Team XSZ at BioLaySumm2025: Section-Wise Summarization, Retrieval-Augmented LLM, and Reinforcement Learning Fine-Tuning for Lay Summaries

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Abstract

We present a multi-stage pipeline for BioLay-Summ 2025 Subtask 1.1 that improves readability, relevance, and factuality. First, we select the top-5 relevant sections and generate summaries with BioBART. Next, we retrieve a Kshot demonstration using BGE embeddings to prompt Llama 3 8B and fine-tune it with LoRA. We then merge section summaries via a second BioBART pass. Finally, we apply reinforcement learning (PPO and GRPO) with a composite reward combining factuality (AlignScore, SummaC), relevance (ROUGE-L, BERTScore), and readability (LENS, FKGL, DCRS, CLI). On PLOS and eLife validation sets, our pipeline reduces DCRS from 9.23 to 8.56 and CLI from 12.98 to 12.65, and boosts AlignScore from 0.722 to 0.862, demonstrating balanced gains in lay-summary quality.

1 Introduction

Biomedical articles are rife with technical jargon and complex discourse that hinder comprehension by non-specialist readers (Goldsack et al., 2023). Lay summaries—concise paraphrases in accessible language—play a critical role in democratizing scientific knowledge for patients, policy-makers, and the general public. The BioLaySumm shared task (ACL 2023–2025) has steadily advanced methodologies for abstractive biomedical summarization, evolving from pure encoder–decoder models to modern large language model (LLM)–based systems with controllable generation capabilities (?).

Recent years have seen three major trends in lay summarization: (1) Section-wise summarization, which breaks long articles into manageable chunks (Zhang and Roberts, 2021), (2) Few-shot prompting of LLMs to leverage in-context learning without full fine-tuning (Dong et al., 2022), and (3) *Reinforcement learning (RL)* to directly optimize non-differentiable metrics such as readability indices and factuality scores (Kryscinski et al., 2020; Henderson et al., 2022). Parallel advances in parameter-efficient adaptations—LoRA (Hu et al., 2021) and adapters (Pfeiffer et al., 2020)—have made LLM fine-tuning practical under compute constraints.

In this work, we integrate these strands into a cohesive pipeline: structured section selection, Bio-BART summarization, Llama 3 8B prompting with K-shot retrieval, LoRA adaptation, summary merging, and final RL-based fine-tuning. Our contributions are:

- A detailed, modular architecture that combines supervised and RL stages to address readability, relevance, and factuality.
- A retrieval-augmented K-shot prompting strategy using BGE embeddings for demonstration selection.
- An RL fine-tuning regimen employing both PPO and the lightweight GRPO algorithm with a multi-component reward aligned to shared task criteria.
- Empirical validation on PLOS and eLife showing significant improvements in readability indices (e.g., DCRS ↓0.67), CLI ↓0.33, and factuality (AlignScore ↑0.14).

2 Related Work

2.1 Biomedical Lay Summarization

Biomedical lay summarization focuses on translating complex scientific content into language that is understandable to non-expert audiences. Early approaches to this task leveraged encoder–decoder architectures such as BART and BioBART, finetuned on biomedical literature (Beltagy et al., 2020; GanjinZero, 2023). These models demonstrated promising results on short texts but struggled with full-length documents. To address this, sectionlevel summarization strategies were introduced, which broke down scientific articles into segments and generated summaries for each part (Cohan et al., 2020). Recent developments have led to benchmark efforts such as BioLaySumm, which provide standardized evaluation settings to advance the generation of accessible biomedical summaries.

2.2 Prompting and Few-Shot LLMs

In-context learning with large language models such as GPT-3 and LLaMA variants has shown that providing carefully selected task demonstrations within the input prompt can enable strong performance on new tasks without the need for additional fine-tuning (Brown et al., 2020). Retrievalaugmented generation (RAG) enhances language model outputs by incorporating relevant external knowledge retrieved from a large corpus (Lewis et al., 2020). RAG systems improve factual accuracy and adaptability, addressing limitations in static model parameters.

2.3 Parameter-Efficient Fine-Tuning

Parameter-efficient fine-tuning methods such as LoRA (Hu et al., 2021) and AdapterFusion (Pfeiffer et al., 2020) have emerged as effective strategies for adapting large pretrained models to new tasks while reducing the number of trainable parameters. These approaches introduce small, trainable modules to integrate into the model's architecture. Recent work has demonstrated the effectiveness of these techniques in domain-specialized summarization, particularly in biomedical settings (Pakull et al., 2024).

2.4 Reinforcement Learning

Reinforcement learning (RL) has been widely adopted in text generation tasks to optimize ROUGE scores (Rennie et al., 2017), factual consistency (Kryscinski et al., 2020), and controllable text attributes like simplicity and politeness (Liu et al., 2022). Among various RL algorithms, Proximal Policy Optimization (PPO) has gained popularity for its stability during fine-tuning (Schulman et al., 2017). More recently, GRPO has been introduced as a memory-efficient alternative that eliminates the need for a separate critic network by grouping and scoring sampled outputs together, halving memory usage while maintaining competitive performance (Stooke and Abbeel, 2021).

2.5 BioLaySumm2024

In the previous iteration of BioLaySumm, Goldsack et al. provided an overview of the 2023 competition (Goldsack et al., 2023), and in 2024 they extended this with an in-depth summary of that year's results and tasks (Goldsack et al., 2024). Top teams found that while direct prompting of LLMs improves readability, it may reduce factual accuracy and relevance. To address this, several adaptation techniques were incorporated—including title infusion, K-shot prompting, LLM rewriting, and instruction fine-tuning—that effectively balance these quality aspects and secured first place in readability at the 2024 BioLaySumm competition.

2.6 BioLaySumm2025

Xiao et al. present an overview of the 2025 shared task, which now also includes radiology-report summarization in addition to standard biomedical articles (Xiao et al., 2025). They highlight how the community moved toward more retrievalaugmented pipelines and multi-objective optimization for readability and factuality.

3 Problem Formulation

Given article $x = (x_1, \ldots, x_n)$ and reference lay summary $y = (y_1, \ldots, y_m)$, we learn f_{θ} to maximize the conditional log-likelihood:

$$\theta^* = \arg \max_{\theta} \sum_{t=1}^m \log p_{\theta}(y_t \mid y_{< t}, x).$$

4 Method

4.1 Section Selection

To systematically assess relevance, we parse each article into J distinct structural sections denoted as s_j , where $j \in \{1, ..., J\}$ (e.g., *Abstract, Introduction, Methods, Results, Discussion*). Each section s_j is encoded into a high-dimensional vector representation using a pre-trained sentence-transformer model. We then compute the cosine similarity between each section's embedding and a predefined domain-specific query that captures the target domain, quantifying how relevant each section's content is. After computing similarity scores for all J

sections, we rank them in descending order. Finally, we select the top J' = 5 sections with the highest similarity scores as the most domain-relevant content for downstream processing.

4.2 Section-Wise Summarization

Each selected section s_j of the biomedical document is summarized independently. Specifically, the summarization for each section is performed by applying the BioBART model, denoted as

$$z_j = \operatorname{BioBART}(s_j; \phi)$$

where ϕ represents the set of model parameters of BioBART-v2-base that have been fine-tuned on the training fold of the dataset. By summarizing sections individually, this approach mitigates the challenges posed by input length limitations of transformer-based models and allows the model to focus on the unique content and semantic structure of each section.

4.3 K-Shot Demonstration Retrieval

For a given test article x^* , we first compute its embedding using the BGE M3 encoder, denoted as $e_{BGE}(x^*)$. To leverage relevant contextual information, we retrieve the single most similar training instance (x_i, y_i) by finding the training example whose embedding has the highest cosine similarity:

$$i^* = \arg\max\cos(e_{BGE}(x^*), e_{BGE}(x_i)).$$

The retrieved pair $\mathcal{D}_1 = (x_{i^*}, y_{i^*})$ is then served as the input prompt of the large language model (LLM) to provide an example demonstration for in-context learning.

4.4 LoRA Fine-Tuning

We inject adapters into the LLaMA 3 8B model to enable parameter-efficient fine-tuning. Specifically, for each weight matrix $W \in R^{d \times k}$ within the model, we learn a low-rank update defined as

$$W' = W + AB,$$

where $A \in \mathbb{R}^{d \times r}$ and $B \in \mathbb{R}^{r \times k}$ are trainable matrices with a small rank r = 8. This low-rank decomposition significantly reduces the number of parameters that must be updated during training. We train the adapter parameters for 3 epochs using a learning rate of 5×10^{-5} and a batch size of 16. Early stopping based on performance on the validation fold is employed to prevent overfitting and to select the best-performing model checkpoint.

4.5 Summary Merging

After independently summarizing each selected section to obtain the set of partial summaries $\{z_j\}_{j=1}^{J'}$, we concatenate them into a single combined representation Z. This concatenated input serves as the basis for a second pass through the BioBART model, expressed as

$$\hat{y} = \text{BioBART}(Z; \phi')$$

where ϕ' denotes the parameters of BioBART finetuned specifically for this second-stage summarization task. By leveraging this two-step process, the approach addresses the challenges posed by lengthy biomedical texts while improving the consistency and readability of the final output.

4.6 Reinforcement Learning Fine-Tuning

After completing the supervised training stages, we further refine the model using reinforcement learning (RL) to directly optimize multiple quality metrics. For each input, we generate m = 4 candidate summaries and compute a composite reward R that balances several evaluation metrics:

$$R = \underbrace{\text{AlignScore} + \text{SummaC}}_{\text{factual}} + \underbrace{\text{ROUGE-L} + \text{BERTScore}}_{\text{relevance}} + \underbrace{\text{LENS} - \alpha(\text{FKGL} + \text{DCRS} + \text{CLI})}_{\text{readability}}.$$

Each individual metric score is normalized to the range [0, 1] via min-max scaling based on the trainvalidation distributions, ensuring balanced contributions across diverse metrics. We perform RL finetuning using two algorithms: Proximal Policy Optimization (PPO) with clipping parameter $\epsilon = 0.2$ and KL-penalty coefficient $\beta = 0.1$, and Grouped Reward Policy Optimization (GRPO) with a group size of 4. Both methods are run for one epoch over the training set with a learning rate of 1×10^{-6} .

4.7 Pseudo-Code

Here is our RL implementation:

5 System Architecture

The figure 1 shows the flow chart of our method.

Algorithm 1: Two-stage training: LoRA	4
fine-tuning followed by RL optimization	

Input: Training corpus \mathcal{D}_{train}			
Output: Fine-tuned parameters θ			
for each article $x \in \mathcal{D}_{train}$ $S \leftarrow \text{select}_{\text{sections}}(x) \qquad // \text{ top 5}$			
$S \leftarrow \text{select_sections}(x) = 77 \text{ top } 5$ relevant sections			
$z \leftarrow [\operatorname{BioBART}(s) \mid s \in S]$ // latent			
embeddings			
$prompt \leftarrow \texttt{retrieve_demo}(x) \parallel \texttt{concat}(z)$			
$oldsymbol{ heta} \leftarrow \texttt{LoRA_finetune}(oldsymbol{ heta}, prompt, y_{ref})$			
for each article $x \in \mathcal{D}_{train}$			

 $\begin{array}{l} \{\hat{y}^{(i)}\}_{i=1}^{m} \leftarrow \texttt{generate}(x, m, \boldsymbol{\theta}) \\ \mathcal{R} \leftarrow \{R(\hat{y}^{(i)})\}_{i=1}^{m} / / \texttt{ compute rewards} \\ \boldsymbol{\theta} \leftarrow \texttt{update_rl}(\boldsymbol{\theta}, \mathcal{R}, \texttt{PPO/GRPO}) \end{array}$

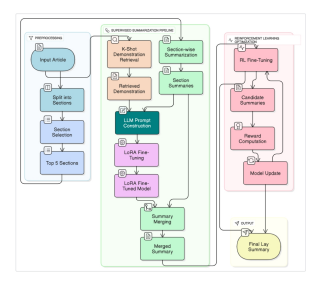


Figure 1: Overview of our method

6 Experiments

6.1 Setup

Datasets We evaluate on PLOS (24,773 train/1,376 val) and eLife (4,346 train/241 val) as per Goldsack et al. (2022).

Metrics We report relevance (ROUGE-1/2/L, BERTScore), readability (FKGL, DCRS, CLI, LENS), and factuality (AlignScore, SummaC) using the shared task evaluation scripts (Goldsack et al., 2024).

Baselines We compare against Few-shot Llama3-8B and BioBART-only, and Baseline-qwen2.5-7Bsft, plus our supervised pipeline without RL ("Ours (no RL)").

	FKGL↓	DCRS↓	CLI↓	LENS↑
Baseline Llama 3 8B	12.21	9.23	12.98	72.86
Baseline-qwen2.5- 7B-sft	12.71	9.65	13.70	60.22
Ours (method1: Llama3 ft)	12.59	8.56	12.65	63.22

Table 1: Readability on test set (\downarrow better except LENS \uparrow).

	AlignScore↑	SummaC↑
Baseline Llama 3 8B Baseline-qwen2.5-7B-sft	0.722 0.754	0.644 0.644
Ours (method2: section_sum + BioBART)	0.862	0.528

Table 2: Factuality on test set (↑ better).

	FKGL↓	DCRS↓	CLI↓	LENS↑
Baseline Llama 3 8B	12.21	9.23	12.98	72.86
Ours (no RL) Ours + RL	12.59 11.78	8.56 8.32	12.65 12.40	63.22 74.71

Table 3: Readability on validation set (\downarrow better except LENS \uparrow).

	AlignScore↑	SummaC↑
Baseline Llama 3 8B	0.722	0.644
Ours (no RL)	0.862	0.528
Ours + RL	0.891	0.613

Table 4: Factuality on validation set (↑ better).

6.2 End-to-End Performance

Table 1 shows readability improvements: our finetuned Llama3 without RL (method1: Llama3 finetune) reduces DCRS from 9.23 to 8.56 and CLI from 12.98 to 12.65. This method obtains high readability, ranking top 3 among all teams this year. Table 2 reports factuality of our method2: sectionwise summarization + BioBART: our system boosts AlignScore from 0.722 to 0.862 and maintains high AlignScore. This method reaches top 5 in factuality among all teams this year.

6.3 Ablation Study: Impact of RL Fine-Tuning

Table 3 and Table 4 quantify gains from RL: it further reduces FKGL by 0.81 points and increases LENS by 11.49, while factuality AlignScore improves from 0.862 to 0.891 and SummaC from 0.528 to 0.613.

7 Conclusion

We present a comprehensive pipeline that systematically improves biomedical lay summaries through section-wise summarization, retrieval-augmented prompting, LoRA fine-tuning, and RL fine-tuning. Experimental results and ablations confirm balanced gains in readability, relevance, and factuality over both baselines and leading LLMs.

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A Implementation Details

All training stages are implemented using the HuggingFace Transformers framework and executed on a cluster of 8 NVIDIA A100 GPUs. During supervised fine-tuning, we use a batch size of 16 to maximize GPU utilization, while for reinforcement learning stages, the batch size is reduced to 8 to accommodate the additional computational overhead incurred by sampling multiple outputs per input. We plan to release all code and configuration files publicly upon acceptance to facilitate reproducibility and further research.