

Singular Value Few-shot Adaptation of Vision-Language Models

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Paper under double-blind review

Abstract

Vision-language models (VLMs) like CLIP have shown impressive zero-shot and few-shot learning capabilities across diverse applications. However, adapting these models to new fine-grained domains remains difficult due to reliance on prompt engineering and the high cost of full model fine-tuning. Existing adaptation approaches rely on augmented components, such as prompt tokens and adapter modules, which could limit adaptation quality, destabilize the model, and compromise the rich knowledge learned during pretraining. In this work, we present **CLIP-SVD**, a novel *multi-modal* and *parameter-efficient* adaptation technique that leverages Singular Value Decomposition (SVD) to modify the internal parameter space of CLIP without injecting additional modules. Specifically, we fine-tune only the singular values of the CLIP parameter matrices to rescale the basis vectors for domain adaptation while retaining the pretrained model. This design enables enhanced adaptation performance using only **0.04%** of the model’s total parameters and better preservation of its generalization ability. CLIP-SVD achieves state-of-the-art classification results on 11 natural and 10 biomedical datasets, outperforming previous methods in both accuracy and generalization under few-shot settings. Additionally, we leverage a natural language-based approach to analyze the effectiveness and dynamics of the CLIP adaptation to allow interpretability of CLIP-SVD.

1 Introduction

Vision-language models (VLMs), such as CLIP (Radford et al., 2021), have demonstrated remarkable versatility and generalization by aligning images and text through large-scale contrastive pretraining. These models enable powerful zero-shot and few-shot capabilities for various applications, but adapting them effectively to downstream tasks remains non-trivial. Full model fine-tuning is often computationally infeasible, while prompt learning strategies (e.g., CoOp (Zhou et al., 2022b) and CoCoOp (Zhou et al., 2022a)) may be limited by heavy dependency on handcrafted or learned text prompts. Adapter-based methods, such as CLIP-Adapter (Gao et al., 2024) and MaPLe (Khattak et al., 2023a), infuse additional modules to improve adaptation quality, but this increases model complexity, lowers inference efficiency, and sometimes degrades zero-shot performance by destabilizing pretrained representations (Khattak et al., 2023b). Recent efforts in parameter-efficient fine-tuning (PEFT) aim to overcome these limitations. Among them, Singular Value Fine-Tuning (SVF) (Sun et al., 2022) has emerged as a compelling strategy by modifying only the singular values of model weight matrices without changing the original model. While Sun et al. (2022) and Meng et al. (2024) showed that SVF has promise in CNNs and large language models (LLMs), it has not been fully explored in Transformer-based, multi-modal settings. Furthermore, despite the popularity of CLIP adaptation methods, very few have attempted to interpret the dynamics and effectiveness of model adaptation. Lastly, most CLIP adaptation techniques focus on natural domains alone, with few specialized in biomedical applications (Bie et al., 2024; Koleilat et al., 2025b) due to distinctive visual features and complex clinical descriptions. This creates a gap for a universal strategy that can generalize effectively across natural and biomedical domains without high computational complexity or tailored adjustments (e.g., specialized prompt engineering (Bie et al., 2024; Koleilat et al., 2025b)).

Table 1: Natural language-based interpretations for the top 3 Attention Heads associated with the highest normalized changes (sorted in descending order) in the Output-Value circuit after CLIP adaptation for different datasets. Here, “**L**” denotes layer while “**H**” denotes attention head.

EuroSAT (Satellite Images)	DTD (Texture Images)
(L10.H0): Aerial Landscapes & Environments (L10.H10): Mood, Atmosphere & Highlights (L11.H0): Semantic Layout & Spatial Context	(L8.H6) Refined Textural Details of Everyday Objects (L10.H2) Cultural & Textural Scenes (L9.H4) Natural Landscapes & Textures
SUN397 (Scene Understanding)	UCF101 (Action Recognition)
(L11.H0): Semantic Layout & Spatial Context (L11.H2): Numbers, Symbols & Temporal Cues (L11.H3): Lifestyle & Tranquil Activities	(L10.H5) Action, Emotion & Faces (L10.H6) Organic Flow & Movement (L10.H1) Human Experiences & Objects in Action
BUSI (Breast Ultrasound)	BTMRI (Brain MRI)
(L11.H8): Converging Edges & Cluster Markers (L8.H6): Contour Irregularity & Internal Spread (L8.H3): Radiologic Artifacts & Diffuse Shapes	(L8.H9) Scattered Highlights & Artifactual Spots (L8.H0) Focal Markers & Shape Cues (L9.H5) Streaks, Texture, & Soft Borders
COVID-QU-Ex (Chest X-ray)	CTKIDNEY (Kidney CT)
(L11.H0): Signal Voids & Shifts (L9.H1): Ring-Like Structures & Localized Spread (L11.H3): Cross-Lobe Flow & Density Buildup	(L8.H6): Contour Irregularity & Internal Spread (L8.H3): Radiologic Artifacts & Diffuse Shapes (L8.H10): Diffuse Zones & Overlapping Shapes

To address the aforementioned challenges, we present **CLIP-SVD**, a novel parameter-efficient few-shot framework for unified CLIP adaptation across both natural and biomedical domains. While many relevant methods (Zhou et al., 2022b) primarily focus on the text branch, recent ones (Khattak et al., 2023a) have shown the benefit of adapting image and text branches jointly, at the cost of heavy “add-on” modules. For example, the popular MaPLe requires additional trainable parameters for 2.85% of the CLIP model. In contrast, our approach leverages Singular Value Decomposition (SVD) to decompose the projection weights in CLIP’s attention and feedforward layers into corresponding singular values and singular vectors, with only the first fine-tuned for both image and text encoders. We hypothesize that this allows the model to rescale the basis vectors for each downstream task with superior adaptation quality and generalizability. Furthermore, our combination of CLIP’s multi-head attention and SVD-based weight adaptation invites an opportunity for a natural language-based paradigm to localize, rank, and semantically “describe” the most significant dynamic shifts in the adapted model. Here, we probe the best text basis to map the semantic meaning of the attention heads (Gandelsman et al., 2023) with the most significant updates through CLIP-SVD, as shown in Table 1 for 16-shot CLIP adaptation on distinct tasks. Compared with previous efforts (Sun et al., 2022) that rely on visual interpretation and/or model weight statistics, this approach offers more intuitive and granular insights into CLIP adaptation. Yet, a related text corpus for analyzing biomedical data is still unavailable.

Our study has four major contributions: **First**, we proposed an SVD-based few-shot adaptation framework for Transformer-based multi-modal settings (e.g., CLIP and BiomedCLIP) for the first time, requiring just **0.04%** of the model’s total parameters, significantly lower than other multi-modal methods. **Second**, we performed comprehensive validation with 11 natural and 10 biomedical domain datasets, demonstrating CLIP-SVD’s superior performance against the state-of-the-art (SOTA) methods in both accuracy and generalization. **Third**, with ranked weight changes associated with our method, we adopted a natural language-facilitated approach to intuitively interpret the effectiveness and dynamics of task-specific CLIP adaptation. **Lastly**, to meet an urgent need for semantic interpretation of attention heads in CLIP for biomedical applications (e.g., analysis of CLIP-SVD), we built the first corpus of biomedical image descriptions.

2 Related Works

2.1 Parameter-efficient Fine-tuning

Fine-tuning large VLMs is often computationally prohibitive for domain-specific adaptation. Parameter-efficient fine-tuning addresses this by updating only a small subset of parameters while keeping the backbone frozen (Lialin et al., 2023). Selective tuning methods like BitFit (Zaken et al., 2022) adjust only bias terms, while pruning and sparsity techniques can further reduce trainable parameters (Guo et al., 2021; Holmes et al., 2021), though they often compromise robustness in zero-shot settings. Adapter-based tuning offers a more robust alternative by inserting lightweight modules (Rebuffi et al., 2017; Houlsby et al., 2019; Karimi Mahabadi et al., 2021; Chen et al., 2022b; Lian et al., 2022), but can introduce inference latency (Pfeiffer et al., 2021). Popular prompt tuning (Lester et al., 2021; Li & Liang, 2021; Jia et al., 2022) and Low-Rank Adaptation (LoRA) (Hu et al., 2021) provide other PEFT strategies, with LoRA inserting low-rank matrices to minimize overfitting (Li et al., 2018; Aghajanyan et al., 2021). However, these methods typically rely on external, randomly initialized modules, risking destabilization and forgetting of original CLIP knowledge (Zhang et al., 2023b; Zhu et al., 2024b; Shuttleworth et al., 2025). Designing PEFT methods that enhance adaptation without compromising pre-trained strengths remains a major goal. Recently, Singular Value Fine-Tuning (SVF) (Sun et al., 2022) has emerged as a promising alternative. Initially applied to CNNs for segmentation with strong results, SVF modifies only the singular values of weight matrices while preserving their directions without introducing new modules. Later, SAM-PARSER (Peng et al., 2024) extended SVF to large vision Transformer models, but limited its application to the vision encoder without fine-tuning query, key, and value (Q, K, V) matrices crucial for cross-modal tasks. Additionally, SVD-based methods have shown effectiveness in LLMs (Meng et al., 2024), but their potential for multi-modal VLM adaptation remains largely unexplored, offering an exciting direction for future research.

2.2 Adapting Vision-Language Models

Vision-language models, such as CLIP (Radford et al., 2021) and ALIGN (Jia et al., 2021), have significantly advanced multi-modal learning by aligning image and text embeddings in a shared space using self-supervised contrastive training. These models perform well on general-domain tasks like zero-shot classification and cross-modal retrieval, but their reliance on broad, non-specialized datasets limits their effectiveness in expert domains, such as healthcare, where nuanced visual cues and domain-specific semantics are critical. To address this, recent research has explored adapting VLMs to specialized settings using techniques like prompt learning, which offers a lightweight alternative to full fine-tuning. Methods such as CoOp (Zhou et al., 2022b) and CoCoOp (Zhou et al., 2022a) learn optimized prompts while keeping the VLM backbone frozen, with extensions like MaPLe (Khattak et al., 2023a) and PromptSRC (Khattak et al., 2023b) improving robustness through encoder tuning and self-regularization. Adapter-based strategies like CLIP-Adapter (Gao et al., 2024) and Tip-Adapter (Zhang et al., 2021) modify the visual branch or incorporate support-set features to boost few-shot performance, although they may face optimization hurdles. Enhanced probing methods such as LP++ (Huang et al., 2024) further refine adaptation by balancing modality-specific features with adaptive learning dynamics. In the biomedical domain, adaptations of CLIP like BioViL (Boecking et al., 2022), PubMedCLIP (Eslami et al., 2021), and BiomedCLIP (Zhang et al., 2024) leverage domain-specific corpora to improve relevance, with specific methods bridging general-purpose biomedical VLMs and specialized clinical tasks (Koleilat et al., 2024; 2025a; Spiegler et al., 2025; Rasaei et al., 2025). Yet, these models still struggle with fine-grained clinical understanding (Xu et al., 2024; Zhao et al., 2023). Prompt-learning methods, including XCoOp (Bie et al., 2024) and DCPL (Cao et al., 2024), extend CoOp-style tuning to medical applications, but often demand relatively large training sets. In comparison, BiomedCoOp (Koleilat et al., 2025b) demonstrates that prompt tuning can preserve generalization across diverse medical tasks even in low-resource conditions. Despite this broad range of techniques, no method has yet achieved robust performance across both natural and biomedical domains.

3 Method

3.1 CLIP Preliminaries

CLIP consists of a vision encoder E_v and a text encoder E_t that project images and text into a shared embedding space. Given a batch of B images and C distinct classes, image inputs $\mathbf{X}_v \in \mathbb{R}^{B \times 3 \times H \times W}$ are RGB images of height H and width W , and text inputs $\mathbf{X}_t \in \mathbb{R}^{C \times L}$ are tokenized sequences of length L , where each sequence serves as a text prompt representing a single class.

The encoders generate modality-specific features:

$$\mathbf{V} = E_v(\mathbf{X}_v) \in \mathbb{R}^{B \times D}, \quad \mathbf{T} = E_t(\mathbf{X}_t) \in \mathbb{R}^{C \times D} \quad (1)$$

where D is the embedding dimension. Both \mathbf{V} and \mathbf{T} are L2-normalized onto the unit hypersphere.

In zero-shot classification, CLIP matches an image to C class descriptions (e.g., "a photo of a [CLASS]"). The probability of assigning image embedding $\hat{\mathbf{V}}$ to class k is:

$$p(Y = k | \hat{\mathbf{V}}, \hat{\mathbf{T}}) = \frac{\exp(\hat{\mathbf{V}}^\top \hat{\mathbf{T}}^{(k)} / \tau)}{\sum_{j=1}^C \exp(\hat{\mathbf{V}}^\top \hat{\mathbf{T}}^{(j)} / \tau)} \quad (2)$$

where τ is a learnable temperature parameter. The predicted class \hat{k} is:

$$\hat{k} = \arg \max_k p(Y = k | \hat{\mathbf{V}}, \hat{\mathbf{T}}) \quad (3)$$

This formulation enables CLIP to generalize to unseen categories by leveraging the alignment between images and natural language descriptions.

3.2 Singular Value Decomposition of Weight Matrices

SVD-Based Decomposition of Pre-trained Weights: The overall framework of CLIP-SVD is shown in Fig. 1. For our proposed CLIP-SVD technique, we decompose the weight matrices in the **Multi-Head Self-Attention (MHSA)** and **Multi-Layer Perceptron (MLP)** blocks of each Transformer layer in CLIP’s text and image encoders with SVD. Specifically, each weight matrix W in the MHSA and MLP blocks can be factorized using SVD as follows:

$$W = USR^\top \quad (4)$$

where $U \in \mathbb{R}^{d \times r}$ is the left singular vector matrix, $S = \text{diag}(\lambda_1, \lambda_2, \lambda_3, \dots, \lambda_r) \in \mathbb{R}^{r \times r}$ is a diagonal matrix containing the singular values ($\lambda_1 \geq \lambda_2 \geq \dots \geq \lambda_r \geq 0$) arranged in a descending order, $R \in \mathbb{R}^{m \times r}$ is the right singular vector matrix, and $r = \min(d, m)$ is the rank of W . Instead of fully modifying W directly, we **freeze the singular vectors** U and R and **fine-tune only the singular values** λ_i . Note that we adapt the vector of full-rank singular values for our application. We further analyze the effect of different rank configurations in Appendix D.

Multi-Head Self-Attention Computation: Each Transformer layer in CLIP-type models applies MHSA using the Query (Q), Key (K), Value (V), and Output (O) projection matrices:

$$W_Q, W_K, W_V \in \mathbb{R}^{D \times d}, \quad W_O \in \mathbb{R}^{d \times D}. \quad (5)$$

Given an input $X \in \mathbb{R}^{B \times L \times D}$, self-attention for the h^{th} head is computed as:

$$Q_h = XW_{Q_h} = X(U_{Q_h}S_{Q_h}R_{Q_h}^\top), K_h = XW_{K_h} = X(U_{K_h}S_{K_h}R_{K_h}^\top) \quad (6)$$

$$Z_h = \text{softmax}\left(\frac{Q_h K_h^\top}{\sqrt{d}}\right)(XW_{V_h}) = \text{softmax}\left(\frac{Q_h K_h^\top}{\sqrt{d}}\right)(XU_{V_h}S_{V_h}R_{V_h}^\top) \quad (7)$$

where B is the batch size, L is the sequence length, D is the embedding dimension, and d is the dimension of each attention head.

For G attention heads of a Transformer layer:

$$Z_{\text{MHSA}} = \text{Concat}(Z_1, \dots, Z_G)(W_O) = \text{Concat}(Z_1, \dots, Z_G)(U_O S_O R_O^\top). \quad (8)$$

The output of the multi-head self-attention is then combined with the input via a residual connection:

$$X' = X + Z_{\text{MHSA}}. \quad (9)$$

Feed-forward Network: Following the self-attention block, the updated representation X' is passed through a feedforward MLP block ($\{W_{in}, W_{out}\}$), which is also decomposed using SVD. The MLP applies two linear transformations with an activation function in between, and finally, a residual connection is applied:

$$H = \text{ReLU}(X'W_{in}) = \text{ReLU}(X'U_{in}S_{in}R_{in}^\top) \quad (10)$$

$$X'' = H(W_{out}) = H(U_{out}S_{out}R_{out}^\top) \quad (11)$$

$$X_{\text{out}} = X' + X''. \quad (12)$$

With CLIP-SVD, each Transformer layer maintains its original representational capacity by rescaling the singular vectors, thus allowing robust adaptation and retention of pretrained knowledge.

3.3 Semantic interpretation of CLIP-SVD with TextSpan

To understand how our adaptation reshapes CLIP’s internal representations, we utilize **TextSpan** (Gandelsman et al., 2023), which aligns text descriptions to each attention head h of layer l in the ViT image encoder to reveal their semantic roles. This is achieved by decomposing the MHSA outputs $x^{l,h}$ into a summation of contributions from image tokens through SVD:

$$x^{l,h} = \sum_{i=0}^N x_i^{l,h}, \quad x_i^{l,h} = \alpha_i^{l,h} W_O^{l,h} W_V^{l,h} z_i^{l-1} = \alpha_i^{l,h} (U^{l,h} S^{l,h} V^{l,h\top}) z_i^{l-1}, \quad (13)$$

where $\alpha_i^{l,h}$ denotes the attention weight from the class token to token i , and z_i^{l-1} is the input token representation. **TextSpan** interprets the semantic roles of heads by projecting candidate text features $\mathbf{T}_{\text{corpus}}$ into the span of $x^{l,h}$:

$$\tilde{\mathbf{T}}^{l,h} = x^{l,h} (x^{l,h\top} x^{l,h})^{-1} x^{l,h\top} \mathbf{T}_{\text{corpus}}, \text{ with } P^{l,h} = x^{l,h} \cdot \tilde{\mathbf{T}}^{l,h\top}, \quad (14)$$

and identifies directions that maximize explained variance in $P^{l,h}$. Importantly, this procedure depends only on the span of the singular vectors $U^{l,h}$, not their magnitudes. In contrast, CLIP-SVD finetunes only the singular values $S^{l,h}$:

$$\tilde{x}_i^{l,h} = \alpha_i^{l,h} U^{l,h} \tilde{S}^{l,h} V^{l,h\top} z_i^{l-1} = \alpha_i^{l,h} \sum_j \tilde{s}_j^{l,h} \langle v_j^{l,h}, z_i^{l-1} \rangle u_j^{l,h}, \quad (15)$$

which preserves the Output-Value (OV) subspace $\text{span}(U^{l,h})$ while reweighting its basis directions through $\tilde{s}_j^{l,h}$. Thus, **TextSpan** and CLIP-SVD are complementary: the former reveals *which* semantic directions are encoded in $\text{span}(U^{l,h})$, while the latter modulates *how much* each direction contributes after adaptation. To quantify these effects, we rank heads by the normalized change in their singular values $S_O^{l,h}$ and $S_V^{l,h}$, summing absolute changes across both matrices. This provides a direct measure of the functional shifts in heads that **TextSpan** already grounds in interpretable text semantics.

4 Experiments and Results

4.1 Benchmark evaluation settings

Few-Shot Learning: To assess the model’s performance under limited supervision, we conduct few-shot image classification experiments with varying numbers of labeled examples per class ($K = 1, 2, 4, 8$, and 16

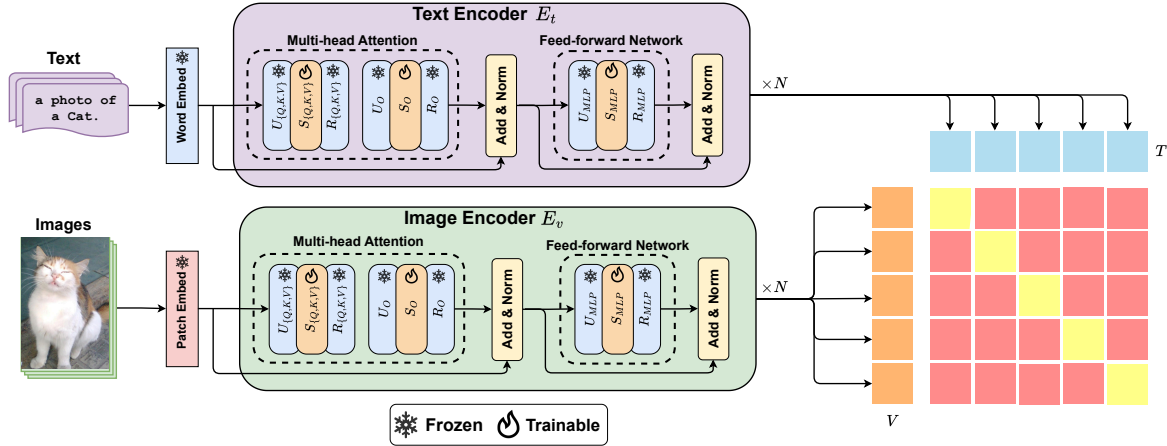


Figure 1: The overall framework of CLIP-SVD. We decompose the Query, Key, Value, and Output projection weights (W_Q , W_K , W_V and W_O) of the MHSA blocks in both vision and text encoders E_v and E_t , as well as the linear weights of the Feed-forward Networks (W_{MLP}) in all layers. We finetune only the singular values S of the SVD decomposed weights.

shots) to assess the robustness of our method across both natural and biomedical domains. This is critical for evaluating the method’s ability to learn effectively from sparse data by obtaining task-specific knowledge while retaining general domain comprehension.

Base-to-Novel Generalization: We evaluate the generalizability of CLIP-SVD in natural and biomedical domains, and follow a zero-shot setting, where the datasets are split into base and novel classes for classification tasks. Here, the model is trained only on the base classes in a few-shot setting and evaluated on both base and novel categories. Additionally, we compute the harmonic mean (HM) of both base and novel class prediction accuracies.

Cross-dataset Evaluation: To validate the performance of our approach in cross-dataset transfer, we evaluate our ImageNet-trained model directly on other datasets in the natural domain. Consistent with previous methods, our model is trained on all 1000 ImageNet classes in a few-shot manner. Due to the lack of a similar ImageNet-like dataset and large domain shifts across datasets, we didn’t perform cross-dataset evaluation for the biomedical domain.

Datasets: For the natural domain, we follow Zhou et al. (2022b;a) and evaluate the performance of our method on 11 image classification datasets that cover a wide range of recognition tasks. This includes two generic-objects datasets, ImageNet (Deng et al., 2009) and Caltech101 (Fei-Fei et al., 2004); five fine-grained class-specific datasets, OxfordPets (Parkhi et al., 2012), StanfordCars (Krause et al., 2013), Flowers102 (Nilsback & Zisserman, 2008), Food101 (Bossard et al., 2014), and FGVCAircraft (Maji et al., 2013); a scene recognition dataset SUN397 (Xiao et al., 2010); an action recognition dataset UCF101 (Soomro et al., 2012); a texture dataset DTD (Cimpoi et al., 2014); and a satellite-image dataset EuroSAT (Helber et al., 2019). For the biomedical domain, we follow Koleilat et al. (2025b) and evaluate the performance of our method on 10 diverse medical imaging datasets covering 9 different organs and 8 imaging modalities: Computerized Tomography (CTKidney (Islam et al., 2022)), Endoscopy (Kvasir (Pogorelov et al., 2017)), Fundus Photography (RETINA (Porwal et al., 2018; Köhler et al., 2013)), Histopathology (LC25000 (Borkowski et al., 2019), CHMNIST (Kather et al., 2016)), brain tumor Magnetic Resonance Imaging (BTMRI (Nickparvar, 2021)), Optical Coherence Tomography (OCTMNIST (Kermany et al., 2018)), breast ultrasound (BUSI (Al-Dhabyani et al., 2020)), and chest and knee X-Ray (COVID-QU-Ex (Tahir et al., 2021), KneeXray (Chen, 2018)).

Implementation Details We use a few-shot training strategy in all experiments, with random sampling for each class. We use the ViT-B/16 CLIP and BiomedCLIP models for natural and biomedical domains, respectively. All models are trained using a batch size of 32 and the AdamW optimizer (Loshchilov & Hutter, 2017) with a weight decay of 0.01 on a single NVIDIA A100 GPU (40GB RAM). For natural image datasets,

we set the learning rate at 5×10^{-4} for few-shot classification, 6×10^{-4} for base-to-novel evaluations, and 5×10^{-4} for cross-dataset transfer tasks. In the biomedical domain, learning rates are optimized per dataset based on validation performance, depending on task complexity and data modality. We report classification accuracies averaged over three independent runs. Prompt templates and full hyperparameter configurations are detailed in Appendix A and B, respectively.

4.2 Few-shot Evaluation

Our method demonstrates superior performance in few-shot learning across both natural and biomedical domains, as shown in Tables 2 and 3. In the natural domain, CLIP-SVD achieves a +1.00% improvement over the second-best method (CLIP-LoRA) in the 1-shot setting (73.20% vs. 72.20%). In the biomedical domain, it surpasses the second-best approach (BiomedCoOp) by +4.28% in the 8-shot setting (73.24% vs. 68.96%). These consistent gains highlight the robustness and effectiveness of our SVD-based tuning strategy across diverse domains.

Table 2: **Evaluation against state-of-the-art techniques for natural domain:** The average classification accuracy (%) obtained from 11 benchmarks derived from 3 sampled support sets for each dataset. The top-performing results are in bold, and the second-best are underlined.

Method	$K = 1$	$K = 2$	$K = 4$	$K = 8$	$K = 16$
Zero-shot CLIP (Radford et al., 2021)			65.36		
CoOp (Zhou et al., 2022b)	68.09	70.13	73.59	76.45	79.01
CoCoOp (Zhou et al., 2022a)	66.95	67.63	71.98	72.92	75.02
ProGrad (Zhu et al., 2024a)	68.20	71.78	74.21	77.93	79.20
KgCoOp (Yao et al., 2023a)	69.51	71.57	74.48	75.82	77.26
MaPLe (Khattak et al., 2023a)	69.27	72.58	75.37	78.89	81.79
Linear Probing (Radford et al., 2021)	45.77	56.92	66.79	73.43	78.39
LP++ (Huang et al., 2024)	70.35	72.93	75.77	77.94	80.32
CLIP-Adapter (Gao et al., 2024)	67.87	70.20	72.65	76.92	79.86
Tip-Adapter (Zhang et al., 2021)	68.89	70.42	72.69	74.41	76.44
Tip-Adapter-F (Zhang et al., 2021)	70.62	73.08	75.75	78.51	81.15
GDA (Wang et al., 2024)	69.39	73.09	76.24	79.71	81.70
ProKeR (Bendou et al., 2025)	71.32	73.74	76.23	79.84	82.01
AdaLoRA (Zhang et al., 2023a)	69.04	72.21	75.50	78.13	80.95
TCP (Yao et al., 2024)	70.63	73.59	76.07	78.39	80.98
CLIP-LoRA (Zanella & Ben Ayed, 2024)	<u>72.20</u>	<u>75.41</u>	<u>77.32</u>	<u>80.10</u>	<u>82.89</u>
CLIP-SVD (Ours)	73.20	76.06	78.18	80.55	82.97

4.3 Base-to-Novel Generalization

Our method demonstrates strong base-to-novel generalization across both natural and biomedical domains, as shown in Tables 4 and 5. In the natural domain, CLIP-SVD improves over MaPLe by +1.67% on base accuracy, +1.06% on novel accuracy, and +1.58% on the harmonic mean, despite MaPLe being approximately 38× more computationally expensive. In the biomedical domain, CLIP-SVD achieves substantial gains over BiomedCoOp, improving base accuracy by +4.04%, novel accuracy by +0.41%, and the harmonic mean by +4.21%. These results highlight the robustness and scalability of our SVD-based tuning approach across domains. In addition, they also demonstrate the benefit of multi-modal tuning. The proposed CLIP-SVD enhances generalization without compromising the powerful representations learned during CLIP’s pretraining.

4.4 Cross-dataset Transfer

Table 6 shows that CLIP-SVD achieves the highest average accuracy of 66.99%, slightly outperforming MaPLe’s 66.30%. It obtains the best performance on several target datasets, including Aircraft (+1.29%),

Table 3: **Evaluation against state-of-the-art techniques for biomedical domain:** The average classification accuracy (%) obtained from 10 benchmarks derived from 3 sampled support sets for each dataset. The top-performing results are in bold, and the second-best are underlined.

Method	$K = 1$	$K = 2$	$K = 4$	$K = 8$	$K = 16$
Zero-shot BiomedCLIP (Zhang et al., 2024)			42.38		
CoOp (Zhou et al., 2022b)	52.59	55.71	61.35	67.74	71.48
CoCoOp (Zhou et al., 2022a)	50.88	53.91	57.63	63.15	67.51
ProGrad (Zhu et al., 2023)	53.67	56.42	62.10	67.06	69.21
KgCoOp (Yao et al., 2023b)	54.31	55.79	60.92	66.00	67.71
Linear Probing (Radford et al., 2021)	48.91	55.82	62.12	67.33	70.81
LP++ (Huang et al., 2024)	49.27	55.88	61.30	65.48	70.09
CLIP-Adapter (Gao et al., 2024)	45.53	44.70	45.30	46.54	48.46
Tip-Adapter (Zhang et al., 2021)	50.35	53.50	58.33	62.01	67.60
Tip-Adapter-F (Zhang et al., 2021)	52.55	54.17	62.30	68.12	68.12
GDA (Wang et al., 2024)	49.56	58.39	63.41	70.60	72.86
ProKeR (Bendou et al., 2025)	49.40	58.84	63.72	70.98	71.86
XCoOp (Bie et al., 2024)	52.50	55.39	60.87	66.37	71.04
DCPL (Cao et al., 2024)	49.65	58.65	62.62	68.65	70.79
CLIP-LoRA (Zanella & Ben Ayed, 2024)	48.31	57.63	62.31	68.16	70.31
MaPLe (Khattak et al., 2023a)	37.99	40.89	44.09	47.37	52.93
BiomedCoOp (Koleilat et al., 2025b)	56.87	<u>59.32</u>	<u>64.34</u>	<u>68.96</u>	<u>73.41</u>
CLIP-SVD (Ours)	<u>56.35</u>	62.63	68.02	73.26	76.46

Table 4: **Base-to-novel generalization** comparison measured by classification accuracy (%) between CLIP-SVD and SOTA methods on 11 natural domain datasets.

Acc.	CLIP	CoOp	CoCoOp	KgCoOp	ProGrad	MaPLe	IVLP	GDA	TCP	CLIP-LoRA	CLIP-SVD
Base	69.34	82.69	80.47	80.73	82.48	82.28	84.21	83.96	84.13	84.10	84.38
Novel	74.22	63.22	71.69	73.60	70.75	<u>75.14</u>	71.79	74.53	75.36	74.80	76.29
HM	71.70	71.66	75.83	77.00	76.16	78.55	77.51	78.72	79.51	<u>79.18</u>	80.13

SUN397 (+0.73%), and UCF101 (+1.22%). These results suggest that CLIP-SVD offers strong cross-dataset transfer capabilities, confirming its potential for effective generalization.

4.5 Ablation Experiments: Selective Model Component Fine-tuning

Effect of Tuning Different Weights: We ablated CLIP-SVD’s components (W_Q , W_K , W_V , W_O , and W_{MLP}) under a 4-shot setting in natural and biomedical domains (see Table 7). Without adaptation, the accuracy was 65.36% (natural) and 42.38% (biomedical). Adding W_O alone substantially improved performance (75.45% and 62.27%), highlighting its importance. Including W_{MLP} further boosted accuracy (77.78% and 66.40%), showing its role in fine-grained transformation. Adding W_Q , W_K , or W_V individually on top of W_O and W_{MLP} led to near-identical results in the natural domain, but slight gains in biomedical, up to 67.40% with W_V . Using all components yielded the best performance (natural: 78.18% and biomedical: 68.02%), confirming the benefit of full adaptation. Previously, Biderman et al. (2024) observed similar trends when finetuning LLMs in math and coding tasks with LoRA, where a more prominent impact is also seen with adapting MLP than the attention layers.

Table 5: **Base-to-novel generalization** comparison measured by classification accuracy (%) between CLIP-SVD and SOTA methods on 10 biomedical domain datasets.

Acc.	BiomedCLIP	CoOp	CoCoOp	KgCoOp	ProGrad	MaPLe	XCoOp	BiomedCoOp	GDA	DCPL	CLIP-LoRA	CLIP-SVD
Base	49.27	76.71	75.52	71.90	75.69	65.40	74.62	<u>78.60</u>	57.70	73.70	70.56	82.64
Novel	67.17	65.34	67.74	65.94	67.33	49.51	63.19	<u>73.90</u>	64.66	69.35	59.84	74.31
HM	55.23	68.80	69.11	67.22	69.86	53.10	68.43	<u>74.04</u>	60.98	71.46	64.76	78.25

Table 6: **Cross-dataset natural image benchmark** with classification accuracy (%)

	Source											Target
	ImageNet	Caltech101	OxfordPet	StanfordCars	Flowers102	Food101	Aircraft	SUN397	DTD	EuroSAT	UCF101	
CLIP	66.72	92.98	89.13	65.29	71.30	86.11	<u>24.90</u>	62.59	44.56	<u>47.84</u>	66.83	65.15
CoOp	71.51	93.70	89.14	64.51	68.71	85.30	18.47	64.15	41.92	46.39	66.55	63.88
Co-CoOp	71.02	94.43	90.14	65.32	71.88	86.06	22.94	<u>67.36</u>	45.73	45.37	68.21	65.74
KgCoOp	70.66	<u>93.92</u>	89.83	<u>65.41</u>	70.01	86.36	22.51	66.16	<u>46.35</u>	46.04	68.50	65.51
ProGrad	72.24	91.52	89.64	62.39	67.87	85.40	20.61	62.47	39.42	43.46	64.29	62.71
MaPLe	70.72	93.53	90.49	65.57	<u>72.23</u>	86.20	24.74	67.01	46.49	48.06	<u>68.69</u>	<u>66.30</u>
CLIP-SVD	<u>72.15</u>	93.68	91.06	65.00	72.45	<u>86.21</u>	26.03	67.74	45.15	47.51	69.91	66.99

Table 8: Top 3 descriptions returned by TextSpan (Gandelsman et al., 2023) applied to the attention head with the greatest adaptation-related change for different datasets.

EuroSAT (L10.H0)	BUSI (L11.H8)	DTD (L8.H6)
Aerial view of an agricultural field Image taken in the Namibian desert Picture taken in the Brazilian rainforest	A low-contrast region in a clustered pattern A double-density sign suggesting benignity A solid-cystic component suggesting malignancy	Collage of textures Close-up of a textured bark Mesmerizing kinetic sculpture
BTMRI (L8.H9)	SUN397 (L11.H0)	COVID-QU-Ex (L11.H0)
An area with decreased perfusion in the left hemisphere A bright spot artifact in a clustered pattern A contrast-enhanced region on axial view	Mysterious day scene Urban rooftop panorama A zoomed out photo	A collapsed lung lobe A low signal-to-noise ratio in the upper lobe A lesion crossing compartments
UCF101 (L10.H5)	CTKIDNEY (L8.H6)	RETINA (L8.H4)
Dynamic action Energetic children Playful winking facial expression	An anatomical displacement A spiculated margin A zone of tissue infiltration	An area with decreased perfusion A vascular displacement A vascular structure with sharp borders

Effect of Tuning Image and Text Encoders: The *right panel* of Fig. 2 illustrates how different input modality tuning setups affect classification accuracy. Multi-modal tuning (text+image) consistently outperforms unimodal cases in both domains. In the natural domain, text-only and image-only achieve 75.78% and 74.31% accuracy, while their combination reaches 78.18%. In the biomedical domain, multi-modal tuning yields 68.02%, compared to 64.21% (text) and 65.94% (image), suggesting the complementary nature of visual and textual cues, especially valuable in biomedical settings with limited data and higher complexity.

Effect of Tuning Different Layers: The *left panel* of Fig. 2 shows how freezing different sets of Transformer layers that are matched in both text and image encoders during SVD-based adaptation affects few-shot accuracy in natural and biomedical domains. In the natural domain, accuracy stays relatively stable, dropping only slightly from 78.18% (all layers adapted) to 77.35% (first four layers frozen) and 75.36% (first and last four frozen), suggesting robust, distributed representations. In contrast, the biomedical domain is more sensitive: accuracy drops from 68.02% (all layers adapted) to 66.68% (last four frozen), 64.38% (first and last four), and 62.87% (top eight frozen), indicating that deeper layers are more critical for capturing domain-specific complexity.

4.6 Natural Language-based Interpretation of CLIP-SVD

Natural language offers a powerful but under-explored lens into the conceptual space of VLMs. To gain insights into the impact of CLIP-SVD, we investigate how textual descriptions align with CLIP’s internal representations. Besides the natural domain, we present the first **systematic, text-based analysis** of a

Table 7: Impact of each component of the proposed CLIP-SVD on the 4-shot accuracy (%) of natural and biomedical domain benchmarks.

W_Q	W_K	W_V	W_O	W_{MLP}	Natural	Biomedical
×	×	×	×	×	65.36	42.38
×	×	×	×	✓	76.69	65.27
×	×	×	✓	×	75.45	62.27
×	×	×	✓	✓	77.78	66.40
✓	✓	✓	×	×	77.15	65.35
✓	✓	✓	×	✓	77.83	67.50
✓	✓	✓	✓	×	77.88	66.74
✓	×	×	✓	✓	78.12	66.99
×	✓	×	✓	✓	78.12	67.09
×	×	×	✓	✓	78.11	67.40
✓	✓	✓	✓	✓	78.18	68.02

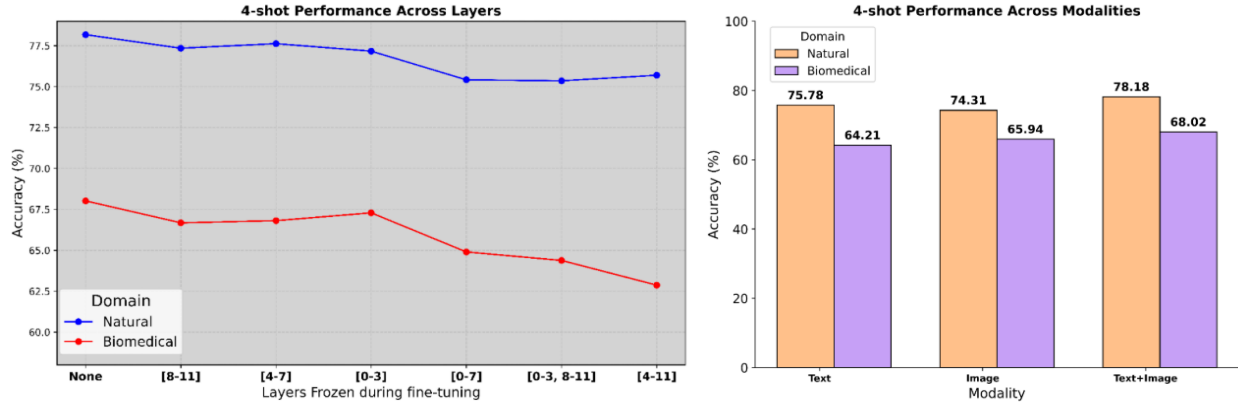


Figure 2: 4-shot performance by freezing certain layers during finetuning (*left*) and by adapting text encoder and/or image encoder (*right*) for natural and biomedical domains.

biomedical VLM at the **attention head level**, focusing on how fine-tuning, such as in BiomedCLIP, reshapes semantic VLM’s representations. To support this, similar to Gandelsman et al. (2023), we constructed a new *biomedical caption corpus* of 300 clinically relevant text elements using GPT-4 (Achiam et al., 2023), describing features like contrast, shape, and texture. This enables interpretable alignment between vision and language representations and domain-targeted probing of attention heads. Our framework combines **CLIP-SVD** with **TextSpan** (Gandelsman et al., 2023) to quantify semantic shifts in Output-Value circuits of ViT backbones, focusing on the last four layers (Gandelsman et al., 2023). By ranking the attention heads via singular value shift magnitude from CLIP-SVD and extracting aligned text spans, we gain an intuitive interpretation for the importance of different attention heads during model adaptation and their roles in the finetuned tasks (see Tables 1 and 8). These analyses highlight how finetuning steers VLMs toward task-relevant, domain-specific understanding. Additionally, the insights could allow debugging and further refinement (e.g., with prompt engineering) of task/domain-specific CLIP models.

5 Discussion

Compared with SVF (Sun et al., 2022) that targeted CNNs and single-modal learning, our CLIP-SVD introduces the first SVD-based adaptation for multi-modal Transformers to modulate attention jointly in vision and text to offer excellent performance while better preserving model generalization. Furthermore, by rescaling semantic subspace vectors of pretrained CLIP models via singular value modulation, CLIP-SVD explicitly reveals which pretrained feature directions are reused, suppressed, or amplified. This subspace-level interpretability offers a new lens into VLM adaptation mechanisms, allowing diagnosis of fine-grained adaptation dynamics, which is an aspect absent from prior works like CLIP-LoRA that introduce new degrees of freedom and thus obscure these observations.

Recent studies (Shuttleworth et al., 2025) on LLM finetuning reveal that LoRA and its variants could result in shifted singular vectors of the model parameter space (called intruder dimensions), potentially causing forgetting of past knowledge. On the other hand, our CLIP-SVD leverages the rich semantic representation of VLM models by freezing the pretrained singular vectors U and V of the parameter space and tuning only the singular values Σ . This allows us to recalibrate the “importance” of task-relevant subspaces without distorting the original model geometry. While freezing U and V may potentially limit the expressivity of the parameter space, we find that this is both suitable and effective in the few-shot setting in our work, provided that CLIP-like models have been trained extensively with fine-grained representations. Unlike fully supervised scenarios common in LLM fine-tuning, our adaptation method assumes access to only a handful of labeled examples per class. In such cases, introducing large numbers of randomly initialized or fully tunable parameters will be suboptimal. Since many downstream tasks lie in a low intrinsic dimension, and pretraining implicitly shapes these subspace vectors, fine-tuning via a low-dimensional subspace often

suffices (Aghajanyan et al., 2021). Additionally, our interpretability analysis using TextSpan (Gandelsman et al., 2023) also shows that tuning only Σ results in semantically meaningful shifts in attention.

In our experiments, few-shot VLM adaptation in the biomedical vision domain is notably more challenging than in the natural domain, likely due to factors like complex and unintuitive image features and ambiguous boundaries, as shown in previous works (Koleilat et al., 2025b). We show that CLIP-SVD bridges this domain gap, achieving strong performance in both biomedical and natural vision settings. In contrast, as shown in Tables 2 and 3, methods like CLIP-LoRA that were proposed for natural domain benchmarks do not generalize as well to biomedical tasks.

We adopted a natural language-based technique to understand the impact and insights of CLIP-SVD in model adaptation. For the related analyses in the biomedical domain, we constructed a new corpus of biomedical image descriptions by leveraging the large language models. Although it is shown to facilitate the interpretation, further validation is still required in broader applications, particularly with domain experts. We will investigate this in the near future.

6 Conclusion

In conclusion, we introduced CLIP-SVD, a novel parameter-efficient adaptation method for CLIP models that finetunes only the singular values of the weight matrices while preserving its pre-trained structure. Our approach enables effective few-shot learning with minimal computational overhead, achieving state-of-the-art results across both natural and biomedical domains. Through extensive ablation studies, we demonstrated the critical role of singular value adaptation in enhancing task-specific feature extraction. Further analyses of the Output-Value circuit with a natural-language-based approach revealed that adapting singular values steers attention heads toward more specialized roles, thus opening doors for further investigation of CLIP’s characteristics in broader applications.

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A Dataset Details

Following Zhou et al. (2022b) and Koleilat et al. (2025b), we conducted extensive experiments on 11 natural and 10 biomedical classification benchmark datasets to evaluate the effectiveness of the proposed CLIP-SVD. The natural datasets include ImageNet (Deng et al., 2009), Caltech101 (Fei-Fei et al., 2004), OxfordPets (Parkhi et al., 2012), StanfordCars (Krause et al., 2013), Flowers102 (Nilsback & Zisserman, 2008), Food101 (Bossard et al., 2014), FGVCAircraft (Maji et al., 2013), SUN397 (Xiao et al., 2010), DTD (Cimpoi et al., 2014), EuroSAT (Helber et al., 2019), and UCF101 (Soomro et al., 2012). The biomedical datasets consist of CTKidney (Islam et al., 2022), DermaMNIST (Tschandl et al., 2018; Codella et al., 2019), Kvasir (Pogorelov et al., 2017), RETINA (Porwal et al., 2018; Köhler et al., 2013), LC25000 (Borkowski et al., 2019), CHMNIST (Kather et al., 2016), BTMRI (Nickparvar, 2021), OCTMNIST (Kermany et al., 2018), BUSI (Al-Dhabyani et al., 2020), COVID-QU-Ex (Tahir et al., 2021), and KneeXray (Chen, 2018). For distribution shift experiments, we also included ImageNetV2 (Recht et al., 2019), ImageNet-Sketch (Wang et al., 2019), ImageNet-A (Hendrycks et al., 2021b), and ImageNet-R (Hendrycks et al., 2021a). Dataset statistics are provided in Tables S3 and S4.

B Detailed Hyperparameters

The hyperparameters reported in Tables S1 and S2 across both natural and biomedical datasets were carefully tuned based on benchmark type and dataset characteristics. A consistent batch size (BS) of 32 was maintained throughout to ensure uniformity in training dynamics. For natural datasets, the learning rate (LR) was set to 5×10^{-4} for few-shot settings and reduced to 6×10^{-4} for base-to-novel evaluations. For the natural few-shot setting, we follow Zanella & Ben Ayed (2024) and set the number of iterations to $200 \times K$ (number of shots), whereas we use epochs for the base-to-novel setting, with the number of training epochs (EP) varying between 2 and 20 depending on the dataset’s size. In biomedical datasets, LR values were more diverse, ranging from 0.5×10^{-3} to 20×10^{-3} , reflecting the varying difficulty and modality of the medical tasks, while EP was generally higher (up to 200) to accommodate the typically smaller dataset sizes and slower convergence. Notably, BUSI (Al-Dhabyani et al., 2020) lacked base-to-novel evaluation due to the absence of appropriate class splits.

C Domain Generalization

We evaluate the robustness of our method on out-of-distribution datasets. Similar to cross-dataset evaluation, we test our ImageNet-trained model directly on four other ImageNet datasets with different types of distribution shifts, including ImageNetV2 (Recht et al., 2019), ImageNet-Sketch (ImageNet-S) (Wang et al., 2019), ImageNet-A (Hendrycks et al., 2021b), and ImageNet-R (Hendrycks et al., 2021a). Table S5 presents the domain generalization results, where methods are trained on ImageNet and evaluated on datasets with various domain shifts (-V2, -S, -A, and -R). CLIP-SVD achieves the highest average accuracy of 62.55%, outperforming MaPLe (62.36%) and other methods. It also leads in two target domains, namely ImageNet-S (-S) (+0.47%) and ImageNet-V2 (-V2) (+0.28%), demonstrating its robustness to domain shifts. These results highlight the effectiveness of CLIP-SVD in generalizing across domains.

D Effect of Rank-based Singular Value Selection for CLIP-SVD

For CLIP-SVD, selective finetuning of the singular values could further reduce the computational requirement. With singular value decomposition, the singular values are naturally ranked in descending order with respect to their magnitude. In this experiment, we varied the number of non-zero singular values out of the full rank to be adapted according to their magnitude ordering, in order to investigate their impact on 4-shot model performance across both natural and biomedical domains. Specifically, we tested two configurations: one where only the top singular values (in descending order) are fine-tuned, and the other where only the bottom singular values (in ascending order) are adjusted. The results shown in Fig. S1 reveal that, in the natural domain, both top and bottom configurations exhibit a steep initial accuracy increase starting from a small proportion of adjustable singular values, stabilizing near 78.18% as the ratio approaches full-rank,

Dataset	Benchmark	BS	LR (10^{-4})	EP/IT
ImageNet	Few-shot	32	5	200
	Base-to-Novel	32	6	10
	Cross-dataset Transfer	32	5	2
	Domain Generalization	32	5	2
Caltech101	Few-shot	32	5	200
	Base-to-Novel	32	6	20
DTD	Few-shot	32	5	200
	Base-to-Novel	32	6	20
EuroSAT	Few-shot	32	5	200
	Base-to-Novel	32	6	20
StanfordCars	Few-shot	32	5	200
	Base-to-Novel	32	6	10
Flowers102	Few-shot	32	5	200
	Base-to-Novel	32	6	10
FGVCAircraft	Few-shot	32	5	200
	Base-to-Novel	32	6	20
SUN397	Few-shot	32	5	200
	Base-to-Novel	32	6	10
OxfordPets	Few-shot	32	5	200
	Base-to-Novel	32	6	10
UCF101	Few-shot	32	5	200
	Base-to-Novel	32	6	10
Food101	Few-shot	32	5	200
	Base-to-Novel	32	6	10

Table S1: Hyperparameter values across natural datasets and benchmarks. (BS = Batch Size, LR = Learning Rate, EP = Epochs, IT = Iterations)

Dataset	Benchmark	BS	LR (10^{-3})	EP
BTMRI	Few-shot	32	7	100
	Base-to-Novel	32	9	100
BUSI	Few-shot	32	2	100
	Base-to-Novel	-	-	-
COVID-QU-Ex	Few-shot	32	0.5	100
	Base-to-Novel	32	2	50
CTKIDNEY	Few-shot	32	1	200
	Base-to-Novel	32	10	200
Kvasir	Few-shot	32	5	100
	Base-to-Novel	32	3	60
CHMNIST	Few-shot	32	1	60
	Base-to-Novel	32	7	100
LC25000	Few-shot	32	3	100
	Base-to-Novel	32	10	100
RETINA	Few-shot	32	7	100
	Base-to-Novel	32	20	60
KneeXray	Few-shot	32	8	100
	Base-to-Novel	32	2	60
OCTMNIST	Few-shot	32	10	100
	Base-to-Novel	32	20	100

Table S2: Hyperparameter values across biomedical datasets and benchmarks. (BS = Batch Size, LR = Learning Rate, EP = Epochs)

suggesting a limited sensitivity to the ranking of the selected singular values to be adapted. In contrast, the biomedical domain shows a more pronounced dependency on the ranking of the adjustable singular values. Here, finetuning the top singular values consistently outperforms the bottom configuration, reaching 68.02% at a full rank ratio, while the bottom approach lags behind by approximately 5% at the 0.5 ratio, highlighting the importance of prioritizing top singular values for specialized medical contexts.

E Experiments with Other Backbones

In this experiment, we evaluated our proposed method, CLIP-SVD, using the alternative ViT-B/32 backbone in a few-shot learning setting for the natural domain. Table S6 presents the average classification accuracy (%) across 11 natural domain benchmarks derived from three randomly sampled support sets for each dataset. Our method consistently outperforms the state-of-the-art techniques across different few-shot settings. Notably, CLIP-SVD achieves the highest performance in all cases, surpassing the second-best method, Tip-Adapter-F, by a significant margin, with an improvement of approximately 1.2% at $K = 1$ and 1.4% at $K = 16$. These results highlight the strength of our singular value decomposition-based adaptation strategy in enhancing generalization under limited data conditions. Furthermore, the effectiveness of CLIP-SVD with the ViT-B/32 backbone demonstrates that our approach is adaptable to different model backbones, making it broadly applicable across various vision-language models.

F TextSpan Analysis Details

We analyze singular value shifts in the Output-Value (OV) circuit using TextSpan (Gandelsman et al., 2023), focusing on the last four layers (L8–L11) of CLIP and BiomedCLIP ViT-B/16 (*Note: layer and head*

Table S3: **Summary of natural datasets:** Overview of the datasets used in the natural domain, including the number of classes, dataset splits (train, validation, test), and the corresponding hand-crafted text prompts used for classification.

Dataset	Classes	Train	Val	Test	Hand-crafted prompt
ImageNet	1,000	1.28M	N/A	50,000	"a photo of a [CLASS]."
Caltech101	100	4,128	1,649	2,465	"a photo of a [CLASS]."
OxfordPets	37	2,944	736	3,669	"a photo of a [CLASS], a type of pet."
StanfordCars	196	6,509	1,635	8,041	"a photo of a [CLASS]."
Flowers	102	4,093	1,633	2,463	"a photo of a [CLASS], a type of flower."
Food101	101	50,500	20,200	30,300	"a photo of [CLASS], a type of food."
FGVCAircraft	100	3,334	3,333	3,333	"a photo of a [CLASS], a type of aircraft."
SUN397	397	15,880	3,970	19,850	"a photo of a [CLASS]."
DTD	47	2,820	1,128	1,692	"[CLASS] texture."
EuroSAT	10	13,500	5,400	8,100	"a centered satellite photo of [CLASS]."
UCF101	101	7,639	1,898	3,783	"a photo of a person doing [CLASS]."
ImageNetV2	1,000	N/A	N/A	10,000	"a photo of a [CLASS]."
ImageNet-Sketch	1,000	N/A	N/A	50,889	"a photo of a [CLASS]."
ImageNet-A	1,000	N/A	N/A	50,889	"a photo of a [CLASS]."
ImageNet-R	1,000	N/A	N/A	50,889	"a photo of a [CLASS]."

Table S4: **Summary of biomedical datasets:** Overview of the datasets used in the biomedical domain, including the number of classes, dataset splits (train, validation, test), and the corresponding hand-crafted text prompts used for classification.

Dataset	Classes	Train	Val	Test	Hand-crafted prompt
CTKIDNEY	4	6,221	2,487	3,738	"a photo of a [CLASS]."
Kvasir	8	2,000	800	1,200	"a photo of a [CLASS]."
RETINA	4	2,108	841	1,268	"a photo of a [CLASS]."
LC25000	5	12,500	5,000	7,500	"a photo of a [CLASS]."
CHMNIST	8	2,496	1,000	1,504	"a photo of [CLASS]."
BTMRI	4	2,854	1,141	1,717	"a photo of a [CLASS]."
OCTMNIST	4	97,477	10,832	1,000	"a photo of a [CLASS]."
BUSI	3	389	155	236	"a photo of a [CLASS]."
COVID-Qu-Ex	4	10,582	4,232	6,351	"a chest xray of [CLASS]."
KneeXray	5	5,778	826	1,656	"a photo of a [CLASS]."

indexing in these experiments begins at 0.). The ViT-B/16 vision encoder employs 12 attention heads per MHSA block. For the natural image domain, we use ImageNet (Deng et al., 2009), while for the biomedical domain, we aggregate all relevant datasets (Islam et al., 2022; Pogorelov et al., 2017; Porwal et al., 2018; Köhler et al., 2013; Borkowski et al., 2019; Kather et al., 2016; Nickparvar, 2021; Kermayn et al., 2018; Al-Dhabyani et al., 2020; Tahir et al., 2021; Chen, 2018) into a single comprehensive dataset. Images from both domains are fed into their corresponding vision encoder network. For the natural domain, we utilize the text corpus from Gandelsman et al. (2023), which consists of approximately 3,000 GPT-3.5-generated general image descriptions. In contrast, no large-scale text corpus exists for biomedical images. To address this, we generated a new corpus comprising 300 general medical image descriptions and relevant terminologies using GPT-4 (Achiam et al., 2023). We use the following prompt to query GPT-4: "Generate 300 distinct descriptive prompts for medical images that capture specific visual features commonly found in medical scans, such as contrast, shape, color, location, and texture." Using TextSpan, we extract the top three text descriptions from the corpus that best characterize the role

Table S5: **Domain generalization:** Methods are trained on ImageNet using 16-shots and evaluated on datasets with domain shifts (ImageNet-V2, -S, -A, and -R) for classification accuracy (%).

	Source	Target				Avg.
	ImageNet	-V2	-S	-A	-R	
CLIP	66.73	60.83	46.15	47.77	73.96	59.09
CoOp	71.51	64.20	47.99	49.71	75.21	61.72
Co-CoOp	71.02	64.07	48.75	50.63	76.18	62.13
CLIP-Adapter	68.46	59.55	39.88	38.83	64.62	54.27
TIP-Adapter	53.81	45.69	29.21	36.04	55.26	44.00
TIP-Adapter-F	51.71	43.07	27.13	27.04	45.07	38.80
TaskRes	70.84	62.15	43.76	43.91	71.59	58.45
KgCoOp	71.20	64.10	48.97	50.69	76.70	62.33
ProGrad	72.24	64.73	47.61	49.39	74.58	61.71
MaPLe	70.72	64.07	49.15	50.90	76.98	62.36
CLIP-SVD	72.15	65.03	49.62	49.17	76.79	62.55

Table S6: **Evaluation against state-of-the-art techniques for natural domain with ViT-B/32 backbone:** This table presents the average classification accuracy (%) obtained from 11 natural domain benchmarks derived from 3 sampled support sets for each dataset. The top-performing results are in bold, and the second-best are underlined.

Method	$K = 1$	$K = 2$	$K = 4$	$K = 8$	$K = 16$
Zero-shot CLIP (Radford et al., 2021)			61.8		
CoOp (Zhou et al., 2022b)	62.8	65.3	68.6	72.2	74.7
CoCoOp (Zhou et al., 2022a)	63.1	64.8	66.7	68.1	70.7
ProGrad (Zhu et al., 2024a)	64.7	67.1	69.8	73.2	75.9
KgCoOp (Yao et al., 2023a)	64.9	66.6	68.4	70.5	72.1
MaPLe (Khattak et al., 2023a)	61.5	65.2	68.7	71.6	74.1
CLIP-Adapter (Gao et al., 2024)	62.5	63.5	64.3	67.6	71.6
Tip-Adapter-F (Zhang et al., 2021)	<u>66.5</u>	<u>69.2</u>	<u>71.6</u>	<u>74.1</u>	<u>77.1</u>
CLIP-SVD (Ours)	67.7	71.0	73.4	75.8	78.5

of each attention head in each of the final four layers. Based on these top-3 descriptions, GPT-4 is used to assign a concise and descriptive title to each head’s function: "Assign a concise and descriptive title that best captures the primary function represented by these descriptions.". The identified roles and associated top-3 descriptions for each head in layers L8–L11 for the natural and biomedical domains are detailed in Tables S10 and S11, respectively.

G t-SNE Visualization

We used t-SNE (van der Maaten & Hinton, 2008) to visualize the image embeddings generated by the fine-tuned model using our CLIP-SVD and compared them with the embeddings from the pre-trained CLIP and BiomedCLIP models. As shown in Figure S3, the embeddings produced by CLIP-SVD exhibit significantly better clustering, indicating that our method enables the model to generate more distinct and well-separated feature representations. This experiment was conducted on the EuroSAT, OxfordPets, CTXkidney, and LC25000 datasets, where the improved clustering patterns highlight the enhanced feature extraction capabilities of our approach.

H Segmentation Performance

We evaluated the segmentation performance of our fine-tuned model using the TextSpan algorithm (Gandelsman et al., 2023) on the ImageNet-Segmentation dataset (Guillaumin

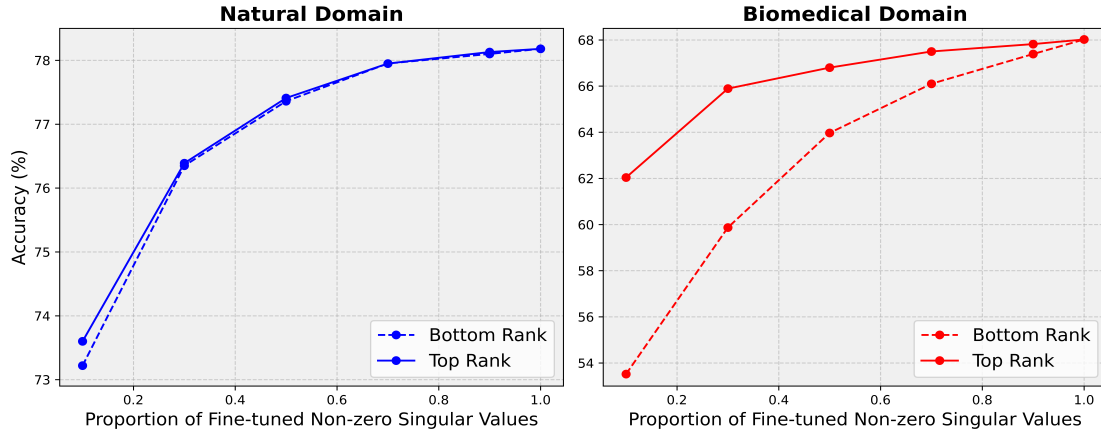


Figure S1: Accuracy comparison for models finetuned with varying rank ratios in the natural (left) and biomedical (right) domains

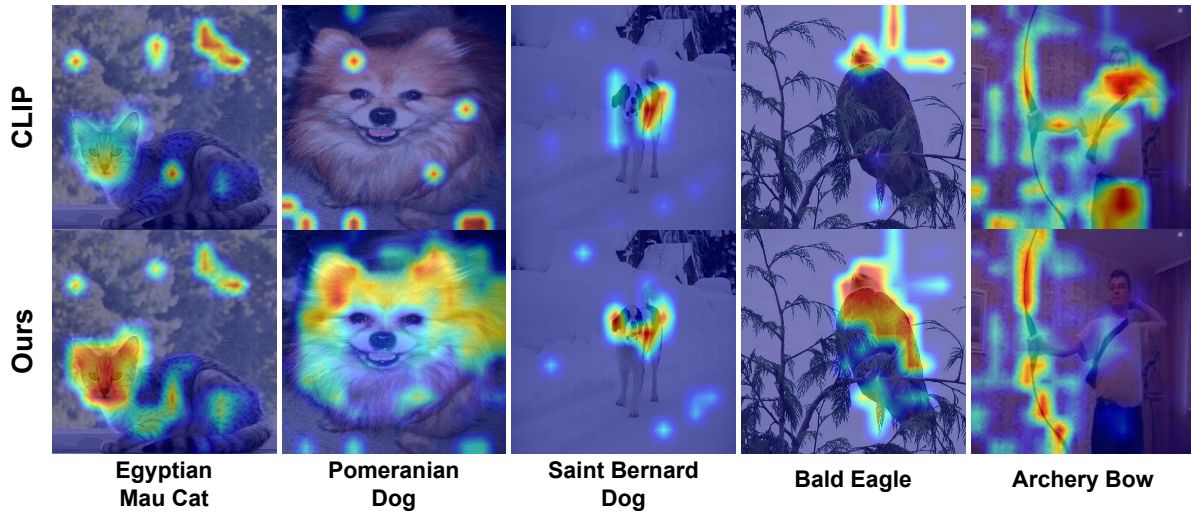


Figure S2: Saliency map comparison between the pre-trained CLIP model and the CLIP-SVD fine-tuned model. The adapted model shows more focused and semantically aligned attention, highlighting improved localization of relevant image regions.

Table S7: **Natural Domain Efficiency comparison** of different parameter-efficient tuning methods. We report trainable parameter counts, training, and inference time.

Method	Trainable Params	Training Time (min)	Inference Time (s)
CoOp	2.0K	1.84	7.34
CoCoOp	35.4K	2.25	18.52
MaPLe	3.5M	2.95	7.40
CLIP-Adapter	131.1K	4.93	7.48
Tip-Adapter	0	–	24.97
TCP	331.9K	1.02	7.58
CLIP-LoRA	184.3K	9.82	7.41
CLIP-SVD (Ours)	92.2K	0.88	7.13

Table S8: **Biomedical Domain Efficiency comparison** of different parameter-efficient tuning methods. We report trainable parameter counts, training, and inference time.

Method	Trainable Params	Training Time (min)	Inference Time (s)
CoOp	2.0K	1.19	7.20
CoCoOp	44.8K	0.58	23.6
MaPLe	5.3M	2.4	19.8
CLIP-Adapter	131.1K	2.85	7.79
Tip-Adapter	0	–	23.52
DCPL	5.5M	2.58	177.6
BiomedCoOp	3.1K	1.76	7.20
CLIP-LoRA	221.2K	10.25	7.58
CLIP-SVD (Ours)	110.6K	1.78	6.76

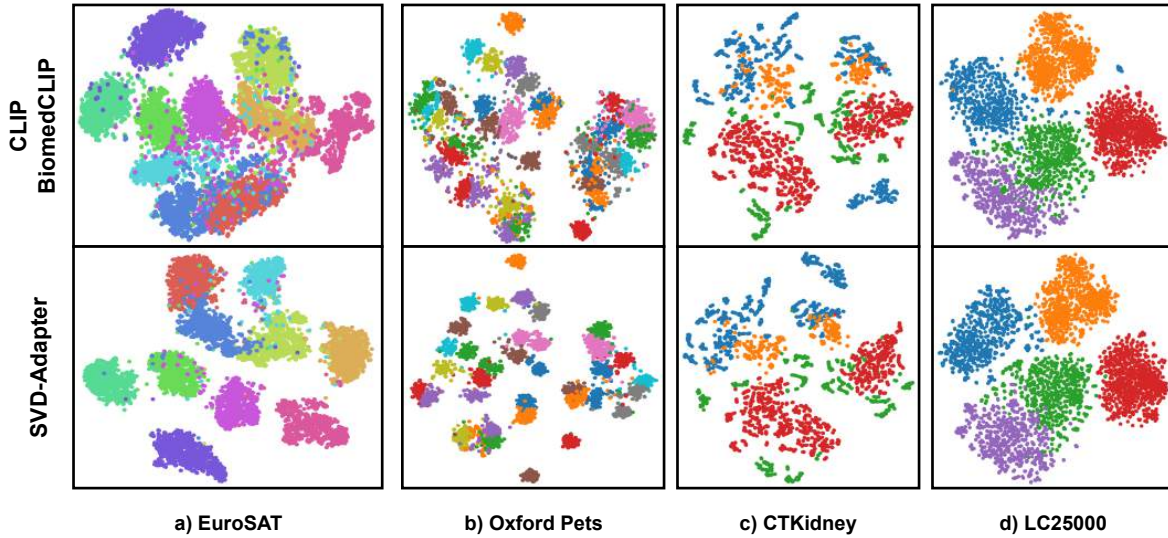


Figure S3: **t-SNE visualization of image embeddings:** Comparison of embeddings produced by the pre-trained CLIP/BiomedCLIP model and the fine-tuned CLIP-SVD on (a) EuroSAT, (b) Oxford Pets, (c) CTKidney, and (d) LC25000. CLIP-SVD generates more compact and well-separated clusters, indicating improved feature extraction and task-specific alignment.

et al., 2014), a curated subset of 4,276 ImageNet validation images with pixel-level annotations, and compared the results against those of the original pre-trained CLIP model.

To perform segmentation, we follow the approach introduced in the TextSpan framework (Gandelsman et al., 2023), which builds on a fine-grained decomposition of CLIP’s image representation. Specifically, we use the fact that the CLIP image encoder’s output can be expressed as a sum over attention head contributions across spatial positions. Each image patch contributes a vector in the shared image-text embedding space. Given a text prompt corresponding to the object class (i.e. "an image of a [CLASS]"), we compute a heatmap by measuring the similarity between each patch’s contribution and the CLIP embedding of the text description. These similarity scores are then aggregated into a spatial map that highlights the regions most semantically aligned with the prompt. We also used gScoreCAM (Chen et al., 2022a)

Model	Pixel Acc.	mIoU	mAP
CLIP	77.24	57.73	82.62
CLIP-SVD (Ours)	77.55	58.27	82.82
Δ	+0.31	+0.54	+0.20

Table S9: Segmentation Performance in terms of pixel accuracy, mean IoU, and mean average precision (%).

to visualize saliency maps for both the pre-trained CLIP model and the version fine-tuned with CLIP-SVD. Table S9 shows the results in terms of pixel accuracy, mean intersection over union (mIoU), and mean average precision (mAP). Our method achieves a performance boost across all metrics, with a gain of +0.31% in pixel accuracy, +0.54% in mIoU, and +0.20% in mAP. These improvements demonstrate that CLIP-SVD enhances the model’s ability to localize and segment fine-grained regions in complex natural images, leading to more accurate and consistent segmentation results. This boost in segmentation performance stems from improved feature extraction and attention alignment through singular value adaptation as indicated in Section 4.6. Fine-tuning the singular values helps the model’s attention heads capture task-relevant features, such as geographic cues in EuroSAT and color-related patterns in Oxford Pets, as shown in Figure S2. This enables better foreground-background differentiation and more precise segmentation boundaries, explaining the observed performance gains.

I Computational Cost

We present detailed results in Tables S7 and S8, comparing various PEFT methods in terms of total trainable parameters, training time, and inference time on the DTD and RETINA datasets, respectively. This analysis highlights the performance–efficiency trade-off across methods. All experiments were conducted on a single NVIDIA A100 GPU with 40GB of RAM, using a batch size of 8 for inference for consistency and fairness. The results demonstrate that CLIP-SVD strikes a favorable balance: it requires relatively few trainable parameters, converges more quickly, and introduces no inference overhead, while achieving high accuracy in few-shot settings. Moreover, CLIP-SVD avoids reliance on external modules or architectural modifications, simplifying deployment and improving efficiency, particularly in low-resource or latency-sensitive scenarios.

J Additional Per-dataset Results

We provide the complete per-dataset few-shot results for both the natural and biomedical domains to offer a detailed evaluation of our method’s performance. The natural and biomedical domain results are summarized in Tables S12 and S13, respectively, which report the classification accuracy across different few-shot settings for each dataset. On the other hand, we offer the per-dataset base-to-novel generalization results in Table S14 and S15. This detailed breakdown highlights the consistent improvement achieved by CLIP-SVD over state-of-the-art methods, demonstrating its ability to generalize effectively across diverse datasets and domains.

Table S10: Natural Domain Analysis of the attention heads of each layer with the top 3 descriptions returned by TextSpan(Gandelsman et al., 2023). Here, “**L**” denotes layer while “**H**” denotes attention head.

L8.H0: Foundational Shapes & Silhouettes	L8.H1: Motion & Directional Cues
Rural windmill silhouette A plank Intricate clock mechanism	Dynamic sports action shot Tranquil boating on a lake Lively city parade
L8.H2: Contrast & Light-Dark Regions	L8.H3: Geometric Patterns & Grids
Photograph with the artistic style of fisheye lens Psychedelic color swirls Sunlit meadow path	Artwork featuring Morse code typography Cubist still life painting Miniature diorama photography
L8.H4: Object Boundaries & Contours	L8.H5: Repeated Structures & Textures
Cinematic portrait with dramatic lighting A whirlpool Image with harlequin patterns	Plaid pattern Reflective surfaces Image of a scooter
L8.H6: Refined Textural Details of Everyday Objects	L8.H7: Positional Layout & Spatial Balance
Collage of textures Close-up of a textured bark Mesmerizing kinetic sculpture	Cozy cabin interior Minimalist white backdrop Rolling vineyard landscapes
L8.H8: Edge Detection & Detail Refinement	L8.H9: Primitive Forms & Visual Anchors
A scalene quadrilateral Minimalist architectural photography Detailed charcoal sketch	Dappled sunlight A cloverleaf Ocean sunset silhouette
L8.H10: Coarse-to-Fine Visual Transitions	L8.H11: Structural Symmetry & Alignment
Intricate wood carving Contemplative monochrome portrait Aerial view of a marketplace	Birds-eye view Detailed illustration of a futuristic brain-computer interface Tranquil garden pathway
L9.H0: Scene Composition & Perspective	L9.H1: Depth Perception & Visual Planes
Aerial view of a marketplace Glowing neon cityscape Playful siblings	A zoomed out photo Classic black and white cityscape Antique historical artifact
L9.H2: Interaction Between Objects	L9.H3: Material Surfaces & Reflection
An image of a Gymnast Emotional and heartfelt human connection Detailed illustration of a futuristic brain-computer interface	Aerial view of a teeming rainforest Inviting reading nook Dynamic and high-energy dance competition
L9.H4: Natural Landscapes & Textures	L9.H5: Architectural Layouts & Forms
Aerial view of a vineyard Enchanting forest nymph aesthetic Delicate and intricate lace patterns	Photograph showcasing laughter Minimalist architectural photography Urban labyrinth
L9.H6: Facial Features & Gaze Cues	L9.H7: Semantic Grouping of Elements
Detailed charcoal sketch Intense water sports moment Photograph with the artistic style of light trails	Urban rooftop panorama Cozy cabin interior Photo taken in the Japanese tea gardens
L9.H8: Soft Lighting & Shadow Play	L9.H9: Organic Flow & Movement
Impressionist portrait painting Intriguing and enigmatic passageway Ephemeral soap bubble pattern	Photograph with abstract geometric overlay Artwork featuring Morse code typography Abstract oil painting
L9.H10: Conceptual Boundaries & Themes	L9.H11: Temporal Sequences & Visual Narratives
Miniature diorama photography emotional candid moment Artwork featuring Morse code typography	Colorful hot air balloons Picture taken in Rwanda Joyful family picnic scene

Table S10 (continued): Natural Domain Analysis of the attention heads of each layer with the top 3 descriptions returned by TextSpan (Gandelsman et al., 2023). Here, “**L**” denotes layer while “**H**” denotes attention head.

L10.H0: Aerial Landscapes & Environments	L10.H1: Human Experiences & Objects in Action
Aerial view of an agricultural field Image taken in the Namibian desert Picture taken in the Brazilian rainforest	Hands in an embrace Dynamic and high-energy music concert Emergency Medical Technician at work
L10.H2: Cultural & Textural Scenes	L10.H3: Colorful Festivities & Patterns
Picture taken in the Spanish Flamenco festivals Ornate wood carving Image captured in the Peruvian Andes	Lively and colorful parade Tie-dye design Soft pastel tones
L10.H4: Cozy Indoors & Historical Locations	L10.H5: Action, Emotion & Faces
Photograph taken in a retro diner Picture taken in the Scottish castles Cozy living room ambiance	Dynamic Action Energetic children Playful winking facial expression
L10.H6: Organic Forms & Flow	L10.H7: Geographic & Cultural Diversity
Graceful wings in motion Curved organic designs Ethereal double exposure photography	Picture taken in Pakistan Bustling cultural market Image snapped in the Swiss chocolate factories
L10.H8: Color & Design Aesthetics	L10.H9: Sketches, Illustrations & Concepts
Photograph with a yellow color palette Photo taken in the Brazilian samba parade Colorful hot air balloons	Collage of vintage magazine clippings Detailed charcoal sketch Impressionist-style digital painting
L10.H10: Mood, Atmosphere & Highlights	L10.H11: Human-Centered Realism & Objects
Image with holographic retro synthwave aesthetics Serene lakeside cabin Whirlpool of brimstone	Urban labyrinth Image of a police car Intricate gemstone arrangement
L11.H0: Semantic Layout & Spatial Context	L11.H1: Human Relationships & Portraiture
Mysterious day scene Urban rooftop panorama A zoomed out photo	Images of people and emotional interaction Family portraits and group dynamics Visuals of children, couples, and professionals
L11.H2: Numbers, Symbols & Temporal Cues	L11.H3: Lifestyle & Tranquil Activities
Digits and symbolic representations in images Temporal cues like clocks, sunsets, motion blur Confetti, reflections, and expressive timing	Calm urban and rural lifestyle scenes Tranquil locations: diners, libraries, markets Cozy homes, beach sunsets, fairgrounds
L11.H4: Visual Symbols & Cultural Motifs	L11.H5: Rare Words & Abstract Concepts
Cultural symbols and calligraphy art Caricatures, retro TV test patterns, mystical motifs Architectural and folkloric aesthetics	Abstract words (quasar, zephyr), letter imagery Rare geographic terms, numerals, and geometry Conceptual elements and visual abstractions
L11.H6: Global Geographic Locations	L11.H7: Color Themes & Tonal Palettes
Images tied to global cities and landmarks Natural and urban landscapes across continents Famous locations: Machu Picchu, Santorini, NYC	Dominant color schemes (red, blue, gray) Pastel, sepia tones, artistic overlays Color-coded compositions and political art
L11.H8: Weather & Natural Forces	L11.H9: Everyday Objects & Refined Details
Storms, fog, lightning, and natural disasters Atmospheric effects and visual phenomena Wildlife amidst dynamic weather elements	Common items like scarves, bowls, wallets Mechanisms, furniture, modern artifacts Textures: wood grain, bubble, fabric
L11.H10: Natural Flora, Landscapes & Tranquility	L11.H11: Living Creatures & Natural Forms
Floral and plant life in serene settings Natural scenes: cherry blossoms, forests Geographic tranquility and biodiversity	Wide range of animals: dogs, elephants, fish Insects, feathers, natural surface textures Shells, fur, fruit, snowflakes

Table S11: Biomedical Domain Analysis of the attention heads of each layer with the top 3 descriptions returned by TextSpan (Gandelsman et al., 2023). Here, “L” denotes layer while “H” denotes attention head.

L8.H0: Focal Markers & Shape Cues	L8.H1: Peripheral Edges & Enhancement Rings
Target signs and air bronchograms Geographic zones of tissue involvement Microcalcifications and pseudocapsule patterns	Ring-enhancing lesions and peripheral edema Pseudocapsules and symmetric deviations Finger-in-glove signs and disruption of growth plates
L8.H2: Lobe Collapse & Serpentine Patterns	L8.H3: Radiologic Artifacts & Diffuse Shapes
Collapsed lobes and reverse halo signs Serpiginous lesions and high signal regions Growth plate disruptions and cortical thinning	Comet-tail artifacts and nodular regions Fluid levels suggesting benignity Asymmetries and contrast-enhanced zones
L8.H4: Flow Voids & Vascular Boundaries	L8.H5: Textural Shifts & Patchy Regions
An area with decreased perfusion A vascular displacement A vascular structure with sharp borders	Mosaic attenuation and necrotic centers Patchy regions with blurry margins Air bronchograms and target-like patterns
L8.H6: Contour Irregularity & Internal Spread	L8.H7: Symmetry Deviation & Dense Regions
An anatomical displacement A spiculated margin A zone of tissue infiltration	Mass effect and asymmetry in dense zones Patchy consolidations and star-shaped opacities Air bronchograms and peri-lesional edema
L8.H8: Clustered Signals & Midline Shifts	L8.H9: Scattered Highlights & Artifactual Spots
Midline shifts with hyperintense clustering Target signs and septated fluid collections Cortical thinning and pseudocapsule formations	An area with decreased perfusion in the left hemisphere A bright spot artifact in a clustered pattern A contrast-enhanced region on axial view
L8.H10: Diffuse Zones & Overlapping Shapes	L8.H11: Dense Regions & Local Signal Buildup
Donut and target signs in scattered areas Diffuse abnormalities and cross-compartment lesions Hypo- and hyperintense signals across lobes	Zones of infiltration with surrounding edema Septated collections and mass effects Contrast enhancement and vascular structures
L9.H0: Symmetry Shifts & Linear Features	L9.H1: Ring-Like Structures & Localized Spread
A deviation from expected symmetry in the left hemisphere A serpiginous lesion in the left hemisphere A well-defined border suggesting benignity	A central necrosis in a scattered distribution A crescent-shaped fluid collection in a scattered distribution A deviation from anatomical landmarks consistent with inflammation
L9.H2: Irregular Densities & Textured Borders	L9.H3: Small Patterned Anomalies
A beam-hardening artifact A collapsed lung lobe without enhancement A peripherally enhancing collection with sharp borders	A tree-in-bud pattern without enhancement A cavitary lesion without enhancement A high signal-to-noise ratio with sharp borders
L9.H4: Defined Edges & Subpleural Zones	L9.H5: Streaks, Texture, & Soft Borders
A subpleural nodule A central necrosis in a scattered distribution A lesion crossing compartments	A zone of tissue infiltration with surrounding edema A tram-track sign in the left hemisphere A striated muscle texture
L9.H6: Sharp Edges & Tissue Mismatch	L9.H7: Thickened Zones & Star-Like Forms
An anatomical displacement An irregular border with surrounding edema A streak artifact	A region of tissue thickening A star-shaped opacity in the upper lobe A contrast-enhanced region
L9.H8: Homogeneous Textures & Clustered Lines	L9.H9: Contour Disruptions & Shape Markers
A lesion with fluid level A homogeneous texture A disruption of normal anatomy	A cystic lesion suggesting benignity A reverse halo sign A triangular-shaped defect
L9.H10: Circumscribed Lesions & Flow Paths	L9.H11: Signal Artifacts & Internal Septations
A well-circumscribed lesion A cluster of microcalcifications suggesting malignancy A serpiginous lesion in the left hemisphere	A comet-tail artifact A lesion with internal septations without enhancement A motion artifact

Table S11 (continued): Biomedical Domain Analysis of the attention heads of each layer with the top 3 descriptions returned by TextSpan (Gandelsman et al., 2023). Here, “**L**” denotes layer while “**H**” denotes attention head.

L10.H0: Low-Contrast Regions & Collapse Patterns	L10.H1: Localized Irregularities & Shadowing
A collapsed lung lobe A vascular structure with sharp borders A focal area of calcification in the left hemisphere	A deviation from anatomical landmarks A lytic lesion consistent with inflammation A finger-in-glove sign in the left hemisphere
L10.H2: Contrast Rings & Interface Blur	L10.H3: Multi-Compartment Spread & Fine Edges
An air bronchogram suggesting malignancy A blurring of tissue interfaces A peripherally enhancing collection consistent with infection	A lesion crossing compartments in the left hemisphere A miliary pattern A vascular structure with sharp borders
L10.H4: Fused Regions & Curved Forms	L10.H5: Sharp Transitions & Highlighted Foci
A finger-in-glove sign in the left hemisphere A region with cortical thinning with surrounding edema A lesion with fluid level suggesting benignity	A streak artifact A targetoid lesion A cluster of microcalcifications suggesting malignancy
L10.H6: Circular Opacities & Clean Boundaries	L10.H7: Textured Rings & Septated Centers
A circular opacity A septated fluid collection A clean fluid interface	A central necrosis A lesion with calcified rim A reverse halo sign
L10.H8: Artifact-Like Patterns & Symmetry Shift	L10.H9: Smooth Outlines & Diffuse Flow
A midline shift in a clustered pattern A beam-hardening artifact A disruption of growth plate suggesting benignity	A serpiginous lesion in the left hemisphere A stenotic segment in the upper lobe A lesion with irregular internal architecture
L10.H10: Star Patterns & Linear Disruptions	L10.H11: Texture Aggregates & Uniform Zones
A star-shaped opacity in the upper lobe A shifting fluid level A motion artifact suggesting benignity	A lytic lesion on axial view A honeycomb pattern A sunburst pattern suggesting benignity
L11.H0: Signal Voids & Midline Shifts	L11.H1: Blurred Regions & Artifact Shapes
A collapsed lung lobe A low signal-to-noise ratio in the upper lobe A lesion crossing compartments	A solid-cystic component suggesting malignancy A finger-in-glove sign in a scattered distribution A double-density sign suggesting benignity
L11.H2: Peripheral Contrast & Mosaic Attenuation	L11.H3: Cross-Lobe Flow & Density Buildup
A mosaic attenuation A geographic area of involvement in the upper lobe A peripherally enhancing collection consistent with infection	A cavitary lesion without enhancement A lesion crossing compartments A shifting fluid level
L11.H4: Sharp Fluid Interfaces & Contrast Pockets	L11.H5: Ringed Zones & Pattern Complexity
A tram-track sign on axial view A clean fluid interface A central necrosis with sharp borders	A targetoid lesion A fluid-fluid level suggesting malignancy A mosaic attenuation
L11.H6: Rounded Opacities & Midline Distortions	L11.H7: Compact Signal Shifts & Grid-Like Forms
A circular opacity A solid-cystic component suggesting malignancy A crescent-shaped fluid collection	A star-shaped opacity in the upper lobe A widened mediastinum A spiculated margin
L11.H8: Converging Edges & Cluster Markers	L11.H9: Scattered Intensity & Symmetry Cues
A low-contrast region in a clustered pattern A double-density sign suggesting benignity A solid-cystic component suggesting malignancy	A midline shift A disruption of normal anatomy A clean fluid interface
L11.H10: Rimmed Signals & Flow Distributions	L11.H11: Sharp Vascular Lines & Compact Zones
A peripherally enhancing collection consistent with infection A change in signal intensity suggesting malignancy A beam-hardening artifact	A vascular structure with sharp borders A cluster of microcalcifications suggesting malignancy A homogeneous texture

Table S12: **Natural per-dataset performance** comparison of with various methods in few-shot setting in terms of classification accuracy (%).

Dataset	Method	$K = 1$	$K = 2$	$K = 4$	$K = 8$	$K = 16$
ImageNet	CLIP			66.72		
	CLIP-Adapter	67.93	68.60	69.56	70.20	71.60
	Tip-Adapter	68.80	68.90	69.68	70.01	70.36
	Tip-Adapter-F	67.43	68.60	69.86	71.40	73.10
	Linear Probing	31.30	43.83	54.20	61.93	67.27
	LP++	69.29	69.63	70.80	72.10	72.95
	CoOp	65.77	68.17	69.38	70.83	71.51
	CoCoOp	69.51	69.84	70.55	70.77	71.02
	KgCoOp	69.03	69.63	70.19	70.23	71.2
	ProGrad	64.33	66.12	70.21	71.10	72.68
	MaPLe	62.67	65.10	67.70	70.30	72.33
	CLIP-SVD (Ours)	70.03	70.68	71.46	72.36	73.25
Caltech101	CLIP			93.31		
	CLIP-Adapter	93.30	93.67	94.00	94.27	94.57
	Tip-Adapter	93.08	93.43	94.19	94.21	94.50
	Tip-Adapter-F	93.56	94.22	95.06	95.20	95.79
	Linear Probing	80.08	87.00	92.27	93.85	95.35
	LP++	92.85	94.14	94.87	95.44	95.94
	CoOp	92.37	92.83	94.44	94.10	95.50
	CoCoOp	93.80	94.92	94.98	95.11	95.19
	KgCoOp	94.13	94.20	94.65	94.97	95.03
	ProGrad	90.96	93.21	94.93	94.92	95.80
	MaPLe	92.57	93.97	94.43	95.20	96.00
	CLIP-SVD (Ours)	93.91	94.44	95.12	95.85	96.17
DTD	CLIP			44.09		
	CLIP-Adapter	45.20	47.87	53.83	66.40	71.17
	Tip-Adapter	50.24	52.44	57.64	61.92	65.68
	Tip-Adapter-F	53.25	56.32	62.09	67.28	72.05
	Linear Probing	36.03	46.02	56.01	63.71	69.82
	LP++	52.34	56.05	62.17	67.30	71.41
	CoOp	49.00	51.70	58.57	64.70	68.63
	CoCoOp	48.51	52.02	54.79	58.92	63.78
	KgCoOp	52.50	55.73	58.31	65.87	69.37
	ProGrad	52.79	54.35	57.72	62.13	65.92
	MaPLe	52.13	55.50	61.00	66.50	71.33
	CLIP-SVD (Ours)	56.05	60.60	64.83	68.28	72.42
EuroSAT	CLIP			48.37		
	CLIP-Adapter	61.70	66.07	66.80	73.23	81.87
	Tip-Adapter	62.46	64.65	70.39	71.57	78.44
	Tip-Adapter-F	63.95	70.38	76.43	81.11	87.46
	Linear Probing	56.36	59.56	71.90	77.93	85.45
	LP++	65.02	67.99	74.00	76.32	84.63
	CoOp	54.80	61.20	68.62	75.53	83.60
	CoCoOp	55.71	46.24	63.83	68.26	73.82
	KgCoOp	60.83	68.97	71.06	72.37	74.93
	ProGrad	55.10	66.19	70.84	79.22	84.38
	MaPLe	71.80	78.30	84.50	87.73	92.33
	CLIP-SVD (Ours)	75.06	83.30	84.65	89.74	92.68

Table S12 (continued): **Natural per-dataset performance** comparison of with various methods in few-shot setting in terms of classification accuracy (%).

Dataset	Method	$K = 1$	$K = 2$	$K = 4$	$K = 8$	$K = 16$
StanfordCars	CLIP			65.63		
	CLIP-Adapter	67.10	68.97	71.13	76.47	80.90
	Tip-Adapter	66.47	68.03	70.08	71.61	74.08
	Tip-Adapter-F	67.88	71.39	74.59	78.34	83.09
	Linear Probing	33.70	48.93	62.34	71.21	80.10
	LP++	66.31	70.41	74.57	76.12	80.77
	CoOp	67.10	69.37	72.73	76.57	78.89
	CoCoOp	67.92	68.77	69.10	70.19	71.68
	KgCoOp	67.03	68.13	71.98	73.53	74.87
	ProGrad	67.11	71.94	71.75	78.78	81.46
	MaPLe	66.60	71.60	75.30	79.47	83.57
	CLIP-SVD (Ours)	70.56	73.15	77.25	81.10	84.98
Flowers102	CLIP			70.81		
	CLIP-Adapter	54.83	54.83	54.83	56.08	56.50
	Tip-Adapter	80.74	86.59	89.62	92.15	93.68
	Tip-Adapter-F	86.63	90.21	91.50	94.98	96.18
	Linear Probing	72.50	85.48	91.65	95.89	97.36
	LP++	86.22	90.38	92.96	95.25	96.44
	CoOp	71.98	76.12	82.56	84.17	87.64
	CoCoOp	82.20	88.47	91.14	94.37	96.10
	KgCoOp	74.63	79.47	90.69	89.53	92.90
	ProGrad	83.81	88.62	89.98	93.51	94.87
	MaPLe	83.30	88.93	92.67	95.80	97.00
	CLIP-SVD (Ours)	83.25	90.32	93.44	95.49	97.50
FGVCAircraft	CLIP			24.81		
	CLIP-Adapter	27.60	29.60	31.10	37.20	42.67
	Tip-Adapter	27.22	28.08	30.19	34.12	37.64
	Tip-Adapter-F	29.58	32.51	35.15	40.61	45.59
	Linear Probing	20.13	23.78	29.46	37.18	42.85
	LP++	29.11	32.32	34.31	38.83	41.46
	CoOp	15.03	26.53	26.73	33.18	40.93
	CoCoOp	13.20	30.87	31.29	29.57	37.63
	KgCoOp	26.90	28.07	32.47	34.97	36.27
	ProGrad	27.97	30.84	32.93	37.89	40.39
	MaPLe	26.73	30.90	34.87	42.00	48.40
	CLIP-SVD (Ours)	31.11	34.85	39.42	45.91	52.49
SUN397	CLIP			62.60		
	CLIP-Adapter	67.10	69.00	71.30	73.13	75.30
	Tip-Adapter	65.37	66.46	68.39	70.12	71.47
	Tip-Adapter-F	64.49	66.74	70.29	73.60	74.99
	Linear Probing	39.83	52.70	63.07	69.54	73.29
	LP++	67.56	70.09	73.22	75.14	76.09
	CoOp	64.70	66.40	70.13	72.50	74.50
	CoCoOp	68.19	69.11	70.50	70.62	72.05
	KgCoOp	68.43	69.53	71.79	72.50	73.40
	ProGrad	64.54	68.51	71.17	72.91	75.00
	MaPLe	64.77	67.10	70.67	73.23	75.53
	CLIP-SVD (Ours)	70.37	71.93	73.75	75.16	76.85

Table S12 (continued): **Natural per-dataset performance** comparison of with various methods in few-shot setting in terms of classification accuracy (%).

Dataset	Method	$K = 1$	$K = 2$	$K = 4$	$K = 8$	$K = 16$
OxfordPets	CLIP			89.10		
	CLIP-Adapter	89.03	89.73	90.87	91.77	92.03
	Tip-Adapter	89.32	88.52	89.36	90.69	91.32
	Tip-Adapter-F	90.72	91.10	91.58	92.09	92.70
	Linear Probing	40.70	55.03	69.96	79.99	85.80
	LP++	89.52	89.94	91.09	91.91	92.80
	CoOp	92.20	89.20	91.30	90.83	91.80
	CoCoOp	91.17	92.14	93.01	93.44	93.25
	KgCoOp	91.97	92.13	93.20	93.10	93.23
	ProGrad	89.01	90.55	93.21	92.18	92.13
	MaPLe	89.10	90.87	91.90	92.57	92.83
	CLIP-SVD (Ours)	92.67	92.09	92.83	92.57	93.77
UCF101	CLIP			67.64		
	CLIP-Adapter	69.67	74.10	77.30	81.87	84.53
	Tip-Adapter	68.92	71.50	74.07	75.65	77.35
	Tip-Adapter-F	73.27	76.11	80.24	82.24	84.50
	Linear Probing	48.24	61.08	70.87	77.51	81.27
	LP++	72.36	75.13	79.49	82.17	83.75
	CoOp	24.96	25.89	23.85	26.23	28.48
	CoCoOp	25.42	28.85	30.66	21.78	24.86
	KgCoOp	72.93	74.83	78.40	80.03	81.43
	ProGrad	71.91	74.39	77.82	88.64	81.59
	MaPLe	71.83	74.60	78.47	81.37	85.03
	CLIP-SVD (Ours)	76.34	79.69	81.80	83.95	86.58
Food101	CLIP			85.87		
	CLIP-Adapter	85.90	86.10	86.47	86.67	86.83
	Tip-Adapter	85.14	86.03	85.99	86.42	86.36
	Tip-Adapter-F	86.01	86.26	86.48	86.74	87.24
	Linear Probing	44.63	62.66	72.96	78.98	83.69
	LP++	83.23	86.12	85.94	86.76	87.28
	CoOp	82.07	80.80	82.58	83.37	85.17
	CoCoOp	86.10	86.21	86.64	86.92	87.19
	KgCoOp	86.27	86.60	86.59	86.90	87.03
	ProGrad	82.75	84.81	85.77	85.91	87.01
	MaPLe	80.50	81.47	81.77	83.60	85.33
	CLIP-SVD (Ours)	85.82	85.58	85.47	85.59	86.00
Average	CLIP			65.36		
	CLIP-Adapter	67.87	70.20	72.65	76.92	79.86
	Tip-Adapter	68.89	70.42	72.69	74.41	76.44
	Tip-Adapter-F	70.62	73.08	75.75	78.51	81.15
	Linear Probing	45.77	56.92	66.79	73.43	78.39
	LP++	70.35	72.93	75.77	77.94	80.32
	CoOp	68.09	70.13	73.59	76.45	79.01
	CoCoOp	66.95	67.63	71.98	72.92	75.02
	KgCoOp	69.51	71.57	74.48	75.82	77.26
	ProGrad	68.20	71.78	74.21	77.93	79.20
	MaPLe	69.27	72.58	75.37	78.89	81.79
	CLIP-SVD (Ours)	73.20	76.06	78.18	80.55	82.97

Table S13: **Biomedical per-dataset performance** comparison of CLIP-SVD with various methods in few-shot setting in terms of classification accuracy (%).

Dataset	Method	$K = 1$	$K = 2$	$K = 4$	$K = 8$	$K = 16$
BTMRI	BiomedCLIP			56.79		
	CLIP-Adapter	56.80	57.13	56.80	57.15	60.16
	Tip-Adapter	66.66	67.77	76.37	73.75	78.97
	Tip-Adapter-F	59.60	61.94	77.90	79.18	82.27
	Standard LP	62.24	72.45	75.98	77.63	81.24
	LP++	64.72	71.69	75.48	77.11	81.61
	CoOp	63.82	68.82	74.68	79.27	82.37
	CoCoOp	59.47	64.14	67.83	71.69	78.45
	KgCoOp	63.33	70.16	75.40	79.79	81.07
	ProGrad	66.92	71.46	76.24	78.82	82.84
	MaPLe	38.01	37.02	42.36	50.75	56.22
	BiomedCoOp	65.08	70.57	77.23	78.55	83.30
	CLIP-SVD (Ours)	63.25	73.62	78.43	83.21	84.64
BUSI	BiomedCLIP			59.75		
	CLIP-Adapter	61.44	61.01	61.72	61.86	63.55
	Tip-Adapter	62.71	61.44	59.03	55.93	68.78
	Tip-Adapter-F	61.86	56.35	64.54	68.50	71.89
	Standard LP	51.41	47.88	53.38	65.53	68.78
	LP++	51.12	55.50	60.31	66.10	70.05
	CoOp	48.73	53.53	60.17	64.69	69.49
	CoCoOp	52.26	49.15	59.75	65.82	70.2
	KgCoOp	53.39	55.51	62.01	67.37	70.62
	ProGrad	46.33	49.15	62.29	64.83	71.47
	MaPLe	41.38	33.47	47.74	42.65	45.62
	BiomedCoOp	50.71	50.71	59.32	63.27	70.34
	CLIP-SVD (Ours)	61.58	63.56	68.12	73.87	74.29
COVID-QU-Ex	BiomedCLIP			43.8		
	CLIP-Adapter	50.42	43.04	46.28	48.68	49.55
	Tip-Adapter	62.13	58.72	63.84	66.77	73.05
	Tip-Adapter-F	54.89	54.01	69.97	69.89	76.07
	Standard LP	49.91	48.06	60.55	68.29	71.98
	LP++	46.41	56.42	62.32	66.19	72.79
	CoOp	58.82	58.37	67.03	74.66	76.37
	CoCoOp	69.36	68.80	63.70	69.36	74.52
	KgCoOp	61.68	54.68	65.91	74.86	75.65
	ProGrad	60.42	64.22	68.56	74.65	74.93
	MaPLe	35.86	38.99	33.32	36.43	40.89
	BiomedCoOp	72.64	71.53	73.28	76.26	78.72
	CLIP-SVD (Ours)	71.15	71.08	70.47	73.97	73.14
CTKIDNEY	BiomedCLIP			42.43		
	CLIP-Adapter	47.17	41.94	42.19	44.64	47.28
	Tip-Adapter	45.85	51.65	55.33	69.89	73.38
	Tip-Adapter-F	46.68	58.99	60.18	75.24	82.07
	Standard LP	43.82	59.35	69.54	78.89	82.50
	LP++	57.70	61.57	65.73	77.06	79.07
	CoOp	54.51	60.57	68.12	77.40	83.52
	CoCoOp	47.88	52.71	61.07	73.93	77.70
	KgCoOp	58.92	62.81	68.68	77.43	77.67
	ProGrad	54.65	64.66	67.90	78.23	81.13
	MaPLe	30.62	38.98	38.00	39.67	51.06
	BiomedCoOp	56.13	64.21	66.50	77.16	83.20
	CLIP-SVD (Ours)	55.64	58.88	74.31	78.33	86.34

Table S13 (continued): **Biomedical per-dataset performance** comparison of CLIP-SVD with various methods in few-shot setting in terms of classification accuracy (%).

Dataset	Method	$K = 1$	$K = 2$	$K = 4$	$K = 8$	$K = 16$
Kvasir	BiomedCLIP			54.58		
	CLIP-Adapter	54.83	54.83	54.83	56.08	56.50
	Tip-Adapter	56.72	60.94	69.61	69.13	74.22
	Tip-Adapter-F	59.19	64.22	69.94	75.86	78.00
	Standard LP	54.30	62.00	72.38	78.88	79.00
	LP++	58.27	60.47	69.36	72.52	75.41
	CoOp	58.2	64.86	70.78	77.14	77.88
	CoCoOp	59.45	65.50	68.94	72.92	75.22
	KgCoOp	61.67	65.67	68.28	72.05	72.95
	ProGrad	60.78	64.70	70.00	76.03	75.88
	MaPLe	41.06	45.17	53.22	56.03	63.50
	BiomedCoOp	62.17	67.25	74.08	77.72	78.89
	CLIP-SVD (Ours)	65.08	72.08	74.92	80.08	82.83
CHMNIST	BiomedCLIP			30.65		
	CLIP-Adapter	31.27	31.67	33.26	36.48	42.06
	Tip-Adapter	46.14	63.32	70.05	69.57	77.68
	Tip-Adapter-F	52.81	58.90	71.74	74.51	80.43
	Standard LP	58.44	64.42	71.07	76.30	80.34
	LP++	57.18	60.61	67.79	72.40	78.32
	CoOp	57.34	59.68	68.66	75.00	79.63
	CoCoOp	49.07	50.82	58.58	66.58	72.16
	KgCoOp	59.02	60.06	68.77	69.50	73.58
	ProGrad	60.15	59.60	69.13	70.99	75.11
	MaPLe	48.05	57.76	65.87	68.88	71.99
	BiomedCoOp	59.82	59.79	71.19	74.78	79.05
	CLIP-SVD (Ours)	54.94	66.13	75.69	81.05	84.75
LC25000	BiomedCLIP			50.03		
	CLIP-Adapter	54.83	53.47	52.91	56.33	57.56
	Tip-Adapter	75.37	72.73	83.32	87.25	89.17
	Tip-Adapter-F	74.21	71.82	79.57	90.41	92.35
	Standard LP	74.50	78.40	85.30	90.24	92.77
	LP++	63.05	71.42	82.61	89.14	92.58
	CoOp	71.90	76.55	84.66	87.50	92.19
	CoCoOp	63.66	71.76	77.44	85.57	87.38
	KgCoOp	71.80	75.18	82.10	84.63	86.79
	ProGrad	72.48	74.76	84.72	87.86	90.70
	MaPLe	67.13	69.80	76.73	82.19	86.73
	BiomedCoOp	77.56	77.74	85.60	88.77	92.68
	CLIP-SVD (Ours)	72.34	82.31	89.01	89.98	91.52
RETINA	BiomedCLIP			26.26		
	CLIP-Adapter	25.49	25.49	26.07	25.84	26.05
	Tip-Adapter	26.52	31.07	43.42	48.08	54.23
	Tip-Adapter-F	39.53	33.07	47.37	56.07	62.85
	Standard LP	39.35	46.03	51.31	53.94	62.27
	LP++	35.77	39.37	46.95	53.44	60.62
	CoOp	35.02	35.26	42.22	51.87	59.38
	CoCoOp	32.94	36.43	39.75	48.45	53.91
	KgCoOp	33.54	35.17	42.61	49.97	51.18
	ProGrad	33.49	36.49	43.09	52.26	50.47
	MaPLe	27.73	31.07	30.89	41.80	53.78
	BiomedCoOp	36.64	38.67	45.58	56.47	61.28
	CLIP-SVD (Ours)	41.46	44.93	52.63	62.33	68.06

Table S13 (continued): **Biomedical per-dataset performance** comparison of CLIP-SVD with various methods in few-shot setting in terms of classification accuracy (%).

Dataset	Method	$K = 1$	$K = 2$	$K = 4$	$K = 8$	$K = 16$
KneeXray	BiomedCLIP			29.53		
	CLIP-Adapter	29.00	28.66	28.96	28.80	29.08
	Tip-Adapter	29.04	33.55	24.19	25.76	33.17
	Tip-Adapter-F	30.01	28.38	26.59	26.46	27.67
	Standard LP	26.02	26.57	27.83	22.20	23.97
	LP++	21.25	26.40	28.92	23.75	26.38
	CoOp	24.96	25.89	23.85	26.23	28.48
	CoCoOp	25.42	28.85	30.66	21.78	24.86
	KgCoOp	29.07	28.14	22.44	23.37	24.80
	ProGrad	30.09	23.83	23.95	24.78	26.27
	MaPLe	23.41	22.67	22.56	24.78	25.87
	BiomedCoOp	36.13	37.72	35.91	37.7	39.69
	CLIP-SVD (Ours)	27.38	27.05	27.67	30.70	38.37
OCTMNIST	BiomedCLIP			30.00		
	CLIP-Adapter	44.00	49.73	49.96	49.50	52.73
	Tip-Adapter	32.36	33.8	38.10	53.93	53.33
	Tip-Adapter-F	46.66	53.93	55.20	65.00	72.50
	Standard LP	47.25	54.21	61.00	65.85	69.40
	LP++	47.24	53.18	59.02	63.69	68.35
	CoOp	52.63	53.57	53.37	63.67	65.47
	CoCoOp	49.33	50.93	48.57	55.40	60.67
	KgCoOp	50.63	50.53	52.97	61.03	62.80
	ProGrad	51.40	55.33	55.07	62.17	63.33
	MaPLe	26.63	34.00	30.17	30.53	33.63
	BiomedCoOp	51.83	55.03	54.73	58.87	66.93
	CLIP-SVD (Ours)	50.63	66.67	68.97	79.07	80.67
Average	BiomedCLIP			42.38		
	CLIP-Adapter	45.53	44.70	45.30	46.54	48.46
	Tip-Adapter	50.35	53.50	58.33	62.01	67.60
	Tip-Adapter-F	52.55	54.17	62.30	68.12	68.12
	Standard LP	48.91	55.82	62.12	67.33	70.81
	LP++	49.27	55.88	61.30	65.48	70.09
	CoOp	52.59	55.71	61.35	67.74	71.48
	CoCoOp	50.88	53.91	57.63	63.15	67.51
	KgCoOp	54.31	55.79	60.92	66.00	67.71
	ProGrad	53.67	56.42	62.10	67.06	69.21
	MaPLe	37.99	40.89	44.09	47.37	52.93
	BiomedCoOp	56.87	59.32	64.34	68.96	73.41
	CLIP-SVD (Ours)	56.35	62.63	68.02	73.26	76.46

Table S14: **Natural Base-to-novel generalization** comparison of CLIP-SVD with previous methods

Dataset		CLIP (Radford et al., 2021)	CoOp (Zhou et al., 2022b)	CoCoOp (Zhou et al., 2022a)	KgCoOp (Yao et al., 2023a)	ProGrad (Zhu et al., 2024a)	MaPLe (Khattak et al., 2023a)	CLIP-SVD (Ours)
Average on 11 datasets	Base	69.34	82.69	80.47	80.73	82.48	82.28	84.38
	Novel	74.22	63.22	71.69	73.60	70.75	75.14	76.29
	HM	71.70	71.66	75.83	77.00	76.16	78.55	80.13
ImageNet	Base	72.43	76.47	75.98	75.83	77.02	76.66	78.01
	Novel	68.14	67.88	70.43	69.96	66.66	70.54	70.71
	HM	70.22	71.92	73.10	72.78	71.46	73.47	74.18
Caltech101	Base	96.84	98.00	97.96	97.72	98.02	97.74	98.54
	Novel	94.00	89.81	93.81	94.39	93.89	94.36	94.00
	HM	95.40	93.73	95.84	96.03	95.91	96.02	96.21
OxfordPets	Base	91.17	93.67	95.20	94.65	95.07	95.43	96.28
	Novel	97.26	95.29	97.69	97.77	97.63	97.76	97.60
	HM	94.12	94.47	96.43	96.18	96.33	96.58	96.93
Stanford Cars	Base	63.37	78.12	70.49	71.76	77.68	72.94	78.89
	Novel	74.89	60.40	73.59	75.04	68.63	74.00	74.03
	HM	68.65	68.13	72.01	73.36	72.88	73.47	76.38
Flowers102	Base	72.08	97.60	94.87	95.00	95.54	95.92	96.30
	Novel	77.80	59.67	71.75	74.73	71.87	72.46	76.19
	HM	74.83	74.06	81.71	83.65	82.03	82.56	85.07
Food101	Base	90.10	88.33	90.70	90.50	90.37	90.71	90.52
	Novel	91.22	82.26	91.29	91.70	89.59	92.05	91.74
	HM	90.66	85.19	90.99	91.09	89.98	91.38	91.13
FGVC Aircraft	Base	27.19	40.44	33.41	36.21	40.54	37.44	44.00
	Novel	36.29	22.30	23.71	33.55	27.57	35.61	37.01
	HM	31.09	28.75	27.74	34.83	32.82	36.50	40.20
SUN397	Base	69.36	80.60	79.74	80.29	81.26	80.82	82.56
	Novel	75.35	65.89	76.86	76.53	74.17	78.70	78.98
	HM	72.23	72.51	78.27	78.36	77.55	79.75	80.73
DTD	Base	53.24	79.44	77.01	77.55	77.35	80.36	82.91
	Novel	59.90	41.18	56.00	54.99	52.35	59.18	65.42
	HM	56.37	54.24	64.85	64.35	62.45	68.16	73.13
EuroSAT	Base	56.48	92.19	87.49	85.64	90.11	94.07	94.53
	Novel	64.05	54.74	60.04	64.34	60.89	73.23	74.03
	HM	60.03	68.69	71.21	73.48	72.67	82.35	83.03
UCF101	Base	70.53	84.69	82.33	82.89	84.33	83.00	85.66
	Novel	77.50	56.05	73.45	76.67	74.94	78.66	79.45
	HM	73.85	67.46	77.64	79.65	79.35	80.77	82.44

Table S15: **Biomedical Base-to-novel generalization** comparison of CLIP-SVD with previous methods

Dataset		CLIP (Radford et al., 2021)	CoOp (Zhou et al., 2022b)	CoCoOp (Zhou et al., 2022a)	KgCoOp (Yao et al., 2023a)	ProGrad (Zhu et al., 2024a)	MaPLe (Khattak et al., 2023a)	BiomedCoOp (Koleilat et al., 2025b)	CLIP-SVD (Ours)
Average on 9 datasets	Base	49.27	76.71	75.52	71.90	75.69	65.40	78.60	82.64
	Novel	67.17	65.34	67.74	65.94	67.33	49.51	73.90	74.31
	HM	55.23	68.80	69.11	67.22	69.86	53.10	74.04	78.25
BTMRI	Base	40.88	82.25	77.88	78.00	82.13	66.17	82.42	88.83
	Novel	96.18	94.51	94.84	95.05	94.98	49.58	96.84	94.98
	HM	57.37	87.95	85.53	85.69	88.09	56.69	89.05	91.80
COVID-QU-Ex	Base	53.96	75.92	77.28	75.42	75.19	61.36	75.91	75.39
	Novel	89.43	90.07	87.61	89.61	90.34	71.25	91.63	89.56
	HM	67.31	82.39	82.12	81.90	82.07	65.94	83.03	81.87
CTKIDNEY	Base	38.55	82.24	81.96	81.67	83.86	63.99	86.93	88.31
	Novel	52.99	67.92	56.56	58.45	63.01	63.51	78.94	68.02
	HM	44.63	74.40	66.93	68.14	71.96	63.75	82.74	76.85
Kvasir	Base	75.00	86.22	85.94	81.56	82.89	76.66	86.50	86.78
	Novel	60.50	58.06	53.95	59.00	60.45	26.17	61.83	61.89
	HM	66.97	69.39	66.29	68.47	69.91	39.02	72.11	72.25
CHMNIST	Base	37.63	89.41	87.77	75.45	82.98	89.19	88.87	90.38
	Novel	40.69	35.11	42.51	38.70	44.19	23.76	42.73	35.42
	HM	39.10	50.42	57.28	51.16	57.67	37.52	57.71	50.89
LC25000	Base	59.73	90.12	88.33	88.13	90.29	87.16	93.77	96.68
	Novel	87.60	87.55	95.02	86.44	85.47	52.66	97.00	96.36
	HM	71.03	88.82	91.55	87.28	87.81	65.65	95.36	96.52
RETINA	Base	45.18	70.98	66.88	60.77	68.77	57.40	68.46	84.62
	Novel	55.28	56.90	65.56	54.91	58.43	53.33	67.72	84.88
	HM	49.72	63.16	66.21	57.69	63.18	55.29	68.09	84.75
KneeXray	Base	35.89	38.28	34.08	37.94	40.88	36.44	44.23	43.73
	Novel	71.90	47.69	63.14	61.19	59.12	55.35	78.35	76.52
	HM	47.88	42.47	44.27	46.84	48.34	43.95	56.54	55.65
OCTMNIST	Base	56.60	75.00	79.60	68.20	74.20	50.27	80.33	89.00
	Novel	50.00	50.23	50.47	50.13	50.02	50.00	50.07	61.13
	HM	53.10	60.17	61.77	57.79	59.76	50.13	61.69	72.48