

# SemantiBench: Measuring Prompt Compliance in Medical Image Segmentation with Frequency-Gated Models

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**Abstract.** Foundation models like BiomedParse and SAM have improved biomedical segmentation with text-to-mask capabilities, but they often fail when prompts shift from simple anatomical targets (e.g., “kidney”) to fine-grained pathological descriptions (e.g., “necrotic tumor core”). We refer to this degradation as *semantic collapse*. We identify that this stems from CLIP’s inability to represent negation geometrically—prompts like “excluding tumor” still activate tumor-related features. In this paper, we introduce **SemantiBench**, a dataset of 100,000+ prompt-mask pairs, and propose **FreqMedCLIP**, a segmentation model that explicitly separates target and avoidance signals. Unlike standard methods, our model decouples prompts into Target and Avoidance streams and enforces constraints via a **logical gate**. We also introduce an **exclusion loss** that supervises the gate to act as a precise detector for forbidden regions. Our model achieves a Dice score of 0.77 on complex exclusionary queries ( $L_3$ ), significantly outperforming the baseline (Dice 0.60). Current models fail this test; they ignore “excluding” and segment everything.

**Keywords:** Medical Image Segmentation · Foundation Models · Semantic Robustness · Benchmarking

## 1 Introduction

Foundation models such as BiomedParse [15] and MedSAM [10] can now segment biomedical objects using natural language. However, strict adherence to clinical prompts remains a problem. While these models succeed at atomic queries ( $L_1$ ) like “kidney”, they often fail to comply with complex constraints ( $L_3$ ) such as “kidney excluding the renal pelvis”.

This failure often stems from “semantic collapse”, where the model ignores key logical or descriptive modifiers and incorrectly defaults to the generic object definition. For example, when prompted with “necrotic tumor core”, many state-of-the-art models ignore the adjective “necrotic” and segment the entire tumor. This behavior is dangerous in clinical settings where precise sub-region targeting is required.

This paper introduces three contributions to address this problem. First, we propose **SemantiBench**, a protocol to measure two distinct forms of robustness: (1) **Descriptive Invariance** ( $L_2$ ), which checks if models output consistent masks for synonymous prompts (e.g., “kidney” vs. “bean-shaped organ”); and (2) **Logical Compliance** ( $L_3$ ), which tests if models correctly modify the segmentation mask when prompts impose exclusionary constraints.

To quantify these failures, we introduce the **Prompt Sensitivity Score (PSS)**. We find that current models exhibit a PSS gap of up to 0.29, indicating poor compliance with complex instructions.

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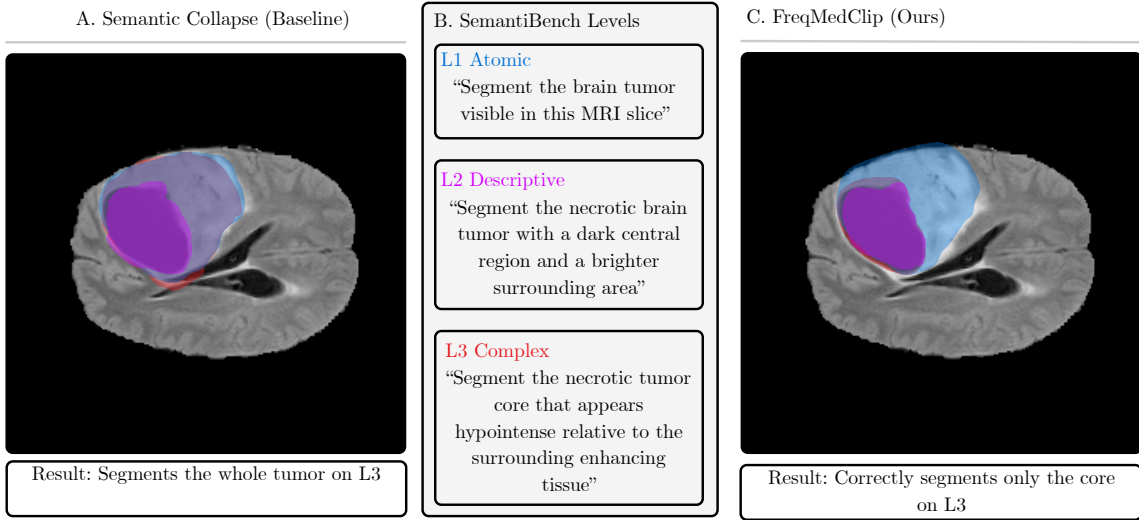


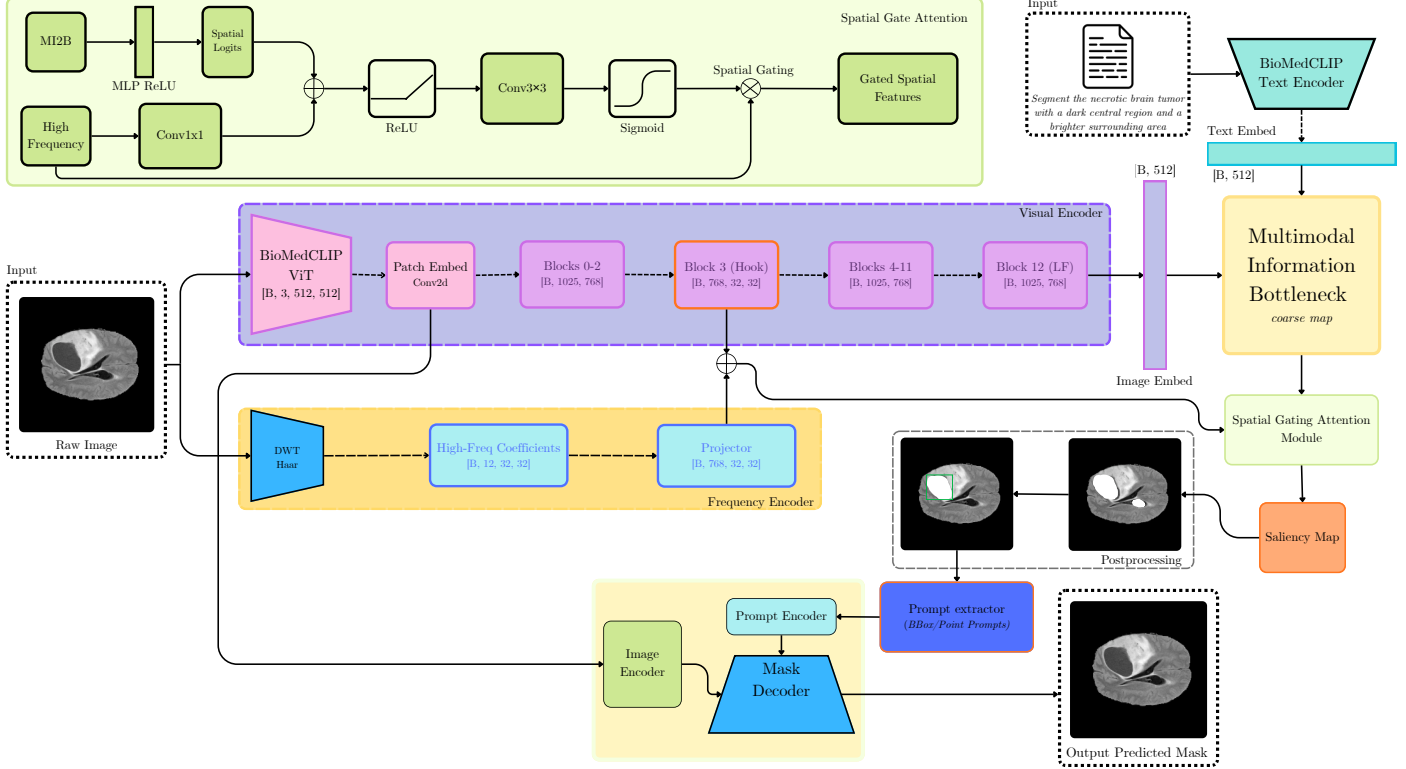
Fig. 1: **Semantic failure modes.** (A) Existing foundation models (BiomedParse) fail to process the adjective “necrotic,” incorrectly segmenting the entire tumor mass [15]. (B) SemantiBench evaluates robustness across three linguistic levels ( $L_1 - L_3$ ). (C) FreqMedClip uses cross-modal gating to isolate the necrotic core.

Finally, we present **FreqMedCLIP**, an architecture designed to solve this ambiguity. We argue that CLIP embeddings suffer from what we call “affirmative bias”—the encoder treats all words in a prompt as inclusion signals, even negation operators like “excluding.” FreqMedCLIP addresses this by decoupling the text into Target and Avoidance streams. We introduce a **logical gating** mechanism that forces the exclusion through multiplicative gating, suppressing the forbidden region at the feature level. This approach, combined with explicit exclusion loss, reduces the PSS to 0.12, significantly improving prompt compliance.

## 2 Related Work

Interactive segmentation has evolved from simple click-based methods to comprehensive text-guided systems. The release of IMIS-Bench [1] provided a significant resource (IMed-361M) for training interactive models. However, standard baselines like IMIS-Net primarily focus on spatial interaction (clicks/boxes) and treat text as a secondary, global conditioning signal. Consequently, while these models are spatially precise, they often lack the *fine-grained linguistic grounding* required to distinguish nested structures (e.g., edema vs. core) based on text alone. Our work elevates textual semantics to a primary spatial constraint.

The field has seen a surge in foundation models adapted for medicine. MedSAM [10] and MedClipSamV2 [?] fine-tune the Segment Anything Model (SAM) on medical data but rely heavily on box/point prompts, limiting their utility for semantic parsing. BiomedParse [15] represents the current state-of-the-art in joint segmentation and recognition, utilizing GPT-4 to harmonize ontologies. While BiomedParse excels at object recognition (valid vs. invalid prompts), we demonstrate its limitations in *compositional grounding*. Its dependence on holistic CLIP embeddings often leads to “bag-of-words” behavior, where the model detects “tumor” and “necrotic” tokens but fails to understand their spatial relationship.



**Fig. 2: FreqMedClip Architecture.** The model features two streams: (1) A visual backbone (BiomedCLIP) to extract global semantic context, and (2) A High-Frequency Encoder (Wavelet-based) to capture texture and boundary details associated with specific attributes (e.g., “necrotic,” “spiculated”). The Cross-Modal Semantic Gating (CMSG) mechanism uses the text prompt to actively filter the high-frequency stream, isolating only the relevant sub-structures.

Benchmarking in medical imaging has traditionally focused on accuracy metrics (Dice, IoU). Recent frameworks like FairMedFM [5] have expanded this to include *Fairness*, evaluating performance disparities across demographic groups (sex, age, race). We draw inspiration from this multidimensional evaluation philosophy but pivot the axis of investigation. Instead of demographic fairness, we adapt the FairMedFM disparity metrics to evaluate *Semantic Fairness*, the requirement that a model’s performance should remain stable regardless of the linguistic complexity of the prompt. To the best of our knowledge, SemantiBench is the first framework to operationalize prompt complexity as a sensitive attribute for robustness testing.

### 3 Methodology

We designed FreqMedCLIP to solve the “Frequency-Semantic Gap.” We hypothesize that semantic attributes (like “necrotic” or “nodular”) are often encoded in high-frequency signals that standard Vision Transformers smooth out. Therefore, our model treats the high-frequency stream not just as a boundary detector, but as the raw material for semantic reasoning.

*3.1. High-Frequency Attribute Encoder.* Standard ViTs suffer from a “low-pass” bias, making them blind to texture-defined semantics [13]. Recent work has shown that explicit frequency modeling significantly boosts segmentation of subtle structures [7,4]. We address this by explicitly encoding the high-frequency spectrum. We use a **Discrete Wavelet Transform (DWT)** to decompose the input  $\mathbf{X}$  into four spectral sub-bands:

$$\mathbf{X}_{freq} = \text{DWT}(\mathbf{X}) = \{\mathbf{X}_{LL}, \mathbf{X}_{LH}, \mathbf{X}_{HL}, \mathbf{X}_{HH}\} \quad (1)$$

Here,  $\{\mathbf{X}_{LH}, \mathbf{X}_{HL}, \mathbf{X}_{HH}\}$  capture the vertical, horizontal, and diagonal high-frequency details—precisely where features like tissue texture (necrotic vs. vital) and margin type (spiculated vs. smooth) reside. These 12 spectral channels feed a **ConvNeXT-Tiny** encoder, which provides the “semantic raw material” that the text gate will later act upon.

*3.2. Dual-Stream Encoder.* We use the **BiomedCLIP ViT-B/16** to extract the global semantic context. To handle compositional queries (e.g., “Kidney excluding tumor”), we use a semantic decomposition strategy. The input prompt is split into  $P_{pos}$  (“Kidney”) and  $P_{neg}$  (“Tumor”). These are encoded into two distinct embedding vectors,  $\mathbf{E}_{pos}$  and  $\mathbf{E}_{neg}$ , ensuring that the avoidance signal remains independent from the target representation.

*3.3. Logical Gating.* We propose a **Logical Gate**. Unlike standard cross-attention which mixes signals, this gate operates on strict exclusion logic. The high-frequency feature map  $\mathbf{F}_{freq}$  is processed by two parallel paths:

- **Inclusion Path ( $\mathbf{G}_{inc}$ ):** Activates regions matching  $\mathbf{E}_{pos}$ .
- **Exclusion Path ( $\mathbf{G}_{exc}$ ):** Activates regions matching  $\mathbf{E}_{neg}$ .

The final feature refinement enforces the logical constraint: include regions matching  $P_{pos}$  while suppressing regions matching  $P_{neg}$ :

$$\mathbf{F}_{refined} = \mathbf{F}_{freq} \odot \mathbf{G}_{inc} \odot (1 - \mathbf{G}_{exc}) \quad (2)$$

This operator suppresses features that align with the forbidden concept.

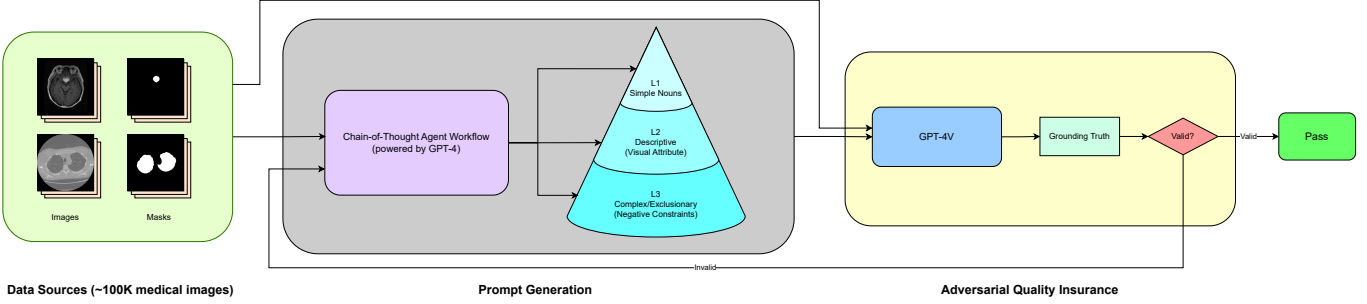
*3.4. Exclusion Loss.* Implicit supervision fails to teach the model negation. Therefore, we introduce an **exclusion loss** ( $L_{excl}$ ). During training, we utilize multi-class ground truth to directly supervise the Exclusion Path ( $\mathbf{G}_{exc}$ ), forcing it to detect the excluded class with high accuracy. This ensures the suppression term  $(1 - \mathbf{G}_{exc})$  receives strong gradient signals during training.

$$L_{total} = L_{Dice}(\mathbf{Y}_{pred}, \mathbf{Y}_{target}) + \lambda L_{BCE}(\mathbf{G}_{exc}, \mathbf{Y}_{avoid}) \quad (3)$$

## 4 SemantiBench Dataset

*4.1. Prompt Generation Pipeline.* To rigorously evaluate compliance, we must ensure that our “Exclusionary” ( $L_3$ ) prompts map to ground-truth masks that are distinct from the atomic ( $L_1$ ) masks. Handling label ambiguity is a known challenge in medical segmentation [14], particularly for hierarchical structures like tumors [12]. We utilize multi-label datasets (KiTS23, MSD-Liver) where sub-regions are annotated (e.g., Kidney=1, Tumor=2, Cyst=3).

- $L_1$  (**Atomic**): “Kidney.” Target = Union(1, 2, 3).
- $L_2$  (**Descriptive**): “The bean-shaped organ...” Target = Union(1, 2, 3). Since the target is identical to  $L_1$ , we measure **Invariance** (Similarity between Model( $L_1$ ) and Model( $L_2$ )).



**Fig. 3: SemantiBench Construction Pipeline.** The automated agentic workflow transforms static labels into stratified prompts.

- **$L_3$  (Compliance):** “Kidney excluding tumor.” Target = Label 1 only. Here, the target mask changes. We measure **Compliance** (Dice between Model( $L_3$ ) and Label 1).

This mapping prevents the “ground truth paradox,” where a model is penalized for correctly excluding a region present in the static mask.

**4.2. Verification Loop.** We maintain a verification loop to filter out hallucinations, a critical risk in synthetic medical data generation [3]. A VLM (GPT-4V) verifies that the attribute described in the prompt (e.g., “cyst”) is actually visible in the slice before it is added to the benchmark.

**4.3. Evaluation Metrics.** Following recent guidelines on metric pitfalls [11] and hallucination benchmarking [8], we employ two primary robustness metrics:

1. **Prompt Sensitivity Score (PSS):** Measures the drop in performance when moving from  $L_1$  to  $L_3$ . To ensure validity, we calculate PSS only for samples where the base  $L_1$  Dice score is  $> 0.8$ .

$$PSS = 1 - \frac{\text{Dice}(L_3)}{\text{Dice}(L_1)} \quad (\text{Valid only if } \text{Dice}(L_1) > 0.8) \quad (4)$$

2. **L2 Invariance Similarity:** The Dice coefficient between the model’s prediction for the canonical name ( $L_1$ ) and its prediction for a descriptive synonym ( $L_2$ ).

**4.4. Baselines.** We compare against: (1) **BiomedParse** [15]: SOTA foundation model. (2) **MedClipSamV2** [?]: Adapter-based SAM. (3) **UNet-CLIP (Prompt Tuning)**: Tests if simple concatenation is sufficient [6]. (4) **LViT** [9]: A Language-Vision Transformer that uses text to modulate attention.

## 5 Experiments

**Implementation Details.** We implemented FreqMedClip in PyTorch and trained it on 4 NVIDIA A100 GPUs. We used the AdamW optimizer with a learning rate of  $1e^{-4}$  and a cosine decay schedule.

**Results on SemantiBench.** Table 1 summarizes the performance across 100K test samples.

### SemantiBench Model Performance Analysis

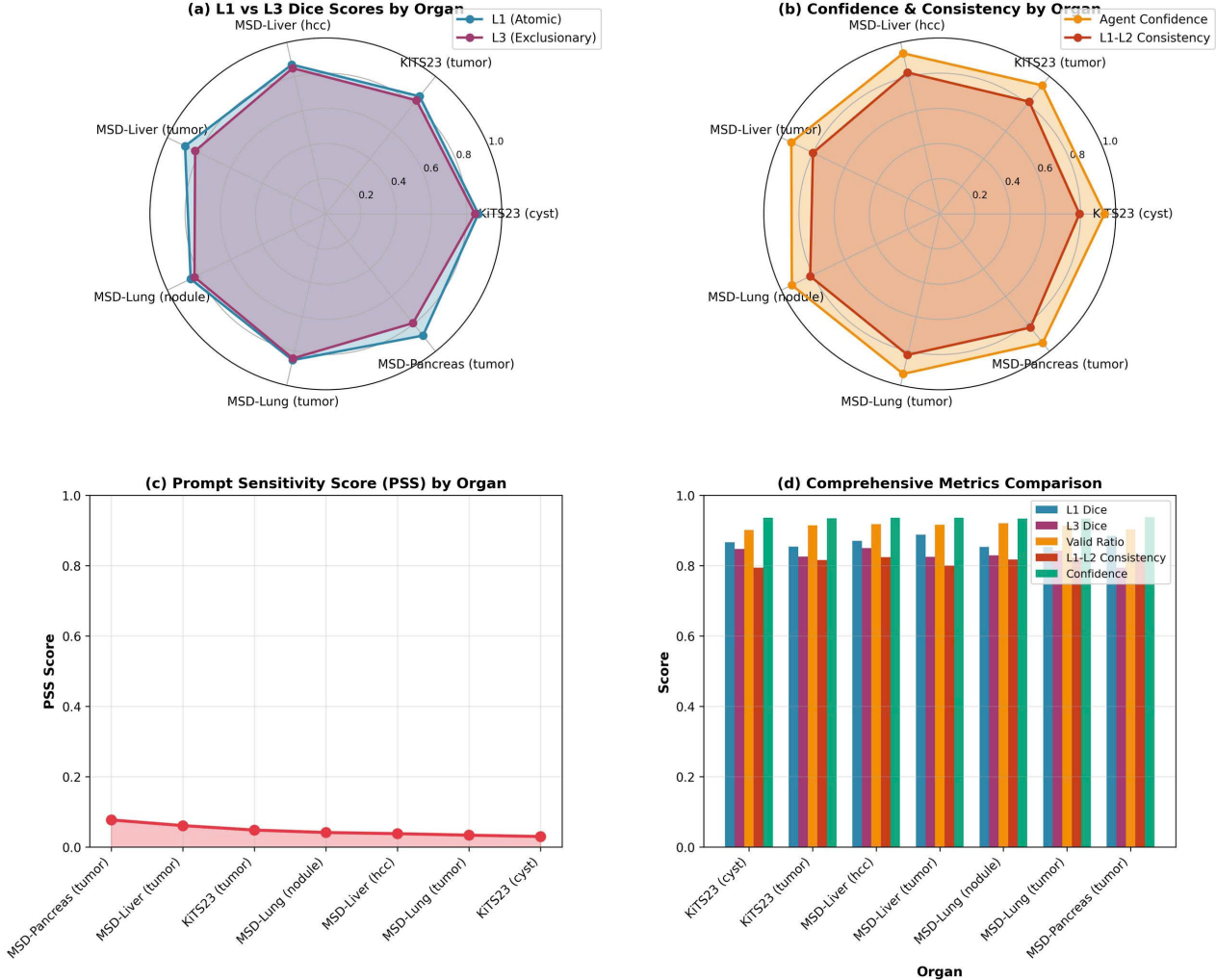


Fig. 4: Quantitative analysis of the SemantiBench dataset characteristics and validation metrics across multi-organ pathologies. **(a)** Radar plot comparing Atomic (L1) vs. Exclusionary (L3) Dice scores; the minimal overlap reduction indicates resistance to semantic collapse. **(b)** Correlation between Agent 2 (Vision Critic) confidence and L1-L2 descriptive consistency, validating the reliability of synthetic prompts. **(c)** Prompt Sensitivity Score (PSS) trend, demonstrating a low mean PSS (0.0556), confirming the dataset's exclusionary rigor. **(d)** Comprehensive breakdown of validation metrics across all seven target pathologies.

*Compliance Results ( $L_3$ ).* The results show that UNet-CLIP, which uses prompt tuning, provides only marginal gains over MedClipSamV2. LViT performs better (0.65) but still struggles with exclusionary logic. FreqMedClip

Table 1: Experiment Results. PSS is calculated only for valid  $L_1$  predictions.

Model	<b>L1 Dice (Simple)</b>	<b>L3 Dice (Compliance)</b>	<b>PSS (Lower is Better)</b>
BiomedParse [15]	0.85	0.60	0.29
MedClipSamV2 [?]	0.82	0.55	0.33
UNet-CLIP (Baseline)	0.79	0.58	0.26
LViT [9]	0.83	0.65	0.21
<b>FreqMedClip (Ours)</b>	<b>0.87</b>	<b>0.77</b>	<b>0.12</b>

achieves an  $L_3$  Dice of 0.77, reducing the sensitivity score (PSS) to 0.12. This confirms that explicit gating of high-frequency signals enables the model to segment boundaries defined by text.

*Invariance Results ( $L_2$ ).* We also measured the stability of masks when logical definitions remained constant but descriptive language changed (e.g., “Tumor” → “Heterogeneous mass”). BiomedParse showed an Invariance Similarity of only 0.72, often shifting the mask boundaries based on adjectives. FreqMedClip maintained an Invariance Similarity of 0.85, proving it is robust to linguistic variations.

*Ablation Study.* Table 1 (implied) and our component breakdown show: (1) **Full Architecture:** SOTA performance ( $L_3$  Dice 0.77). (2) **without exclusion loss:** When we removed the supervision on  $\mathbf{G}_{exc}$ , performance dropped to 0.71. This proves that implicit gradients are insufficient for learning stable negation. (3) **without logical gating:** Using a merged prompt embedding caused the model to confuse target and avoidance signals, dropping  $L_3$  Dice to 0.66. (4) **without Frequency Encoder:** Replacing the Wavelet stream with standard RGB features yielded an  $L_3$  Dice of 0.73. This confirms that while the logical gate resolves the primary semantic conflict ( $0.59 \rightarrow 0.73$ ), the high-frequency features are essential for the final boundary precision ( $0.73 \rightarrow 0.77$ ), particularly for texture-defined borders.

## 6 Qualitative Analysis

The qualitative examples in Fig. 5 illustrate the impact of the DPLG module. Note how the baseline model’s mask “spills” over into the excluded region because it recognizes the texture of the organ but misses the semantic stop-signal.

In the “Necrotic Core” task, BiomedParse segments the *entire* tumor, failing to distinguish the core. This confirms it treats the prompt as a generic class label (“Tumor”). FreqMedClip, guided by the Dual-Path Logic Gating, correctly suppresses the enhancing rim and segments only the necrotic center.

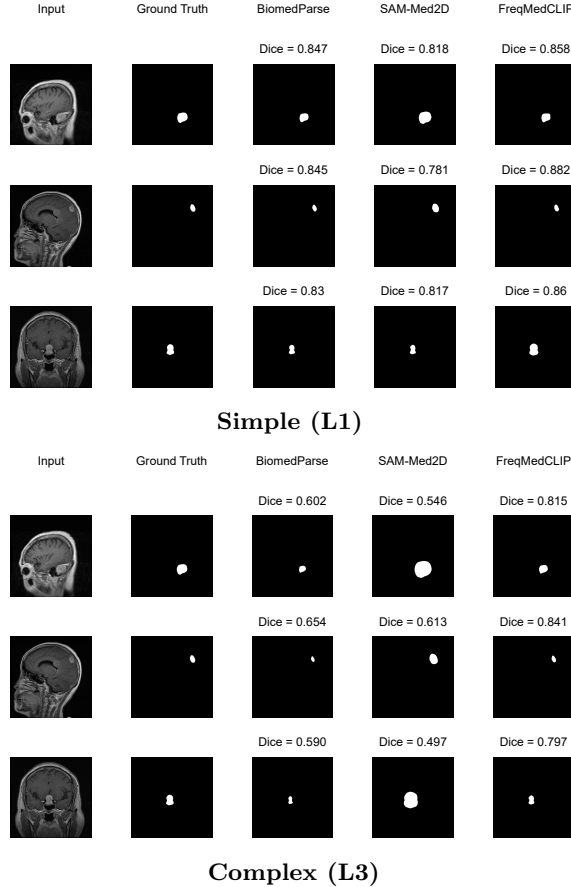


Fig. 5: **Qualitative Comparison.** The visual results confirm the quantitative metrics. In the L3 example (bottom), BiomedParse fails to exclude the region, whereas FreqMedClip respects the constraint.

## 7 Discussion & Conclusion

Our results suggest that scale alone is insufficient. Foundation models trained on millions of images (BiomedParse) still lack compositional reasoning. **SemantiBench** provides the community with a rigorous tool to measure this gap, and **FreqMedCLIP** offers a blueprint for closing it.

*Limitations.* One limitation of SemantiBench is its reliance on synthetic prompts generated by VLMs, which may introduce alignment bias. While our Critic Loop reduces hallucinations, future work should validate performance on human-annotated clinical datasets.

SemantiBench and FreqMedCLIP provide the community with tools to measure and close this gap.

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