ReCAP: Report-Conditioned Attentive Patching for Gigapixel Histopathology

Tawsifur Rahman¹

ARAHMA34@JHU.EDU

¹ Biomedical Engineering, Johns Hopkins University

Rama Chellappa²

RCHELLA4@JHU.EDU

² Johns Hopkins University

Alexander S. Baras³

BARAS@JHMI.EDU

³ School of Medicine, Johns Hopkins University

Editors: Under Review for MIDL 2026

Abstract

Vision-language learning has become a powerful framework for multimodal representation, achieving exceptional performance across diverse image—text tasks. However, in histopathology, existing methods often rely on high-resolution region-level annotations to achieve fine-grained visual-textual alignment—an assumption that is impractical for Whole Slide Image (WSI) classification due to the gigapixel scale of pathology images and the weak supervision provided by slide-level labels. To address this challenge, we propose Report-Conditioned Attentive Patching for Weakly Supervised WSI Classification (ReCAP), a novel approach that leverages slide-level pathology reports to enrich patchlevel feature learning without requiring localized supervision. Instead of relying on explicit region annotations, ReCAP adopts a hybrid multimodal contrastive MIL framework in which report-conditioned text embeddings guide cross-attention to highlight semantically discriminative tissue regions. We further introduce a self-normalizing cross-modal attended similarity function that enhances the robustness and stability of patch-text alignment under weak supervision. In addition, our approach incorporates an efficient report-aware patch aggregation strategy that suppresses redundant or noisy regions while retaining the most diagnostically informative patterns within the vision-language context. Across multiple cancer subtype classification and survival prediction tasks, ReCAP consistently improves performance by 2–5%, demonstrating the effectiveness of report-conditioned cross-modal alignment for scalable and annotation-efficient WSI understanding.

Keywords: Whole Slide Image, Vision–Language Learning, Pathology Reports, Weakly Supervised Learning, Cross-Modal Alignment, Contrastive Learning.

1. Introduction

In histopathology, Whole Slide Image (WSI) analysis requires more than isolated patch examination—it demands a holistic and context-aware approach. Pathologists navigate WSIs dynamically, integrating morphological cues across multiple regions to form a diagnosis (1; 2; 3). This reasoning process unfolds like a multi-turn dialogue, where different tissue regions contribute varying degrees of diagnostic evidence (4; 5; 6). Importantly, pathology reports play a crucial role in this workflow: they summarize salient morphological findings, highlight suspicious patterns, and provide semantic cues that complement visual inspection. These textual insights often capture diagnostic signals—such as tumor descriptors,

cellular abnormalities, or clinically relevant context—that may not be uniformly present across sampled WSI patches. Incorporating such report-derived information into computational models therefore offers an avenue to enhance cancer subtype prediction by helping the model attend to patches that are more likely to carry diagnostic significance.

However, computational WSI analysis struggles to replicate this nuanced diagnostic behavior. Classical approaches treat WSI patches independently, overlooking their spatial relationships and failing to utilize the additional semantic information encoded in corresponding pathology reports (7; 8; 9). Weakly supervised learning settings further complicate this task, as slide-level labels provide no explicit supervision for identifying the most informative regions (10; 11; 12). As a result, models often aggregate thousands of patches indiscriminately, incorporating noise and diluting the discriminative power of key morphological patterns.

Recent advances in vision—language models (VLMs) have transformed multimodal representation learning, excelling in tasks such as image captioning, visual question answering, and cross-modal retrieval (13; 14). Yet, applying these methods directly to histopathology is non-trivial. Existing VLMs frequently assume the availability of region-level annotations or explicit image—text alignment, which is impractical for gigapixel WSIs annotated only at the slide level (15; 16; 17). Similarly, conventional multiple instance learning (MIL) frameworks lack mechanisms for selectively attending to patches that gain relevance only when contextualized with complementary textual information (18; 19; 20; 21). Domain-specific models such as CONCH and CPLIP partially mitigate domain shifts by leveraging large biomedical image—text corpora (13; 16), but they still aggregate patches uniformly and do not adaptively integrate report information to guide patch selection. This often leads to redundant or noisy patch inclusion and computational inefficiencies (22; 23; 24; 25).

To address these limitations, we propose **Report-Conditioned Attentive Patching** (**ReCAP**), a weakly supervised multimodal framework that leverages pathology reports to provide additional contextual information that improves patch-level attention and aggregation. Rather than assuming direct correspondence between individual patches and textual descriptions, ReCAP uses report information to modulate patch relevance, enabling the model to focus on tissue regions that better reflect the semantic content of the report. The main contributions of ReCAP include:

- (1) **Report-Guided Patch Prioritization:** ReCAP leverages pathology reports as complementary semantic information to modulate patch-level importance under weak supervision, improving downstream performance by emphasizing clinically meaningful regions.
- (2) **Hybrid Contrastive MIL With Cross-Attention:** Our framework integrates cross-attention-based contrastive learning to refine patch embeddings using report-conditioned context, enhancing semantic consistency between modalities without requiring region-level supervision.
- (3) Efficient Report-Aware Patch Aggregation: By suppressing redundant or non-informative regions, ReCAP achieves computational efficiency while preserving diagnostically relevant visual patterns.
- (4) Improved Performance Across Cancer Subtypes and Survival Tasks: Re-CAP demonstrates consistent gains of 2–5% across multiple cancers, highlighting the value of using pathology reports to enhance multimodal WSI understanding.

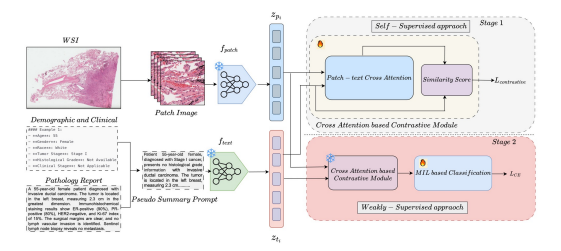


Figure 1: The overall architecture of our ReCAP framework (details in Section 2). ReCAP integrates pathology report information into WSI modeling through a two-stage design. Stage 1 uses report-conditioned cross-attention to refine patch representations, while Stage 2 aggregates the attended patches via a MIL-based predictor for classification and survival analysis.

2. Method

Our proposed ReCAP framework operates in two stages to incorporate pathology report information into weakly supervised WSI analysis. In the first stage, we use a multimodal contrastive learning module in which report-conditioned text embeddings guide cross-attention to provide additional contextual cues for identifying relevant patches. This enables the model to refine patch representations without requiring region-level annotations. In the second stage, the attended patch features are aggregated using a MIL-based predictor for downstream tasks, including classification and survival analysis formulated as an ordinal regression problem. The overall architecture of ReCAP is shown in Figure 1.

2.1. Extract Patch and Text Embeddings

Each WSI is divided into non-overlapping patches, yielding N_p total patches. Each patch p_i is encoded into a latent feature vector z_{p_i} using a frozen PLIP (14) image encoder:

$$z_{p_i} = f_{\text{patch}}(p_i; \theta_p) \in \mathbb{R}^{d_p},$$
 (1)

where f_{patch} denotes the patch encoder and θ_p are its fixed parameters.

To incorporate report-derived context, the pathology report and demographic information are combined into a pseudo-prompt and tokenized into N_t text tokens. Each token t_j is encoded using the frozen PLIP text encoder:

$$z_{t_j} = f_{\text{text}}(t_j; \theta_t) \in \mathbb{R}^{d_t}, \tag{2}$$

providing complementary semantic information that ReCAP later uses to guide patch-level attention.

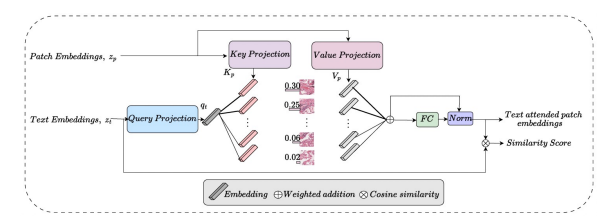


Figure 2: ReCAP cross-attention contrastive module.

2.2. Report-Conditioned Contrastive Learning with Cross-Attention

In the first stage of ReCAP, we incorporate report-derived context to refine patch representations using a contrastive learning module guided by cross-attention. Text embeddings provide additional cues that help highlight patches more likely to carry diagnostic information. As shown in Figure 2, text tokens act as queries while patch embeddings serve as keys and values. The attention weight between text token t_i and patch p_i is:

$$A_{ij} = \operatorname{softmax}\left(\frac{q_{t_j} K_{p_i}^T}{\sqrt{d}}\right),\tag{3}$$

where q_{t_i} and K_{p_i} denote the query and key vectors, respectively.

The attended patch embedding is computed as:

$$\tilde{z}_{p_i} = \sum_j A_{ij} V_{p_i}. \tag{4}$$

2.3. Similarity-Based MIL Aggregation

To estimate how well attended patches align with report context, we compute cosine similarity:

$$S_{ij} = \frac{\tilde{z}_{p_i} \cdot z_{t_j}}{\|\tilde{z}_{p_i}\| \|z_{t_j}\|}.$$
 (5)

Instead of learned attention, ReCAP directly uses similarity scores to weight patch contributions. From all patches, we retain the top-k% based on similarity:

$$P_{\text{top-k}} = \{ \tilde{z}_{p_i} \mid i \in \text{Top-k}(S_{ij}) \}.$$
 (6)

The final slide-level representation is:

$$\hat{z} = \sum_{i \in P_{\text{top-k}}} \alpha_i g(\tilde{z}_{p_i}), \tag{7}$$

where the normalized similarity weight is:

$$\alpha_i = \frac{\exp(S_i)}{\sum_{j \in P_{\text{top-k}}} \exp(S_j)}.$$
 (8)

Using similarity as instance weights ensures that patches better aligned with report context contribute more to the downstream prediction.

2.4. Loss Function

ReCAP is trained with a two-stage objective. The first stage uses a contrastive loss to align attended patches with relevant textual cues:

$$\mathcal{L}_{\text{Align}} = -\sum_{(i,j)\in P} \log \frac{\exp(S_{ij}/\tau)}{\sum_{k} \exp(S_{ik}/\tau)}.$$
 (9)

The second stage optimizes downstream classification or survival prediction using crossentropy:

$$\mathcal{L}_{\text{Predict}} = -\sum_{c} [y_c \log \hat{y}_c + (1 - y_c) \log(1 - \hat{y}_c)].$$
 (10)

The final objective combines both terms:

$$\mathcal{L}_{\text{Total}} = \lambda \mathcal{L}_{\text{Align}} + (1 - \lambda) \mathcal{L}_{\text{Predict}}, \tag{11}$$

where based on empirical evaluation, we found that setting $\lambda = 0.4$ yielded optimal performance.

Table 1: Performance of ReCAP and state-of-the-art methods on cancer subtype classification and survival prediction tasks.

Method	Image	Text	Cancer Subtype Classification				Survival Prediction									
			TCGA-RCC TCGA-NSCLC		TCGA-BRCA BF		BRCA	BRCA-IDC BRO		A-ILC	NSCLC-LUAD		NSCLC-LUSC			
			Acc	AUC	Acc	AUC	Acc	AUC	Acc	AUC	Acc	AUC	Acc	AUC	Acc	AUC
ABMIL (17)	✓	-	90.18	92.86	91.63	93.41	87.21	89.67	83.02	85.33	84.12	86.01	72.45	74.28	71.38	76.22
CLAM-SB (19)	✓	-	92.06	94.11	93.83	94.95	90.72	92.89	84.22	86.78	85.67	87.45	75.34	77.11	73.22	78.45
DS-MIL (26)	✓	-	91.41	93.13	92.73	94.20	89.41	91.82	83.71	86.34	85.23	87.01	74.11	76.45	70.68	77.23
TransMIL (27)	✓	-	91.92	94.06	93.01	94.23	89.93	92.61	84.45	86.92	85.78	87.56	74.78	77.56	72.34	78.11
CONCH (13)	✓	✓	94.29	96.31	95.36	96.73	93.11	94.88	86.12	88.67	87.23	89.12	78.45	80.11	75.11	80.78
PLIP (14)	✓	✓	93.56	95.80	94.71	96.64	92.34	93.78	85.78	88.01	86.67	88.91	77.56	79.34	74.78	79.89
ReCAP (Ours)	✓	✓	96.12	97.84	97.41	98.66	94.89	96.91	88.67	90.98	90.72	92.34	81.88	83.92	80.77	84.11

3. Results

3.1. Dataset

We evaluated our methodology on two downstream tasks cancer subtype classification and cancer survival prediction.

WSI Cancer Subtype Classification Our comprehensive approach was rigorously evaluated on three cancer subtype classification tasks using the TCGA-RCC (28), TCGA-NSCLC (28), and TCGA-BRCA (28) datasets. To avoid data leakage, we ensured that slides from the same patient case were grouped together during partitioning.

TCGA-RCC WSI. The TCGA-RCC WSI dataset includes 884 diagnostic Whole Slide Images (WSIs) from the TCGA RCC repository, covering Kidney Chromophobe (TCGA-KICH), Kidney Clear Cell Renal Cell Carcinoma (TCGA-KIRC), and Kidney Renal Papillary Cell Carcinoma (TCGA-KIRP) projects. Specifically, there are 111 slides from 99 cases of Chromophobe Renal Cell Carcinoma (CRCC), 489 slides from 483 cases of Clear Cell Renal Cell Carcinoma (CCRCC), and 284 slides from 264 cases of Papillary Renal Cell Carcinoma (PRCC). On average, 13,907 patches were extracted per slide at ×20 magnification.

TCGA-NSCLC WSI. The TCGA-NSCLC WSI dataset contains 1020 Formalin-Fixed Paraffin-Embedded (FFPE) WSIs, with 490 associated with lung adenocarcinoma (LUAD) and 530 with lung squamous cell carcinoma (LUSC).

TCGA-BRCA WSI. The TCGA-BRCA WSI dataset consists of 954 FFPE WSIs, including 787 slides with Invasive Ductal Carcinoma (IDC) and 167 slides with Invasive Lobular Carcinoma (ILC).

WSI Cancer Survival Prediction We investigated our model on four cancer subtype survival prediction tasks, considering survival beyond 1, 2, and 5 years. The datasets used include BRCA-IDC (28), BRCA-ILC (28), NSCLC-LUAD (28), and NSCLC-LUSC (28).

NSCLC-LUAD. This dataset comprises 490 whole-slide images (WSIs) of lung adenocarcinoma from Formalin-Fixed Paraffin-Embedded (FFPE) tissue samples. The WSIs are distributed as follows: 248 for survival beyond 1 year, 123 for 2 years, and 34 for 5 years.

NSCLC-LUSC. This dataset includes 530 WSIs of lung squamous cell carcinoma from FFPE samples, with survival-based distribution as follows: 258 WSIs for over 1 year, 171 for 2 years, and 75 for 5 years.

BRCA-IDC. The TCGA-BRCA-IDC dataset consists of 787 WSIs of invasive ductal carcinoma (IDC) from FFPE samples. The survival distribution includes 563 WSIs for more than 1 year, 346 for 2 years, and 136 for 5 years.

BRCA-ILC. The TCGA-BRCA-ILC dataset contains 167 WSIs of invasive lobular carcinoma (ILC) obtained from FFPE samples. The WSIs are distributed as follows: 128 for survival beyond 1 year, 69 for 2 years, and 30 for 5 years.

3.2. Experiment settings and Evaluation metrics

We implemented our approach using PyTorch and conducted all experiments on a server with 4 NVIDIA Tesla V100 GPUs and 32 CPU cores. All models were optimized using the Adam optimizer. For WSI survival prediction tasks, models were trained for up to 1500 steps with a batch size of 256 and an initial learning rate of 0.001. Hyperparameters were fine-tuned for each dataset, and the best model was selected based on validation performance within a 5-fold cross-validation setup. Evaluation metrics included accuracy and area under the curve (AUC), with accuracy in multi-label classification computed as the average across all target classes.

3.3. Performance Analysis

This section evaluates the performance of the proposed **ReCAP** framework against state-of-the-art baselines on two tasks: cancer subtype classification and survival prediction.

Table 2: Performance on different number of patch aggregation strategies in ReCAP across cancer subtype classification and survival prediction tasks.

Aggregation Strategy	Cancer	r Subtype Class	ification	Survival Prediction				
	TCGA-RCC	TCGA-NSCLC	TCGA-BRCA	ΓCGA-BRCA BRCA-IDC		NSCLC-LUAD	NSCLC-LUSC	
Random-k	89.12	91.34	84.98	79.56	80.45	72.12	70.34	
First-k	90.45	93.11	85.93	80.89	81.78	73.34	71.56	
Last-k	92.34	94.01	87.61	82.45	83.12	75.67	74.23	
Top-k (Ours)	96.12	97.41	94.89	88.67	90.72	81.88	80.77	

Results across seven TCGA cohorts are summarized in Table 1. Overall, ReCAP demonstrates strong and consistent improvements across all datasets, highlighting the benefits of integrating pathology report information into weakly supervised WSI analysis.

Cancer Subtype Classification. ReCAP achieves the highest accuracy across all three subtype classification tasks. It obtains 96.12% on TCGA-RCC, 97.41% on TCGA-NSCLC, and 94.89% on TCGA-BRCA—consistent improvements of 1–2% over the strongest multi-modal baselines (CONCH, PLIP) and larger gains over image-only approaches such as ABMIL, CLAM-SB, DS-MIL, and TransMIL. These improvements highlight the effectiveness of incorporating report-conditioned contextual cues during patch refinement. By introducing text-guided similarity signals early in the pipeline, ReCAP is better able to isolate subtype-relevant regions, particularly in morphologically heterogeneous datasets such as BRCA. This demonstrates that pathology reports provide complementary diagnostic information that strengthens patch selection and reduces noise in large, variable tissue regions.

Survival Prediction. Across the four survival cohorts, ReCAP again outperforms all competing models. It achieves 88.67% on BRCA-IDC, 90.72% on BRCA-ILC, 81.88% on NSCLC-LUAD, and 80.77% on NSCLC-LUSC. These values reflect consistent 1–3% gains over multi-modal baselines and larger improvements over single-modality MIL methods. Performance differences are most pronounced in the NSCLC cohorts, where relying solely on image features is insufficient to capture prognostic complexity. Incorporating report-derived cues helps the model better identify subtle histological patterns associated with patient outcomes.

The improvements arise from ReCAP's two-stage design: report-conditioned contrastive learning produces more discriminative patch embeddings, while similarity-driven MIL aggregation selects only the most relevant regions for prediction. This leads to more robust and interpretable patient-level risk estimation, ensuring that the model focuses on clinically meaningful evidence rather than spurious or redundant regions.

Overall, ReCAP establishes new state-of-the-art accuracies across both classification and survival tasks, demonstrating the value of integrating pathology report information as complementary context in computational pathology. These findings highlight the potential of multimodal WSI modeling to enhance diagnostic precision and support clinically informed decision-making.

3.4. Ablation Study

We evaluated the effect of different patch aggregation strategies in **ReCAP** using four methods—Random-k, First-k, Last-k, and Top-k—across multiple cancer subtype and survival prediction tasks. As shown in Table 2, the Top-k strategy clearly outperforms the

Table 3: Ablation study evaluating the impact of cross-attention by comparing ReCAP with image-only and text-only variants across classification and survival tasks.

Model Variant	Cancer Sub	type Classificati	on (Accuracy)	Survival Prediction (Accuracy)				
	TCGA-RCC	TCGA-NSCLC	TCGA-BRCA	BRCA-IDC	BRCA-ILC	NSCLC-LUAD	NSCLC-LUSC	
Image-Only	89.45	91.78	86.03	81.34	83.01	75.12	73.78	
Text-Only	75.12	78.23	72.45	79.01	80.12	72.34	70.89	
ReCAP (Full Model)	96.12	97.41	94.89	88.67	90.72	81.88	80.77	

Table 4: Ablation study on the loss weighting parameter λ in ReCAP across subtype classification and survival prediction tasks.

λ Value	Cancer	r Subtype Class	ification	Survival Prediction					
	TCGA-RCC	TCGA-NSCLC	TCGA-BRCA	BRCA-IDC	BRCA-ILC	NSCLC-LUAD	NSCLC-LUSC		
$\lambda = 0.3$	94.01	95.89	92.12	86.23	88.01	79.12	78.23		
$\lambda = 0.4 \text{ (Best)}$	96.12	97.41	94.89	88.67	90.72	81.88	80.77		
$\lambda = 0.5$	95.23	96.34	93.45	87.12	89.23	80.34	79.45		
$\lambda = 0.6$	94.78	95.89	92.67	86.45	88.67	79.89	78.89		
$\lambda = 0.7$	93.89	95.12	91.78	85.23	87.34	78.45	77.56		

others, achieving the highest accuracies on all datasets. For example, ReCAP with *Top-k* reaches 96.12% (RCC), 97.41% (NSCLC), and 94.89% (BRCA) in classification, and 88.67%, 90.72%, 81.88%, and 80.77% on BRCA-IDC, BRCA-ILC, LUAD, and LUSC survival tasks, respectively. Across strategies, performance generally increases when selecting more informative patches. *Last-k* shows steady gains (e.g., from 93.01% to 94.89% on NSCLC, and 76.12% to 75.01% on LUSC), but consistently remains below *Top-k*. These results indicate that simply taking sequential patches is less effective than using report-conditioned similarity to identify the most relevant regions.

To evaluate the contribution of cross-attention in **ReCAP**, we compared the full model with two ablated variants: an *image-only* model that removes the text branch, and a *text-only* model that excludes patch embeddings. As shown in Table 3, ReCAP consistently outperforms both variants across all classification and survival datasets. In the subtype classification tasks (TCGA-RCC, TCGA-NSCLC, TCGA-BRCA), ReCAP improves accuracy by 5–10% compared to image-only and by 8–14% compared to text-only baselines. Similarly, in survival prediction tasks (BRCA-IDC, BRCA-ILC, NSCLC-LUAD, NSCLC-LUSC), ReCAP shows gains of 3–6% over single-modality models. These results highlight the importance of integrating report-conditioned textual cues with patch-level image features. The cross-attention mechanism enables ReCAP to leverage complementary information from both modalities, resulting in more informative patch selection and stronger downstream performance.

To further examine the contribution of the contrastive alignment loss within **ReCAP**, we conducted an ablation study by varying the loss weighting parameter λ from 0.3 to 0.7. As shown in Table 4, model performance improves steadily as λ increases from 0.3 to 0.4, with $\lambda = 0.4$ yielding the strongest results across all cancer subtype classification and survival prediction tasks. This setting provides the best balance between enforcing meaningful cross-modal alignment and maintaining strong supervision from the prediction objective. When λ is set too low, the model underutilizes the benefits of contrastive alignment, resulting in weaker multimodal fusion. Conversely, larger values such as 0.6 and 0.7 overly

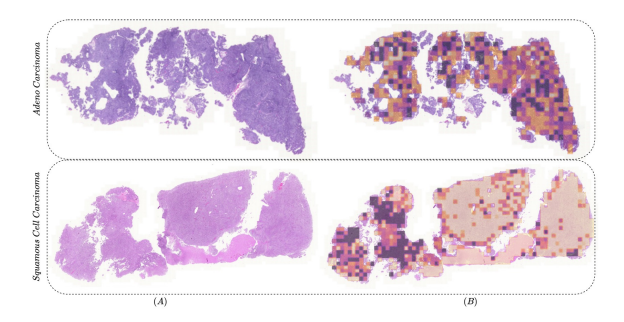


Figure 3: Visualization of ReCAP: (A) Original WSI (B) Report-conditioned attentive patching heatmap on WSI

emphasize the alignment term, which can hinder the predictive component and lead to noticeable drops in accuracy. These outcomes highlight the importance of carefully tuning λ to ensure that ReCAP effectively integrates both visual and textual information. Overall, the ablation demonstrates that performance is highly sensitive to the interplay between alignment and prediction losses, and that a moderate weighting is essential for maximizing the discriminative power of the learned multimodal representations.

3.5. Model Interpretation

To gain deeper insights into how ReCAP leverages pathology reports during WSI analysis, we first visualize its report-conditioned patch attention. As shown in Figure 3, the heatmap is derived from attention scores produced by the cross-attention contrastive module, where textual cues guide the model to prioritize patches that are semantically consistent with the report content. This targeted attention reveals diagnostically meaningful tissue regions and provides a more interpretable view of the model's decision-making process. Compared to uniform or image-only attention mechanisms, ReCAP produces sharper and more localized activation patterns, demonstrating the benefit of incorporating clinical text to refine patch relevance.

We further assess the quality of the learned representations by comparing embeddings generated from image-only, text-only, and multimodal versions of the model. For this analysis, we sampled representative WSIs, extracted aggregated slide-level features for each modality, and projected them into a two-dimensional space using UMAP, as shown in Figure 4. Image-only and text-only embeddings show substantial overlap between cancer subtypes, indicating limited discriminative power when each modality is used in isolation.

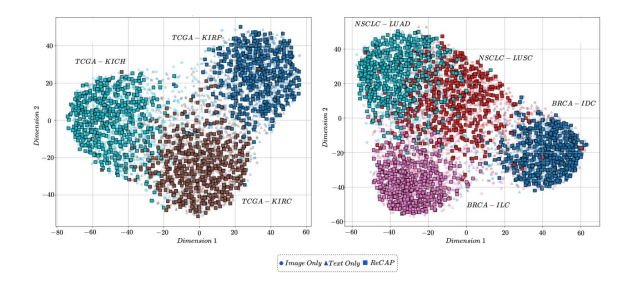


Figure 4: UMAP Projection of Patch Representations from Image-only, Text-only, and Re-CAP Models (TCGA-RCC Cancer Subtype Classification (Left) and Survival Prediction Across Subtypes (Right))

In contrast, the multimodal embeddings produced by ReCAP form clearer and more distinct clusters, reflecting stronger subtype separation. These results confirm that ReCAP's integration of report-guided information not only improves attention quality but also leads to more discriminative and semantically aligned feature representations.

4. Conclusion

In this work, we introduced **ReCAP**, a report-conditioned contrastive learning framework for weakly supervised WSI analysis. By integrating pathology report information through a cross-attention module and refining patch relevance using similarity-based aggregation, ReCAP effectively captures complementary visual—textual cues and improves both classification and survival prediction performance. Comprehensive experiments across multiple TCGA cohorts demonstrate that ReCAP consistently outperforms existing image-only and multimodal baselines, confirming the benefits of using report-derived context to guide patch-level reasoning. Ablation studies further show that the cross-attention mechanism and the top-k aggregation strategy are key contributors to the model's gains, enabling more discriminative feature learning and more informative patch selection. Overall, ReCAP offers a scalable and interpretable approach for multimodal pathology modeling, highlighting the value of incorporating clinical text to enhance WSI-based decision making and pointing toward broader opportunities for vision—language integration in digital pathology.

References

- [1] Gurcan, M.N., Boucheron, L.E., Can, A., Madabhushi, A., Rajpoot, N.M., Yener, B.: Histopathological image analysis: a review. IEEE Rev. Biomed. Eng. 2, 147–171
- [2] Yang, X., et al.: Virtual stain transfer in histology via cascaded deep neural networks. ACS Photonics 9(9), 3134–3143 (2022)
- [3] Cui, M., Zhang, D.Y.: Artificial intelligence and computational pathology. Lab. Invest. 101, 412–422 (2021)
- [4] El Nahhas, Omar SM, Marko van Treeck, Georg Wölflein, Michaela Unger, Marta Ligero, Tim Lenz, Sophia J. Wagner et al. "From whole-slide image to biomarker prediction: end-to-end weakly supervised deep learning in computational pathology." Nature Protocols 20, no. 1 (2025): 293-316.
- [5] Srinidhi, Chetan L., Ozan Ciga, and Anne L. Martel. "Deep neural network models for computational histopathology: A survey." Medical image analysis 67 (2021): 101813.
- [6] Kapse, Saarthak, Srijan Das, Jingwei Zhang, Rajarsi R. Gupta, Joel Saltz, Dimitris Samaras, and Prateek Prasanna. "Attention De-sparsification Matters: Inducing diversity in digital pathology representation learning." Medical Image Analysis 93 (2024): 103070.
- [7] Ahmedt-Aristizabal, David, Mohammad Ali Armin, Simon Denman, Clinton Fookes, and Lars Petersson. "A survey on graph-based deep learning for computational histopathology." Computerized Medical Imaging and Graphics 95 (2022): 102027.
- [8] Yang, X., et al.: Virtual stain transfer in histology via cascaded deep neural networks. ACS Photonics 9(9), 3134–3143 (2022)
- [9] Le Bescond, Loïc, Marvin Lerousseau, Fabrice Andre, and Hugues Talbot. "SparseXMIL: Leveraging spatial context for classifying whole slide images in digital pathology." (2024).
- [10] Tang, Wenhao, Fengtao Zhou, Sheng Huang, Xiang Zhu, Yi Zhang, and Bo Liu. "Feature Re-Embedding: Towards Foundation Model-Level Performance in Computational Pathology." In Proceedings of the IEEE/CVF Conference on Computer Vision and Pattern Recognition, pp. 11343-11352. 2024.
- [11] Rahman, Tawsifur, Alexander S. Baras, and Rama Chellappa. "Evaluation of a Task-Specific Self-Supervised Learning Framework in Digital Pathology Relative to Transfer Learning Approaches and Existing Foundation Models." Modern Pathology 38, no. 1 (2025): 100636.
- [12] Laleh, Narmin Ghaffari, Hannah Sophie Muti, Chiara Maria Lavinia Loeffler, Amelie Echle, Oliver Lester Saldanha, Faisal Mahmood, Ming Y. Lu et al. "Benchmarking weakly-supervised deep learning pipelines for whole slide classification in computational pathology." Medical image analysis 79 (2022): 102474.

- [13] Lu, Ming Y., Bowen Chen, Drew FK Williamson, Richard J. Chen, Ivy Liang, Tong Ding, Guillaume Jaume et al. "A visual-language foundation model for computational pathology." Nature Medicine 30, no. 3 (2024): 863-874.
- [14] Huang, Zhi, Federico Bianchi, Mert Yuksekgonul, Thomas J. Montine, and James Zou. "A visual–language foundation model for pathology image analysis using medical twitter." Nature medicine 29, no. 9: 2307-2316. (2023)
- [15] Ahmed, Faruk, Andrew Sellergren, Lin Yang, Shawn Xu, Boris Babenko, Abbi Ward, Niels Olson et al. "Pathalign: A vision-language model for whole slide images in histopathology." arXiv preprint arXiv:2406.19578 (2024).
- [16] Javed, Sajid, Arif Mahmood, Iyyakutti Iyappan Ganapathi, Fayaz Ali Dharejo, Naoufel Werghi, and Mohammed Bennamoun. "Cplip: zero-shot learning for histopathology with comprehensive vision-language alignment." In Proceedings of the IEEE/CVF conference on computer vision and pattern recognition, pp. 11450-11459. 2024.
- [17] Ilse, M., Tomczak, J., Welling, M.: Attention-based deep multiple instance learning. In: International Conference on Machine Learning, pp. 2127–2136. PMLR (2018)
- [18] Fashi, Parsa Ashrafi, Sobhan Hemati, Morteza Babaie, Ricardo Gonzalez, and H. R. Tizhoosh. "A self-supervised contrastive learning approach for whole slide image representation in digital pathology." Journal of Pathology Informatics 13 (2022): 100133.
- [19] Lu, M.Y., Williamson, Mahmood, F.: Data-efficient and weakly supervised computational pathology on whole-slide images. Nat. Biomed. Eng. 5(6), 555–570 (2021)
- [20] Srinidhi, Chetan L., and Anne L. Martel. "Improving self-supervised learning with hardness-aware dynamic curriculum learning: an application to digital pathology." In Proceedings of the IEEE/CVF International Conference on Computer Vision, pp. 562-571. 2021.
- [21] Adnan, Mohammed, Shivam Kalra, and Hamid R. Tizhoosh. "Representation learning of histopathology images using graph neural networks." In Proceedings of the IEEE/CVF Conference on Computer Vision and Pattern Recognition Workshops, pp. 988-989. 2020.
- [22] Pati, Pushpak, Guillaume Jaume, Lauren Alisha Fernandes, Antonio Foncubierta-Rodríguez, Florinda Feroce, Anna Maria Anniciello, Giosue Scognamiglio et al. "Hactnet: A hierarchical cell-to-tissue graph neural network for histopathological image classification." In Uncertainty for Safe Utilization of Machine Learning in Medical Imaging, and Graphs in Biomedical Image Analysis: Second International Workshop, UNSURE 2020, and Third International Workshop, GRAIL 2020, Held in Conjunction with MICCAI 2020, Lima, Peru, October 8, 2020, Proceedings 2, pp. 208-219. Springer International Publishing, 2020.
- [23] Rahman, T., Baras, A.S. and Chellappa, R., 2025, February. CEMIL: Contextual Attention Based Efficient Weakly Supervised Approach for Histopathology Image Classification. In 2025 IEEE/CVF Winter Conference on Applications of Computer Vision (WACV) (pp. 4248-4257). IEEE.

- [24] Nair, Aravind, Helena Arvidsson, Jorge E. Gatica V, Nikolce Tudzarovski, Karl Meinke, and Rachael V. Sugars. "A graph neural network framework for mapping histological topology in oral mucosal tissue." BMC bioinformatics 23, no. 1 (2022): 506.
- [25] Chen, Richard J., Chengkuan Chen, and Faisal Mahmood. "Scaling vision transformers to gigapixel images via hierarchical self-supervised learning." In Proceedings of the IEEE/CVF Conference on Computer Vision and Pattern Recognition, pp. 16144-16155. 2022.
- [26] Li, Bin, Yin Li, and Kevin W. Eliceiri. "Dual-stream multiple instance learning network for whole slide image classification with self-supervised contrastive learning." In Proceedings of the IEEE/CVF conference on computer vision and pattern recognition, pp. 14318-14328. 2021.
- [27] Shao, Z., Bian, H., Chen, Y., Wang, Y., Zhang, J., Ji, X., et al.: Transmil: transformer based correlated multiple instance learning for whole slide image classification. Adv. Neural Inf. Process. Syst. 34, 2136–2147 (2021)
- [28] Tomczak, K., Czerwińska, P. and Wiznerowicz, M., 2015. Review The Cancer Genome Atlas (TCGA): an immeasurable source of knowledge. Contemporary Oncology/Współczesna Onkologia, 2015(1), pp.68-77. (2015)

Appendix A. Whole Slide Image Preprocessing

Whole slide image (WSI) preprocessing begins with automated tissue segmentation. Each WSI is loaded at a downsampled resolution (e.g., $20\times$) and converted from RGB to HSV color space, where tissue regions are identified by thresholding the saturation channel after applying median blurring to reduce noise. A binary tissue mask is generated and refined using morphological closing to remove small gaps and holes, followed by contour extraction and area-based filtering to retain only meaningful tissue regions. The resulting segmentation mask is saved for optional visual inspection, and a log file is produced to facilitate manual parameter adjustments when needed.

After segmentation, fixed-size patches (256×256) are extracted from within the tissue contours at the desired magnification. Each patch is stored along with its spatial coordinates and slide-level metadata in an HDF5 file for efficient downstream processing. The number of extracted patches varies substantially across slides—ranging from a few hundred in small biopsy samples at $20\times$ to several hundred thousand in large resection slides at higher magnifications.

Appendix B. Performance with Top-k Patch Aggregation.

Performance with Top-k Patch Aggregation. To evaluate the effectiveness of different patch aggregation strategies within our ReCAP framework, we conducted an ablation study comparing four variants: Random-k, First-k, Last-k, and our proposed Top-k strategy. The Top-k approach, which selects the most informative 70% of patches based on attention scores, consistently outperformed the others across both cancer subtype classification and

survival prediction tasks. In cancer subtype classification, Top-k aggregation yielded remarkable performance improvements. For instance, on the TCGA-NSCLC dataset, Top-k achieved an accuracy of 96.33%, surpassing Last-k by 2.32% and Random-k by 4.99%. Similarly, on TCGA-RCC, Top-k reached 94.68%, outperforming Last-k and First-k by 2.34% and 4.23%, respectively. In the TCGA-BRCA dataset, Top-k attained 92.87%, marking a notable gain of 5.26% over Random-k. These results highlight the advantage of attentionguided patch selection over simple heuristics or positional subsets. For survival prediction, Top-k again delivered the best outcomes across all datasets. On BRCA-ILC and BRCA-IDC, Top-k achieved accuracies of 87.45% and 85.74%, improving over the next best (Last-k) by 4.33% and 3.29%, respectively. The improvement trend continued for NSCLC subtypes, where Top-k achieved 79.67% (LUAD) and 78.66% (LUSC), both outperforming Last-k by over 3.5% on average. These consistent gains demonstrate that aggregating the most semantically relevant, text-attended patches allows ReCAP to focus on diagnostically and prognostically critical regions, thereby enhancing its discriminative capacity. Overall, the Top-k strategy significantly boosts performance across tasks and cancer types, validating its generalizability. While we empirically fixed the selection threshold at k = 0.7, future work could explore adaptive selection mechanisms to further optimize the trade-off between computational efficiency and predictive power.

Table 5: Performance of different patch aggregation strategies in ReCAP across cancer subtype classification and survival prediction tasks, under varying k values.

2*k value	2*Aggregation Strategy	Cancer Subt	ype Classification	Accuracy (%)	Survival Prediction Accuracy (%)				
		TCGA-RCC	TCGA-NSCLC	TCGA-BRCA	BRCA-IDC	BRCA-ILC	NSCLC-LUAD	NSCLC-LUSC	
4*0.5	Random-k	88.34	90.12	83.21	78.12	79.45	70.12	68.34	
	First-k	89.01	91.67	84.12	79.34	80.67	71.23	69.01	
	Last-k	90.56	92.89	85.78	80.56	82.34	73.89	72.23	
	Top-k	92.45	94.23	87.65	83.01	84.78	76.56	75.89	
4*0.6	Random-k	88.89	91.01	84.45	78.67	79.89	71.34	69.78	
	First-k	89.67	92.56	85.56	80.12	81.89	72.56	70.45	
	Last-k	91.12	93.34	86.89	81.45	83.45	74.45	73.01	
	Top-k	93.23	95.45	89.12	84.12	85.78	77.34	76.23	
4*0.7	Random-k	89.12	91.34	84.98	79.56	80.45	72.12	70.34	
	First-k	90.45	93.11	85.93	80.89	81.78	73.34	71.56	
	Last-k	92.34	94.01	87.61	82.45	83.12	75.67	74.23	
	Top-k (Ours)	94.68	96.33	92.87	85.74	87.45	79.67	78.66	
4*0.8	Random-k	89.34	91.56	85.21	79.78	80.89	72.45	70.89	
	First-k	90.67	93.23	86.34	81.01	82.23	74.12	72.23	
	Last-k	92.56	94.23	88.23	83.12	84.12	76.12	75.12	
	Top-k	94.12	96.42	91.56	85.12	86.56	78.78	77.89	

Appendix C. Model Interpretation

In addition to evaluating the performance of ReCAP across both cancer subtype classification and survival prediction tasks, we further investigated the quality of the learned feature representations. We assess its interpretability by analyzing the most informative patches selected by the attention module. This module dynamically prioritizes highly text-attended patches, enhancing the efficiency and relevance of survival prediction. To explore how our model distinguishes between cancer subtypes, we visualize the highly attended patches for ductal breast cancer, as shown in Figure 5. The first column displays the most attended patches from WSIs of breast ductal carcinoma, while the second column presents the cor-

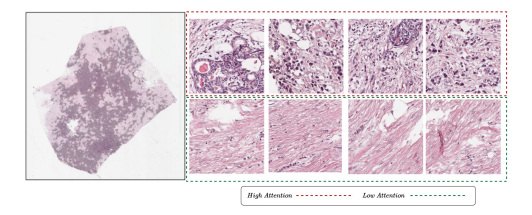


Figure 5: Attention map based WSI visualization using ReCAP framework.

responding low-attended patches. These selected regions highlight the key morphological features that influence the model's predictions.

Appendix D. Algorithm of ReCAP

```
Algorithm 1: Report-Conditioned Contrastive MIL (ReCAP)
Input: Training set \{(X^{(n)}, R^{(n)}, y^{(n)})\}_{n=1}^N of WSIs X, reports R, and labels y;
        Frozen PLIP image encoder f_{\text{patch}}(\cdot; \theta_p); frozen PLIP text encoder f_{\text{text}}(\cdot; \theta_t);
        Trainable cross-attention + projection parameters \Theta_{\text{CaCo}}; MIL head \Theta_{\text{MIL}};
        Temperature \tau; loss weight \lambda; top-k retention ratio k
Output: Trained parameters \Theta_{CaCo}, \Theta_{MIL}
while not converged do
     Sample a mini-batch \mathcal B of slide–report pairs
     foreach (X, R, y) \in \mathcal{B} do
                                                                                                                                     */
          /* 1. Extract patch and text embeddings
          Extract non-overlapping patches \{p_i\}_{i=1}^{N_p} from WSI X for i=1 to N_p do
           z_{p_i} \leftarrow f_{\text{patch}}(p_i; \theta_p) ;
          end
          Build pseudo-prompt from R + demographics and tokenize into \{t_j\}_{j=1}^{N_t} for j=1
           z_{t_i} \leftarrow f_{\text{text}}(t_j; \theta_t) ;
                                                                                                          // text embedding
          /* 2. Report-conditioned cross-attention
          Project embeddings to queries/keys/values (learnable): q_{t_j} \leftarrow W_q z_{t_j}, K_{p_i} \leftarrow W_k z_{p_i}
            V_{p_i} \leftarrow W_v z_{p_i} for j = 1 to N_t do
               \mathbf{for} \ i = 1 \ \mathbf{to} \ N_p \ \mathbf{do}
A_{ij} \leftarrow \mathrm{softmax}_i \left( \frac{q_{t_j} K_{p_i}^\top}{\sqrt{d}} \right)
          end
          \begin{array}{l} \textbf{for } i = 1 \textbf{ to } N_p \textbf{ do} \\ \mid \ \tilde{z}_{p_i} \leftarrow \sum_{j=1}^{N_t} A_{ij} V_{p_i} \ ; \end{array}
                                                                                        // attended patch embedding
          /* 3. Similarity-based Top-k MIL aggregation
                                                                                                                                     */
          for i = 1 to N_p do
               // aggregate similarity over tokens, e.g., max or mean
               S_i \leftarrow \max_j \frac{\tilde{z}_{p_i} \cdot z_{t_j}}{\|\tilde{z}_{p_i}\| \|z_{t_i}\|}
          Select index set P_{\text{top-}k} of top-k\% patches sorted by S_i foreach i \in P_{top-k} do \alpha_i \leftarrow \frac{\exp(S_i)}{\sum_{j \in P_{\text{top-}k}} \exp(S_j)}; // normalized w
                                                                                                    // normalized weight
          end
          Compute slide-level representation: \hat{z} \leftarrow \sum_{i \in P_{\text{top-}k}} \alpha_i g(\tilde{z}_{p_i}; \Theta_{\text{MIL}}) Predict logits /
           risks: \hat{y} \leftarrow h(\hat{z}; \Theta_{\text{MIL}})
          /* 4. Loss computation
                                                                                                                                     */
          Compute prediction loss \mathcal{L}_{Predict}(y, \hat{y}) (classification or ordinal survival) Con-
           struct positive patch–text pairs (i,j) \in P and compute:
            -\sum_{(i,j)\in P} \log \frac{\exp(S_{ij}/\tau)}{\sum_{k} \exp(S_{ik}/\tau)} \mathcal{L}_{\text{Total}} \leftarrow \lambda \mathcal{L}_{\text{Align}} + (1-\lambda)\mathcal{L}_{\text{Predict}}
     end
     // 5. Parameter update
     Update \Theta_{\text{CaCo}}, \Theta_{\text{MIL}} using Adam on \sum_{(X,R,y)\in\mathcal{B}} \mathcal{L}_{\text{Total}}
```

 $\mathbf{return}\ \Theta_{CaCo}, \Theta_{MIL}$