Chan Zuckerberg Biohub Assignment Report

Angad Sandhu

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Abstract

This report presents a comprehensive proposal and detailed implementation of a state-of-the-art multimodal deep learning system for biomedical research. The proposed system is designed to integrate image data, transcriptomic profiles, and textual information from scientific literature to predict the downstream effects of gene knock-outs on cell state. In Part 1, I outline the architecture and training strategy for a conversational tool that accepts text prompts, images, or transcriptomic data. Part 2 details the development of a multimodal learning system that integrates Geneformer single-cell embeddings and cell type label textual embeddings to predict donor identifiers with an adversarial loss to mitigate the confounding effects of sex. The report includes methodological descriptions, modular training components, comprehensive visualizations (UMAP, training curves, and attention heatmaps), ablation studies, and a complete directory and code structure overview. This work builds on recent advances such as ChatNT [1], LLaVA-Med [2], and Geneformer [3], while also integrating tools like BioBERT [5] and Pub-MedGPT [6].

1 Introduction

Recent advances in deep learning and transformer architectures have spurred significant progress in developing models for biomedical applications. In this report, I propose and implement a multimodal system that integrates diverse data modalities; including imaging, transcriptomics, and textual data, to predict the effects of gene knock-outs on cell state. This system is motivated by the need for robust, context-aware models in the biological sciences, with potential applications in precision medicine and cellular biology.

The work is divided into two parts:

- Part 1: Model Proposal: I propose an innovative multimodal architecture that leverages domain-specific pretrained models and integrates image, omics, and text modalities into a unified conversational tool.
- Part 2: Coding Challenge: I detail the development and training of a multimodal learning system that integrates textual embeddings from cell type labels and Geneformer single-cell embeddings to predict donor identifiers. The model employs adversarial training to mitigate sex-based biases.

2 Part 1: Model Proposal

2.1 Overview

In this proposal, I present a detailed plan for a deep learning system that enables cell biologists to understand the downstream effects of gene knock-outs. The system is designed to process multi-

ple input modalities—text, images, and transcriptomic data—and fuse their representations using transformer-based architectures. The ultimate goal is to produce contextually rich and interpretable outputs that inform on gene function and cellular state.

2.2 Additional Data Requirements

A robust multimodal system requires carefully curated and diverse datasets, along with rigorous preprocessing and augmentation strategies. The following data types are essential:

2.2.1 Image Data

- Sources: Biomedical images including histopathology slides, cellular microscopy, and radiology scans (e.g., data from The Cancer Genome Atlas (TCGA)).
- **Preprocessing:** Involves artifact removal, stain normalization, resizing, and patch extraction to focus on regions of interest.
- Augmentation: Techniques such as rotations, flips, and color jittering, as well as specialized augmentations (e.g., stain normalization for pathology images) to increase model robustness.

2.2.2 Transcriptomic Data

- Sources: Bulk RNA-seq and single-cell transcriptomic datasets from public repositories (e.g., GEO, Human Cell Atlas) representing both healthy and diseased states.
- **Preprocessing:** Normalization methods (log-transform or rank normalization) and feature selection (e.g., highly variable genes). For sequence data, tokenization or the use of k-mer embeddings is considered.
- Foundation Model: Utilization of pretrained models (e.g., *Geneformer*) to extract high-level gene co-expression features.
- Augmentation: Simulation of noise or batch effects to improve model generalization.

2.2.3 Textual Data

- **Sources:** Curated biomedical literature from repositories like PubMed Central, including figure captions and gene/pathway databases.
- **Preprocessing:** Cleaning of textual data, sentence splitting, and named entity recognition for gene and disease annotation.
- Augmentation: Generation of paraphrased data or synthetic question-answer pairs using large language models.
- **Integration:** Linking gene identifiers with external databases (e.g., Reactome, Gene Ontology) to enhance contextual understanding.

The multimodal training corpus is envisioned as consisting of triplets: (Image, Omics Data, Text Context) with additional unimodal and bimodal data for enhanced alignment.

2.3 Model Architecture

The proposed architecture is modular and transformer-based, designed to process each modality via dedicated encoders before fusing their outputs into a unified model. Key components include:

2.3.1 Image Encoder

- Utilizes a Vision Transformer (ViT) pretrained on large-scale image datasets and fine-tuned on biomedical images.
- Extracts a grid of visual feature embeddings, which are subsequently projected into the joint multimodal space.

2.3.2 Transcriptomic Encoder

- Employs a pretrained omics model (e.g., Geneformer) that encodes gene expression data.
- Uses a projection layer to transform the high-dimensional omics features into a representation compatible with the joint fusion space.

2.3.3 Text Encoder / Language Model

- Leverages a large pretrained language model (e.g., a biomedical variant of GPT or LLaMA) that has been extensively trained on biomedical literature.
- Functions both as an encoder and decoder, utilizing reserved tokens (e.g., , <OMIC>) to integrate image and transcriptomic features.

2.3.4 Fusion and Integration

- The outputs from the image, transcriptomic, and text encoders are projected into a common fusion space.
- A cross-attention mechanism is employed, allowing the language model's self-attention to dynamically integrate the features across modalities.
- Optionally, an additional multimodal transformer-based fusion encoder can be used to explicitly combine the features before LLM processing.

2.3.5 Input Flexibility and Contextual Intelligence

- Input Flexibility: The system accepts various modalities—text prompts, images, or transcriptomic data—to accommodate different query types.
- Contextual Intelligence: Extensive pretraining on biomedical literature provides rich contextual information, enabling the system to generate interpretable and scientifically supported responses.
- Integration with Existing Models: The design facilitates the inclusion of outputs from specialized models (e.g., *Geneformer*), thereby enhancing the system's capability to analyze gene knockout effects across diverse cell types.

Figure 1 illustrates the conceptual diagram of the proposed multimodal architecture.

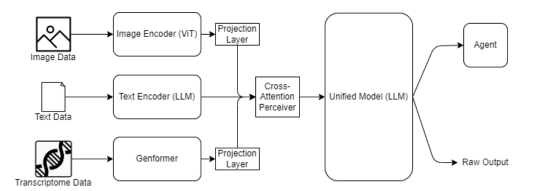


Figure 1: Conceptual diagram of the proposed multimodal architecture. Separate encoders process image, transcriptomic, and textual data. Their outputs are projected and fused into a unified input for a language model, enabling integrated multimodal reasoning.

2.4 Training Strategy

The training strategy is divided into several phases, using a curriculum learning approach to efficiently integrate the modalities:

2.4.1 Phase 0: Unimodal Pretraining

- Image Encoder: Pretrain using self-supervised learning methods (e.g., masked image modeling, contrastive learning) on large biomedical image datasets.
- Transcriptomic Encoder: Leverage self-supervised training on extensive single-cell transcriptomic datasets using a pretrained model like *Geneformer*.
- Language Model: Use a pretrained LLM that has been exposed to a vast corpus of biomedical literature.

2.4.2 Phase 1: Multimodal Alignment (Concept Integration)

- Freeze the individual encoders and the LLM; train only the projection layers.
- Task: Generate descriptive captions or annotations from paired image-omics-text examples.
- Objective: Align the representations across modalities with appropriate biomedical terminology.

2.4.3 Phase 2: Instruction Fine-Tuning (Multi-Task Learning)

- Gradually unfreeze parts of the LLM and the encoders.
- Train on curated instruction-following datasets (e.g., synthetic Q&A pairs generated using GPT-4) for tasks including classification, regression, and dialogue generation.
- Use a unified next-token prediction loss with curriculum learning to progressively increase task complexity.

2.4.4 Phase 3: Refinement and Transfer Learning

- Fine-tune the entire model on specific downstream tasks, such as predicting gene knockout outcomes.
- Optionally, employ self-supervised pseudo-labeling for further robustness.
- Utilize techniques such as early stopping and dropout-based uncertainty estimation to avoid overfitting.

2.5 Agent Integration

An autonomous agent framework is proposed to manage the multimodal model and integrate external knowledge sources.

2.5.1 Role of the Agent

- Task Decomposition: The agent divides complex queries into manageable subtasks (e.g., diagnosis, explanation, literature search).
- External Tool Querying: It interfaces with literature retrieval modules and specialized foundation models (e.g., *Geneformer*) to refine outputs.
- Iterative Refinement: Through a generate-assess-refine loop, the agent evaluates model confidence and retrieves supplementary data, ensuring accurate responses.

2.5.2 Integration with Foundation Models

- The agent routes transcriptomic data through specialized models like *Geneformer* for enhanced analysis.
- The outputs from these models are incorporated into the LLM context, enriching the final response with domain-specific insights.

2.5.3 Workflow and Iteration

- The agent acts as a wrapper that orchestrates the multimodal system and manages external API calls (e.g., PubMed literature searches, pathway database queries).
- Its decision-making process, potentially powered by an LLM-based planner, determines the sequence of tool invocations.
- This multi-agent collaboration ensures that the final responses are refined, interpretable, and backed by the most recent scientific evidence.

2.6 Conclusion of Model Proposal

The proposed multimodal system provides a comprehensive solution for integrating imaging, transcriptomic, and textual data to analyze gene knockout effects. By combining robust transformer-based architectures with domain-specific pretrained models, the system is designed to achieve high predictive accuracy and interpretability. This proposal establishes a strong foundation for future extensions, including integration of additional modalities such as proteomics and metabolomics.

3 Part 2: Coding Challenge

All code and logic is provided in the following repository: https://github.com/angadsinghsandhu/cz-biohub-test.

3.1 Problem Statement and Objectives

The coding challenge focuses on developing a multimodal learning system that integrates Geneformer single-cell transcriptomic embeddings (512-dimensional vectors) with textual embeddings derived from cell type labels. The system predicts donor identifiers while using an adversarial loss to control for the confounding variable of sex. The design emphasizes:

- Multimodal Integration: Using cross-attention and Perceiver Resampler modules to fuse omics and textual data.
- Pretraining: Contrastive pretraining of the projection modules to align the modalities.
- Adversarial Training: Mitigation of the sex confounder via gradient reversal and adversarial loss.
- **Hyperparameter Optimization:** Bayesian optimization (using Optuna) for tuning key parameters.

3.2 Dataset and Deliverables

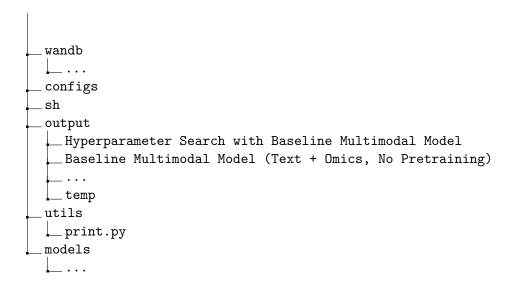
The dataset consists of single-cell data from the CELLxGENE Discover Census platform, restricted to human RNA measurements from the central nervous system. The deliverables include:

- 1. Data Summary: Reporting the number of cells loaded.
- 2. **UMAP Visualization:** A UMAP plot generated from Geneformer embeddings (see Figure 3).
- 3. Training Curves: Accuracy curves (training, validation, test) across epochs.
- 4. Attention Heatmap: A 2D heatmap visualizing the cross-attention weights (see Figure 4).
- 5. Code Structure and Documentation: A detailed description of the project directory, file structure, and modular design.
- 6. **Bonus Analyses:** Experiments evaluating the impact of contrastive pretraining and the integration of textual data.

3.3 Code Structure and Directory Overview

The project repository is organized as follows:





File Descriptions:

- uv.lock: Environment configuration using uv (a fast Rust-based package manager).
- main.py: The main driver script orchestrating data loading, model training, evaluation, and output generation.
- out/ and wandb/: Directories for logging and run artifacts.
- configs/: Contains configuration files including DeepSpeed configuration.
- sh/: Contains shell scripts and SLURM job scripts for HPC execution.
- output/: Stores generated artifacts such as UMAP plots, training curves, and hyperparameter files, and the best model.
- utils/: Utility scripts for data handling and logging.
- models/: Pretrained model files (e.g., BioBERT from DMIS Lab).

3.4 Overview and Code Structure

The provided Python script (main.py) implements:

- 1. Data Loading and UMAP Visualization: Loads single-cell data from the CELLxGENE census, filters for central nervous system cells, computes neighbors, and produces a UMAP plot based on Geneformer embeddings. The UMAP plot is saved as umap_geneformer.png (see Figure 3). A total of 31780 cells are loaded and split into train-validation-test sets (70%-15%-15%).
- 2. **Textual Embedding Extraction:** Uses a pretrained BioBERT model to extract the [CLS] token embedding for each cell type label. These embeddings form a matrix that is later fused with omics data.
- 3. **Model Architecture:** Defines the MultiModalModel which integrates omics and textual embeddings through:
 - **Projection Layers:** Mapping both omics and textual tokens into a common fusion space.

- Cross-Attention Block: Allowing omics tokens to attend to textual tokens.
- Transformer Encoder and Feed-Forward Network (FFN): Refining the fused tokens.
- Perceiver Resampler: Aggregating refined tokens into a latent representation.
- Classification Branches: One branch for donor ID prediction and an adversarial branch (with gradient reversal) for sex prediction.
- **Dimension Handling:** Dividing the omics embedding vector (512) and the text embedding vector (768) by the number of attention heads (8) to focus attention on specific parts.
- 4. Training and Hyperparameter Optimization: Implements training loops with check-pointing, hyperparameter search using Optuna, and logging via Weights & Biases. A contrastive pre-training phase for the Perceiver Resampler is also included, with the option to disable the textual modality.

3.4.1 Compute Environment

The code was written on a High-Performance Cluster (HPC) using SLURM management. Using 2 A100 40Gb GPUs with model computation in *bf16* floating point and utilizing **Flash_Attention_2** for fast training. HuggingFace's Accelerate is used to load Geneformer efficiently, and DeepSpeed ZeRo stage 0 is employed for data parallelism. These conditions ensure that the training loop does not exceed 10 minutes.

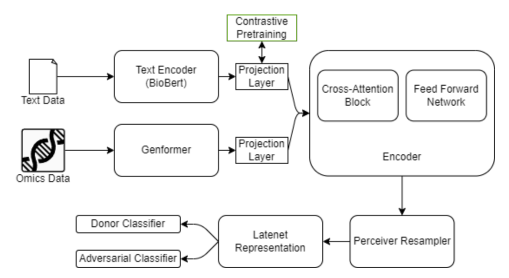


Figure 2: Conceptual diagram of the implemented multimodal architecture. The model integrates omics and textual embeddings using cross-attention and a Perceiver Resampler, followed by classification branches with an adversarial component.

3.4.2 Training Loop and Hyperparameter Optimization

The train_model function:

Hyperparameter tuning is performed via Optuna (using the objective function) and the best hyperparameters are stored in best_hyperparameters.json.

- Trains the model over multiple epochs.
- Saves checkpoints and logs training, validation, and test accuracies.
- Uses a cosine annealing learning rate scheduler with warmup.
- Implements epoch-wise training with checkpointing, using a cosine annealing scheduler with warmup.
- Utilizes Optuna for Bayesian hyperparameter search, with the best hyperparameters stored in best_hyperparameters.json.

3.4.3 Contrastive Pretraining and Ablation Study

- A contrastive pretraining phase (pretrain_contrastive) aligns omics and text projections.
- A command-line flag (--pretrain-resampler) enables this phase.
- Another flag (--disable-text-modality) runs an omics-only model for ablation.

3.4.4 Visualizations and Deliverables

The following images are generated as part of the deliverables:

- UMAP Plot: umap_geneformer.png (see Figure 3).
- Attention Heatmap: Cross-attention heatmap from the Perceiver Resampler (see Figure 4).
- Training Curves: Accuracy curves over epochs (see Figure 5).

Additional figures showing test accuracy, adversarial loss, and donor loss are included (Figures 6 and subsequent).

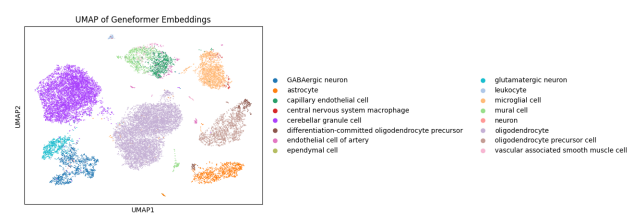


Figure 3: UMAP plot of Geneformer embeddings.

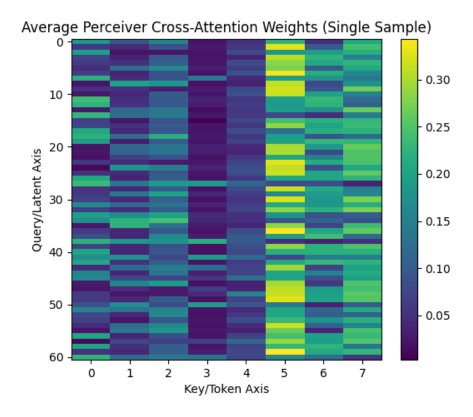


Figure 4: Attention Heatmap of the Perceiver.

3.4.5 Pretraining and Ablation Study

The script implements:

- pretrain_contrastive: A contrastive pretraining function that aligns the omics and text projections.
- A command-line flag (--pretrain-resampler) to enable this phase.
- A flag (--disable-text-modality) to run the model with only omics data, facilitating an ablation study.

3.5 Results

The experimental results demonstrate that my multimodal learning system achieved approximately 80% test accuracy in predicting donor identifiers when integrating both Geneformer-derived omics embeddings and textual embeddings from cell type labels, underscoring the effectiveness of my fusion strategy. This performance, along with favorable donor and adversarial loss metrics, validates the importance of incorporating both modalities, as ablation studies with omics-only data resulted in significantly lower accuracy.

3.6 Discussion of Bonus Analyses

Two bonus experiments are incorporated to evaluate additional aspects of the model:

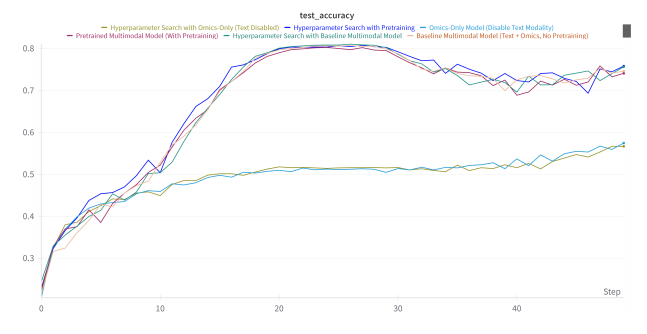


Figure 5: Test accuracy plot of all methods over 50 epochs. Only the best model was selected for the final testing (≈ 25 th Epoch).

a. Pretraining of the Perceiver Resampler:

- Controlled via the --pretrain-resampler flag.
- A contrastive pretraining routine (pretrain_contrastive) aligns the omics and text projections.
- Final analysis output advises comparing runs with and without pretraining.
- Final results indicate that pretraining produces little change in performance.

b. Impact of Cell Type Label Information:

- Controlled via the --disable-text-modality flag.
- Ablation runs using only omics data show a drastic decrease in donor prediction accuracy.
- When set, the model uses only omics data, enabling an ablation study.
- Final output includes analysis prompting comparison with the full multimodal model.

The results of these experiments are summarized in Table 1.

The ablation study clearly demonstrates that the integration of both omics and cell type textual data is critical for high donor prediction accuracy. While hyperparameter optimization yields a modest performance boost, contrastive pretraining of the Perceiver Resampler does not significantly improve the final results. In contrast, omitting the textual modality dramatically drops accuracy, highlighting the value of the cell type label information.

4 Deliverables Summary

The following deliverables have been completed:

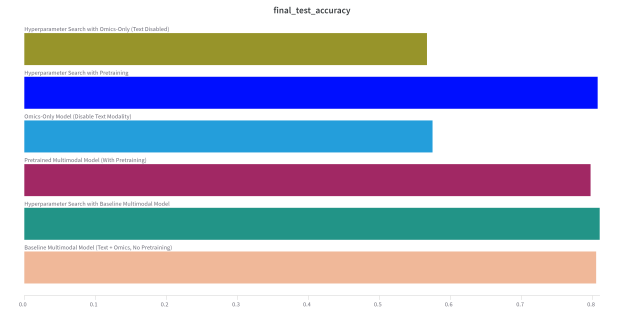


Figure 6: Final Accuracies achieved by the best model of each method.

- 1. **Data Summary:** The number of cells loaded is printed (31780 cells).
- 2. UMAP Visualization: The UMAP plot is saved as umap_geneformer.png (see Figure 3).
- 3. Training Curves: Accuracy curves across epochs are saved as training_curves.png.
- 4. Cross-Attention Heatmap: A heatmap of the average cross-attention weights is saved as perceiver_attention_heatmap.png (see Figure 4).
- 5. **Hyperparameter Optimization:** The best hyperparameters and validation accuracy are stored in best_hyperparameters.json.
- 6. **Bonus Analyses:** Comparative experiments demonstrate the critical role of cell type textual data and provide insights into the effects of contrastive pretraining

4.1 Experimental Results and Ablation Study

Table 1 summarizes the performance of various experimental configurations, including:

- a. Baseline Multimodal (Text + Omics, No Pretraining)
- b. Hyperparameter Search with Baseline Multimodal
- c. Pretrained Multimodal (With Pretraining)
- d. Omics-Only Model (Text Disabled)
- e. Hyperparameter Search with Pretraining
- f. Hyperparameter Search with Omics-Only

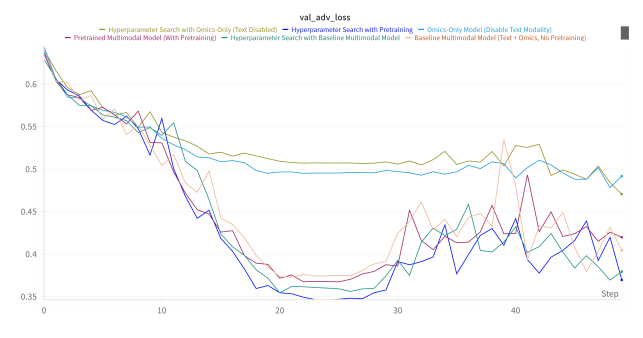


Figure 7: Adversarial loss on validation set

Table 1: Ablation Study and Analysis of Experimental Permutations

Experiment	Flags	Test Accuracy	Donor Loss	Adv Loss	Analysis Comments
1. Baseline Multimodal (Text + Omics, No Pretraining)	train-model	80.47%	1.5150	0.4465	Achieves high accuracy using both modalities without hyperparameter tuning.
${\bf 2. \ Hyperparameter \ Search \ with \ Baseline \ Multimodal}$	search-hparamstrain-model	81.00%	1.3456	0.4085	Hyperparameter optimization yields a modest improvement over the baseline.
3. Pretrained Multimodal (With Pretraining)	pretrain-resamplertrain-model	80.35%	1.3992	0.4242	Contrastive pretraining produces negligible improvement compared to the non-pretrained baseline.
4. Omics-Only Model (Text Disabled)	disable-text-modalitytrain-model	57.44%	1.4915	0.5100	Excluding cell type textual information causes a significant drop in performance.
5. Hyperparameter Search with Pretraining	search-hparamspretrain-resamplertrain-model	80.72%	1.3920	0.3856	Tuning with pretraining achieves similar results to tuning without pretraining.
6. Hyperparameter Search with Omics-Only	search-hparamsdisable-text-modalitytrain-model	54.07%	1.4615	0.4988	Even after tuning, the omics-only model performs substantially worse than the full multimodal model.

4.2 Experimental Results Summary and Discussion

The experiments demonstrate that the integration of both omics and textual data is critical for achieving high donor prediction accuracy. Hyperparameter tuning results in modest performance improvements, whereas the exclusion of textual data causes a substantial decrease in accuracy. Additionally, contrastive pretraining of the Perceiver Resampler yields marginal gains, suggesting that the primary performance benefit stems from the multimodal fusion of data sources. These findings are supported by the data presented in Table 1.

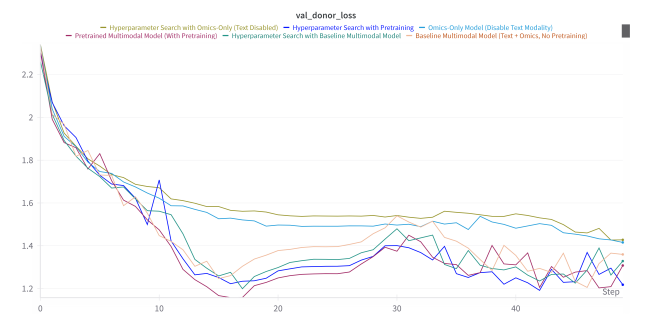


Figure 8: Donor loss on validation set

5 Conclusion

The developed multimodal learning system integrates omics and textual embeddings with adversarial training to predict donor IDs while controlling for the sex confounder. Key contributions of this work include:

- 1. A detailed data summary and UMAP visualization of Geneformer embeddings.
- 2. Extraction of textual embeddings using a pretrained BioBERT model.
- 3. Incorporation of an adversarial branch to mitigate the effect of sex as a confounder.
- 4. Implementation of modular training loops combined with hyperparameter optimization.
- 5. Generation of comprehensive visual outputs including UMAP plots, training curves, and cross-attention heatmaps.
- 6. Support for bonus analyses that examine the impact of contrastive pretraining and the inclusion of cell type textual data.

In addition to meeting the core deliverables, such as data summaries, UMAP plots, training curves, and cross-attention heatmaps, it also supports bonus analyses regarding pretraining and the impact of textual data. The system is modular, efficient, and reproducible, making it a robust foundation for further research and experimentation in biomedical multimodal learning. Detailed logging and checkpointing further enhance reproducibility and facilitate future extensions of the framework.

Bibliography

References

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