

University of Bonn

MASTER'S THESIS FOR OBTAINING THE ACADEMIC DEGREE "MASTER OF SCIENCE (M.Sc.)"

Detection of Focal Cortical Dysplasia Type II Using Text Descriptions

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Submitted: December 20, 2025

Declaration of Authorship

I declare that the work presented here is original and the result of my own inves-
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has not been submitted, either in part or whole, for a degree at this or any other
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Abstract

Numerous methods have been developed for the detection of tumors in internal organs such as the lungs, brain, kidneys, and breast. However, detecting epileptogenic lesions remains significantly more challenging. Unlike tumors, these lesions do not typically increase in size over time, and there is a severe shortage of publicly available annotated datasets. As a result, researchers often need to contact hospitals and clinical centers directly to obtain even a minimal number of scans. Although recent studies have demonstrated that deep learning models can detect epileptogenic lesions, their performance remains limited, highlighting the ongoing difficulty of this task.

At the same time, recent work on tumor detection using text-guided approaches has shown promising results, demonstrating that incorporating textual descriptions can significantly improve segmentation accuracy. Inspired by these advances, this study proposes a new method that combines visual and textual features to enhance Focal Cortical Dysplasia (FCD) detection under limited data conditions. We further present a systematic comparison of multiple types of textual annotations and analyze their influence on model performance.

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1 Introduction

Automatic detection of tumors using medical imaging is widely studied in many internal organs. Recent advances in deep learning and transformer-based approaches have led to impressive results in the diagnosis of lung cancer [1], brain tumor [2], kidney Computed Tomography (CT) [3], and breast cancer [4]. These studies demonstrate the potential of modern computer vision techniques to achieve clinically significant results in various diagnostic tasks.

On the contrary, the detection of epileptogenic lesions, in particular FCD, remains a much more difficult task. Unlike tumors, such lesions usually do not change in size over time, and their inconspicuous appearance makes it difficult even for experienced neuroradiologists to identify them. In addition, the lack of large publicly available annotated datasets is hindering progress in this area. To address this problem, within the framework of the Multicentre Epilepsy Lesion Detection (MELD) project a large-scale collaborative dataset has recently been created and graph neural network—based approaches have been developed for epileptogenic lesion detection [5]. Although this is the most advanced solution to date, performance remains limited, underscoring the continued complexity of this task.

At the same time, text-guided and multimodal approaches have gained momentum in medical image analysis. Integrating textual prompts or language-guided embeddings with visual features has been shown to substantially enhance segmentation accuracy in chest X-ray infection detection [6], language-guided multi-level alignments [7], organ-aware segmentation [8], multimodal tumor analysis [9], and pneumothorax segmentation [10]. These advances demonstrate that textual annotations, ranging from atlas-based region names to descriptive clinical reports, provide valuable complementary information. By projecting textual and visual features into a shared latent space, such information can be effectively aligned and leveraged to enhance model performance.

We systematically evaluate the impact of different types of textual information, as well as alternative strategies for combining visual features, including the integration of an additional Graph Neural Network (GNN) blocks on top of MELD-based representations. We analyze how these design choices influence segmentation performance, demonstrating that the joint integration of language and visual features

can enhance both the accuracy and the sensitivity of epileptogenic lesion detection.

2 Related Works

This section reviews existing approaches to FCD detection and text-guided medical image segmentation for tumor detection tasks, focusing on their architecture, fusion strategies, and limitations.

Early work on FCD detection focused exclusively on vision-based methods [1–4]. It is noteworthy that one of the newest models, the MELD Graph [5], represents the surface of the cerebral cortex as a graph with multiple resolutions and applies a GNN-U-Net to segment lesions. It provides high sensitivity and specificity by identifying characteristic peaks (more than 20% in saliency) and determining calibration reliability using the expected calibration error. However, it has not been tested on patients with multiple FCDs, it lacks cross-attentional mechanisms for more complete integration of features, and it does not integrate textual clinical information or evaluate zero-shot generalization across FCD subtypes.

A second line of research explores language-guided segmentation by embedding text semantics into the segmentation pipeline. Early methods [11, 12] simply tokenized text and merged embeddings with image features via attention, but struggled to capture high-level semantics. More recent approaches [6, 13] employ deep pretrained text encoders (e.g., CXR-BERT [14]) fused with ConvNeXt-Tiny [15] image features. In [7], the authors introduce two novel modules: the Target-sensitive Semantic Distance Module (TSDM), which computes contrastive distances between segmentation masks to focus on disease-related regions, and the Language-guided Target Enhancement Module (LTEM), which applies crossattention to reinforce critical image areas. A bi-directional contrastive loss (averaged cross-entropy in both image \rightarrow text and text \rightarrow image directions) yields more fine-grained guidance and enables models trained on 25% of the data to outperform single-modal baselines trained on full datasets. However, these improvements come at the cost of higher architectural complexity and a strong reliance on highquality textual annotations, which may limit robustness and generalizability in real-world clinical settings.

To address unpaired multi-modal data, MulModSeg [9] conditions text embeddings on imaging modality using frozen CLIP [16]/BioBERT [17]/Med-CLIP [18] encoders combined with medical prompts, and alternates training between vision and text branches (3D-UNet[19] or SwinUNETR [20]). This scheme improves

generalization without requiring paired CT/MR scans. Experiments show that varying the CT:MR ratio in training data shifts performance, underscoring the importance of balanced modality representation. However, effectiveness depends critically on prompt template design and alternating-training convergence.

Finally, weakly supervised methods such as SimTxtSeg [21] leverage simple text cues (e.g., "lesion in the left hemisphere") to guide segmentation without requiring pixel-level annotations. SimTxtSeg was evaluated using standard segmentation metrics, including Dice similarity coefficient (DSC), Intersection over Union (IoU), Positive Predictive Value (PPV), Normalized Surface Dice (NSD), and the 95% Hausdorff Distance (HD95), alongside cross-entropy loss. These results demonstrate that even coarse textual hints can substantially improve mask quality over vision-only baselines. However, such approaches typically rely on consistent, high-quality text labels and have not yet been validated on rare pathologies such as FCD.

Building on these insights, we draw inspiration from Ariadne's Thread [6], which employs lightweight text prompts and a GuideDecoder to segment infected regions in chest X-rays. Remarkably, this method achieves over a 6% Dice improvement compared to unimodal baselines while using only 10% of the training set, high-lighting the potential of multimodal prompting in low-data regimes. For our FCD task, we adopt the MELD Graph model as a pretrained backbone, since it was trained on the largest dataset among the methods surveyed, providing a robust basis for feature extraction and downstream adaptation.

To the best of our knowledge, no prior study has applied text-guided segmentation methods to the FCD detection task, underscoring the novelty of our approach. In summary, while graph-based GNNs excel at modeling cortical geometry and text-guided methods enrich segmentation with semantic context, no existing approach simultaneously addresses surface-space lesion detection, integration of fine-grained clinical narratives, robustness to scanner-induced domain shifts, and zero-shot generalization to unseen FCD subtypes. In this work, we aim to demonstrate that incorporating textual information is also beneficial for the FCD detection task. To this end, we integrate different types of text descriptions derived from anatomical atlases with vision-based features into a multi-resolution GNN framework, using cross-attention fusion and contrastive alignment to enhance robustness and detection accuracy across heterogeneous datasets.

3 Method

As mentioned earlier, the design of our architecture was inspired by the work of [6]. The key differences in our implementation are threefold: (1) we increase the number of feature extraction layers, (2) we introduce a GNN-based block to better aggregate visual features at the top layers, and (3) we disable the text branch in the final decoder layers to reduce overfitting. The overall architecture is illustrated in Figure 3.1, and each component is described in more detail below.

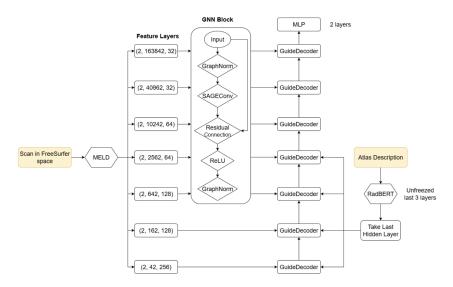


Figure 3.1: Overview of the proposed multimodal segmentation architecture. Visual features extracted from FreeSurfer surface space are processed through hierarchical GNN layers, while textual embeddings from RadBERT guide the decoder via cross-attention.

3.1 Visual Feature Extraction

As input to the vision model, structural MRI scans are first mapped into the FreeSurfer surface space. From the MELD preprocessing pipeline, we obtain a multi-resolution set of surface-based features across seven hierarchical levels. Each level is represented as a tensor of shape (H, N, C), where:

- H number of hemispheres (left/right),
- \bullet N number of vertices on the cortical mesh per hemisphere,
- \bullet C number of features per vertex.

To aggregate higher-order geometric information, we process the **five top feature layers** individually through a dedicated **GNN Block**. This choice is motivated by empirical findings (see Chapter 5), where including the lowest two layers led to oversmoothing and diluted discriminative information, while the top five layers yielded the best trade-off between expressivity and stability.

GNN Block. Formally, let

$$\mathbf{X}^{(l)} \in \mathbb{R}^{(B \cdot H \cdot N_l) \times C_l}$$

denote the input feature matrix at layer l, where B is the batch size, H is the number of hemispheres, N_l is the number of vertices per hemisphere at layer l, and C_l is the number of input channels.

Each GNN Block then applies the following sequence of operations:

$$\mathbf{H}_0 = \operatorname{GraphNorm}(\mathbf{X}^{(l)}, \mathtt{batch}), \qquad (3.1)$$

$$\mathbf{H}_1 = \mathrm{SAGEConv}(\mathbf{H}_0, \mathtt{edge_index}_l),$$
 (3.2)

$$\mathbf{H}_2 = \mathbf{H}_1 + \mathbf{H}_0$$
 (residual connection), (3.3)

$$\mathbf{H}_3 = \text{ReLU}(\mathbf{H}_2), \tag{3.4}$$

$$\mathbf{H}_4 = \text{Dropout}(\mathbf{H}_3), \tag{3.5}$$

$$\mathbf{Z}^{(l)} = \operatorname{GraphNorm}(\mathbf{H}_4, \mathtt{batch}). \tag{3.6}$$

where:

- $\mathbf{X}^{(l)} \in \mathbb{R}^{(B \cdot H \cdot N_l) \times C_l}$ input feature matrix at level l;
- edge_index_l $\in \mathbb{N}^{2 \times |E_l|}$ adjacency structure of the cortical surface graph;
- batch batch vector;
- $\mathbf{Z}^{(l)} \in \mathbb{R}^{(B \cdot H \cdot N_l) \times C_l}$ resulting representation for level l;

• SAGEConv implements the GraphSAGE [22] update rule

$$\mathbf{x}_i' = \mathbf{W}_1 \mathbf{x}_i + \mathbf{W}_2 \cdot \underset{j \in \mathcal{N}(i)}{\text{mean }} \mathbf{x}_j,$$

where $\mathcal{N}(i)$ is the neighborhood of node i;

- **GraphNorm** $(x) = \gamma \cdot \frac{x \mu_g}{\sqrt{\sigma_g^2 + \epsilon}} + \beta$, where μ_g, σ_g^2 are mean and variance per graph, and γ, β are learnable parameters;
- $\mathbf{ReLU}(x) = \max(0, x)$ rectified linear activation function;
- **Dropout**(x) randomly sets a subset of elements in x to zero with probability p.

After passing through the GNN Block, the feature dimensionality remains unchanged, ensuring consistency with subsequent blocks. The resulting features are then passed to the GuideDecoder for multimodal fusion.

3.2 Textual Feature Extraction

The text branch of our architecture leverages the pretrained **RadBERT** model [23], which is specifically designed for radiology reports. Prior studies have demonstrated that selectively unfreezing the last few layers of transformer models can improve downstream performance compared to freezing the entire model [24]. Motivated by this insight, we unfreeze the last 3 hidden layers of RadBERT—this specific choice is justified in Chapter 5. The output of the (now trainable) final hidden layer is then forwarded to each GuideDecoder for multimodal fusion, except for the topmost three layers.

3.3 GuideDecoder

We adopt the GuideDecoder architecture proposed by Zhong et al. [6], which fuses visual and textual features in a multi-modal fashion. The decoder first projects the textual tokens to align their dimensionality with that of the visual tokens, then applies multi-head self-attention and cross-attention to exchange semantic information across modalities. Finally, the fused features are upsampled and combined with skip connections from the visual encoder at the same resolution before the final prediction.

In our implementation, the overall design of the GuideDecoder is preserved, but we introduce two modifications to adapt the architecture to surface—based representations. First, we replace the standard 2D upsampling with the HexUnpool operator (see Appendix HexUnpool), which performs mean unpooling on the icosphere mesh. Second, after upsampling and fusion with skip connections, the features are processed by a convolutional block based on DynUNetBlock (Code link) from the MONAI framework, which applies a sequence of convolution, normalization, and activation layers. In our architecture, we replace this module with a custom SpiralConv-based block that operates on the surface mesh representation of the cortex.

4 Dataset

We gratefully acknowledge the **MELD Project** for providing access to the dataset used in this work. Without this resource, it would not have been possible to conduct a systematic evaluation of our model.

4.1 Bonn Dataset

The Bonn scientific group has published a publicly available presurgical MRI dataset on **OpenNeuro**, entitled "An open presurgery MRI dataset of people with epilepsy and focal cortical dysplasia type II" [25]. This dataset comprises 170 participants, including 85 individuals with FCD type II and 85 healthy controls. For each participant, the following data are available:

- High-resolution **3D T1-weighted** MRI scans (isotropic voxels, 1 mm³ or 0.8 mm³ resolution, depending on the subject);
- Corresponding **isotropic 3D FLAIR** imaging, available for most participants;
- Manually delineated regions of interest (ROIs) identifying FCD lesions, provided for patients only;
- A set of clinical and demographic variables (including age, sex, lesion laterality and location, histopathological subtype IIa/IIb, MRI-negative status, and postoperative outcome according to Engel classification).

Dataset Organization

The dataset follows the BIDS standard. Each subject folder contains an anat subfolder with NIfTI files and corresponding JSON metadata. For participants with FCD, lesion ROI masks are also provided. Sequence acquisition parameters are documented in the JSON sidecar files.

4.2 MELD Dataset

The MELD Project provides a large-scale neuroimaging dataset comprising Magnetic Resonance Imaging (MRI) scans and clinical data of patients with FCD, as well as healthy controls. In total, the dataset includes 1185 participants from 23 international epilepsy surgery centers. Due to missing or corrupted files in the obtained release, we used a subset of 960 participants in our experiments. The dataset is not publicly available; access must be requested directly from the study authors [5].

For each participant, the dataset includes:

- Structural MRI: 3D T1-weighted (all participants) and, when available,
 3D FLAIR;
- Lesion annotations: manually delineated Region of InterestRegions of Interest (ROIs) identifying FCD lesions (patients only). For MRI-negative cases, postsurgical resection cavities were used to guide ROI definition;
- Demographic and clinical metadata;
- Predefined splits for training, validation, and test cohorts.

4.3 FreeSurfer Processing

Each image was processed with the FreeSurfer framework [26], from which **11 core** surface-based features were extracted:

- Morphometric features: cortical thickness, sulcal depth, curvature, and intrinsic curvature;
- Intensity features: gray—white matter intensity contrast, and FLAIR intensity sampled at 6 intracortical and subcortical depths.

In addition to the raw measurements, features were further processed into raw values, control-normalized features, and asymmetry features (left vs. right hemisphere). Cortical thickness was additionally adjusted by regressing out curvature. Altogether, this yielded 34 input features per participant, computed at 163,842 vertices on a bilaterally symmetrical cortical surface template [5].

Since MRI features vary across scanners and sites, potentially impairing generalization, an **intersite harmonization** procedure was applied using the distributed **ComBat** algorithm, a well-established statistical method for removing scanner/site effects [5].

For conversion from volume space to FreeSurfer surface space, detailed instructions are available in the MELD documentation [27]. In practice, however, it is often more practical to request preprocessed files directly from the authors, since converting a single image to surface space is computationally expensive: approximately 6–7 hours per scan (even with FastSurfer [28] on an NVIDIA A100 GPU, the process required 3–4 hours).

4.4 Data Augmentation

In the original study MELD authors applied augmentations in three stages:

- Lesion augmentation modifying the lesion mask to create new training samples;
- Mesh-space transforms applying geometric transformations to the cortical surface mesh;
- Intensity transforms modifying the intensity features at each vertex.

Each transform is applied independently with probability p defined in the experiment configuration. The order is fixed: lesion augmentation \rightarrow mesh transforms \rightarrow intensity transforms.

Lesion–mask augmentation. Given a geodesic distance map D on the cortical surface (negative inside the lesion), we first normalise it by $|\min D|$ and add low–frequency noise, which is generated on a low–resolution icosphere (level 2) and then upsampled to the target resolution using the predefined unpool operators:

$$\tilde{D} \; = \; \mathtt{Unpool}\Big(\tfrac{D}{|\min D|} \; + \; \mathcal{N}(0,\sigma^2) \Big) \,, \qquad L' \; = \; \mathbb{1}\{\tilde{D} \leq 0\}.$$

In our implementation, we use $\sigma = 0.5$ by default. And as Unpool we use the HexUnpool operator (see Appendix HexUnpool). After modifying the binary lesion mask L, the geodesic distances and smoothed labels are **recomputed**:

$$D' = \texttt{fast_geodesics}(L'), \qquad \widetilde{L} = \texttt{smoothing}(L', \text{ iteration} = 10).$$

This procedure is applied only if the lesion mask is non–empty. In our configurations, however, the recomputed distances and smoothed labels were not used further.

Mesh-space transforms (icosphere re-indexing). We use precomputed index maps (provided by the MELD authors in their GitHub repository) and apply them once per sample to all vertex-wise tensors (features, labels, distances, etc.). The MELD framework defines three types of such maps:

- **Spinning** index remapping that corresponds to a rigid rotation of the icosphere.
- Warping smooth non-rigid remapping that locally compresses or stretches the mesh.
- Flipping mirror reflection of the mesh, e.g. exchanging left and right hemispheres. *Note:* because SpiralConv is order—sensitive, flipping requires reversing the spiral neighbour order; in our final experiments we set p(flip)=0 to avoid this mismatch.

Per-vertex intensity transforms. All intensity transforms act channel-wise on features:

- Gaussian noise: add $\epsilon \sim \mathcal{N}(0, \sigma^2)$ with $\sigma^2 \sim \mathcal{U}(0, 0.1)$.
- Brightness scaling: multiply each channel by $m_c \sim \mathcal{U}(0.75, 1.25)$.
- Contrast adjustment: for each channel c, sample $f \sim \mathcal{U}(0.65, 1.5)$ and apply

$$x_c \leftarrow (x_c - \mu_c) f + \mu_c,$$

$$min_c = \min(x_c), \quad max_c = \max(x_c),$$

$$x_c[x_c < min_c] = min_c, \quad x_c[x_c > max_c] = max_c.$$

• Gamma: for $\gamma \sim \mathcal{U}(0.7, 1.5)$,

$$x' \leftarrow \left(\frac{x - \min(x)}{\max(x) - \min(x) + \varepsilon}\right)^{\gamma} (\max(x) - \min(x) + \varepsilon) + \min(x);$$
$$x'' \leftarrow \frac{x' - \mu}{\sigma + \varepsilon}, \quad \text{where } \varepsilon = 10^{-7}, \ \mu = \text{mean}(x), \ \sigma = \text{std}(x).$$

We also use an *inverted* variant by applying the same operation to -x and negating back.

Disabled/placeholder transforms. The current implementation contains placeholders for Gaussian blur and low-resolution downsampling. These operations are not implemented in the original authors' code and were not used in our reported results.

4.5 Types of Atlas Descriptions

We generated textual descriptions using AtlasReader [29], which registers the input (statistical) MRI map to a template and, for user-specified atlases, produces region names and coordinate tables. In this work we used the Harvard–Oxford (cortical and subcortical) probabilistic atlas [30] and the AAL atlas [31].

However, even strong validation results with full descriptions do not guarantee robustness in real-world settings: in practice, exact overlap percentages and full region names are rarely available. Therefore, in chapter 5 we additionally evaluate reduced-text settings: (i) hemisphere only (Left/Right), (ii) gross lobar labels only (e.g., Frontal Lobe, Temporal Lobe, Parietal lobe . . .), and (iii) their combination.

5 Experiments

We investigate three factors: (i) how many MELD feature *stages* are fed into the GNN block; (ii) how many hidden layers from the text encoder are injected into the GuideDecoder; and (iii) the effect of including textual descriptions.

We freeze the MELD backbone. In all variants, we *train* the same geometry-based upsampling path (HexUnpool + SpiralConv) and the segmentation head. The variants differ only in whether and how the GuideDecoder is used before upsampling:

- 1. **Unpool+Spiral (no text).** No GuideDecoder is used; features are fed directly into the upsampling path (HexUnpool + SpiralConv) and the segmentation head; no textual input.
- 2. **GuideDecoder** (self-attention only). A GuideDecoder layer is inserted before upsampling, but the text branch is disabled, so only self-attention operates; the same Unpool+Spiral path and segmentation head are trained.
- 3. **GuideDecoder** + **Text** (full). The full GuideDecoder is used (self-attention plus cross-attention to the text encoder); textual descriptions are provided, while the same Unpool+Spiral path and segmentation head are trained.

5.1 Connecting MELD Feature Stages into the GNN Block

TODO

5.2 Text-Decoder Connections (Layer Injection)

TODO

5.3 Effect of Textual Descriptions

TODO

6 Discussion

7 Conclusion

Appendix

HexUnpool We implement surface upsampling with a custom HexUnpool operator. Given features $X \in \mathbb{R}^{B \times H \times N_{\text{from}} \times C}$ on a coarse icosphere with N_{from} vertices, HexUnpool produces $X' \in \mathbb{R}^{B \times H \times N_{\text{to}} \times C}$ on a denser icosphere $(N_{\text{to}} > N_{\text{from}})$. The operation works in two steps:

- 1. Existing vertices are copied to the corresponding positions in X'.
- 2. Each new vertex is assigned the mean of its parent vertices from the coarse level, according to predefined upsampling indices \mathcal{I} .

Formally, for each new vertex v, we compute

$$X'_v = \frac{1}{|\mathcal{I}(v)|} \sum_{u \in \mathcal{I}(v)} X_u.$$

List of Acronyms

CT Computed Tomography

FCD Focal Cortical Dysplasia

GNN Graph Neural Network

MELD Multicentre Epilepsy Lesion Detection

MRI Magnetic Resonance Imaging

ROI Region of Interest

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