

Distilling Protein Language Models with Complementary Regularizers

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Abstract

Large autoregressive protein language models (700M+ parameters) generate novel sequences *de novo*, yet their computational demands limit throughput in biopharma workflows such as antibody affinity maturation and enzyme engineering, where millions of candidates must be evaluated. We present a distillation framework combining temperature-scaled knowledge distillation with two protein-specific enhancements: uncertainty-aware position weighting, which emphasizes biologically variable regions via teacher entropy, and calibration-aware label smoothing. A central finding is *complementary regularizers*: each enhancement individually degrades distillation quality, yet their combination yields 53% perplexity improvement over baseline distillation—a result we explain through information-theoretic analysis of noise amplification and signal filtering. We train three student models at 3.8–20× compression, all outperforming baselines (31–87% perplexity improvement). The resulting models achieve 2.4–5.3× speedup while preserving amino acid distributions consistent with natural proteins (KL divergence < 0.015), enabling on-premise screening of proprietary sequences without cloud API dependencies—a key requirement in pharmaceutical settings. When fine-tuned on domain-specific protein families (50–1,000 sequences), distilled students outperform the teacher on conotoxin perplexity at every training set size and generate lysozyme sequences with 94% Pfam family match rate versus the teacher’s 69%—demonstrating that smaller distilled models are superior starting points for domain adaptation on scarce data.

1 Introduction

Protein language models (pLMs) trained on evolutionary sequence data now enable computational protein design [1–3]. By learning the statistical patterns of natural protein sequences, autoregressive pLMs can generate novel sequences *de novo* with properties resembling those found in nature [1]. Among these, ProtGPT2—a GPT-2 architecture model [4] with 738 million parameters trained on UniRef50 [5]—has demonstrated the ability to produce sequences with natural amino acid distributions, plausible secondary structure content, and globular characteristics.

However, the computational cost of large pLMs creates a significant barrier to practical deployment. ProtGPT2 requires high-end GPUs for inference, generates sequences at limited throughput (~3 seconds per sequence), and cannot be deployed on edge devices or in resource-constrained laboratory settings. These constraints are particularly acute in biopharma applications such as ML-guided antibody affinity maturation [6] and protein engineering campaigns [3], where evaluating thousands to millions of candidate sequences demands both high throughput and cost-effective deployment. Beyond inference efficiency, a critical question for such applications is whether distilled models can serve as effective starting points for domain-specific fine-tuning on proprietary datasets—typically comprising only 50–1,000 sequences from a target protein family.

Knowledge distillation [7] offers a principled approach to model compression by training a smaller student model to mimic the probability distributions of a larger teacher model. The key

insight of Hinton et al. is that temperature-softened output distributions encode rich inter-class relationships—“dark knowledge”—that one-hot labels cannot convey. For protein sequences, these soft distributions capture amino acid substitution patterns: the teacher’s prediction that position t should be leucine, with isoleucine and valine as secondary preferences, conveys evolutionary constraints that a hard label alone cannot express.

In natural language processing, distillation has produced compact yet performant models such as DistilBERT [8] and TinyBERT [9], motivating similar approaches for protein language models. DistilProtBert [10] compressed ProtBert using response-based distillation, and MTDP [11] introduced multi-teacher distillation for protein representations. For causal protein LMs, Dubey et al. [12] applied distillation to a domain-specific model trained on 572 spider silk sequences. However, *no systematic study has addressed distillation for general-purpose autoregressive protein language models*—despite these being the models required for open-ended *de novo* sequence design.

We address this gap with a distillation framework that combines standard Hinton-style knowledge distillation with two protein-specific enhancements: (1) *uncertainty-aware position weighting*, which uses teacher entropy to emphasize biologically variable regions during distillation, and (2) *calibration-aware label smoothing*, which applies confidence-dependent smoothing to teacher distributions to improve student calibration [13, 14]. Our central finding is a *complementary regularizers*: uncertainty weighting alone increases perplexity by 95% and calibration smoothing alone increases it by 109%, yet their combination improves perplexity by 53% over baseline distillation. We provide a mechanistic explanation grounded in information theory: smoothing acts as a noise filter on teacher distributions, while weighting amplifies the cleaned signal at biologically important positions.

Our contributions are as follows:

1. The first systematic study of knowledge distillation for general-purpose autoregressive protein language models.
2. Two protein-specific distillation enhancements: uncertainty-aware position weighting and calibration-aware label smoothing.
3. Discovery and mechanistic explanation of complementary regularizers, where individually harmful modifications combine for substantial improvement.
4. A comprehensive evaluation framework spanning perplexity, calibration (ECE), amino acid distributional fidelity, and inference benchmarks.
5. Open-source compressed models at three scales (37M, 78M, 194M parameters) available on HuggingFace.
6. Demonstration that distilled students are superior fine-tuning starting points on scarce protein family data, achieving higher sample efficiency and better family-specific generation than the teacher.

2 Results

2.1 Ablation reveals complementary regularizers

To assess the contribution of each enhancement, we conducted a 2×2 ablation study using a Micro architecture (4 layers, 4 heads, 256 embedding dimensions; ~ 16 M parameters, $\sim 46 \times$ compression), toggling uncertainty-aware position weighting and calibration-aware label smoothing independently

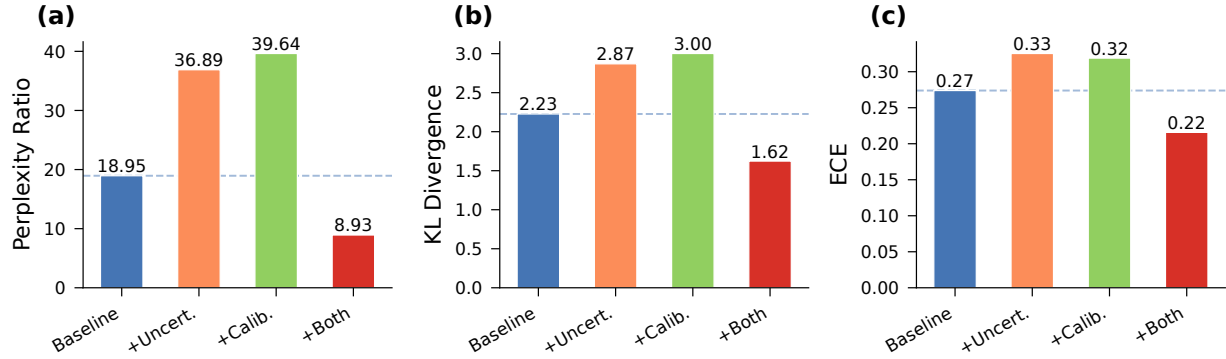


Figure 1: Ablation study showing the complementary-regularizer effect. Each enhancement individually degrades distillation quality (higher perplexity, higher KL divergence, higher ECE), but their combination yields a 53% perplexity improvement over baseline.

Table 1: Ablation study results on Micro architecture (4L/4H/256E, ~16M parameters). Each enhancement individually degrades distillation quality, but their combination yields a 53% improvement over baseline. PPL: perplexity; KL: KL divergence from teacher; ECE: expected calibration error.

Configuration	Uncertainty	Calibration	PPL	KL Div	ECE	vs. Baseline
Baseline (standard KD)	✗	✗	18.95	2.23	0.274	—
+Uncertainty only	✓	✗	36.89	2.87	0.325	+95%
+Calibration only	✗	✓	39.64	3.00	0.319	+109%
+Both (synergy)	✓	✓	8.93	1.62	0.216	-53%

(Table 1). This smaller architecture was chosen to reduce the computational cost of running all four configurations under identical training hyperparameters ($T = 2.0$, $\alpha = 0.5$, learning rate = 10^{-3} , 3 epochs).

The baseline (standard Hinton-style distillation) achieved a perplexity of 18.95. Applying uncertainty weighting alone degraded perplexity to 36.89 (+95%), while calibration smoothing alone degraded it further to 39.64 (+109%). Both individual enhancements also increased KL divergence from the teacher and worsened expected calibration error (ECE). Taken individually, neither enhancement appears beneficial.

However, when both enhancements are applied simultaneously, perplexity drops to 8.93—a 53% improvement over baseline and a 75–77% improvement over either individual enhancement. KL divergence decreases from 2.23 to 1.62, and ECE improves from 0.274 to 0.216. This complementary-regularizer effect, where two individually harmful modifications combine to produce substantial improvement, is the central finding of this work (Fig. 1).

2.2 Scaling across model sizes

To test whether the complementary-regularizer effect generalizes beyond the Micro ablation architecture, we trained paired baseline and synergy models at three larger scales: Tiny (4L/4H/512E, 20× compression), Small (9.4×), and Medium (3.8×). Note that the Tiny model here uses a 512-dimensional embedding, distinct from the 256-dimensional Micro model used for the ablation. Based on the ablation finding that synergy training requires careful learning rate selection, we

Table 2: Scaling results across three model sizes. Synergy models use both uncertainty weighting and calibration smoothing with adjusted learning rates and warmup. All synergy models outperform their respective baselines.

Scale	Method	Compression	PPL	ECE	Improvement
Tiny (512E)	Baseline	20×	39.91	0.345	—
	Synergy	20×	5.06	0.183	87%
Small (768E)	Baseline	9.4×	15.19	0.235	—
	Synergy	9.4×	7.05	0.259	54%
Medium (1024E)	Baseline	3.8×	3.72	0.169	—
	Synergy	3.8×	2.58	0.135	31%

adopted a protocol using approximately half the baseline learning rate with 500 steps of linear warmup (see Methods for details).

Table 2 and Fig. 2 show that synergy models outperform baselines at all three scales. The improvement is largest at the highest compression ratio (87% at 20× compression for Tiny) and decreases with scale (54% for Small, 31% for Medium). This trend is expected: as student capacity approaches teacher capacity, the marginal benefit of enhanced distillation diminishes because standard KD already transfers knowledge effectively.

2.3 Calibration analysis

Expected calibration error (ECE) measures the alignment between predicted confidence and empirical accuracy across binned probability intervals [13, 15]. We computed ECE with 10-bin quantization on held-out protein sequences (Fig. 3).

Synergy models improve calibration at the Tiny scale (ECE 0.183 vs. 0.345, a 47% reduction) and at the Medium scale (ECE 0.135 vs. 0.169, a 20% reduction). At the Small scale, however, the synergy model shows a minor ECE regression (0.259 vs. 0.235). This anomaly likely reflects the fact that the Small model was the only scale where no learning rate reduction was applied; the warmup schedule alone may not fully optimize calibration. Overall, synergy distillation improves student calibration at 2 of 3 scales, with the largest gains at higher compression ratios where miscalibration risk is greatest.

2.4 Biological validity

For protein language models intended for *de novo* sequence design, preserving biologically realistic amino acid usage is essential. We evaluated the amino acid frequency distributions of generated sequences against the natural distribution observed in UniProt [5] (Fig. 4).

All student models—both baseline and synergy—produce amino acid distributions closely matching the natural UniProt distribution, with KL divergence below 0.015 in all cases. Quantifying per-residue deviations via mean absolute deviation (MAD) from UniProt frequencies reveals a clear ordering: the teacher deviates least (MAD = 0.0034), followed by the synergy student (MAD = 0.0073), then the baseline student (MAD = 0.0089). The synergy model is closer to natural on 13 of 20 amino acids, indicating that the combined enhancements improve distributional fidelity relative to standard distillation.



Figure 2: Perplexity ratio (student/teacher) across model scales. Synergy models outperform baselines at all three compression ratios, with the largest improvement at the highest compression ($20\times$).

2.5 Compression-quality tradeoff

Plotting perplexity against compression ratio for all models reveals a Pareto frontier (Fig. 5). Synergy models dominate baseline models at every compression level tested, offering strictly better perplexity at the same model size. The improvement is most pronounced at high compression ratios, suggesting that complementary-regularizer distillation is especially valuable when aggressive compression is required.

2.6 Practical deployment

Inference benchmarks on an NVIDIA L40S GPU show that student models achieve substantial speedups over the teacher (Fig. 6): $5.3\times$ for Tiny, $4.1\times$ for Small, and $2.4\times$ for Medium models. These speedups, combined with dramatically reduced memory requirements (Fig. 8), enable protein sequence generation on consumer-grade GPUs. Peak GPU memory drops from 3.2 GB for the teacher to just 170 MB for the Tiny model—a $19\times$ reduction. At the Tiny model’s throughput of 111 sequences per minute, screening a library of 10^6 candidate sequences—typical of combinatorial antibody or enzyme engineering campaigns—requires approximately 6 GPU-hours on a single consumer-grade device, compared to ~ 48 GPU-hours for the full teacher. The Tiny model’s $\sim 37M$ parameters fit within 170 MB of GPU memory, permitting deployment on shared laboratory workstations without dedicated accelerator infrastructure.

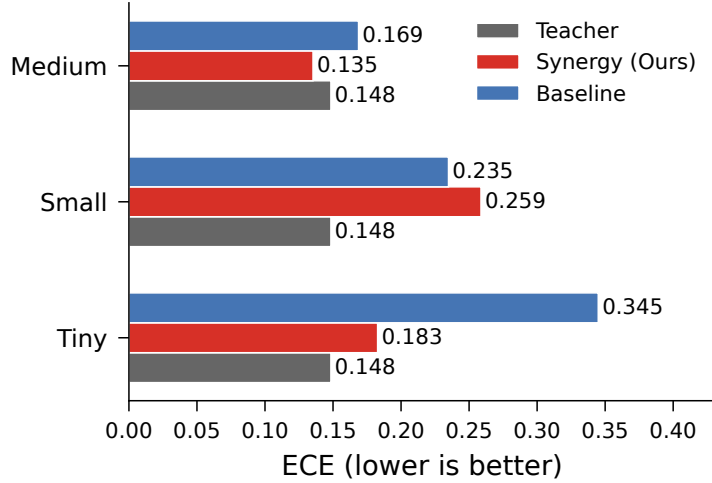


Figure 3: Expected calibration error across model scales. Synergy (red) achieves lower ECE than Baseline (blue) at Medium and Tiny scales, with the largest improvement at Tiny (0.183 vs. 0.345, 47% reduction). At Small scale, Synergy shows a minor regression (0.259 vs. 0.235). The teacher ECE (0.148) is shown for reference.

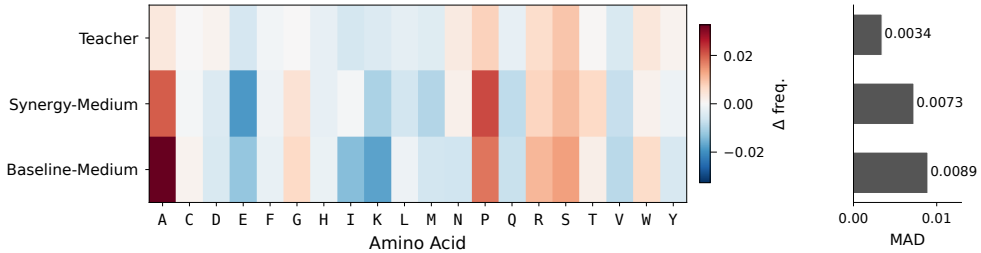


Figure 4: Amino acid frequency deviation from the natural UniProt distribution (left) with mean absolute deviation summary (right). The synergy student ($\text{MAD} = 0.0073$) deviates 18% less than the baseline ($\text{MAD} = 0.0089$), confirming that the combined enhancements improve distributional fidelity.

2.7 Structural quality

To assess whether distilled models generate structurally plausible proteins, we scored 50 sequences per model using ESMFold predicted local distance difference test (pLDDT) [16] (Fig. 7). The teacher achieved a mean pLDDT of 51.2 with 16% of sequences exceeding the confident prediction threshold of 70. Student models scored lower (mean pLDDT 38–40), consistent with the reduced capacity of smaller architectures. Importantly, synergy and baseline models at the same scale produce comparable pLDDT distributions (synergy-medium: 38.1 vs. baseline-medium: 38.1), confirming that complementary-regularizer distillation does not sacrifice structural quality for improved perplexity. The pLDDT gap between teacher and students reflects model capacity rather than a training methodology deficit.

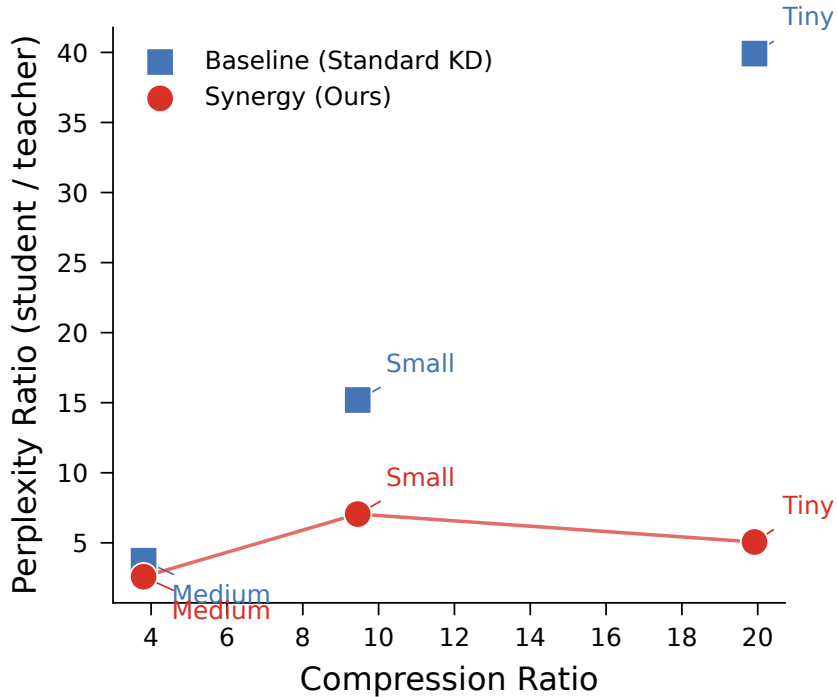


Figure 5: Compression–quality Pareto frontier. Synergy models (filled) dominate baseline models (open) at every compression ratio, achieving strictly better perplexity at the same model size.

2.8 Domain-specific fine-tuning

A critical question for biopharma applications is whether distilled models can serve as effective starting points for fine-tuning on proprietary datasets. We fine-tuned all five models—Teacher, Medium, Small, Tiny (synergy), and Baseline-Tiny (standard KD)—on conotoxin (PF02950, median 68 AA) and lysozyme (PF00959, median 170 AA) at five training set sizes ($N \in \{50, 100, 200, 500, 1000\}$), totaling 75 runs. We evaluated test perplexity and HMMER hit rate (sequences matching the family Pfam profile at $E < 10^{-5}$). AMP results, where the teacher dominates and no HMMER profile is available, are reported in the Appendix (Table 8).

On conotoxin, all distilled students achieve lower test perplexity than the teacher at every training set size (Fig. 9a). The gap is largest at small N : at $N = 50$, the teacher reaches a perplexity of 1,659 while the Medium student achieves 372 ($4.5\times$ lower). Medium at $N = 50$ outperforms the teacher at $N = 100$ (perplexity 1,153), demonstrating $2\times$ sample efficiency. The advantage persists at large N : at $N = 1,000$, Medium achieves a perplexity of 30 versus the teacher’s 54.

On lysozyme, the teacher achieves lower test perplexity at every N (Fig. 9b). Yet students generate sequences that more frequently match the PF00959 HMM profile (Fig. 10a). At $N = 1,000$, Small achieves a 94% HMMER hit rate versus the teacher’s 69% (+25 percentage points). At $N = 200$, the gap is even wider: Small 73% versus teacher 28%. This decoupling between perplexity and family-specific generation quality suggests that distilled representations capture family-level structural patterns more effectively during fine-tuning.

On conotoxin HMMER hit rate, Medium dominates: 42.5% at $N = 1,000$ versus the teacher’s 8.0% (Fig. 10b). Hit rates are lower overall for conotoxin, reflecting the short peptide length

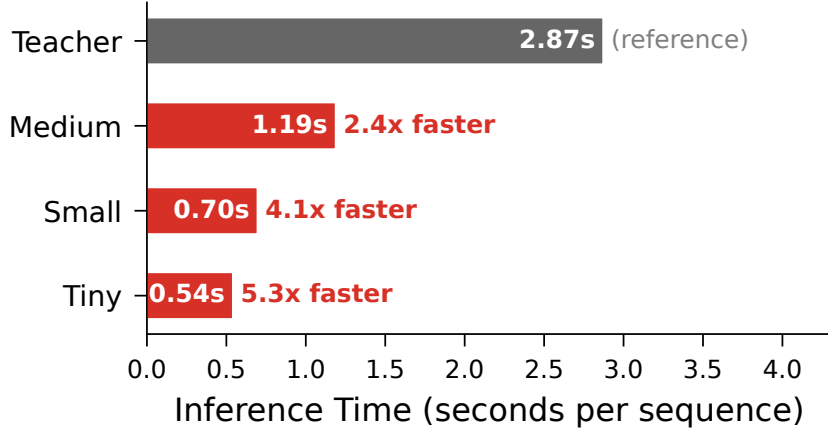


Figure 6: Inference speedup on an NVIDIA L40S GPU. Student models achieve 2.4–5.3× speedup over the ProtGPT2 teacher, enabling deployment on consumer-grade hardware.

Table 3: Synergy-Tiny versus Baseline-Tiny at $N = 1,000$: same 37M architecture, different distillation method. Synergy-Tiny outperforms the teacher on conotoxin PPL and lysozyme HMMER, while Baseline-Tiny performs near teacher level, confirming that the synergy distillation procedure drives the fine-tuning advantage.

Model	Conotoxin		Lysozyme	
	PPL	HMMER (%)	PPL	HMMER (%)
Teacher (738M)	54	8.0	187	69.0
Synergy-Tiny (37M)	40	8.0	327	84.0
Baseline-Tiny (37M)	52	13.0	569	71.0

(median 68 AA) relative to the generation length (200 tokens); Medium nonetheless leads on both perplexity and HMMER for this family.

The fine-tuning advantage is not limited to one model size: all three synergy students (Medium, Small, Tiny) outperform the teacher on conotoxin perplexity and lysozyme HMMER hit rate at every N tested. At $N = 1,000$, conotoxin perplexity is 30 (Medium), 39 (Small), and 40 (Tiny), all below the teacher’s 54; Baseline-Tiny (52) barely edges out the teacher (Table 3). On lysozyme HMMER at $N = 1,000$, Small reaches 94%, Tiny 84%, and Medium 83.5%, all far exceeding the teacher’s 69%; Baseline-Tiny (71%) performs at teacher level. The controlled comparison—Synergy-Tiny versus Baseline-Tiny (same 37M architecture, different distillation method)—isolates the source of the advantage: Baseline-Tiny performs near teacher level while Synergy-Tiny far exceeds both, with 15 out of 15 perplexity wins across all families. This confirms that the synergy distillation procedure, not just model compression, drives the fine-tuning advantage.

Students fine-tune 20–162× faster than the teacher in wall-clock time. The Tiny model completes AMP fine-tuning ($N = 1,000$) in 25 seconds versus 66 minutes for the teacher. No gradient checkpointing is required for any student, making fine-tuning feasible on consumer GPUs.

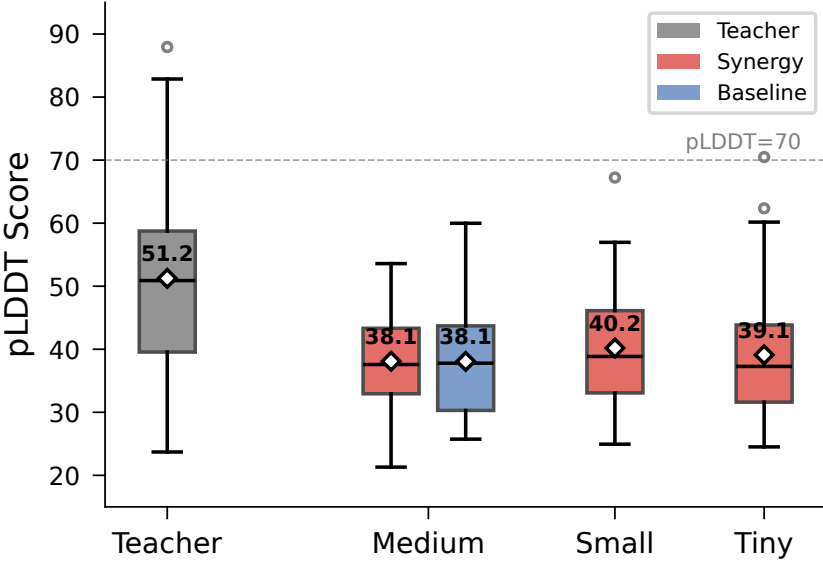


Figure 7: ESMFold pLDDT scores for 50 generated sequences per model. The teacher achieves higher structural confidence (mean 51.2) due to greater capacity. At the Medium scale, synergy and baseline produce indistinguishable pLDDT distributions (both 38.1), confirming that complementary-regularizer distillation preserves structural quality—the capacity gap is a compression effect, not a method deficit.

3 Discussion

Mechanistic explanation of complementary regularizers. The central finding of this work—that individually harmful modifications combine to produce substantial improvement—has a mechanistic explanation grounded in information theory. Consider the teacher’s probability distribution at each sequence position as containing both *signal* (genuine amino acid preferences reflecting protein biology) and *noise* (miscalibration artifacts from the teacher’s own training).

Uncertainty-aware position weighting increases the loss contribution at high-entropy positions, effectively amplifying both signal and noise. Because noise dominates at high-entropy positions (where the teacher is uncertain and potentially miscalibrated), the net effect of weighting alone is to amplify noise more than signal, degrading distillation quality.

Calibration-aware label smoothing acts as a low-pass filter on teacher distributions, blending predictions toward the uniform distribution in proportion to teacher uncertainty. Applied alone, this attenuates both signal and noise, but signal is disproportionately affected because the fine-grained probability structure at uncertain positions—which encodes biologically meaningful substitution preferences—is smoothed away.

When both enhancements operate simultaneously, they address each other’s failure mode. Calibration smoothing removes the noise that uncertainty weighting would otherwise amplify, while uncertainty weighting compensates for the signal attenuation introduced by smoothing by directing additional learning capacity toward the affected positions. The combined effect is *amplified but regularized* attention to variable positions: the student is instructed to “pay extra attention here” (weighting) while matching a denoised target (smoothing). This is analogous to a standard signal

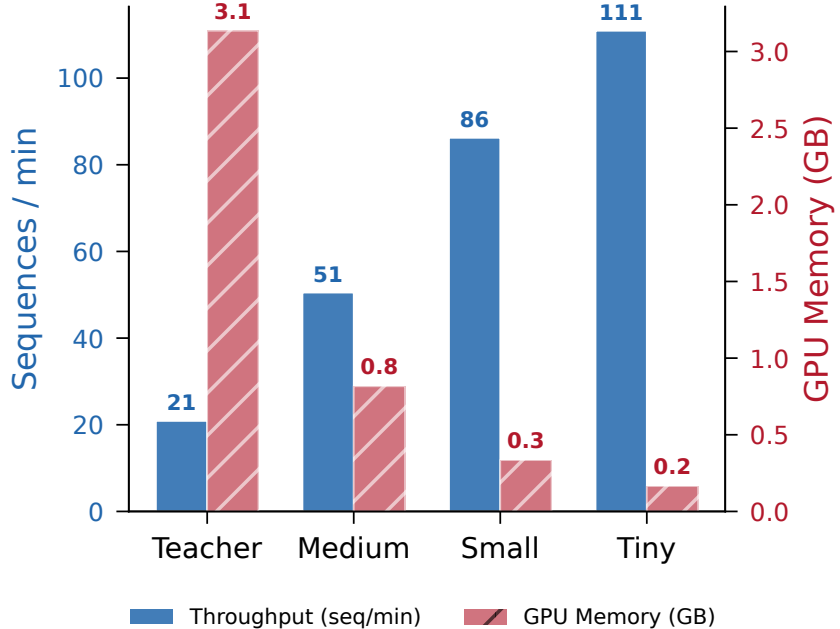


Figure 8: Generation throughput and GPU memory usage on an NVIDIA L40S. The Tiny model generates 111 sequences per minute while requiring only 170 MB GPU memory ($19\times$ reduction from the teacher’s 3.2 GB), enabling large-scale screening on consumer hardware.

processing pipeline where amplification followed by filtering improves reception quality, whereas either operation alone degrades the signal-to-noise ratio.

Formally, the two enhancements modify different components of the per-position loss: weighting acts as an outer multiplier on the loss magnitude, while smoothing modifies the inner KL divergence target distribution. Because they operate on orthogonal aspects of the loss, their effects compose multiplicatively rather than additively, enabling synergistic interaction when the modifications address complementary failure modes.

Training dynamics and the role of warmup. Analysis of training logs reveals that the first \sim 500 steps constitute a critical window for synergy training (Fig. 11). Without warmup, the modified objective—which is inherently easier to minimize from random initialization due to smoothed targets—allows the student to rapidly converge toward a degenerate minimum that achieves low training loss but poor generalization. This is evidenced by anomalously low initial loss values (6.62 vs. 7.94 for baseline at the Tiny scale) followed by severe train–evaluation misalignment.

With linear warmup over 500 steps, the near-zero initial learning rate forces the student to make incremental updates, learning basic token frequency patterns before encountering the full modified objective. By the time the learning rate reaches its target value, the student has already formed preliminary representations that constrain it to a generalizable region of the loss landscape. The enhanced objective then *refines* this foundation rather than corrupting it.

Scale-dependent effects. The synergy improvement decreases with model scale ($87\% \rightarrow 54\% \rightarrow 31\%$), which we attribute to three factors. First, larger students have less to gain from regularization because they already approach teacher capacity ($3.8\times$ compression for Medium vs. $20\times$ for Tiny).

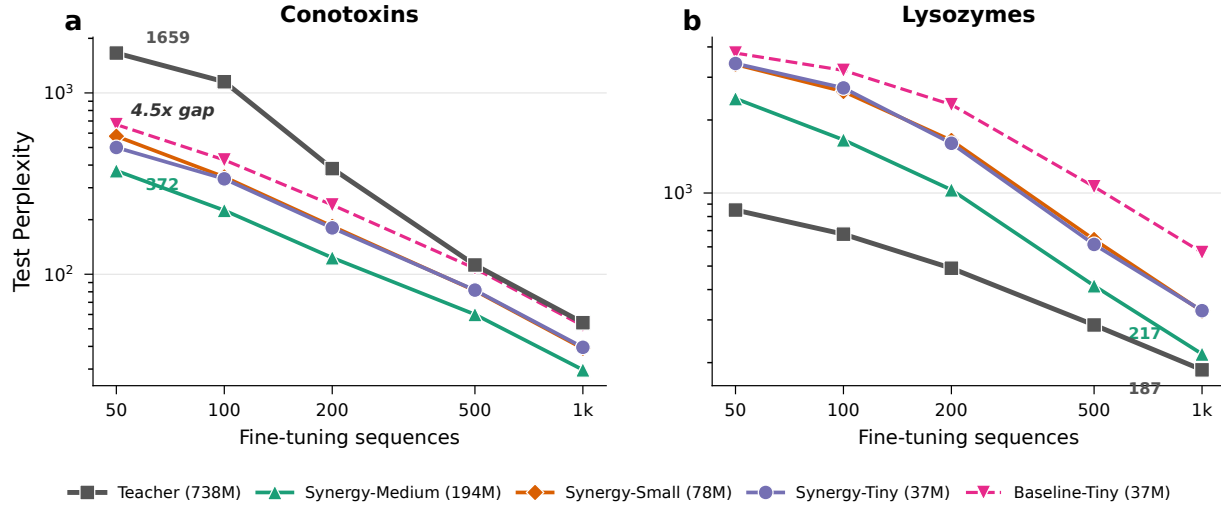


Figure 9: Test perplexity versus training set size for domain-specific fine-tuning. (a) Conotoxin: all distilled students achieve lower perplexity than the teacher at every N , with Medium at $N = 50$ outperforming the teacher at $N = 100$ ($2\times$ sample efficiency). (b) Lysozyme: the teacher achieves lower perplexity, but students generate more family-matching sequences (see Fig. 10).

Second, baseline distillation improves approximately exponentially with scale, narrowing the gap. Third, larger students can better model the true distribution at variable positions natively, reducing the marginal benefit of the noise-filtering mechanism.

PPL-vs-hit-rate decoupling. On lysozyme, the teacher achieves lower test perplexity than all students at every training set size, yet students generate sequences that more frequently match the PF00959 HMM profile (94% for Small versus 69% for the teacher at $N = 1,000$). We interpret this as evidence that distilled representations capture family-level structural patterns more effectively during fine-tuning: the compressed parameter space imposes an inductive bias that prevents overfitting to non-family sequence patterns, channeling limited capacity toward family-specific features. This challenges the assumption that larger models are always superior fine-tuning starting points. Importantly, the advantage holds across all three synergy scales (Medium, Small, Tiny), not just the smallest model—and the Baseline-Tiny control confirms it is the distillation method, not compression ratio alone, that enables superior domain adaptation.

Learning rate scaling. A practical finding is that synergy training requires approximately half the baseline learning rate at matching scales, plus warmup. The smoothed targets create a loss landscape where the same nominal learning rate produces effectively larger functional steps; halving the learning rate compensates for this effect. This $0.5\times$ scaling rule held at the Tiny and Medium scales. At the Small scale, where the baseline learning rate (5×10^{-4}) happened to already be appropriate, no reduction was needed.

Small model ECE regression. The Small synergy model shows a minor ECE regression (0.259 vs. 0.235 for baseline), the only scale where calibration worsened. This model was the only one trained without an explicit learning rate reduction, suggesting that the warmup schedule alone is

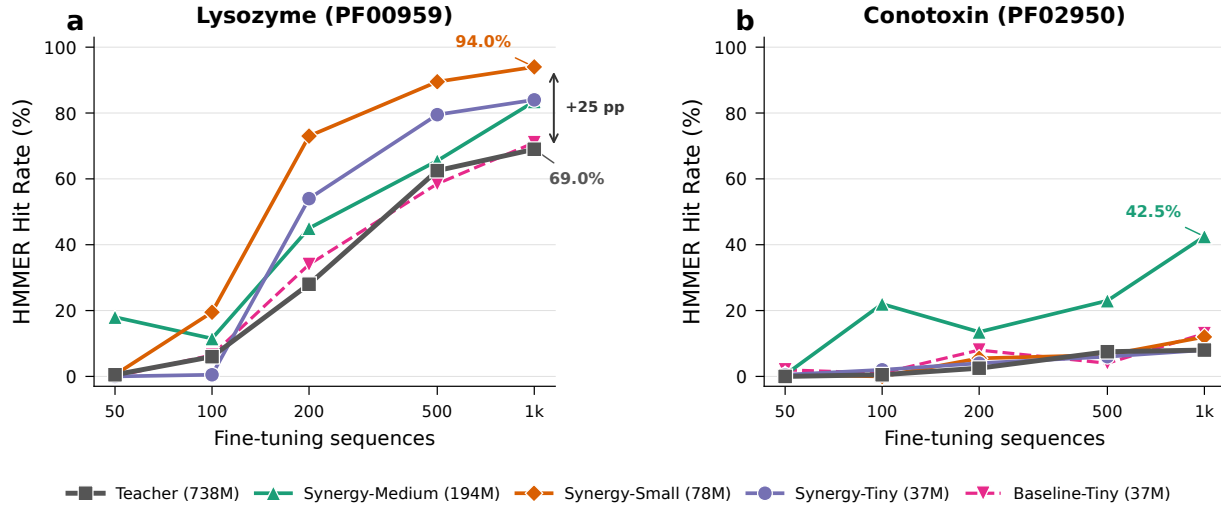


Figure 10: HMMER hit rate versus training set size. (a) Lysozyme: Small achieves 94% hit rate at $N = 1,000$ versus the teacher’s 69%, despite having higher perplexity. (b) Conotoxin: Medium reaches 42.5% versus the teacher’s 8.0%; lower absolute rates reflect the generation length mismatch relative to the short peptide family.

not universally sufficient to optimize calibration. A targeted learning rate sweep at this scale would likely resolve the regression.

Practical implications for biopharma. The compressed models address concrete deployment bottlenecks in protein engineering pipelines. In antibody discovery, where language-model-guided affinity maturation has shown promise [6], the $5.3\times$ speedup of the Tiny model reduces the cost of scoring large variant libraries by an order of magnitude, making iterative design-build-test cycles feasible on commodity hardware. For enzyme engineering campaigns that screen combinatorial sequence libraries [3], on-premise deployment of compact models avoids transmitting proprietary sequences to cloud APIs, satisfying data-governance requirements common in pharmaceutical settings. More broadly, the 170 MB memory footprint of the Tiny model enables integration into automated laboratory workflows where GPU resources are shared across instruments. The fine-tuning results further strengthen this case: distilled students not only generate faster at inference time but also serve as superior starting points for domain adaptation, achieving better family-specific generation with fewer training sequences and $20\text{--}162\times$ faster fine-tuning wall-clock time.

Limitations. This work has several limitations. First, we evaluate a single teacher model (ProtGPT2); generalization to other protein LMs such as ProGen [3] or non-protein causal LMs remains to be established. Second, ECE is computed at the token level and may not fully capture sequence-level calibration relevant to downstream applications. Third, structural plausibility is assessed via predicted metrics (pLDDT from ESMFold [16]) rather than experimental validation. Fourth, all experiments use a fixed smoothing factor ($\lambda = 0.1$) and temperature ($T = 2.0$); a joint hyperparameter search over the enhanced distillation objective could yield further improvements. Fifth, the fine-tuning evaluation used a single learning rate per model size without grid search, and the generation length (`max_length=200` tokens) produces sequences 2–15× longer than the training families, which depresses HMMER hit rates and makes the reported student advantage a conser-

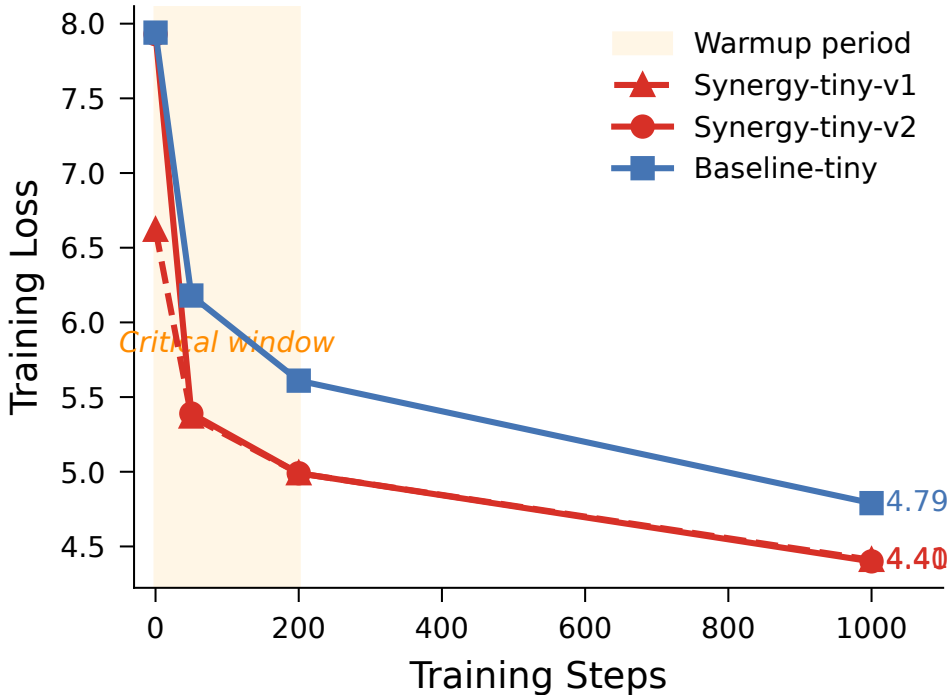


Figure 11: Training loss dynamics during the first 500 steps. Without warmup, the synergy objective allows rapid convergence to a degenerate minimum. Linear warmup over 500 steps constrains early optimization, enabling the student to reach a generalizable region of the loss landscape.

vative estimate.

Future directions. We identify three extensions. Multi-teacher distillation, combining signals from diverse protein LMs, could provide more robust soft targets. The complementary-regularizer approach should be tested on other autoregressive protein LMs and on non-protein biological sequence models. Finally, experimental validation of generated sequences—via wet-lab synthesis and characterization—would provide the strongest evidence of preserved biological function. For fine-tuning, length-constrained generation with family-aware stopping criteria would likely improve HMMER hit rates further, and evaluation on additional protein families would test the generality of the fine-tuning advantage.

4 Methods

4.1 Standard distillation framework

We adopt the response-based knowledge distillation framework of Hinton et al. [7]—as opposed to feature-based [17] or relational [18] approaches—adapted for autoregressive protein language modeling. Given a protein sequence $x = (x_1, \dots, x_n)$ over vocabulary \mathcal{V} , the teacher and student models produce logit vectors $z_t^T, z_t^S \in \mathbb{R}^{|\mathcal{V}|}$ at each position t .

Temperature-scaled softmax. To reveal inter-class relationships in the teacher’s predictions, logits are softened with temperature $\tau > 1$:

$$p_i^{(\tau)} = \frac{\exp(z_i/\tau)}{\sum_j \exp(z_j/\tau)} \quad (1)$$

Higher temperatures produce smoother distributions that expose the relative preferences among amino acids [7].

Soft loss. The soft loss measures the Kullback–Leibler divergence between temperature-scaled teacher and student distributions, averaged over sequence positions:

$$\mathcal{L}_{\text{soft}} = \frac{1}{n-1} \sum_{t=1}^{n-1} D_{\text{KL}}\left(p_T^{(\tau)}(\cdot|x_{\leq t}) \parallel p_S^{(\tau)}(\cdot|x_{\leq t})\right) \quad (2)$$

Hard loss. The hard loss is the standard cross-entropy on ground-truth next-token labels:

$$\mathcal{L}_{\text{hard}} = - \sum_{t=1}^{n-1} \log p_S(x_{t+1}|x_{\leq t}) \quad (3)$$

Combined loss. The total distillation loss balances hard and soft objectives with coefficient $\alpha \in [0, 1]$, applying a τ^2 correction to maintain gradient magnitude under temperature scaling:

$$\mathcal{L} = \alpha \cdot \mathcal{L}_{\text{hard}} + (1 - \alpha) \cdot \tau^2 \cdot \mathcal{L}_{\text{soft}} \quad (4)$$

The τ^2 factor compensates for the $1/\tau^2$ gradient attenuation introduced by temperature scaling in the softmax [7].

4.2 Uncertainty-aware position weighting

Protein sequences exhibit heterogeneous predictability: conserved structural positions (e.g., hydrophobic core residues) are highly predictable, while variable positions (loops, linkers, surface residues) admit multiple plausible amino acids. We exploit this structure by weighting each position’s contribution to the soft loss in proportion to the teacher’s prediction entropy.

Shannon entropy. At each position t , the teacher’s uncertainty is quantified as:

$$u_t = H(p_T(\cdot|x_{<t})) = - \sum_{v \in \mathcal{V}} p_T(v) \log p_T(v) \quad (5)$$

where p_T uses temperature $\tau = 1$ (unscaled) to reflect the teacher’s true predictive uncertainty.

Position weights. Entropies are min-max normalized per sequence and mapped to the range $[0.5, 1.0]$:

$$w_t = 0.5 + 0.5 \cdot \frac{u_t - \min(\mathbf{u})}{\max(\mathbf{u}) - \min(\mathbf{u})} \quad (6)$$

The floor of 0.5 ensures that even highly predictable positions contribute to the distillation loss, preventing the student from ignoring conserved regions entirely.

Weighted soft loss. The uncertainty-weighted soft loss replaces the uniform average in Eq. 2:

$$\mathcal{L}_{\text{soft}}^{\text{weighted}} = \frac{1}{|\mathcal{T}|} \sum_{t \in \mathcal{T}} w_t \cdot D_{\text{KL}}\left(p_T^{(\tau)}(\cdot|x_{\leq t}) \parallel p_S^{(\tau)}(\cdot|x_{\leq t})\right) \quad (7)$$

where \mathcal{T} denotes the set of non-padded positions.

4.3 Calibration-aware distillation

Well-calibrated confidence estimates are critical for protein engineering applications where model predictions guide experimental prioritization [13]. Neural networks, including large language models, tend to be poorly calibrated, and this miscalibration can be transferred during distillation. We introduce dynamic label smoothing [14] applied to teacher distributions, with smoothing intensity inversely proportional to teacher confidence.

Dynamic smoothing. At each position t , the smoothing intensity is computed on the temperature-scaled teacher distribution—the same distribution used as the KL divergence target:

$$\epsilon_t = \lambda \cdot \left(1 - \max_{v \in \mathcal{V}} p_T^{(\tau)}(v|x_{<t})\right) \quad (8)$$

where λ is a base smoothing factor. Because temperature scaling ($\tau > 1$) flattens the teacher distribution and reduces peak probabilities, the effective smoothing is amplified relative to what unscaled ($\tau = 1$) confidence would produce. This is by design: the softened distribution better exposes positions where the teacher’s distributional uncertainty warrants regularization. When the teacher is confident ($\max_v p_T^{(\tau)}(v)$ remains high despite softening), smoothing is minimal ($\epsilon_t \approx 0$). When the teacher is uncertain, smoothing increases, regularizing the distribution. Note that this differs from the entropy computation in uncertainty-aware weighting (Eq. 5), which uses unscaled ($\tau = 1$) probabilities to reflect the teacher’s true predictive uncertainty; the smoothing intensity instead operates in the temperature-scaled space where the KL target is defined, ensuring that the degree of regularization is calibrated to the actual target distribution the student must match.

Smoothed targets. The smoothed teacher distribution blends the temperature-scaled prediction with a uniform distribution:

$$\bar{p}_T^{(\tau)}(v) = (1 - \epsilon_t) \cdot p_T^{(\tau)}(v) + \frac{\epsilon_t}{|\mathcal{V}|} \quad (9)$$

Expected calibration error. We evaluate calibration using ECE with $B = 10$ equal-width bins [15]:

$$\text{ECE} = \sum_{b=1}^B \frac{|B_b|}{N} |\text{acc}(B_b) - \text{conf}(B_b)| \quad (10)$$

where B_b is the set of predictions falling in bin b , and $\text{acc}(B_b)$ and $\text{conf}(B_b)$ are the average accuracy and confidence within the bin, respectively.

4.4 Model architectures

All models use the GPT-2 architecture [4, 19] with varying depth and width. The teacher is ProtGPT2 (738M parameters) [1]. Student architectures span a $20\times$ compression range (Table 4).

Table 4: Model architectures and compression ratios. All models use the GPT-2 architecture with the ProtGPT2 tokenizer ($|\mathcal{V}| = 50,257$). [†]Micro is used only for the ablation study; scaling experiments use Tiny–Medium.

Model	Layers	Heads	Embedding dim	Parameters	Compression
Teacher (ProtGPT2)	36	20	1280	738M	$1\times$
Medium	12	16	1024	$\sim 194\text{M}$	$3.8\times$
Small	6	8	768	$\sim 78\text{M}$	$9.4\times$
Tiny	4	4	512	$\sim 37\text{M}$	$20\times$
Micro [†]	4	4	256	$\sim 16\text{M}$	$\sim 46\times$

4.5 Training details

Data. We use a 10% subset of UniProt [5] protein sequences stored in Parquet format. Sequences are tokenized using the ProtGPT2 tokenizer with a maximum length of 1024 tokens.

Optimization. All models are trained with the AdamW optimizer for 3 epochs. Following common practice in knowledge distillation, we set temperature $\tau = 2.0$ and balancing coefficient $\alpha = 0.5$. For calibration smoothing, the base smoothing factor is $\lambda = 0.1$.

Baseline models use learning rates of 10^{-3} (Tiny and Small) and 10^{-4} (Medium) without warmup. Synergy models use approximately half the baseline learning rate with 500 steps of linear warmup: 5×10^{-4} (Tiny and Small) and 5×10^{-5} (Medium). This learning rate reduction compensates for the smoother loss landscape created by label smoothing, which causes the same nominal learning rate to produce effectively larger optimization steps.

Hardware. The Medium model was trained on an NVIDIA L40S GPU (48 GB). Smaller models were trained on various NVIDIA GPUs with at least 24 GB of memory. Gradient accumulation (4 steps) was used to achieve an effective batch size of 32.

4.6 Fine-tuning evaluation

To assess whether distilled models serve as effective starting points for domain adaptation, we fine-tuned all five models on two protein families: conotoxin (PF02950, median 68 AA) and lysozyme (PF00959, median 170 AA), at five training subset sizes ($N \in \{50, 100, 200, 500, 1000\}$). All models were trained with AdamW, cosine learning rate schedule, early stopping (patience 3), effective batch size 8, and FP16 mixed precision; learning rates were scaled by model size (Table 5). For each fine-tuned model, we generated 200 sequences and evaluated them with HMMER [20] against family Pfam HMM profiles ($E < 10^{-5}$). The teacher required gradient checkpointing on a 24 GB L4 GPU; students did not. Full hyperparameter, dataset, and generation configuration details are provided in Tables 5–7.

4.7 Data and code availability

Training code, evaluation scripts, and trained model weights are available at <https://github.com/ewijaya/protein-lm-distill>. Compressed models are hosted on HuggingFace under `littleworth/protgpt2-distilled-tiny`, `-small`, and `-medium`. Training data were derived from UniProt [5], which is freely available at <https://www.uniprot.org>.

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A Fine-tuning experimental details

This appendix provides full experimental details for the domain-specific fine-tuning evaluation described in Section 2.8.

Table 5: Fine-tuning hyperparameters for each model. All models use the same optimizer, scheduler, and early stopping configuration; only learning rate, batch size, and gradient checkpointing differ across scales.

Parameter	Teacher	Medium	Small	Tiny	Baseline-Tiny
Parameters	738M	194M	78M	37M	37M
Learning rate	2e-5	5e-5	1e-4	2e-4	2e-4
Batch size	2	8	8	8	8
Grad accumulation	4	1	1	1	1
Effective batch	8	8	8	8	8
Grad checkpointing	Yes	No	No	No	No
Max epochs	20	20	20	20	20
Early stopping	3	3	3	3	3
Optimizer	AdamW	AdamW	AdamW	AdamW	AdamW
Scheduler	Cosine	Cosine	Cosine	Cosine	Cosine
Warmup steps	100	100	100	100	100
Precision	FP16	FP16	FP16	FP16	FP16

Table 6: Dataset summary for fine-tuning evaluation. Training subsets of size $N \in \{50, 100, 200, 500, 1000\}$ were sampled from the full training set. Validation and test sets were held constant across all runs.

Family	Train (full)	Val	Test	Median length (AA)	Pfam ID
Conotoxin	6,164	770	770	68	PF02950
Lysozyme	10,740	1,342	1,342	170	PF00959
AMP	2,501	313	313	29	None

Table 7: Generation and evaluation configuration. All models used identical sampling parameters. HMMER evaluation was performed against family-specific Pfam HMM profiles where available.

Parameter	Value
Sequences per run	200
Max length	200 tokens
Top- k	950
Temperature	1.0
Repetition penalty	1.2
HMMER E -value	$< 10^{-5}$

Table 8: AMP fine-tuning results. The teacher dominates on perplexity at all training set sizes, consistent with the absence of a family-specific HMM profile for evaluation. AMP is a functionally diverse class rather than a single sequence family, making perplexity an incomplete measure of domain adaptation. AA KL divergence values show students converge to competitive amino acid composition by $N = 1,000$.

AMP — Test Perplexity					
N	Teacher	Medium	Small	Tiny	Baseline-Tiny
50	2,576	3,003	3,005	3,090	3,169
100	2,060	2,895	2,845	2,886	2,997
200	1,664	2,611	2,524	2,610	2,787
500	1,036	2,073	2,293	2,354	2,534
1000	829	1,375	1,764	1,782	2,048
AMP — Amino Acid KL Divergence					
N	Teacher	Medium	Small	Tiny	Baseline-Tiny
50	0.021	0.196	0.122	0.132	0.158
100	0.047	0.211	0.126	0.178	0.181
200	0.074	0.215	0.144	0.103	0.142
500	0.081	0.146	0.139	0.157	0.158
1000	0.068	0.055	0.048	0.037	0.061