

Model Generalizability Investigation for GFCE-MRI Synthesis in Radiotherapy of NPC patients Using Multi-institutional Data and Patient-based Data Normalization

Wen Li, Saikit Lam, Tian Li, Jens Kleesiek, Andy Lai-Yin Cheung, Ying Sun, Francis Kar-ho Lee, Kwok-hung Au, Victor Ho-fun Lee, and Jing Cai

Abstract—Recently, deep learning has been demonstrated to be feasible to eliminate the use of gadolinium-based contrast agents (GBCAs) through synthesizing gadolinium-free contrast-enhanced MRI (GFCE-MRI) from contrast-free MRI sequences, providing the community with an alternative to get rid of GBCAs-associated safety issues in patients. Nevertheless, generalizability assessment of the GFCE-MRI model has been largely challenged by the high inter-institutional heterogeneity of MRI data, on top of the scarcity of multi-institutional data itself. Although various data normalization methods have been adopted in previous studies to address the heterogeneity issue, it has been limited to single-institutional investigation and there is no standard normalization approach presently. In this study, we aimed at investigating generalizability of GFCE-MRI model using data from seven institutions by manipulating heterogeneity of training MRI data under two popular normalization approaches. A multimodality-guided synergistic neural network (MMgSN-Net) was applied to map from T1-weighted and T2-weighted MRI to contrast-enhanced MRI (CE-MRI) for GFCE-MRI synthesis in patients with nasopharyngeal carcinoma. MRI data from three institutions were used separately to generate three uni-institution models and jointly for a tri-institution model. Min-Max and Z-Score were applied for data normalization of each model. MRI data from the remaining four institutions served as external cohorts for model generalizability assessment. Quality of GFCE-MRI was quantitatively evaluated against ground-truth CE-MRI using mean absolute error (MAE) and peak signal-to-noise ratio (PSNR). Results showed that performance of all uni-institution models remarkably dropped on the external cohorts. By contrast, model trained using

multi-institutional data with Z-Score normalization yielded significantly improved model generalizability.

Index Terms—Contrast enhanced MRI, data normalization, model generalizability, nasopharyngeal carcinoma

I. INTRODUCTION

Nasopharyngeal carcinoma (NPC), a highly aggressive epithelial carcinoma originating in the mucosal lining of the nasopharynx, has long been prevalent in the population of East and Southeast Asia [1]. Radiotherapy (RT) is currently the mainstay treatment modality for NPC, which achieved 66%-83% 5-year survival rate for NPC patients with RT alone [2]. Precise tumor delineation is the most critical prerequisite for a successful RT treatment. Contrast-enhanced MRI (CE-MRI), using gadolinium-based contrast agents (GBCAs), has become an indispensable part in accurate NPC tumor delineation [3] in routine RT treatment planning practice. Nevertheless, emerging evidence has shown that nephrogenic systemic fibrosis (NSF), a severe disease that can lead to joint contractures and immobility, has been strongly linked with the administration of GBCAs in renal failure patients [4]. Further evidence has shown that gadolinium accumulation in the dentate nucleus and globus pallidus has been observed in paediatric patients [5]. Apart from this, gadolinium deposition was also observed in patients with normal renal function [6]. The mechanism of gadolinium deposition in patients has not been fully elucidated, and the underlying long-term effects remain unclear. Therefore, there is a global consensus to minimize or avoid GBCA exposure to patients whenever possible [4]. Considering this, a GBCA-based CE-MRI alternative is desperately demanded.

Numerous efforts have been made to address the GBCA-associated safety issues. Worldwide interests have sparked recently in synthesizing gadolinium-free contrast-enhanced MRI (GFCE-MRI), which serves similar purposes as the CE-MRI, through deep learning approaches [7-15]. However, current works have focused on model development or feasibility studies at different tumor sites using in-house datasets. It has been reported that the models trained with in-house dataset may perform poorly on datasets from external institutions [16-18], which largely limits the wide application of proposed approaches. Therefore, a generalizable GFCE-MRI model is highly demanded in clinical practice, which extends the GFCE-MRI technique to a considerably wider range of hospitals for use.

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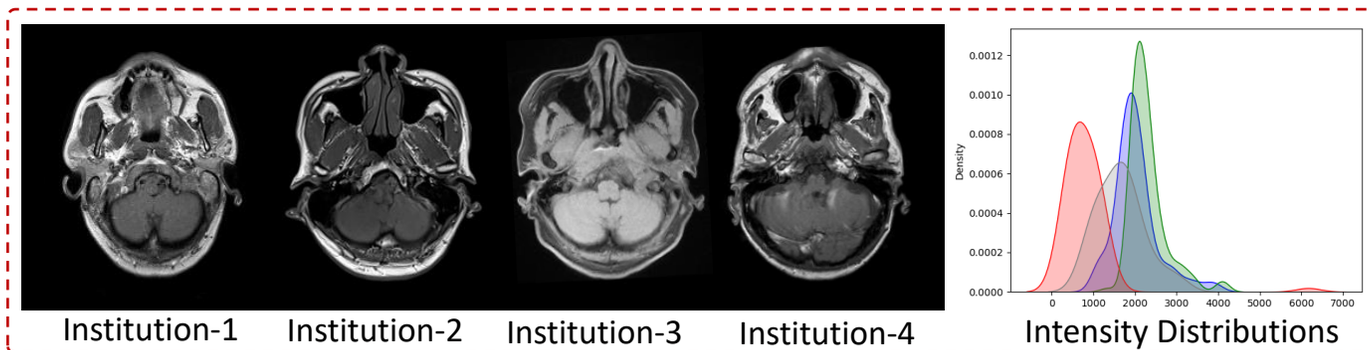


Fig. 1. Illustration of heterogeneous multi-institutional MRI data.

TABLE I
THE OVERALL STUDY DESIGN

Normalization	Model name	Training				Testing			
		Institution-1	Institution-2	Institution-3	Institution-4	Institution-5	Institution-6	Institution-7	
Min-Max	Uni-m1	✓			✓	✓	✓	✓	
	Uni-m2		✓		✓	✓	✓	✓	
	Uni-m3			✓	✓	✓	✓	✓	
	Tri-M	✓	✓	✓	✓	✓	✓	✓	
Z-Score	Uni-z1	✓			✓	✓	✓	✓	
	Uni-z2		✓		✓	✓	✓	✓	
	Uni-z3			✓	✓	✓	✓	✓	
	Tri-Z	✓	✓	✓	✓	✓	✓	✓	

Despite the urgent need for generalizable models, limited research has been conducted to investigate the underlying mechanism of model generalizability and the methods to improve the model generalizability, especially for the multi-parametric MRI images, presumably due to two key challenges: 1) high inter-institutional heterogeneity of MRI data; 2) scarcity of multi-institutional MRI data. The MRI images from different institutions often suffer from large domain shifts due to the use of diverse scanning parameters, scanners of different field strengths, as well as different patient demographics, leading to large distribution divergences such as means, standard deviations, and intensity ranges (Fig. 1). These challenges have raised a growing concern of model generalizability developed using deep learning algorithms, which strongly rely on the assumption that the training data and testing data are independent and identically distributed (i.i.d.) [19]. In reality, however, the external MRI datasets are typically out-of-distribution (OOD) due to the abovementioned domain shift, incurring tremendous performance degradation of the trained models [19]. To tackle this, a potential remedy to improve model generalizability is to integrate multi-institutional MRI images during model training to enlarge view of deep learning models [20, 21], which has been rarely reported in the literature, probably due to the scarcity of multi-institutional data for patient privacy protection. Another potential solution is to develop a generalizable network architecture by mapping data distributions from source domain to target domain [19, 22], while these approaches are limited to specific domain datasets. As such, data normalization techniques have been widely used to improve the model performances in a range of application areas. Nevertheless, related research in multi-institutional setting that contain

various real-world distributions of MRI data is severely scarce in the body of literature.

We consider minimize the distribution variations between training and external testing MRI data by using data normalization should be a practical approach to improve the model generalizability since it requires no model architecture improvement and retraining the model. In this study, we included MRI data from seven different institutions, aiming at investigating the GFCE-MRI model generalizability influenced by distribution difference between training and external testing data. Specially, we investigated: (i) how significant is the influence of different data normalization methods on the model generalizability; (ii) how significant is the degradation of external performance for models trained with single-institution MRI; and (iii) how significant is the improvement of external performance when using multi-institutional MRI for model development.

Compared to other tumor types such as brain and liver tumors, NPC is highly infiltrative with ill-defined tumor-to-normal tissue interface, which presents challenges to oncologists in NPC contouring. Hence, the success of this study may not only provide the medical community with better insights into the issue of GFCE-MRI model generalizability of NPC patients, but also may potentially be translated to other cancer types as well. To the best of our knowledge, this is the first multi-institutional investigation for GFCE-MRI synthesis. As a result, this study may have a far-reaching impact on the medical community to better understand the issue of model generalizability, establish a standard multi-institutional data normalization method, and further facilitate the development of generalizable GFCE-MRI models in the future.

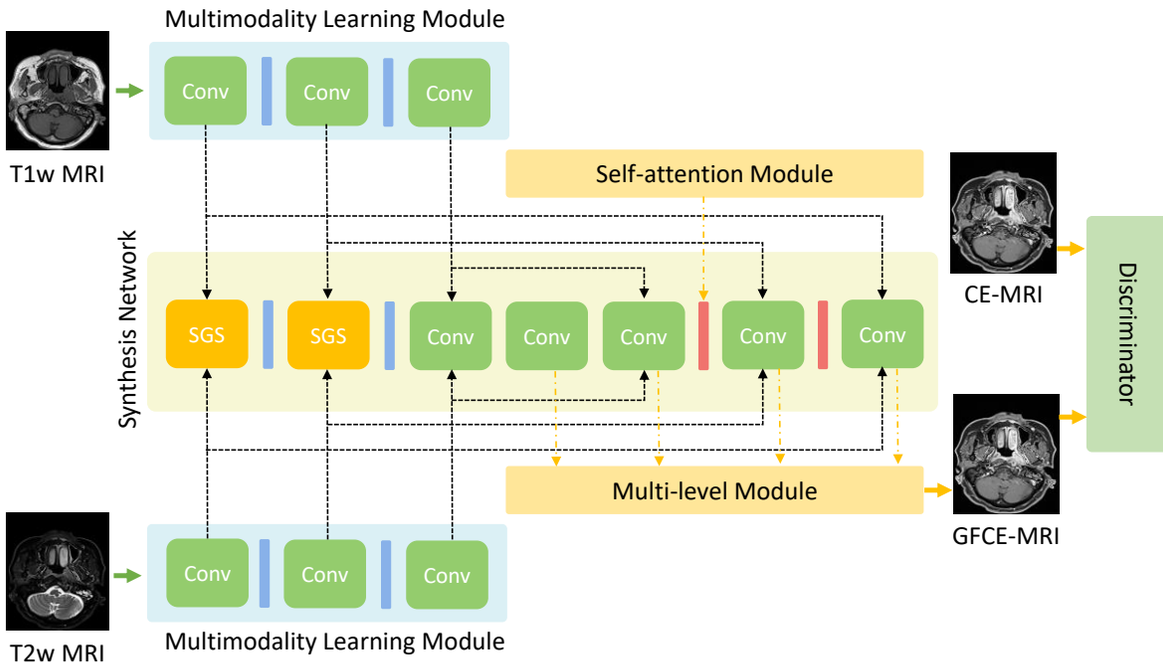


Fig. 2. The architecture of MMgSN-Net. It is a two-inputs network consisting of five key components: multimodality learning module, synthesis network, self-attention module, multi-level module, and a discriminator. T1-weighted MRI and T2-weight MRI were used as inputs, gadolinium-based contrast-enhanced MRI was used as the learning target. SGS, synergistic guidance system; Conv, convolutional layers.

II. METHODS AND MATERIALS

A. Patient Data

A total of 256 patients from seven medical institutions were retrospectively collected in this study. For fair comparisons, same number of patients (71 patients) were retrieved from Institution-1, Institution-2, and Institution-3, respectively for uni-institution and joint-institution model development, 18 patients, 9 patients, 9 patients, and 7 patients were retrieved from Institution-4, ..., Institution-7, respectively for external testing to evaluate the model generalizability. T1-weighted (T1w) MRI, T2-weighted (T2w) MRI, and CE-MRI were collected for each patient. This study was approved by the Institutional Review Board of the University of Hong Kong/Hospital Authority Hong Kong West Cluster (HKU/HA HKW IRB), reference number UW21-412 and the Research Ethics Committee (Kowloon Central/Kowloon East), reference number KC/KE-18-0085/ER-1. Due to the retrospective nature of this study, patient consent was waived. All images were acquired in the same position and automatically aligned. For model training, all images were resampled to the size of 256*224 using bilinear interpolation [23]. For Institution-1, Institution-2, and Institution-3, the 71 patients were randomly divided into 53 and 18 for model training and validation, respectively.

B. Study design

The overall idea of this study was first using the data collected from three different centers (i.e., Institution-1, Institution-2, and Institution-3) to develop a series of separately and jointly trained models using different data normalization methods for investigating the GFCE-MRI model generalizability. The separately and jointly trained models were

referred to as uni-institution models and tri-institution models, respectively. **Table 1** illustrated the overall study design.

1) Neural network

The multimodality-guided synergistic neural network (MMgSN-Net) was used as the base network in this study. The MMgSN-Net is a 2D deep learning algorithm [15], which consists of five key modules: multimodality learning module, synthesis network, self-attention module, multi-level module, and a discriminator. The structure of the MMgSN-Net is illustrated in **Fig. 2**. The T1w and T2w MRI were put into the multimodality learning module separately. The multimodality learning module was used to extract the modality-specific features. The extracted modality-specific features were put into the synergistic guidance system (SGS) in synthesis network for complementary feature selection and fusion. In the decoder of synthesis network, the fused features and the learned features from multimodality learning modules were concatenated to different channels. The self-attention module and multi-level module were applied to capture the long-term dependencies and detect the edge information of the high-level features, respectively. A discriminator was utilized to distinguish the synthetic GFCE-MRI from ground-truth CE-MRI, thus encouraging the synthesis network to generate more realistic GFCE-MRI.

2) Data Normalizations

Data normalization plays a pivotal role in model development [24]. It minimizes feature bias by transforming the features into a common space so that larger numeric feature values cannot dominate smaller numeric feature values [25]. Currently different data normalization methods are applied in medical image translation tasks. The most popular two normalization methods are Min-Max (also called scaling) [26] and Z-Score [27]. These two normalization methods are also applied to different objects, i.e., dataset-based, patient-based,

TABLE II. INTERNAL AND EXTERNAL QUANTITATIVE RESULTS USING MIN-MAX NORMALIZATION

Model	Testing	MAE \pm SD ($\times 10^3$)	PSNR \pm SD
Uni-m1	<i>Institution-1</i>	25.39 \pm 3.59	33.45 \pm 1.38
	Institution-4	52.12 \pm 10.89	27.65 \pm 1.72
	Institution-5	35.03 \pm 6.56	30.47 \pm 1.24
	Institution-6	34.97 \pm 4.02	31.65 \pm 0.67
	Institution-7	40.80 \pm 9.12	29.35 \pm 1.51
	<i>Overall</i>	40.73 \pm 7.65	29.78 \pm 1.29
Uni-m2	<i>Institution-2</i>	24.45 \pm 3.67	32.17 \pm 0.89
	Institution-4	50.26 \pm 7.11	27.50 \pm 0.95
	Institution-5	51.76 \pm 6.28	27.83 \pm 1.02
	Institution-6	58.74 \pm 19.93	27.05 \pm 2.13
	Institution-7	45.27 \pm 3.83	28.41 \pm 0.73
	<i>Overall</i>	51.51 \pm 9.29	27.70 \pm 1.21
Uni-m3	<i>Institution-3</i>	25.56 \pm 6.92	31.30 \pm 1.72
	Institution-4	44.53 \pm 7.63	28.51 \pm 1.32
	Institution-5	35.67 \pm 5.09	30.09 \pm 0.78
	Institution-6	45.36 \pm 15.96	29.41 \pm 2.08
	Institution-7	33.30 \pm 7.81	30.69 \pm 1.48
	<i>Overall</i>	39.72 \pm 9.12	29.68 \pm 1.42
Tri-M	Institution-1	26.27 \pm 4.01	33.06 \pm 1.30
	Institution-2	26.27 \pm 4.19	31.74 \pm 0.86
	Institution-3	28.91 \pm 6.38	31.45 \pm 2.05
	<i>Overall</i>	27.15 \pm 4.86	32.08 \pm 1.40
	Institution-4	41.82 \pm 7.82	28.97 \pm 1.20
	Institution-5	41.55 \pm 9.04	29.19 \pm 1.51
	Institution-6	46.12 \pm 13.55	29.29 \pm 1.84
	<i>Overall</i>	40.76 \pm 9.66	29.51 \pm 1.53

and single-image based normalizations. In natural image tasks, most studies are 2D-based networks, and they normalize their data using statistical values of each single image or the whole dataset [18]. For medical images, however, image and dataset-based normalizations may not appropriate for clinical applications, especially for 3D volumes since the image-based normalization ignores the inter-slice continuous information within a volume, which leads to contrast bias between nearby-slices, while dataset-based normalization brings challenge during model inference for a new patient as only statistical values of this specific patient could be used for data normalization. Herein, we consider that patient-based normalization is proper in medical image studies, which is more applicable to clinical setting. In this study, the patient-based Min-Max normalization and patient-based Z-Score normalizations were applied to evaluate the model generalizability affected by data normalization. The two patient-based normalization methods could be mathematically described as:

$$x_{min_max} = \frac{x - x_{min}}{x_{max} - x_{min}} \quad (1)$$

$$x_{z_score} = \frac{x - \mu_x}{\sigma_x} \quad (2)$$

Where x represent the intensities of each patient volume, while x_{min} , x_{max} , μ_x , and σ_x are minimum value, maximum value, mean value and standard deviation of the patient. x_{min_max} and x_{z_score} are the patient data after Min-Max and Z-Score normalization, respectively. The Min-Max normalization rescales the intensity range to [0, 1] and preserves the relationship among the original data values, while Z-Score normalize the mean value and standardization of the patient to 0 and 1 respectively, which enables the comparison of two datasets with different distributions.

TABLE III. INTERNAL AND EXTERNAL QUANTITATIVE RESULTS USING Z-SCORE NORMALIZATION

Model	Testing	MAE \pm SD ($\times 10^3$)	PSNR \pm SD
Uni-z1	<i>Institution-1</i>	23.03 \pm 3.18	34.21 \pm 1.58
	Institution-4	43.10 \pm 5.91	28.96 \pm 1.20
	Institution-5	32.74 \pm 6.27	31.03 \pm 1.16
	Institution-6	32.07 \pm 5.05	32.36 \pm 1.07
	Institution-7	38.22 \pm 8.77	29.84 \pm 1.42
	<i>Overall</i>	36.53 \pm 6.5	30.55 \pm 1.21
Uni-z2	<i>Institution-2</i>	24.87 \pm 4.64	32.28 \pm 1.10
	Institution-4	48.47 \pm 7.30	27.62 \pm 1.22
	Institution-5	31.35 \pm 7.52	31.33 \pm 1.51
	Institution-6	33.27 \pm 5.23	31.68 \pm 1.14
	Institution-7	37.27 \pm 9.36	29.76 \pm 1.57
	<i>Overall</i>	37.59 \pm 7.35	30.10 \pm 1.36
Uni-z3	<i>Institution-3</i>	26.84 \pm 6.17	31.97 \pm 2.09
	Institution-4	38.30 \pm 5.53	29.50 \pm 1.21
	Institution-5	31.92 \pm 7.32	31.06 \pm 1.42
	Institution-6	30.78 \pm 4.70	32.52 \pm 1.08
	Institution-7	33.51 \pm 8.08	30.95 \pm 1.50
	<i>Overall</i>	33.63 \pm 6.41	31.01 \pm 1.30
Tri-Z	Institution-1	23.71 \pm 3.12	33.72 \pm 1.43
	Institution-2	25.74 \pm 4.80	32.01 \pm 1.10
	Institution-3	27.36 \pm 6.80	31.87 \pm 2.23
	<i>Overall</i>	25.60 \pm 4.91	32.53 \pm 1.59
	Institution-4	37.20 \pm 5.14	29.72 \pm 1.21
	Institution-5	29.94 \pm 6.43	31.69 \pm 1.25
	Institution-6	29.60 \pm 4.94	32.78 \pm 1.12
	<i>Overall</i>	32.45 \pm 6.22	31.27 \pm 1.04

3) Uni-institution models

To investigate how significant is the external performance degradation for models trained with single-institution MRI, we first trained three uni-institution models using data from Institution-1, Institution-2, and Institution-3 for each normalization method separately. 53 patients were used for training of each uni-institution model. The three uni-institution models were labeled as Uni-m1, Uni-m2, and Uni-m3 for Min-Max normalization and Uni-z1, Uni-z2, and Uni-z3 for Z-Score normalization, respectively. We tested each uni-institution model using four external testing datasets (i.e., Institution-4 to Institution-7).

4) Tri-institution models

To investigate how significant is the external performance improvement for models trained with multi-institution MRI, we trained the model jointly with data from three institutions. Considering the number of training samples may influence the tri-institution model assessment since we cannot determine whether the model generalizability improvement is caused by a diverse dataset or an increasement of training samples. Therefore, we randomly selected 18 patients from each institution's training dataset. Then randomly discarded one sample to ensure training samples were the same as the numbers for uni-institution models. The two tri-institution models with different normalization methods are labeled as Tri-M (with Min-Max normalization) and Tri-Z (with Z-Score normalization), respectively. The four testing datasets from Institution-4 to Institution-7 were used for external testing to evaluate the model generalizability.

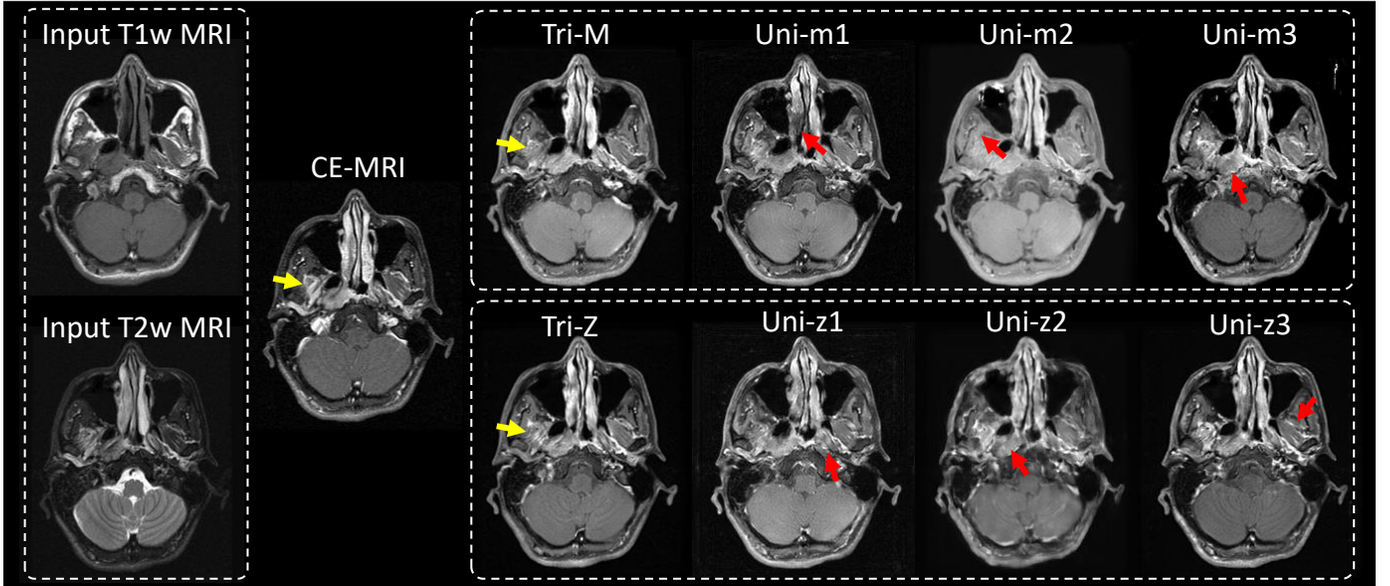


Fig. 3. Illustration of GFCE-MRI generated from uni-institution and tri-institution models using Min-Max normalization and Z-Score normalization.

C. Evaluations

1) Quantitative evaluation

To quantitatively evaluate the performance of uni- and tri-institution models, mean absolute error (MAE) and peak signal-to-noise ratio (PSNR) between the synthetic GFCE-MRI and ground-truth CE-MRI were calculated. The MAE and PSNR have been widely employed for medical image analysis tasks. MAE measures pixel-wise differences while PSNR measures the ratio between the maximum power of a signal and the power of noise [15, 28, 29]. Smaller MAE and larger PSNR values indicate better quantitative results. Prior to quantitative evaluation, we rescaled the CE-MRI and predicted GFCE-MRI intensities to $[0, 1]$ to compute the percentage differences between GFCE-MRI and CE-MRI. Paired two-tailed t-test was performed to analysis if there is significant difference between results from different models.

$$MAE = \frac{\sum_{i=1}^n |y_i - f(x_i)|}{n} \quad (3)$$

$$PSNR = 20 \cdot \log_{10} \frac{\max(y_i) \cdot \sqrt{n}}{\|y_i - f(x_i)\|_2} \quad (4)$$

Where y_i and $f(x_i)$ are intensities of real CE-MRI and GFCE-MRI, n is the number of intensities. Here $\max(y_i)$ is 1 as we have rescaled the CE-MRI and GFCE-MRI intensities to $[0, 1]$.

2) Qualitative evaluation

To visually assess the performance of the models on external datasets, we directly applied the trained uni- and tri-institution models to the external datasets for comparison. The input T1w, T2w MRI and ground-truth CE-MRI were shown alongside the GFCE-MRI generated from different models.

III. RESULTS

A. Quantitative results

1) Generalizability of single-institution models

All uni-institution models suffered from dramatically performance drop on external MRI data for both Min-Max and Z-Score normalizations. **Table 2** and **Table 3** summarize the

quantitative comparisons between the synthetic GFCE-MRI and ground-truth CE-MRI using Min-Max and Z-Score, respectively. As MAE and PSNR have the similar trend, we use the MAE as an indicator to illustrate the results. The average MAE increased from 25.39 ± 3.59 to 40.73 ± 7.65 for Uni-m1, 24.45 ± 3.67 to 51.51 ± 9.29 for Uni-m2, 25.56 ± 6.92 to 39.72 ± 9.12 for Uni-m3, and from 23.03 ± 3.18 to 36.53 ± 6.5 for Uni-z1, 24.87 ± 4.64 to 37.59 ± 7.35 for Uni-z2, 26.84 ± 6.17 to 33.63 ± 6.41 for Uni-z3, respectively, indicating the model trained with single-institution MRI data failed to generalize to external MRI datasets.

2) Generalizability of tri-institution models

The model generalizability improved when training the model with more diverse MRI data for both Min-Max and Z-Score normalization methods. As shown in **Table 4**, the overall external performance obtained 7.34% improvement for Tri-M model and 9.66% improvement for Tri-Z model in MAE and 1.57% improvement for Tri-M model and 2.36% improvement for Tri-Z model in PSNR.

TABLE IV. EXTERNAL PERFORMANCE IMPROVEMENT OF TRI-INSTITUTION MODELS

Model	MAE	PSNR
Tri-M	7.34%	1.57%
Tri-Z	9.66%	2.36%

TABLE V. EXTERNAL PERFORMANCE DROP OF UNI-INSTITUTION MODELS

Model	Min-Max		Z-Score		
	MAE	PSNR	MAE	PSNR	
Uni-m1	60.42%	12.32%	Uni-z1	58.62%	10.70%
Uni-m2	110.67%	13.89%	Uni-z1	51.15%	6.75%
Uni-m3	34.37%	5.18%	Uni-z1	25.30%	3.00%
Overall	68.49%	10.46%	Overall	44.42%	6.82%

3) Influence of normalization methods to model generalizability

The quantitative results from **Table 4** and **Table 5** indicate that Z-Score normalization outperformed the Min-Max

normalization on external datasets, with less performance drop for uni-institution models (44.42% v.s. 68.49% for MAE and 6.82% v.s. 10.46% for PSNR, respectively). From **Table 4**, Z-Score normalization-based Tri-Z model obtained better external performance than Min-Max normalization-based Tri-M model in both MAE (9.66% v.s. 7.34%) and PSNR (2.36% v.s. 1.57%), which suggest that Z-Score normalization outperforms Min-Max normalization in model generalizability improvement.

B. Qualitative results

To visually evaluate the external generalization performance of uni-institution and tri-institution models with different normalization methods, the external results of different models are illustrated in **Fig. 3**. The generalizability of uni-institution models varies greatly. For the uni-institution models for both Min-Max and Z-Score normalization, all uni-institution models showed worse generalizability to external MRI data with various tumor-to-normal tissue contrast and different degrees of contrast-enhancement failure (indicated with red arrows), especially the model trained with Institution-2 data (i.e., Uni-m2 and Uni-z2). The model trained with Institution-1 data (i.e., Uni-m1 and Uni-z1) showed overall image contrast difference compared with ground truth CE-MRI while the models trained with Institution-3 data showed tumor (Uni-m3) and normal vessel (Uni-z3) contrast enhancement failure.

Both the two tri-institution models achieved promising generalizability to external data. The generated GFCE-MRI from both Tri-M and Tri-Z models achieved a better visual approximation of tumor contrast enhancement. Compared with the Tri-M model, the Tri-Z model obtained a better approximation of tumor surrounding structures (indicated with yellow arrows).

IV. DISCUSSION

In radiotherapy, CE-MRI is commonly used for accurate tumor delineation, especially for the highly infiltrative NPC [15]. However, GBAs-associated safety issues have stimulated the medical community to eliminate the use of GBAs. Recently, a worldwide interest has been promoted to synthesize the GFCE-MRI for providing a gadolinium-free alternative for precision tumor delineation [7-15]. Nevertheless, the model generalizability on external institution data remains unexplored and there is no standard multi-institutional MRI normalization method has been established. Herein, for the first time, we investigated the model generalizability using different data normalizations for GFCE-MRI synthesis in NPC patients using MRI data retrieved from seven institutions. In this discussion, we attempted to summarize key findings, discuss the potential underlying mechanisms, and provide the research community with our perspectives in future directions.

The uni-institution models suffered from various degrees of degradation on external MRI datasets. As shown in **Table 2** and **Table 3**, the quantitative results of uni-institution models show that the uni-institution models performed well on internal testing datasets with lower MAE and higher PSNR but failed to

generalize to external unseen data. The visual comparisons (**Fig. 3**) of synthetic GFCE-MRI among different models also showed that uni-institution models failed to predict the correct contrast enhancement, both in tumor and surrounding normal tissues. These results suggest that there exist significant MRI data bias across institutions, resulting in a phenomenon that performance of well-trained in-house models cannot generalize to external MRI datasets.

By involving diverse MRI data from multiple institutions, the Tri-Z model achieved improved external performance than uni-institution models, as shown in **Table 4**. This result indicates that involving diverse MRI data from multiple institutions is more capable of achieving a better external performance, possibly due to the view of the model has been enlarged. From **Table 2** and **Table 3**, the Tri-M and Tri-Z that trained with same number of training samples as uni-institution models did not obtain obvious performance degradation in the three intra-institution datasets, indicating that involving diverse MRI data from multiple institutions for model development is capable of maintain the intra-institution accuracy, though the two tri-institution models were trained with 1/3 number of samples from each institution.

Z-Score normalization outperformed Min-Max normalization in improving the model generalizability, for both uni-institution models and the tri-institution model. As shown in **Table 4** and **Table 5**, Z-Score normalization achieved 24.07% and 3.64% less drop of MAE and PSNR respectively than Min-Max normalization for uni-institution models. Z-Score normalization also obtained additional 2.32% and 0.79% performance gain in MAE and PSNR for the tri-institution model from **Table 4**. This is possibly due to Z-Score normalizes all the patients' mean and standard deviation to the same value (0 and 1, respectively), which minimize the distribution variations among all training patients and external testing patients, while Mix-Max normalization preserves the relationship (i.e., the intensity ratio) among the original data values. Moreover, Min-Max normalization does not help in results interpretation in the multi-institutional setting as the data distribution among different institutions is not identical. As demonstrated in [21], the model trained with smaller mean intensity dataset would obtain significantly better quantitative results, even with same number of training samples. Herein, the Z-Score normalization is more appropriate in multi-institutional setting as it normalize the mean intensities of multi-institutional datasets to the same value.

Our study has several limitations. Firstly, since our findings are based on MMgSN-Net [15], applicability of our results using other deep-learning models deserves future investigation. Secondly, this work takes into account the diversity of MRI images and signal intensities of MRI between institutions, other MRI characteristic, such as image texture, artifacts, and tumor size should also be considered to further improve the model generalizability.

V. CONCLUSION

In this study, we investigated the model generalizability for GFCE-MRI synthesis in NPC using data from seven

institutions and explored potential methods to improve the model generalizability. Results of the present work showed that the tri-institution models developed from multi-institutional MRI generally resulted in higher generalizability than the uni-institution models developed from single-institution datasets. Application of the Z-Score normalization was capable of improving the model generalizability in multi-institutional MRI setting, which outperformed Min-Max normalization.

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