# Focus on What Matters: Enhancing Medical Vision-Language Models with Automatic Attention Alignment Tuning

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#### Abstract

Medical Large Vision-Language Models (Med-LVLMs) often exhibit suboptimal attention distribution on visual inputs, leading to hallucinated or inaccurate outputs. Existing mitigation methods primarily rely on inference-time interventions, which are limited in attention adaptation or require additional supervision. To address this, we propose  $A^{3}TUNE$ , a novel fine-tuning framework for Automatic Attention Alignment Tuning. A<sup>3</sup>TUNE leverages zeroshot weak labels from SAM, refines them into prompt-aware labels using BioMedCLIP, and then selectively modifies visually-critical attention heads to improve alignment while minimizing interference. Additionally, we introduce a A<sup>3</sup>MOE module, enabling adaptive parameter selection for attention tuning across diverse prompts and images. Extensive experiments on medical VQA and report generation benchmarks show that A<sup>3</sup>TUNE outperforms state-of-the-art baselines, achieving enhanced attention distributions and performance in Med-LVLMs.<sup>1</sup>

## 1 Introduction

While medical Large Vision-Language Models (Med-LVLMs) have shown significant progress in the medical domain (Li et al., 2024; Chen et al., 2024d; Thawkar et al., 2023; Moor et al., 2023), they often produce inaccurate or hallucinated outputs that deviate from the provided visual medical information, as revealed by recent benchmarks (Xia et al., 2024; Gu et al., 2024; Chen et al., 2024a). An example of medical visual question answering (VQA) extracted from the SLAKE (Liu et al., 2021) dataset is shown in Figure 1(A), where we visualize the average attention map on the image inputs during generating answers. We can observe that LLaVA-Med in Figure 1(A.2) generates the hallucinated response "*Alzheimer's disease*", neglecting

the tumor region and over-focusing on irrelevant background areas, as shown in the corresponding attention map. This reveals a significant **bias in attention distribution** on visual inputs that limits the model's effectiveness, which have been identified in general LVLMs (Gong et al., 2024; Woo et al., 2024; Liu et al., 2025).

Unfortunately, none of the specified bias mitigation strategies have been proposed in the medical domain. In the general domain, research primarily focuses on inference-time interventions to reduce attention biases, employing two main approaches. The first approach, contrastive decoding (Leng et al., 2024; Favero et al., 2024; Liu et al., 2025; Woo et al., 2024; Gong et al., 2024), introduces a contrastive adjustment to the decoding logits. However, this method does not directly modify the attention distribution, meaning it cannot guarantee that the model attends to diagnostically critical regions. The second approach directly modifies attention maps during inference, as seen in ControlMLLM (Wu et al., 2024), which enforces the model to focus on pre-annotated regions of interest (RoIs). However, as shown in Figure1(A.3), this method still generates partially hallucinated content. Besides, it requires additional tuning and ground truth RoIs for each inference process, making it impractical for real-world applications.

To overcome the limitations of inference-time intervention, an ideal solution is to automatically adjust attention maps towards RoIs during fine-tuning for downstream medical tasks. This approach enables the model to place more attention on critical regions during inference, eliminating the need for additional labels or interventions. However, implementing such a method poses several challenges:

(1) Limited availability of medical segmentation labels. As shown in Figure 1(A.3), using segmentation labels (RoIs) as guidance, as done in ControlMLLM, can enhance the learning of accurate attention maps and answer correctness. How-

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<sup>&</sup>lt;sup>1</sup>Source code is available at https://github.com/ Aofei-Chang/A3Tune



Figure 1: (A) Examples of medical VQA and attention maps on medical images. In this example of Brain MRI from the SLAKE dataset, red box denotes the RoI of the brain tumor that LLaVA-Med should focus on. Red texts and green texts indicate wrong answers and correct answers, respectively. (B) Example of ground truth RoIs for different prompts on an Abdomen CT from SLAKE.

ever, such labels are often unavailable in medical datasets.

(2) Trade-off between attention alignment and model stability. Assuming that RoI labels are available, directly modifying the attention maps of all attention heads towards the labels without any strategy is still risky. This may lead to the over-alignment issue that potentially impacts the output stability and overall performance. Therefore, achieving the right balance between attention alignment and model stability is essential.

(3) Adapting attention to diverse prompts and images. Even if attention alignment and model stability are balanced, the parameter-sharing strategy in fine-tuning remains a limitation for adaptive attention alignment. For example, as shown in Figure 1, the optimal RoIs can vary significantly based on the input prompt and image, requiring dynamic alignment. While the computation of attention maps adjusts to input representations, the shared parameters in fine-tuning limit the model's ability to flexibly learn attention distribution across diverse inputs.

To address these challenges simultaneously, we propose  $A^{3}TUNE$ , a novel fine-tuning framework designed for Automatic Attention Alignment **Tun**ing. As shown in Figure 2,  $A^{3}TUNE$  integrates a set of cooperative strategies to ensure that attention in Med-LVLMs is well aligned, minimally disruptive, and highly adaptable across diverse medical tasks. Firstly, to overcome the lack of segmentation labels (Challenge 1),  $A^{3}TUNE$  utilizes zero-shot segmentation labels generated by SAM (Roy et al., 2023) and further refines them into prompt-aware weak labels using BioMedCLIP (Zhang et al., 2023). These weak labels serve as guidance for attention alignment, eliminating the need for manual annotations. However, weak labels alone are not sufficient — uncontrolled modifications on all attention heads can disrupt model stability (Challenge 2). To further balance attention alignment with model stability, A<sup>3</sup>TUNE selectively modifies only the most "visually-critical" attention heads, minimizing the risk of over-alignment and instability. Furthermore, the parameter-sharing strategy in alignment tuning remains a limitation, as RoIs vary significantly based on the input prompt and image (Challenge 3). To address this, we incorporate a custom-designed **M**ixture-**o**f-Experts A<sup>3</sup>MOE into A<sup>3</sup>TUNE on attention modules, allowing the model to dynamically select parameters and adjust attention maps for different images and prompts.

In summary, this work makes the following contributions: (1) We propose  $A^{3}TUNE$ , a novel visual attention tuning approach that utilizes zeroshot weak labels to refine the visual focus of Med-LVLMs and enhance their performance. (2) We develop a set of designs for cooperative attention alignment tuning: (i) weak label supervision, (ii) selective tuning of visually-critical attention heads, and (iii) a A<sup>3</sup>MOE module for adaptive attention adjustment. (3) We conduct extensive experiments on five medical VQA and two report generation benchmarks against ten baselines, demonstrating that A<sup>3</sup>TUNE outperforms state-of-the-art methods in both effectiveness and interpretability, improving visual grounding and overall model performance.

## 2 Related Work

While some efforts, such as CoMT (Jiang et al., 2024), have attempted to reduce hallucinations in Med-LVLMs for report generation by training on hierarchical QA pairs derived from real clinical



Figure 2: (A) The overview of A<sup>3</sup>TUNE and (B) the details of the designed visual attention alignment tuning.

image reports, mitigation strategies specifically designed for Med-LVLMs remain largely underexplored. Since Med-LVLMs share the same structure and training process as general LVLMs, hallucination issues are a common challenge across both. As a result, many inference-time mitigation strategies developed for LVLMs are also applicable to Med-LVLMs, including: (1) Enhancing visual information and mitigating text over-reliance: VCD (Leng et al., 2024), M3ID (Favero et al., 2024), PAI (Liu et al., 2025), HELPD (Yuan et al., 2024), RBD (Liang et al., 2024). (2) Mitigating visual token attention bias, with approaches such as AVISC (Woo et al., 2024) and DAMRO (Gong et al., 2024). (3) Refining and modifying LLM generation patterns during decoding, OPERA (Huang et al., 2024), DoLa (Chuang et al., 2023). In addition, other mitigation methods, such as Truth-Flow (Wang et al., 2025), leverage trained steering vectors to guide the model toward more truthful outputs in LLMs. Despite these advancements, significant challenges remain in mitigation attention biases in Med-LVLMs.

## **3** Preliminaries

## 3.1 Background of Med-LVLMs

Med-LVLMs share the same fundamental architecture as general LVLMs (Liu et al., 2024; Chen et al., 2023), consisting of three primary components: the Visual Encoder, the Visual Alignment Layer, and the Large Language Model (LLM). The input image is firstly divided into N patches and processed by the Visual Encoder, which converts it into a sequence of visual tokens  $\mathcal{T}_v$  with embeddings represented as  $\mathbf{X}_v \in \mathbb{R}^{N \times d_v}$ , where  $d_v$  denotes the dimension of the visual hidden representation. The visual tokens are then projected through the Visual Alignment Layer into  $\mathbf{X}'_v \in \mathbb{R}^{N \times d_p}$ , where  $d_p$  denotes the dimension of the LLM token space. These aligned visual tokens are subsequently forwarded to the LLM along with the embeddings  $\mathbf{X}_p \in \mathbb{R}^{V \times d_p}$  of the tokenized textual prompts  $\mathcal{T}_p$ , where V is the number of text tokens.

The final objective of the LLM in Med-LVLMs is to predict the next token  $y_t$  based on the current visual input  $\mathbf{X}'_v$ , prompt input  $\mathbf{X}_p$ , and previously generated tokens  $y_{< t}$ , formulated as:

$$p_t = p(y_t \mid \mathbf{X}'_v, \mathbf{X}_p, y_{< t}; \Theta^*), \tag{1}$$

where  $\Theta^*$  denotes the parameters of the LLM.

#### 3.2 Visual Attention Map

Most Med-LVLMs adopt a Transformer (Vaswani et al., 2017) decoder-based LLM, which processes inputs through L decoder layers, each equipped with a multi-head attention module. In layer l, the module includes H attention heads, where each head h (with  $1 \le h \le H$ ) computes attention separately using their corresponding attention map  $\mathbf{M}_{lh}$ . This attention mechanism models relationships between visual and textual tokens.

To analyze how visual tokens contribute to text generation, we focus on the attention scores between visual tokens and subsequent text tokens, referred to as visual attention map  $\mathbf{M}_{lh}^v$ , a submatrix of the overall attention map  $\mathbf{M}_{lh}^v$ . To gain a global understanding of attention distribution on visual tokens, the averaged visual attention map  $\mathbf{M}^v$ can be obtained by aggregating attention across all heads and layers (Wu et al., 2024):

$$\mathbf{M}^{v} = \frac{1}{LH} \sum_{l=1}^{L} \sum_{h=1}^{H} \mathbf{M}_{lh}^{v}.$$
 (2)

# 4 The Proposed A<sup>3</sup>TUNE

A<sup>3</sup>TUNE aims to automatically align attention to enhance visual grounding and improve the performance of Med-LVLMs. The fine-tuning pipeline is illustrated in Figure 2. Given an input image Iand prompt P, we firstly generate a set of promptaware weak labels S using a zero-shot method (Section 4.1). To achieve effective attention alignment (Section 4.2) with the guidance of S, we first identify visually-critical attention heads, then integrate an A<sup>3</sup>MOE design into each decoder layer to enable flexible attention distribution learning.

## 4.1 Prompt-Aware Weak Labels Generation

Given a medical image I, we first use SAM (Kirillov et al., 2023) to generate a set of candidate segments  $S^*$  following (Yang et al., 2024). However, using all segments in  $S^*$  for attention tuning introduces noise, as only a subset is relevant to each prompt. Therefore, it is necessary to select promptaware segments as weak labels to adaptively guide attention tuning. To filter prompt-aware weak labels, we embed each segment  $s \in S^*$  into a feature representation  $\mathbf{E}_s$  using BioMedCLIP's vision encoder, while the text prompt P is embedded into  $\mathbf{E}_P$  using its text encoder. We then select an adaptive threshold  $\tau_K$  to select K segments that are most similar to the text prompt based on cosine similarity (Sim) in the embedding space:

$$\mathcal{S} = \{ s \in \mathcal{S}^* \mid \operatorname{Sim}(\mathbf{E}_s, \mathbf{E}_P) \ge \tau_K \}.$$
(3)

## 4.2 Visual Attention Alignment Tuning

The goal of visual attention alignment tuning is to align averaged visual attention map  $\mathbf{M}^v$  with fixed weak labels S (i.e., masking regions) via Eq. (3) during fine-tuning. To achieve this goal, we first design a new parameter-efficient fine-tuning strategy and select "visually-critical" attention heads to the alignment of attention maps with S.

# 4.2.1 A<sup>3</sup>MOE Design

We build  $A^{3}TUNE$  based on a parameter-efficient fine-tuning technique, LoRA (Hu et al., 2021), and apply it to all linear modules of the LLM in the Med-LVLM, including the Query and Key matrices in attention modules. In standard LoRA fine-tuning, the trainable LoRA parameters  $\Delta W_q$ and  $\Delta W_k$  for Query and Key matrices are shared across all training instances. However, this static parameter-sharing strategy in attention is insufficient for  $A^{3}TUNE$ . As shown in the abdominal



Figure 3: Motivation for using A<sup>3</sup>MOE. The second column shows prompt-aware weak labels, with red bounding boxes and green inner segments. The third column shows the attention maps generated using shared parameters for the Query and Key matrices.

organ analysis tasks (Figure 3), weak labels S vary with the prompt, yet attention maps generated with shared  $\Delta W_q$  and  $\Delta W_k$  lack the flexibility to adapt effectively to different prompts and images.

To address this limitation, we introduce  $A^3MOE$ , a **M**ixture-of-Experts mechanism specifically designed for  $A^3TUNE$ , with two sub-modules: Q-MoE and K-MOE, applied to the LoRA parameters  $\Delta W_q^{(l)}$  and  $\Delta W_k^{(l)}$  in each LLM decoder layer l. For clarity, we omit the explicit annotation of l in subsequent equations, as  $A^3MOE$  is applied consistently across all decoder layers. The following sections describe these sub-modules in detail.

**Q-MoE: Prompt-Level MoE on Query Matrix.** As shown in Figure 3, the target RoIs often depend on the text prompt, even for the same image. To handle this, we introduce Q-MoE, a prompt-level MoE on  $\Delta \mathbf{W}_q$ , consisting of  $O^q$  experts. For the *o*-th expert  $(1 \le o \le O^q)$ , we define  $\mathcal{E}_o^q = \mathbf{B}_o \mathbf{A}_o$ , where  $\mathbf{B}_o \in \mathbb{R}^{d_p \times r}$  and  $\mathbf{A}_o \in \mathbb{R}^{r \times d_p}$  are matrices with a low rank *r*.

To dynamically route experts based on the prompt, we apply a prompt-level gating mechanism that generates router weights  $\alpha$  based on the hidden states of the prompt  $\mathbf{H}_p$ , which is obtained at each layer l after processing  $\mathbf{X}_p$  from the previous decoder layers. To capture the prompt's overall context,  $\mathbf{H}_p$  is first averaged using a pooling operation. The routing and parameter computation are formulated as:

$$\boldsymbol{\alpha} = \operatorname{softmax}(\operatorname{MLP}(\operatorname{Pooling}(\mathbf{H}_p))), \quad (4)$$

$$\Delta \mathbf{W}_q = \sum_{o=1}^{O^q} \alpha_o \mathcal{E}_o^q, \tag{5}$$

where  $\boldsymbol{\alpha} \in \mathbb{R}^{O^q}$  represents the router weights, with each  $\alpha_o$  determining the contribution of the

corresponding expert  $\mathcal{E}_o^q$ . MLP denotes the multilayer perceptron used in the gating mechanism.

K-MoE: Visual Token-Level Sparse MoE in Key Matrix. Unlike text prompts, which can be summarized with pooling, visual inputs contain finegrained information at the token level, where subtle differences can indicate abnormalities that require varied attention. To capture these nuances, we introduce K-MoE, a visual token-level MoE applied on  $\Delta W_k$ . K-MoE includes  $O^k$  experts, each implemented using LoRA.

For each visual token c, the gating mechanism dynamically select experts based on its hidden states  $\mathbf{H}_v^c$ , and the LoRA parameters  $\Delta \mathbf{W}_k^c$  for visual token c are computed as:

$$\boldsymbol{\beta}^{c} = \operatorname{softmax}(\operatorname{MLP}(\mathbf{H}_{v}^{c})), \quad (6)$$

$$\Delta \mathbf{W}_{k}^{c} = \sum_{o=1}^{O_{k}} \mathbb{1}_{o} \boldsymbol{\beta}_{o}^{c} \boldsymbol{\mathcal{E}}_{o}^{k}, \tag{7}$$

where  $\beta^c \in \mathbb{R}^{O^k}$  and  $\mathbb{1}_o$  is a binary indicator enforcing sparsity by retaining only the top-*B* gating weights. It is set to 1 if expert  $\mathcal{E}_o^k$  is among the top-*B* most relevant experts, otherwise it is set to 0. This mechanism ensures that for each visual token, only the most relevant experts contribute, minimizing interference among visual tokens while improving efficiency in alignment tuning.

## 4.2.2 "Visually-Critical" Heads Selection

Each layer l in the LLM decoder fine-tuned with A<sup>3</sup>MOE learns H attention heads and the total number of attention heads is  $L \times H$ . Among them, the attention heads that assign higher weights to visual information are more crucial in processing visual features. Thus in layer l, the importance of each head h can be quantified using the visual attention ratio  $r_{lh}$ , which measures the proportion of attention allocated to visual tokens relative to all tokens as follows:

$$r_{lh} = \frac{\sum_{c \in \mathcal{T}_v} \mathbf{M}_{lh}^v[c]}{\sum_{c' \in \mathcal{T}_v \cup \mathcal{T}_p} \mathbf{M}_{lh}^v[c']},$$
(8)

where  $\mathbf{M}_{lh}^{v}[c]$  represents the attention score assigned by attention head h in layer l to token c.

To refine the visual attention matrix  $\mathbf{M}^{v}$ , we select top-R "visually-critical" heads based on the visual attention ratios  $r_{lh}$ . To achieve this, we use a binary indicator  $\mathbb{1}_{lh}$  to identify and retain only the most influential attention heads. Specifically, it is set to 1 if attention head (l, h) is among the R

heads with the highest visual attention ratios and 0 otherwise. The updated visual attention matrix is then computed as:

$$\widetilde{\mathbf{M}}^{v} = \frac{1}{R} \sum_{l=1}^{L} \sum_{h=1}^{H} \mathbb{1}_{lh} \mathbf{M}_{lh}^{v}.$$
(9)

## 4.2.3 Attention Alignment Function

After obtaining refined  $\mathbf{M}^v$ , we define how it is adjusted to align with the weak labels S. Intuitively, in the averaged attention map  $\mathbf{M}^v$ , tokens within the masking regions S should receive higher attention values. Motivated by the cross-attention control design for image generation in (Chen et al., 2024c), we use a mask-based energy function to guide the attention alignment tuning:

$$\mathcal{L}_{\text{align}} = \sum_{s \in \mathcal{S}} \left( 1 - \frac{\sum_{c \in s} \widetilde{\mathbf{M}}_{c}^{v}}{\sum_{c'=1}^{N} \widetilde{\mathbf{M}}_{c'}^{v}} \right)^{2}, \qquad (10)$$

where s represents each referring segment in S, c and c' is the visual token index, and N is the number of visual tokens. Among all N visual tokens, this function encourages higher attention on visual tokens within each s, minimizing the loss and guiding  $\widetilde{\mathbf{M}}^{v}$  to effectively focus on the prompt-aware regions S.

## 4.3 Final Objective

In the fine-tuning process, we incorporate the loss of proposed visual attention alignment tuning  $\mathcal{L}_{align}$ (Eq. 10) as a regularization term for downstream medical tasks. This term is combined with the language modeling objective  $\mathcal{L}_{LLM}$ :

$$\mathcal{L}_{LLM} = -\sum_{t=1}^{T} \log p_t(y_t \mid \mathbf{X}'_v, \mathbf{X}_p, y_{< t}; \Theta, \Theta^*),$$
(11)

where  $\Theta^*$  denotes the frozen parameters of LLM,  $\Theta$  denotes the trainable parameters in LoRA and  $A^3MOE$ , and  $y_{<t}$  denotes the generated tokens before time step t. The final objective loss is:

$$\mathcal{L} = \mathcal{L}_{LLM} + \lambda \mathcal{L}_{align}, \qquad (12)$$

where  $\lambda > 0$  is a hyperparameter that controls the strength of attention tuning.

## **5** Experiments

We evaluate our method on representative Med-LVLMs, including LLaVA-Med (Li et al., 2024) and LLaVA-Med-1.5. The experimental results for LLaVA-Med-1.5 are provided in Appendix G.

Madal	Mathad	SI	ake	VQA	-RAD	Path	VQA	IU-Xray	OmniVQA
Model	Method	Open	Closed	Open	Closed	Open	Closed	Closed	Closed
	LLaVA-Med	41.28	57.75	32.48	68.90	10.34	52.49	72.83	31.79
	Greedy	42.81	60.00	35.68	68.11	11.35	52.34	75.00	31.11
	Beam (Sutskever et al., 2014)	42.31	61.69	33.57	66.93	9.86	53.79	73.59	30.89
	Nucleus (Holtzman et al., 2020)	41.06	61.41	32.64	68.11	9.93	53.17	73.09	30.96
	VCD (Leng et al., 2024)	39.76	59.44	33.93	66.54	11.17	53.52	74.36	32.89
LLaVA-Med	DoLa (Chuang et al., 2023)	42.37	59.72	35.68	68.50	11.38	52.55	74.87	31.04
	OPERA (Huang et al., 2024)	40.44	60.00	35.54	68.50	9.77	53.17	75.00	31.72
	AVISC (Woo et al., 2024)	41.46	59.72	35.43	64.17	11.03	52.20	73.85	32.48
	M3ID (Favero et al., 2024)	38.85	60.28	35.31	62.60	9.80	52.93	72.32	31.11
	DAMRO (Gong et al., 2024)	41.33	59.72	32.73	66.54	10.80	51.84	72.19	31.45
	PAI (Liu et al., 2025)	43.18	60.28	35.10	68.50	11.09	52.46	74.87	32.13
	LLaVA-Med + LoRA	80.65	82.82	33.37	66.54	31.92	90.95	83.29	90.65
	Greedy	81.12	85.07	31.88	68.50	33.90	91.86	83.93	90.73
	Beam (Sutskever et al., 2014)	81.32	86.76	32.55	68.90	33.60	91.92	84.06	90.58
	Nucleus (Holtzman et al., 2020)	80.18	85.35	31.34	68.50	30.14	90.92	83.41	90.05
	VCD (Leng et al., 2024)	79.58	84.23	32.96	67.72	30.20	90.86	83.93	90.73
LLaVA-Med	DoLa (Chuang et al., 2023)	81.84	86.48	31.94	68.50	34.00	91.86	84.06	90.69
+ LoRA	OPERA (Huang et al., 2024)	81.25	86.48	33.18	68.90	33.64	91.80	84.06	90.42
	AVISC (Woo et al., 2024)	80.15	85.63	33.66	69.29	32.37	90.62	84.06	90.54
	M3ID (Favero et al., 2024)	79.83	84.79	31.40	68.90	32.47	91.15	84.95	90.73
	DAMRO (Gong et al., 2024)	82.19	83.66	32.41	66.14	32.27	90.12	84.69	90.08
	PAI (Liu et al., 2025)	81.02	86.76	32.14	68.11	33.46	91.77	84.31	90.95
	A <sup>3</sup> TUNE (ours)	82.36	86.76	36.97	70.87	34.61	92.19	85.97	91.98

Table 1: Performance comparison on medical VQA benchmarks using LLaVA-Med. The best results are highlighted in **bold**, and the second-best results are underlined. OmniVQA corresponds to the OmniMedVQA dataset.

#### 5.1 Settings

**Datasets.** We evaluate our method on two key tasks in medical application of Med-LVLMs: medical VQA and medical report generation. For medical VQA, we use diverse datasets including SLAKE (Liu et al., 2021), VQA-RAD (Lau et al., 2018), PathVQA (He et al., 2020), IU-Xray (Demner-Fushman et al., 2016) and OmniMedVQA (Hu et al., 2024). For medical report generation, we use MIMIC-CXR (Johnson et al., 2019) and IU-Xray. Details of dataset processing and settings are provided in Appendix A.

**Baselines.** We compare our approach with widely used hallucination mitigation methods with or without fine-tuning, including decoding strategies and contrastive decoding techniques<sup>2</sup>. The **decoding** baselines include Greedy decoding, Nucleus sampling (Holtzman et al., 2020), Beam search (Sutskever et al., 2014). For **contrastive decoding** techniques, we specifically compare with: VCD (Leng et al., 2024), OPERA (Huang et al., 2024), DoLa (Chuang et al., 2023), AVISC (Woo et al., 2024), M3ID (Favero et al., 2024), DAMRO (Gong et al., 2024) and PAI (Liu et al., 2025). Additionally, we include ControlM-LLM (Wu et al., 2024) as a baseline only when the ground truth RoIs are available. Detailed settings

Table 2: Comparison of Visual Attention Distribution.

Mathad	Metrics					
Method	Coverage ↑	Intensity $\uparrow$				
LLaVA-Med	0.122	0.076				
LLaVA-Med + LoRA	0.132	0.076				
A <sup>3</sup> TUNE	0.275	0.147				

of baselines and implementation are in Appendix B and Appendix C, respectively.

Metrics. (1) Metrics for Performance Evaluation. For *medical VOA*, we report Accuracy for close-ended questions and Recall for openended questions, following LLaVA-Med (Li et al., 2024). For medical report generation, we use standard metrics for generation tasks, including BLEU (Papineni et al., 2002), ROUGE-L (Lin, 2004), METEOR (Banerjee and Lavie, 2005), and BERTScore (Zhang et al., 2019). Additionally, we report domain-specific metrics designed for medical report generation: CheXbert (Smit et al., 2020), RadGraph (Jain et al., 2021) and RaTEScore (Zhao et al., 2024). More details of these metrics are provided in Appendix D.1. (2) Metrics for Attention Maps Evaluation. For datasets with ground truth RoIs (e.g., SLAKE), we design two metrics to evaluate the attention distribution on images: (1) Coverage Score (Coverage), which measures spatial alignment, and (2) Intensity Alignment (Intensity), which assesses the degree of focus. Details of these two metrics are in Appendix D.2.

<sup>&</sup>lt;sup>2</sup>We do not compare with the methods requiring tailored de-hallucination training pipelines such as CoMT (Jiang et al., 2024) and HELPD (Yuan et al., 2024).

Datasat	Metric		Method										
Dataset		LLaVA-Med + LoRA	Greedy	Beam	Nucleus	VCD	DoLa	OPERA	AVISC	M3ID	DAMRO	PAI	A <sup>3</sup> Tune
	BLEU	7.70	8.86	9.34	7.80	8.83	8.93	8.27	8.52	8.63	7.29	8.87	11.05
	ROUGE-L	26.15	27.09	27.56	26.72	27.36	26.94	27.14	26.97	27.79	25.61	26.74	30.00
	METEOR	29.50	26.01	26.44	30.33	31.77	25.74	29.66	31.14	31.65	30.03	25.99	34.26
IU-Xray	BERTScore	88.36	88.50	88.52	88.28	88.30	88.42	88.41	88.47	88.52	88.14	88.39	89.05
	CheXbert	53.81	52.55	52.88	52.73	51.86	52.27	56.26	53.33	54.45	51.50	52.45	57.19
	RadGraph	20.44	20.76	21.29	20.85	22.02	20.63	21.87	22.27	22.22	20.75	20.48	24.24
	RaTEScore	58.37	58.24	58.77	57.84	58.93	58.10	58.73	59.21	59.37	59.19	57.81	63.03
	BLEU	3.28	4.07	3.39	3.29	3.53	3.99	3.14	3.34	3.62	3.37	4.32	4.56
	ROUGE-L	16.54	18.75	17.25	16.14	16.58	18.62	15.52	16.79	16.67	15.75	18.64	19.03
MIMIC	METEOR	17.90	18.81	17.36	17.98	18.68	18.68	14.70	18.29	18.22	17.03	19.78	20.23
CYP	BERTScore	85.57	86.14	85.78	85.42	85.57	86.14	84.75	85.59	85.58	85.27	86.10	86.17
CAR	CheXbert	22.14	24.24	23.05	21.76	23.38	25.14	20.07	22.74	22.95	22.76	25.78	24.93
	RadGraph	9.43	10.73	9.78	9.20	9.93	10.75	7.74	9.52	9.93	9.40	11.17	11.55
	RaTEScore	40.00	41.06	38.59	39.26	40.95	40.73	35.72	40.08	39.87	39.62	41.03	42.73

Table 3: Performance on report generation benchmarks using LLaVA-Med fine-tuned with LoRA.

# 5.2 Medical VQA Results

The performance of  $A^{3}TUNE$  on diverse medical VOA benchmarks is presented in Table 1, indicating A<sup>3</sup>TUNE maintains its superiority across all datasets and its effectiveness across diverse medical images and VQA tasks. In addition, we also analyze visual attention distribution on the test set of SLAKE with annotated RoIs (Table 2). Since the baselines in Table 1 improve the model only on the decoding side without modifying attentions, their attention distributions remain similar to the base model, LLaVA-Med and we compare only against LLaVA-Med and LLaVA-Med + LoRA. We observe that A<sup>3</sup>TUNE achieves the highest scores in both coverage (0.275) and intensity (0.147), outperforming baselines. These results highlight A<sup>3</sup>TUNE's ability to focus more effectively on RoIs in medical images, explaining its better VQA performance and improved interpretability.

# 5.3 Medical Report Generation Results

Table 3 presents the evaluation of  $A^{3}TUNE$  on medical report generation tasks using traditional metrics (e.g., BLEU, ROUGE-L) and domain-specific metrics such as RaTEScore. The baselines are applied on fine-tuned LLaVA-Med using LoRA, as the original LLaVA-Med performs significantly worse on this task, as shown in the full results in Appendix E. Similar to the results in Table 1,  $A^{3}TUNE$ outperforms all baselines across both datasets and almost all metrics.

## 5.4 Module Effectiveness Analysis

## 5.4.1 Weak Labels Generation

(1) Quality of Weak Labels. This experiment evaluates the upper bound of  $A^3TUNE$ 's performance by upgrading weak labels to high-quality ground



Figure 4: Effectiveness analysis of RoIs labels. **Base** is the base model, **Control** means adding ControlMLLM to align attention maps with ground truth labels, **Weak** uses weak labels, and **GT** uses ground truth labels.



Figure 5: Analysis of (a) the number of selected attention heads R in A<sup>3</sup>TUNE and (b) the number of weak labels K, evaluated on the SLAKE dataset.

truth labels (**GT**). We use a subset of SLAKE with RoIs annotations, including 992 training samples and 206 test samples and take ControlMLLM (Wu et al., 2024) as the baseline.

Figure 4 (B.2) shows that replacing weak labels (**Weak**) with high-quality labels (**GT** in  $A^{3}TUNE$  further improves performance, validating the reasonableness of using weak labels. Additionally, while ControlMLLM (**Control**) improves performance in LLaVA-Med (Figure 4 (A)), it negatively impacts fine-tuned LLaVA-Med+LoRA in Figure 4 (B.1), whereas  $A^{3}TUNE$  achieve improved results. These results highlight both the effectiveness of  $A^{3}TUNE$  and its potential for further improvement when ground truth RoI labels are available.

(2) Selection of Weak Labels. To obtain promptaware weak labels, we select K segments that are most similar to the text prompt. In this experiment,



Figure 6: Case study for fine-grained effectiveness analysis. The red box in the first column (not provided as input to the model) highlights the RoI that LLaVA-Med should focus on. The second column shows weak segmentation labels, with red bounding boxes and green inner segments generated using the method described in Section 4.1.



Figure 7: Analysis of (a) the contribution of different modules in  $A^3MOE$ , and (b) the number of experts in K-MoE within  $A^3MOE$ . In (a), each module's impact is evaluated by removing it from  $A^3TUNE$ , denoted "w/o".

we analyze the impact of different values of K on model performance using the SLAKE dataset. As shown in Figure 5 (a) increasing K initially improves performance by adding more relevant labels. However, performance peaks at K = 4, after which additional segments introduce noise, leading to a performance decline.

## 5.4.2 Selection of Attention Heads

In A<sup>3</sup>TUNE, we select R most "visually-critical" attention heads to balance model stability and effectiveness of A<sup>3</sup>TUNE. As shown in Figure 5 (b), disabling attention tuning (R = 0) results in noticeably lower performance. Performance peaks at R = 128, which strikes a balance between the strength of attention tuning and stability. Thus, we set R = 128 in our experiments.

# 5.4.3 A<sup>3</sup>MOE Design

(1) Ablation Study of  $A^3MOE$ . Figure 7 (a) shows the impact of removing key components of  $A^3MOE$  from  $A^3TUNE$ : Q-MOE, K-MOE and  $A^3MOE$  as a whole. The removal of each module leads to a noticeable performance drop, particularly for K-MOE. This highlights the importance of K-MOE, which is designed for fine-grained attention

tuning at the visual token level and is critical for datasets with diverse image modalities like SLAKE. Additionally, the removal of the entire  $A^3MOE$  leads to a further decline in performance. However, even with these components removed, the performance of  $A^3TUNE$  remains above the baseline LoRA tuning on LLaVA-Med, demonstrating the effectiveness of the overall framework.

(2) The Number of Selected Experts in K-MoE. As shown in Figure 7(b), performance peaks at B = 3. Beyond this point, additional experts lead to diminishing and unstable returns, likely due to saturation and increased interference among experts handling different visual tokens.

## 5.5 Case Study

Figure 6 presents a case study for three image modalities. The attention map visualizations demonstrate that our method effectively redirects the model's focus to relevant regions, mitigating hallucination issues compared to baselines. In addition, we provide fine-grained effectiveness evaluation across Chest X-ray, Abdomen CT, and Brain MRI images from SLAKE in Appendix F Table 10.

#### 5.6 Generalization to Other Medical LVLMs

We further evaluate our approach on medical VQA (Table 4) and report generation (Table 5) using a more recent and stronger Med-LVLM, HuatuoGPT-Vision-7B (Chen et al., 2024b). As shown in the results, transferring  $A^{3}TUNE$  to this new backbone consistently achieves the best performance across all metrics on IU-Xray for report generation and SLAKE for VQA. These results demonstrate the flexibility and strong generalization ability of  $A^{3}TUNE$ , along with its capacity to further enhance

Table 4: Performance on SLAKE VQA benchmark using HuatuoGPT-Vision-7B (denoted as **HuatuoGPT-V**). We report both open-ended and close-ended performance.

Madal		Motui							Ν	Aethod					
Model	Mouri		Base	Greedy	Beam	Nucleus	VCD	DoLa	OPERA	AVISC	M3ID	DAMRO	PAI	A <sup>3</sup> Tune	
HuatuoG	PT-V	Open-	ended	85.46	84.89	85.62	85.57	85.57	85.19	85.07	84.42	85.41	85.57	83.85	86.77
+ LoF	RA	Close-	ended	91.27	89.86	89.86	90.99	90.99	90.42	89.86	90.70	91.27	90.99	90.99	91.55
Table 5: Performance on report generation benchmarks using HuatuoGPT-Vision-7B fine-tuned with LoRA.															
Dataset	Motr	ic							Met	hod					
Dataset Metric Huatu + I		Huatuo + L	oGPT-V oRA	Greedy	Beam	Nucleus	VCD	DoLa	OPERA	AVISC	M3ID	DAMRO	PAI	A <sup>3</sup> TUNE	
	BLE	U	8.	.65	9.34	10.21	8.19	9.10	9.03	10.01	7.47	8.35	9.10	6.92	10.52
	ROU	GE-L	27	.34	28.17	28.64	26.28	27.80	27.50	28.57	24.78	27.05	27.80	24.36	28.85
	MET	EOR	31	.48	31.76	34.23	30.72	32.13	31.24	34.10	30.88	32.17	32.13	30.84	36.30
IU-Xray	BER	ГScore	88	.35	88.53	88.60	88.19	88.36	88.39	88.51	87.77	88.19	88.36	87.44	88.67
	CheX	bert	54	.21	55.16	55.84	53.86	54.34	53.89	55.01	52.11	53.78	54.34	50.09	56.27
	Rade	Fraph	21	.35	21.86	22.47	20.17	21.65	21.04	22.59	19.71	21.19	21.65	18.26	23.51
	RaTE	EScore	58	.21	58.66	59.78	58.29	58.29	57.84	59.33	56.49	57.86	58.29	55.16	60.51

performance when built upon more capable backbone models.

## 6 Conclusion

In this work, we present  $A^{3}TUNE$ , a novel finetuning framework designed to enhance the visual grounding capabilities of Med-LVLMs. By leveraging prompt-aware weak labels and integrating a  $A^{3}MOE$  design,  $A^{3}TUNE$  dynamically aligns attention distributions to RoIs across diverse medical tasks and datasets, without requiring inferencetime adjustments. Extensive experiments highlights  $A^{3}TUNE$  as a promising direction for enhancing Med-LVLMs in downstream applications.

## Limitations

While the use of weak labels in  $A^{3}TUNE$  demonstrates its effectiveness, it also introduces noise that can limit performance. In some cases, the model can only focus on generally correct regions, lacking accuracy but providing directions for future research. As shown in Section 5.4.1, using high-quality labels leads to further performance improvements. Additionally, our framework is currently restricted to fine-tuning for downstream tasks, limiting its broader applicability. Furthermore, the metrics used to evaluate visual attention distribution are constrained to patch-level granularity due to the inherent design of Med-LVLMs, rather than achieving pixel-level precision.

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## A Dataset Processing and Statistics

For IU-Xray and OmniVQA in medical VQA task, we utilize the preprocessed datasets provided by the CARES benchmark (Xia et al., 2024), splitting each dataset into training and test sets with a 7:3 ratio. For the MIMIC-CXR dataset used in the report generation task, we randomly sample 2,000 imagereport pairs from the preprocessed MIMIC-CXR-JPG dataset (Johnson et al., 2019) for the training

Task	Dataset	Training	Test
	SLAKE	4,919	1,061
Medical	VQA-RAD	1,797	451
VOA	PathVQA	19,755	6,761
VQA	IU-Xray	1,789	784
	OmniMedVQA	6,155	2,642
Report	IU-Xray	2,069	590
Generation	MIMIC-CXR	1,902	441

Table 6: Dataset statistics for the Medical VQA and Report Generation tasks.

set and 500 pairs for the test set. We extract the "Findings" and "Impression" sections from each report in sampled MIMIC-CXR reports, filtering out those with an extremely low word count. The statistics of those datasets are shown in Table 6.

## **B** Implementation Details of Baselines

Generally, we follow the recommended settings for all baselines while making necessary adjustments to adapt them to Med-LVLMs. The detailed settings are listed as follows:

- Beam Search (Sutskever et al., 2014): The number of beams is set to 5.
- Nucleus Sampling (Holtzman et al., 2020): The top-*p* value for sampling is 0.9.
- VCD (Leng et al., 2024): The contrastive decoding parameters are set to  $\alpha = 1$  and  $\beta = 0.1$ . Diffusion noise is added to images using 500 steps.
- DoLa (Chuang et al., 2023): The mature layer is set to 32, while the early candidate mature layers are [0, 2, 4, 6, 8, 10, 12, 14].
- OPERA (Huang et al., 2024): The number of beams is set to 5, with a scale factor of 50, threshold of 15, and num-attn-candidates = 5. The penalty weight is set to 1. Notably, for LLaVA-Med-1.5 in the report generation task, the scale factor is set to 25 and the threshold is adjusted to 25, as the default values result in nonsensical decoded content.
- AVISC (Woo et al., 2024): We select the top-10 outlier image tokens to construct the negative decoding object. The contrastive decoding parameters are set to  $\alpha = 1$  and  $\beta = 0.1$ .
- M3ID (Favero et al., 2024): The contrastive decoding parameters are set as follows: λ =

Table 7: Fine-tuning epochs and  $\lambda$  values for different datasets.

Dataset	Epochs	$\lambda$							
Medical VQA									
SLAKE	6	0.1							
VQA-RAD	9	0.06							
PathVQA	3	0.02							
IU-Xray	6	0.12							
OmniMedVQA	3	0.03							
Medical Repo	Medical Report Generation								
IU-Xray	12	0.08							
MIMIC-CXR	12	0.05							

0.02 and  $\gamma_t = \exp(-\lambda \cdot t)$ , where t denotes the current decoding step.

- DAMRO (Gong et al., 2024): We select the top-10 tokens with the highest attention to the [CLS] token in the visual encoder as outlier tokens. The contrastive decoding parameters are set to  $\alpha = 0.5$  and  $\beta = 0.1$ .
- PAI (Liu et al., 2025): In the inference intervention, the start layer and end layer are set to 2 and 32, respectively, γ = 1.1 and α = 0.2.
- ControlMLLM (Wu et al., 2024): In inferencetime tuning, it is configured with β = 0.5, α = 400, and a learning rate of 4. For LLaVA-Med-1.5, the same parameters are applied but with a reduced learning rate of 1.

## **C** Implementation Details

## C.1 Hyperparameter setting

All fine-tuning tasks are performed using the same seed for LoRA initialization. The LoRA rank is set to 64 and the rank of each expert in  $A^3MOE$  is set to 16, with a learning rate of 2e-4. For  $A^3MOE$ , the default number of experts in K-MOE  $O^k$  is 8, while the default number of experts in Q-MOE  $O^q$  is 4. For example, we use these settings in all report generation tasks for Chest X-rays. However, for large datasets with diverse image modalities such as SLAKE, PathVQA and OmniMedVQA, the number of experts is increased to 16 and 8, respectively. Some other key hyperparameters are: K = 4, R = 128, B = 3 when  $O^k = 16$  and B = 2 when  $O^k = 8$ .

Madal	Mathad	IU-Xray									
woder	Wiethou	BLEU	ROUGE-L	METEOR	BERTScore	CheXbert	RadGraph	RaTEScore			
	LLaVA-Med	1.30	11.55	16.09	84.12	35.21	6.46	42.32			
	Greedy	1.21	13.97	19.12	84.61	35.82	6.74	42.32			
	Beam	1.19	13.42	19.49	84.24	39.41	8.69	44.56			
	Nucleus	1.33	11.15	16.28	83.89	32.39	6.39	41.39			
	VCD	1.19	10.26	15.46	83.49	33.25	5.67	41.20			
LLaVA-Med	DoLa	1.20	13.93	19.14	84.60	35.61	6.74	42.31			
	OPERA	0.89	8.46	13.90	82.52	33.19	3.48	39.19			
	AVISC	1.16	11.12	17.34	83.86	30.99	5.08	43.30			
	M3ID	1.40	12.01	16.48	84.07	34.32	6.71	42.02			
	DAMRO	1.23	10.69	16.68	83.72	30.21	5.63	43.62			
	PAI	1.14	12.90	18.46	84.33	38.64	6.96	42.92			
	LLaVA-Med + LoRA	7.70	26.15	29.50	88.36	53.81	20.44	58.37			
	Greedy	8.86	27.09	26.01	88.50	52.55	20.76	58.24			
	Beam	9.34	27.56	26.44	88.52	52.88	21.29	58.77			
	Nucleus	7.80	26.72	30.33	88.28	52.73	20.85	57.84			
	VCD	8.83	27.36	31.77	88.30	51.86	22.02	58.93			
LLaVA-Med	DoLa	8.93	26.94	25.74	88.42	52.27	20.63	58.10			
+ LoRA	OPERA	8.27	27.14	29.66	88.41	56.26	21.87	58.73			
	AVISC	8.52	26.97	31.14	88.47	53.33	22.27	59.21			
	M3ID	8.63	27.79	31.65	88.52	54.45	22.22	59.37			
	DAMRO	7.29	25.61	30.03	88.14	51.50	20.75	59.19			
	PAI	8.87	26.74	25.99	88.39	52.45	20.48	57.81			
	A <sup>3</sup> TUNE (ours)	11.05	30.00	34.26	89.05	57.19	24.24	63.03			

Table 8: Full results of report generation on IU-Xray, based on LLaVA-Med and the LoRA fine-tuned LLaVA-Med.



Figure 8: Analysis of the value of  $\lambda$  in A<sup>3</sup>TUNE on the SLAKE dataset.

## C.2 $\lambda$ Selection

In our final loss in Eq. (12), we use a key hyperparameter  $\lambda$  to balance the two loss terms. Here, we conduct an analysis to select the optimal value of  $\lambda$ . Figure 8, illustrate the impact of  $\lambda$ , which controls the strength of attention alignment tuning. We can observe that on SLAKE, performance peaks at  $\lambda = 0.1$ . Beyond this point, the performance declines due to over-alignment and extreme attention distributions. Notably,  $\lambda$  varies with the scale of datasets and the training epoches, and the values of  $\lambda$  and fine-tuning epoches for all the datasets are as shown in Table 7.

# All the experiments are conducted using four A6000 GPUs.

## **D** Metrics

#### **D.1** Metrics for Report Generation

We evaluate model performance using commonly used metrics for generation tasks. These include BERTScore (Zhang et al., 2019), which measures the similarity between the embeddings of predicted and reference texts, and METEOR (Banerjee and Lavie, 2005), which evaluates alignment between generated answers and reference texts, accounting for synonyms and stemming. Additionally, we employ ROUGE-L (Lin, 2004), which measures n-gram overlap and the longest common subsequence, and BLEU (Papineni et al., 2002), which calculates n-gram precision in the predicted text relative to the reference, focusing on exact matches. In addition, we include the following domain-specific metrics designed for medical report generation:

- CheXbert (Smit et al., 2020) is an automatic labeler that extracts pathology indicators from radiology reports. We follow (Yu et al., 2023) to calculate the CheXbert vector similarity that measures the cosine similarity between pathology indicator vectors derived from ground truth and model-generated reports.
- RadGraph (Jain et al., 2021) is a tool that extracts entity and relation from radiology reports. We use RadGraph to specifically

M- J-1	Mathad	MIMIC-CXR									
Model	Method	BLEU	ROUGE-L	METEOR	BERTScore	CheXbert	RadGraph	RaTEScore           32.91           29.81           34.30           34.17           34.06           29.59           36.12           35.22           33.68           35.71           30.57			
	LLaVA-Med	1.38	12.28	13.20	84.24	15.45	3.14	32.91			
	Greedy	0.42	9.23	6.77	83.94	14.23	1.42	29.81			
	Beam	1.20	13.06	13.00	83.25	10.90	3.16	34.30			
	Nucleus	1.22	12.49	11.99	83.65	15.79	3.31	34.17			
	VCD	1.12	12.21	11.53	83.53	14.25	2.83	34.06			
LLaVA-Med	DoLa	0.39	9.13	6.54	83.96	14.26	1.38	29.59			
	OPERA	1.06	12.90	11.93	83.11	12.90	2.80	36.12			
	AVISC	1.20	11.90	13.08	83.20	13.09	2.62	35.22			
	M3ID	1.10	12.13	11.47	83.69	15.08	2.77	33.68			
	DAMRO	1.27	12.40	12.84	83.40	13.76	3.36	35.71			
	PAI	0.48	9.74	7.22	83.81	14.39	1.45	30.57			
	LLaVA-Med + LoRA	3.28	16.54	17.90	85.57	22.14	9.43	40.00			
	Greedy	4.07	18.75	18.81	86.14	24.24	10.73	41.06			
	Beam	3.39	17.25	17.36	85.78	23.05	9.78	38.59			
	Nucleus	3.29	16.14	17.98	85.42	21.76	9.20	39.26			
	VCD	3.53	16.58	18.68	85.57	23.38	9.93	40.95			
LLaVA-Med	DoLa	3.99	18.62	18.68	86.14	25.14	10.75	40.73			
+ LoRA	OPERA	3.14	15.52	14.70	84.75	20.07	7.74	35.72			
	AVISC	3.34	16.79	18.29	85.59	22.74	9.52	40.08			
	M3ID	3.62	16.67	18.22	85.58	22.95	9.93	39.87			
	DAMRO	3.37	15.75	17.03	85.27	22.76	9.40	39.62			
	PAI	4.32	18.64	19.78	86.10	25.78	11.17	41.03			
	A <sup>3</sup> TUNE (ours)	4.56	19.03	20.23	86.17	24.93	11.55	42.73			

Table 9: Full results of report generation on MIMIC-CXR, based on LLaVA-Med and the LoRA fine-tuned LLaVA-Med.

indicate RadGraph F1, which measures the overlap of clinical entities and their relations extracted from ground truth and model-generated reports.

• RaTEScore (Zhao et al., 2024) is a recently proposed metric that prioritizes crucial medical entities, including diagnostic outcomes and anatomical details. This metric is robust to complex medical synonyms and sensitive to negation expressions, aligning more closely with human judgment compared to existing metrics.

#### **D.2** Metrics for Attention Distribution

(a) Coverage Score. The Coverage Score measures the proportion of the ground truth region that is covered by the attention map. Let **B** denote the binary mask of the ground truth segment (where  $\mathbf{B}(i, j) = 1$  for pixels belonging to the ground truth and  $\mathbf{B}(i, j) = 0$  otherwise) and **M** denote the attention map output by the model. The score is defined as:

$$\text{Coverage} = \frac{\sum_{i,j} \mathbf{B}(i,j) \cdot \mathbf{M}_{\tau}(i,j)}{\sum_{i,j} \mathbf{B}(i,j)},$$

where  $\mathbf{M}_{\tau}$  is the thresholded attention map, i.e.,  $\mathbf{M}_{\tau}(i, j) = 1$  if  $\mathbf{M}(i, j) \geq \tau$ , and  $\mathbf{M}_{\tau}(i, j) = 0$  otherwise. This metric quantifies how well the attention aligns spatially with the ground truth. In our experiments, we set  $\tau$  as 0.15.

(b) Intensity Alignment. The Intensity Alignment metric evaluates the average attention intensity within the ground truth region. It is computed as:

Intensity Alignment = 
$$\frac{\sum_{i,j} \mathbf{B}(i,j) \cdot \mathbf{M}(i,j)}{\sum_{i,j} \mathbf{B}(i,j)}$$

This score reflects the degree to which the model focuses its attention on the ground truth segment, considering the intensity values of the attention map.

Table 10: Results on different medical image modalities.

Mathad	Chest	t X-ray	Abdor	nen CT	Brai	n MRI
Wiethou	Open	Closed	Open	Closed	Open	Closed
LLaVA-Med + LoRA	76.96	89.08	79.67	81.15	79.15	79.37
Greedy	79.53	89.92	81.50	83.61	80.57	82.54
Beam	79.83	89.92	81.76	84.43	78.90	84.12
Nucleus	80.95	87.40	79.49	79.51	79.72	80.95
DoLa	78.53	89.92	81.50	83.61	79.89	80.95
VCD	81.39	86.55	80.54	80.33	79.62	69.84
M3ID	79.54	85.71	76.70	80.33	80.57	73.01
AVISC	80.52	87.40	81.41	78.69	80.49	85.71
OPERA	80.19	89.92	81.15	83.61	77.55	82.54
DAMRO	79.23	88.24	80.19	83.61	75.60	80.95
PAI	79.78	89.92	80.45	81.15	81.95	80.95
A <sup>3</sup> TUNE (ours)	82.21	89.92	83.33	84.43	82.41	87.30

# E Full Results on Report Generation Benchmarks

We present the comparison results of report generation using LLaVA-Med in Table 8 (IU-Xray) and Table 9 (MIMIC-CXR). These tables include the original results of baselines applied to LLaVA-Med

Model	Method	DI DI	DOUGE I	METEOR	IU-Xray		<b>D</b> 10	D TEC	
		BLEU	ROUGE-L	METEOR	BERIScore	CheXbert	RadGraph	RaTEScore	
	LLaVA-Med-1.5	1.40	12.41	16.30	84.55	38.20	7.77	41.15	
	Greedy	1.04	12.15	9.87	85.43	38.04	5.43	34.97	
	Beam	1.09	11.17	19.59	83.43	40.13	9.42	48.64	
	Nucleus	1.44	12.10	15.60	81.45	38.04	6.51	40.41	
	VCD	1.42	12.29	15.72	84.54	36.57	6.49	39.93	
LLaVA-Med-1.5	DoLa	0.99	12.15	9.36	85.64	38.22	5.40	34.93	
	OPERA	1.13	11.49	14.63	83.76	37.38	1.41	35.96	
	AVISC	1.18	11.32	16.66	83.76	35.83	6.63	40.36	
	M3ID	1.33	12.31	16.31	84.45	37.54	6.42	40.41	
	DAMRO	1.27	11.56	16.42	84.08	35.60	6.80	40.08	
	PAI	1.11	12.05	10.99	85.03	37.56	5.21	34.83	
	LLaVA-Med-1.5 + LoRA	8.04	26.52	30.37	88.24	51.32	20.35	56.97	
	Greedy	9.36	27.57	27.91	88.55	52.44	21.28	58.61	
	Beam	9.54	28.41	35.40	88.45	53.70	22.43	59.65	
	Nucleus	7.80	26.72	30.33	88.28	52.73	20.85	57.84	
	VCD	8.83	27.36	31.77	88.30	51.86	22.02	58.93	
LLaVA-Med-1.5	DoLa	8.93	26.94	25.74	88.42	52.27	20.63	58.10	
+ LoRA	OPERA	9.23	27.48	34.17	88.17	51.65	21.37	57.89	
	AVISC	5.57	21.71	26.84	87.34	47.32	16.87	53.66	
	M3ID	8.44	26.21	30.86	88.20	51.13	20.77	59.37	
	DAMRO	8.21	25.77	30.58	88.09	50.10	22.33	57.31	
	PAI	8.52	26.97	28.63	88.42	52.22	20.99	58.21	
	A <sup>3</sup> TUNE (ours)	10.51	28.76	35.74	88.51	53.88	23.10	59.66	
A°1 UNE (ours)         10.51         28.76         35.74         88.51         53.88         23.10         59.66           Table 12: Results of Report Generation on MIMIC-CXR									
	Table 12. K	esuits (	of Report	Generation		C-CAR			
	Mathad			Generation	MIMIC-CX				
Model	Method	BLEU	ROUGE-L	METEOR	MIMIC-CX BERTScore	C-CAR KR CheXbert	RadGraph	RaTEScore	
Model	Method LLaVA-Med-1.5	BLEU	ROUGE-L 10.73	METEOR 10.95	MIMIC-CX BERTScore 82.29	<b>KR</b> CheXbert 14.11	RadGraph	RaTEScore 31.90	
Model	Method LLaVA-Med-1.5 Greedy	BLEU 0.98 0.93	ROUGE-L 10.73 10.90	METEOR 10.95 9.45	MIMIC-CX BERTScore 82.29 83.09	<b>C-CAR</b> <b>CheXbert</b> 14.11 14.11	<b>RadGraph</b> 1.83 1.09	<b>RaTEScore</b> 31.90 28.07	
Model	Method LLaVA-Med-1.5 Greedy Beam	BLEU           0.98           0.93           1.13	ROUGE-L 10.73 10.90 10.95	METEOR 10.95 9.45 13.01	MIMIC-CX BERTScore 82.29 83.09 82.34	<b>C-CAR</b> <b>CheXbert</b> 14.11 14.11 12.51	<b>RadGraph</b> 1.83 1.09 1.87	<b>RaTEScore</b> 31.90 28.07 32.86	
Model	Method LLaVA-Med-1.5 Greedy Beam Nucleus	BLEU           0.98           0.93           1.13           0.96	ROUGE-L 10.73 10.90 10.95 10.55	METEOR 10.95 9.45 13.01 10.70	MIMIC-CX BERTScore 82.29 83.09 82.34 78.53	C-CAR CheXbert 14.11 14.11 12.51 14.11	RadGraph 1.83 1.09 1.87 1.64	<b>RaTEScore</b> 31.90 28.07 32.86 31.54	
Model	Method LLaVA-Med-1.5 Greedy Beam Nucleus VCD	BLEU           0.98           0.93           1.13           0.96           1.00	ROUGE-L 10.73 10.90 10.95 10.55 10.93	METEOR 10.95 9.45 13.01 10.70 11.31	MIMIC-CX BERTScore 82.29 83.09 82.34 78.53 83.11	C-CAR           CheXbert           14.11           14.11           12.51           14.11           13.46	<b>RadGraph</b> 1.83 1.09 1.87 1.64 1.97	<b>RaTEScore</b> 31.90 28.07 32.86 31.54 31.75	
Model	Method LLaVA-Med-1.5 Greedy Beam Nucleus VCD DoLa	BLEU           0.98           0.93           1.13           0.96           1.00           0.65	ROUGE-L 10.73 10.90 10.95 10.55 10.93 9.88	METEOR 10.95 9.45 13.01 10.70 11.31 7.83	MIMIC-CX BERTScore 82.29 83.09 82.34 78.53 83.11 83.47	C-CXK           CheXbert           14.11           14.11           12.51           14.11           13.46           14.07	RadGraph 1.83 1.09 1.87 1.64 1.97 1.09	<b>RaTEScore</b> 31.90 28.07 32.86 31.54 31.75 28.07	
Model LLaVA-Med-1.5	Method LLaVA-Med-1.5 Greedy Beam Nucleus VCD DOLa OPERA	BLEU           0.98           0.93           1.13           0.96           1.00           0.65           1.09	ROUGE-L 10.73 10.90 10.95 10.55 10.93 9.88 11.51	METEOR 10.95 9.45 13.01 10.70 11.31 7.83 12.22	MIMIC-CX BERTScore 82.29 83.09 82.34 78.53 83.11 83.47 82.69	C-CAX           CheXbert           14.11           14.11           12.51           14.11           13.46           14.07           13.00	RadGraph 1.83 1.09 1.87 1.64 1.97 1.09 0.52	<b>RaTEScore</b> 31.90 28.07 32.86 31.54 31.75 28.07 27.77	
Model LLaVA-Med-1.5	Method LLaVA-Med-1.5 Greedy Beam Nucleus VCD DoLa OPERA AVISC	BLEU 0.98 0.93 1.13 0.96 1.00 0.65 1.09 1.16	<b>ROUGE-L</b> 10.73 10.90 10.95 10.55 10.93 9.88 11.51 11.14	METEOR 10.95 9.45 13.01 10.70 11.31 7.83 12.22 12.30	MIMIC-CX BERTScore 82.29 83.09 82.34 78.53 83.11 83.47 82.69 82.61	C-CAX           CheXbert           14.11           14.11           12.51           14.11           13.46           14.07           13.00           14.11	RadGraph 1.83 1.09 1.87 1.64 1.97 1.09 0.52 1.97	RaTEScore           31.90           28.07           32.86           31.54           31.75           28.07           27.77           32.20	
Model	Method LLaVA-Med-1.5 Greedy Beam Nucleus VCD DoLa OPERA AVISC M3ID	BLEU           0.98           0.93           1.13           0.96           1.00           0.65           1.09           1.16           1.06	ROUGE-L 10.73 10.90 10.95 10.55 10.93 9.88 11.51 11.14 11.50	METEOR 10.95 9.45 13.01 10.70 11.31 7.83 12.22 12.30 11.54	MIMIC-CX BERTScore 82.29 83.09 82.34 78.53 83.11 83.47 82.69 82.61 83.15	Chexbert           14.11           14.11           12.51           14.11           13.46           14.07           13.00           14.11           13.98	RadGraph 1.83 1.09 1.87 1.64 1.97 1.09 0.52 1.97 2.24	RaTEScore           31.90           28.07           32.86           31.54           31.75           28.07           27.77           32.20           32.74	
Model LLaVA-Med-1.5	Method LLaVA-Med-1.5 Greedy Beam Nucleus VCD DoLa OPERA AVISC M3ID DAMRO	BLEU           0.98           0.93           1.13           0.96           1.00           0.65           1.09           1.16           1.06           1.14	ROUGE-L 10.73 10.90 10.95 10.55 10.93 9.88 11.51 11.14 11.50 11.01	METEOR 10.95 9.45 13.01 10.70 11.31 7.83 12.22 12.30 11.54 12.35	MIMIC-CX BERTScore 82.29 83.09 82.34 78.53 83.11 83.47 82.69 82.61 83.15 82.80	C-CAR           CheXbert           14.11           14.11           12.51           14.11           13.46           14.07           13.00           14.11           13.98           13.73	RadGraph 1.83 1.09 1.87 1.64 1.97 1.09 0.52 1.97 2.24 2.26	RaTEScore           31.90           28.07           32.86           31.54           31.75           28.07           27.77           32.20           32.74           32.18	
Model	Method LLaVA-Med-1.5 Greedy Beam Nucleus VCD DoLa OPERA AVISC M3ID DAMRO PAI	BLEU           0.98           0.93           1.13           0.96           1.00           0.65           1.09           1.16           1.06           1.14	<b>ROUGE-L</b> 10.73 10.90 10.95 10.55 10.93 9.88 11.51 11.14 11.50 11.01 11.67	METEOR           10.95           9.45           13.01           10.70           11.31           7.83           12.22           12.30           11.54           12.35           10.63	MIMIC-CX BERTScore 82.29 83.09 82.34 78.53 83.11 83.47 82.69 82.61 83.15 82.80 82.80	C-CAR           CheXbert           14.11           12.51           14.11           12.51           14.11           13.46           14.07           13.00           14.11           13.98           13.73           14.05	RadGraph 1.83 1.09 1.87 1.64 1.97 1.09 0.52 1.97 2.24 2.26 0.98	RaTEScore           31.90           28.07           32.86           31.54           31.75           28.07           27.77           32.20           32.74           32.18           28.07	
Model LLaVA-Med-1.5	Method LLaVA-Med-1.5 Greedy Beam Nucleus VCD DoLa OPERA AVISC M3ID DAMRO PAI LLaVA-Med-1.5 + LoRA	BLEU           0.98           0.93           1.13           0.96           1.00           0.65           1.09           1.16           1.06           1.14           1.12           3.50	ROUGE-L 10.73 10.90 10.95 10.55 10.93 9.88 11.51 11.14 11.50 11.01 11.67 16.38	METEOR 10.95 9.45 13.01 10.70 11.31 7.83 12.22 12.30 11.54 12.35 10.63 18.95	MIMIC-CX BERTScore 82.29 83.09 82.34 78.53 83.11 83.47 82.69 82.61 83.15 82.80 82.80 82.80 85.56	C-CAR CheXbert 14.11 14.11 12.51 14.11 13.46 14.07 13.00 14.11 13.98 13.73 14.05 21.45	RadGraph 1.83 1.09 1.87 1.64 1.97 1.09 0.52 1.97 2.24 2.26 0.98 9.54	RaTEScore           31.90           28.07           32.86           31.54           31.75           28.07           27.77           32.20           32.74           32.18           28.07           40.45	
Model	Method LLaVA-Med-1.5 Greedy Beam Nucleus VCD DoLa OPERA AVISC M3ID DAMRO PAI LLaVA-Med-1.5 + LoRA Greedy	BLEU           0.98           0.93           1.13           0.96           1.00           0.65           1.09           1.16           1.06           1.14           1.12           3.50	ROUGE-L 10.73 10.90 10.95 10.55 10.93 9.88 11.51 11.14 11.50 11.01 11.67 16.38 16.49	METEOR 10.95 9.45 13.01 10.70 11.31 7.83 12.22 12.30 11.54 12.35 10.63 18.95 18.71	MIMIC-CX BERTScore 82.29 83.09 82.34 78.53 83.11 83.47 82.69 82.61 83.15 82.80 82.80 82.80 85.56 85.54	C-CAR           CheXbert           14.11           14.11           12.51           14.11           12.51           14.11           13.46           14.07           13.00           14.11           13.98           13.73           14.05           21.45           23.43	RadGraph 1.83 1.09 1.87 1.64 1.97 1.09 0.52 1.97 2.24 2.26 0.98 9.54 9.63	RaTEScore           31.90           28.07           32.86           31.54           31.75           28.07           27.77           32.20           32.74           32.18           28.07           40.45           40.49	
Model LLaVA-Med-1.5	Method       LLaVA-Med-1.5       Greedy       Beam       Nucleus       VCD       DoLa       OPERA       AVISC       M3ID       DAMRO       PAI       LLaVA-Med-1.5 + LoRA       Greedy       Beam	BLEU           0.98           0.93           1.13           0.96           1.00           1.6           1.06           1.12           3.50           3.66	<b>ROUGE-L</b> 10.73 10.90 10.95 10.55 10.93 9.88 11.51 11.14 11.50 11.01 11.67 16.38 16.49 16.85	METEOR 10.95 9.45 13.01 10.70 11.31 7.83 12.22 12.30 11.54 12.35 10.63 18.95 18.71 20.68	MIMIC-CX BERTScore 82.29 83.09 82.34 78.53 83.11 83.47 82.69 82.61 83.15 82.80 82.80 82.80 85.56 85.54 85.54	C-CXK           CheXbert           14.11           14.11           12.51           14.11           13.46           14.07           13.00           14.11           13.98           13.73           14.05           21.45           23.43           25.00	RadGraph           1.83           1.09           1.87           1.64           1.97           1.09           0.52           1.97           2.24           2.26           0.98           9.54           9.63           9.91	RaTEScore           31.90           28.07           32.86           31.54           31.75           28.07           27.77           32.20           32.74           32.18           28.07           40.45           40.49           41.46	
Model LLaVA-Med-1.5	Method         LLaVA-Med-1.5         Greedy         Beam         Nucleus         VCD         DoLa         OPERA         AVISC         M3ID         DAMRO         PAI         LLaVA-Med-1.5 + LoRA         Greedy         Beam         Nucleus	BLEU           0.98           0.93           1.13           0.96           1.00           0.65           1.09           1.16           1.06           1.14           3.50           3.66           3.48	<b>ROUGE-L</b> 10.73 10.90 10.95 10.55 10.93 9.88 11.51 11.14 11.50 11.01 11.67 16.38 16.49 16.85 16.35	METEOR           10.95           9.45           13.01           10.70           11.31           7.83           12.22           12.30           11.54           12.35           10.63           18.95           18.71           20.68           18.93	MIMIC-CX BERTScore 82.29 83.09 82.34 78.53 83.11 83.47 82.69 82.61 83.15 82.80 82.80 82.80 85.56 85.54 85.51 85.50	C-CAR           CheXbert           14.11           14.11           12.51           14.11           12.51           14.11           13.46           14.07           13.00           14.11           13.98           13.73           14.05           21.45           23.43           25.00           22.21	RadGraph           1.83           1.09           1.87           1.64           1.97           1.09           0.52           1.97           2.24           2.26           0.98           9.54           9.63           9.91           9.27	RaTEScore           31.90           28.07           32.86           31.54           31.75           28.07           27.77           32.20           32.74           32.18           28.07           40.45           40.45           40.45	
Model LLaVA-Med-1.5	Method LLaVA-Med-1.5 Greedy Beam Nucleus VCD DoLa OPERA AVISC M3ID DAMRO PAI LLaVA-Med-1.5 + LoRA Greedy Beam Nucleus VCD	BLEU           0.98           0.93           1.13           0.96           1.00           0.65           1.09           1.16           1.06           1.14           1.12           3.50           3.66           3.48           3.74	<b>ROUGE-L</b> 10.73 10.90 10.95 10.55 10.93 9.88 11.51 11.14 11.50 11.01 11.67 16.38 16.49 16.85 16.35 16.88	METEOR           10.95           9.45           13.01           10.70           11.31           7.83           12.22           12.30           11.54           12.35           10.63           18.95           18.71           20.68           18.93           19.03	MIMIC-CX BERTScore 82.29 83.09 82.34 78.53 83.11 83.47 82.69 82.61 83.15 82.80 82.60 82.61 83.15 82.80 82.80 85.56 85.54 85.51 85.50 85.56	C-CAR           CheXbert           14.11           14.11           12.51           14.11           12.51           14.11           13.46           14.07           13.00           14.11           13.98           13.73           14.05           21.45           23.43           25.00           22.21           22.98	RadGraph           1.83           1.09           1.87           1.64           1.97           1.09           0.52           1.97           2.24           2.26           0.98           9.54           9.54           9.27           9.56	RaTEScore           31.90           28.07           32.86           31.54           31.75           28.07           27.77           32.20           32.74           32.18           28.07           40.45           40.45           40.45           40.08           40.93	
Model	Method       LLaVA-Med-1.5       Greedy       Beam       Nucleus       VCD       DoLa       OPERA       AVISC       M3ID       DAMRO       PAI       LLaVA-Med-1.5 + LoRA       Greedy       Beam       Nucleus       VCD       DoLa	BLEU           0.98           0.93           1.13           0.96           1.00           0.65           1.09           1.16           1.06           1.14           1.12           3.50           3.66           3.48           3.74           3.48	ROUGE-L 10.73 10.90 10.95 10.55 10.93 9.88 11.51 11.14 11.50 11.01 11.67 16.38 16.49 16.85 16.35 16.45 16.45	METEOR           10.95           9.45           13.01           10.70           11.31           7.83           12.22           12.30           11.54           12.35           10.63           18.95           18.71           20.68           18.93           19.03           18.66	MIMIC-CX BERTScore 82.29 83.09 82.34 78.53 83.11 83.47 82.69 82.61 83.15 82.80 82.60 82.61 83.15 82.80 82.80 85.56 85.54 85.56 85.56 85.54	C-CAR           CheXbert           14.11           14.11           12.51           14.11           13.46           14.07           13.00           14.11           13.98           13.73           14.05           21.45           23.43           25.00           22.21           22.98           23.34	RadGraph           1.83           1.09           1.87           1.64           1.97           0.52           1.97           2.24           2.26           0.98           9.54           9.63           9.91           9.27           9.56           9.52	RaTEScore           31.90           28.07           32.86           31.54           31.75           28.07           27.77           32.20           32.74           32.18           28.07           40.45           40.45           40.08           40.93           40.49	
Model LLaVA-Med-1.5	Method         LLaVA-Med-1.5         Greedy         Beam         Nucleus         VCD         DoLa         OPERA         AVISC         M3ID         DAMRO         PAI         LLaVA-Med-1.5 + LoRA         Greedy         Beam         Nucleus         VCD         DoLa         OPERA	BLEU           0.98           0.93           1.13           0.96           1.00           0.65           1.09           1.16           1.06           1.14           1.12           3.50           3.66           3.48           3.74           3.48	<b>ROUGE-L</b> 10.73 10.90 10.95 10.55 10.93 9.88 11.51 11.14 11.50 11.01 11.67 16.38 16.49 16.85 16.35 16.45 16.45 16.45	METEOR           10.95           9.45           13.01           10.70           11.31           7.83           12.22           12.30           11.54           12.35           10.63           18.95           18.71           20.68           18.93           19.03           18.66           20.10	MIMIC-CX BERTScore 82.29 83.09 82.34 78.53 83.11 83.47 82.69 82.61 83.15 82.80 82.60 82.80 82.80 85.56 85.54 85.50 85.54 85.54 85.54	C-CAR           CheXbert           14.11           14.11           12.51           14.11           12.51           14.11           13.46           14.07           13.00           14.11           13.98           13.73           14.05           21.45           23.43           25.00           22.21           22.98           23.34           24.31	RadGraph  1.83  1.09  1.87  1.64  1.97  1.09  0.52  1.97  2.24  2.26  0.98  9.54  9.63  9.91  9.27  9.56  9.52  9.81	RaTEScore           31.90           28.07           32.86           31.54           31.75           28.07           27.77           32.20           32.74           32.18           28.07           40.45           40.49           41.46           40.93           40.49           41.33	
Model LLaVA-Med-1.5 LLaVA-Med-1.5 + LoRA	Method         LLaVA-Med-1.5         Greedy         Beam         Nucleus         VCD         DoLa         OPERA         AVISC         M3ID         DAMRO         PAI         LLaVA-Med-1.5 + LoRA         Greedy         Beam         Nucleus         VCD         DoLa         OPERA         AVISC	BLEU           0.98           0.93           1.13           0.96           1.00           0.65           1.09           1.16           1.06           1.14           1.12           3.50           3.66           3.48           3.54           3.56           3.31	<b>ROUGE-L</b> 10.73 10.90 10.95 10.55 10.93 9.88 11.51 11.14 11.50 11.01 11.67 16.38 16.49 16.85 16.35 16.35 16.45 16.77 16.36	METEOR           10.95           9.45           13.01           10.70           11.31           7.83           12.22           12.30           11.54           12.35           10.63           18.95           18.71           20.68           18.93           19.03           18.66           20.10           18.64	MIMIC-CX BERTScore 82.29 83.09 82.34 78.53 83.11 83.47 82.69 82.61 83.15 82.80 82.60 82.80 85.56 85.54 85.54 85.54 85.54 85.54 85.54 85.54 85.54	C-CAR           CheXbert           14.11           14.11           12.51           14.11           12.51           14.11           13.46           14.07           13.00           14.11           13.98           13.73           14.05           21.45           23.43           25.00           22.21           22.98           23.34           24.31           23.31	RadGraph           1.83           1.09           1.87           1.64           1.97           2.24           2.26           0.98           9.54           9.63           9.91           9.27           9.56           9.52           9.81           9.02	RaTEScore           31.90           28.07           32.86           31.54           31.75           28.07           22.03           32.74           32.18           28.07           40.45           40.45           40.93           40.93           40.49           41.33           40.36	
Model LLaVA-Med-1.5 LLaVA-Med-1.5 + LoRA	Method         LLaVA-Med-1.5         Greedy         Beam         Nucleus         VCD         DoLa         OPERA         AVISC         M3ID         DAMRO         PAI         LLaVA-Med-1.5 + LoRA         Greedy         Beam         Nucleus         VCD         DoLa         OPERA         AVISC         Main	BLEU           0.98           0.93           1.13           0.96           1.00           1.6           1.06           1.14           3.50           3.66           3.48           3.74           3.48           3.50           3.48           3.74           3.48           3.51	<b>ROUGE-L</b> 10.73 10.90 10.95 10.55 10.93 9.88 11.51 11.14 11.50 11.01 11.67 16.38 16.49 16.85 16.35 16.88 16.45 16.77 16.36 16.13	METEOR           10.95           9.45           13.01           10.70           11.31           7.83           12.22           12.30           11.54           12.35           10.63           18.95           18.93           19.03           18.66           20.10           18.64           18.52	MIMIC-CX BERTScore 82.29 83.09 82.34 78.53 83.11 83.47 82.69 82.61 83.15 82.80 82.60 85.56 85.54 85.56 85.54 85.56 85.54 85.54 85.48 85.48 85.39	C-CXX           CheXbert           14.11           14.11           12.51           14.11           13.46           14.07           13.00           14.11           13.98           13.73           14.05           21.45           23.43           25.00           22.21           22.98           23.34           24.31           23.31           22.42	RadGraph           1.83           1.09           1.87           1.64           1.97           1.64           1.97           2.24           2.26           0.98           9.54           9.54           9.52           9.51           9.52           9.81           9.02           9.15	RaTEScore           31.90           28.07           32.86           31.54           31.75           28.07           27.77           32.20           32.74           32.18           28.07           40.49           41.46           40.08           40.93           40.49           41.33           40.36           39.74	
Model LLaVA-Med-1.5 LLaVA-Med-1.5 + LoRA	Method         LLaVA-Med-1.5         Greedy         Beam         Nucleus         VCD         DoLa         OPERA         AVISC         M3ID         DAMRO         PAI         LLaVA-Med-1.5 + LoRA         Greedy         Beam         Nucleus         VCD         DoLa         OPERA         AVISC         Mathematical Structure         VCD         DoLa         OPERA         AVISC         M3ID         DAMRO	BLEU           0.98           0.93           1.13           0.96           1.00           1.6           1.06           1.14           3.50           3.66           3.48           3.74           3.48           3.51           3.14           3.42	<b>ROUGE-L</b> 10.73 10.90 10.95 10.55 10.93 9.88 11.51 11.14 11.50 11.01 11.67 16.38 16.49 16.85 16.35 <u>16.88</u> <u>16.45</u> 16.77 16.36 <u>16.36</u> 16.13 16.63	METEOR           10.95           9.45           13.01           10.70           11.31           7.83           12.22           12.30           11.54           12.35           10.63           18.95           18.71           20.68           18.93           19.03           18.66           20.10           18.64           18.52           18.87	MIMIC-CX BERTScore 82.29 83.09 82.34 78.53 83.11 83.47 82.69 82.61 83.15 82.80 82.60 82.60 85.56 85.54 85.51 85.50 85.56 85.54 85.56 85.54 85.56 85.54 85.48	C-CAR           CheXbert           14.11           14.11           12.51           14.11           12.51           14.11           13.46           14.07           13.00           14.11           13.98           13.73           14.05           21.45           23.43           25.00           22.21           22.98           23.34           24.31           23.31           22.42           23.30	RadGraph           1.83           1.09           1.87           1.64           1.97           1.09           0.52           1.97           2.24           2.26           0.98           9.54           9.63           9.27           9.56           9.52           9.81           9.02           9.15           9.46	RaTEScore           31.90           28.07           32.86           31.54           31.75           28.07           22.03           27.77           32.20           32.74           32.18           28.07           40.45           40.45           40.45           40.45           40.45           40.45           40.45           40.45           40.45           40.33           40.36           39.74           40.80	
Model LLaVA-Med-1.5 LLaVA-Med-1.5 + LoRA	Method         LLaVA-Med-1.5         Greedy         Beam         Nucleus         VCD         DoLa         OPERA         AVISC         M3ID         DAMRO         PAI         LLaVA-Med-1.5 + LoRA         Greedy         Beam         Nucleus         VCD         DoLa         OPERA         AVISC         M3ID         DALa         OPERA         AVISC         M3ID         DAMRO         PAI	BLEU           0.98           0.93           1.13           0.96           1.00           0.65           1.09           1.16           1.06           1.14           1.12           3.50           3.66           3.48           3.74           3.48           3.56           3.31           3.14           3.42           3.63	<b>ROUGE-L</b> 10.73 10.90 10.95 10.55 10.93 9.88 11.51 11.14 11.50 11.01 11.67 16.38 16.49 16.35 16.85 16.35 16.88 16.45 16.36 16.13 16.63 16.65	METEOR           10.95           9.45           13.01           10.70           11.31           7.83           12.22           12.30           11.54           12.35           10.63           18.95           18.71           20.68           18.93           19.03           18.66           20.10           18.64           18.52           18.87           18.61	MIMIC-CX BERTScore 82.29 83.09 82.34 78.53 83.11 83.47 82.69 82.61 83.15 82.80 82.60 85.56 85.54 85.51 85.50 85.56 85.54 85.56 85.54 85.56 85.54 85.56 85.54 85.56 85.54 85.48 85.39 85.48 85.60	C-CAR           CheXbert           14.11           14.11           12.51           14.11           12.51           14.11           13.46           14.07           13.00           14.11           13.98           13.73           14.05           21.45           23.43           25.00           22.21           22.98           23.34           24.31           23.31           22.42           23.30           24.51	RadGraph           1.83           1.09           1.87           1.64           1.97           1.09           0.52           1.97           2.24           2.26           0.98           9.54           9.53           9.54           9.52           9.51           9.52           9.51           9.52           9.51           9.52           9.51           9.52           9.81           9.02           9.15           9.46           9.60	RaTEScore           31.90           28.07           32.86           31.54           31.75           28.07           27.77           32.20           32.74           32.18           28.07           40.45           40.45           40.45           40.45           40.45           40.45           40.45           40.45           40.45           40.45           40.33           40.36           39.74           40.80           40.49	

Table 11: Results on report generation benchmarks, based on LLaVA-Med-1.5.



Figure 9: VQA Types where  $A^{3}TUNE$  outperforms LLaVA-Med + LoRA across three image modalities.

without fine-tuning, where the model performs significantly worse on report generation. For example, in Table 8, the best-performing baseline, M3ID, achieves a BLEU score of 1.40, which is much lower than fine-tuned LLaVA-Med baselines. This highlights the challenge of generating professional medical reports without fine-tuning. As discussed in the experiments section, A<sup>3</sup>TUNE consistently achieves the best performance, outperforming all baselines by a large margin.

## F Fine-grained Effectiveness Analysis

In this experiment, we conduct fine-grained analyses to evaluate the effectiveness of our proposed



Figure 10: Case study on LLaVA-Med-1.5. The red box in the **Input Image** (not provided as input to the model) highlights the RoI that model should focus on. Red texts and green texts indicate wrong answers and correct answers, respectively.

method across different medical image modalities. Specifically, we examine how our approach improves model performance on Chest X-ray, Abdomen CT, and Brain MRI images from SLAKE. As shown in Table 10, our method outperforms all baselines, particularly on Brain MRI. These results demonstrates the effectiveness of  $A^{3}TUNE$  across diverse medical images and its generalization ability in medical applications. To explain this improvement, we also include case studies with attention map visualizations for each modality in Section 5.5.

Furthermore, we analyze the VQA types where our method outperforms fine-tuned LLaVA-Med with LoRA across the three image modalities. As shown in Figure 9, our approach improves the model's performance in three key areas sensitive to image information:

- General Radiology Knowledge: Understanding medical modality types and their features.
- Anatomical Structures: Recognizing features of key anatomical structures, such as organ count, location.
- Abnormalities: Identifying and diagnosing abnormalities based on features like location and color.

These improvements highlight the effectiveness of attention tuning in handling VQA types relying on accurate visual information and interaction.

## G Analysis on LLaVA-Med-1.5

# G.1 Attention Biases and Hallucination Issues in LLaVA-Med-1.5

The attention biases we address are not unique to LLaVA-Med but are prevalent across Med-LVLMs. For instance, we include cases from LLaVA-Med-1.5 in this section. Although LLaVA-Med-1.5 is an enhanced version of LLaVA-Med, with improved training data and an increased number of visual tokens (from 256 to 576), the attention biases and hallucination issues persist.

In Figure 10, we visualize the attention biases and corresponding hallucination issues in LLaVA-Med-1.5. As shown, the model often fails to focus on the correct RoIs and generates hallucinated outputs. Even with attention tuning via ControlM-LLM, hallucinations persist. For example, in Case (B.3), while the model identifies an abnormality in the liver, it incorrectly classifies it as a liver abscess instead of liver cancer.

These attention biases are common issues in Med-LVLMs, underscoring the need for continued research on emergent Med-LVLMs to mitigate such challenges effectively.

## G.2 Experiment Results on LLaVA-Med-1.5

Similar to the LLaVA-Med experiments in Section 5.3, Table 11 and Table 12 show that A<sup>3</sup>TUNE outperforms all baselines across almost all metrics on both IU-Xray and MIMIC-CXR, excelling in both language quality and clinical accuracy. These results underscore A<sup>3</sup>TUNE' effectiveness in medical applications across diverse Med-LVLMs, demonstrating its strong generalization ability.