

Synthetic Data for Robust Stroke Segmentation

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Abstract

Current deep learning-based approaches to lesion segmentation in neuroimaging often depend on high-resolution images and extensive annotated data, limiting clinical applicability. This paper introduces a novel synthetic data framework tailored for stroke lesion segmentation, expanding the SynthSeg methodology to incorporate lesion-specific augmentations that simulate diverse pathological features. Using a modified nnUNet architecture, our approach trains models with label maps from healthy and stroke datasets, facilitating segmentation across both normal and pathological tissue without reliance on specific sequence-based training. Our method achieves robust out-of-domain performance where conventional approaches fail, with in-domain performance of 48.2% Dice compared to 57.5% for conventional training. Crucially, even with oracle knowledge of the optimal domain adaptation method - an unrealistic scenario in practice - conventionally-trained models cannot match our synthetic approach in out-of-domain settings. The framework demonstrates that synthetic pre-training provides fundamental robustness unachievable through test-time adaptation alone. Our approach reduces reliance on domain-specific training data and helps bridge the gap between research-grade and clinical scans to improve clinical stroke neuroimaging workflows. PyTorch training code and weights are publicly available at <https://github.com/liamchalcroft/SynthStroke>, along with an SPM toolbox featuring a plug-and-play model at <https://github.com/liamchalcroft/SynthStrokeSPM>.

Keywords

Machine Learning, Image Segmentation, Domain Adaptation

Article informations

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1. Introduction

Semantic segmentation is a critical component of neuroimaging pipelines, enabling precise quantification of anatomical structures and lesions for applications like tracking disease progression and planning treatments. In research settings, segmentation labels are typically derived from high-quality, standardised structural scans (e.g., MPAGE) that benefit from consistent field-of-view, spacing, orientation, and minimal artifacts. In contrast, clinical scans often exhibit significant variability in these factors, which can severely impact deep learning model performance. Consequently, models trained on homogeneous research-grade data may not generalise well to the diverse, lower-quality images encountered in clinical practice.

Both traditional probabilistic methods (Ashburner and Friston, 2005) and modern deep discriminative methods (Isensee et al., 2020) require prior information to be pro-

vided for a given sequence - in the case of a traditional method this may be an atlas or template, and in modern methods this would come in the form of training data. Atlas-based methods build a template of the anatomical structure of the brain, which may be deformed into alignment with a new subject to assign voxel-wise anatomical classes. This is proven to be robust for delineating healthy structure even with shifts in contrast (Puonti et al., 2016), however it is non-trivial to include classes of pathology such as stroke within such a model, due to the inherent heterogeneity in location and geometric properties. In the context of generative models, lesions may be included in the form of anomaly detection as demonstrated in Seghier et al. (2008). This method is not however directly attempting to label the pathology, and (by design) will label physiological changes such as ventricular enlargement in addition to the responsible infarct.

Deep discriminative models trained using supervised

learning have been able to reach human-level performance when tested in-domain on large datasets for a variety of brain pathologies and imaging modalities (Baid et al., 2021; de la Rosa et al., 2024). There is still however a significant gap when trying to translate these models to clinical data, where each hospital is likely to vary both in the scanning equipment used and the choice of imaging sequences (Nguyen et al., 2024). This poses a significant challenge to the adoption of deep learning for automating the labelling of clinical data, which could greatly help to accelerate the translation of modern research in stroke prognosis (Loughnan et al., 2019).

To extend such methods to the open-ended domain of clinical scans, models often need to perform on sequences for which no training data may be available. To this end, domain randomisation via synthetic data has been shown to give impressive results for healthy brain parcellation in SynthSeg (Billot et al., 2023). In this method, a set of ground truth healthy tissues are used to generate synthetic images, under the assumption that each tissue class' intensity distribution should roughly follow a Gaussian. By assigning random Gaussian distributions to each class, a deep learning model can learn to extract shape information for parcellation in a way that is invariant to the input image's relative tissue contrast, hence allowing the model to be used on any sequence at test-time, without training data or prior knowledge of the sequence. This method of training with synthetic data has since been extended to tasks such as image registration (Hoffmann et al., 2022, 2023, 2024), image super-resolution (Iglesias et al., 2021, 2023), surface estimation (Gopinath et al., 2023, 2024a) and vascular segmentation (Chollet et al., 2024). A comprehensive overview is available in Gopinath et al. (2024b).

An additional benefit of this method is that the 'forward model' of creating an MRI (or CT) image from tissues of different physical properties is a perfect 1:1 mapping to the 'inverse model' of labeling the tissues (i.e. segmentation) from the acquired image. In structures that exhibit a large amount of inter-rater variability, this is likely to help prevent a model from imitating under- or over- segmentation from imperfect ground truths - the images segmented are generated from the corresponding segmentation labels and so labels will always be a consistent method of segmentation.

Prior work to SynthSeg demonstrated the potential of encoding anatomical priors in a neural network through pre-training with unpaired parcellation labels (Dalca et al., 2018). Such methods face significantly larger challenges when applied to the heterogeneous shape and spatial distribution of lesions. In healthy parcellation, anatomical structures have consistent positions across individuals (e.g., the brainstem reliably appears in the same region of the brain), a regularity that has motivated atlas-based approaches to

parcellation. In contrast, lesions are highly variable across individuals - not only in number and size but also in their spatial distribution. Unlike anatomical structures, lesions cannot be reliably mapped to a specific location or shape within an atlas. Although the exact site of lesion initiation is often influenced by the brain's vascular architecture - meaning that certain regions are statistically more susceptible to stroke due to the location of large blood vessels - the resulting lesion's size, shape, and spread are highly variable. Multiple sclerosis serves as an exception, with lesions that are somewhat predictable in their white matter localisation (Lassmann, 2018), enabling modelling through synthetic deep learning frameworks (Billot et al., 2021; Laso et al., 2024) and traditional probabilistic models (Cerri et al., 2021). More recently, Liu et al. (2024) has shown promising results by training a SynthSeg-like model on lesion labels from various pathologies, providing a foundation for fine-tuning on multiple downstream datasets.

Robust open-domain stroke segmentation remains an unsolved challenge. Most domain-specific frameworks for stroke lesion segmentation are targeted towards lesion-pasting (Zhang et al., 2021; Dai et al., 2022; Basaran et al., 2023), aiming to augment the anatomical variety without making any attempt to augment the variety in image contrasts. Likewise, label-conditioned generative models such as TumorGAN (Li et al., 2020) can similarly only generate new lesioned brains within the learned distribution of image intensities. None of these works approach the task of robustness to shifts in image appearance, instead focusing on shape-related augmentation.

In our work, we extend the SynthSeg framework to the task of stroke lesion segmentation via a novel lesion-pasting method that better simulates variety in lesion appearance. Our hybrid approach trades a statistically significant 9.3% median Dice reduction in-domain (57.5% vs 48.2%, $p < 0.001$) for improved out-of-domain robustness. Crucially, we demonstrate that even with oracle knowledge of the optimal domain adaptation method, conventional training cannot match our synthetic approach in out-of-domain scenarios. We validate this on a comprehensive range of lesion datasets with a wide distribution of image characteristics and lesion physiology. To assist in widespread evaluation of this framework, we release PyTorch training code/weights, and a MATLAB toolbox for SPM to reduce the barrier to clinical adoption.

2. Methods

Terminology and Notation

For clarity, we define the key terms used throughout this work:

- **TTA (Test-Time Augmentation):** A procedure at in-

ference where multiple augmented versions (here generated by flips) of an input are processed and their predictions averaged to improve robustness.

- **DA (Domain Adaptation):** Techniques applied at test time to adapt a trained model to new data distributions.
- **Oracle DA:** The hypothetical best-case scenario where the optimal DA method is known a priori for each dataset/modality combination.

2.1 Synthetic Data Generation Framework

Rationale. Our goal is to create a large, diverse and perfectly labelled training set without the labour of voxel-wise annotation. We therefore generate paired *image-label* volumes by composing healthy tissue maps with realistically shaped stroke lesions, followed by intensity synthesis and heavy image-quality augmentation (Fig. 2).

(i) Healthy-tissue label bank. Instead of the 100+ FreeSurfer classes used by SynthSeg, we adopt the nine posterior tissue maps produced by MultiBrain (Brudfors et al., 2020). This reduces memory usage, speeds up sampling and still retains the GM/WM/CSF boundaries that matter for lesion realism (Fig. 1).

(ii) Lesion Copy-Paste (Lesion-CP). We extend Soft-CP (Dai et al., 2022) with random dilate/erode 'feathering' and a spatially varying bias-field multiplier (MONAI Random Bias Field (Cardoso et al., 2022)) to mimic penumbral intensity fall-off (Middleton et al., 2024).

(iii) Intensity sampling. Each tissue class is assigned $\mu \sim \mathcal{U}(0, 255)$, $\sigma \sim \mathcal{U}(0, 16)$ and Gaussian blur FWHM $\sim \mathcal{U}(0, 2)$. For stroke lesions we modulate the copy-pasted mask with the bias field to create intra-lesion heterogeneity. We sample from a single Gaussian distribution, with the implications of this choice examined via Hartigan's dip-test in Appendix A.

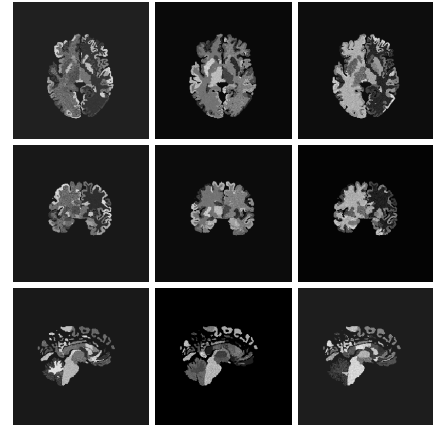
(iv) Image-quality augmentations. Table 1 summarises every random transform (bias field, affine, elastic, skull-strip imperfection, noise, resolution, motion, contrast, etc.).

Future work will explore mixture-of-Gaussians modelling (Ashburner and Friston, 2005) to represent lesions that are simultaneously hyper- and hypo-intense.

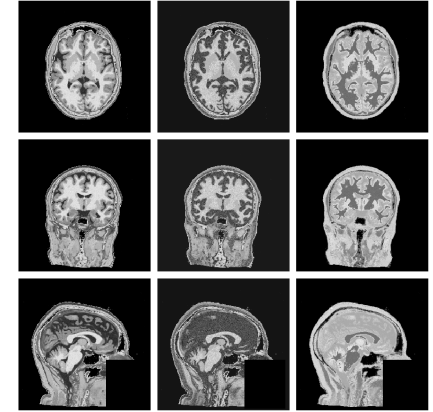
With this synthetic data generation pipeline established, we now describe the datasets, network architecture, and training procedures used in our experiments.

2.2 Training Data and Sampling

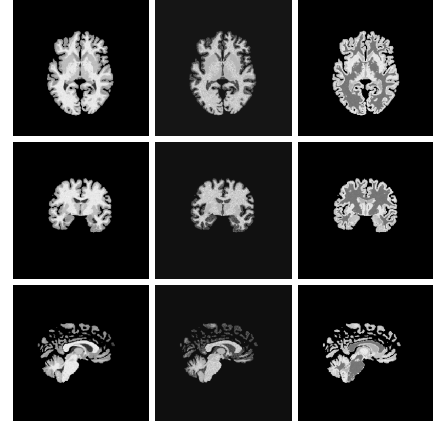
Healthy maps: OASIS-3 ($N=2\,679, 2\,579/100$ train/val) with MultiBrain segmentation, all warped to ICBM space. Lesion masks: ATLAS ($N=655, 419/105/131$ train/val/test)



(a) FreeSurfer (Puonti et al., 2016)



(b) MultiBrain (Brudfors et al., 2020)



(c) MultiBrain (skull-stripped)

Figure 1: Sample generated images using different labels for a single subject. 1a: FreeSurfer anatomical labels. 1b: MultiBrain tissue labels. 1c: MultiBrain tissue labels masked to simulate skull-stripping.

aligned to the same space. During sampling we paste one random ATLAS lesion onto one random OASIS subject and on-the-fly augment as above.

Table 1: Parameter ranges for every image-quality augmentation used during synthetic training. [†] indicates augmentations applied only to synthetic data and not to real ATLAS samples.

Category	Transform	Sampling range / notes
Bias field	Multiplicative bias	Control points $\mathcal{U}(2, 7)$, strength $\mathcal{U}(0, 0.5)$
Affine	Rotation	$\mathcal{U}(-15^\circ, 15^\circ)$ on each axis
	Shear	$\mathcal{U}(0, 0.012)$
	Zoom	$\mathcal{U}(0.85, 1.15)$
	Elastic	Control points $\mathcal{U}(0, 10)$, max disp. $\mathcal{U}(0, 0.05)$
Skull-strip flaw [†]	Dilation	$p = 0.3$, radius = 2 vox.
	Erosion	$p = 0.3$, radius = 4 vox.
	Noise	Gaussian (SNR) SNR $\mathcal{U}(0, 10)$, smoothed by g -factor $\mathcal{U}(2, 5)$
Resolution [†]	Slice anisotropy	Thk. factor $\mathcal{U}(1, 8)$ (base res. 1 mm ³)
Contrast	Gamma	$\gamma = 10^{\mathcal{N}(0, 0.6)}$
Motion blur [†]	PSF width (FWHM)	$\mathcal{U}(0, 3)$ vox.
Flip	Mirror on each axis	$p = 0.8$

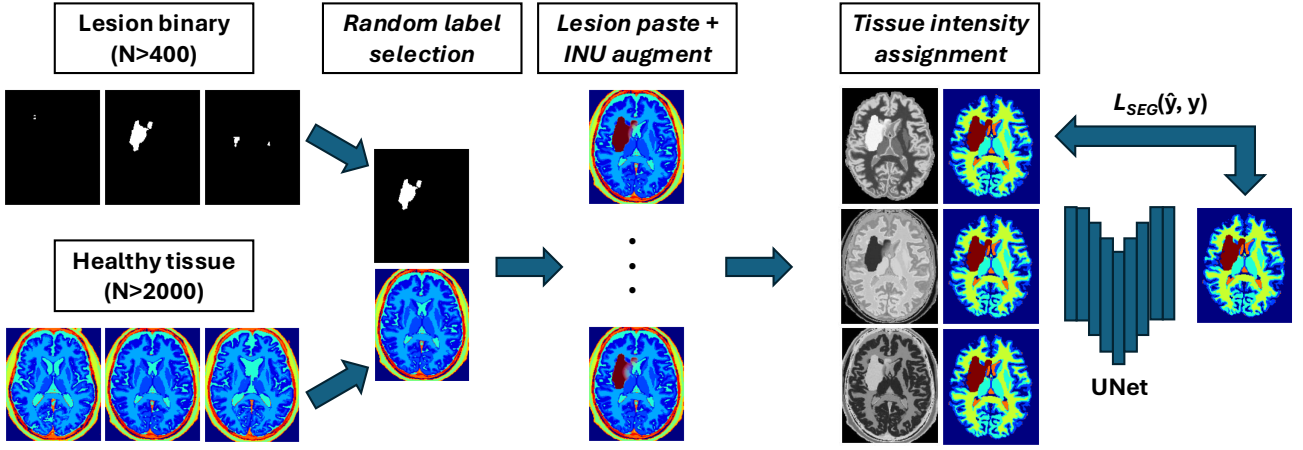


Figure 2: Schematic overview of the data generation process. Lesions are sampled from a template-normalised bank of lesion binary masks, and healthy tissue maps are sampled from a template-normalised bank of MultiBrain segmentations. Pasting of lesions onto healthy tissue maps is performed using a spatially varying lesion intensity to simulate penumbra. Tissue intensities may then be sampled from Gaussian distributions and image-label pairs used to train a segmentation model in a supervised manner.

2.3 Network Architecture and Optimisation

Backbone. 3D U-Net (nnUNet template) with six levels ($16 \rightarrow 32 \rightarrow 64 \rightarrow 128 \rightarrow 320$ channels), PReLU activations and one residual unit per block (Isensee et al., 2024).

Output channels. Background + Gray Matter (GM) + White Matter (WM) + GM/WM Partial Volume + Cerebrospinal Fluid (CSF) + Lesion (total six channels) for **Synth**; Background + Lesion for **Baseline**. When real images (binary GT) enter the mixed loader we mask the loss to the lesion channel only.

Training schedule. 192³ crops, batch 1, 1 200 epochs \times 500 iterations ($=6 \times 10^5$ updates), AdamW ($\eta_0 = 10^{-4}$, weight-decay 0.01, poly LR decay with power 0.9), combined Dice + CE loss, dropout 0.2, gradient-norm clip 12.

Data-loader mixing. Synthetic : Real ratio = 2 579 : 419

(mirrors sample counts). Real MPRAGE images receive the same spatial/intensity transforms as synthetic batches.

Comparison model. WMH-SynthSeg (Laso et al., 2024; Fischl, 2012) is included as an "off-the-shelf" robust-lesion baseline, but its training labels target small, periventricular WMHs and differ substantially from large-cortical stroke lesions.

2.4 Domain Adaptation Methods

At test time we evaluate a diverse set of **unsupervised domain-adaptation (DA)** techniques to determine: (i) whether the **Baseline** model can, under an *oracle* choice of DA method, match the performance of the DA-free **Synth** model, and (ii) whether Synth's inherent robustness offers a better starting point for DA on truly unseen data.

We tested six DA methods (TTA, DAE, TENT, PL, UPL, DPL) and report the best-performing configuration for each modality/dataset combination as **Oracle DA** in Tables 2-5. This represents an upper bound on baseline performance, as in practice one cannot know *a priori* which DA method will work best. Full individual results for all methods appear in Appendix B, Tables 10-13.

DA techniques evaluated.

(1) TTA - Test-Time Augmentation (Wang et al. (2019); denoted **TTA**). Eight mirror-flipped volumes (2^3 axis-flip combinations) are inferred, logits averaged, and softmax/argmax yields the mask. This heuristic is cheap and rarely degrades performance.

(2) DAE - Denoising Auto-Encoder Regularisation (Karar et al., 2021). We train a 3-layer denoising auto-encoder to regularise noisy labels for both Baseline and Synth. At inference a three-layer normalisation network ($3 \times 3 \times 3$ kernels, 16 channels) is *prepended* to the frozen segmentor; its activation is

$$f(x) = \exp - \frac{x^2}{\sigma^2}, \quad (1)$$

where σ is learned. For each test image the network is re-initialised and optimised for 100 steps to minimise Dice + L2 loss between segmentation logits and their DAE-cleaned counterpart.

(3) TENT - Test-Time Entropy Minimisation (Wang et al., 2021). Because 3D memory limits rule out batch-norm adaptation, we instead optimise an identical normalisation network (initialised from scratch per subject) for 100 steps to minimise Shannon entropy

$$\mathcal{H}(\hat{y}) = - \sum_c p_c(\hat{y}) \log p_c(\hat{y}), \quad (2)$$

with p_c the class- c softmax probability.

(4) PL Family - Pseudo-Labeling (Self-Training) (Chen et al., 2021). Three variants are tested:

- **PL**: threshold softmax at $\tau = \frac{1.5}{N_C}$ to keep only high-confidence voxels, where N_C is the total number of output classes.
- **UPL**: PL plus uncertainty masking via 10-sample Monte-Carlo dropout (variance > 0.05 discarded).
- **DPL**: full "prototype-consistency" pipeline that further removes voxels inconsistent with decoder-feature prototypes.

All PL variants fine-tune *all* segmentation weights for 2000 iterations with weighted cross-entropy.

Common optimiser settings. Every trainable DA method (DAE, TENT, PL, UPL, DPL) uses AdamW (Loshchilov and Hutter, 2019) with learning-rate 0.002 and weight-decay 0.01. TTA is inference-only and therefore parameter-free.

2.5 Large-Scale Pseudo-Labeling

Because the synthetic pipeline decouples *image realism* from *label accuracy*, we can tolerate imperfect pseudo-labels. We therefore use Baseline + TTA to annotate 1159 chronic stroke MPRAGE scans from PLORAS Sample 1 (Seghier et al., 2016); this **PLORAS-MPRAGE** cohort feeds the mixed loader exactly like ATLAS.

3. Experiments

3.1 Datasets

Models were validated on four stroke-lesion datasets. We assessed the models' in-domain performance on the hold-out test set for the **ATLAS** dataset (131 subjects, 1 mm isotropic MPRAGE). Out-of-domain (OOD) robustness was evaluated on three additional cohorts: the **ISLES 2015** dataset (Maier et al., 2017) ($N=28$ subjects with skull-stripped T1w/T2w/FLAIR/DWI), the **ARC** dataset (Gibson et al., 2024; Johnson et al., 2024) ($N=229$ T2w, 202 T1w, 85 FLAIR; $N=84$ subjects have all three), and the hospital scans from 661 acute-stroke patients in PLORAS Sample 2 ($N=106$ T2w, 300 FLAIR, 255 CT), collectively referred to as **PLORAS** (Price et al., 2010). ISLES and PLORAS also introduce an acute-versus-chronic shift. There is no overlap in patients between the PLORAS hospital scans (acute) and the PLORAS MPRAGE cohort described in Section 2.5.

For PLORAS, images are resampled from the original 2 mm isotropic resolution to 1 mm to maintain a single preprocessing pipeline, acknowledging that this constitutes a second resampling step.

3.2 Experimental Design

Pre-processing and inference. All test images are re-oriented to RAS, resliced to 1 mm^3 voxels, histogram-normalised and z -scored. Inference uses a 192^3 sliding window with 50 % overlap and a Gaussian blending kernel ($\sigma = 0.125$). Test-time augmentation (**TTA**) averages logits over all eight combinations of left-right, anterior-posterior and inferior-superior flips.

Multi-modal ensembles. When multiple MR sequences are available for the same subject (ISLES 2015, ARC) we average the per-modality logits before the softmax. This simple ensembling mimics a realistic clinical deployment.

Pseudo-label training. Pseudo-labels are generated for the PLORAS MPRAGE cohort with the Baseline + TTA model. A new Baseline and a new Synth model are then re-trained using the union of ATLAS and pseudo-labelled data, following exactly the same optimisation schedule as the originals.

3.3 Evaluation Metrics

Prior to metric computation, predictions and ground truth are resliced to 1 mm and zero-padded to 256^3 voxels. We report Dice and Surface-Dice (Seidlitz et al., 2022) (1 mm tolerance) in the main text; HD95, absolute volume difference (AVD), absolute lesion difference (ALD), lesion-wise F1, true-positive rate (TPR) and false-positive rate (FPR) appear in Appendix B.

Dice quantifies volumetric overlap (1 = perfect, 0 = none), whereas HD95 captures boundary error while being robust to outliers. AVD reports absolute volume mismatch in cm^3 . ALD counts mismatches in the number of connected components, and lesion-wise F1 scores per-lesion detection accuracy. TPR and FPR follow standard definitions.

3.4 Comparison Methods

We evaluate four primary approaches:

(i) Baseline: A standard 3D U-Net trained solely on real MPRAGE images from the ATLAS dataset ($N=419$) using supervised learning. This represents the conventional approach of training on a single modality with manual annotations.

(ii) Oracle DA: The baseline model enhanced with the single best-performing domain adaptation method (selected post-hoc from TTA, TENT, DAE, PL, UPL, DPL) for each modality. This represents the theoretical upper bound of what domain adaptation can achieve when the optimal method is known - an unrealistic scenario in practice.

(iii) WMH-SynthSeg: The pre-trained white matter hyperintensity model from Laso et al. (2024); Fischl (2012), included as an off-the-shelf robust lesion segmentation baseline. Note that this model was trained for small periventricular WMH lesions, which differ substantially from large cortical stroke lesions.

(iv) Synth (Ours): Our proposed approach using synthetic data generation as described in Section 2. The model is trained on a mixture of synthetic data (generated from OASIS healthy maps + ATLAS lesion masks) and real ATLAS MPRAGE images, following the framework illustrated in Figure 2.

For fair comparison, TTA is *only* applied when a model is explicitly labelled "+TTA" or when it is selected as the Oracle DA. All candidate maps are binarised via argmax on posterior probabilities without additional calibration or threshold tuning.

4. Results

Overall results are shown in Figure 3, which compares (i) the Baseline, (ii) the Oracle DA (best-performing do-

main adaptation for Baseline only), (iii) WMH-SynthSeg, and (iv) the DA-free Synth model. Oracle DA represents the hypothetical best-case scenario where the optimal DA method is known *a priori* for each dataset/modality. Tables 2, 3, 4 and 5 provide comprehensive results for all tested domain adaptation methods on both Baseline and Synth models.

4.1 ATLAS

The ATLAS dataset represents the in-domain scenario with T1-weighted images matching our training distribution. Table 2 shows performance on the held-out test set. The conventional **Baseline** achieves a median Dice of **57.5 %**, whereas our **Synth** model reaches **48.2 %**, a 9.3 % gap that represents the 'price of robustness' we accept for the larger out-of-domain gains reported later. Surface Dice follows the same trend (49.4 % vs. 38.1 %).

Applying test-time augmentation (Baseline+TTA) changes Dice by <0.1 %, indicating that TTA adds little benefit when the evaluation domain is already aligned with training.

The off-the-shelf **WMH-SynthSeg**, trained for small periventricular WMH lesions, scores only 7.3 % Dice, confirming that cortical stroke in ATLAS lies well outside its intended scope.

A paired Wilcoxon test (Appendix Figure 7) verifies that the Baseline–Synth Dice difference is statistically significant, underscoring that domain-invariant training still sacrifices some in-domain accuracy. Voxel-level metrics in Appendix Table 10 reveal the Baseline achieves a higher recall but at the cost of more false positives, suggesting a tendency to over-segment.

Worst Scanner Syndrome. This in-domain drop is consistent with the 'Worst Scanner Syndrome' hypothesis (Moyer and Golland, 2021), which posits that many domain-invariance strategies pull feature quality toward the least informative - or noisiest - domain. As the next Results sections demonstrate, that modest in-domain penalty is offset by substantial gains on heterogeneous clinical data.

Table 2: Median results on the ATLAS hold-out set ($N=131$). Best score shown in bold. Student's t distribution 95% confidence intervals given in brackets.

Modality	Model	Dice (%)	Surface Dice (%)
T1w	Baseline	57.5 (52.3-62.7)	49.4 (44.5-54.3)
	Baseline+TTA	57.5 (52.2-62.8)	49.5 (44.5-54.5)
	WMH-SynthSeg	7.3 (4.8-9.7)	8.9 (6.9-10.9)
	Synth (Ours)	48.2 (43.1-53.4)	38.1 (33.4-42.7)

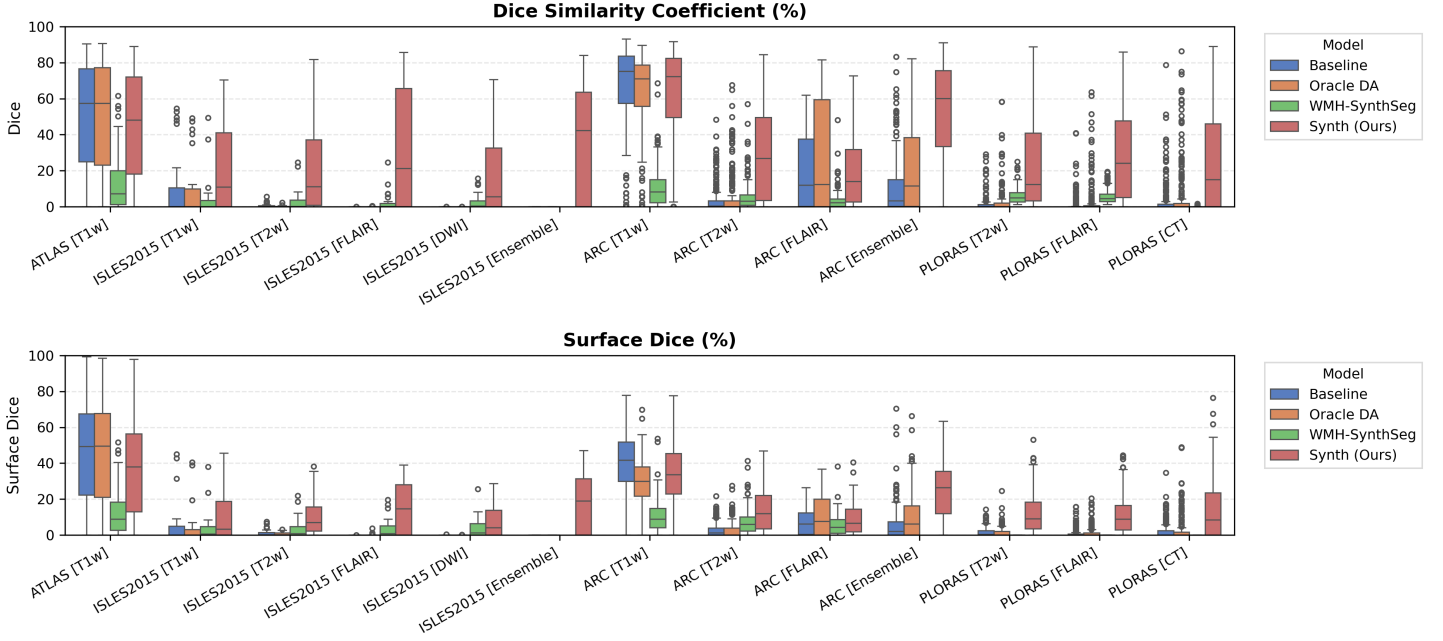


Figure 3: Dice and Surface Dice metrics for all reported datasets. 'Oracle DA' represents the hypothetical best-case scenario where optimal DA method is known a priori for each dataset/modality and applied to the baseline model.

4.2 ARC

The ARC dataset contains research-quality chronic stroke scans, representing a moderate domain shift from our ATLAS training data. We expect T1w performance to be strongest given its proximity to the training domain.

Table 3 shows performance on ARC, comparing Baseline, Oracle DA (best possible domain adaptation for Baseline), WMH-SynthSeg, and our Synth approach. For T1w, both baseline and Oracle DA achieve 75.2% Dice, with Synth at 72.3% - all maintaining strong performance (Appendix Figure 8 shows statistical significance). However, T2w reveals dramatic differences: baseline drops to 0.4% and Oracle DA to 0.1%, while Synth maintains 26.8%. FLAIR shows intermediate performance with Oracle DA at 12.4% versus Synth at 14.1%. The ensemble results are particularly striking: Oracle DA achieves only 11.7% while Synth reaches 60.2%.

This pattern - strong baseline performance on T1w but catastrophic failure on other modalities even with optimal DA - demonstrates that synthetic pre-training provides fundamental robustness unachievable through post-hoc adaptation.

4.3 ISLES 2015

ISLES 2015 contains skull-stripped sub-acute stroke scans, representing our most challenging domain shift with both acquisition and pathology differences from the chronic ATLAS training data.

The results in Figure 3 and Table 4 show that the Synth model outperforms the Baseline model in all modalities in

Table 3: Median results on the ARC dataset ($N=229$). Best score shown in bold. Student's t distribution 95% confidence intervals given in brackets. 'Oracle DA' represents the hypothetical best-case scenario where optimal DA method is known a priori for each dataset/modality and applied to the baseline model.

Modality	Model	Dice (%)	Surface Dice (%)
T1w	Baseline	75.2 (71.5-79.0)	41.7 (39.1-44.4)
	Oracle DA	75.2 (71.3-79.0)	42.1 (39.4-44.8)
	WMH-SynthSeg	8.3 (6.8-9.8)	8.9 (7.7-10.1)
	Synth (Ours)	72.3 (68.4-76.2)	33.7 (31.2-36.2)
T2w	Baseline	0.4 (0.0-1.4)	1.2 (0.8-1.7)
	Oracle DA	0.1 (0.0-0.4)	1.0 (0.8-1.2)
	WMH-SynthSeg	3.2 (2.2-4.2)	6.0 (5.2-6.8)
	Synth (Ours)	26.8 (23.3-30.2)	12.0 (10.4-13.6)
FLAIR	Baseline	12.0 (7.6-16.3)	6.3 (4.7-7.8)
	Oracle DA	12.4 (5.8-19.0)	7.7 (5.2-10.2)
	WMH-SynthSeg	2.4 (0.9-3.9)	4.3 (3.0-5.5)
	Synth (Ours)	14.1 (9.7-18.5)	6.6 (4.8-8.4)
Ensemble	Baseline	3.4 (1.3-5.4)	2.0 (0.8-3.2)
	Oracle DA	11.7 (8.6-14.7)	6.3 (4.7-7.8)
	Synth (Ours)	60.2 (56.6-63.8)	26.3 (24.3-28.4)

regards to both Dice and Surface Dice with high statistical significance in the Dice metric evidenced in Appendix Figures 11-13. Baseline performance is near-zero across all modalities. Oracle DA shows minimal recovery: 10.5% for T1w (still below Synth's 11.0%) and 0.0% for all other modalities (T2w, FLAIR, DWI). In contrast, Synth achieves 11.0% (T1w), 11.1% (T2w), 21.2% (FLAIR), and 5.6% (DWI). The ensemble demonstrates the starkest contrast: 0.0% for both baseline and Oracle DA versus 42.3% for Synth.

The poor performance correlates with high false posi-

tive rates rather than missed detections (Appendix Table 12), suggesting models struggle with tissue discrimination in this domain. When baseline predictions fail catastrophically, no DA method can recover meaningful performance.

It is also evident from Table 4 that the model performance is highly dependent on the choice of image sequence available. The ensemble improves performance over individual sequences in ISLES2015 (Table 4) but shows mixed results in ARC, suggesting dataset-specific benefits rather than universal improvement. Although we only show the upper limit of an ensemble of all four sequences, it is expected that in cases where fewer sequences are available we will still observe constructive gains from post-hoc ensembling.

Table 4: Median results on the ISLES2015 dataset ($N=28$). Best score shown in bold. Student’s t distribution 95% confidence intervals given in brackets. ‘Oracle DA’ represents the hypothetical best-case scenario where optimal DA method is known a priori for each dataset/modality and applied to the baseline model.

Modality	Model	Dice (%)	Surface Dice (%)
T1w	Baseline	0.0 (0.0-7.4)	0.0 (0.0-4.8)
	Oracle DA	10.5 (0.0-21.7)	3.8 (0.0-9.9)
	WMH-SynthSeg	0.0 (0.0-4.5)	0.7 (0.0-3.9)
	Synth (Ours)	11.0 (1.2-20.8)	3.4 (0.0-8.5)
T2w	Baseline	0.0 (0.0-0.5)	0.3 (0.0-1.0)
	Oracle DA	0.0 (0.0-0.2)	0.0 (0.0-0.6)
	WMH-SynthSeg	0.1 (0.0-2.5)	0.8 (0.0-3.0)
	Synth (Ours)	11.1 (0.7-21.6)	7.1 (2.7-11.5)
FLAIR	Baseline	0.0 (0.0-0.0)	0.0 (0.0-0.0)
	Oracle DA	0.0 (0.0-0.0)	0.0 (0.0-0.0)
	WMH-SynthSeg	0.3 (0.0-2.3)	0.8 (0.0-3.0)
	Synth (Ours)	21.2 (8.5-34.0)	14.7 (9.0-20.3)
DWI	Baseline	0.0 (0.0-0.0)	0.0 (0.0-0.0)
	Oracle DA	0.0 (0.0-0.0)	0.0 (0.0-0.0)
	WMH-SynthSeg	0.4 (0.0-2.2)	1.3 (0.0-3.6)
	Synth (Ours)	5.6 (0.0-14.4)	4.1 (0.8-7.4)
Ensemble	Baseline	0.0 (0.0-0.0)	0.0 (0.0-0.0)
	Oracle DA	0.0 (0.0-0.0)	0.0 (0.0-0.0)
	Synth (Ours)	42.3 (30.2-54.5)	19.1 (12.7-25.4)

4.4 PLORAS

The PLORAS dataset represents the most extreme domain shift with real clinical data exhibiting large diversity in acquisition protocols and slice thickness. We expect minimal baseline performance given these challenging conditions. For all available modalities, the Synth model outperforms the baseline with statistical significance in Dice (see Appendix Figures 15 - 17).

Results in Table 5 demonstrate the most extreme performance gap between conventional training and our synthetic approach. On this challenging clinical dataset, Oracle DA achieves at most 0.2% Dice (CT modality), while our Synth model achieves 11.9%, 25.4%, and 11.3% for T2w, FLAIR, and CT respectively. The near-total failure of both baseline and Oracle DA on real clinical data

- where acquisition protocols, slice thickness, and image quality vary substantially - validates our core hypothesis: domain adaptation cannot substitute for domain-invariant training when deployment conditions diverge significantly from training data. Even WMH-SynthSeg, despite being trained for robustness, achieves only 0.0-4.9% Dice, likely due to its focus on small periventricular lesions rather than large cortical strokes. Full results for all individual DA methods are provided in Appendix Table 13. A number of samples are also visualised for this dataset in Figure 4.

Table 5: Median results on the PLORAS dataset ($N=661$). Best score shown in bold. Student’s t distribution 95% confidence intervals given in brackets. ‘Oracle DA’ represents the hypothetical best-case scenario where optimal DA method is known a priori for each dataset/modality and applied to the baseline model.

Modality	Model	Dice (%)	Surface Dice (%)
T2w	Baseline	0.0 (0.0-1.3)	0.1 (0.0-0.6)
	Oracle DA	0.1 (0.0-2.2)	0.1 (0.0-0.8)
	WMH-SynthSeg	4.9 (4.0-5.8)	0.0 (0.0-0.0)
	Synth (Ours)	11.9 (6.7-17.1)	8.4 (6.1-10.7)
FLAIR	Baseline	0.0 (0.0-0.4)	0.0 (0.0-0.2)
	Oracle DA	0.0 (0.0-0.4)	0.0 (0.0-0.2)
	WMH-SynthSeg	4.6 (4.2-5.0)	0.0 (0.0-0.0)
	Synth (Ours)	25.4 (22.5-28.3)	8.5 (7.4-9.6)
CT	Baseline	0.0 (0.0-1.1)	0.0 (0.0-0.5)
	Oracle DA	0.2 (0.0-0.5)	0.3 (0.0-0.7)
	WMH-SynthSeg	0.0 (0.0-0.0)	0.0 (0.0-0.0)
	Synth (Ours)	11.3 (8.0-14.6)	7.9 (6.0-9.8)

4.5 Domain Adaptation

Tables 3, 4 and 5 demonstrate that even with oracle selection of the optimal domain adaptation method, the baseline model cannot match Synth performance in out-of-domain settings. We evaluated six established DA techniques (TTA, TENT, DAE, PL, UPL, DPL) on both baseline and Synth models to determine whether domain adaptation could enable baseline generalisation. Complete results in Appendix Tables 11-13 consistently show that poor initial baseline predictions prevent effective adaptation. In contrast, when applied to our Synth model, several DA methods yield substantial improvements, demonstrating the potential for compound gains when robust pre-training is combined with appropriate post-hoc adaptation.

Analysis of complete DA results (Appendix Tables 11-13) reveals DAE as the most consistently effective method for Synth. DAE achieves substantial improvements: ARC T2w increases from 26.8% to 54.3% Dice, ISLES T1w from 11.0% to 31.1%, and PLORAS CT from 11.3% to 23.9%. TTA provides modest gains for PLORAS FLAIR (25.4% to 29.4%), while TENT benefits ARC FLAIR (14.1% to 36.1%). Pseudo-labeling methods (PL, UPL, DPL) show high variance - occasionally strong but often catastrophic.

The key finding: while DA cannot rescue baseline mod-

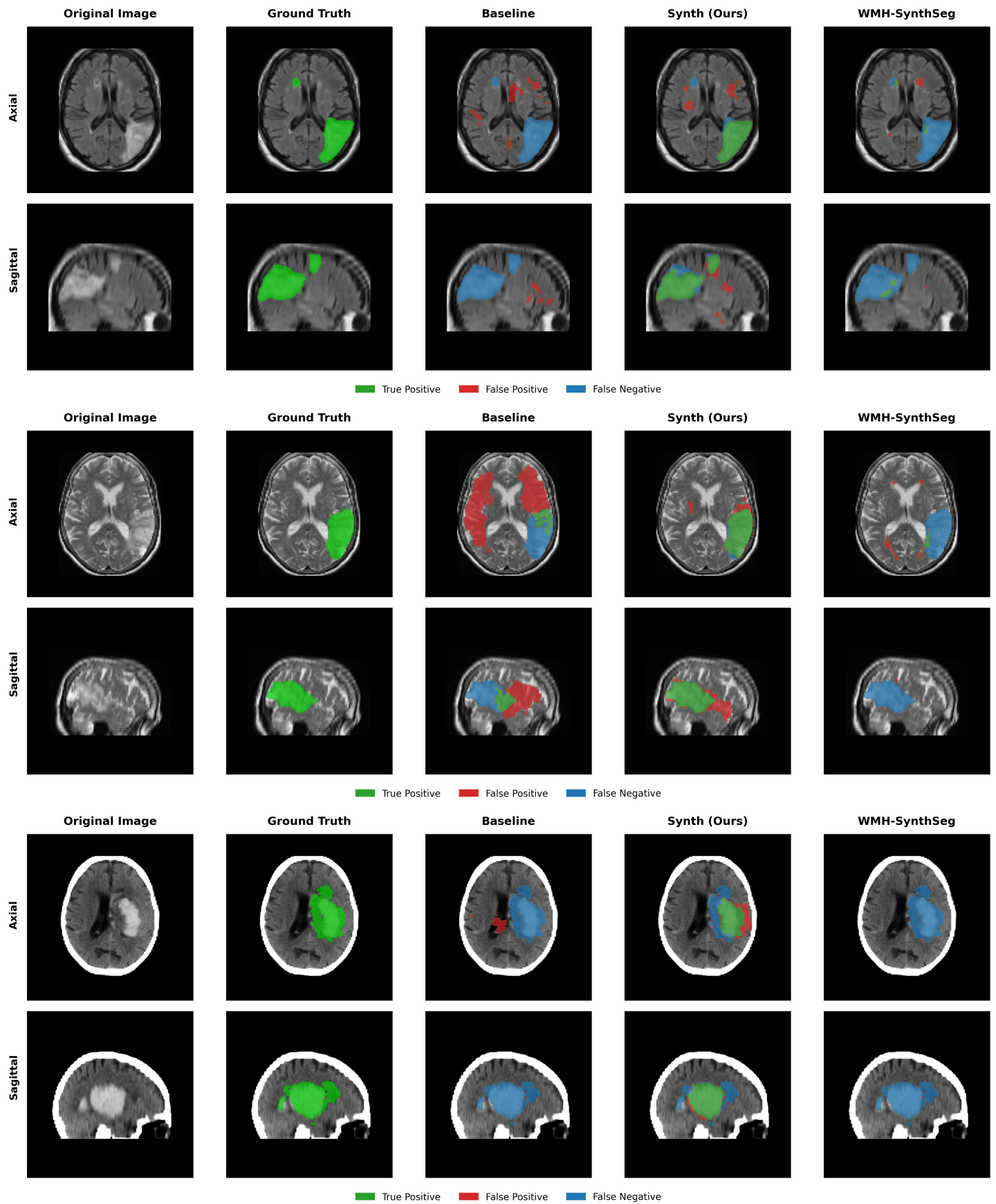


Figure 4: Sample visualisations in the PLORAS dataset. Green indicates a true positive prediction, red a false positive, and blue a false negative.

els trained on narrow data, it can enhance robustly pre-trained models. Synth+DAE consistently outperforms both Synth alone and Baseline+Oracle DA, demonstrating complementary benefits.

Our domain adaptation experiments serve primarily to establish an upper bound on baseline model performance. Even with oracle knowledge of the optimal DA method for each dataset/modality combination - an unrealistic scenario in practice - the baseline model cannot match Synth performance in out-of-domain settings. Full DA results for all methods appear in Appendix Tables 11-13. While individual DA methods show varied effectiveness, the key finding is that synthetic pre-training provides robustness that cannot be recovered through test-time adaptation alone. This underscores the value of appearance-invariant training, even with our current limitations in modelling stroke heterogeneity through single Gaussian distributions. Domain adaptation results varied substantially by method and modality, with no single approach providing consistent improvements. We present all tested combinations rather than cherry-picking optimal results, as real-world deployment would lack oracle knowledge of the best DA method for unseen data.

4.6 Ablation: To mix or not to mix?

For all experiments shown thus far, the Synth model used a mix of both synthetic data and the real ATLAS dataset. In order to evaluate whether this decision is justified, an ablation is performed where the Synth model using mixed real/synthetic data is compared to a model trained with only synthetic data. This model is trained in the exact same manner as described previously for the baseline and Synth models. Results for the test datasets ATLAS, ARC and ISLES 2015 are shown in Tables 6, 7 and 8 respectively.

The results reveal a nuanced picture. For in-domain and near-domain T1w data (ATLAS, ARC), mixing with real data provides clear benefits, with pure synthetic training achieving only 19.7% and 46.7% Dice respectively compared to 48.2% and 72.3% for mixed training. However, for several out-of-domain scenarios, pure synthetic training surprisingly often outperforms mixed training: ARC T2w (62.6% vs 26.8%), ISLES T1w (30.4% vs 11.0%), and ISLES FLAIR (37.2% vs 21.2%). This suggests that including real T1w data may inadvertently bias the model toward T1w-specific features, reducing generalisation to other modalities. The T2w and FLAIR improvements with pure synthetic training may reflect these sequences' different tissue contrast patterns, which are better captured by unbiased synthetic variation than by a model partially trained on T1w-specific features. Further improving the realism of the generated lesions with methods proposed in Liu et al. (2025) may aid in closing the gap for the model

trained only on synthetic data, potentially achieving both in-domain performance and out-of-domain generalisation.

Table 6: Median results for ablation and pseudo-label experiments on the ATLAS test set ($N=131$). Best score shown in bold. Student's t distribution 95% confidence intervals given in brackets.

Modality	Model	Dice (%)	Surface Dice (%)
T1w	Baseline	57.5 (52.3-62.7)	49.4 (44.5-54.3)
	Baseline+Pseudo	59.8 (54.8-64.7)	48.6 (44.0-53.3)
	Synth (no real data)	19.7 (15.5-23.9)	14.6 (12.3-16.9)
	Synth (Ours)	48.2 (43.1-53.4)	38.1 (33.4-42.7)
	Synth+Pseudo	49.6 (44.5-54.7)	38.2 (33.8-42.6)

Table 7: Median results for ablation and pseudo-label experiments on the ARC test set ($N=229$). Best score shown in bold. Student's t distribution 95% confidence intervals given in brackets.

Modality	Model	Dice (%)	Surface Dice (%)
T1w	Baseline	75.2 (71.5-79.0)	41.7 (39.1-44.4)
	Baseline+Pseudo	75.9 (72.1-79.7)	38.2 (35.8-40.7)
	Synth (no real data)	46.7 (43.0-50.4)	18.2 (16.5-20.0)
	Synth (Ours)	72.3 (68.4-76.2)	33.7 (31.2-36.2)
	Synth+Pseudo	74.3 (70.4-78.2)	36.8 (34.3-39.3)
T2w	Baseline	0.4 (0.0-1.4)	1.2 (0.8-1.7)
	Baseline+Pseudo	1.4 (0.3-2.4)	3.0 (2.5-3.4)
	Synth (no real data)	62.6 (58.8-66.5)	30.9 (28.7-33.0)
	Synth (Ours)	26.8 (23.3-30.2)	12.0 (10.4-13.6)
	Synth+Pseudo	37.2 (33.2-41.1)	14.7 (12.7-16.7)
FLAIR	Baseline	12.0 (7.6-16.3)	6.3 (4.7-7.8)
	Baseline+Pseudo	14.5 (10.1-18.9)	7.0 (5.5-8.5)
	Synth (no real data)	9.6 (7.0-12.3)	7.3 (6.1-8.6)
	Synth (Ours)	14.1 (9.7-18.5)	6.6 (4.8-8.4)
	Synth+Pseudo	12.9 (8.4-17.3)	6.3 (4.4-8.1)
Ensemble	Baseline	3.4 (1.3-5.4)	2.0 (0.8-3.2)
	Baseline+Pseudo	12.4 (9.7-15.0)	6.5 (5.1-8.0)
	Synth (no real data)	59.7 (56.2-63.1)	26.0 (24.2-27.9)
	Synth (Ours)	60.2 (56.6-63.8)	26.3 (24.3-28.4)
	Synth+Pseudo	69.4 (65.8-73.0)	34.0 (31.7-36.3)

4.7 Use of Pseudo-labelling

The use of synthetic data for semi-supervised pseudo-label training is also explored through the use of the PLORAS MPAGE dataset ($N=1159$), distinct from the PLORAS hospital scans used in the Experiments. This approach leverages a key advantage of our framework: because synthetic training decouples image generation from label accuracy, the model can tolerate and potentially benefit from imperfect pseudo-labels. Results in Figure 6 and Tables 6, 7 and 8 demonstrate an overall positive effect of pseudo labels in the Synth model, with the exception of FLAIR images in ISLES 2015.

Notable improvements include: ARC T2w increasing from 26.8% to 37.2% Dice, ISLES T1w from 11.0% to 17.9%, and ARC ensemble from 60.2% to 69.4%. The baseline model shows minimal improvement with pseudo-labels in out-of-domain scenarios (e.g., ISLES ensemble re-

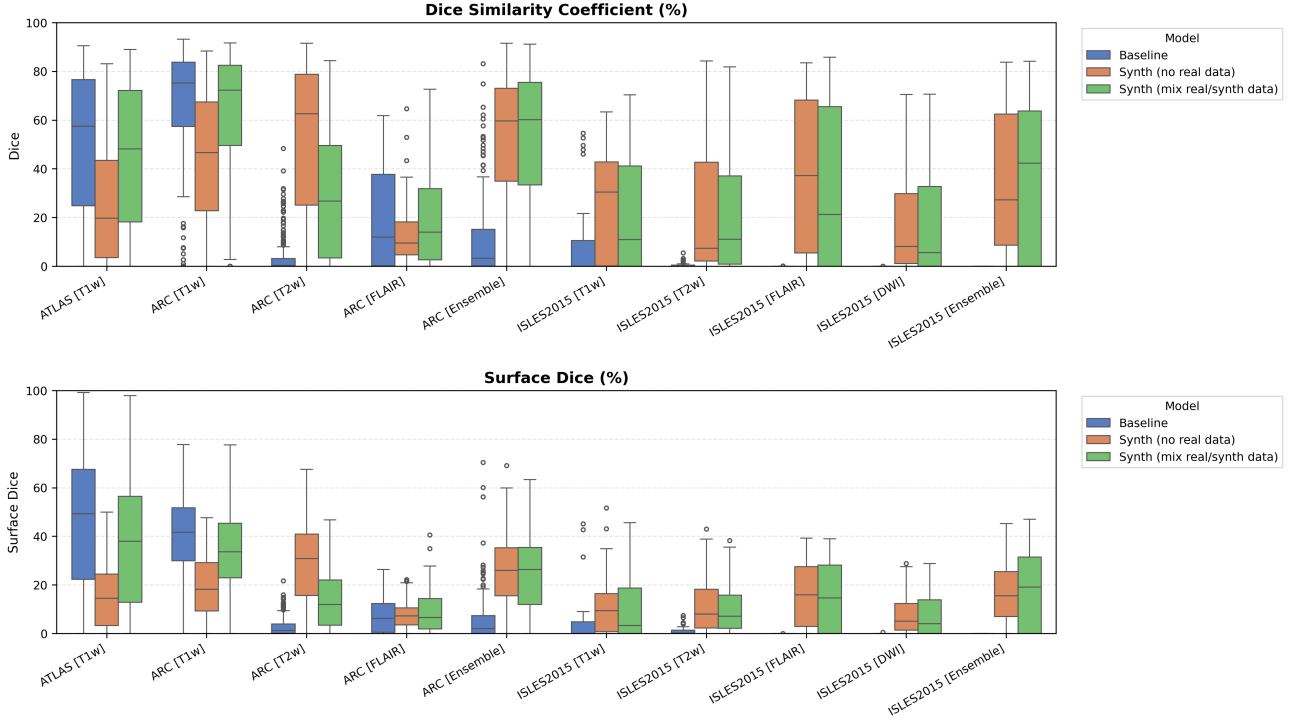


Figure 5: Dice and Surface Dice metrics for all reported datasets, for models trained on different combinations of real/synthetic data.

Table 8: Median results for ablation and pseudo-label experiments on the ISLES 2015 test set ($N=28$). Best score shown in bold. Student’s t distribution 95% confidence intervals given in brackets.

Modality	Model	Dice (%)	Surface Dice (%)
T1w	Baseline	0.0 (0.0-7.4)	0.0 (0.0-4.8)
	Baseline+Pseudo	0.3 (0.0-9.1)	0.0 (0.0-4.0)
	Synth (no real data)	30.4 (21.3-39.6)	9.4 (4.1-14.8)
	Synth (Ours)	11.0 (1.2-20.8)	3.4 (0.0-8.5)
	Synth+Pseudo	17.9 (7.6-28.3)	8.0 (2.1-13.8)
T2w	Baseline	0.0 (0.0-0.5)	0.3 (0.0-1.0)
	Baseline+Pseudo	0.1 (0.0-0.7)	0.4 (0.0-1.1)
	Synth (no real data)	7.4 (0.0-18.4)	8.1 (3.0-13.1)
	Synth (Ours)	11.1 (0.7-21.6)	7.1 (2.7-11.5)
	Synth+Pseudo	18.1 (7.1-29.1)	9.5 (4.4-14.5)
FLAIR	Baseline	0.0 (0.0-0.0)	0.0 (0.0-0.0)
	Baseline+Pseudo	0.0 (0.0-0.3)	0.0 (0.0-0.4)
	Synth (no real data)	37.2 (25.5-48.9)	16.0 (10.8-21.1)
	Synth (Ours)	21.2 (8.5-34.0)	14.7 (9.0-20.3)
	Synth+Pseudo	4.2 (0.0-17.3)	3.9 (0.0-9.8)
DWI	Baseline	0.0 (0.0-0.0)	0.0 (0.0-0.0)
	Baseline+Pseudo	0.0 (0.0-0.4)	0.0 (0.0-0.0)
	Synth (no real data)	8.2 (0.0-16.8)	5.1 (2.0-8.3)
	Synth (Ours)	5.6 (0.0-14.4)	4.1 (0.8-7.4)
	Synth+Pseudo	5.6 (0.0-14.7)	4.3 (0.8-7.8)
Ensemble	Baseline	0.0 (0.0-0.0)	0.0 (0.0-0.0)
	Baseline+Pseudo	0.0 (0.0-0.0)	0.0 (0.0-0.0)
	Synth (no real data)	27.2 (15.7-38.8)	15.6 (10.3-20.9)
	Synth (Ours)	42.3 (30.2-54.5)	19.1 (12.7-25.4)
	Synth+Pseudo	50.1 (37.6-62.6)	24.9 (18.7-31.1)

mains at 0.0%), suggesting that pseudo-labelling cannot overcome fundamental domain shift. The FLAIR degradation in ISLES 2015 (21.2% to 4.2%) may indicate that pseudo-labels from chronic MPRAGE data introduce biases

incompatible with acute FLAIR lesion appearance. The in-domain ATLAS dataset also shows a notable improvement for both models as a result of pseudo labels, with a more marked increase in the baseline model, suggesting pseudo-labels effectively expand the training distribution when the domain gap is small.

These results across four diverse datasets - from research-quality ATLAS to challenging clinical PLORAS scans - reveal consistent patterns that warrant deeper analysis of the fundamental principles underlying domain robustness.

5. Discussion

Our results reveal fundamental principles about domain robustness in medical image segmentation. The consistent failure of domain adaptation to rescue baseline models - even with oracle selection of the optimal method - demonstrates that post-hoc adaptation cannot substitute for domain-invariant training. This finding challenges prevailing assumptions about the sufficiency of test-time adaptation for clinical deployment.

The compound effect observed with Synth+DAE suggests that synthetic pre-training and test-time adaptation address different aspects of domain shift. Synthetic training provides feature representations invariant to appearance changes, while DA methods fine-tune these representations to specific test-time distributions. However, this synergy only emerges when the base model already pos-

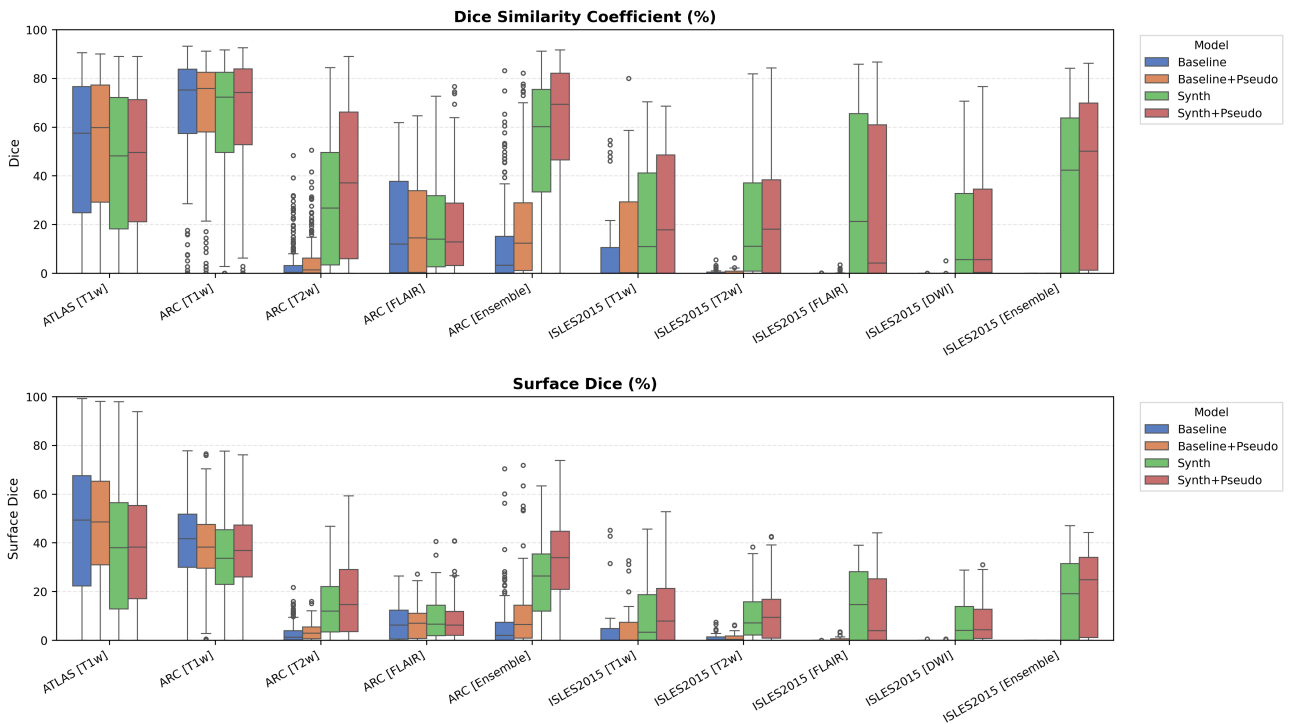


Figure 6: Dice and Surface Dice metrics for all reported datasets, for models trained with/without an additional training dataset of MPRAGE images and pseudo-labels.

sesses sufficient robustness - when initial predictions fail catastrophically (as with baseline models on ISLES 2015 and PLORAS), no amount of adaptation can recover performance.

The variable benefit of multi-modal ensembling - substantial for ISLES 2015 but mixed for ARC - indicates that fusion strategies must consider dataset-specific characteristics rather than assuming universal improvement. Clinical deployment should balance single-modality robustness with opportunistic multi-modal fusion.

6. Limitations and Future Directions

Appendix A confirms that 20–40% of lesions in several cohorts are multimodal, underscoring the central limitation of our single-Gaussian sampling. While our spatially-varying approach introduces intra-lesion heterogeneity, it does not address the multi-modal nature of stroke appearance. Future implementations should explore mixture models to capture lesions that simultaneously exhibit hyper- and hypo-intense regions within the same pathology. It may also be advantageous to paste several lesions per subject, each assigned its own independently sampled intensity profile. This would allow a single synthetic brain to display chronic hypointense scars alongside acute hyper-intense infarcts, better reflecting the heterogeneous lesion chronology typically observed in stroke cohorts.

Additional qualitative results in Appendix Figures 18 -

23 display further positive and negative results across the available image modalities within the PLORAS dataset. A number of failure modes may be observed here, such as missing cerebellar lesions or under-segmenting large hemispheric lesions.

Although results in the paper provide strong evidence of the raw value of the segmentation metrics, it is important to further validate that the predictions made from the trained model are useful for downstream tasks. This could be validated by comparing predictivity of the segmentation masks for functional scores such as the Comprehensive Aphasia Test (CAT) (Hope et al., 2024) or the NIH Stroke Score (NIHSS).

In the context of neuroimaging data, there are many shifts in domain beyond those related to scanner sequence and resolution targeted in this work. Shifts related to anatomical shape, brought on by changes in demographics or the presence of confounding conditions such as atrophy are of equal importance to shifts in image appearance. Prior work has attempted to model such factors through the use of causal inference and counterfactual generation (Pawlowski et al., 2020; Pombo et al., 2023; Ribeiro et al., 2023). It is conceivable that such an approach could be used to introduce morphological changes to the healthy tissue maps to create a model that is agnostic to shifts in domains related to both shape and appearance.

While this study focused on stroke, the results are expected to translate to other similar domains, such as haem-

orrhage and glioblastoma. Future work will compare the impact of the mixing proportions of real and synthetic data. Additionally, the utility of multi-modal data in a multi-channel model will be examined versus post-hoc averaging of individual modality predictions. Lesions often appear differently across modalities, so a model trained with multi-channel inputs is expected to leverage these differences more effectively. This relationship between multi-channel inputs may better be modelled using quantitative MRI data.

7. Conclusion

In this study, we introduced a novel synthetic data generation and training framework for stroke lesion segmentation, building on the success of prior works in healthy brain parcellation. A model trained using this novel framework was evaluated on a wide range of datasets covering research and clinical data, and chronic and acute stroke pathologies. Our experiments demonstrate that synthetic pre-training provides fundamental robustness unachievable through test-time adaptation alone. While accepting a 9.3% median Dice reduction in-domain (48.2% vs 57.5%), our approach maintains performance where conventional methods fail entirely. Even with oracle knowledge of the optimal domain adaptation method - an unrealistic scenario in practice - conventionally-trained models cannot match our synthetic approach in out-of-domain settings.

Our results demonstrate that even imperfect appearance modelling can provide substantial benefits for cross-modality segmentation. Incorporating recent contemporary work on realistic lesion simulation (Liu et al., 2025) may further improve performance both in- and out-of-distribution.

Qualitative results in Figure 4 demonstrate the potential of the model as a starting checkpoint in an active-learning framework such as MONAI Label (Diaz-Pinto et al., 2024), where predictions may be refined to generate training data for task-specific fine-tuning. The uncertainty of healthy-tissue segmentations via MC dropout may also provide an effective heuristic for prioritising items to refine/label in such a framework (Nath et al., 2021).

Results also showed the potential benefit of the model for semi-supervised learning through pseudo labelling, where predicted labels may be fed back into the model with much less caution than is typically required when training with real images (Pham et al., 2021; Xu et al., 2024). The ablation study revealed that pure synthetic training can outperform mixed training for certain out-of-domain modalities, suggesting opportunities for further optimisation of the real/synthetic data balance. A simple solution to further reinforce this may be to train a real/fake discriminator to perform automated quality-control and threshold optimisation for the generated binary maps.

Our approach reduces reliance on domain-specific training data and helps bridge the gap between research-grade and clinical scans to improve clinical stroke neuroimaging workflows, providing a foundation for more robust and widely applicable lesion segmentation tools.

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Ethical Standards

The work follows appropriate ethical standards in conducting research and writing the manuscript, following all applicable laws and regulations regarding treatment of animals or human subjects.

Conflicts of Interest

We declare we don't have conflicts of interest.

Data availability

Authors have released public code and weights to reimplement all experiments. Model weights are also made available as a toolbox for SPM. This toolbox is written in pure MATLAB and requires no external installs or compilation, to minimise the barrier for clinical evaluation. All datasets are public except the PLORAS study data, which is part of an ongoing study and will be published upon completion.

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Appendix A. Lesion-intensity unimodality

Motivation. Most classical stroke-segmentation pipelines, including our synthetic generator, implicitly assume that the grey-level distribution of a lesion is *unimodal*. That assumption underpins common appearance priors such as single-Gaussian modelling and simple intensity normalisation. However, chronic infarcts often contain a mixture of tissue constituents, and in CT the same infarct can include both hypo- and hyper-dense cores. Quantifying how often real lesions violate unimodality therefore informs whether richer mixture-based priors are warranted.

Experiment. For every connected component in each dataset, we extracted the raw voxel intensities and applied Hartigan’s dip test (Hartigan and Hartigan, 1985) to the empirical one-dimensional distribution. A lesion was labelled *unimodal* when the null hypothesis of unimodality could not be rejected at $\alpha = 0.05$; otherwise it was classified as *multimodal*. For efficiency, we randomly subsampled at most 2048 voxels per lesion before testing. Table 9 summarises the counts.

Findings. Across MRI datasets the majority of lesions were indeed unimodal, but a non-negligible tail of multimodal cases emerged:

- **ATLAS** (chronic MPRAGE T1w) showed $> 95\%$ unimodal lesions; only 80 of 1818 connected components exhibited multimodality, in line with mature cavities whose signal is dominated by a single CSF-like class.
- **ISLES 2015** exhibited 12-38% multimodal lesions depending on modality, reflecting mixed-phase tissue in a sub-acute cohort.
- **ARC** (chronic research scans) retained a high unimodal rate in T1w/T2w but FLAIR contained 39% multimodal lesions.
- **PLORAS** revealed the strongest departure: while T2w/FLAIR remained largely unimodal, CT lesions were predominantly multimodal (58%), underscoring how density heterogeneity dominates in CT.

Implications. A single-Gaussian appearance prior is largely adequate for T1w/T2w MRI, where $\geq 80\%$ of lesions were unimodal; it begins to break down for modalities that emphasise subtle tissue heterogeneity - most notably FLAIR and especially CT. Adopting explicit mixture-based priors, or synthetically sampling lesions with mixed intensities, therefore represents a principled next step for improving model robustness across modalities and imaging protocols.

Table 9: Counts of connected-component lesions that are **unimodal** or **multimodal** in each dataset–modality pair. For every lesion larger than 10 voxels we sampled at most 2048 raw intensity values and ran Hartigan’s dip test; failure to reject the null hypothesis of unimodality at $\alpha = 0.05$ yields the **Unimodal** label, otherwise **Multimodal**. ATLAS is split into the subjects used for model development (T1w [Train]) and the held-out evaluation set (T1w [Validation]).

Dataset	Modality	Unimodal	Multimodal
ATLAS	T1w [Train]	1384	65
	T1w [Validation]	354	15
ISLES2015	T1w	52	9
	T2w	46	15
	FLAIR	54	7
	DWI	38	23
ARC	T1w	286	64
	T2w	335	48
	FLAIR	90	57
PLORAS	T2w	203	14
	FLAIR	712	29
	CT	248	342

Appendix B. Further segmentation metrics

Additional metrics are reported below in Appendix Tables 10 - 13 for all datasets. Here we report individual scores for each DA method, for both the baseline model and our Synth model.

Table 10: Mean results on ATLAS hold-out set ($N=131$). Best score shown in bold. FPR values are shown as a multiple of 10^{-3} .

Modality	Model	Dice	HD95	AVD	ALD	LF1	TPR	FPR
T1w	Baseline	0.500	44.3	7.60	3.48	0.480	0.510	0.251
	Baseline+TTA	0.503	41.4	7.34	2.52	0.537	0.502	0.234
	Synth (Ours)	0.456	47.4	8.73	2.26	0.536	0.426	0.200
	Synth+TTA	0.455	48.3	8.51	1.76	0.559	0.424	0.191

Table 11: Mean results on the ARC dataset ($N=229$). Best score per column is shown in bold. FPR values are shown as a multiple of 1000.

Modality	Model	Dice	HD95	AVD	ALD	LF1	TPR	FPR
T1w	Baseline	0.646	24.5	18.04	6.19	0.335	0.600	1.178
	Baseline+TTA	0.648	23.7	18.33	3.42	0.442	0.598	1.126
	Baseline+TENT	0.077	144.9	75.50	3.51	0.180	0.060	0.285
	Baseline+DAE	0.614	26.2	26.53	3.15	0.507	0.685	2.238
	Baseline+PL	0.325	67.5	146.72	175.61	0.017	0.613	10.342
	Baseline+UPL	0.341	74.4	48.10	22.84	0.244	0.262	0.695
	Baseline+DPL	0.340	52.5	52.48	9.53	0.324	0.247	0.465
	Synth (Ours)	0.609	27.9	22.88	1.99	0.613	0.539	0.953
	Synth+TTA	0.611	27.0	24.08	1.84	0.640	0.534	0.885
	Synth+TENT	0.560	33.4	26.71	2.42	0.528	0.604	2.385
	Synth+DAE	0.613	23.4	20.60	2.26	0.581	0.640	1.903
	Synth+PL	0.311	158.6	48.41	260.54	0.017	0.310	2.303
T2w	Baseline	0.040	76.8	73.80	47.12	0.044	0.063	6.778
	Baseline+TTA	0.023	77.0	71.83	26.00	0.054	0.036	6.819
	Baseline+TENT	0.000	148.5	85.40	5.11	0.047	0.000	0.206
	Baseline+DAE	0.056	75.8	123.49	10.09	0.113	0.134	10.938
	Baseline+PL	0.006	78.8	127.78	353.31	0.007	0.014	12.731
	Baseline+UPL	0.000	256.0	85.74	2.57	0.013	0.000	0.185
	Baseline+DPL	0.000	256.0	85.74	2.57	0.013	0.000	0.185
	Synth (Ours)	0.299	66.2	40.79	38.81	0.060	0.304	2.282
	Synth+TTA	0.284	66.2	43.70	16.00	0.117	0.274	1.893
	Synth+TENT	0.321	55.5	43.43	15.09	0.140	0.326	1.663
	Synth+DAE	0.437	40.8	40.19	11.43	0.196	0.410	1.344
	Synth+PL	0.384	65.5	40.26	22.77	0.095	0.475	3.881
FLAIR	Baseline	0.193	66.2	52.15	49.49	0.049	0.205	4.532
	Baseline+TTA	0.202	65.7	47.98	25.98	0.090	0.197	3.719
	Baseline+TENT	0.011	165.6	79.40	4.94	0.070	0.008	0.628
	Baseline+DAE	0.287	54.8	75.47	11.09	0.232	0.327	5.981
	Baseline+PL	0.126	81.9	162.13	382.66	0.007	0.240	13.548
	Baseline+UPL	0.000	256.0	82.36	2.62	0.035	0.000	0.497
	Baseline+DPL	0.000	256.0	82.36	2.62	0.035	0.000	0.497
	Synth (Ours)	0.199	59.3	60.36	17.69	0.150	0.157	1.349
	Synth+TTA	0.190	56.4	62.98	8.36	0.256	0.143	1.185
	Synth+TENT	0.340	49.7	46.05	5.01	0.247	0.286	1.582
	Synth+DAE	0.263	51.5	53.14	5.88	0.248	0.215	1.581
	Synth+PL	0.276	96.8	45.97	93.18	0.027	0.255	2.133
Ensemble	Baseline	0.107	53.3	77.04	22.20	0.309	0.071	1.137
	Baseline+TTA	0.087	65.8	78.91	9.10	0.429	0.057	1.126
	Baseline+TENT	0.003	238.0	85.73	2.60	0.054	0.002	0.185
	Baseline+DAE	0.212	51.3	73.84	39.25	0.168	0.212	2.350
	Baseline+PL	0.016	76.1	70.76	512.92	0.004	0.014	3.869
	Baseline+UPL	0.000	256.0	85.74	2.57	0.013	0.000	0.185
	Baseline+DPL	0.000	256.0	85.74	2.57	0.013	0.000	0.185
	Synth (Ours)	0.525	34.9	35.60	10.09	0.356	0.451	0.770
	Synth+TTA	0.530	33.0	36.51	4.50	0.470	0.452	0.696
	Synth+TENT	0.451	39.1	38.50	11.12	0.370	0.412	0.960
	Synth+DAE	0.579	27.6	27.52	6.77	0.432	0.531	0.923
	Synth+PL	0.529	50.2	33.21	29.02	0.106	0.515	1.551
	Synth+UPL	0.244	53.0	65.77	9.80	0.470	0.159	0.310
	Synth+DPL	0.172	71.4	71.62	13.97	0.414	0.107	0.284

Appendix C. Wilcoxon Significance Tests

Significance measures are provided below in Figures 7 - 17 as Wilcoxon signed-rank test values for pairwise comparisons between models on each dataset. 'Oracle DA' represents the hypothetical best-case scenario where optimal DA method is known a priori for each dataset/modality and applied to the baseline model. Median Dice values are shown along the diagonal.

Table 12: Mean results on the ISLES2015 dataset ($N=28$). Best score per column is shown in bold. FPR values are shown as a multiple of 1000.

Modality	Model	Dice	HD95	AVD	ALD	LF1	TPR	FPR
T1w	Baseline	0.115	84.2	38.14	3.32	0.269	0.076	0.034
	Baseline+TTA	0.091	113.2	40.11	1.96	0.311	0.058	0.012
	Baseline+TENT	0.002	178.2	43.77	2.14	0.161	0.001	0.001
	Baseline+DAE	0.246	68.2	48.07	7.18	0.285	0.191	1.604
	Baseline+PL	0.040	81.6	42.79	643.29	0.003	0.039	1.979
	Baseline+UPL	0.000	256.0	43.79	2.11	0.000	0.000	0.000
	Baseline+DPL	0.000	256.0	43.79	2.11	0.000	0.000	0.000
	Synth (Ours)	0.222	70.5	30.49	7.14	0.208	0.168	0.162
	Synth+TTA	0.230	78.5	31.41	2.18	0.349	0.163	0.085
	Synth+TENT	0.076	108.2	33.79	40.86	0.103	0.058	0.265
	Synth+DAE	0.268	58.1	29.18	3.64	0.316	0.223	0.293
	Synth+PL	0.050	76.9	42.55	3.93	0.321	0.028	0.008
	Synth+UPL	0.000	241.7	43.79	2.04	0.050	0.000	0.000
T2w	Baseline	0.001	221.7	43.77	2.46	0.127	0.000	0.000
	Baseline+TTA	0.007	65.4	50.81	15.00	0.078	0.044	2.786
	Baseline+TENT	0.003	65.5	48.97	6.86	0.095	0.025	2.260
	Baseline+DAE	0.001	142.6	43.38	3.75	0.050	0.001	0.085
	Baseline+PL	0.006	71.5	88.17	9.43	0.071	0.069	5.805
	Baseline+UPL	0.002	74.2	71.28	155.21	0.013	0.005	0.555
	Baseline+DPL	0.000	256.0	43.79	2.11	0.000	0.000	0.000
	Synth (Ours)	0.230	256.0	43.79	2.11	0.000	0.000	0.000
	Synth+TTA	0.001	74.8	52.73	11.82	0.166	0.443	3.534
	Synth+TENT	0.245	74.6	50.99	4.68	0.279	0.454	3.379
	Synth+DAE	0.215	77.4	33.44	20.11	0.109	0.276	1.755
	Synth+PL	0.255	63.1	31.10	4.61	0.280	0.328	1.437
	Synth+UPL	0.245	73.3	37.57	13.71	0.161	0.288	2.280
FLAIR	Baseline	0.152	63.3	31.06	12.43	0.138	0.116	0.631
	Synth+DPL	0.111	58.6	35.25	8.11	0.172	0.073	0.143
	Baseline	0.000	71.4	40.06	10.79	0.018	0.000	0.515
	Baseline+TTA	0.000	112.9	41.24	3.43	0.000	0.000	0.238
	Baseline+TENT	0.000	119.0	43.47	4.32	0.000	0.000	0.021
	Baseline+DAE	0.000	80.1	61.65	15.29	0.015	0.004	3.679
	Baseline+PL	0.000	77.1	43.43	70.43	0.009	0.000	0.490
	Baseline+UPL	0.000	256.0	43.79	2.11	0.000	0.000	0.000
	Baseline+DPL	0.000	256.0	43.79	2.11	0.000	0.000	0.000
	Synth (Ours)	0.314	62.9	18.84	10.61	0.154	0.346	0.975
	Synth+TTA	0.329	82.1	18.37	3.50	0.275	0.359	0.891
	Synth+TENT	0.127	79.0	28.03	34.54	0.031	0.107	0.460
	Synth+DAE	0.274	55.7	20.25	7.86	0.152	0.305	0.989
DWI	Synth+PL	0.252	75.7	50.59	111.43	0.036	0.330	3.176
	Synth+UPL	0.232	137.9	24.88	3.04	0.271	0.187	0.111
	Synth+DPL	0.205	145.6	25.16	7.36	0.236	0.168	0.191
	Baseline	0.000	80.1	42.63	5.54	0.023	0.000	0.138
	Baseline+TTA	0.000	102.8	43.83	2.43	0.000	0.000	0.133
	Baseline+TENT	0.000	103.9	43.52	1.89	0.000	0.000	0.021
	Baseline+DAE	0.000	76.2	40.04	11.21	0.025	0.000	2.123
	Baseline+PL	0.004	77.5	40.33	835.36	0.001	0.004	0.726
	Baseline+UPL	0.000	256.0	43.79	2.11	0.000	0.000	0.000
	Baseline+DPL	0.000	256.0	43.79	2.11	0.000	0.000	0.000
	Synth (Ours)	0.193	82.5	47.64	16.71	0.100	0.352	2.741
	Synth+TTA	0.204	81.1	44.32	8.39	0.187	0.336	2.344
	Synth+TENT	0.104	86.6	30.71	38.50	0.051	0.108	0.882
Ensemble	Synth+DAE	0.185	80.4	39.10	12.36	0.106	0.262	2.139
	Synth+PL	0.186	85.2	81.64	17.43	0.120	0.397	5.759
	Synth+UPL	0.179	60.8	32.42	19.43	0.154	0.155	0.268
	Synth+DPL	0.089	117.4	39.27	3.75	0.300	0.060	0.031
	Baseline	0.000	242.8	43.79	2.00	0.000	0.000	0.000
	Baseline+TTA	0.000	256.0	43.79	2.11	0.000	0.000	0.000
	Baseline+TENT	0.000	256.0	43.79	2.11	0.000	0.000	0.000
	Baseline+DAE	0.000	231.0	43.68	2.00	0.000	0.000	0.007
	Baseline+PL	0.000	198.9	43.79	2.11	0.024	0.000	0.000
	Baseline+UPL	0.000	256.0	43.79	2.11	0.000	0.000	0.000
	Baseline+DPL	0.000	256.0	43.79	2.11	0.000	0.000	0.000
	Synth (Ours)	0.370	60.4	19.68	7.50	0.234	0.339	0.256
	Synth+TTA	0.378	69.7	20.01	2.75	0.348	0.334	0.223
	Synth+TENT	0.111	114.8	32.23	3.21	0.235	0.089	0.047
	Synth+DAE	0.365	51.4	19.84	3.75	0.354	0.331	0.204
	Synth+PL	0.246	60.7	29.01	21.18	0.125	0.188	0.150
	Synth+UPL	0.016	218.2	42.52	5.18	0.131	0.009	0.000
	Synth+DPL	0.033	187.3	41.09	6.04	0.223	0.020	0.000

Appendix D. Additional Qualitative Results

Results are shown below in Figures 18 - 23 for failure and success cases for the three different sequences in the PLO-RAS dataset.

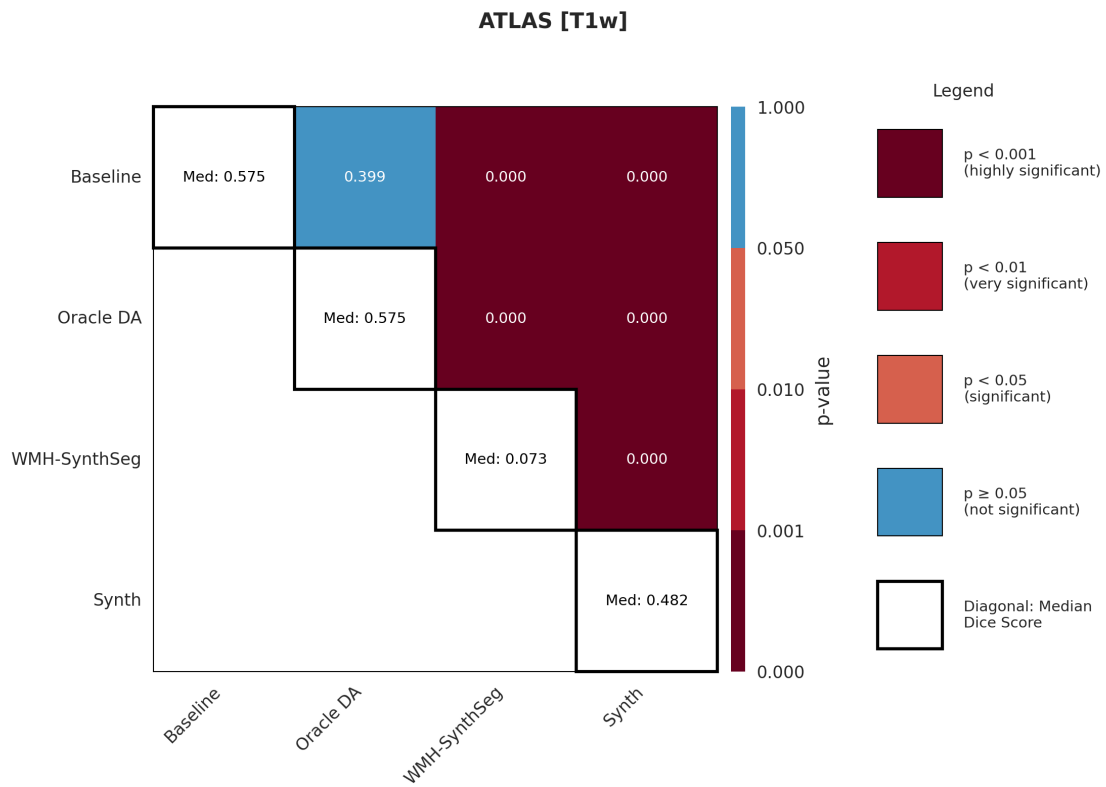


Figure 7: Wilcoxon signed-rank test values for Dice metric measurements in the ATLAS T1w dataset.

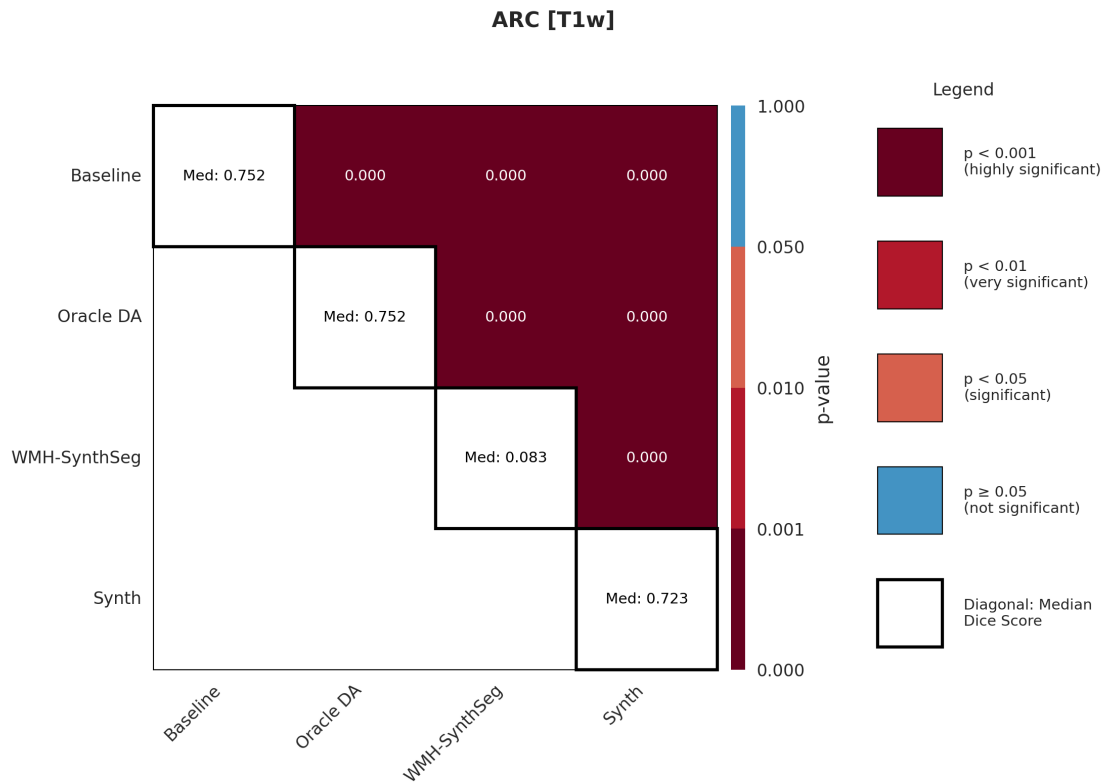


Figure 8: Wilcoxon signed-rank test values for Dice metric measurements in the ARC T1w dataset.

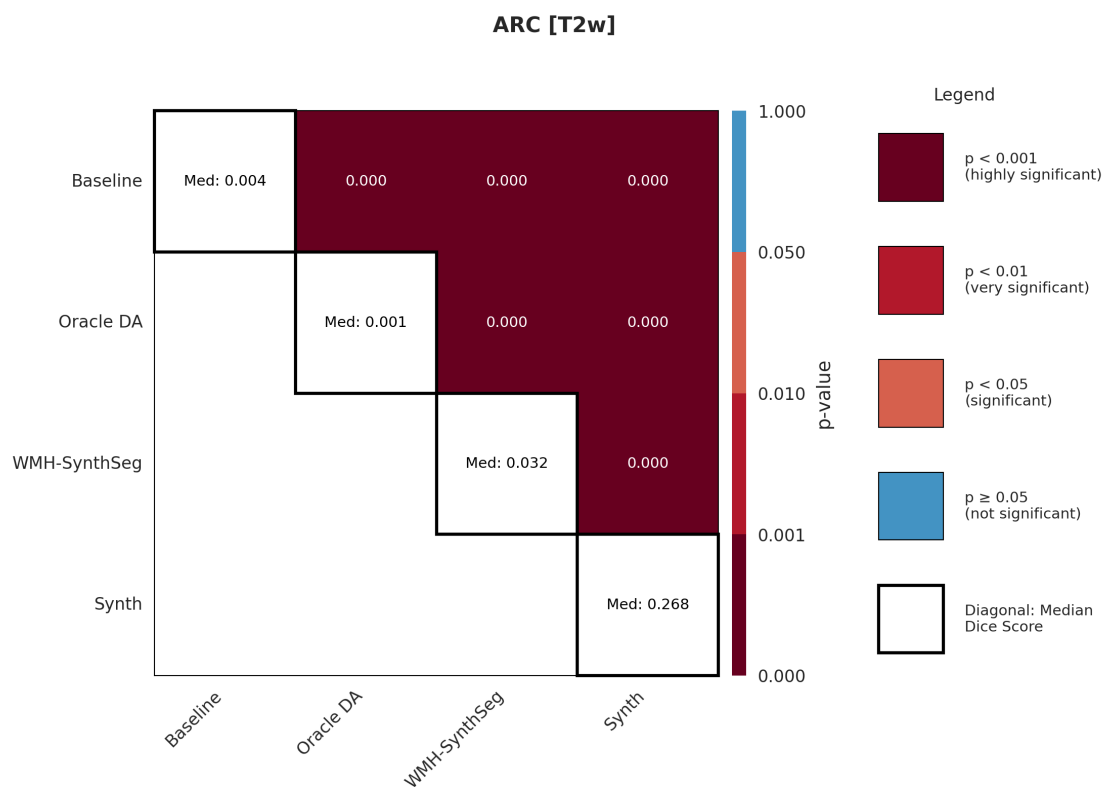


Figure 9: Wilcoxon signed-rank test values for Dice metric measurements in the ARC T2w dataset.

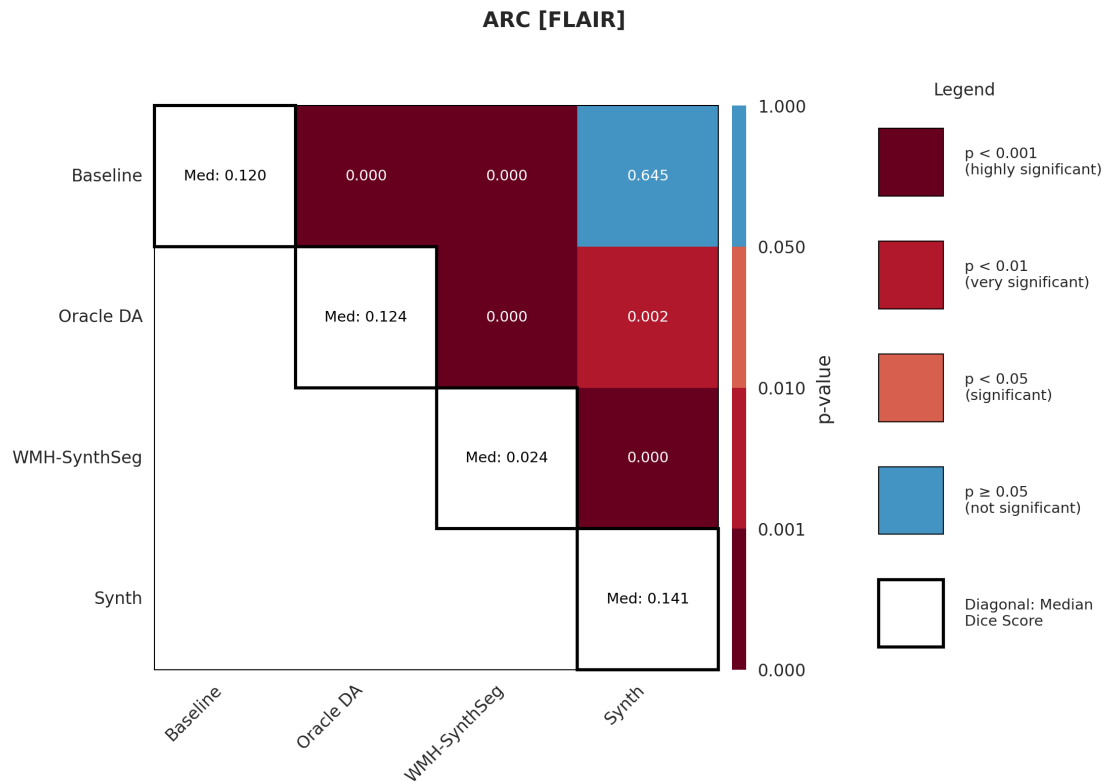


Figure 10: Wilcoxon signed-rank test values for Dice metric measurements in the ARC FLAIR dataset.

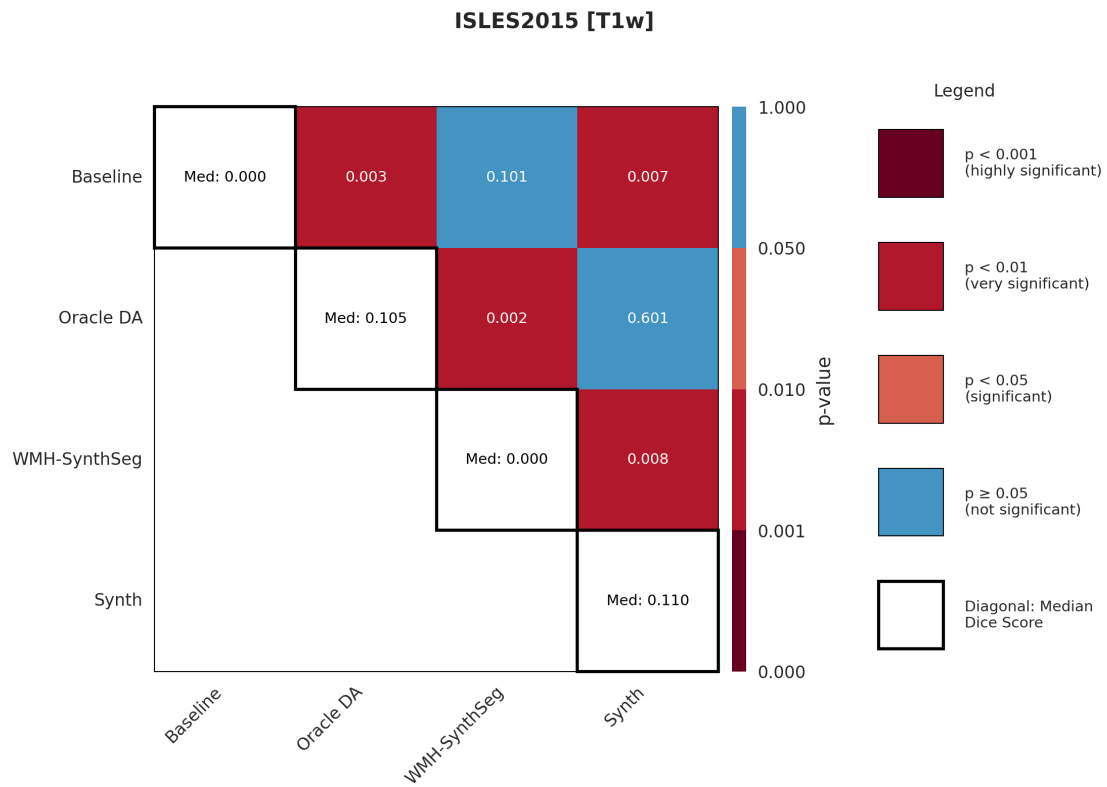


Figure 11: Wilcoxon signed-rank test values for Dice metric measurements in the ISLES 2015 T1w dataset.

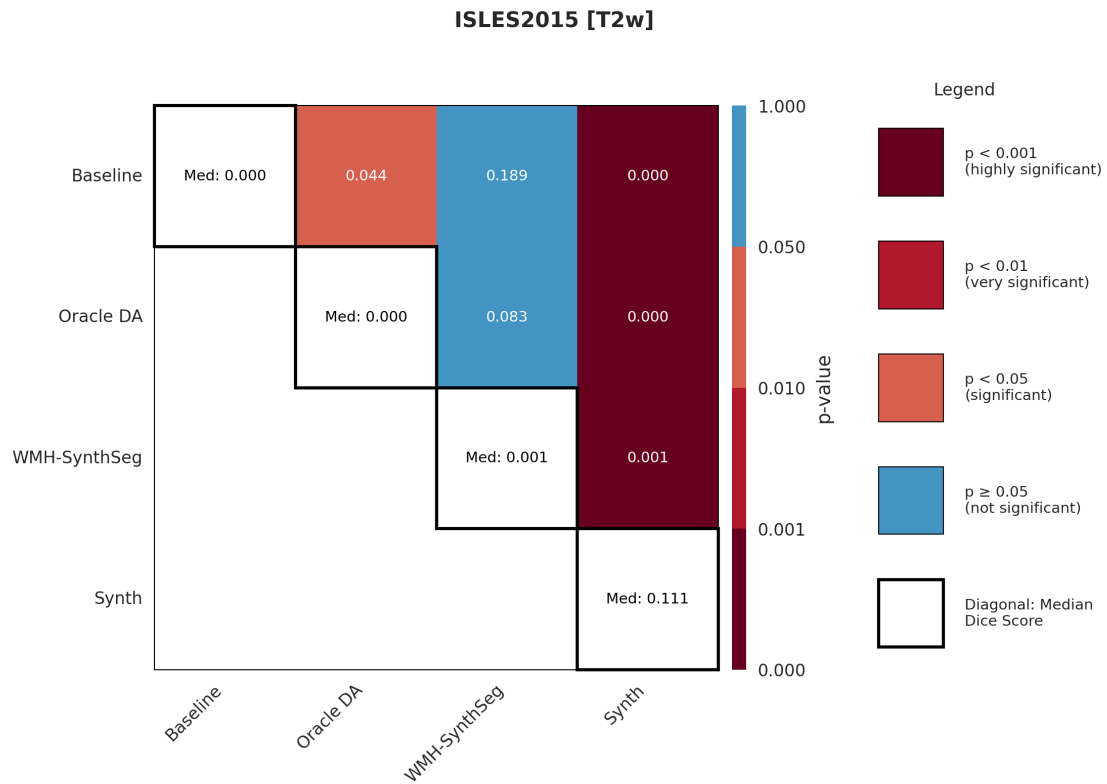


Figure 12: Wilcoxon signed-rank test values for Dice metric measurements in the ISLES 2015 T2w dataset.

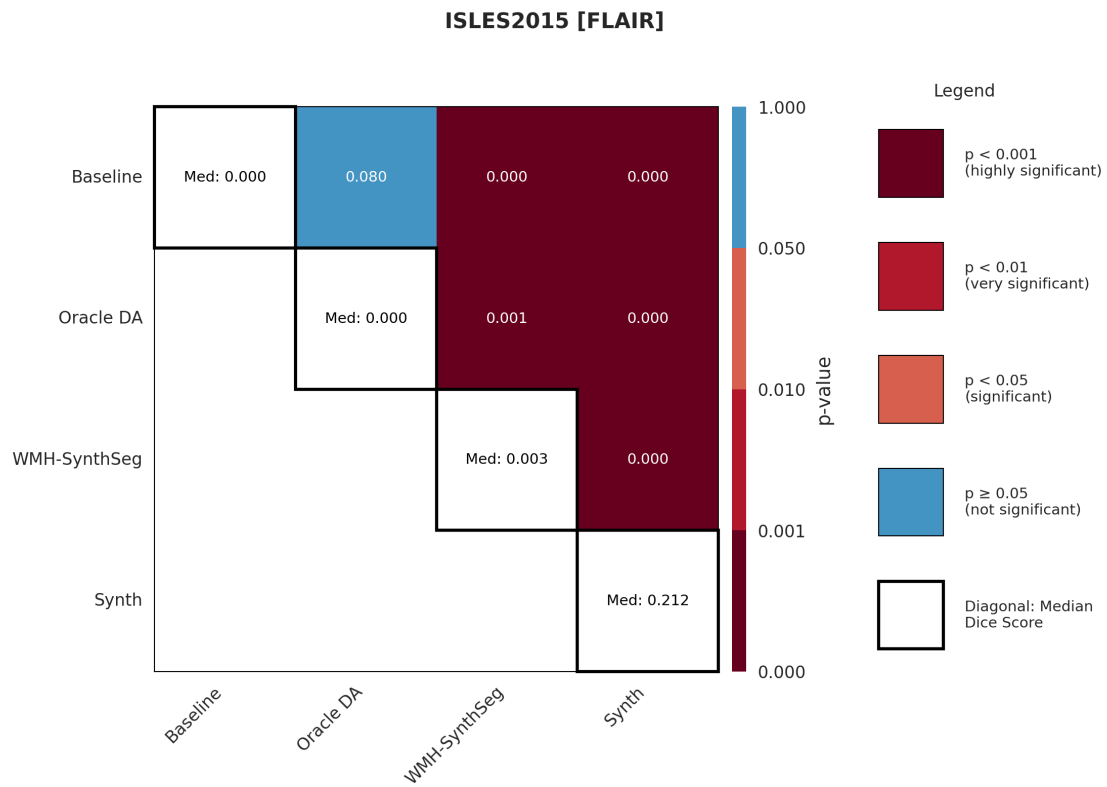


Figure 13: Wilcoxon signed-rank test values for Dice metric measurements in the ISLES 2015 FLAIR dataset.

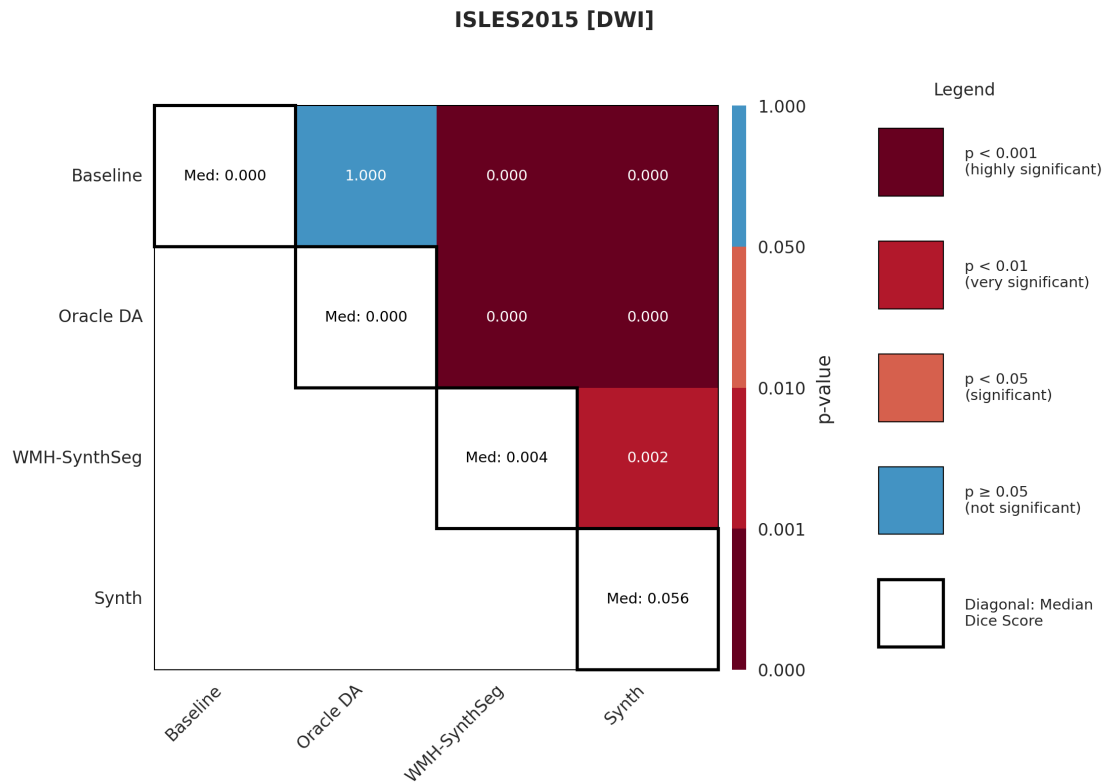


Figure 14: Wilcoxon signed-rank test values for Dice metric measurements in the ISLES 2015 DWI dataset.

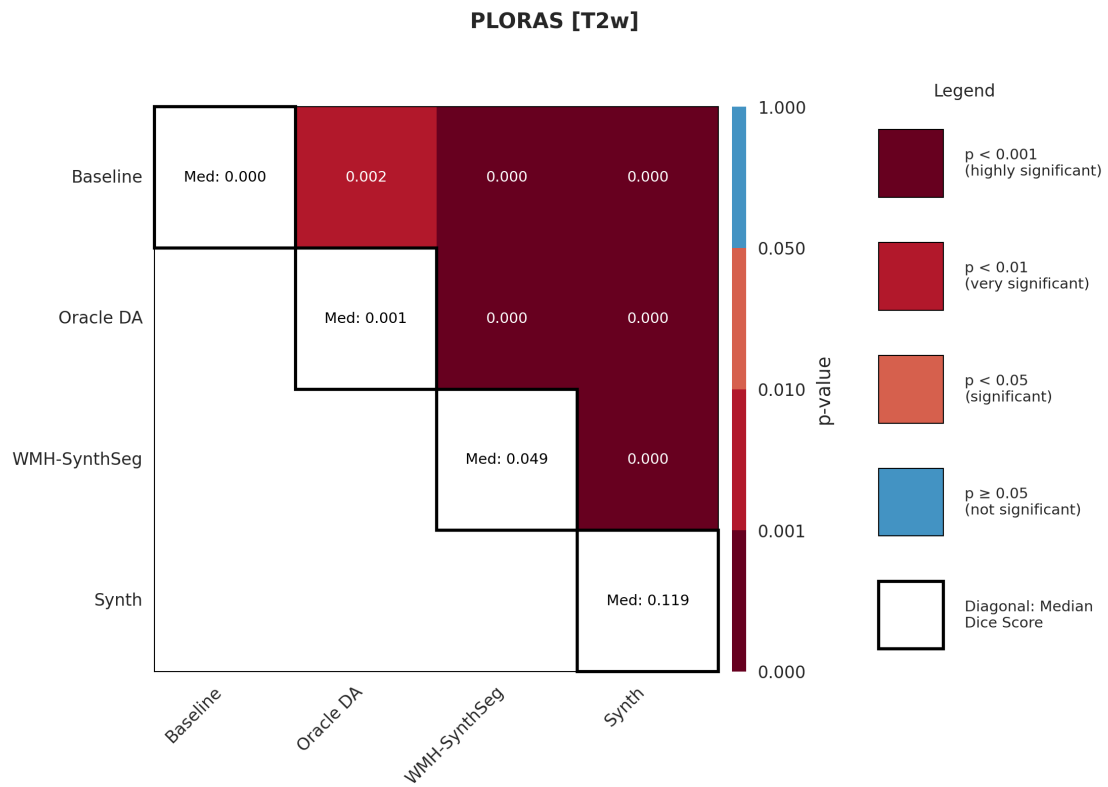


Figure 15: Wilcoxon signed-rank test values for Dice metric measurements in the PLORAS T2w dataset.

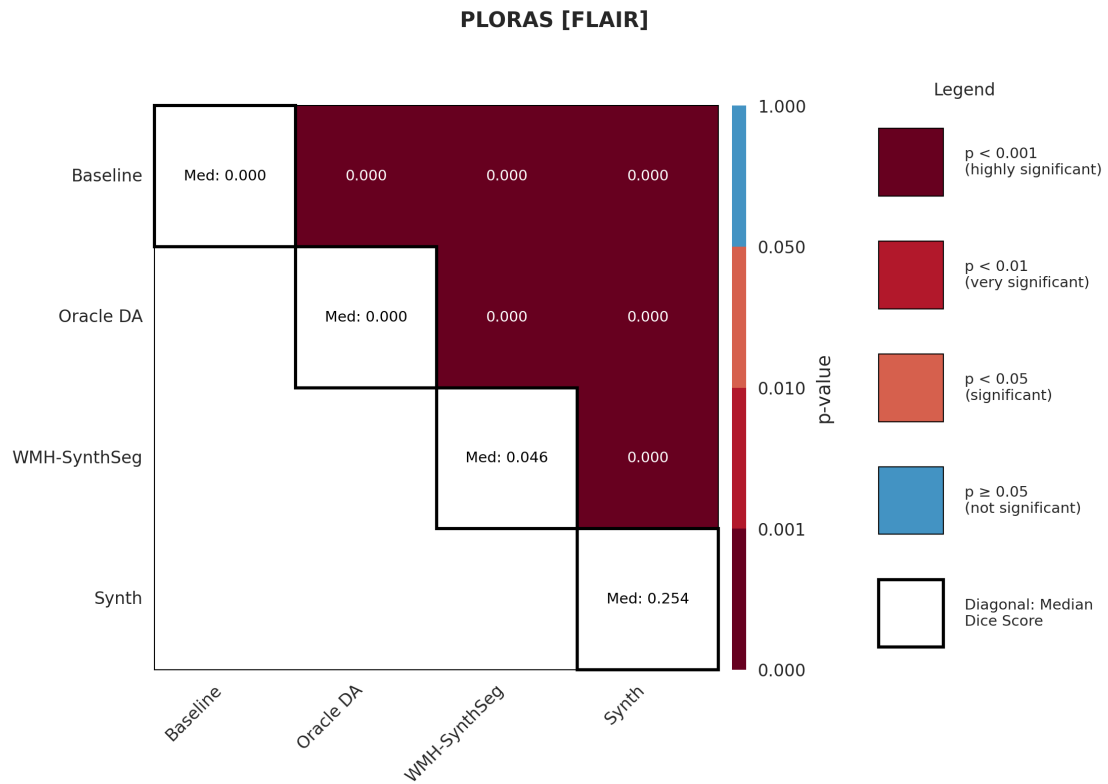


Figure 16: Wilcoxon signed-rank test values for Dice metric measurements in the PLORAS FLAIR dataset.

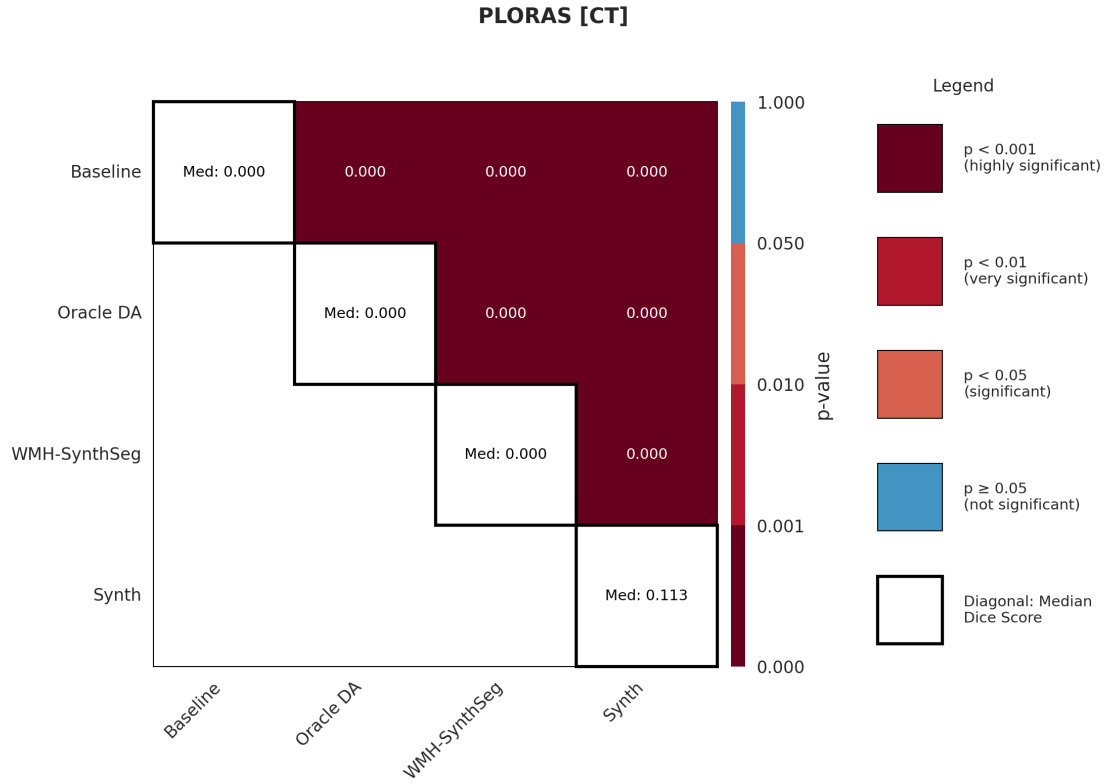


Figure 17: Wilcoxon signed-rank test values for Dice metric measurements in the PLORAS CT dataset.

Table 13: Mean results on the PLORAS dataset ($N=661$). Best score per column is shown in bold. FPR values are shown as a multiple of 1000.

Modality	Model	Dice	HD95	AVD	ALD	LF1	TPR	FPR
T2w	Baseline	0.028	80.3	112.82	12.22	0.101	0.064	9.638
	Baseline+TTA	0.020	79.1	92.69	11.02	0.103	0.045	8.409
	Baseline+TENT	0.005	101.5	30.79	5.12	0.020	0.007	1.849
	Baseline+DAE	0.044	82.7	204.49	2.64	0.276	0.123	14.983
	Baseline+PL	0.000	235.8	32.46	2.13	0.009	0.004	1.696
	Baseline+UPL	0.000	256.0	32.47	2.12	0.009	0.004	1.695
	Baseline+DPL	0.000	256.0	32.47	2.12	0.009	0.004	1.695
	Synth (Ours)	0.250	71.9	40.20	8.42	0.243	0.371	4.156
	Synth+TTA	0.253	70.5	37.00	4.75	0.329	0.361	3.917
	Synth+TENT	0.202	77.0	61.99	21.23	0.118	0.330	5.612
	Synth+DAE	0.291	58.7	23.68	4.28	0.307	0.312	2.969
	Synth+PL	0.162	81.0	33.28	8.58	0.179	0.186	3.737
FLAIR	Synth+UPL	0.101	66.0	25.80	11.07	0.186	0.073	2.014
	Synth+DPL	0.053	72.1	28.44	3.42	0.212	0.036	1.832
	Baseline	0.010	74.2	42.12	18.69	0.033	0.013	2.839
	Baseline+TTA	0.006	74.6	38.04	12.57	0.026	0.008	2.119
	Baseline+TENT	0.000	106.3	37.65	4.99	0.014	0.003	0.718
	Baseline+DAE	0.027	76.2	98.18	3.81	0.155	0.055	7.805
	Baseline+PL	0.000	249.9	39.25	2.46	0.000	0.002	0.582
	Baseline+UPL	0.000	256.0	39.25	2.53	0.000	0.002	0.582
	Baseline+DPL	0.000	256.0	39.25	2.53	0.000	0.002	0.582
	Synth (Ours)	0.296	72.1	48.04	17.93	0.137	0.415	3.443
	Synth+TTA	0.314	71.4	41.15	9.07	0.213	0.412	2.967
	Synth+TENT	0.221	79.4	48.21	9.87	0.140	0.280	3.362
CT	Synth+DAE	0.304	63.6	43.34	8.82	0.199	0.359	3.258
	Synth+PL	0.279	84.3	81.32	20.01	0.156	0.519	5.573
	Synth+UPL	0.145	78.5	31.51	15.50	0.115	0.108	1.028
	Synth+DPL	0.243	80.1	25.71	13.82	0.146	0.224	1.514
	Baseline	0.029	70.9	28.01	8.54	0.084	0.032	1.492
	Baseline+TTA	0.017	125.0	27.63	4.11	0.076	0.011	0.593
	Baseline+TENT	0.000	207.7	28.74	2.59	0.018	0.000	0.550
	Baseline+DAE	0.064	78.2	78.65	2.98	0.198	0.106	5.426
	Baseline+PL	0.011	78.3	79.93	2010.46	0.002	0.032	6.333
	Baseline+UPL	0.000	256.0	28.87	2.84	0.000	0.000	0.542
	Baseline+DPL	0.000	256.0	28.87	2.84	0.000	0.000	0.542
	Synth (Ours)	0.234	66.8	19.71	5.43	0.233	0.204	0.965
	Synth+TTA	0.229	85.2	20.34	2.65	0.317	0.195	0.913
	Synth+TENT	0.222	58.6	20.82	9.17	0.139	0.210	1.245
	Synth+DAE	0.307	51.6	23.81	4.13	0.258	0.298	1.657
	Synth+PL	0.000	110.0	26.69	34.56	0.002	0.001	0.714
	Synth+UPL	0.000	255.0	28.86	2.84	0.000	0.000	0.543
	Synth+DPL	0.000	256.0	28.87	2.84	0.000	0.000	0.542

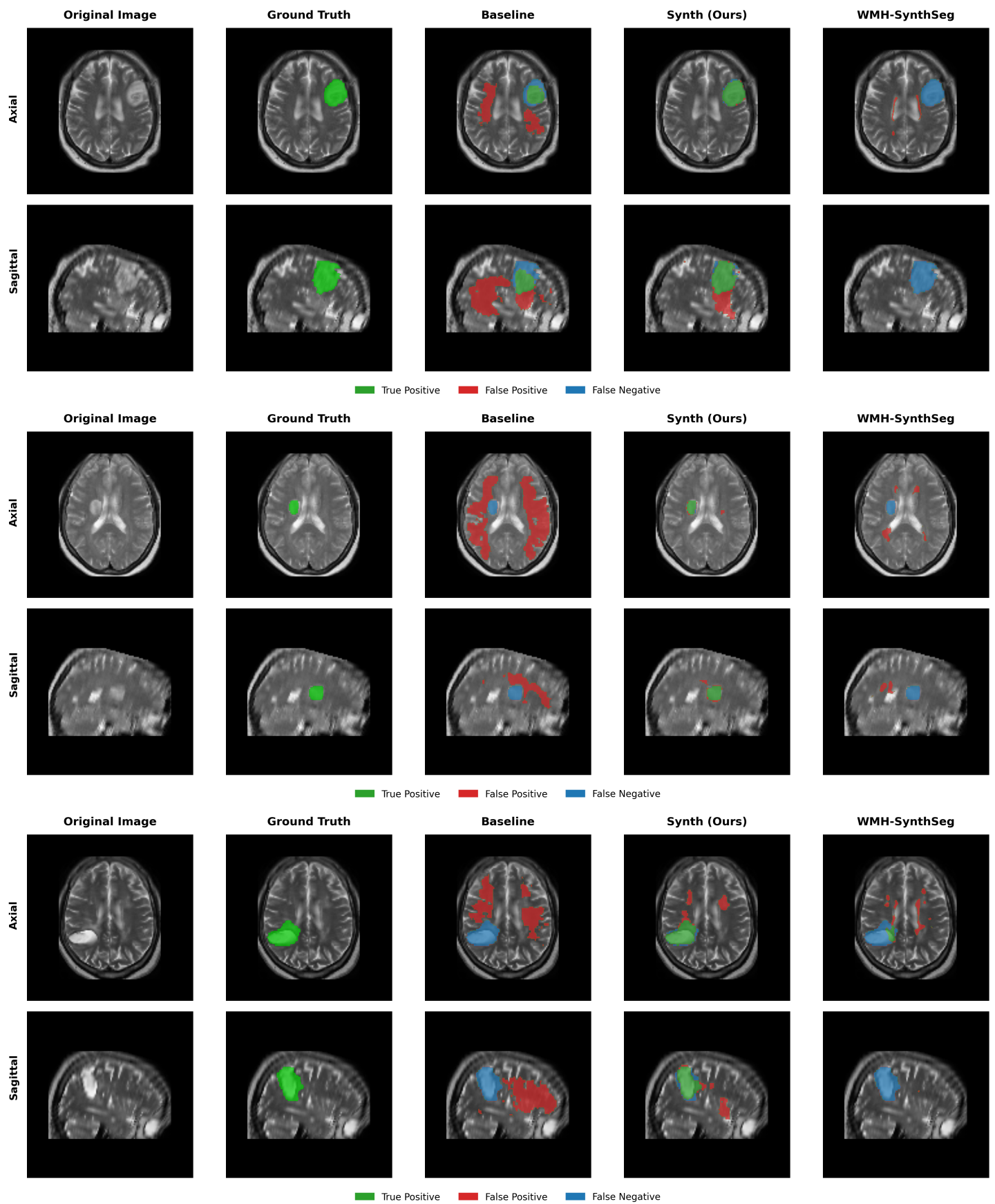


Figure 18: Sample visualisations of successful cases in the PLORAS T2w dataset. Green indicates a true positive prediction, red a false positive, and blue a false negative.

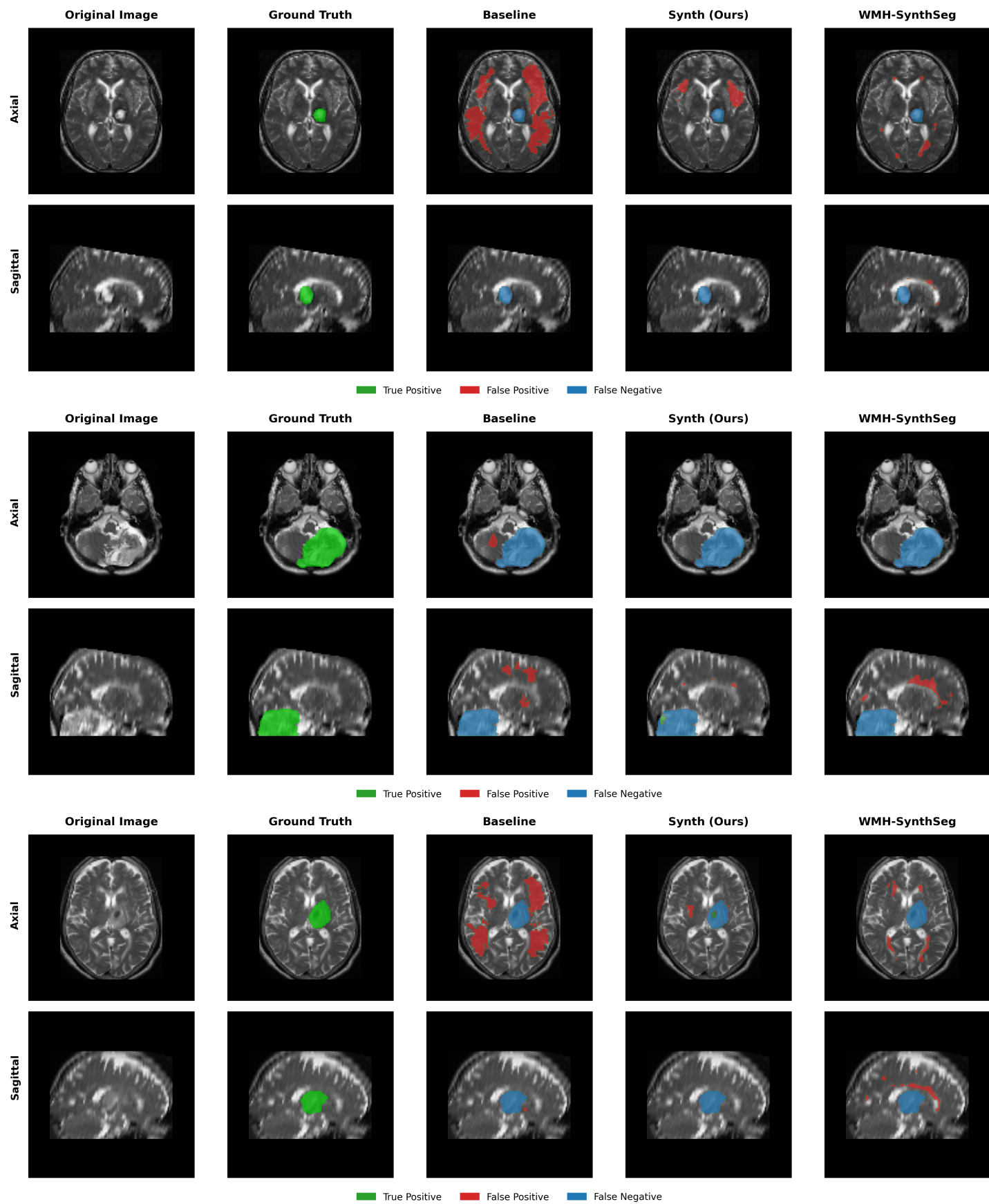


Figure 19: Sample visualisations of failure cases in the PLORAS T2w dataset. Green indicates a true positive prediction, red a false positive, and blue a false negative.

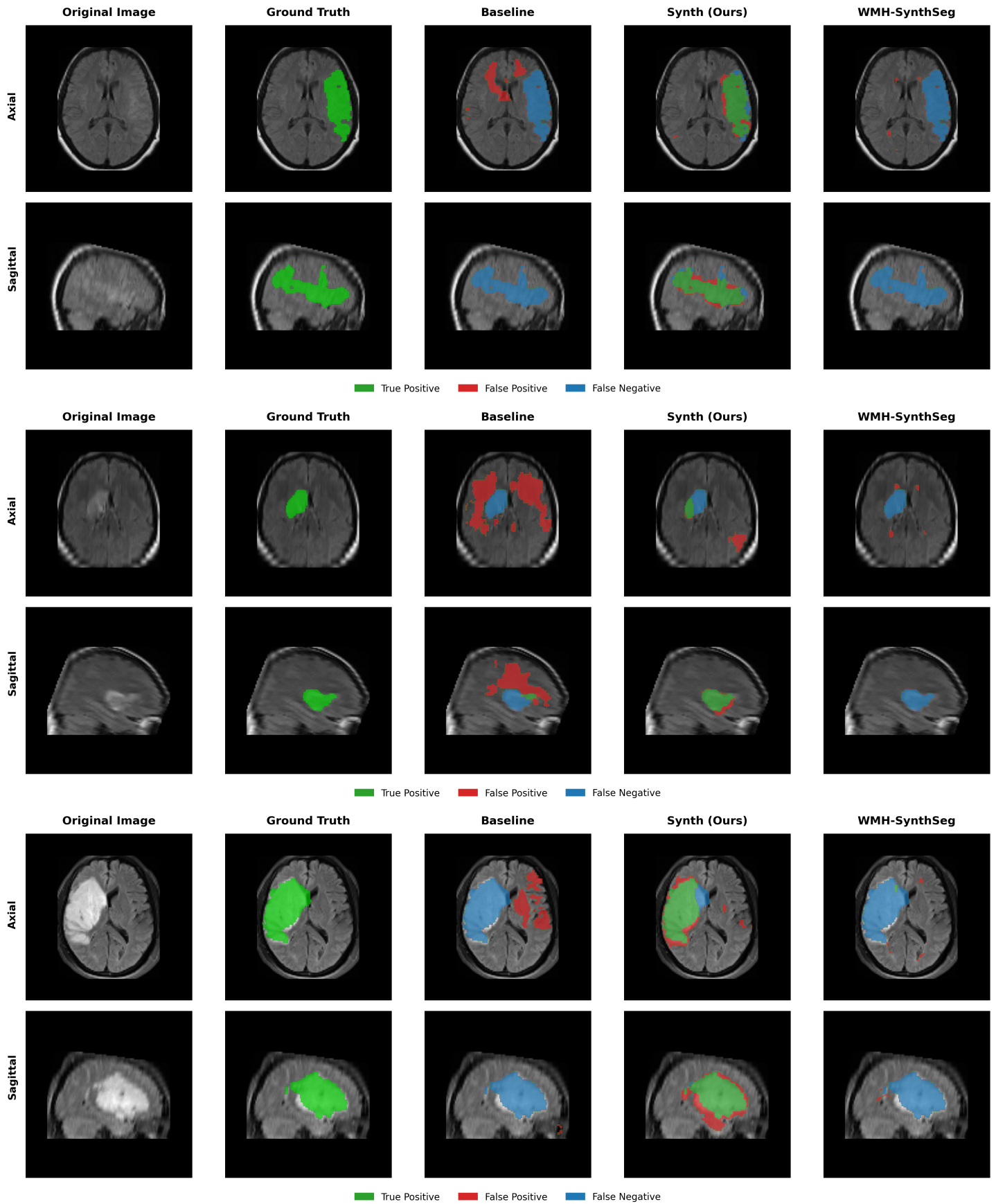


Figure 20: Sample visualisations of successful cases in the PLORAS FLAIR dataset. Green indicates a true positive prediction, red a false positive, and blue a false negative.

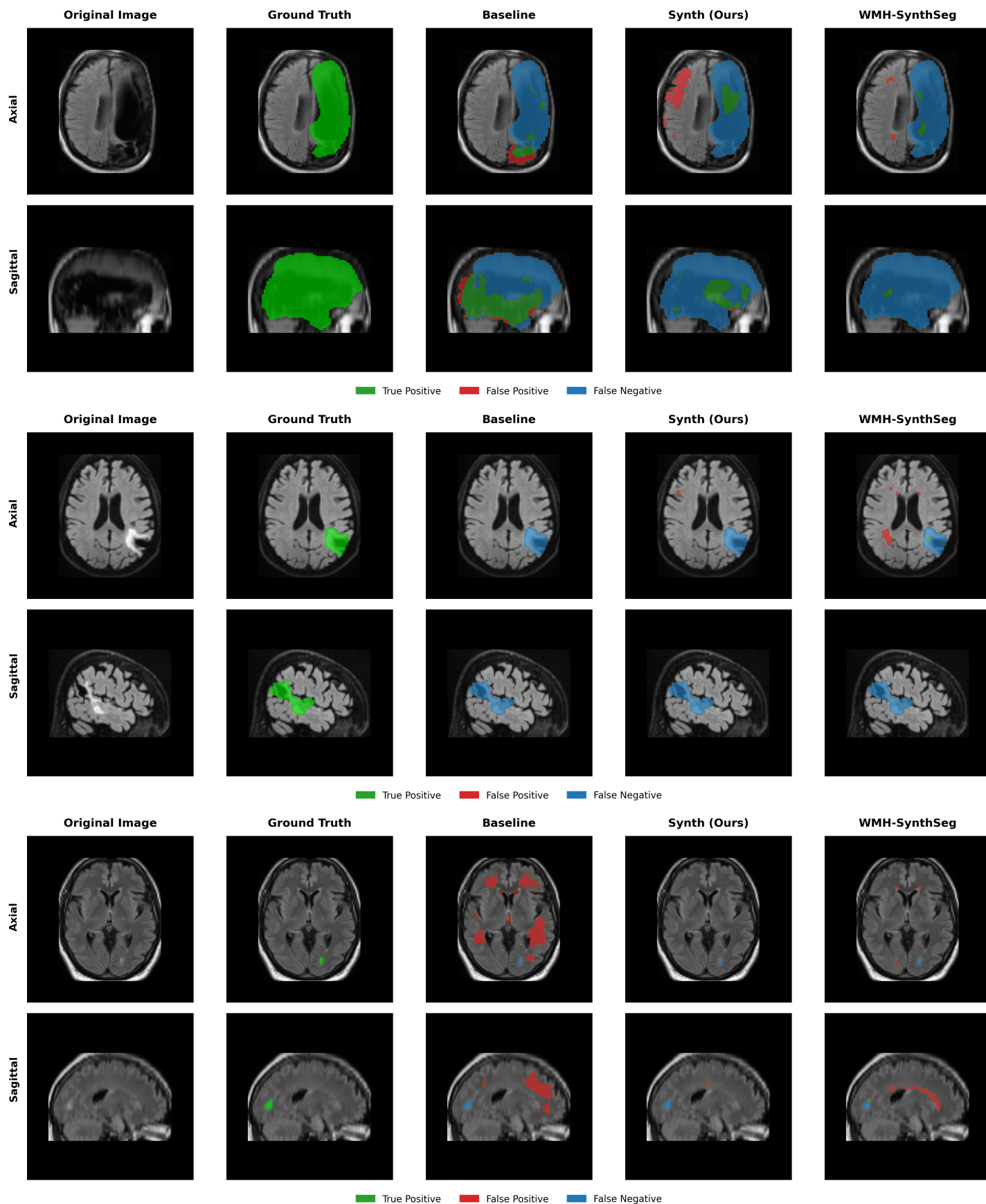


Figure 21: Sample visualisations of failure cases in the PLORAS FLAIR dataset. Green indicates a true positive prediction, red a false positive, and blue a false negative.

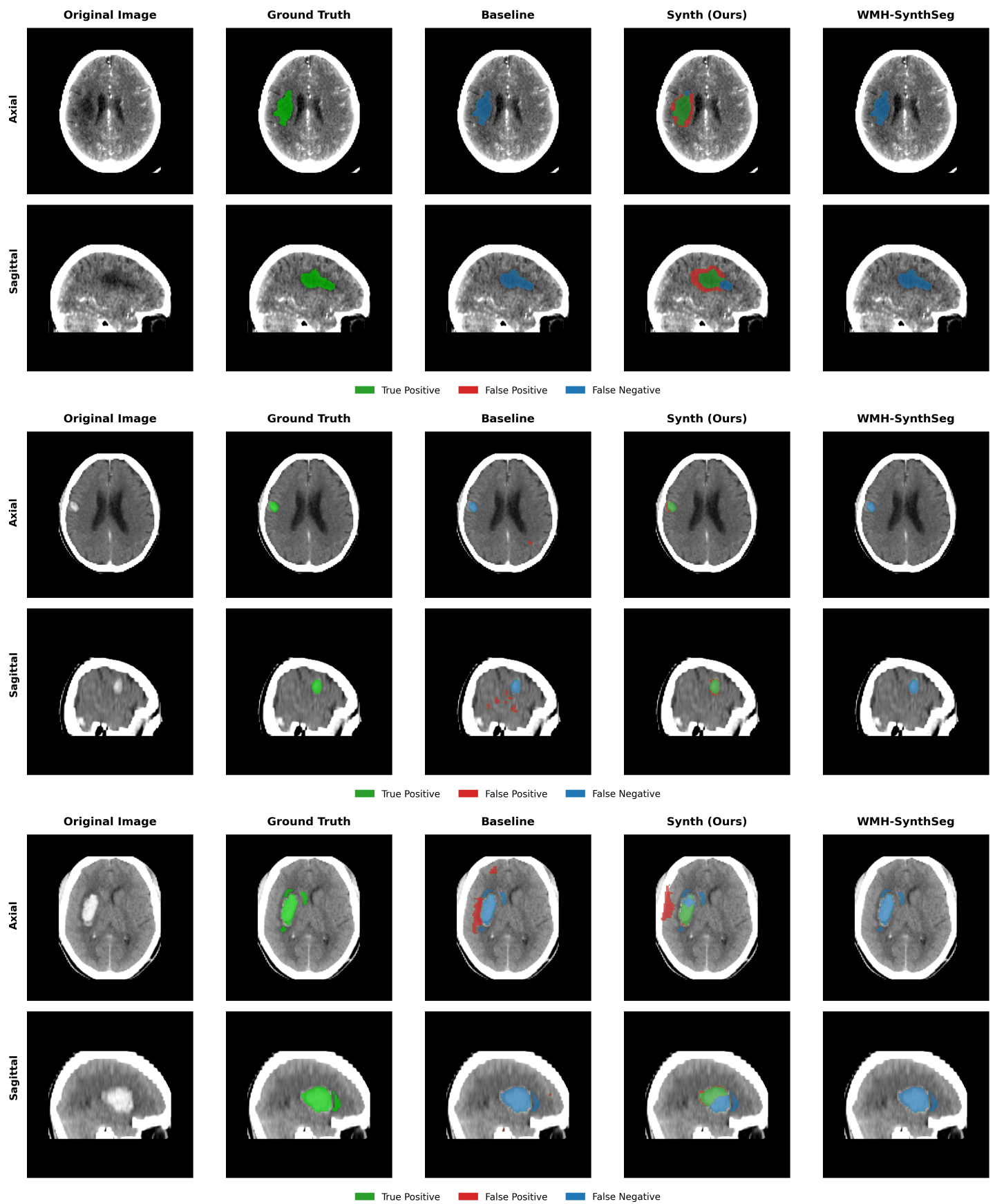


Figure 22: Sample visualisations of successful cases in the PLORAS CT dataset. Green indicates a true positive prediction, red a false positive, and blue a false negative.

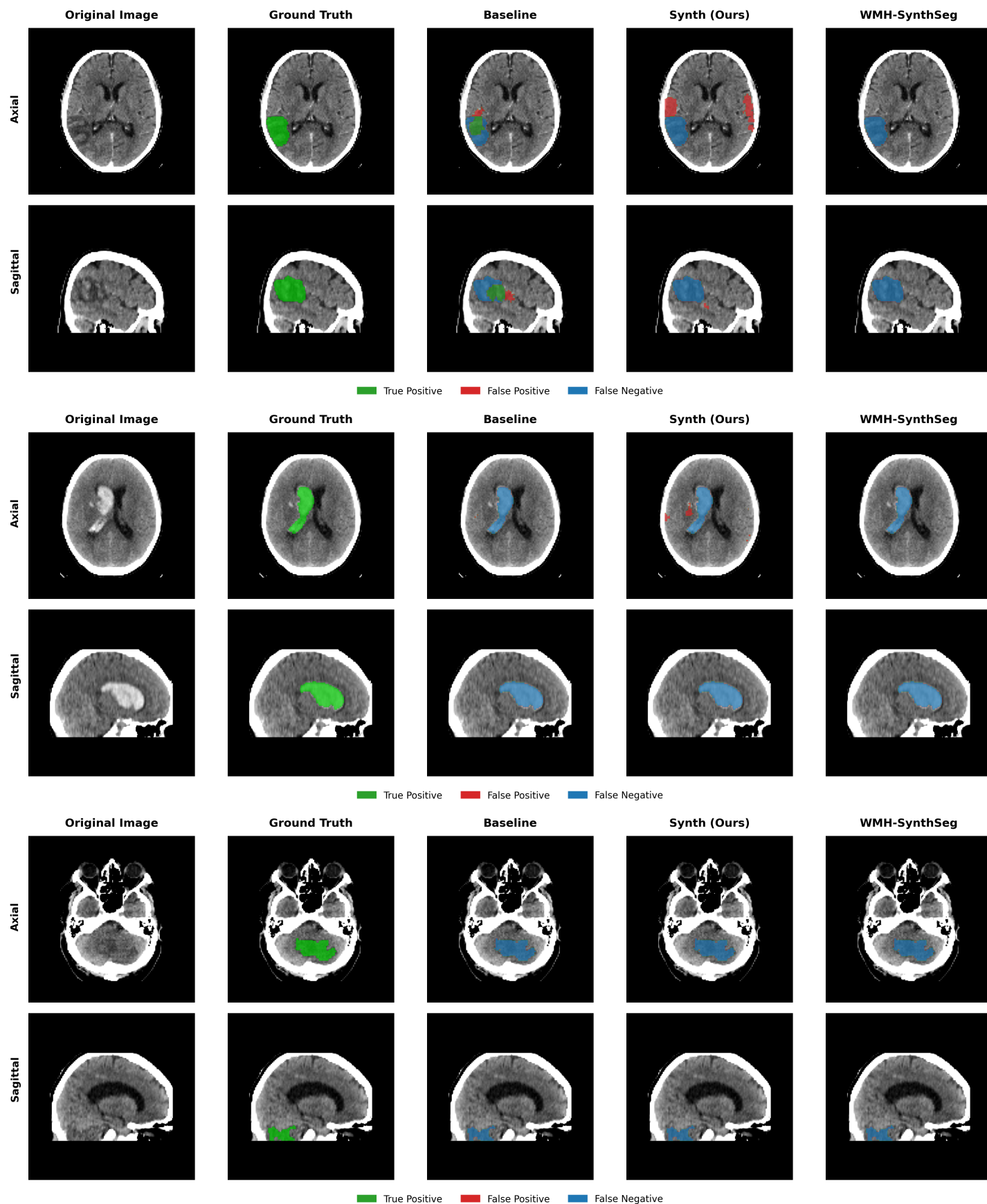


Figure 23: Sample visualisations of failure cases in the PLORAS CT dataset. Green indicates a true positive prediction, red a false positive, and blue a false negative.