DrVD-Bench: Do Vision-Language Models Reason Like Human Doctors in Medical Image Diagnosis?

Abstract

Vision-language models (VLMs) exhibit strong zero-shot generalization on natural images and show early promise in interpretable medical image analysis. However, existing benchmarks do not systematically evaluate whether these models truly reason like human clinicians or merely imitate superficial patterns. To address this gap, we propose DrVD-Bench, the first multimodal benchmark for clinical visual reasoning. DrVD-Bench consists of three modules: Visual Evidence Comprehension, Reasoning Trajectory Assessment, and Report Generation Evaluation, comprising a total of 7,789 image-question pairs. Our benchmark covers 20 task types, 17 diagnostic categories, and five imaging modalities—CT, MRI, ultrasound, radiography, and pathology. DrVD-Bench is explicitly structured to reflect the clinical reasoning workflow from modality recognition to lesion identification and diagnosis. We benchmark 19 VLMs, including general-purpose and medicalspecific, open-source and proprietary models, and observe that performance drops sharply as reasoning complexity increases. While some models begin to exhibit traces of human-like reasoning, they often still rely on shortcut correlations rather than grounded visual understanding. DrVD-Bench offers a rigorous and structured evaluation framework to guide the development of clinically trustworthy VLMs. **Dataset:** Kaggle, Hugging Face Code: GitHub

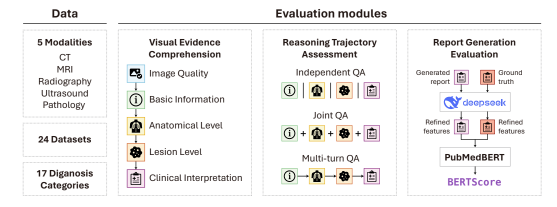


Figure 1: Overview of the DrVD-Bench

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1 Introduction

The lack of interpretability in AI-driven diagnostic systems has long been a critical barrier to their adoption in clinical practice [87, 36]. Vision-Language Models (VLMs) [29, 18, 14, 27, 74, 35, 90] offer a promising direction toward interpretable medical AI, as they can generate outputs that mimic the stepwise clinical reasoning process of physicians [72, 13, 66]. In clinical settings, physicians arrive at diagnosis by progressively integrating information across multiple levels—ranging from basic image features to anatomical structures and lesion characteristics—before synthesizing this evidence step-by-step into a clinical diagnostic report. Despite the increasing adoption of VLMs in analyzing medical images, there exists no benchmark that systematically evaluates whether these models follow a similar reasoning trajectory or merely rely on memorized patterns and spurious shortcuts.

To address this gap, we introduce DrVD-Bench (Doctor-like Visual Diagnosis Benchmark), the first benchmark explicitly designed to evaluate the understanding and reasoning capabilities of VLMs in medical image analysis. DrVD-Bench comprises three complementary modules: (i) *Visual Evidence Comprehension*, containing 4,480 high-quality image–question pairs structured to mirror the each step of clinical reasoning, assessing how well models identify essential visual cues and generate diagnoses; (ii) *Reasoning Trajectory Assessment*, consisting of 487 images with 3,321 question–answer turns that emulate the progressive reasoning process of clinicians and evaluate whether models reason step by step or rely on shortcuts; and (iii) *Report Generation Evaluation*, comprising 475 questions aimed at assessing holistic understanding through free-form clinical report generation.

Modules 1 and 2 are explicitly designed to reflect the clinical reasoning workflow—from modality recognition to anatomical localization, lesion characterization, and final diagnosis. This workflow is defined as follows:

- Level 0: Image Quality Assessment Detecting artifacts or noise; typically handled during acquisition.
- Level 1: Basic Information Extraction Identifying modality, body region, view, etc.; often available as metadata in clinical settings.
- Level 2: Anatomy-Level Recognition Recognizing and localizing anatomical structures (e.g., organs, tissues).
- Level 3: Lesion-Level Identification Detecting, localizing, and describing abnormalities such as lesions or fractures.
- Level 4: Clinical Interpretation Integrating visual findings to produce a diagnosis or generate a structured report.

By organizing tasks into stepwise levels, DrVD-Bench enables fine-grained analysis of VLM performance—not only in terms of final diagnostic predictions, but also in their capacity to extract intermediate visual evidence and engage in coherent reasoning. This structure allows us to examine whether models genuinely analyze medical images by identifying key visual cues and reasoning in a step-by-step manner, akin to human clinicians. In the *Visual Evidence Comprehension* module, we introduce specially designed *organ erasure* and *lesion erasure* tasks, which compel models to rely on visible image content rather than memorized associations, thereby mitigating potential information leakage from pretraining. The *Reasoning Trajectory Assessment* module includes three distinct QA formats—*Independent QA*, *Joint QA*, and *Multi-turn QA*, —with the latter specifically enforcing a stepwise reasoning pattern that mirrors clinical workflows. Except for this module, all other stages adopt an independent QA format, where each question is presented in isolation without shared context. Finally, unlike previous benchmarks that limit report generation to specific modalities [41, 75, 93] such as radiographs or CT scans, our *Report Generation Evaluation* module introduces a setting to assess models' holistic understanding across diverse imaging modalities.

We evaluated 19 publicly available VLMs, including both general purpose and medical-specific models (see Table 3). Our key findings are summarized as follows:

• Reasoning performance declines with task complexity: VLMs perform well on low-level visual tasks such as modality or view recognition, but their accuracy drops substantially on higher-level tasks involving anatomical understanding, lesion localization, and diagnosis.

- Correct answers without supporting evidence: Many models achieve higher diagnostic accuracy than their performance on lesion-level tasks, suggesting they can produce correct diagnoses without fully understanding or localizing the supporting visual evidence.
- Limited capacity for stepwise clinical reasoning: Models perform best when provided with all questions at once (Joint QA), but struggle in Multi-turn QA, suggesting difficulty with maintaining dialogue state and reasoning trajectories.
- Hallucinations in report generation: In free-form generation tasks, models often produce plausible but unsupported statements, revealing challenges in grounding clinical language in image evidence.
- Smaller, specialized models can compete: While larger and newer models generally perform better, domain-optimized models demonstrate strong performance relative to their scale, highlighting the value of medical-specific alignment.

2 Related Works

2.1 Vision-Language Models

Modern vision-language models (VLMs) build upon the reasoning capabilities and world knowledge of large language models (LLMs) by aligning visual inputs with the textual domain [90]. For instance, LLaVA [45] introduces a multi-layer perceptron (MLP)[59] to bridge a vision encoder (e.g., CLIP [62]) with a language model backbone, enabling the system to perform tasks such as interpreting scientific figures [82] and understanding cartoons [47]. VLMs have also been adapted for the medical domain [42, 89, 39]. For example, LLaVA-Med [39], derived from LLaVA, is fine-tuned on medical data to adapt the model for healthcare-specific tasks. Although VLMs have shown impressive performance on general visual reasoning tasks [16, 88, 32, 51], it remains uncertain whether they truly comprehend medical images or merely rely on prior knowledge and pattern matching [17, 77, 60, 37].

2.2 Medical VLM Benchmarks

The application of VLMs in medicine demands benchmarks with broad coverage and fine-grained, clinically relevant evaluation. However, most existing benchmarks remain narrow in scope (see Table 1), typically limited to single modalities or task types. For example, PathMMU [73], VQA-RAD [38], and PMC-VQA [92] focus primarily on visual question answering (VQA) within specific imaging domains, limiting generalizability and diagnostic depth.

Recent benchmarks introduce hierarchical structures [58, 85, 31], but these do not align with the stepwise nature of clinical reasoning. GMAI-MMBench [85], for instance, organizes questions by perceptual complexity rather than reasoning stages. OmniMedVQA [31] covers diverse tasks like modality recognition and diagnosis but lacks a clinically grounded task progression, grouping questions only by type. As a result, current benchmarks fall short in evaluating whether VLMs reason like clinicians. A critical gap remains: we lack a benchmark that not only measures answer correctness but also reveals how and why models succeed or fail—essential for assessing clinical reasoning ability.

Table 1: Comparison of Medical VLM Benchmarks

Benchmark	Imaging modalities	Task hierarchy	Clinical reasoning	Task types
VQA-RAD[38]	Radiography, CT	Х	Х	VQA
SLAKE[44]	Radiography, CT, MRI	X	X	VQA
PMC-VQA [92]	CT, MRI, and others	X	X	VQA
Rad-ReStruct [58]	Radiography	\checkmark	X	VQA
GMAI-MMBench[85]	CT, MRI, Radiography, Ultrasound, Pathology	\checkmark	X	VQA
PathMMU[73]	Pathology	X	X	VQA
OmniMedVQA[31]	12 modalities	\checkmark	X	VQA
CARES[79]	16 modalities	X	X	VQA
MultiMedEval[68]	≥ 11 modalities	X	X	VQA, open QA, and others
DrVD-Bench	5 modalities	✓ (stepwise)	\checkmark	VQA, report generation

3 Design of DrVD-Bench

3.1 Overview

We propose DrVD-Bench, a multi-scale benchmark for systematically evaluating vision—language models (VLMs) in the medical domain. Inspired by the diagnostic workflow of clinicians, DrVD-Bench defines a three-module framework that evaluates VLMs from three aspects: (i) Reliability in visual evidence extraction; (ii) Ability in stepwise clinical reasoning; and (iii) Comprehensive understanding of medical image-revealed by the ability in report generation. Representative examples are shown in Figure 2.

DrVD-Bench comprises three modules: (1) **Visual Evidence Comprehension**, containing 4,480 expert-curated image—question pairs across 16 tasks, structured by the depth of clinical reasoning, from the superficial modality recognition to the deep lesion-level identification and diagnosis. To reduce reliance on shortcuts, we introduce **organ** and **lesion erasure** tasks that force models to reason from visible evidence. (2) **Reasoning Trajectory Assessment**, with 3,321 QA turns on 487 images, evaluates whether models reason step by step using three prompting formats: **Joint QA**, **Independent QA**, and **Multi-turn QA**. (3) **Report Generation Evaluation**, spanning all the five modalities, it contains 475 questions aimed at assessing holistic understanding through free-form clinical report generation. See Appendix A.2 for the detailed composition of our benchmark.

Together, these components enable fine-grained analysis of VLMs' clinical visual understanding and diagnostic reasoning.

3.2 Dataset Collection and Task Construction

3.2.1 Data Collection

This study systematically aggregates multi-modal medical images (CT, MRI, ultrasound, X-ray, and pathology) from 24 publicly available datasets and online repositories (See Table 8 in Appendix A). These sources encompass a wide range of imaging scales, including panoramic, organ-level, and histopathological views. Only images with a resolution of at least 256×256 are retained to ensure sufficient visual clarity for structural and anatomical interpretation;

3.2.2 Task Overview

To enable fine-grained evaluation of models' intermediate reasoning in medical image analysis, DrVD-Bench organizes all QA tasks (except report generation) into a five-level hierarchy that mirrors the cognitive stages of clinical diagnosis. Each level targets a distinct reasoning step:

- Level 0 (Image Quality): noise and artifact detection.
- Level 1 (Basic Information): modality, view, body part, magnification, stain, and imaging technique recognition.
- Level 2 (Anatomy Level): organ/tissue identification, localization, and organ-erasure detection.
- Level 3 (Lesion Level): lesion recognition, lesion-erasure detection, and morphological description.
- Level 4 (Clinical Interpretation): diagnostic classification.

The *Visual Evidence Comprehension* module includes tasks from all levels, with each image paired with a single QA focused on one aspect of visual understanding. In contrast, the *Reasoning Trajectory Assessment* module evaluates multi-level reasoning per image by combining one task from each level (e.g., modality, body part, organ, lesion, diagnosis). It offers three prompting formats: **Joint QA** presents all questions at once; **Independent QA** asks them sequentially; and **Multi-turn QA** incorporates the model's prior response into subsequent prompts, simulating stepwise reasoning.

3.2.3 Dataset and QA Pair Construction

To support diverse evaluation objectives, we employ task-specific dataset construction strategies. For noise robustness, we simulate modality-specific clinical noise at three PSNR levels—15, 25, and 35

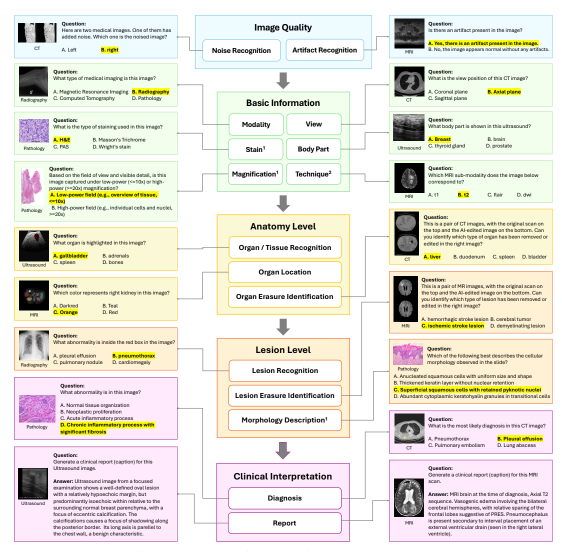


Figure 2: Hierarchical five-level evaluation framework for medical imaging diagnostics. Representative tasks—spanning CT, MRI, ultrasound, radiography, and pathology—are shown alongside their corresponding stage in the clinical reasoning cascade. Tasks labeled with ¹ are exclusive to pathology, and those with ² are exclusive to MRI.

dB—representing severe, moderate, and mild interference per clinical standards. Recognition and localization tasks include bounding box and region annotations for relevant anatomical or pathological areas. To assess sensitivity to local visual cues, we introduce an erasure task: using the Bailian high-throughput image platform, targeted anatomical or lesion regions are digitally removed while preserving structural continuity. All modified images are manually verified to ensure artifact-free erasure and reliable evaluation of visual evidence dependence.

The *Reasoning Trajectory Assessment* module builds on the *Visual Evidence Comprehension* question format but varies in structure and prompting. Independent QA pairs one image with multiple questions at different reasoning levels. Joint QA bundles sub-questions from multiple levels into a single prompt for simultaneous response. Multi-turn QA sequentially passes earlier responses into higher-level questions, simulating stepwise reasoning (See Appendix Figure 12).

For *Report Generation Evaluation*, we frame the task as medical captioning, using curated ground-truth references from PubMedVision [15] and PathMMU [73] that provide detailed clinical explanations.

All OA pairs are authored by physicians based on task objectives (e.g., modality recognition, localization, diagnosis) and span both closed-form and open-ended formats. Each is blind-reviewed by a senior clinician to ensure clinical relevance, visual grounding, and diagnostic realism. Distractors are carefully curated to remain plausible, and all questions are self-contained, unambiguous, and visually answerable. Data distribution is summarized in Table 2.

Experiments

4.1 Experiment Setup

In this study, we evaluated a total of 19 models (Table 3), including general-purpose open-source models, proprietary models accessed via API calls, and fine-tuned medical vision models. The parameter sizes of open-source models range from 7B to 72B. All experiments were conducted under a standardized zero-shot evaluation framework using system prompts on 8×NVIDIA A100 GPUs (80GB each). This setup ensures consistent and fair comparisons across different model architectures while maintaining reliable performance benchmarking.

Table 2: Statistics of DrVD Panch

Table 2: Statistics of I	DrVD-Bench		Table 3: VLMs benchmar	ked in our sti	ıdy
Category	Metric	Count	Model	Developer	Year
Module 1: Visual Evidence Co	mprehension		Proprietary		
Total QA pairs		4,480	GPT-4o[54]	OpenAI	2024.11
Level 0: Image Quality	QA pairs	591	GPT-o1[55]	OpenAI	2024.12
Level 1: Basic Information	QA pairs	1,400	GPT-o3[53]	OpenAI	2025.04
Level 2: Anatomy	QA pairs	1,151	Gemini 2.5 Pro[25]	Google	2025.03
Level 3: Lesion	QA pairs	890	Grok-3[78]	xAI	2025.02
Level 4: Clinical Interpretation	QA pairs	923	Doubao1.5-VisionPro [11]	ByteDance	2025.01
Module 2: Reasoning Trajecto	rv Assessment		Claude 3.7 Sonnet[4]	Anthropic	2025.02
Total images		487	Open-source		
Independent QA	QA pairs	2,347	Qwen2.5-VL [6]	Alibaba	2025.01
Joint QA	QA pairs	487	Phi-4 14B [1]	Microsoft	2024.12
Multi-turn QA	QA pairs	487	GLM-4V [23]	Tsinghua	2024.06
CT/MRI/Radiography/US	images each	100	Janus-Pro-7B [19]	DeepSeek	2025.01
Pathology	images	87	Medical-specific		
Module 3: Report Generation	Evaluation		HuatuoGPT-Vision-34B [15]	CUHK	2024.06
Open-ended QA pairs		475	HealthGPT-L14B [42]	ZJU	2025.02
			RadFM-14B [76]	PJLAB	2023.12
Global Dataset Statistics			LLaVA-Med-7B [39]	Microsoft	2024.04
Total QA pairs (all modules)		7,789			
Organ/Tissue classes		38			
Lesion classes		27			
Diagnosis categories		17			

4.2 Evaluation

In DrVD-Bench, we use accuracy to evaluate multiple-choice tasks, each with a single correct answer. If the model's output conforms to the expected format, we directly compare it with the ground truth to determine correctness. For responses that deviate from the format, we apply DeepSeek-V3 [43] to extract the selected option. If extraction fails, the answer is marked as incorrect (See Appendix D). All results are averaged over five independent runs to ensure robustness. Accuracy is computed per question and then averaged within each level to obtain level-specific scores.

For open-ended tasks such as report generation, we employ DeepSeek-V3 [43] to extract key features from both the model's response and the reference text. We then adopt BERTScore [91] with PubMedBERT [26] to capture biomedical semantics of the generated data. To enable consistent comparison, we normalize BERTScore using the baseline (See Appendix B) and best-performing model. Let s_{model} be the model's BERTScore, s_{baseline} the score of an irrelevant response, and s_{best}

the highest score observed. The normalized score is:

$$\label{eq:Normalized BERTScore} \text{Normalized BERTScore} = \frac{s_{\text{model}} - s_{\text{baseline}}}{s_{\text{best}} - s_{\text{baseline}}}$$

In addition, to further evaluate factual consistency and hallucination in report generation, we incorporate FActScore [50] as a complementary metric. FActScore first extracts atomic-level clinical facts from both the reference and generated reports using DeepSeek-v3, followed by fact-level matching with GPT-4o. We compute two indicators: recall, which measures the proportion of true facts correctly captured by the model, and specificity, which measures the proportion of generated facts supported by the reference. Low recall and specificity respectively reflect insufficient factual coverage and the presence of hallucinated content.

5 Results Analysis

5.1 Models' Performance on Visual Evidence Comprehension

5.1.1 Models Perform Differently Across Task Levels

Table 4 presents the evaluation results for CT modality tasks, with results for other modalities provided in Appendix B. Across all models, we observe a consistent decline in performance as the reasoning level increases. While most models perform well on basic recognition tasks such as identifying the imaging modality or view, their accuracy drops markedly on tasks requiring organ-level understanding, and declines even further on lesion-level reasoning. For instance, GPT-o3 achieves 86% accuracy on Basic Information tasks, decreases to 66% on Organ-Level tasks, and further falls to 41% on Lesion-Level tasks. This trend reflects a clear gap between surface-level visual parsing and clinically meaningful reasoning. As tasks increasingly demand multi-step inference and integration of both global context and local features, model performance becomes less stable and less reliable. These results highlight the limitations of current VLMs in replicating the fine-grained, layered reasoning processes central to clinical diagnosis.

Table 4: Accuracy of different VLMs across different task levels. The best-performing scores are highlighted in red, and the second-best in blue. Due to the large number of tasks across imaging modalities, we only present the performance of different models on task levels here. Results for the detailed subtasks among five modalities (CT, MRI, Ultrasound, Radiography, and Pathology) are provided in Appendix B. These are recognition subtasks across reasoning levels.

Model	Image Quality	Basic Info	Anatomy Level	Lesion Level	Diagnosis								
Random	49	29	27	25	24								
Proprietary													
GPT-40	68	84	57	50	54								
GPT-o1	56	71	44	37	39								
GPT-o3	69	86	66	41	48								
Claude 3.7 Sonnet	68	83	60	42	48								
Gemini 2.5 Pro	76	88	65	52	54								
Grok-3	63	78	56	45	51								
Doubao-VisionPro	63	82	52	59	52								
Qwen-VL-MAX	65	78	54	56	53								
		Open-source	e										
Qwen2.5-VL-72B	65	77	54	56	52								
LLaVA-1.6-34B	61	60	38	49	46								
Qwen2.5-VL-32B	61	73	51	48	56								
Phi-4-14B	70	68	39	44	47								
GLM-4V-9B	65	70	43	32	36								
Qwen2.5-VL-7B	68	69	41	45	38								
Janus-Pro-7B	59	68	44	39	5 6								
		Medical-spec	ific										
HuaTuoGPT-Vision-34B	61	85	58	54	59								
HealthGPT-L14B	56	77	46	41	53								
RadFM-14B	52	61	33	38	31								
LLaVA-Med-7B	52	49	34	32	29								

5.1.2 Overdiagnosis without Understanding

Notably, as shown in Table 4, many models perform better on diagnosis tasks than on lesion recognition, revealing a disconnect between output accuracy and reasoning fidelity. In other words, models can produce clinically plausible diagnostic results without actually identifying the supporting lesion evidence, a phenomenon we term "overdiagnosis without understanding".

This gap likely stems from biases in training data. Most VLMs are trained on image—report pairs that provide final diagnoses (e.g., "pneumonia", "fracture") but omit intermediate steps such as lesion localization or characterization. As a result, models tend to learn global pattern-to-label mappings, bypassing the step-by-step reasoning process that underpins clinical decision-making.

The issue becomes especially pronounced in zero-shot settings that demand fine-grained lesion-level reasoning. As shown in Table 11 (Appendix B), in CT lesion erasure detection, GPT-o3 and Gemini 2.5 Pro perform worse than random (19% and 16%, respectively), and achieve only 28% accuracy on lesion-level tasks—indicating a failure to recognize missing lesion evidence. Yet, on CT diagnosis tasks, they attain substantially higher accuracy (52% and 71%, respectively). This stark discrepancy suggests that diagnostic conclusions can be produced without properly grounding in anatomically or lesion-relevant visual features.

This behavior raises fundamental concerns about the reliability and clinical validity of current models. Despite producing seemingly accurate outputs, many models struggle with evidence-based reasoning, especially when visual grounding is essential to safe and explainable diagnosis.

5.2 Models Prefer Global Context over Step-by-Step Clinical Reasoning

Our benchmark reveals a clear performance hierarchy across the three reasoning settings (Table 5): Joint QA achieves the best overall results, followed by Independent QA, while Multi-turn QA performs the worst. In the Joint QA setting, where the model receives the full sequence of questions at once, it can reason more effectively by integrating global context and avoiding cumulative errors. Independent QA, where each question is asked separately without memory of previous turns, provides more stable but fragmented reasoning, leading to limited performance in higher-order tasks such as lesion identification and diagnosis. Surprisingly, Multi-turn QA—which retains previous questions and answers to simulate a realistic step-by-step clinical reasoning process—results in the weakest performance. This suggests that current models struggle with managing dialogue state and are vulnerable to propagating errors over turns. Overall, these findings indicate that today's VLMs benefit more from static, comprehensive context than from dynamic, trajectory-based reasoning.

To further verify this, we conducted a controlled experiment to isolate the effect of error accumulation from other factors such as missing image references or prompt formatting. We designed three conditions:

- (1) **Baseline** the model reasons based on its own previous answers;
- (2) **Insert-Correct** lower-level answers are replaced with ground-truth labels;
- (3) **Insert-Error** lower-level answers are replaced with random incorrect labels.

As shown in Table 6, results on three representative models (Qwen2.5-VL-72B, Claude 3.7 Sonnet, and GPT-40) reveal that diagnosis accuracy improves notably when earlier answers are corrected (e.g., +28 for Qwen) and drops when errors are inserted (-15). Intermediate tasks (Body Part and Organ) change only slightly (±6%), suggesting that dependencies intensify at deeper reasoning stages. These results confirm that cumulative errors—rather than prompt design or forgetting—are the main cause of performance degradation in multi-turn reasoning.

5.3 Hallucinations and Reasoning Errors in Clinical Report Generation

Figure 9 and Table 17 (Appendix B) shows the report scores of various models across five imaging modalities. Gemini 2.5 Pro, HuatuoGPT, and Claude 3.7 Sonnet demonstrate strong performance across multiple modalities, with leading overall scores. Figure 3 further illustrates the performance of the highest-scoring model, Gemini 2.5 Pro, under different scenarios: examples A and B are high-scoring cases that successfully identify key lesions and generate structured descriptions, but still exhibit occasional diagnostic errors or hallucinations. In contrast, example C is a low-scoring

Table 5: **Performance across modality, bodypart, organ, lesion, and diagnosis (recognition subtasks across levels in module 1).** Each cell shows accuracy in the format: **Independent/Multiturn/Joint**. The highest value(s) are **bolded**, second highest are <u>underlined</u>. Missing values are shown as –.

Model	Modality	Bodypart	Organ	Lesion	Diagnosis
GPT-40	99/99/99	86 / <u>85</u> /–	<u>58</u> /55/ 65	<u>45/45/54</u>	<u>40/40/41</u>
Claude	99/99/99	<u>82</u> / 83 /–	<u>53</u> /52/ 70	38/36/ 51	33/30/ 38
Gemini 2.5 Pro	100/100/ 99	90 / <u>78</u> /–	<u>62</u> /47/ 76	<u>51</u> /38/ 61	<u>48</u> /35/ 55
Qwen2.5-VL-72B	99/99/99	85/85/-	<u>57</u> /56/ 68	<u>44</u> / 47/47	<u>36</u> /34/ 38
Grok-3	99/99/99	87 / <u>59</u> /–	<u>56</u> /37/ 62	<u>38</u> /20/ 46	<u>33</u> /18/ 35

Table 6: **Impact of low-level answer accuracy on multi-turn QA performance.** Each cell shows accuracy under three conditions: **Baseline/Insert-Correct/Insert-Error**. The highest value(s) are **bolded**, and the second highest are <u>underlined</u>.

Model	Modality	Body Part	Organ	Lesion	Diagnosis
Qwen2.5-VL-72B	99/99/99	85 / <u>84</u> / <u>84</u>	57/58/55	47/ 60 /43	37/ 65 /22
Claude 3.7 Sonnet	99/99/99	<u>83</u> / 84 /77	52/52/46	36/ 50 /32	30/ 60 /19
GPT-40	99/99/99	<u>85</u> / 86 /85	55/55/54	45/ 52 /44	40/ 67 /33

case, where most of the generated content consists of hallucinated findings unrelated to the reference diagnosis. These results suggest that while current models have made noticeable progress in medical image-based reasoning and diagnosis, they still struggle to fully eliminate inaccuracies and hallucinations during the reasoning process, limiting their clinical reliability.

To quantify this issue, we evaluate four representative models—Gemini 2.5 Pro, HuaTuoGPT-Vision-34B, Qwen2.5-VL-72B, and Claude 3.7 Sonnet—using FActScore. As shown in Table 7, all models exhibit low recall and low specificity, indicating that they capture only a small fraction of clinically relevant facts while generating substantial hallucinated content unrelated to the reference reports or underlying images. These findings further demonstrate that factual consistency remains a key bottleneck for current VLMs in clinical report generation.

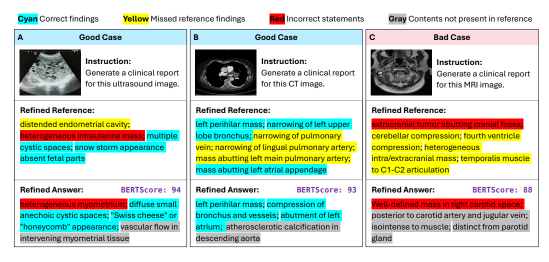


Figure 3: Report generation examples of Gemini 2.5 Pro. A and B represent high-scoring examples, while C represents bad-scoring examples

5.4 Bigger, Newer, or Specialized: Factors Drive Model Performance

Model performance on DrVD-Bench is shaped by both scale and recency. Large proprietary models such as Gemini 2.5 Pro and GPT-o3 lead the overall leaderboard, confirming the advantages of

Table 7: FActScore results: factual consistency of clinical report generation. Recall measures the coverage of true facts, while specificity reflects the proportion of generated facts supported by the reference report. Low values indicate persistent hallucination issues across models.

Model	Recall (std)	Specificity (std)
Gemini 2.5 Pro	0.13 (0.16)	0.07 (0.10)
Claude 3.7 Sonnet	0.06 (0.11)	0.03 (0.05)
Qwen2.5-VL-7B	0.08 (0.13)	0.05 (0.09)
HuatuoGPT-Vision-34B	0.13 (0.16)	0.09 (0.17)

scale—especially for complex reasoning and fine-grained visual tasks. Meanwhile, newer models consistently outperform older ones at similar sizes (Figure 4), highlighting the impact of improved architectures, training pipelines, and data quality on overall effectiveness.

Yet scale and recency are not the whole story. While proprietary models dominate, open-source models like Qwen2.5-VL-72B perform competitively despite having fewer parameters. More notably, HuaTuoGPT-Vision-34B achieves the second-highest accuracy with fewer than half the parameters of top-tier models. Its success demonstrates that domain-specific optimization—when aligned with medical structure and semantics—can enable smaller models to rival or even surpass larger general-purpose systems in clinical reasoning.

6 Conclusion

We present DrVD-Bench, a hierarchical, multimodal benchmark designed to assess whether VLMs reason like human clinicians. Covering five imaging modalities, 20 task types, and 7,789 QA pairs, DrVD-Bench captures the full spectrum of clinical visual reasoning—from basic information tasks like modality recognition to lesion identification and diagnosis. While many VLMs perform well on surface-level recognition tasks, their accuracy drops sharply as reasoning complexity increases. Notably, some models generate plausible diagnoses without correctly identifying supporting visual evidence, revealing a disconnect between diagnostic output and evidence-based understanding. By explicitly targeting intermediate reasoning steps and simulating clinical workflows, DrVD-Bench shows that current VLMs show early signs of clinical reasoning, but still far from truly interpreting medical images like human doctors. Our benchmark offers a structured foundation for developing clinically reliable and visually grounded medical AI systems.

7 Limitations

Although DrVD-Bench is clinically inspired, it remains a controlled evaluation without real-world patient context (e.g., notes, disease progression), limiting its reflection of actual diagnostic workflows. Thus, results should not be overinterpreted for clinical translation, and future refinements should better capture real-world complexity.

As DrVD-Bench uses public datasets, some recent models may have seen parts of them during pre-training, leading to potential train–test overlap and affecting performance trends. Besides, given current VLM limitations, selecting one "most informative" 2D slice from 3D or 4D volumes is a practical compromise but remains a common constraint. Moreover, most training datasets (Table 10 in Appendix A) originate from large institutions,

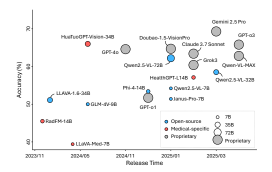


Figure 4: Performance of VLMs on DrVD-Bench visual evidence comprehension tasks across different scales and recencies.

underrepresenting minority groups and smaller or resource-limited hospitals. Future benchmarks should better control data leakage, enable volumetric reasoning, and reduce data bias to improve fairness and generalizability.

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A Technical Appendices and Supplementary Material

A.1 Dataset Availability

DrVD-Bench is publicly available at the following links:

- Kaggle: https://www.kaggle.com/datasets/tianhongzhou/drvd-bench
- Hugging Face: https://huggingface.co/datasets/jerry1565/DrVD-Bench

The code used to conduct our experiments is released at https://github.com/Jerry-Boss/DrVD-Bench.

A.2 Detailed Composition of Our Benchmark

In this section, we describe the task compositions and dataset usage in more detail. Our benchmark contains 3 modules, Module 1 (Visual Evidence Comprehension) contains 4480 QA pairs, Module 2 (Reasoning Trajectory Assessment) contains 3321 QAs, and Module 3 (Report Generation Evaluation) contains 475 questions (See Figure 5). The compositions of Module 1 is illustrated in Figure 6, and that of Module 2 is illustrated in Figure 7. The levels and tasks of Module 1 are listed in detail in Table 9.

Tasks of Module 2 (Reasoning Trajectory Assessment) are adapted from Module 1 (Visual Evidence Comprehension), But they are differently organized. They are organized into 3 forms using different prompts (See Appendix C). Notably, there are 4 questions per Joint QA, and 5 questions per Multi-turn QA.

In Joint QA, the number of answer options is carefully designed to reduce prompt-induced information leakage. Both modality and organ questions contain 4 options each, while lesion and diagnosis questions are expanded to 8 options (i.e., 4×2), mitigating the information leak from our prompts.

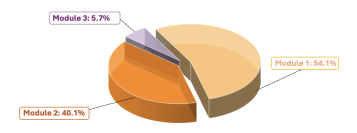


Figure 5: Diagram for the composition of our benchmark, which contains 3 modules



Figure 6: Diagram for the composition of Module 1. Module 1 spans all five modalities, and is designed to contain tasks of five levels



Figure 7: Diagram for the composition of Module 2. Module 2 contains three parts: Independent QA, Joint QA and Multi-turn QA.

Table 8: Datasets used in DrVD-Bench, organized by imaging modality.

Modality Dataset Name Radiography (X-ray) MIMIC-CXR v4[34] MURA[64] VinDr-SpineXR[52]	
MURA[64]	
Panoramic Dental Radiograph SA-Med2D-20M [84] RSNA-Pneumonia[3]	hy / Tufts Dental Dataset[56]
CT SA-Med2D-20M [84] AMOS 2022 [33] DeepLesion[81] CT-RATE[28] PubMedVision [15] LiTS (Liver Tumor Segmenta COVID-CTset [63] CTPelvic1k [46] CT-ORG [65] StructSeg2019-subtask1 [70] KiTS 2021 [30] LNDb [57] MSD-Liver [5, 71] COVID-19	
MRI TotalSegmentator MRI[21] BraTS 2020[49, 7, 8] BraTS 2021 [49, 7, 8] PI-CAI [69] LLD-MMRI [48] MICCAI 2024 CARE MyoPS ISPY1-Tumor-SEG-Radiomic	
Ultrasound BUSI[2] CardiacUDA[83] CuRIOUS2022[80, 9] TG3K [24] Abdominal Ultrasound Image Annotated Ultrasound Liver I	
Pathology PathMMU[73]	

Table 9: Task QA numbers in Module 1 (continued on next page)

Modality	Level	Task	Number
	Image Quality	Noise Recognition Artifact Recognition	100 100
	Basic Information	Modality View Body Part	100 100 100
CT	Anatomy Level	Organ Recognition Organ Location Organ Erasure Identification	100 99 98
	Lesion Level	Lesion Recognition Lesion Erasure Identification	95 100
	Clinical Interpretation	Report Diagnosis	100 90
	Image Quality	Noise Recognition Artifact Recognition	100 61
Radiography	Basic Information	Modality View Body Part	100 100 100
	Anatomy Level	Organ Recognition Organ Location Organ Erasure Identification	100 100 68
	Lesion Level	Lesion Recognition Lesion Erasure Identification	95 100
	Clinical Interpretation	Report Diagnosis	100 100
	Image Quality	Artifact Recognition	46
	Basic Information	Modality Body Part	100 100
Ultrasound	Anatomy Level	Organ Recognition Organ Erasure Identification	100 100
	Lesion Level	Lesion Recognition Lesion Erasure Identification	100 100
	Clinical Interpretation	Report Diagnosis	75 63
	Basic Information	Stain Magnification	100 100
Pathology	Anatomy Level	Organ/Tissue Recognition	100
	Lesion Level	Morphology Description	100
	Clinical Interpretation	Report Diagnosis	100 100

(continued)

Modality	Level	Task	Number
	Image Quality	Noise Recognition	100
	image Quanty	Artifact Recognition	84
MRI		Modality	100
	Basic Information	View	100
	Basic Illiorniation	Technique	100
		Body Part	100
WIN		Organ Recognition	100
	Anatomy Level	Organ Location	86
	·	Organ Erasure Identification	100
	Lesion Level	Lesion Recognition	100
	Lesion Level	Lesion Erasure Identification	100
	Clinical Intermediation	Report	100
	Clinical Interpretation	Diagnosis	95

Table 10: Training datasets for medical-Specific VLMs. List of domain-specific vision—language models and their corresponding medical training datasets.

Medical-specific VLM	Medical training datasets
HuatuoGPT-Vision-34B	PubMedVision
HealthGPT-L14B	PubMedVision, LLaVA-Med, PathVQA, MIMIC-CXR-VQA, SLAKE, VQA-RAD
RadFM-14B	Rad3D-series, MPx-series, PMC-Inline, PMC-CaseReport, VinDr-Mammo, VinDr-SpineXR, VinDr-PCXR, PMC-OA, PMC-VQA, VQA-RAD, SLAKE, MIMIC-CXR, VinDr-CXR, NIH ChestXray14, CheXpert, Covid-CXR2, NLM-TB, Object-CXR, OpenI
LLaVA-Med-7B	PMC-15M

B Detailed Results

In this section, we show our benchmark results in detail.

For Module 1 (Visual Evidence Comprehension), the result for CT is in Table 11; the result for Radiography is in Table 12; the result for MRI is in Table 13; the result for Ultrasound is in Table 14; the result for Pathology is in Table 15.

We also analyzed model performance across different imaging modalities (Figure 8). VLMs perform best on Pathology and worst on Ultrasound. The strong results in Pathology may stem from the fact that we did not use whole-slide images; instead, human annotators selected diagnostically relevant regions and zoomed in, effectively reducing task complexity. In contrast, Ultrasound poses a unique challenge due to its inherently dynamic nature—clinical interpretation typically relies on real-time video sequences rather than static frames, making single-image reasoning considerably more difficult.

We evaluated the performance of different models on the report generation task by comparing their outputs against clinically relevant references (See Figure 9 and Table 17). The performance differences across modalities is minimal.

To establish a baseline, we introduced a set of medically plausible but image-irrelevant texts and computed their BERTScores against the ground-truth references. Each reference was paired with a randomly selected sentence from the list below, and the resulting BERTScore served as the normalization baseline:

- No focal consolidation is seen. However, based on clinical history, the findings may suggest a prior episode of viral gastroenteritis.
- There is no acute intracranial hemorrhage. The patient's recent weight loss should be evaluated further with laboratory studies.
- No pulmonary embolism is identified. The patient's chronic insomnia is unlikely to be explained by these imaging findings.
- Normal appearance of abdominal organs. Note: elevated serum calcium should be correlated with parathyroid hormone levels.
- No significant degenerative changes are observed. Patient's dizziness may be related to recent changes in medication dosage.
- Imaging of the chest is unremarkable. Recommend thyroid function tests given the history of fatigue and cold intolerance.
- No mass lesion is detected. Given the positive ANA, autoimmune evaluation is advised.
- The study is negative for acute pathology. Further investigation is warranted for the reported night sweats and low-grade fever.
- No obstructive uropathy is evident. Patient's lab findings of hematuria require correlation with urine cytology.
- Brain MRI is within normal limits. Symptoms of memory loss may be functional in origin or related to recent psychosocial stressors.

Table 11: Results of different VLMs across different VQA tasks in visual evidence tasks for the CT modality. The best-performing scores are bolded, and the second-best are underlined.

<u>'</u>															
Model	In Overall	nage Qualit Artifact	y Noise	Overall	Basic Info Modality	rmation Bodypart	View	Overall	Organ Recognition	Level Location	Erasure	Overall	Lesion Level Recognition	Erasure	Clinical Interpretation Diagnosis
Random	48	51	45	30	25	27	39	29	29	27	30	26	25	26	24
							I	Proprietary							
GPT-4o	72	52	92	89	100	85	92	50	60	46	44	38	39	36	67
GPT-o1	62	51	72	86	94	78	87	35	39	34	33	29	30	27	46
GPT-o3	80	65	94	94	100	90	92	70	76	77	56	28	37	19	52
Claude 3.7 Sonnet	75	52	97	84	100	84	92	47	60	46	36	29	28	29	46
Gemini 2.5 Pro	82	64	99	93	100	88	92	<u>67</u>	<u>67</u>	73	62	28	39	16	71
Grok-3	65	54	76	66	100	88	43	44	58	33	41	35	36	33	58
Doubao1.5-VisionPro	78	68	87	90	98	81	90	50	59	52	38	55	61	49	59
Qwen-VL-MAX	74	61	87	87	100	85	89	54	55	58	49	<u>45</u>	42	<u>47</u>	68
							C	pen-source							
Qwen2.5-VL-72B	75	59	90	88	100	87	89	54	57	61	45	45	42	47	69
LLaVA-1.6-34B	62	53	70	72	80	85	51	32	42	16	38	43	41	45	54
Qwen2.5-VL-32B	59	50	68	88	97	80	86	45	48	41	47	39	46	32	68
Phi-4-14B	75	50	99	67	99	76	57	36	45	27	36	41	35	46	64
GLM-4V-9B	66	49	83	74	86	83	52	36	44	27	36	33	29	36	38
Qwen2.5-VL-7B	71	57	84	76	100	79	73	37	40	34	36	40	42	38	39
Janus-Pro-7B	71	52	90	82	94	88	63	37	45	32	34	41	40	42	73
							Ме	dical-speci	fic						
HuaTuoGPT-Vision-34B	67	60	74	94	100	91	92	52	63	48	44	33	47	33	68
HealthGPT-L14B	51	49	53	84	100	90	84	38	52	30	32	32	41	22	<u>72</u>
RadFM-14B	52	47	56	72	92	56	67	30	39	29	22	44	45	42	35
LLaVA-Med-7B	57	53	60	55	82	57	26	33	32	36	32	34	33	35	30

Table 12: Results of different VLMs across different VQA tasks in visual evidence tasks for the radiography modality. The best-performing scores are bolded, and the second-best are <u>underlined</u>.

Model		nage Qualit			Basic Info			1	Organ				Lesion Level		Clinical Interpretation
	Overall	Artifact	Noise	Overall	Modality	Bodypart	View	Overall	Recognition	Location	Erasure	Overall	Recognition	Erasure	Diagnosis
Random	49	51	46	27	29	25	28	30	18	44	27	25	28	22	21
Proprietary															
GPT-4o	69	70	67	92	100	95	81	59	46	79	53	36	45	27	27
GPT-o1	60	61	59	82	91	74	81	51	49	57	48	33	40	25	18
GPT-o3	72	74	69	92	100	98	79	76	80	94	53	33	41	25	24
Claude 3.7 Sonnet	62	62	61	95	100	91	93	66	81	65	53	38	47	28	36
Gemini 2.5 Pro	79	70	88	88	100	97	68	71	80	89	43	53	57	48	25
Grok-3	64	72	55	91	99	85	90	55	37	74	53	32	36	28	32
Doubao-VisionPro	53	44	61	90	100	89	82	60	84	77	18	37	43	30	42 27
Qwen-VL-MAX	66	68	64	85	100	93	63	71	80	78	56	37	37	36	27
							()pen-source	e						
Qwen2.5-VL-72B	65	63	67	76	100	63	64	70	79	78	52	41	48	34	28
LLaVA-1.6-34B	62	72	51	69	96	65	46	37	38	62	10	25	27	23	31
Qwen2.5-VL-32B	64	62	66	67	100	63	37	68	75	78	52	33	38	27	29
Phi-4-14B	77	70	84	75	100	90	34	43	39	73	16	31	34	27	37
GLM-4V-9B	66	64	68	81	100	89	53	50	56	66	28	28	29	26	20
Qwen2.5-VL-7B	75	78	72	73	98	89	33	46	53	61	25	33	36	30	37
Janus-Pro-7B	57	46	67	73	99	88	33	39	37	73	6	33	35	30	24
							Me	dical-speci	fic						
HuaTuoGPT-Vision-34B	63	66	59	81	100	63	79	58	67	75	32	35	42	28	37
HealthGPT-L14B	60	70	49	75	98	60	68	39	32	66	19	32	37	27	34
RadFM-14B	60	70	49	66	72	58	68	39	34	51	31	24	17	30	44
LLaVA-Med-7B	51	49	52	51	70	51	32	44	28	60	44	28	25	31	24

Table 13: **Results of different VLMs across different VQA tasks in visual evidence tasks for the MRI modality.** The best-performing scores are **bolded**, and the second-best are <u>underlined</u>.

	•														
Model	In Overall	nage Qualit	y Noise	Overall	Basic In Modality	formation Bodypart	Imaging	Overall	Organ Recognition	Level Location	Erasure	Overall	Lesion Level Recognition	Erasure	Clinical Interpretation
Random	49	55	42	27	21	29	24	34	24	22	29	20	28	25	30
								oprietary							
GPT-40	66	51	80	72	100	68	55	66	52	66	45	46	58	53	57
GPT-o1	50	33	67	56	78	46	36	65	35	38	29	38	44	46	40
GPT-o3	62	40	84	76	100	69	62	71	53	66	49	43	51	50	48
Claude 3.7 Sonnet	70	54	86	65	100	56	43	60	49	56	38	<u>53</u>	56	61	43
Gemini 2.5 Pro	70	45	94	80	99	74	81	67	54	58	46	57	55	55	57
Grok-3	60	45	75	64	98	65	37	55	51	64	35	53	50	51	50
Doubao-VisionPro	62	56	68	64	99	53	45	60	46	61	38	40	61	54	45
Qwen-VL-MAX	62	45	79	61	96	58	37	54	43	48	36	46	65	68	60
							Op	en-source							
Qwen2.5-VL-72B	65	50	79	63	96	60	40	54	46	54	36	49	67	71	58
LLaVA-1.6-34B	62	50	74	46	57	43	29	56	35	36	24	45	68	60	76
Qwen2.5-VL-32B	60	49	71	62	91	60	39	57	44	45	38	48	63	61	<u>70</u>
Phi-4-14B	66	43	88	55	97	47	25	51	31	33	24	36	57	49	44
GLM-4V-9B	65	50	79	50	93	55	17	34	39	39	30	47	35	29	37
Qwen2.5-VL-7B	62	45	78	51	95	52	24	34	42	49	29	47	54	46	39
Janus-Pro-7B	59	38	79	57	87	68	22	50	43	45	44	40	36	31	79
							Med	ical-specifi	e						
HuaTuoGPT-Vision-34B	57	40	73	77	100	<u>70</u>	65	73	49	63	36	47	68	67	67
HealthGPT-L14B	60	67	53	61	98	63	25	58	43	50	35	43	54	54	42
RadFM-14B	46	50	42	49	94	31	38	33	43	58	32	40	55	50	35
LLaVA-Med-7B	51	59	43	43	71	34	28	38	25	31	17	27	36	34	32

Table 14: Results of different VLMs across different VQA tasks in visual evidence tasks for the ultrasound modality. The best-performing scores are bolded, and the second-best are underlined.

Model	Image Quality	В	asic Informa	rmation		Organ Level			Lesion Level		Clinical Interpretation
	Artifact	Overall	Modality	Bodypart	Overall	Recognition	Erasure	Overall	Recognition	Erasure	Diagnosis
Random	53	25	22	27	24	23	25	22	23	20	20
					Propri	etary					
GPT-40	50	76	99	53	44	37	51	54	76	31	<u>62</u>
GPT-o1	35	61	89	32	35	27	43	35	44	26	43
GPT-o3	46	82	99	65	46	36	55	36	36	36	48
Claude 3.7 Sonnet	48	71	100	41	54	46	61	28	36	19	49
Gemini 2.5 Pro	<u>50</u>	<u>81</u>	100	<u>61</u>	48	<u>40</u>	<u>56</u>	59	58	59	51
Grok-3	54	75	98	52	45	36	54	53	74	32	64
Doubao-VisionPro	48	71	100	42	35	31	39	83	87	78	49
Qwen-VL-MAX	39	67	98	35	37	33	40	<u>69</u>	76	62	52
					Open-s	ource					
Qwen2.5-VL-72B	30	66	98	33	36	33	38	63	74	52	46
LLaVA-1.6-34B	<u>50</u>	54	77	31	32	22	42	58	<u>83</u>	32	51
Qwen2.5-VL-32B	<u>50</u>	59	94	24	32	28	36	34	36	32	52
Phi-4-14B	37	66	95	36	26	25	27	43	49	36	41
GLM-4V-9B	<u>50</u>	75	100	49	29	30	28	17	23	10	33
Qwen2.5-VL-7B	<u>50</u>	47	92	2	30	32	28	46	36	56	25
Janus-Pro-7B	15	69	98	40	30	29	30	26	47	4	60
					Medical-	specific					
HuaTuoGPT-Vision-34B	46	82	98	65	44	33	55	64	63	64	64
HealthGPT-L14B	<u>50</u>	67	94	40	35	36	34	25	4	45	51
RadFM-14B	<u>50</u>	47	60	33	20	25	14	36	60	12	11
LLaVA-Med-7B	44	50	58	41	23	21	25	31	30	32	30

Table 15: Results of different VLMs across different VQA tasks in visual evidence tasks for the pathology modality. The best-performing scores are bolded, and the second-best are <u>underlined</u>.

Model	Ba	asic Information		Organ Level	Lesion Level	Clinical Interpretation	
	Overall	Magnification	Stain	Recognition	Morphology	Diagnosis	
Random	34	46	21	30	23	25	
			Proprie	etary			
GPT-40	88	88	88	80	62	57	
GPT-o1	72	73	71	65	43	50	
GPT-o3	88	89	87	<u>83</u>	56	<u>66</u>	
Claude 3.7 Sonnet	<u>94</u>	<u>96</u>	91	<u>83</u>	61	67	
Gemini 2.5 Pro	96	99	93	86	63	67	
Grok-3	83	83	83	<u>83</u>	55	49	
Doubao-VisionPro	93	94	91	69	58	63	
Qwen-VL-MAX	86	84	87	65	63	58	
		(Open-so	ource			
Qwen2.5-VL-72B	86	85	87	66	<u>64</u>	59	
LLaVA-1.6-34B	60	56	63	53	51	49	
Qwen2.5-VL-32B	88	89	87	66	69	59	
Phi-4-14B	69	65	72	59	46	47	
GLM-4V-9B	69	73	65	61	48	51	
Qwen2.5-VL-7B	89	93	84	50	50	50	
Janus-Pro-7B	58	70	46	73	57	44	
		M	edical-s	specific			
HuaTuoGPT-Vision-34B	93	93	93	86	61	57	
HealthGPT-L14B	91	93	89	75	<u>64</u>	64	
RadFM-14B	73	60	85	32	33	30	
LLaVA-Med-7B	47	56	38	43	32	27	

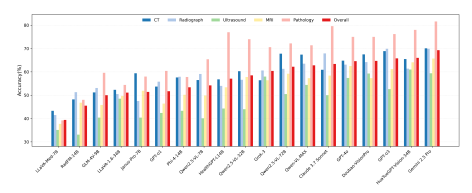


Figure 8: Performance of VLMs across five medical imaging modalities in visual evidence tasks, sorted by overall accuracy.

Table 16: **Performance of VLMs across five medical imaging modalities in visual evidence tasks.** The best-performing scores are **bolded**, and the second-best are <u>underlined</u>.

Model	СТ	Radiography	Ultrasound	MRI	Pathology	Overall		
		Propriet	ary					
GPT-40	64.8	63.1	57.4	62.5	75	64.6		
GPT-o1	53.7	55.8	42.4	46.4	60.4	51.7		
GPT-o3	68.9	69.9	52.6	61.2	76.2	65.8		
Claude 3.7 Sonnet	60.9	<u>67.9</u>	50	58.4	<u>79.6</u>	63.4		
Gemini 2.5 Pro	70.1	69.9	<u>59.4</u>	65.7	81.6	69.3		
Grok-3	56.4	60.6	58	56.4	70.6	60.4		
Doubao-1.5-VisionPro	67.5	64.2	59.3	57.3	75	64.7		
Qwen-VL-MAX	67.4	63.5	54.4	57.3	71.4	62.8		
Open-source								
Qwen2.5-VL-72B	67.8	61.3	50.5	59.2	72.2	62.2		
LLaVA-1.6-34B	52.3	50.5	48.5	49.7	54.4	51.1		
Qwen2.5-VL-32B	60.3	56.6	44	57.8	74	58.5		
Phi-4-14B	57.6	57.9	43.3	50.2	57.8	53.4		
GLM-4V-9B	51.2	53.1	40.4	45.8	59.6	50.0		
Qwen2.5-VL-7B	56.5	59.1	40.1	49.9	65.4	54.2		
Janus-Pro-7B	59.4	47.5	40.4	51.9	58	51.4		
Medical-specific								
HuaTuoGPT-Vision-34B	65.5	61.4	61	64.1	78	<u>66</u>		
HealthGPT-L14B	56.8	54	44.3	53.4	77	57.1		
RadFM-14B	48.2	51.3	33.1	46.8	48	45.5		
LLaVA-Med-7B	43.3	41.5	35.1	37.7	39.2	39.4		

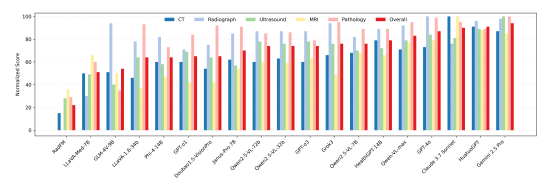


Figure 9: Normalized scores of report generation across five imaging modalities (sorted by overall scores).

Table 17: **BERTScore performance of vision-language models across five imaging modalities.** Best-performing scores are bolded, and second-best scores are underlined. Models follow visual grouping.

Model	CT	Radiography	Ultrasound	MRI	Pathology	Overall			
Proprietary									
GPT-4o	73/89.42	100/92.09	84/89.98	79/89.11	99/90.20	87/90.16			
GPT-o1	60/88.96	71/90.79	69/89.39	42/87.90	84/89.69	65/89.35			
GPT-o3	60/88.97	87/91.52	78/89.73	63/88.60	79/89.52	74/89.67			
Claude 3.7 Sonnet	100/90.35	76/91.02	81/89.85	100/89.80	95/90.08	90/90.22			
Gemini 2.5 Pro	87/89.89	98/91.98	100/90.61	85/89.30	100/90.25	94/90.41			
Grok3	66/88.90	94/91.07	76/89.35	49/87.52	95/89.99	76/89.77			
Doubao1.5-VisionPro	54/87.67	75/89.76	64/88.91	42/87.45	92/89.73	65/88.70			
Qwen-VL-max	71/89.71	92/91.01	79/89.62	77/89.38	95/89.84	83/89.91			
Open-source									
Qwen2.5-VL-72b	60/87.85	87/89.99	78/89.53	60/87.50	85/89.71	74/88.92			
LLaVA-1.6-34b	46/85.32	78/88.66	64/88.34	37/86.76	93/89.81	64/87.78			
Qwen2.5-VL-32b	63/88.26	87/89.99	76/89.31	59/87.43	86/89.72	74/88.94			
Phi-4-14B	60/87.84	82/89.12	58/87.46	47/87.04	73/88.26	64/87.74			
GLM-4V-9B	51/86.37	94/91.05	40/86.36	50/87.23	35/85.48	54/87.30			
Qwen2.5-VL-7B	68/88.98	82/89.12	70/88.91	68/88.61	89/89.88	76/89.50			
Janus-Pro-7B	62/88.15	85/89.79	58/87.10	54/87.28	91/89.89	70/88.84			
Medical-specific									
HuatuoGPT	91/91.40	96/91.65	89/89.06	88/88.38	89/89.37	91/90.94			
HealthGPT-14B	79/90.08	89/88.79	72/88.26	66/88.09	89/89.08	79/88.86			
RadFM	15/81.46	0/80.00	29/82.85	36/83.64	29/82.87	22/82.17			
LLaVA-Med-7B	50/85.04	30/83.01	49/84.94	66/88.09	60/86.01	51/85.13			

C Prompts

In this section, we present the exact prompts used in our experiments. To ensure model compliance, particularly with recent models that may reject clinical queries, all prompts are framed within an educational context.

C.1 Multiple-Choice Problems

System prompt:

You are a helpful assistant participating in an educational visual reasoning task. Always follow the instructions strictly.

User prompt:

 $\textbf{Question:}\ \{\,\}$

Options:

A) {}

B) {}

C) {}
D) {}

Instructions:

Choose the SINGLE best answer by replying with one capital letter (A–D). Do not explain. Do not add extra text.

C.2 Independent QA

You are participating in an educational exercise based on visual information.

Below is an image typically used in medical training, followed by a question.

Question: {}

Options:

A) {}

B) {}

C) {}

D) {}

Instructions: Based only on the visible structures and general anatomical knowledge, choose the most appropriate answer by replying with a single capital letter (A, B, C, or D). Do not explain your reasoning.

C.3 Joint QA

System prompt:

You are participating in a step-by-step medical reasoning diagnosis task based on interpretation of a medical image.

User prompt:

Please answer the following four questions sequentially. Each question builds upon the reasoning of the previous one. Carefully analyze the image and select the most appropriate answer at each step. For each question, choose one capital letter (A, B, C, or D). Do not skip any step.

1. What imaging modality is used in this image?

Options:

A. {}

B. {}

C. {}

D. (

D. {}

2. Which organ appears to be abnormal in this image?

Options:

A. {}

B. {}

C. {} D. {} 3. Based on the abnormal organ, what lesion or finding is most clearly visible? Options: A. {} B. {} C. {} D. {} E. {} F. {} G. {} H. {}
4. Considering all the above findings, what is the most likely diagnosis? Options: A. {} B. {} C. {} D. {} E. {} F. {} G. {} H. {} Instructions:
Please reply with your four selected letters in order, separated by commas (e.g. A,C,B,A). Do not provide explanations.
Multi-turn QA
System prompt: You are an expert medical AI. You will answer several step-by-step questions about the same medical image. Respond with one capital letter (A–H).
User (Round 1): (with image) 1. What imaging modality is used in this image? Options: A) CT B) MRI C) Ultrasound D) Radiography Assistant: A
User: [Same image QA record] 1. What imaging modality is used in this image? Assistant: CT
User (Round 2): 2. Which organ is shown in this medical image? Options: A) Liver B) Pancreas C) Spleen D) Kidney Assistant: A
User: [Same image QA record] 2. Which organ is shown in this medical image? Assistant: Liver
User (Round 3): 3. What lesion is visible in this image? Options: A) Target sign B) Hepatic steatosis C) Pancreatic pseudocyst D; Splenic infarct

C.4

Assistant:

Α

User (Round 4):

4. What is the most likely diagnosis?

Options: A) Crohn's disease B) Acute pancreatitis C) Splenic rupture D) Diverticulitis

C.5 Report Generation

System prompt:

Generate a clinical report based on the image. This is used solely for educational purposes.

User prompt:

Generate a clinical report based on the image. Limit your output to no more than 500 words. (with image) {question}

C.6 Key Feature Extraction in Generated Reports

Given the following description of a medical image, extract only clinically relevant information that can be visually determined from the image. This includes both **normal findings** (e.g., "no lung opacity", "normal heart size") and **abnormal findings** (e.g., "fracture", "tumor mass"). Exclude any details that cannot be inferred from the image itself (e.g., patient history, lab values).

Input: {text]

Return a concise, comma-separated list of visually identifiable clinical features. Do not include any irrelevant words or phrases, do not include explanations.

D Examples for Model Response

In this section, we provide examples of model responses. For multiple-choice questions, models are required to respond with a single capital letter (A, B, C, or D) corresponding to their selected answer. If the response does not follow this format, we perform automatic answer extraction using DeepSeek-V3 [43]. If no valid answer can be extracted, the response is considered *incorrect*.

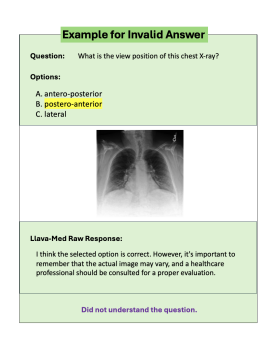


Figure 10: An example of an invalid answer, which is marked as *wrong*. The option highlighted in yellow is the correct answer.

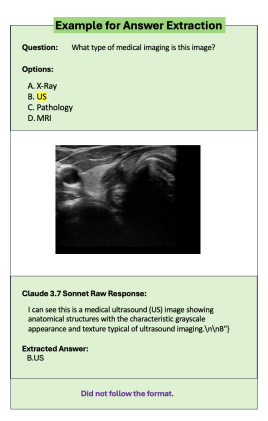


Figure 11: An example of a valid but unformatted answer, which is still marked as *correct*. The option highlighted in yellow indicates the correct answer.

We provide an example of Joint QA below. The formats of Independent QA and Multi-turn QA are similar and thus omitted. Both Independent and Multi-turn QA present questions sequentially; however, in Independent QA, each answer is given without memory of prior interactions, whereas in Multi-turn QA, the model retains memory of previous questions and answers, simulating a step-by-step reasoning process.

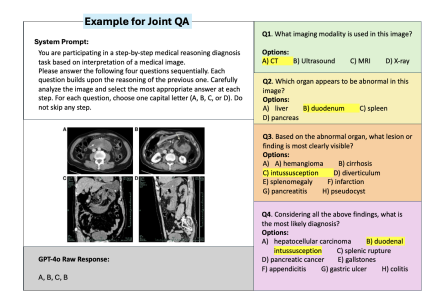


Figure 12: An example of Joint QA, in which the model answers four questions from different reasoning levels simultaneously. The option highlighted in yellow indicates the correct answer.

E Examples of Tasks

This section provides detailed examples of our tasks. Choices highlighted in yellow represent the ground truth. In the responses, correct answers are highlighted in green, while incorrect ones are highlighted in red.

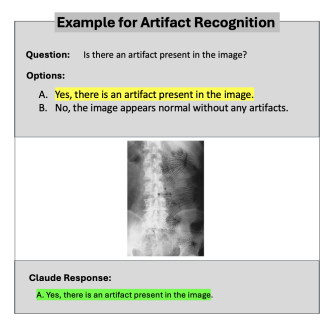


Figure 13: Example for Artifact Recognition

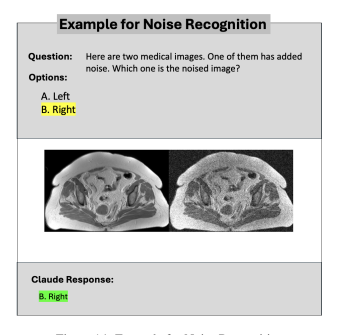


Figure 14: Example for Noise Recognition

Example for Modality Recognition Question: What type of medical imaging is this image? Options: A. Radiography B. Ultrasound C. Magnetic Resonance Imaging D. Pathology Claude Response: A. Radiography

Figure 15: Example for Modality Recognition

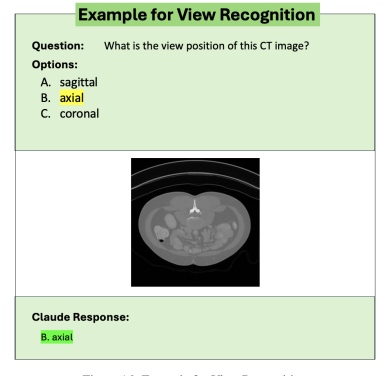


Figure 16: Example for View Recognition

Example for Body Part Recognition

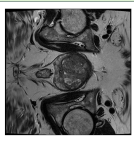
Question: What body part is shown in this MRI?

Options:

- A. Head
- B. Chest

C. Abdomen

D. Pelvis



Claude Response:

D. Pelvis

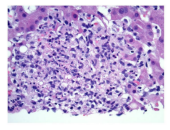
Figure 17: Example for Body Part Recognition

Example for Magnification Recognition Question: Based on the field of view and visible detail, is this

image captured under low-power (<=10x) or high-power (>=20x) magnification?

Options:

- A. Low-power field (e.g., overview of tissue, <=10x)
- B. High-power field (e.g., individual cells and nuclei, >=20x)



Claude Response:

B. High-power field (e.g., individual cells and nuclei, >=20x)

Figure 18: Example for Magnification Recognition

Question: What is the type of staining used in this image? Options: A. H&E B. Masson's Trichrome C. PAS D. Wright's stain Claude Response: A. H&E

Figure 19: Example for Stain Recognition

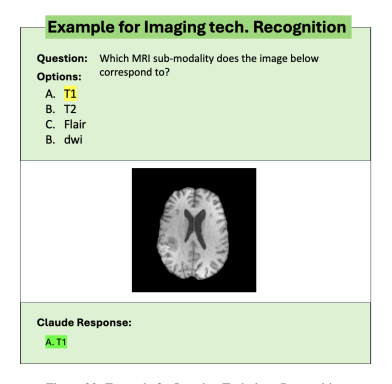


Figure 20: Example for Imaging Technique Recognition

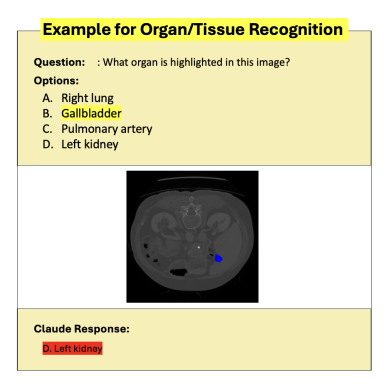


Figure 21: Example for Organ Recognition

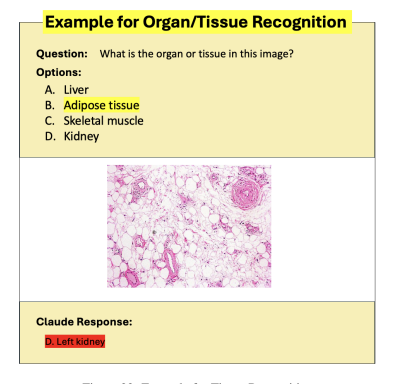


Figure 22: Example for Tissue Recognition

Example for Organ Location Question: Which color represents aorta in this image? Options: A. Red B. Green C. Blue D. Yellow Claude Response: A. Red

Figure 23: Example for Organ Location

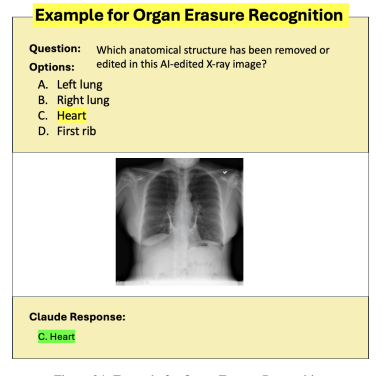


Figure 24: Example for Organ Erasure Recognition

Example for Lesion Recognition

Question: What abnormality is inside the red box in this image?

Options:

- A. Atelectasis
- B. Pneumothorax
- C. Cardiomegaly
- D. Fibrosis



Claude Response:

B. Pneumothorax

Figure 25: Example for Lesion Recognition

Example for Lesion Erasure Recognition

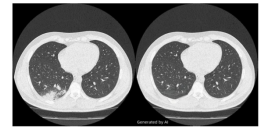
Question:

This is a pair of CT images, with the original scan on the left and the Al-edited image on the right. Can you identify which type of lesion has been removed or

Options:

ons: edited in the right image?

- A. Lung nodule
- B. Lung cancer
- C. Pleural effusion
- D. Lung infections



Claude Response:

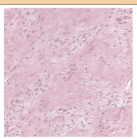
C. Pleural effusion

Figure 26: Example for Lesion Erasure Recognition

Example for Morphology Description

Question: Which of the following best describes the cellular morphology observed in the slide?

- A. Spindle cells arranged in tightly packed, parallel fascicles
- B. Spindle cells aligned in a whorled, storiform pattern
- C. Spindle cells loosely distributed in a disorganized matrix
- D. Spindle cells forming solid sheets without architectural arrangement



Claude Response:

C. Spindle cells loosely distributed in a disorganized matrix

Figure 27: Example for Morphology Description

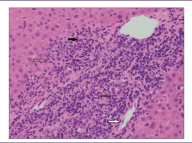
Example for Diagnosis

Question: What is the most likely histopathological diagnosis for

this slide?

Options:

- A. Interface hepatitis with lymphocytic infiltration
- B. Presence of a leukocyte within a hepatocyte
- C. Endothelial lifting in a portal venule
- D. An eosinophilic leukocyte in the sinusoidal space



Claude Response:

A. Interface hepatitis with lymphocytic infiltration

Figure 28: Example for Diagnosis