, 48G RAM and a NVIDIA Titan XP 12GB GPU. The commonly used criteria on the overall accuracy and cross-entropy loss were used for the performance evaluation. The size of the input image slice was 128*128. Class-weight was set to 2 for the class IDH1 mutation and 1 for the class IDH1 wild-type in the multi-stream CNN training, since the number of samples in the IDH1 mutation class was about half in that of IDH1 wild-type class. For the pairwise GAN, 1000 epochs were used for the training, where the learning rate was set to 0.0002 with an Adam optimizer. For the multi-stream...
CNN pre-training, the number of epochs was set to 100, the optimizer Adagrad was used, and a step-wise learning rate was set, i.e., to 0.0001 for epochs $\in [1, 30]$, 0.00001 for epochs $\in [31, 60]$, and 0.000001 for epochs $\in [61, 100]$. For refined training of multi-stream CNNs, the learning rate was set to 0.000001 using 50 epochs. Early stopping strategy was adopted during the CNN training process, where parameters of the network were fixed from a certain epoch when a best validation performance was achieved. Simple data augmentation approaches were used as well, including random horizontal flipping and shifting (maximum 10% of width and height). They were performed only on the training dataset in real time to minimize the memory usage.

2) DATASET

The dataset used in our experiments contains 3D brain volume images from TCGA-GBM [29] and TCGA-LGG [30]. In the dataset, the MR images of each patient consist of four modalities (T1, T1e, T2, FLAIR), the tumor segmentation results, as well as the corresponding molecular-based IDH1 genotype labels as the tumor subtypes.

**TABLE 1.** (a) Dataset information. Females/males: F/M. Four age groups were included: ($\leq 30$) / [30,60) / [60,80) / [$\geq 80$]. Noting that one patient of IDH1 wild-type in the dataset lacks age and gender information. (b) Partitioned dataset.

| IDH1 genotype | # Patients (F/M) | Age group | #Slices in each modality | #Slices in each modality |
|---------------|-----------------|-----------|--------------------------|--------------------------|
| Mutation      | 55(32/23)       | (13/35/7/0) | 55                       | 330                      |
| Wild-type     | 112(44/67)      | (5/55/48/3) | 112                      | 672                      |

(a) Dataset information

| IDH1 genotype | # Slices | Training dataset (Real/GAN Augmented) | Validation dataset | Testing dataset |
|---------------|----------|---------------------------------------|--------------------|-----------------|
| Mutation      | 792 (396/396) | 56 (396/55/188) | 66                 | 66              |
| Wild-type     | 1384 (396/1188) | 138 (396/1188) | 138                | 138             |

(b) Partitioned dataset

Detailed information of the dataset is given in Table 1(a). Observing that the volume of tumor is usually small/medium in size, six slices that contain glioma were extracted from each individual scan. This was done for both classes. For the focused feature learning on the tumor areas instead of the whole brain, tumor masks were applied to enhance the tumor feature learning similar to [24]. For our experiments, the dataset was partitioned into 3 subsets: training, validation and testing, detailed information is shown in Table1(b). All 2D image slices in these 3 subsets were partitioned according to patients, i.e., images from the same patient were kept together in either training subset or the testing subset, as such partition is clinically important.

We define 2 case studies, Case-A is for enlarged training dataset containing fake patients, Case-B is for enlarged training dataset including augmentation of both fake patients and missing scans from some modalities. For Case-A study, the training dataset was formed by the combination of ($S1 + S2$), for Case-B study, the training dataset was formed by the combination of ($S3 + S4 + S5$), as shown in Figure 6.

In Case-A and Case-B studies, $S1$-$S5$ are defined as follows:

$S1$: A subset of original training MR images from all modalities (i.e., 60% MRIs from the dataset in Table 1);

$S2$: A subset of GAN-generated synthetic MRIs for all modalities, which was equivalent to generating synthetic MRIs for fake patients. This subset consisted of 297 fake patients with 1782 MRIs;

$S3$: A subset of original training MRIs in ($S1$) minus 24% of MRIs from four scanner modalities, where 6% of patients’ images were removed in each modality;

$S4$: A subset of GAN-generated synthetic MRIs that were used to replace the missing 24% MRIs in ($S3$);

$S5$: A subset of GAN-generated synthetic MRIs for fake patients in four modalities. It consists of a total of 225 fake patients with 1350 MRIs.

3) METRICS FOR PERFORMANCE EVALUATION

Objective metrics were used to evaluate the performance of the glioma classification, based on the following four kinds of samples.

**True positive**: the IDH1 mutation glioma, and was correctly classified as the IDH1 mutation.

**False positive**: the IDH1 wild-type glioma, but was incorrectly classified as the IDH1 mutation.

**True negative**: the IDH1 wild-type glioma, and was correctly classified as the IDH1 wild-type.

**False negative**: the IDH1 mutation glioma, but was incorrectly classified as the IDH1 wild-type.

Let TP, FP, TN and FN be the number of true positives, false positives, true negatives and false negatives, the three metrics overall accuracy, sensitivity and specificity were defined as follows respectively:

\[
\text{Accuracy} = \frac{TP + TN}{TP + FP + TN + FN}
\]

\[
\text{Sensitivity} = \frac{TP}{TP + FN} \quad \text{Specificity} = \frac{TN}{FP + TN}
\]
B. PERFORMANCE OF THE PROPOSED METHOD: 2 CASE STUDIES

To test the effectiveness of the proposed method for classifying the glioma subtype IDH1 mutation, experiments were conducted with 5 runs, where the partitions of training, validation and test subsets in the dataset were done randomly in each of the 5 runs. Table 2 shows the performance of the 5 runs as well as the average performance based on testing for patient-level diagnosis based on 3D volume images.

| Methods | Mean acc. (%) | Mean sen. (%) | Mean spe (%) |
|---------|--------------|---------------|--------------|
| Prop-A  | 88.82(6.37)  | 81.81(11.13)  | 92.17(4.77)  |
| Prop-B  | 88.23(6.57)  | 79.99(14.94)  | 92.17(4.77)  |

Observing Table 2, the proposed method is shown to be effective on the testing dataset. For Case-A study, a relatively high averaging classification accuracy 88.82% was achieved in 5 runs ($|\sigma| = 6.03\%$), with a sensitivity value 81.81% ($|\sigma| = 11.13\%$) and specificity value 92.17% ($|\sigma| = 4.77\%$). For Case-B study, averaging classification accuracy was 88.23%, with a sensitivity value 79.99% ($|\sigma| = 14.94\%$) and specificity value 92.17% ($|\sigma| = 4.77\%$). Further comparing with the results on Case-A and Case-B, one can see that Case-B has a slightly reduced accuracy ($-0.59\%$) and sensitivity ($-1.82\%$) and a similar specificity value on the average of 5 runs. This slightly reduced performance is expected as Case-B also included missing scanner modalities in some patients’ training data.

C. VISUAL EXAMPLES OF CROSS-MODALITY MRI AUGMENTATION FROM PAIRWISE-GANS

The pairwise GAN is shown to work well both empirically, and from visual observations of randomly selected augmented MR images, where the augmented brain MRIs seem to closely resemble the real ones containing tumors. Some cross-modality synthetic image examples generated by the pairwise GANs are shown in Figure 7, where each row contains the real MRI (in red box) and the remaining 3 augmented MRIs in different modalities (e.g., T1, T1e, T2, or FLAIR).

Observing Figure 7, the augmented images are of good quality, and look rather similar to the real ones. Although the distributions could still be somewhat different from the real images.

We also randomly picked up a small percentage of GAN-generated images, and provided them to a radiologist, who considered them highly resemble the real brain images with glioma. It is worth mentioning that the GAN-generated data was used for generic feature learning in the pre-training stage, while the refined training used the real MRIs for learning more specific features of gliomas.

D. IMPROVED TESTING PERFORMANCE OF CLASSIFIERS TRAINED BY ENLARGED Datasets WITH REAL AND AUGMENTED MRIS

1) COMPARISON OF THE PROPOSED METHOD ON CASE-A WITH BASELINE-1 METHOD

In this set of experiments, we evaluate the impact of adding synthetic MRIs of fake patients to the training dataset. The proposed method was tested using the enlarged training dataset by combining ($S_1 + S_2$). We then compared our proposed method with the baseline-1 method. The baseline-1 method was defined the same as the multi-stream 2D CNN followed by post-processing, however, the training dataset only consisted of the real MRIs, i.e., ($S_1$). Table 3 shows the glioma classification results on the testing set, including the overall performance, sensitivity and specificity on the testing set, both for the proposed method and the baseline method.

Observing Case-A results in Table 3, one can see that the proposed method has obtained improved the accuracy on the test set (average 88.82%, about 2.94% improvement over the baseline-1 method), with a slightly increased standard deviation (3.15%) indicating more variance in the estimation. The results have also shown an increased sensitivity (average 81.81% about 12.73% improvement over the baseline-1 method) indicating increased positive classification rate, and slightly decreased specificity (average 92.17% about 1.94% lower) indicating a slightly more increase on false alarm.
2) COMPARISON OF THE PROPOSED METHOD ON CASE-B WITH BASELINE-2 METHOD
In this set of experiments, we evaluate the impact of adding synthetic MRIs from missing modalities and from fake patients. The proposed method was then tested using the above training data combination ($S_3 + S_4 + S_5$). The baseline-2 method was defined as the same as the multi-stream 2D CNN followed by post-processing. For each run, the training, validation and testing subsets were randomly re-partitioned according to the patients. The mean value and standard deviation were also included. Prop-A: the proposed method on Case-A, Bas-1: baseline-1 method, STD: standard deviation.

TABLE 3. Test results on the dataset in the 5 runs from the proposed method where the enlarged training dataset consisted of ($S_1$)+($S_2$); and from the baseline-1 method where the training dataset only consisted of ($S_1$). Both the baseline and proposed method use multi-stream 2D CNNs for glioma classification followed by post-processing. For each run, the training, validation and testing subsets were randomly re-partitioned according to the patients. The mean value and standard deviation were also included. Prop-A: the proposed method on Case-A, Bas-1: baseline-1 method, STD: standard deviation.

| Run | Accuracy (%) | Sensitivity (%) | Specificity (%) |
|-----|--------------|----------------|-----------------|
| Prop-A/Bas-1 | Prop-A/Bas-1 | Prop-A/Bas-1 | Prop-A/Bas-1 |
| 1 | 88.23/65.29 | 81.81/63.65 | 91.30/95.65 |
| 2 | 91.17/91.17 | 90.90/90.90 | 91.30/91.30 |
| 3 | 79.41/85.29 | 63.63/72.72 | 86.95/91.30 |
| 4 | 97.05/88.23 | 90.90/63.63 | 100.00/100.00 |
| 5 | 88.23/65.29 | 81.81/63.65 | 91.30/91.30 |
| Mean | 88.82(6.37)/ | 81.81(11.13)/ | 92.17(4.77)/ |
| (STD $\sigma$) | 85.88(3.22) | 69.08(13.79) | 93.91(3.89) |

TABLE 4. Test results on the dataset in the 5 runs from the proposed method where the training dataset consisted of ($S_3 + S_4 + S_5$); and from the baseline-2 method where the training dataset only consisted of ($S_3$). Both the baseline and proposed method use 2D multi-stream CNNs for glioma classification followed by post-processing. For each run, the training, validation and testing subsets were randomly re-partitioned according to the patients. The mean value and standard deviation were also included. Prop-B: the proposed method on Case-B, Bas-2: baseline-2 method, STD: standard deviation.

| Run | Accuracy (%) | Sensitivity (%) | Specificity (%) |
|-----|--------------|----------------|-----------------|
| Prop-B/Bas-2 | Prop-B/Bas-2 | Prop-B/Bas-2 | Prop-B/Bas-2 |
| 1 | 88.23/65.29 | 81.81/72.72 | 91.30/91.30 |
| 2 | 91.17/91.17 | 90.90/90.90 | 91.30/91.30 |
| 3 | 79.41/73.52 | 54.54/36.36 | 91.30/91.30 |
| 4 | 97.05/88.23 | 90.90/63.63 | 100.00/100.00 |
| 5 | 88.23/65.29 | 81.81/72.72 | 86.95/88.95 |
| Mean | 88.23(6.57)/ | 79.99(14.94)/ | 92.17(4.77)/ |
| (STD $\sigma$) | 84.09(6.78) | 67.27(19.92) | 92.17(4.77) |

E. EVALUATION OF TUMOR MASK IN GLIOMA CLASSIFICATION AND GAN-BASED MRI AUGMENTATION

1) EFFECT OF MASKS FOR CLASSIFIER
To examine the effect of tumor masks for glioma classification, a comparison was made on the proposed method using MRIs with and without tumor mask enhancement. Table 5 shows the comparison on accuracy, sensitivity and specificity from the proposed method on the testing set.

Observing Table 5, the proposed method using tumor mask enhancement has led to improved classification accuracy (by 7.67%), with much higher sensitivity rate and specificity rate. This also indicates that tumor mask enhancement has led to better glioma classification performance on IDH1 mutation subtype and more balanced results on two subtype classes.

TABLE 5. Test accuracy for overall classification performance, sensitivity and specificity from different methods: with and without tumor mask enhancement. Results of 5 runs are shown in mean value (standard deviation $\sigma$). Acc: accuracy, sen: sensitivity, spe: specificity.

| Mask | Mean acc. (%) | Mean sen. (%) | Mean spe (%) |
|------|---------------|---------------|--------------|
| With | 88.82(6.37) | 81.81(11.13) | 92.17(4.77) |
| Without | 81.15(3.34) | 65.44(9.95) | 88.69(4.95) |

2) EFFECT OF MASKS ON AUGMENTED DATA
To examine the effect of tumor masks on GAN-augmented data, comparisons were made on pairwise GAN generated images with and without tumor mask enhancement. The augmented images were then compared with the real MRIs by using peak signal-to-noise ratio. In addition, the images were projected to the latent feature space by convolutional autoencoder similar to [16]. The encoder had 8 convolutional layers with 3x3 filter size. The number of filters was set to 64, 128, 128, 256, 256, 512 and 512 respectively, and the stride was set to 2 every other layer. The decoder had a reverse structure of the encoder and each convolution was replaced by deconvolution. The input image size was 128*128*4 with 4 modalities were stacked together. The latent feature vector with dimensionality 512 was obtained from the globally max pooled feature map of the last convolutional layer in the decoder. The Euclidean distance between centroids of the feature space was applied to compare the real and GAN-augmented images from methods with and without tumor mask enhancement. Table 6 shows the comparison on two metrics, peak signal-to-noise ratio (PSNR) and the distance to the real images based on autoencoder features (DAEF), on the whole image and on the tumor region only.

TABLE 6. Comparison of real and generated MRIs using peak signal-to-noise ratio (PSNR) and distance to the real images based on autoencoder features (DAEF) from methods with and without tumor mask enhancement. The larger the value of PSNR, the better the performance. The smaller the value of DAEF, the better the performance. Results of 5 runs are shown in mean value (standard deviation $\sigma$).

| Mask | Whole image | Tumor region | Whole image | Tumor region |
|------|-------------|--------------|-------------|--------------|
| With | 24.39 (0.06) | 26.14 (0.13) | 112.40 (17.05) | 275.96 (7.93) |
| Without | 25.99 (0.07) | 25.69 (0.10) | 137.25 (16.33) | 277.00 (8.18) |

Observing Table 6, the proposed method using tumor mask enhancement has led to higher PSNR on the tumor region, although on the whole image PSNR of the method without
mask was higher. It indicated that the proposed pairwise GAN with tumor mask enhancement was able to generate more precise tumor regions. DAEF results in Table 6 showed that the method with mask had the encoded feature closer to that of real images both on the whole image and on the tumor region. It is worth noting that the comparison on the augmented data only gave indications since the two metrics used here did not reflect the image quality from the molecular level.

F. COMPARISONS WITH STATE-OF-THE-ART METHODS

Comparisons were made with existing methods for classifying gliomas subtypes with IDH1 mutation/wild-type. Results are shown in Table 7. It is worth emphasizing that these comparisons can only be used as an indication as the datasets used in these methods were different, with exception of the method [15] in Table 7.

| Method          | # Patients IDH1 mutation/wild-type | Accuracy (%) |
|-----------------|-----------------------------------|--------------|
| Liang [15]      | 55/112                            | 84.60        |
| Proposed        | 55/112                            | 88.82        |
| Yu [10]         | 76/34                             | 80.00        |
| Zhang [11]      | 70/33                             | 80.00        |
| Chang [14]      | 233/263                           | 85.70        |

Observing Table 7, the proposed method was comparable to others, indicating relatively good performance. The proposed method achieved a better result than [15] on the same dataset. Other methods [10], [11], [14] only give indications on relative performance due to using different datasets, which also indicated that the proposed scheme has the performance comparable to the state-of-the-art.

G. DISCUSSIONS

From different sets of experimental results described above, the following insights may be gained on the proposed scheme:

- Overall performance: The proposed scheme is effective, with an excellent classification performance of glioma subtypes (88.82% on the testing set);
- Training the proposed scheme by enlarged datasets with real + pairwise GAN augmented MRIs across different modalities from fake patients has led to improved classification performance (increased 2.94% on the testing set for glioma subtypes of IDH1 mutation/wild-type). This indicates that the pairwise GAN is robust and effective, and can be used as a tool for enlarging the training dataset with mixed real and augmented data with further enhanced generalization performance of glioma subtype classification;
- Tumor masks are effective for glioma subtype classification and GAN-based MRI augmentation. They lead to an increase of classification performance by 7.67%, and more precise tumor regions in the GAN augmented images.

Future work includes applying multiple datasets with cross-domain cross-modality GAN data augmentation, classifying gliomas with additional subtypes of 1p/19q codeletion classes which are important to glioma prognosis, and incorporating patient side information (e.g. ages, survival years), for predicting more glioma subtypes as well as patient survival.

IV. CONCLUSION

The proposed scheme has been tested using real and pairwise GAN-augmented MRIs as training data, results obtained on the testing dataset have demonstrated that the scheme is effective and robust (average 88.82% test accuracy for gliomas subtypes of IDH1 mutation/wild-type). Using two enlarged datasets containing real and GAN-augmented MRIs for training the proposed scheme, has both resulted in increased generalization performance of the classifier on the testing set. This indicates that the proposed pairwise GAN is effective and robust, and is useful for augmenting MRIs when the size of brain tumor training dataset is not sufficiently large for deep learning. The post-processing step is essential for diagnosis based on 3D volume images, and the two-stage training strategy is useful for real and GAN-augmented MRIs. Finally, comparisons with several existing methods, though based on different datasets, have indicated that the proposed scheme using mixed real and GAN-augmented training datasets has reached comparable performance to the state-of-the-art.

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