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# ProtQueSt: Query-Conditioned Retrieval-Augmented Generation for Protein Function Annotation

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## Abstract

Protein function prediction from sequence is inherently query-dependent: the same protein may be characterized by its catalytic activity, domain architecture, or cellular localization depending on the biological question. Prior work has shown that large language models consistently underperform retrieval-based methods on this task, yet simple retrieval transfers annotations from embedding-space neighbors without adapting to the query. We introduce **ProtQueSt**, a retrieval-augmented framework that pairs a structure-aware retriever with a query-conditioned contrastive retriever that aligns protein and annotation representations via Feature-wise Linear Modulation (FiLM). FiLM conditioning and query-pooled negative sampling prove jointly essential, as neither alone improves over the structural baseline. ProtQueSt achieves state-of-the-art Entity-BLEU (48.79, +37% over RAPM) and LLM-as-a-judge score on Prot-Inst-OOD. This result supports reframing text-based protein understanding as a query-conditioned retrieval problem rather than a single fixed sequence-to-text mapping.

## 1. Introduction

Accurate protein functional annotation is fundamental to biomedicine and drug discovery. Recent protein-language models such as Evolla (Zhou et al., 2025), ProteinGPT (Xiao et al., 2024), and BioReason-Pro (Fallahpour et al., 2026) have demonstrated strong capabilities, but they typically require predicted 3D structures, interaction networks, or Gene Ontology terms beyond the raw sequence. For novel or orphan proteins, amino acid sequence is often the only reliable input, making sequence-based prediction a practically critical setting. Meanwhile, LLMs consistently underperform

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retrieval-based methods on sequence-only benchmarks (Wu et al., 2025), a finding our own results confirm (Table 1).

While this sequence-only constraint poses a challenge for current models, it is theoretically sufficient for functional annotation. As Anfinsen’s dogma (Anfinsen, 1973) states, protein functionality arises from its amino acid sequence. However, high-quality annotation relies on *text-based protein sequence understanding*: the ability to translate a raw amino acid sequence directly into the nuanced natural language concepts that define its actual function. While prior work like Prot2Text-V2 (Fei et al., 2025) advances this by reformulating function prediction as free-text generation, it assumes a single fixed description. In practice, a protein’s function is rarely context-independent; the same sequence may be queried for catalytic activity, domain architecture, or cellular localization depending on the specific context, making the ability to yield distinct descriptions essential.

Therefore, we rethink text-based protein sequence understanding as a query-driven ability: dynamically mapping sequences to biological entities conditioned on the query.

While retrieval-based methods outperform LLMs on protein function prediction, they do not achieve true sequence understanding. Systems based on sequence alignment or embedding similarity transfer annotations from similar proteins without adapting to query. To move beyond this nearest-neighbor transfer, the retriever must learn query-conditioned alignment between protein sequences and annotations. This draws on the broader idea behind conditional representation learning (Liu et al., 2025) and feature modulation (Perez et al., 2018), where the same input amplifies different features depending on the conditioning signal. However, any conditioning module can only emphasize signals already present in the base embedding.

We introduce ProtQueSt (Protein Query-conditioned Structure-aware retrieval-augmented framework) with the following contributions:

- We reapproach protein function annotation as text-based protein sequence understanding and reframe the latter as a query-driven ability, where the functional meaning of a sequence is conditioned on the specific biological question asked.

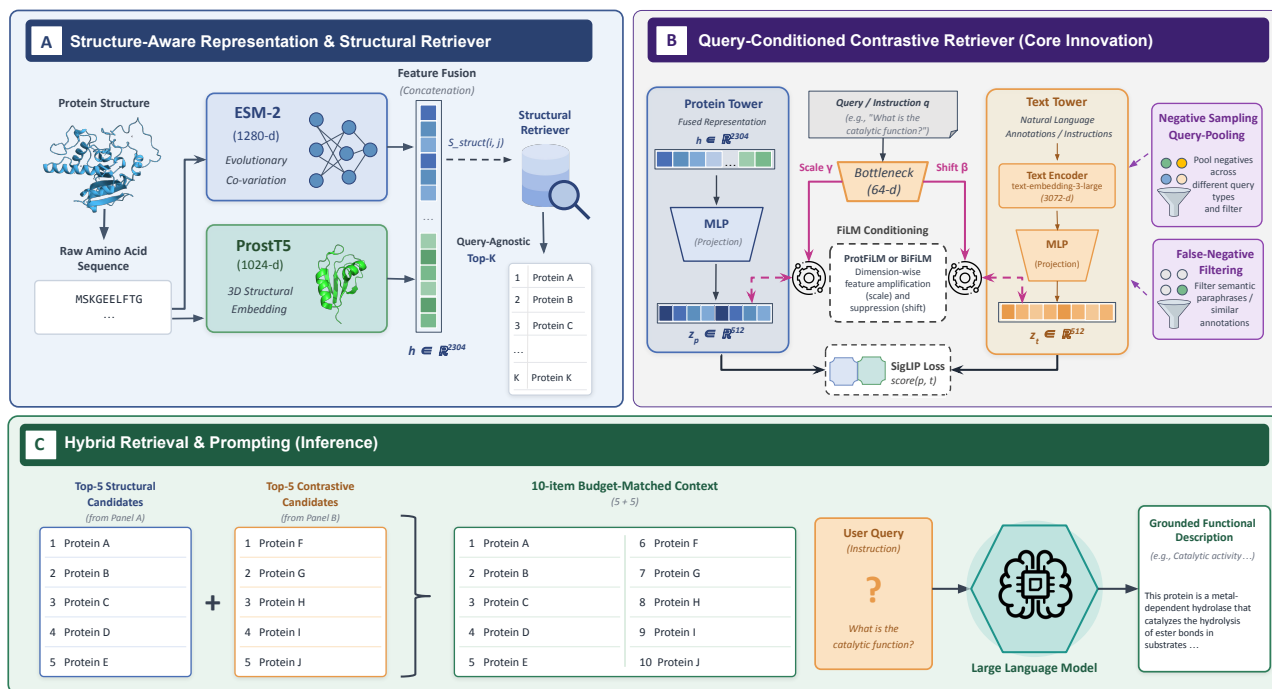


Figure 1. ProtQueSt Architecture. (A) A structure-aware retriever fuses ESM-2 and ProstT5 embeddings for query-independent top-K candidate retrieval. (B) A query-conditioned contrastive retriever uses FiLM conditioning and query-pooled negative sampling to align protein-text representations via SigLIP loss. (C) Hybrid retrieval are combined for grounded functional description generation by an LLM.

- We realize this through a contrastive retriever that applies Feature-wise Linear Modulation (FiLM) to select query-relevant features from frozen protein embeddings, trained with query-pooled negative sampling that provides truly query-specific contrastive signal.
- We present **ProtQueSt**, a hybrid retrieval system that combines structure-aware and query-conditioned channels to achieve state-of-the-art results on the Prot-InstOOD benchmark.

## 2. Background and Related Work

**LLMs for protein function prediction.** BioT5+ (Pei et al., 2024) aligns small molecules and proteins with natural language via IUPAC nomenclature and multi-task tuning. InstructProtein (Wang et al., 2024) uses knowledge-graph-grounded instruction tuning for bidirectional sequence-to-text generation. BioReason-Pro (Fallahpour et al., 2026) adds reinforcement learning over biological knowledge.

**Retrieval-augmented protein annotation.** RAPM (Wu et al., 2025) pioneered retrieval-augmented protein function description by combining MMseqs2 and ESM-2 similarity to retrieve neighbor proteins and prepend their annotations to the LLM prompt. Evolla-10B (Zhou et al., 2025) instead internalizes protein-text alignment in a 10B-parameter end-to-end fusion model, removing explicit retrieval at a heavy parameter cost. Beyond functional annotation, PoET-

2 (Truong Jr & Bepler, 2025) and Protriever (Weitzman et al., 2025) apply retrieval augmentation to fitness and variant effect prediction by retrieving homologous sequences.

**Contrastive protein-text alignment.** ProteinCLIP (Wu et al., 2024) and ProtST (Xu et al., 2023) align protein sequences and text via contrastive objectives over (sequence, text) pairs. Prot2Text-V2 (Fei et al., 2025) extends this with a stronger encoder stack and a contrastive auxiliary objective for protein-to-text generation.

## 3. Methods

Our system retrieves protein functional annotations through two complementary channels, a *structural retriever* and a *contrastive metadata retriever*, whose outputs are merged and passed to an LLM for retrieval-augmented generation.

### 3.1. Structure-Aware Protein Representation and Structural Retriever

Protein function is largely determined by 3D structure, so a representation encoding structural signals should capture richer functional information than sequence data alone. At the same time, any downstream conditioning module can only expose signals already present in the base embedding (Section 1). To raise this representational ceiling, we fuse frozen ESM-2 (Lin et al., 2023) (1,280-d, evolutionary co-

variation) with ProstT5 (Heinzinger et al., 2023) (1,024-d, 3D fold information predicted from sequence) to yield  $\mathbf{h} \in \mathbb{R}^{2304}$ , providing both the structural retriever and the contrastive module with a structure-aware input.

The structural retriever scores candidates by a weighted cosine fusion of L2-normalized ProstT5 and ESM-2 embeddings ( $\alpha=0.7$ ; sweep in Appendix K.1), replacing RAPM’s dual-key pipeline with a unified dense score. The top- $K$  candidates are retrieved from the full training corpus, and their curated annotations are retained.

### 3.2. Query-Conditioned Contrastive Retriever

The structural retriever is query-independent by design: the same protein always yields the same neighbors regardless of whether the user asks about catalytic activity or cellular localization. The contrastive channel closes this gap by learning a query-conditioned compatibility function. On the protein side, we reuse the same fused ESM-2 and ProstT5 representation ( $\mathbf{h} \in \mathbb{R}^{2304}$ ). On the text side, following the ProteinCLIP (Wu et al., 2024) paradigm of aligning protein representations with natural-language descriptions, we use the OpenAI TEXT-EMBEDDING-3-LARGE model to generate 3072-d embeddings as the text representation for both annotation descriptions and instruction queries.

**Architecture.** We adopt a two-tower contrastive model with lightweight two-layer MLP adapters that project both protein and text embeddings to a shared 512-d space (full specification in Appendix B). The compatibility score is cosine similarity scaled by a learnable temperature  $\tau$ .

**FiLM query conditioning.** By design, the base protein adapter yields a fixed embedding  $\mathbf{z}_p$ , forcing it to serve all query types during direct contrastive training. Yet, the fused embedding encodes a rich set of functional signals across its dimensions. We apply Feature-wise Linear Modulation (FiLM) (Perez et al., 2018) as a lightweight conditioning mechanism that selectively emphasizes the dimensions relevant to a given query type.

Concretely, the instruction embedding  $\mathbf{q} \in \mathbb{R}^{3072}$  is first compressed to a 64-d bottleneck  $\tilde{\mathbf{q}} = \text{GELU}(\text{LN}(\mathbf{q})\mathbf{W}_q)$ . A 64-d bottleneck suffices because protein function queries span only a handful of distinct question types, so this compression discards noise while preserving coarse query identity. From  $\tilde{\mathbf{q}}$ , two linear maps produce a scale vector and a shift vector, each in  $\mathbb{R}^{512}$ , which modulate the adapter embedding  $\mathbf{z} \in \{\mathbf{z}_p, \mathbf{z}_t\}$  dimension-by-dimension:

$$\mathbf{z}_{\text{out}}[d] = \mathbf{z}[d] \cdot (1 + \text{scale}(\tilde{\mathbf{q}})[d]) + \text{shift}(\tilde{\mathbf{q}})[d]. \quad (1)$$

Initialization and training details are in Appendix C.

We explore two variants: **ProtFiLM**, which conditions only the protein encoder, and **BiFiLM**, which conditions both encoders (the query additionally modulates which text fea-

tures are matched). ProtFiLM adds  $\sim 6.8\text{M}$  parameters and BiFiLM  $\sim 7.1\text{M}$ ; all base encoder weights remain frozen.

**Negative sources and query pooling.** Training negatives include both in-batch annotations and annotations from the protein’s structural neighbors (details in Appendix C). In a multi-task batch, negatives from different query types are trivially distinguishable. Query pooling addresses this: for each anchor with instruction  $\mathbf{q}_i$ , we mask out any candidate whose instruction satisfies  $\cos(\mathbf{q}_i, \mathbf{q}_j) < \tau_{\text{query}}$ , ensuring only same-type comparisons provide gradient.

**False-negative filtering.** In-batch annotations may be paraphrases of the anchor’s gold; we mask candidates whose annotation embedding exceeds a cosine threshold  $\tau_{\text{fn}}$  against the gold (details in Appendix C).

**Training objective.** Because query pooling and false-negative filtering mask different subsets per anchor, the effective negative set varies across proteins in the batch. This is incompatible with InfoNCE, which requires a shared softmax denominator. We therefore optimize a pairwise sigmoid loss (SigLIP) (Zhai et al., 2023):

$$\mathcal{L}_{ij} = -\log \sigma(y_{ij}(\text{score}(p_i, t_j) + b)) \quad (2)$$

Details and a Soft-SigLIP variant are in Appendix C.

### 3.3. Hybrid Retrieval and Prompting

At inference, we retrieve the top-5 candidates from each channel (structural and contrastive) and concatenate them into a 10-item context, budget-matched to the structural-only baseline for fair comparison. The contrastive channel applies the same query mask used during training ( $\tau_{\text{query}}=0.65$ ) to filter the annotation pool. Both source lists are presented with source labels in a structured prompt to Claude Sonnet 4.6, which generates the final functional description (prompt template in Figure 2; generation configuration in Appendix D).

## 4. Experiments and Results

### 4.1. Experimental Setup

**Dataset.** We evaluate all models on Prot-Inst-OOD (Wu et al., 2025), the out-of-distribution evaluation split of a UniProt-derived protein-instruction corpus (Appendix A).

**Metrics.** Our primary lexical metric is Entity-BLEU (Wu et al., 2025), which restricts BLEU to tokens present in the metadata so the score reflects biologically-relevant content. We also report ROUGE-L (Lin, 2004) and METEOR (Banerjee & Lavie, 2005), and an LLM-as-a-judge metric (full rubric in Appendix E). We also report Recall@10 for retrieval (Appendix F). ROUGE-L is reported in the appendix.

**Baselines.** We compare against 4 families in the protein-text

Table 1. Evaluation for ProtQueSt and baselines on Prot-Inst-  
 OOD. Lexical metrics: Entity-BLEU (E-BL) and METEOR. LLM-  
 based metric: LLM-as-judge Final score (Judge-F, 0–10 scale).  
 ProtQueSt uses ProtFiLM with SigLIP loss,  $\tau_{\text{query}}=0.65$ ,  $\tau_{\text{fn}}=0.70$ ,  
 and hybrid 5+5 merging (Section 4.3; Appendix I). Only-Structure  
 uses 10 structural candidates. Both use Claude Sonnet 4.6; full-  
 dataset results in Appendix H.

Method	E-BL	METEOR	Judge-F
<i>LLM-only (3-shot)</i>			
Llama-3.1-8B	3.21	18.70	3.31
Qwen3-8B	1.09	24.27	3.30
Sonnet 4.6	2.96	28.96	4.24
GPT-5.4	0.30	17.84	3.88
<i>Fine-tuned on OOD training data</i>			
Qwen3-8B-FT	2.20	34.72	4.07
BioT5+	4.37	32.64	4.49
InstructProtein	5.96	31.68	3.87
Prot2Text-V2	19.35	49.81	6.23
<i>Domain-specific pretrained</i>			
Evolla-10B-DPO	10.54	19.05	3.96
<i>Retrieval-based</i>			
MMseqs2	21.91	44.52	5.60
RAPM with GPT-4.1	24.11	37.30	5.89
RAPM with Sonnet 4.6	35.56	42.48	6.06
Only-Structure	44.26	46.42	6.19
<b>ProtQueSt</b>	<b>48.79</b>	<b>50.84</b>	<b>6.40</b>

prediction regime (Table 1; implementation details and fine-tuning hyperparameters in Appendix L): *LLM-only (3-shot)* prompts frontier general-purpose models without protein-specific input; *domain-specific pretrained* uses protein-text foundation models with end-to-end alignment; *fine-tuned on the OOD train split* provides direct supervision on the target task; and *retrieval-based* transfers annotations from neighbor proteins, the family that ProtQueSt belongs to.

## 4.2. Results

ProtQueSt achieves the highest Entity-BLEU (48.79) and Judge-F (6.40) across all models (Table 1), with large gains over the strongest prior methods (per-task breakdowns in Tables 15 and 16). Bootstrap 95% CIs are tight (Entity-BLEU [48.44, 50.16], Judge-F [6.31, 6.49]; full per-task intervals in Appendix Table 27). Per-task analysis confirms that the contrastive channel contributes complementary candidates that the structural retriever misses (Appendix F).

## 4.3. Ablation Studies

We conduct ablations to isolate the effect of each design choice. Reported metrics include Entity-BLEU (end-to-end) and Recall@10 (deterministic retrieval quality). Full tables with all model variants are in Appendix I.

**Query pooling and FiLM are jointly essential** (Table 2). Without query pooling, ProtFiLM *degrades* below the structural-only baseline (Recall@10: 0.229 vs. 0.231). With

Table 2. Ablation study ( $n=256$  per task, masked contrastive pool). R@10: Recall@10; E-BL: Entity-BLEU. All use hybrid 5+5 merging except Struct-only (10 structural).

FiLM	Q-pool	R@10	E-BL
<i>Struct-only baseline</i>		0.231	44.95
None	None	0.238	44.24
ProtFiLM	None	0.229	42.49
BiFiLM	None	0.264	44.19
None	0.65	0.256	45.87
ProtFiLM	0.65	<b>0.324</b>	<b>47.46</b>
BiFiLM	0.65	0.303	45.96

query pooling ( $\tau_{\text{query}}=0.65$ ), the same model jumps to 0.324, a 40% improvement. Query pooling alone (no FiLM) yields a smaller gain (0.256). The two mechanisms are complementary: query pooling provides clean training signal by restricting comparisons to the same question type, and FiLM gives the adapter the capacity to act on that signal by selecting query-relevant feature channels (Appendix Figure 6). Neither alone is sufficient.

**False-negative filtering.** Removing the filter degrades Bi-FiLM Recall@10 by 9% and ProtFiLM by 4% (Appendix Table 22), confirming that unmasked paraphrases corrupt the training signal. The effect is larger for BiFiLM, which modulates the text side and is therefore more exposed to annotation-side noise (detailed analysis in Appendix C).

The query-conditioned design also remains effective when swapping to a fully open-source encoder and generator (Appendix Table 19; qualitative analysis in Appendix J).

## 5. Conclusion and Future Directions

Text-based protein sequence understanding should be query-conditioned: the functional meaning of a sequence emerges from the interaction between the sequence and a specific biological question. ProtQueSt demonstrates this by adding a contrastive channel that learns query-dependent protein-text correspondence through FiLM conditioning, complementing the structural retriever. Our ablations confirm that this requires both query-conditioned representation (FiLM) and query-conditioned training signal (query pooling), as neither alone improves over structural retrieval.

Our approach currently relies on a single benchmark and a fixed set of frozen encoders, which limits the scope of our evaluation. Future work could extend the query-conditioned paradigm to richer conditioning signals such as compositional multi-axis queries and multi-turn refinement. Advances in protein encoders would directly raise the representational ceiling that conditioning can expose, and additional benchmarking would test how broadly the paradigm generalizes. Computational cost analysis, broader impact, and additional limitations are discussed in Appendices K and M.

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## A. Dataset Statistics and Characterization

### A.1. Per-Task Sample Counts

Table 3 summarizes the Prot-Inst-OOD benchmark composition. The full dataset contains 299,029 protein-annotation pairs: 284,526 in the retrieval pool (train split) and 14,503 in the OOD test set. Of the pool, 270,300 serve as contrastive training anchors; the remaining 14,226 (5% per task,  $dev\_frac=0.05$ ,  $split\_seed=42$ ) are held out for early-stopping validation. Dev proteins remain in the retrieval pool but are not used as contrastive training anchors.

Table 3. Prot-Inst-OOD dataset split statistics.

Task	Pool	Train	Dev	Test
Catalytic activity	51,187	48,628	2,559	1,987
Domain/motif	42,368	40,250	2,118	2,732
General function	82,275	78,161	4,114	4,297
Protein function	108,696	103,261	5,435	5,487
<b>Total</b>	<b>284,526</b>	<b>270,300</b>	<b>14,226</b>	<b>14,503</b>

Prot-Inst-OOD is derived from a UniProt-based protein-description corpus following the out-of-distribution split protocol introduced by Wu et al. (2025). The benchmark is constructed in three steps: (1) all sequences are clustered using MMseqs2 (Steinegger & Söding, 2017) with no minimum sequence identity threshold, (2) clusters are randomly partitioned into training and test sets at an 80/20 ratio, and (3) any test sample whose annotation can be directly retrieved from the training set is reassigned to training, repeated twice to minimize label leakage. Crucially, OOD-ness is defined operationally by this leakage-elimination procedure rather than by an explicit sequence identity cut-off or novel GO term constraint, meaning that test proteins are guaranteed to lack directly retrievable training annotations but may still share sequence similarity with training examples.

### A.2. Sequence Length Distribution

Table 4. Protein sequence length statistics per task (test set).

Task	Min length	Max length
Catalytic activity	12	768
Domain/motif	37	768
General function	11	768
Protein function	11	768

No proteins in the test set exceed the ESM-2 or ProtT5 maximum input length of 1,024 tokens (one amino acid corresponds to one token in ESM-2’s vocabulary). The maximum observed sequence length across all tasks is 768 residues (Table 4), so truncation is not an issue for this dataset.

### A.3. Query Templates and Instruction Embedding Analysis

Each task draws from a pool of paraphrased natural-language instruction templates: 5 for catalytic activity, 10 for domain/motif, 7 for general function, and 38 for protein function (60 total). Table 5 shows one representative template per task. Instructions within a task are semantically coherent (intra-task mean cosine similarity 0.74–0.87), while covering diverse phrasing that helps the model generalize across surface-level variation.

Table 5. Representative instruction templates for the four Prot-Inst-OOD tasks. Each task has multiple paraphrased variants (5, 10, 7, and 38 respectively).

Task	#	Example instruction template
Catalytic activity	5	“Please evaluate the following protein sequence and provide an explanation of the enzyme’s catalytic activity, including the chemical reaction it facilitates: ”
Domain/motif	10	“Please conduct a domain/motif search on the given protein sequence and provide your findings. The sequence is: ”
General function	7	“Assess the following protein sequence and provide a brief report on its primary characteristics: ”
Protein function	38	“Considering the protein sequence below, predict its biological function by examining its structural features and comparing it to functionally characterized proteins: ”

To validate that the query-pooling threshold  $\tau_{query}=0.65$  cleanly separates task types, we report the mean pairwise cosine similarity between all instruction embeddings of each task pair, computed with TEXT-EMBEDDING-3-LARGE (Table 6). All six cross-task means fall below 0.66, with catalytic activity vs. domain/motif the most distinct (0.551)

and general function vs. protein function the closest (0.656). Individual cross-task pairs can exceed 0.65 (e.g., 58.6% of general–protein pairs), reflecting genuine semantic overlap between these broader functional queries. This means query pooling treats some instruction pairs from these task categories as interchangeable during negative sampling, which is acceptable given their overlapping annotation space.

Table 6. Mean pairwise cosine similarity of instruction embeddings across task pairs (TEXT-EMBEDDING-3-LARGE). Off-diagonal: averaged over all template pairs between the two tasks. Diagonal: mean intra-task similarity (excluding self-pairs).

	Cat. act.	Dom./mot.	Gen. func.	Prot. func.
Cat. act.	0.867	0.551	0.600	0.595
Dom./mot.	0.551	0.793	0.654	0.639
Gen. func.	0.600	0.654	0.780	0.656
Prot. func.	0.595	0.639	0.656	0.741

#### A.4. Query-Filtered Pool Sizes

Query filtering at  $\tau_{\text{query}}=0.65$  restricts which pool candidates provide gradient for each anchor during training. It does not remove proteins from the training set. Because each task has multiple instruction templates, the effective pool size varies per anchor depending on which instruction it carries. Table 7 reports the mean number of eligible pool candidates per task, averaged over all anchors (and thus all instruction variants).

Table 7. Effective contrastive pool sizes after query filtering ( $\tau_{\text{query}}=0.65$ ). Mean  $\pm$  std computed over all anchors in each task; min/max reflect variation across the different instruction templates.

Task	Mean	Std	Min	Max	% of pool
Cat. act.	69,194	20,442	51,187	109,005	24.3
Dom./mot.	128,333	45,365	43,930	196,590	45.1
Gen. func.	177,747	38,507	126,626	241,421	62.5
Prot. func.	170,654	35,482	33,373	258,202	60.0

Catalytic activity retains only 24.3% of the pool on average, compared to 62.5% for general function, a  $2.6\times$  disparity. The wide min–max ranges (driven by the diversity of instruction templates within each task) show that some instruction variants are much more selective than others; for example, certain protein–function instructions retain as few as 33 K candidates while others retain 258 K. Despite this variability, our per-task results (Section F.3) suggest that intrinsic task difficulty (precise reaction chemistry vs. broad functional descriptions) is likely the dominant factor in performance differences.

#### A.5. Gold Reference Structure

Each sample has two gold components: (1) a compact *meta-data* string containing GO-style annotations (e.g., “*lipid*

*binding*” or “*plastid, ribonucleoprotein complex, ribosome — rRNA binding*”), and (2) a longer natural-language *description*. The contrastive retriever is trained to match protein embeddings with annotation metadata embeddings. The LLM generator produces free-text descriptions. Entity-BLEU bridges the two by evaluating whether biologically relevant entities from the metadata appear in the generated text.

## B. Model Architecture Details

### B.1. Frozen Encoder Details

**ESM-2.** We use FACEBOOK/ESM2\_T33\_650M\_UR50D (650M parameters). The protein representation is extracted via the CLS token, yielding a 1,280-d embedding that captures evolutionary co-variation patterns.

**ProstT5.** A 1,024-d embedding is obtained via masked mean pooling over the encoder’s *last\_hidden\_state*, weighted by the attention mask. ProstT5 provides fold-aware representations derived from sequence via structure-conditioned language modeling; “structural” in our context refers to these learned fold representations, not actual 3D coordinate comparison.

**Text encoder.** We use OpenAI TEXT-EMBEDDING-3-LARGE (3,072-d) for all text embeddings (instruction queries and annotation metadata). All embeddings are pre-cached offline as .npy files keyed by SHA1 hashes; no text encoder runs during training. The cached files are archived for reproducibility.

**Fusion.** The protein representation is the simple concatenation of ESM-2 and ProstT5 embeddings:  $\mathbf{h} \in \mathbb{R}^{2304}$ . The adapter MLP learns to fuse them in the shared space.

### B.2. Adapter MLP Specification

Each tower uses the same adapter architecture (Table 8). Both outputs are L2-normalized to a shared 512-d embedding space.

Table 8. Adapter MLP specification for protein and text towers.

Layer	Protein tower	Text tower
LayerNorm	2,304	3,072
Linear	2,304 $\rightarrow$ 1,024	3,072 $\rightarrow$ 1,024
Activation	GELU	GELU
Dropout	$p=0.1$	$p=0.1$
Linear	1,024 $\rightarrow$ 512	1,024 $\rightarrow$ 512
LayerNorm	512	512
L2 normalize	unit sphere	

### B.3. FiLM Conditioning Details

The instruction embedding  $\mathbf{q} \in \mathbb{R}^{3072}$  is first compressed to a 64-d bottleneck:  $\tilde{\mathbf{q}} = \text{GELU}(\text{LN}(\mathbf{q})\mathbf{W}_q)$ , where  $\mathbf{W}_q \in \mathbb{R}^{3072 \times 64}$ . The bottleneck is deliberately low-dimensional because protein function queries occupy a narrow subspace of the text embedding space (only a handful of distinct question types exist). From  $\tilde{\mathbf{q}}$ , two linear maps (bias=False, zero-initialized) produce scale and shift vectors in  $\mathbb{R}^{512}$ .

The zero initialization ensures that FiLM begins as an identity transform ( $\mathbf{z}_{\text{out}} = \mathbf{z}$ ), so any deviation from identity is learned. The evidence that FiLM learns meaningful query-specific modulation (rather than generic capacity) is the comparison: FiLM *degrades* without query pooling (R@10: 0.229 vs. 0.231 structural-only), but with query pooling the same model jumps to 0.324. This pattern (FiLM is harmful without clean signal, beneficial with it) strongly suggests query-dependent modulation rather than mere parameter overhead. Direct inspection of the learned vectors confirms this: scale vectors are weakly or negatively correlated across tasks (cosines  $-0.28$  to  $+0.35$ ), showing that FiLM amplifies different dimensions for different instructions, while shift vectors act as a largely shared offset (Figure 6).

**Parameter counts.** Table 9 summarizes trainable parameter counts for each FiLM configuration.

Table 9. Trainable parameter counts by configuration.

Configuration	Parameters
No FiLM (unconditional)	6.57M
ProfFiLM (protein tower only)	6.84M
BiFiLM (both towers)	7.11M

### B.4. Notation Summary

Table 10 consolidates the mathematical notation used throughout the paper.

Table 10. Summary of notation.

Symbol	Dim	Description
$\mathbf{h}$	$\mathbb{R}^{2304}$	Fused ESM-2 + ProstT5 protein repr.
$\mathbf{z}_p$	$\mathbb{R}^{512}$	Adapted protein embedding
$\mathbf{z}_t$	$\mathbb{R}^{512}$	Adapted text embedding
$\mathbf{q}$	$\mathbb{R}^{3072}$	Instruction query embedding
$\tilde{\mathbf{q}}$	$\mathbb{R}^{64}$	Compressed query bottleneck
$\mathbf{p}$	$\mathbb{R}^{1024}$	L2-normalized ProstT5 embedding
$\mathbf{e}$	$\mathbb{R}^{1280}$	L2-normalized ESM-2 embedding
$\mathbf{t}$	$\mathbb{R}^{3072}$	Text (annotation) embedding
$\tau$	scalar	Learnable temperature
$\alpha$	scalar	Structural retriever mixing weight
$\tau_{\text{query}}$	scalar	Query-pooling cosine threshold
$\tau_{\text{in}}$	scalar	False-negative cosine threshold
$b$	scalar	SigLIP bias term

## C. Training Details

### C.1. Optimization Hyperparameters

Table 11 consolidates all training hyperparameters.

Table 11. Consolidated training hyperparameters.

Category	Parameter	Value
Optimization	Optimizer	AdamW
	Learning rate	$10^{-4}$
	Weight decay	$2 \times 10^{-2}$
	Scheduler	OneCycleLR (cosine)
	Warmup	3% of total steps
	Gradient clipping	1.0 (global norm)
	Seed	42
Batching	Micro batch size	64
	Gradient accumulation	4 steps
	Effective batch size	256
	DataLoader workers	4
Early Stopping	Max epochs	50
	Patience	6 (dev MRR@10)
Architecture	$d_{\text{hidden}}$	1,024
	$d_{\text{model}}$	512
	$d_{\text{query\_proj}}$	64
	Dropout	0.1
Temperature	$\tau_0$ (initial)	0.07
	$\tau_{\text{min}}$	0.01
	$\tau_{\text{max}}$	0.20
Negatives	Pool neg. per anchor	15
	SigLIP bias $b$	$-10.0$

**Hardware and compute.** All models are trained on a single NVIDIA A6000 GPU. Training takes approximately 5 minutes per epoch; typical runs early-stop at 15–30 epochs, yielding  $\sim 1.5$ – $2.5$  hours per model. With 21 contrastive model variants, total compute for the full ablation is approximately 40–50 GPU-hours.

### C.2. FiLM Initialization and Temperature

The scale and shift maps in the FiLM conditioning layer are initialized to zero, so training begins from the unconditional baseline ( $\mathbf{z}_{\text{out}} = \mathbf{z}$ ) and gradually introduces query-dependent modulation. The bottleneck projection  $\mathbf{W}_q$  and all scale/shift parameters are learned end-to-end via the contrastive loss. The learnable temperature  $\tau$  is parameterized as  $\tau = \exp(\log \tau)$  and clamped to  $[0.01, 0.20]$ , initialized at  $\tau_0 = 0.07$ .

### C.3. Negative Sampling Strategy

Each protein’s training negatives come from two sources:

- In-batch negatives:** Gold annotations of other proteins in the same mini-batch. Proteins sharing the same gold metadata SHA1 hash are treated as co-positives rather

than negatives.

2. **Pool negatives:** Up to 15 annotations sampled from the protein’s structural retrieval candidates (top-100 by weighted cosine  $\alpha \cdot \text{cos}_{\text{ProstT5}} + (1-\alpha) \cdot \text{cos}_{\text{ESM-2}}$  over the cross-task training corpus). These yield harder comparisons than random sampling because they come from the protein’s structural neighborhood. Pool candidates are pre-filtered to exclude (a) the anchor’s own gold annotation (exact SHA1 match) and (b) annotations with  $\text{cos}(\mathbf{g}_{\text{gold}}, \mathbf{g}_j) \geq \tau_{\text{fn}}$ , where  $\mathbf{g}$  denotes the metadata embedding.

**Query pooling.** For each anchor with instruction  $\mathbf{q}_i$ , any batch or pool candidate  $j$  is masked (excluded from gradient) if  $\text{cos}(\mathbf{q}_i, \mathbf{q}_j) < \tau_{\text{query}}$ . This ensures only candidates with instructively similar queries provide gradient signals, preventing trivial cross-task shortcuts.

#### C.4. False-Negative Filtering

False-negative protection operates at three layers:

1. **Candidate building** (offline, train split only): Drop candidates matching the anchor’s gold SHA1 hash or exceeding the semantic cosine threshold  $\tau_{\text{fn}}$  against the gold metadata embedding.
2. **Dataset pool sampling** (per-sample): Defense-in-depth cosine filter on subsampled pool negatives, handling cases where manifests were built with a different threshold than training.
3. **Collate-time valid mask** (per-batch): Same cosine threshold applied to the full batch matrix, including in-batch negatives. This layer is not redundant with layers 1–2 because in-batch negatives are arbitrary gold annotations from other proteins that never went through per-anchor candidate construction.

#### C.5. Data Balancing

A `TaskBalancedSampler` groups dataset indices by task, pads each task’s shuffled index list to the length of the largest task (by cycling), then interleaves one index per task per round, flattens, and shuffles. Smaller tasks are oversampled so that all tasks have equal representation per epoch.

#### C.6. Loss Functions

**SigLIP (primary).** A pairwise sigmoid loss with bias  $b = -10.0$ , adopted from the SigLIP paper’s recommendation for settings with low positive-pair rates ( $\sim 0.4\%$  per row in a batch of 256 + 15 candidates):

$$\mathcal{L}_{ij} = -\log \sigma(y_{ij}(\text{score}(p_i, t_j) + b)) \quad (3)$$

Row reweighting ensures that positives and negatives each contribute exactly 50% of each row’s loss: positives receive weight  $0.5/n_{\text{pos}}$  and negatives  $0.5/n_{\text{neg}}$ . The loss is symmetrized (protein→text + text→protein) for in-batch negatives and protein→text only for pool negatives.

**Soft-SigLIP (variant).** Negatives are weighted by inverse metadata similarity to the gold:  $w_j = \text{clip}(1 - \text{cos}(\mathbf{g}_{\text{gold}}, \mathbf{g}_j), 0, 1)$ , where  $\mathbf{g}$  denotes the raw metadata embedding. This *down-weights* negatives that are semantically close to the gold (potential false negatives), providing a softer alternative to the hard threshold.

#### C.7. Stage 2: Fine-Tuning

An optional second stage loads a Stage 1 checkpoint and switches from the contrastive SigLIP objective to a pointwise ranking loss. For each anchor protein, we form a candidate list consisting of the gold annotation plus up to 49 negatives drawn from the same per-protein candidate pool used in Stage 1 (not re-mined). The loss is softmax cross-entropy that maximizes the probability of the gold annotation scoring highest:

$$\mathcal{L}_{\text{rank}} = -\log \frac{\exp(s_{\text{gold}})}{\sum_j \exp(s_j)},$$

where  $s_j = \mathbf{z}_p \cdot \mathbf{z}_{t_j} / \tau$  is the cosine similarity score. The entire protein adapter (including any protein-side FiLM layers) is frozen; only the text adapter (with any text-side FiLM layers) and the learnable temperature  $\log \tau$  receive gradients.

Table 12. Stage 2 hyperparameters.

Parameter	Value
Learning rate	$2 \times 10^{-5}$
Weight decay	$10^{-2}$
LR schedule	OneCycleLR (cosine, 5% warmup)
Micro batch size	8
Effective batch size	64 ( $8 \times$ grad. accum.)
Max candidates / protein	1 gold + 49 negatives
FN threshold	$\tau_{\text{fn}}=0.7$
Max epochs	6
Early stopping patience	3 (dev MRR@10)
Frozen components	Protein adapter (incl. FiLM)

Our ablations (Table 23) show that Stage 2 provides marginal and inconsistent gains over Stage 1 alone.

#### C.8. Dev Evaluation During Training

During training, validation uses `dev.max_per_task=512` samples per task with `dev.n_neg=9` negatives per anchor, computing MRR@10 over 10 candidates (1 positive + 9 negatives). We assume that ranking ability in a 10-candidate setting transfers to larger pools.

## D. LLM Generation Prompt and Setup

### D.1. Generation Prompt Template

The full generation prompt is shown in Figure 2. The `contexts.json` payload labels each item with "source": "structural" or "source": "contrastive", so the LLM sees accurate source metadata.

The `contexts.json` field is structured as {"structural": [...], "contrastive": [...]}, where each item contains `db_label` (the annotation text), `confidence_level` (the raw numeric similarity score), and `source`.

#### System prompt:

You are a protein function prediction expert with deep knowledge of biological terminology.

**User prompt template** (placeholders {instruction}, {sequence}, {contexts.json}, and {few\_shot.json} are filled per sample):

**Instruction:** {instruction}

**Protein sequence:** {sequence}

**Retrieved protein annotations:** {contexts.json}

**Example input–output pairs for this task:** {few\_shot.json}

Based on the instruction, the protein sequence, the retrieved information, and the examples:

- Use **PRECISE biological terminology** from the retrieved annotations
- **Prioritize high-confidence matches**—they are most similar
- **Avoid generic descriptions**—be specific

Output **ONLY** the functional description of this protein in the following JSON format:

```
{"description": "..."}

```

Do not output any other text or explanation. Only output the JSON answer.

Figure 2. Generation prompt template.

### D.2. Generation Configuration

Table 13. LLM generation configuration.

Parameter	Value
Primary model	Claude Sonnet 4.6
Temperature	1.0
Max tokens	2,048
Parallelism	30 threads
Retries	3 (exponential backoff)
Few-shot examples per task	3

Temperature 1.0 is the Anthropic API default and the recommended setting for generative (as opposed to analytical) tasks (Anthropic, 2025). Since our task produces free-text protein descriptions, we use the default without further tun-

ing.

### D.3. Few-Shot Example Selection

For each task, 3 few-shot examples are sampled from the training set using a fixed random seed (`np.random.default_rng(42)`). The same 3 examples are reused for every test sample within that task. Protein sequences are truncated to 80 characters followed by “...” in the prompt.

## E. LLM-as-Judge Evaluation

### E.1. Judge Configuration

We evaluated model predictions using Claude Sonnet 4.6 as an LLM-as-judge evaluator. The judge was run with extended thinking enabled (`budget_tokens=4,000`), temperature 1.0, and effective `max_tokens=5,024` (combining a 1,024-token output budget on top of the 4,000-token thinking budget). For each model, we scored 500 randomly sampled predictions per task using `seed=42`, across four tasks, for a total of 2,000 nominal judged samples per model. All baselines and our system were evaluated under the same judge configuration to ensure internally consistent cross-model comparisons.

### E.2. Scoring Rubric

The judge assigns scores along four discrete axes using the fixed 5-point scale {0, 3, 5, 7, 10}:

- **Recall:** fraction of reference bio-entities recovered in the prediction.
- **Precision:** whether the prediction remains focused on reference-supported or biologically reasonable claims.
- **Specificity:** granularity of biological content, distinguishing concrete entities from vague functional language.
- **Plausibility:** biological coherence and freedom from contradictions or fabrications.

The final per-sample score is the mean of all active axis scores. Model-level results are macro-averaged across tasks.

### E.3. Full Judge Prompt

The full judge prompt is shown in Figure 7.

## F. Recall@10 Definition and Retrieval Pool Analysis

### F.1. Recall@10 Definition

For each test sample, the system produces a merged retrieval list of 10 candidates (5 from the structural channel and 5

from the contrastive channel). We compute Recall@10 by checking whether the gold annotation’s exact text (identified by its SHA1 hash) appears anywhere in this merged list:

$$\text{Recall@10} = \frac{1}{N} \sum_{i=1}^N \mathbf{1}[\text{gold}_i \in \text{merged}_{10}(i)] \quad (4)$$

where  $N$  is the number of test samples. This is a strict exact-match criterion: semantically equivalent paraphrases of the gold do not count as hits. We report the macro-average across the four tasks. Because Recall@10 depends only on the retriever’s ranked output and not on LLM generation, it is fully deterministic and complements the LLM-dependent Entity-BLEU metric in our ablation comparisons.

## F.2. Hybrid Merge Mechanics

In the hybrid 5+5 scheme, structural candidates always occupy positions 1–5 and contrastive candidates positions 6–10 in the merged list. Consequently, R@1 is always determined by the structural channel alone and is identical across all contrastive model variants. The contrastive channel contributes exclusively to R@10 improvement (positions 6–10).

At inference, contrastive scores are computed as  $\text{score} = 0.5 \cdot ((\mathbf{z}_{\text{corpus}} \cdot \mathbf{z}_p) + 1.0)$ , normalizing cosine similarity from  $[-1, 1]$  to  $[0, 1]$ . When  $\tau_{\text{query}} > 0$ , scores are masked ( $-\infty$ ) for corpus entries whose training queries have cosine similarity below  $\tau_{\text{query}}$  with the current instruction.

## F.3. Per-Task Retrieval Pool Coverage

Figure 3 reports per-task Recall@10 for the structural-only baseline and ProtQueSt hybrid on the full dataset (14,503 samples).

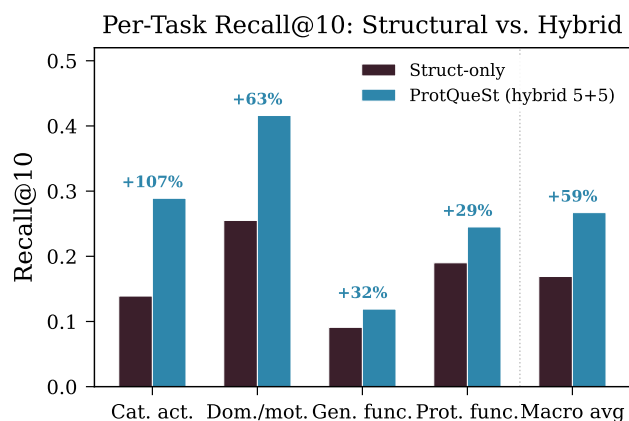


Figure 3. Per-task Recall@10 on the full dataset (14,503 samples). Percentages indicate the relative gain from the contrastive channel. Catalytic activity benefits most (+107%).

The contrastive channel’s relative contribution is *largest* on catalytic activity (+107%) despite its smaller absolute

Entity-BLEU, suggesting the contrastive retriever finds relevant catalytic annotations but the LLM struggles to synthesize them into correct descriptions.

## G. Full Baseline Comparison

Tables 14, 15, and 16 report the full per-metric breakdown, per-task Entity-BLEU, and per-task Judge-F scores for all baselines and our model on the full Prot-Inst-OOD benchmark.

ProtQueSt achieves the highest Judge-F on catalytic activity (6.67) and protein function (6.82), while Prot2Text-V2 leads on domain/motif (6.92) and general function (6.41). This suggests ProtQueSt’s advantage is strongest on tasks requiring precise functional descriptions (catalytic reactions, protein function) rather than structural motif annotation where fine-tuned sequence-to-text models excel at reproducing training-set vocabulary.

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Table 14. Full evaluation of all baselines and our model on Prot-Inst-OOD (14,503 samples per model for lexical metrics; 500 predictions per task, 2,000 nominal per model for the judge). **Lexical metrics (0–100 scale):** Entity-BLEU (E-BL), METEOR, and ROUGE-L (RG-L). **LLM-as-judge axes (0–10 scale):** Recall (J-Rec), Precision (J-Prec), Specificity (J-Spec), Plausibility (J-Plaus), and Final (Judge-F). Bold marks the top score in each column.

Method	E-BL	METEOR	RG-L	J-Rec	J-Prec	J-Spec	J-Plaus	Judge-F
<i>LLM-only (3-shot)</i>								
Llama-3.1-8B	3.21	18.70	11.27	0.14	3.01	6.69	3.41	3.31
Qwen3-8B	1.09	24.27	23.52	0.27	2.13	7.90	2.91	3.30
Sonnet 4.6	2.96	28.96	28.57	1.36	2.67	8.13	4.79	4.24
GPT-5.4	0.30	17.84	16.83	0.83	3.16	7.68	3.85	3.88
<i>Domain-specific pretrained</i>								
Evolla-10B-DPO	10.54	19.05	10.18	1.58	3.31	6.37	4.59	3.96
<i>Fine-tuned on OOD training data</i>								
Qwen3-8B-FT	2.20	34.72	37.20	0.74	3.28	8.95	3.31	4.07
BioT5+	4.37	32.64	37.68	1.24	3.99	8.45	4.27	4.49
InstructProtein	5.96	31.68	34.15	0.69	3.57	7.39	3.85	3.87
Prot2Text-V2	19.35	49.81	<b>50.00</b>	3.34	<b>6.39</b>	8.82	<b>6.38</b>	6.23
<i>Retrieval-based</i>								
MMseqs2	21.91	44.52	43.65	2.79	4.92	9.30	5.37	5.60
RAPM (GPT-4.1)	24.11	37.30	28.83	4.33	4.74	8.77	5.73	5.89
RAPM (Sonnet 4.6)	35.56	42.48	32.75	4.57	4.86	9.00	5.79	6.06
Only-Structure	44.26	46.42	27.72	5.34	3.70	9.59	6.13	6.19
<b>ProtQueSt</b>	<b>48.79</b>	<b>50.84</b>	31.43	<b>5.59</b>	4.04	<b>9.61</b>	6.33	<b>6.40</b>

Table 15. Per-task Entity-BLEU on the full Prot-Inst-OOD benchmark (14,503 samples). Bold marks the top score per task.

Method	Cat. act.	Dom./mot.	Gen. func.	Prot. func.	Avg
<i>LLM-only (3-shot)</i>					
Llama-3.1-8B	2.80	0.01	8.93	1.08	3.21
Qwen3-8B	0.01	0.00	3.95	0.41	1.09
Sonnet 4.6	0.17	0.49	8.43	2.75	2.96
GPT-5.4	0.01	0.02	0.20	0.96	0.30
<i>Domain-specific pretrained</i>					
Evolla-10B-DPO	7.97	4.32	16.69	13.20	10.54
<i>Fine-tuned on OOD training data</i>					
Qwen3-8B-FT	0.75	0.25	1.87	5.92	2.20
BioT5+	0.10	2.34	0.24	14.80	4.37
InstructProtein	1.02	0.00	0.31	22.51	5.96
Prot2Text-V2	18.51	13.94	22.54	22.39	19.35
<i>Retrieval-based</i>					
Only-Structure	45.47	43.84	30.64	57.09	44.26
RAPM (GPT-4.1)	31.29	23.28	8.87	33.02	24.12
RAPM (Sonnet-4.6)	31.53	35.13	24.15	51.43	35.56
MMseqs2	12.79	12.87	16.76	45.22	21.91
<b>ProtQueSt</b>	<b>50.84</b>	<b>52.17</b>	<b>32.63</b>	<b>59.51</b>	<b>48.79</b>

Table 16. Per-task Judge-F (0–10 scale, 500 samples/task) on Prot-Inst-OOD. Bold marks the top score per task.

Method	Cat. act.	Dom./mot.	Gen. func.	Prot. func.	Avg
<i>LLM-only (3-shot)</i>					
Llama-3.1-8B	2.74	3.62	4.02	2.87	3.31
Qwen3-8B	2.94	4.58	2.51	3.18	3.30
Sonnet 4.6	4.05	5.30	2.96	4.66	4.24
GPT-5.4	3.19	4.76	3.40	4.19	3.88
<i>Domain-specific pretrained</i>					
Evolla-10B-DPO	3.53	4.29	3.61	4.41	3.96
<i>Fine-tuned on OOD training data</i>					
Qwen3-8B-FT	3.35	5.57	2.90	4.45	4.07
BioT5+	3.58	6.02	3.49	4.87	4.49
InstructProtein	2.99	5.06	2.42	5.02	3.87
Prot2Text-V2	5.15	<b>6.92</b>	<b>6.41</b>	6.45	6.23
<i>Retrieval-based</i>					
MMseqs2	4.37	6.70	4.96	6.36	5.60
RAPM (GPT-4.1)	6.17	6.05	5.14	6.22	5.89
RAPM (Sonnet 4.6)	5.84	6.68	5.31	6.40	6.06
Only-Structure	6.39	6.12	5.46	6.79	6.19
<b>ProtQueSt</b>	<b>6.67</b>	6.46	5.63	<b>6.82</b>	<b>6.40</b>

## H. Full-Dataset Per-Task Results

### H.1. Per-Task Lexical Results

Table 17 shows per-task results for ProtQueSt on the full Prot-Inst-OOD benchmark (14,503 samples), using the Prot-FiLM + SigLIP model ( $\tau_{\text{query}}=0.65$ ) with Claude Sonnet 4.6 generation.

Table 17. Per-task full-dataset results for ProtQueSt (ProtFiLM + SigLIP,  $\tau_{\text{query}}=0.65$ ).

Task	$n$	E-BL	METEOR	RG-L
Catalytic activity	1,987	50.84	53.81	48.25
Domain/motif	2,732	52.17	49.22	21.25
General function	4,297	32.63	41.09	27.54
Protein function	5,487	59.51	59.24	28.70
<b>Macro avg</b>	<b>14,503</b>	<b>48.79</b>	<b>50.84</b>	<b>31.43</b>

### H.2. Per-Task Retrieval Metrics

Table 18 reports full-dataset per-task retrieval metrics for ProtQueSt and the structural-only baseline. MRR (Mean Reciprocal Rank) is  $\frac{1}{N} \sum_{i=1}^N \frac{1}{\text{rank}_i}$ , where  $\text{rank}_i$  is the position of the gold annotation in the ranked list (0 if absent).  $R@k$  is the fraction of samples whose gold annotation appears in the top  $k$ .

$R@1$  is identical per task across all models because position 1 is always the top structural candidate. The contrastive channel contributes exclusively to  $R@10$  improvement.

### H.3. Encoder × Generator Ablation

Table 19 crosses two text encoders with two generators, holding all upstream retrieval logic byte-identical.

Table 18. Full-dataset per-task retrieval metrics (14,503 samples). ProtQueSt uses hybrid 5+5 merging.

Retriever	Task	MRR	R@1	R@10
Struct-only	Cat. act.	0.050	0.017	0.139
	Dom./mot.	0.096	0.038	0.255
	Gen. func.	0.037	0.016	0.091
	Prot. func.	0.088	0.050	0.190
	<b>Macro</b>	<b>0.068</b>	<b>0.030</b>	<b>0.169</b>
ProtQueSt	Cat. act.	0.072	0.017	0.289
	Dom./mot.	0.120	0.038	0.416
	Gen. func.	0.041	0.016	0.119
	Prot. func.	0.096	0.050	0.245
	<b>Macro</b>	<b>0.082</b>	<b>0.030</b>	<b>0.267</b>

The fully open-source pipeline (BGE-M3 + Llama-3.1-8B-Instruct) achieves Judge-F 5.93, preserving 85% of the 3.09-point retrieval gain over the no-retrieval Llama-3.1-8B 3-shot baseline (3.31). The BGE-M3 variant was *retrained from scratch* with BGE-M3 text embeddings and rebuilt query/metadata matrices, not used as a drop-in replacement at inference.

Table 19. Ablation of ProtQueSt’s text encoder and generation model. We cross OpenAI’s TEXT-EMBEDDING-3-LARGE (closed-source, 3072-d) with BAAI’s BGE-M3 (open-source, 1024-d) on the encoder side, and Llama-3.1-8B-Instruct (open-weight) with Claude Sonnet 4.6 (proprietary) on the generator side.

Encoder	Generator	E-BL	METEOR	RG-L	J-Rec	J-Prec	J-Spec	J-Plaus	Judge-F
BGE-M3	Llama-3.1-8B-Instruct	19.04	32.31	26.91	3.97	4.62	9.22	5.90	5.93
	Claude Sonnet 4.6	43.68	46.89	30.35	5.27	4.03	9.54	6.22	6.27
TEXT-EMBEDDING-3-LARGE	Llama-3.1-8B-Instruct	23.24	35.61	28.15	4.32	<b>4.82</b>	9.22	6.15	6.13
	Claude Sonnet 4.6	<b>48.79</b>	<b>50.84</b>	<b>31.43</b>	<b>5.59</b>	4.04	<b>9.61</b>	<b>6.33</b>	<b>6.40</b>

## I. Comprehensive Ablation Tables

### I.1. Ablation Subset Characterization

All ablation experiments use  $n=256$  samples per task (1,024 total). This subset consists of the *first* 256 rows in JSONL file order per task (a deterministic prefix, not a random sample). Empirically, absolute metric values on this prefix differ from the full 14,503-sample benchmark, and the direction of the gap varies by metric and model. For example, structural-only R@10 is higher on the prefix (0.231 vs. 0.169), while ProtQueSt Entity-BLEU is higher on the full set. This means the subset is not uniformly easier or harder, but simply not representative of the full distribution. **Relative model rankings are preserved** (all models use the same prefix), so the ablations remain valid for comparing design choices.

### I.2. Merging Mode Sweep

This appendix expands the body ablation summary (Table 2) with full sweeps over merging modes (Figure 4), architecture (Table 20), false-negative threshold (Table 21), stage-2 fine-tuning (Table 23), and Recall@10 retrieval metrics (Table 24).

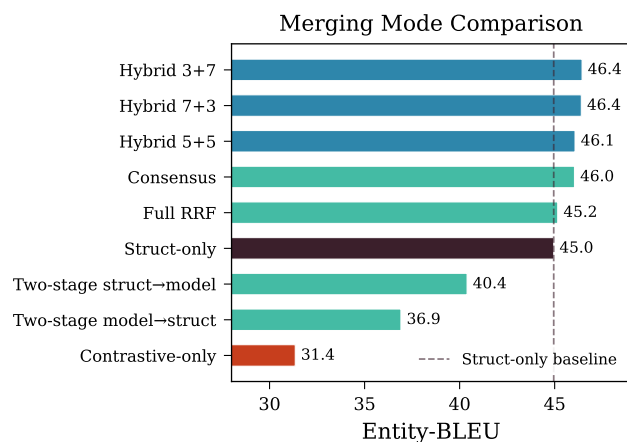


Figure 4. Merging mode sweep with BiFiLM + SigLIP ( $\tau_{\text{query}}=0.65$ ,  $n=256$ ). All modes use budget  $K=10$ . Dashed line marks the structural-only baseline. Hybrid modes (blue) cluster at the top; two-stage re-ranking degrades performance.

Table 20. Architecture ablation ( $n=256$ , masked / unmasked Entity-BLEU). Structural-only baseline: 44.95.

Loss	FiLM	Q-pool	E-BL (m/u)	$\Delta$
SigLIP	None	—	44.24	-0.71
SigLIP	None	0.65	45.87 / 44.04	+0.92
SigLIP	BiFiLM	—	44.19	-0.76
SigLIP	BiFiLM	0.65	45.96 / 45.75	+1.01
SigLIP	ProtFiLM	—	42.49	-2.46
SigLIP	ProtFiLM	0.65	<b>47.46</b> / 46.85	<b>+2.51</b>
Soft	BiFiLM	0.65	45.97 / 45.60	+1.02
Soft	ProtFiLM	0.65	47.07 / 47.18	+2.12

Table 21. False-negative threshold ablation ( $n=256$ , masked Entity-BLEU).

FiLM	Loss	$\tau_{\text{fn}}=0.60$	0.65	0.70
BiFiLM	SigLIP	46.35	46.51	45.96
BiFiLM	Soft	46.82	<b>47.88</b>	45.97
ProtFiLM	SigLIP	—	47.11	47.46
ProtFiLM	Soft	—	47.35	47.07

Table 22. False-negative threshold ablation (Recall@10, masked, hybrid 5+5, query-pooled). Best per row in bold.

FiLM	Loss	None	$\tau_{\text{fn}}=0.65$	$\tau_{\text{fn}}=0.70$
BiFiLM	SigLIP	0.285	<b>0.313</b>	0.303
ProtFiLM	SigLIP	0.310	0.308	<b>0.324</b>

Table 23. Softmax ranking stage-2 fine-tuning ( $n=256$ , masked Entity-BLEU).  $\Delta$ : Stage 1+2 minus Stage 1.

Loss	FiLM	fn	Stage 1	Stage 1+2	$\Delta$
SigLIP	BiFiLM	0.70	45.96	46.43	+0.47
SigLIP	BiFiLM	0.65	46.51	46.48	-0.03
Soft	BiFiLM	0.65	47.88	47.44	-0.44
SigLIP	ProtFiLM	0.70	47.46	46.90	-0.56
Soft	ProtFiLM	0.70	47.07	<b>47.90</b>	+0.83
SigLIP	ProtFiLM	0.65	47.11	47.22	+0.11
Soft	ProtFiLM	0.65	47.35	46.67	-0.68

Table 24. Recall@10 retrieval metrics ( $n=256$ , deterministic). Structural-only Recall@10: 0.231.

FiLM	Config	fn	R@10
<i>With query pooling (<math>\tau_{query}=0.65</math>)</i>			
ProtFiLM	SigLIP	0.70	0.324
ProtFiLM	Soft	0.70	0.309
BiFiLM	Soft	0.65	<b>0.328</b>
BiFiLM	SigLIP	0.65	0.313
ProtFiLM	SigLIP	0.65	0.308
ProtFiLM	Soft	0.65	0.306
<i>Without query pooling</i>			
BiFiLM	SigLIP	0.70	0.264
None	SigLIP	0.70	0.238
None	SigLIP	0.65	0.256
ProtFiLM	SigLIP	0.70	0.229

### I.3. Per-Task Merging Sweep

Figure 5 breaks down Entity-BLEU by task for the main merging strategies ( $n=256$ ), revealing that catalytic activity and general function are consistently the hardest tasks for the contrastive channel.

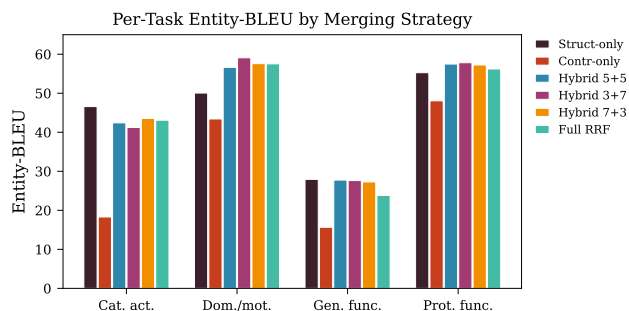


Figure 5. Per-task Entity-BLEU by merging strategy ( $n=256$ , BiFiLM + SigLIP,  $\tau_{query}=0.65$ ). The contrastive channel benefits domain/motif and protein function most; catalytic activity and general function remain structurally dominated.

The key pattern across merging strategies is that the contrastive channel helps most on domain/motif and protein function, where it lifts scores well above structural-only (e.g., domain/motif: 50.05  $\rightarrow$  56.63 with hybrid 5+5), but offers little on catalytic activity and general function, where structural-only already dominates the contrastive-only channel by a wide margin (46.58 vs. 18.29 and 27.93 vs. 15.65). Consequently, hybrid 5+5 balances the two channels well overall: it matches or exceeds structural-only on every task except catalytic activity (where replacing 5 structural slots with contrastive candidates costs  $\sim 4$  points). Shifting the budget toward contrastive (hybrid 3+7) further benefits domain/motif (+2.5) but widens the catalytic activity gap. Full RRF underperforms hybrid merging on general function (23.80 vs. 27.75), suggesting that interleaving by reciprocal rank can dilute the structural signal on tasks where contrastive retrieval is weak.

### I.4. SigLIP vs. Soft-SigLIP Comparison

Table 25 consolidates the comparison between SigLIP and Soft-SigLIP across all FiLM configurations and false-negative thresholds (all with query pooling at  $\tau_{query}=0.65$ ).

Table 25. SigLIP vs. Soft-SigLIP comparison ( $n=256$ , masked Entity-BLEU, query-pooled).

FiLM	$\tau_{fn}$	SigLIP	Soft	$\Delta$
BiFiLM	0.60	46.35	46.82	+0.47
BiFiLM	0.65	46.51	<b>47.88</b>	+1.37
BiFiLM	0.70	45.96	45.97	+0.01
ProtFiLM	0.65	47.11	47.35	+0.24
ProtFiLM	0.70	<b>47.46</b>	47.07	-0.39

Soft-SigLIP tends to help BiFiLM more consistently than ProtFiLM. BiFiLM applies query conditioning to the text encoder, making it more sensitive to annotation similarity; the soft weighting of near-gold negatives may be more beneficial in this setting. For ProtFiLM, the pattern is inconsistent (Soft helps at  $\tau_{fn}=0.65$  but hurts at 0.70).

### I.5. Masked vs. Unmasked Evaluation

For models trained with query pooling ( $\tau_{query}=0.65$ ), Table 20 reports both masked (contrastive pool filtered at  $\tau_{query}=0.65$ , matching training) and unmasked (full pool) Entity-BLEU. The gap is generally  $<1$  Entity-BLEU point, suggesting the contrastive model learns robust embeddings that transfer beyond the query-filtered training pool.

## J. Qualitative Examples

### J.1. Success Cases

We present three test samples where ProtQueSt produces correct predictions, each illustrating a different channel-contribution pattern: structural-only success (S1), contrastive-only rescue (S2), and complementary contribution from both channels (S3).

**Example S1: Domain/motif (GIY-YIG endonuclease).** *Structural channel succeeds; contrastive channel misses the gold annotation.*

- **Gold:** *GIY-YIG*
- **Structural top-3:** (1) *GIY-YIG* ( $\cos=0.975$ ); (2) “endo-deoxyribonuclease activity” (0.945); (3) “endonuclease activity” (0.933). The structural channel retrieves the exact gold domain at rank 1 and functionally related endonuclease annotations at ranks 2–5.
- **Contrastive top-3:** (1) Integrase catalytic (0.663); (2) SAM-dependent MTase C5-type (0.640); (3) viral DNA recombinase (0.631). All five contrastive candi-

825 dates are DNA-processing domains but none is the gold  
826 GIY-YIG motif.

- 827
- 828 • **Prediction:** "...GIY-YIG endonuclease domain, consistent with endodeoxyribonuclease activity and DNA binding functionality. This protein is predicted to function as a strand-specific endonuclease..."
  - 829
  - 830
  - 831
  - 832 • **Analysis:** The GIY-YIG superfamily has a distinctive fold that structural retrieval identifies directly. The contrastive channel retrieves DNA-associated domains (integrases, methyltransferases) that share functional keywords but not the specific motif. The LLM correctly anchors on the structural rank-1 annotation and integrates the endonuclease context from structural ranks 2–3 to produce a precise, detailed prediction.
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841 **Example S2: Domain/motif (FHA domain).** *Contrastive rescues; structural channel misses the gold annotation.*

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- 845 • **Gold:** *FHA* (Forkhead-Associated domain)
  - 846
  - 847 • **Structural top-3:** (1) "Plays a role in microtubule organization. Required for centriole subdistal appendage assembly" ( $\cos = 0.996$ ); (2) "Plays a role in microtubule organization" (0.993); (3) same (0.988). All five structural candidates describe biological processes (microtubule organization, cell growth regulation) but none names the FHA domain.
  - 848
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  - 850
  - 851
  - 852
  - 853 • **Contrastive top-3:** (1) *FHA, PH* (0.734); (2) *FHA* (0.725); (3) DWD box (0.671). The contrastive channel retrieves the exact gold domain at ranks 1–2.
  - 854
  - 855
  - 856
  - 857 • **Prediction:** "...FHA (Forkhead-Associated) domain, PH (Pleckstrin Homology) domain, and SH3-binding motifs. The protein likely plays a role in microtubule organization and centriole subdistal appendage assembly..."
  - 858
  - 859
  - 860
  - 861 • **Analysis:** The structural channel retrieves structurally homologous proteins but their annotations describe cellular functions, not domain identity. The contrastive channel, conditioned on a domain-prediction instruction, retrieves domain-level labels. The LLM synthesizes both: domain names from contrastive candidates and functional context from structural candidates, producing a comprehensive answer.
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871 **Example S3: Catalytic activity (glutamyl-tRNA synthetase).** *Both channels contribute; contrastive provides the exact reaction, structural provides enzymatic context.*

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- 874
- 875 • **Gold:**  $ATP + L\text{-glutamate} + tRNA(Glu) = AMP + diphosphate + L\text{-glutamyl-tRNA}(Glu)$
  - 876
  - 877 • **Structural top-3:** (1) "Catalyzes the attachment of glutamate to tRNA(Glu) in a two-step reaction" ( $\cos = 0.999$ );
  - 878
  - 879

(2) same description (0.998); (3) "HIGH region, KMSKS region" (0.998). Gold reaction appears at structural rank 5 (0.994).

- **Contrastive top-3:** (1)  $ATP + L\text{-glutamate} + tRNA(Glu) = \dots L\text{-glutamyl-tRNA}(Glu)$  (0.724); (2)  $ATP + L\text{-glutamine} + tRNA(Gln) = \dots$  (0.692); (3)  $ATP + L\text{-cysteine} + tRNA(Cys) = \dots$  (0.655). All five contrastive candidates are aminoacyl-tRNA synthetase reactions.
- **Prediction:** "...catalytic activity in the form of this chemical reaction:  $ATP + L\text{-glutamate} + tRNA(Glu) = AMP + diphosphate + L\text{-glutamyl-tRNA}(Glu)$ ."
- **Analysis:** The structural channel correctly identifies the enzyme family (glutamyl-tRNA synthetase) and retrieves the gold at rank 5, but its top candidates emphasize the two-step mechanism and conserved sequence motifs. The contrastive channel places the exact reaction equation at rank 1 with high confidence. Together, the channels provide both the mechanistic context and the precise reaction formula, enabling the LLM to produce an exact match.

## J.2. Failure Cases

We present two failure cases with different root causes.

**Example F1: Catalytic activity (collagenase → phospholipase A2).** Failure type: **retrieval failure** (gold not in top-10).

- **Gold:** Cleavage of the triple helix of collagen at 775-Gly|Ile-776 in the  $\alpha 1(I)$  chain.
- **Structural top-3:** (1) calcium ion binding / protein stabilization (0.922); (2) snake venom PLA2 / phospholipid hydrolysis (0.918); (3) lactose synthase regulatory subunit (0.911).
- **Contrastive top-3:** (1) hydroquinone oxidation (0.653); (2) retinal oxidation (0.643); (3) phosphate monoester hydrolysis (0.629).
- **Prediction:** "...Snake venom phospholipase A2 (PLA2) catalyzes the calcium-dependent hydrolysis of the 2-acyl groups in 3-sn-phosphoglycerides..."
- **Analysis:** The gold annotation (collagenase) does not appear in any of the 10 retrieved candidates. The structural channel returns calcium-binding proteins (collagenases are calcium-dependent metalloproteinases, explaining the structural similarity), while the contrastive channel retrieves generic hydrolysis reactions. The LLM faithfully synthesizes the retrieved PLA2 annotation, producing a confident but incorrect prediction.

**Example F2: Protein function (chloroplast thylakoid → DNA regulation).** Failure type: **generation failure** (correct annotation retrieved but LLM ignores it).

- **Gold:** *chloroplast thylakoid*
- **Structural top-3:** (1) regulation of DNA-templated transcription (0.926); (2) regulation of DNA-templated transcription (0.925); (3) H<sub>2</sub>O<sub>2</sub> removal / lignin degradation (0.924).
- **Contrastive top-3:** (1) **chloroplast thylakoid** (0.706); (2) extracellular region, mitochondrion (0.706); (3) chloroplast thylakoid lumen (0.704).
- **Prediction:** "...primary function related to regulation of DNA-templated transcription, with potential involvement in oxidative stress responses..."
- **Analysis:** The contrastive channel correctly retrieves "chloroplast thylakoid" at rank 1, but the structural channel (positions 1–5) provides DNA regulation annotations. Because structural candidates occupy the first five positions in the prompt, the LLM exhibits primacy bias and follows the structural channel's narrative, ignoring the correct contrastive retrieval at position 6. This highlights a limitation of the fixed 5+5 ordering.

### J.3. Query-Conditioning Demonstration

A central design feature of ProtQueSt is that the contrastive retriever conditions on the instruction, producing different retrievals for the same protein depending on the task. We verify this concretely using a 768-residue protein that appears in both the *domain/motif* and *general function* OOD test splits:

#### Protein sequence (768 residues)

```
MEQTFNPLQNKRSVVGQQVILECQLEGQDPDPVIKWLKDGQNVNCPDYQ
IFQEGNKFKLIIPSVQADAGRFDCAINAAGSKSSSCLIVAPPPSPGERQ
FMTAPSPRPPPTPVGPAAPYVVKELKHQMLKIGSSARFECRITAFPAEITW
LKNKGPIINTSKYKIDNDPDSGICSLTIAMMFAEDVGGYSCSARNAHQAV
TSAEILYKDKYNEWLREEQIKITQEKRRSMMEELDNAVQQPRKQKGTFTY
PHSQRLLEQLYTEEKVETDIKINESEASVENVPFQGVPPQVIRPLQPVSI
EGQKAELTCQIKGNPTPKVRWMKNGVPVQNSQLRQTSYNGAVASLIKITF
AEDAGMYTLVAENQFGRTNQSANIQLTQNSVNGVVRKKNNAVEQQKLSR
DLAVTPDLLGQGRQKPIFQQPLCDLQVAENQNVRFVDRISGRPYPIQWL
KNGVLLQHGHRKLLSSQNDLNTLIYMATVEDSGTYTCVATNESGQAQCS
CELTVKAHSQGSAAHFTEKFNSLIVHPGDSVELKCSAVGQPRPTYHWYKD
DEELIPGQCPYDIVNMPNGTRLRINNVRLDDSGCFQCAVAVMYGTAVHKA
PVKVQMKSSSQITPFRDSTSDQALIASATPELATIRHRQRAAGLSNAAWF
VEAASAEPPKIIGLSADHLSLLEGLAHVEVRFQPENDQNLKVVWFKDDL
PLEQKARLMFNIEAGRASFIDIVYQLSDGGGLYKLVVNVKQKAEATFTISV
TELPEV
```

**Structural channel (query-independent).** Because the structural retriever uses only protein embeddings, both queries return identical candidates: (1–2) 5'-flap endonuclease / exonuclease annotations (cos = 1.000, 0.989); (3–5) Ig-like / SH3 / Fibronectin type-III domain lists (0.984 each).

**Contrastive channel with domain/motif instruction.** Instruction: "Please examine the following protein sequence and predict any domains or motifs you can discern."

- **Contrastive top-5:** (1) Ig-like, SH3, Fibronectin type-III (0.684); (2) SH3, IQ, PH, Ig-like, DH, Fibronectin type-III, Protein kinase (0.680); (3) Ig-like, Fibronectin type-III, Protein kinase (0.675); (4) Protein kinase, Fibronectin type-III, Ig-like C2-type (0.668); (5) Fibrinogen C-terminal, Laminin G-like, EGF-like (0.654).
- All five contrastive candidates are **domain-level annotations**, directly matching the instruction type.
- **Gold:** Ig-like, Fibronectin type-III, Protein kinase. **Prediction:** Correct.

**Contrastive channel with general function instruction.** Instruction: "Could you evaluate the protein with this amino acid sequence and present a summary of its features?"

- **Contrastive top-5:** (1) "Controls actin polymerization and depolymerization" (0.685); (2) "Functions as a component of the nuclear pore complex (NPC)" (0.679); (3) "Plays an important role in regulation of muscle cell contractile activity" (0.675); (4) "Necessary for signaling by class 3 semaphorins and subsequent remodeling of the cytoskeleton" (0.675); (5) "Involved in cytoskeleton remodeling" (0.674).
- All five contrastive candidates are **functional descriptions**, matching the general-function instruction type. None overlaps with the domain-query retrievals above.

**Analysis.** The structural candidates are identical across both queries, confirming that the structural channel is query-independent. In contrast, the contrastive channel produces entirely disjoint candidate sets: domain annotations for the domain query and functional summaries for the general-function query. This demonstrates that the FiLM-conditioned contrastive model learns to route the same protein to different regions of the annotation embedding space depending on the instruction, validating the query-conditioning mechanism.

## K. Additional Analyses

### K.1. Structural Retriever $\alpha$ Ablation

The structural retriever uses a fixed mixing weight  $\alpha=0.7$  (ProstT5 weight) in the weighted cosine score  $s_{\text{struct}} = \alpha \cdot \text{cos}_{\text{ProstT5}} + (1-\alpha) \cdot \text{cos}_{\text{ESM-2}}$ . We sweep  $\alpha \in \{0.0, 0.2, 0.5, 0.7, 0.9, 1.0\}$  evaluated by Recall@10 on the test set to validate this choice. Table 26 reports per-task and macro-averaged results.

Table 26. Structural retriever Recall@10 as a function of the ProST5 mixing weight  $\alpha$ . The pool is the full cross-task train set ( $\sim 284\text{K}$  proteins); queries are test-split proteins. Bold indicates the best value per column.

$\alpha$	Cat. act.	Dom./mot.	Gen. func.	Prot. func.	Macro
0.0	0.1112	0.2042	0.0854	0.1795	0.1451
0.2	0.1389	0.2467	<b>0.0931</b>	<b>0.1919</b>	0.1677
0.5	0.1389	0.2518	0.0901	0.1914	0.1680
<b>0.7</b>	0.1394	<b>0.2548</b>	0.0905	0.1899	<b>0.1686</b>
0.9	0.1399	0.2544	0.0901	0.1874	0.1679
1.0	<b>0.1404</b>	0.2537	0.0901	0.1859	0.1675

The results confirm that  $\alpha=0.7$  achieves the highest macro Recall@10 (0.1686), though the landscape is flat for  $\alpha \geq 0.5$ , with all values in  $[0.5, 1.0]$  within 0.0011 of each other. ProST5 dominates the structural signal. Using ProST5 alone ( $\alpha=1.0$ ) drops macro recall by only 0.0011 relative to the best mixture. Conversely, ESM-2 alone ( $\alpha=0.0$ ) loses 0.0235, confirming that structure-derived embeddings (ProST5) carry the majority of the retrieval-relevant information but sequence embeddings (ESM-2) provide a complementary boost, particularly for domain/motif (+0.0506 from  $\alpha=0.0$  to  $\alpha=0.7$ ).

## K.2. FiLM Conditioning Analysis

To verify that FiLM genuinely adapts to the instruction embedding rather than learning a task-independent correction, we extract the learned scale and shift vectors for a representative instruction from each of the four tasks. Since both vectors are computed solely from the instruction embedding (not from the protein input), a given instruction produces exactly one scale vector and one shift vector in  $\mathbb{R}^{512}$ . Figure 6 reports the pairwise cosine similarity of these vectors across tasks. Scale cosines range from  $-0.28$  (domain/motif vs. general function) to  $+0.35$  (catalytic activity vs. domain/motif), with several pairs near zero or negative, showing that FiLM amplifies different embedding dimensions depending on the instruction. Shift vectors are more correlated across tasks (cosines 0.58–0.90), acting primarily as a shared offset rather than a task-specific routing mechanism.

## K.3. Confidence Intervals

We compute 95% bootstrap confidence intervals by resampling predictions 1,000 times (seed 42). Entity-BLEU is a corpus-level metric, so each resample recomputes corpus BLEU; Judge-F and METEOR are per-sample means. Recall@10 is deterministic and requires no CI. Tables 27 and 28 report the results.

## K.4. Computational Cost Summary

Protein embedding extraction dominates single-sample latency because both ESM-2 (650M parameters) and ProST5

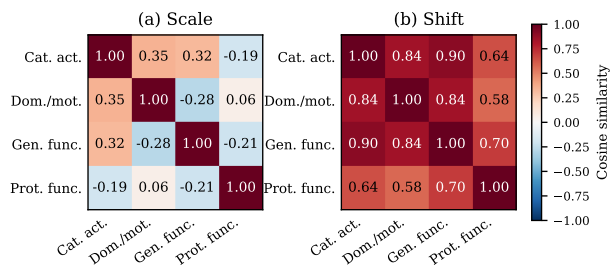


Figure 6. Cross-task cosine similarity of the learned FiLM vectors. (a) Scale vectors are weakly or negatively correlated across tasks, showing that FiLM amplifies different embedding dimensions for different instructions. (b) Shift vectors are highly correlated, acting as a largely shared offset.

Table 27. Bootstrap 95% confidence intervals for ProtQueSt. Entity-BLEU and METEOR: full 14,503-sample benchmark. Judge-F: 500 samples per task.

Task	E-BL 95% CI	METEOR 95% CI	Judge-F 95% CI
Cat. act.	[48.65, 53.28]	[52.44, 55.17]	[6.45, 6.91]
Dom./mot.	[51.71, 56.36]	[47.77, 50.72]	[6.33, 6.68]
Gen. func.	[31.31, 33.95]	[40.39, 41.84]	[5.46, 5.82]
Prot. func.	[58.51, 60.15]	[58.47, 60.00]	[6.62, 6.94]
Macro	[48.44, 50.16]	[50.23, 51.43]	[6.31, 6.49]

must process the full sequence. However, embeddings are cached for the entire corpus, so retrieval at inference requires only the query protein’s embeddings (computed once) plus fast Faiss lookup and MLP scoring. The LLM generation step is the primary cost driver for batch evaluation due to API rate limits and per-token pricing.

Table 28. Recall@10 hit counts for ProtQueSt (deterministic).

Task	Hits	$n$	Recall@10
Cat. act.	575	1,987	0.2894
Dom./mot.	1,136	2,732	0.4158
Gen. func.	513	4,297	0.1194
Prot. func.	1,343	5,487	0.2448
Total	3,567	14,503	0.2459

Table 29. Computational cost breakdown.

Component	Cost
<i>Training</i>	
GPU hardware	NVIDIA A6000
Time per model	~1.5–2.5 h
Total (21 variants)	~40–50 GPU-h
<i>Inference (per sample)</i>	
Protein emb. (ESM-2+ProtT5)	~0.5–2.0 s
Structural retrieval (Faiss)	<10 ms
Contrastive scoring (MLP)	<5 ms
LLM generation (Sonnet 4.6)	~2–5 s
<i>Storage</i>	
Protein emb. cache	~4 GB
Text emb. cache	~2 GB
Faiss HNSW index	~500 MB
Adapter weights	~27 MB

## L. Baseline Details

### L.1. Baseline Implementation Details

**LLM-only (3-shot).** Each LLM receives 3 in-context exemplars per task (same seed-42 selection as ProtQueSt) with protein sequences truncated to 80 characters. GPT-5.4 uses the OpenAI Responses API (GPT-5.4-2026-03-05, *reasoning\_effort=none*, *max\_output\_tokens=8192*, concurrency 8). Llama-3.1-8B-Instruct and Qwen3-8B use greedy decoding (*do\_sample=False*, seed 42). Claude Sonnet 4.6 uses temperature 1.0 (matching our generation setup). All LLM-only baselines are evaluated on the full 14,503-sample test set for lexical metrics; judge scores use 500 samples per task.

**Fine-tuned models.** Prot2Text-V2, BioT5+, InstructProtein, and Qwen3-8B-FT are fine-tuned on the OOD training split. We use published checkpoints where available; Qwen3-8B-FT is fine-tuned in-house. **BioT5+ caveat:** The BioT5+ checkpoint (QIZHIPEI/BIOT5-PLUS-BASE) was pretrained on Mol-Instructions, which overlaps with ProtInst-OOO annotations; for fair comparison, see RAPM Table 3 (Wu et al., 2025).

**Domain-specific pretrained.** Evolla-10B-DPO is evaluated zero-shot (3-shot prompting) without fine-tuning on our training data.

**Retrieval-based.** MMseqs2 retrieves the top-10 most similar training-set proteins by sequence identity and passes

their annotations to the generation LLM (same Sonnet 4.6 prompt). RAPM (Wu et al., 2025) uses MMseqs2 + ESM-2 dual-key retrieval with GPT-4.1 (temperature 0.7, top- $p$  0.9) or Sonnet 4.6 (temperature 0.7) as generators. Only-Structure uses our ProtT5 + ESM-2 weighted-cosine structural retriever ( $\alpha=0.7$ ) with 10 structural candidates (no contrastive channel).

### L.2. Per-Task Baseline Breakdown

Per-task Entity-BLEU and Judge-F for all baselines are reported in Appendix G, Tables 15 and 16.

### L.3. Fine-Tuning Hyperparameters

All four OOD-fine-tuned baselines (Qwen3-8B-FT, BioT5+, InstructProtein, Prot2Text-V2) are fine-tuned in-house on the ProtInst-OOO training split (284,526 examples). Common settings across all four runs: AdamW optimizer, learning rate  $1 \times 10^{-4}$ , cosine learning-rate schedule with warmup ratio 0.1, weight decay 0.01, bf16 mixed precision, and seed 42. Table 30 reports the per-baseline hyperparameters that vary across runs. Hardware:  $4 \times$  NVIDIA A6000 GPUs; training durations were approximately 15.7 h (Qwen3-8B-FT), 2.6 h (BioT5+), 8.9 h (InstructProtein), and 11.5 h (Prot2Text-V2).

**Generation configuration for fine-tuned baselines.** For all OOD-fine-tuned baselines at evaluation time, we use greedy decoding (*do\_sample=False*, seed 42) with each model’s published prompt format. Prot2Text-V2 uses its native sequence-only input; BioT5+ and InstructProtein follow their published instruction templates; Qwen3-8B-FT uses our standard 3-shot prompt template (Appendix D).

Table 30. Per-baseline fine-tuning hyper-parameters. Shared settings (AdamW, lr  $10^{-4}$ , cosine schedule, warmup 0.1, weight decay 0.01, bf16, seed 42) are listed in the §L.3 prose.

Baseline	Parameter	Value	
Qwen3-8B-FT	Base model	Qwen3-8B	
	Method	LoRA: $q, k, v, o$	
	LoRA config	$r=8, \alpha=32$ , dropout 0.05	
	Per-device batch	2	
	Grad accumulation	8	
	GPUs	4	
	Effective batch	64	
	Max seq. length	2,048	
	Epochs	2	
	Optimizer steps	8,892	
	Grad checkpointing	Yes	
	Distributed	DeepSpeed ZeRO-2	
	BioT5+	Base model	BIOT5-PLUS-BASE
Method		Full fine-tuning	
Per-device batch		4	
Grad accumulation		8	
GPUs		2	
Effective batch		64	
Max src. / tgt. length		2,048 / 512	
Epochs		2	
Optimizer steps		8,890	
Grad checkpointing		No	
Distributed		DeepSpeed ZeRO-2	
InstructProtein		Base model	InstructProtein
		Method	LoRA: $q, k, v, out$
	LoRA config	$r=8, \alpha=32$ , dropout 0.05	
	Per-device batch	2	
	Grad accumulation	8	
	GPUs	2	
	Effective batch	32	
	Max seq. length	2,048	
	Epochs	2	
	Optimizer steps	17,782	
	Grad checkpointing	Yes	
	Distributed	DeepSpeed ZeRO-2	
	Prot2Text-V2	Base model	Prot2Text-V2-11B
Architecture		ESM-2 3B + Llama-3.1-8B	
Method		LoRA: Llama (attn + MLP)	
LoRA config		$r=8, \alpha=32$ , dropout 0.05	
Per-device batch		1	
Grad accumulation		4	
GPUs		4	
Effective batch		16	
Max prot. / desc. length		1,021 / 512	
Epochs		1	
Optimizer steps		17,471	
Grad checkpointing		No	
Distributed		PyTorch DDP	

## M. Broader Impact and Limitations

### M.1. Limitations

- **Retrieval ceiling equals corpus ceiling.** ProtQueSt can only surface annotations present in the training corpus. Proteins with genuinely novel functions not yet characterized in any database fall outside the system’s reach. This constraint is shared by all current approaches, including generative models, since no system can predict functions for which no training signal exists.
- **Frozen encoder bottleneck.** FiLM can only re-weight features already encoded by frozen ESM-2 and ProtT5 embeddings. If a functional signal is absent from these representations, no amount of conditioning can surface it.
- **Single-vector protein representation.** CLS-token and mean-pooled embeddings collapse variable-length sequences into fixed-size vectors, discarding residue-level information. Functions governed by specific local motifs or binding sites may be underrepresented.
- **Benchmark scope.** All evaluations use the Prot-Inst-OOD benchmark; generalization to other protein function corpora and newly deposited proteins remains to be tested.

### M.2. Broader Impact

Automated protein function annotation can accelerate characterization of the rapidly growing “dark proteome,” the millions of sequenced proteins with no experimentally verified function. By providing retrieval-augmented functional descriptions grounded in curated databases, ProtQueSt could assist biocurators in prioritizing and drafting annotations for poorly characterized proteins.

However, automated annotation at scale carries the risk of error propagation: if the retrieval corpus contains incorrect annotations or the LLM generator introduces hallucinations, these errors could be amplified across downstream analyses. We emphasize that ProtQueSt is designed as a *recommendation engine* for expert biocurators, not as a replacement for experimental validation or expert review.

### M.3. Reproducibility Statement

We plan to release all source code for training, evaluation, and the retrieval pipeline, along with trained adapter weights (~6.8M parameters). We may additionally release pre-computed Faiss indices, cached protein embeddings, and archived text embeddings for exact reproducibility. The fully open-source configuration (BGE-M3 encoder + Llama-3.1-8B generator) achieves Judge-F 5.93 without any proprietary API dependencies.

1100  
1101 **System prompt:**

1102 You are an expert biocurator evaluating protein function predictions against curated UniProt ground truth. Apply the rubric strictly  
1103 and consistently. Biological knowledge matters—recognize synonyms, equivalent rephrasings, and EC-class relationships. Identical  
1104 inputs must yield identical scores. Scores are chosen from a fixed 5-point scale: {0, 3, 5, 7, 10}. Do not output other values for any  
1105 axis.

1106 **User prompt template** (placeholders {instruction}, {description}, {metadata}, {prediction} are filled per sample):

1107 Evaluate a PREDICTION against a curated GROUND TRUTH for a protein function prediction task.

1108 ---  
1109 **INPUT**

1110  
1111 **TASK INSTRUCTION:**

1112 {instruction}

1113 **GROUND TRUTH DESCRIPTION** (curated from UniProt):

1114 {description}

1115  
1116 **KEY BIO-ENTITIES** (atomic entities pre-extracted from UniProt annotations; top-level separators are “—” and “;” while “+”  
1117 and “=” within a single entity denote reaction components):

1118 {metadata}

1119 **PREDICTION:**

1120 {prediction}

1121 ---  
1122 **STEP 1: Parse reference entities.**

1123 Split KEY BIO-ENTITIES into atomic entities using “—” and “;” as top-level separators. Keep reaction components joined:  
1124 “(R)-prunasin + H<sub>2</sub>O = D-glucose + mandelonitrile” is one catalytic-reaction entity. For concept lists like “lipid binding” or “nucleus,  
1125 cytoplasm”, each comma-separated item is one entity unless the comma is inside a reaction.

1126 Let  $N_{\text{ref}}$  be the count of atomic reference entities.

1127  
1128 **STEP 2: Match reference entities—5 match types.**

1129 For each reference entity, determine whether it is expressed in PREDICTION using these match types, in priority order:

1130 **Type 1 Exact:** case-insensitive substring match of the entity or core term.

1131  
1132 **Type 2 Lexical variant:** morphological or formatting variant. For example, “mitochondrion”  $\equiv$  “mitochondrial”; “DNA-binding”  
1133  $\equiv$  “binds DNA”; “ATP hydrolysis”  $\equiv$  “hydrolyzes ATP”.

1134 **Type 3 Biochemical synonym:** different terms for the same biological entity. For example, “peptidase”  $\equiv$  “protease”  $\equiv$  “pro-  
1135 teinase”; “plasma membrane”  $\equiv$  “cell membrane”; “endoplasmic reticulum”  $\equiv$  “ER”. Do not count superclass-only  
1136 matches.

1137 **Type 4 Reaction equivalence:** catalytic reaction described with different syntax. For example, “(R)-prunasin + H<sub>2</sub>O = D-glucose +  
1138 mandelonitrile”  $\equiv$  “hydrolyzes prunasin to produce glucose and mandelonitrile”.

1139 **Type 5 Not a match:** superclass-only, same family but different member, opposite direction, or related but distinct biology.

1140 Let  $N_{\text{matched}}$  be the count of reference entities matched via Types 1–4. For each match, record the type as 1, 2, 3, or 4.

1141 **STEP 3: Count contradictions.**

1142 A contradiction is a prediction claim mutually exclusive with the reference, such as kinase vs. phosphatase, mitochondrion vs.  
1143 nucleus, catalytic reaction A vs. catalytic reaction B, or DNA-binding vs. RNA-binding only. Additional compatible or unrelated  
1144 predictions are not contradictions.

1145 Let  $N_{\text{contradicted}}$  be the count of distinct contradictions.  
1146  
1147  
1148  
1149  
1150  
1151  
1152  
1153  
1154

Figure 7. Prompt instructions for LLM-as-a-judge evaluation of protein function predictions.

1155  
1156 **Figure 7, continued.**

1157 **STEP 4: Score four axes.** Each axis score must be one of {0, 3, 5, 7, 10}.

- 1158  
1159 • **Axis 1—Recall:** fraction of reference entities recovered via Types 1–4. Let  $r = N_{\text{matched}}/N_{\text{ref}}$ .

1160 10 :  $r = 1.00$ ;

1161 7 :  $r \geq 0.60$ ;

1162 5 :  $r \geq 0.40$ ;

1163 3 :  $r \geq 0.15$ ;

1164 0 :  $r < 0.15$ .

- 1165  
1166 • **Axis 2 (Precision):** prediction stays on-topic to reference biology. Let  $U$  be the count of biological claims unsupported by the  
1167 reference or reasonable inference.

1168  
1169 10 :  $U = 0$ ;

1170 7 :  $U \leq 2$ ;

1171 5 :  $U \leq 5$ ;

1172 3 :  $U \leq 10$ ;

1173 0 :  $U > 10$  or prediction describes a different protein.

- 1174  
1175 • **Axis 3—Specificity:** granularity of biological content, independent of correctness. A prediction can be specifically wrong or  
1176 generally right.

1177  
1178 10 : concrete entities, substrates, products, EC numbers, domains,  
1179 localizations, named reactions, or specific residues;

1180 7 : specific biology one step coarser;

1181 5 : family or class level only;

1182 3 : very general biology;

1183 0 : generic platitudes with no specific biological content.

- 1184  
1185 • **Axis 4—Plausibility:** free of contradictions and fabrications.

1186 10 :  $N_{\text{contradicted}} = 0$  and all claims are grounded;

1187 7 :  $N_{\text{contradicted}} = 0$  with some speculation;

1188 5 :  $N_{\text{contradicted}} = 1$ ;

1189 3 :  $N_{\text{contradicted}} = 2$ ;

1190 0 :  $N_{\text{contradicted}} \geq 3$  or describes a different protein.

1191  
1192 **STEP 5: Final score.**

1193  
1194  
1195 
$$\text{FINAL\_SCORE} = \text{round} \left( \frac{\text{Axis1} + \text{Axis2} + \text{Axis3} + \text{Axis4}}{N_{\text{active axes}}} \right),$$

1196 where  $N_{\text{active axes}}$  excludes any axis scored  $-1$ .

1197  
1198 **OUTPUT FORMAT—return exactly these 8 lines, no preamble, no commentary:**

1199  
1200 N\_ref: <integer>

1201 N\_matched: <integer>

1202 Match\_types: <comma-separated type integers per matched entity>

1203 N\_contradicted: <integer>

1204 N\_unsupported: <integer U from Axis 2>

1205 Axis\_scores: <recall>,<precision>,<specificity>,<plausibility>

1206 Critique: <one sentence: key mismatch, synonym match, or strength>

1207 Final\_score: <integer 0--10>

1208  
1209