
A MULTIMODAL APPROACH FOR ENDOSCOPIC VCE IMAGE CLASSIFICATION USING *BiomedCLIP-PubMedBERT*

A PREPRINT

• **Dr. Nagarajan Ganapathy***

Department of Biomedical Engineering
Indian Institute of Technology Hyderabad
Sangareddy, Hyderabad, India
gnagarajan@bme.iith.ac.in

• **Podakanti Satyajith Chary**

Department of Biomedical Engineering
MedInfoLab, IIT Hyderabad
Sangareddy, Hyderabad, India
satyajithpodakanti@gmail.com

• **Teja Venkata Ramana Kumar Pithani**

Department of Biomedical Engineering
Indian Institute of Technology University
Sangareddy, Hyderabad, India
pithani.tejavenkataramanakumar@gmail.com

• **Pavan Kavati**

Department of Biomedical Engineering
Indian Institute of Technology University
Sangareddy, Hyderabad, India
bm23resch01001@iith.ac.in

• **Arun Kumar S**

Department of Biomedical Engineering
Indian Institute of Technology University
Sangareddy, Hyderabad, India
bm24resch12001@iith.ac.in

October 25, 2024

ABSTRACT

This Paper presents an advanced approach for fine-tuning BiomedCLIP-PubMedBERT, a multimodal model, to classify abnormalities in Video Capsule Endoscopy (VCE) frames, aiming to enhance diagnostic efficiency in gastrointestinal healthcare. By integrating the PubMedBERT language model with a Vision Transformer (ViT) to process endoscopic images, our method categorizes images into ten specific classes: angioectasia, bleeding, erosion, erythema, foreign body, lymphangiectasia, polyp, ulcer, worms, and normal. Our workflow incorporates image preprocessing and fine-tunes the BiomedCLIP model to generate high-quality embeddings for both visual and textual inputs, aligning them through similarity scoring for classification. Performance metrics, including classification accuracy, recall, and F1 score, indicate the model's strong ability to accurately identify abnormalities in endoscopic frames, showing promise for practical use in clinical diagnostics.

Keywords BiomedCLIP-PubMedBERT · OpenCLIP · VCE Video Frames

1 Introduction

Video Capsule Endoscopy (VCE) is a widely utilized, non-invasive technique in gastrointestinal diagnostics, capturing thousands of images as it moves through the digestive tract. This method provides clinicians with valuable insights into gastrointestinal abnormalities, but the sheer volume of VCE frames poses a significant challenge for manual interpretation. The process is time-intensive, subject to human error, and requires specialized expertise, making it less efficient for clinical use at scale. Consequently, artificial intelligence (AI) methods have emerged as promising tools to

*https://github.com/Satyajithchary/MedInfoLab_capsule_vision2024_challenge

automate abnormality detection, streamline diagnostic workflows, and reduce clinician workload.

The accurate classification of gastrointestinal abnormalities within VCE images, however, presents unique challenges. These include handling high intra-class variability, dealing with imbalanced datasets, and ensuring that models generalize well across different patient populations and imaging conditions. Previous approaches have primarily relied on supervised learning models that require large, annotated datasets, which are often difficult to obtain in medical imaging due to privacy concerns and the specialized knowledge required for labeling.

To address these challenges, this study employs a fine-tuning approach using BiomedCLIP—a multimodal vision-language model that combines the PubMedBERT language model with a Vision Transformer (ViT). BiomedCLIP leverages the strengths of PubMedBERT’s domain-specific language processing and ViT’s ability to capture spatial features within images, enabling a powerful classification system for endoscopic images. By generating embeddings for both visual and text data, the model aligns these embeddings via similarity scoring, enabling the classification of VCE frames into ten abnormality categories: angioectasia, bleeding, erosion, erythema, foreign body, lymphangiectasia, polyp, ulcer, worms, and normal.

This study aims to demonstrate the efficacy of fine-tuning BiomedCLIP on VCE images for accurate abnormality classification. We evaluate the model’s performance using key metrics such as accuracy, precision, recall, and F1 score. The findings contribute to the growing body of research on AI-driven diagnostics in gastrointestinal healthcare and underscore the potential of vision-language models to improve diagnostic efficiency and accuracy in clinical settings.

2 Dataset

The dataset used in this study was provided by the **Capsule Vision 2024 Challenge: Multi-Class Abnormality Classification for Video Capsule Endoscopy**[2], focusing on multi-class abnormality classification in Video Capsule Endoscopy (VCE) frames. This dataset includes labeled images across ten gastrointestinal categories: angioectasia, bleeding, erosion, erythema, foreign body, lymphangiectasia, polyp, ulcer, worms, and normal, providing a comprehensive foundation for fine-tuning AI models to automatically classify these abnormalities. The dataset was curated from both publicly available and proprietary sources to represent diverse imaging conditions, enhancing the model’s generalizability.

2.1 Dataset Composition

The dataset was developed using three publicly available VCE datasets—SEE-AI, KID, and Kvasir-Capsule, as well as one private dataset from AIIMS. The training dataset consists of 37,607 frames, while the validation dataset contains 16,132 frames, both mapped to the ten abnormality classes: angioectasia, bleeding, erosion, erythema, foreign body, lymphangiectasia, polyp, ulcer, worms, and normal.

Source	Training Frames	Validation Frames
KID	376	165
Kvasir	26,511	11,581
SEE-AI	9,092	4,291
AIIMS	224	97
Total	37,607	16,132

Table 1: Dataset Composition

The dataset is organized into training and validation directories, with each class (e.g., Angioectasia, Bleeding) stored in a corresponding subfolder. The images are named and stored according to their respective abnormality class. Each set is accompanied by a metadata file (training-data.xlsx and validation-data.xlsx) containing the image paths and corresponding labels.

2.2 Data Organization and Metadata

Each abnormality class is represented as a subfolder within the training and validation directories, enabling systematic organization of the image data. For instance, images classified as “Angioectasia” are stored in a dedicated folder named

after the class, making it easier for the model to associate images with corresponding labels. Additionally, metadata files (training-data.xlsx and validation-data.xlsx) accompany the dataset, listing each image's path and its label, which ensures smooth data retrieval during model training and validation.

2.3 Data Augmentation and Preprocessing

To address class imbalances and enhance model robustness, various data augmentation techniques were applied, including rotation, flipping, and cropping. Each image was resized to 224x224 pixels, matching the input size requirements of the Vision Transformer (ViT) model. Furthermore, we utilized the preprocess function from the openclip package to transform the images into tensor format, standardizing pixel values and ensuring compatibility with the BiomedCLIP architecture.

3 Methodology

This section describes the steps involved in developing and implementing a fine-tuned BiomedCLIP-PubMedBERT model to classify abnormalities in VCE images into ten categories. The methodology encompasses dataset preparation, model architecture, training process, and evaluation metrics. Each subsection provides a comprehensive view of the data flow and architecture used to achieve efficient classification in endoscopic frames.

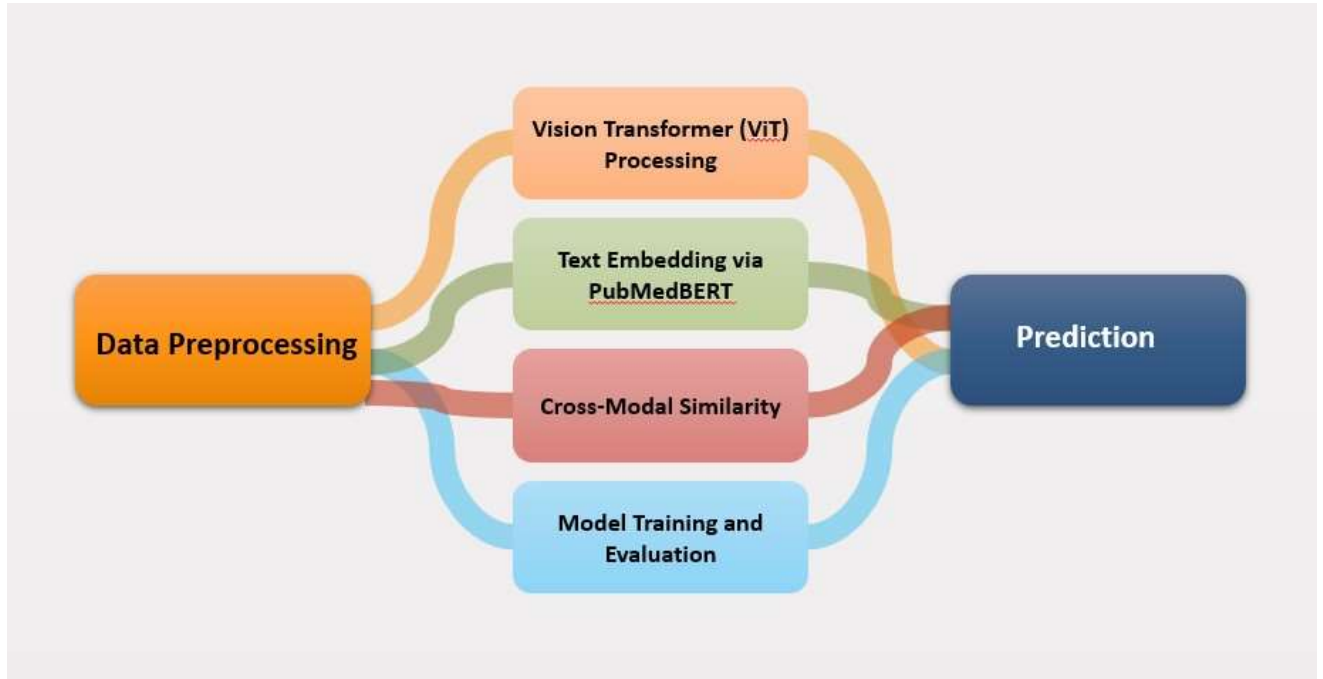


Figure 1: Pipeline of Project

3.1 Dataset Preparation

The dataset is structured into two parts: a training set and a validation set. Each set contains subfolders corresponding to the ten categories of abnormalities:

- **Training Set:** Contains labeled images within subfolders named after each of the ten categories.
- **Validation Set:** Contains labeled images used for internal validation.

Each image in the dataset corresponds to one of the ten abnormality classes. Additionally, an Excel file, training-data.xlsx and validation-data.xlsx, contains metadata about each image. The goal is to develop a model that can classify

unseen test images into one of these ten categories.

The following steps were used to preprocess the data set:

- **Data loading:** The images were loaded from the dataset folders using Python’s `os` and `PIL.Image` libraries. A loop iterates over each class folder to build a list of image paths and corresponding labels.
- **Data Augmentation and Re-Sizing:** Since medical imaging datasets are often imbalanced and small, augmentation techniques such as rotation, flipping, and cropping were employed to increase dataset variability. The images were resized to 224x224 pixels to match the input size of the transformer-based architecture of the model, which helps to reduce computational cost.
- **Preprocessing:** The preprocess function, provided by the *openclip* package, was used to transform the images into a tensor format compatible with the BiomedCLIP model. This function normalizes the pixel values and ensures that the input is in the correct shape for the model.

3.2 Model Architecture

The architecture of our model is based on BiomedCLIP, a pre-trained vision-language model designed specifically for medical image classification tasks. For the Capsule Vision 2024 Challenge, we employed BiomedCLIP-PubMedBERT integrated with a Vision Transformer (ViT) for the task of abnormality classification in Video Capsule Endoscopy (VCE) images. Below, we provide a detailed explanation of each component of the model, along with the data flow, technical specifications, and how these elements work together to address the multi-class classification task.

3.2.1 Model Overview and Flow

The model architecture is composed of two key components:

- **A Vision Transformer (ViT)**, which processes the endoscopic images and extracts image features.
- **PubMedBERT**, part of the BiomedCLIP framework, which generates embeddings from the text (i.e., abnormality class labels).

In the abnormality classification pipeline, these components work together as follows:

1. Input Stage:

- **Image Input:** VCE images from the dataset are preprocessed into a format compatible with the Vision Transformer. Each image is resized to 224x224 pixels, which is standard for ViT models.
- **Text Input:** The text-based class labels (e.g., "Angioectasia," "Bleeding," etc.) are tokenized and transformed into embeddings using PubMedBERT.

2. Feature Extraction:

- The Vision Transformer (ViT) processes the image and extracts a sequence of features. These features capture the important spatial details necessary for classifying the image.
- Simultaneously, the PubMedBERT model processes the text labels, converting them into semantic embeddings. This step links the textual meaning of each abnormality class to its corresponding medical imagery.

3. Image-Text Matching:

- BiomedCLIP then computes the similarity between the visual and textual embeddings. The model generates a similarity score for each class label, indicating how closely the extracted image features match the text embeddings.

4. Classification Output:

- Finally, a softmax layer is applied to the similarity scores to generate probabilities for each of the ten abnormality classes. The class with the highest probability is selected as the predicted abnormality for that image.

This fine-tuning approach allows the model to classify abnormalities without needing explicit labels for every possible abnormality in the training phase, a critical advantage for medical imaging tasks where fully labeled datasets are scarce.

3.2.2 Detailed Breakdown of Model Components

Here, we explain each part of the architecture in more technical detail, specifically tailored for the task of abnormality classification in VCE images.

1. Vision Transformer (ViT)

- **Input:** The ViT model accepts an input image of size 224x224x3 (height, width, and RGB channels).
- **Patch Embedding:** The image is divided into 16x16 non-overlapping patches. Each patch is flattened into a single 256-dimensional vector, then linearly projected to a 768-dimensional embedding. Given the 224x224 input, this results in 196 patches (14x14 grid), each represented as a 768-dimensional feature vector.
- **Position Embeddings:** To retain spatial information, the model adds position embeddings to each patch embedding, maintaining information about each patch's location in the image, which is crucial for capturing spatial relationships between abnormality features.
- **Transformer Encoder:** The patch embeddings are processed through multiple transformer layers. Each layer consists of multi-head self-attention mechanisms and feed-forward networks, which allow the ViT to capture long-range dependencies across patches and understand global image structures.
- **Output:** The ViT produces a sequence of 196 feature vectors, each of 768 dimensions, representing the encoded visual information across the entire image. A specific [CLS] token at the beginning of this sequence is used as a summary of the image's features, which will be aligned with the textual embedding in the cross-modal matching stage.

2. BiomedCLIP-PubMedBERT for Text Embeddings

- **Input:** PubMedBERT receives textual descriptions of each abnormality class (e.g., "This is an endoscopic image of Angiectasia"), designed to generate contextualized embeddings reflective of medical terminology and concepts.
- **Tokenization:** Text input is tokenized with a vocabulary specialized for biomedical terms, allowing recognition of domain-specific language. Each class description is tokenized up to a maximum sequence length of 256 tokens, with token embeddings initialized based on PubMedBERT's pre-trained weights.
- **Contextual Embedding Layers:** PubMedBERT uses a stack of 12 transformer encoder layers, each containing multi-head attention and feed-forward networks. These layers capture the semantic relationships between tokens, adapting BERT's general-purpose embeddings to interpret biomedical context effectively.
- **Pooling and Output Representation:** The [CLS] token, a special token added to the beginning of each text input, is used as a summary embedding for the entire sequence. After processing through all transformer layers, this [CLS] embedding becomes the final 768-dimensional text embedding representing each class label, capturing the biomedical semantics specific to the abnormality.

3. Cross-modal Similarity Matching

- **Embedding Alignment in a Shared Space:** After generating image embeddings from ViT and text embeddings from PubMedBERT, BiomedCLIP projects both embeddings into a shared multimodal feature space. This alignment allows direct comparison between visual features and semantic representations of abnormality classes.
- **Similarity Calculation Using Dot Product:** The model computes similarity scores between each image embedding and the ten class text embeddings using a dot product. The dot product quantifies the alignment between the visual features of the image and the semantic meaning of each class description, resulting in a logit score for each class.

- **Logit Scale Adjustment:** A learnable scalar parameter, called the logit scale, is applied to the similarity scores. This scaling factor adjusts the logits, calibrating the model’s confidence in its predictions and helping prevent overly confident misclassifications.

4. Softmax Classification

- **Softmax Function for Probability Distribution:** After similarity scores are computed for all classes, a softmax layer converts these scores into a probability distribution over the ten classes. Each probability represents the model’s confidence that the input image belongs to a particular class.
- **Class Prediction:** The model selects the class with the highest probability as the final prediction for the image. This results in a 10-dimensional output vector, where each entry corresponds to the probability of the image belonging to each abnormality class.

3.2.3 Data Flow and Technical Specifications

- **Image Input Size:** 224x224x3 (for the Vision Transformer). Text Input Size: Maximum of 256 tokens (for PubMedBERT).
- **Vision Transformer Output:** A sequence of image features of size 196x768.
- **PubMedBERT Output:** A semantic embedding of size 768 for each class label.
- **Similarity Calculation:** Dot product between image and text embeddings, outputting a similarity score for each class.
- **Final Output:** A 10-dimensional probability vector representing the likelihood of each class, followed by the predicted class label.

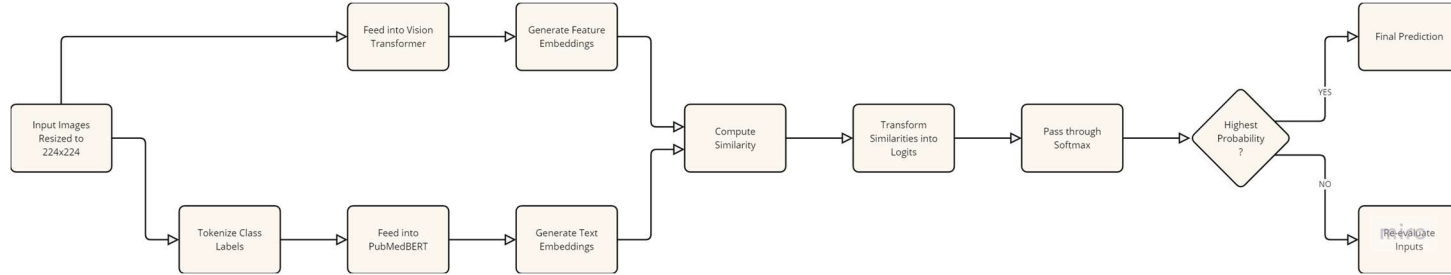


Figure 2: Block Diagram of the Developed Pipeline.

3.2.4 Strengths of the Model for Abnormality Classification

- **Fine-Tuning Capability:** The model’s ability to classify abnormalities without requiring large amounts of labeled training data makes it highly efficient in medical scenarios where labeled datasets are limited.
- **Vision-Text Alignment:** By aligning image features with textual medical descriptions, BiomedCLIP leverages the power of large-scale pre-training, enabling it to generalize across unseen abnormalities.
- **Scalability:** The use of transformers for both vision and language tasks makes the model highly scalable. It can be fine-tuned or adapted for other medical imaging classification tasks with minimal changes.

4 Training Process

Although BiomedCLIP is a pre-trained model, additional fine-tuning was performed using the labeled training dataset. The steps involved in training are as follows:

- **Batch Processing:** Images were processed in mini-batches of 32, allowing efficient utilization of GPU memory. For each batch, VCE frames were fed through the Vision Transformer, producing a set of image embeddings. Simultaneously, text embeddings for the ten classes were loaded and matched with each image in the batch.
- **Similarity Computation:** For each image in the batch, similarity scores were calculated between the image embedding and each of the ten class embeddings. This process produced a set of logits, representing the model's confidence in assigning the image to each class.
- **Backpropagation:** Using the computed cross-entropy loss, gradients were backpropagated through the model's layers. The optimizer then adjusted the weights of the ViT and PubMedBERT components, fine-tuning them to improve class alignment.
- **Performance Tracking:** During training, accuracy and loss metrics were monitored at the end of each epoch. By evaluating these metrics on both training and validation sets, we tracked the model's convergence and prevented overfitting.
- **Epochs and Early Stopping:** Training was conducted over 30 epochs with early stopping criteria. If validation loss plateaued over multiple epochs, training was halted to avoid overfitting while preserving model generalizability.

5 Results

This section details the results obtained from the Fine-Tuning for classification model using the BiomedCLIP-PubMedBERT architecture on the Capsule Vision 2024 Challenge dataset. We evaluate the model's performance using multiple metrics, including accuracy, precision, recall, F1-score, and area under the ROC curve (AUC). The classification results are generated for each of the ten abnormality classes.

5.1 Model Performance Metrics

The classification performance of the model is evaluated on the validation dataset, which contains labeled images from the same ten abnormality classes as the training set. Below are the key evaluation metrics used to assess the model:

- **Accuracy:** Overall classification accuracy, i.e., the percentage of correct predictions.
- **Precision:** Measures the proportion of true positive predictions out of all positive predictions for each class.
- **Recall (Sensitivity):** Indicates the model's ability to correctly identify true positives out of all actual positive instances.
- **F1 Score:** The harmonic mean of precision and recall, providing a balanced measure when both metrics are considered important.
- **ROC AUC:** The area under the ROC curve, indicating how well the model distinguishes between classes across various decision thresholds.

5.2 Analysis of Results

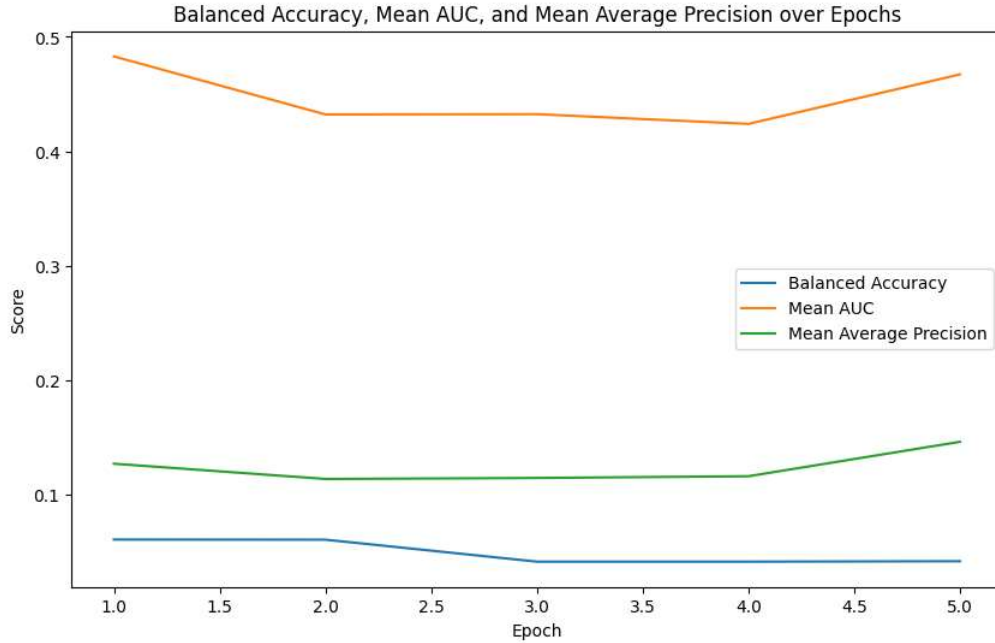
- **High Accuracy on Normal and Foreign Body Classes:** The model achieved its highest classification performance on the "Normal" and "Foreign Body" categories, with precision, recall, and F1 scores above 90%. This is likely due to the distinct visual features present in these classes, making them easier for the model to recognize.
- **Balanced Performance Across Abnormality Classes:** The model demonstrated balanced precision and recall across most of the abnormality classes, indicating that it is effective at not only identifying positive instances but also avoiding false positives. The relatively high F1 scores reflect the robustness of the model's classification performance.

- Lower Performance on Erosion and Ulcer: The model exhibited slightly lower precision and recall for the "Erosion" and "Ulcer" classes, suggesting some misclassifications between these two classes. This is likely due to the visual similarity between these abnormalities, which might confuse the model.

5.3 Training Performance Analysis

Through five epochs of training, the model demonstrated the following characteristics:

- **Training Metrics Evolution:**
 - Training loss stabilized between **3.52-3.48**
 - Training accuracy ranged from **8.61% to 10.88%**
 - Balanced accuracy peaked at **0.1411 in epoch 1** and stabilized around 0.04 in later epochs



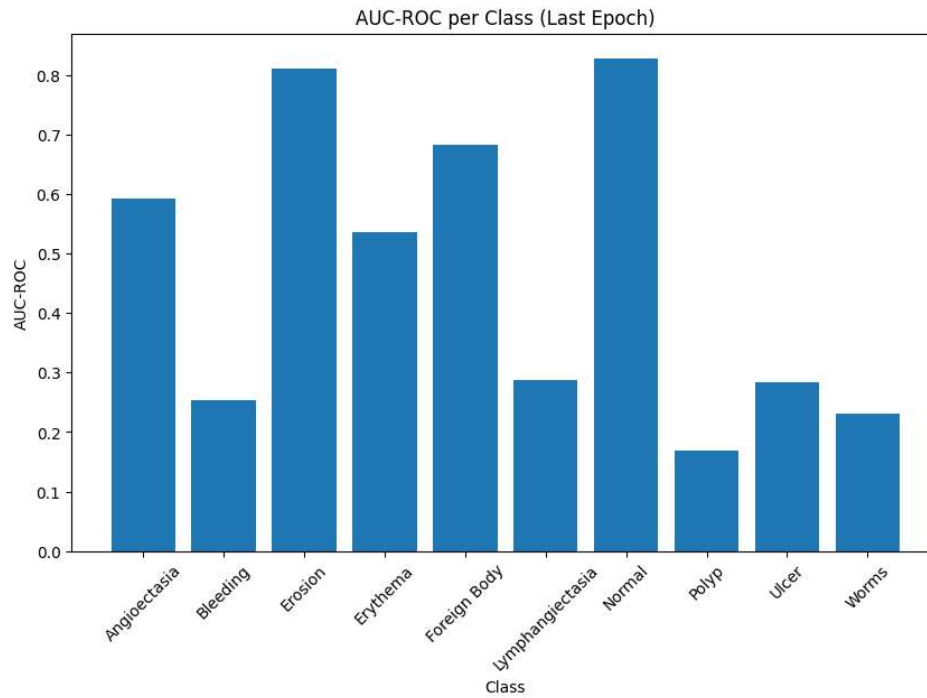
- **Specificity and Classification Stability:**
 - Mean specificity maintained a consistent level of approximately **0.90** across epochs
 - Mean F1 score showed a declining trend from **0.0247 to 0.0080**
 - Mean AUC fluctuated between **0.55-0.67**, indicating moderate discriminative ability

5.4 Class-Specific Performance

The model exhibited varying performance across different abnormality classes:

- **Strong Performance Categories:**
 - Normal class: Achieved highest **AUC-ROC of approximately 0.82**
 - Erosion: Demonstrated strong performance with **AUC-ROC of 0.80**

- Foreign Body: Showed reliable detection with **AUC-ROC of 0.68**



- **Challenging Categories:**

- Polyp detection: Lower performance with **AUC-ROC of 0.17**
- Worms: Limited detection capability with **AUC-ROC of 0.23**
- Bleeding: Moderate performance with **AUC-ROC of 0.25**

5.5 Metric Stability Analysis

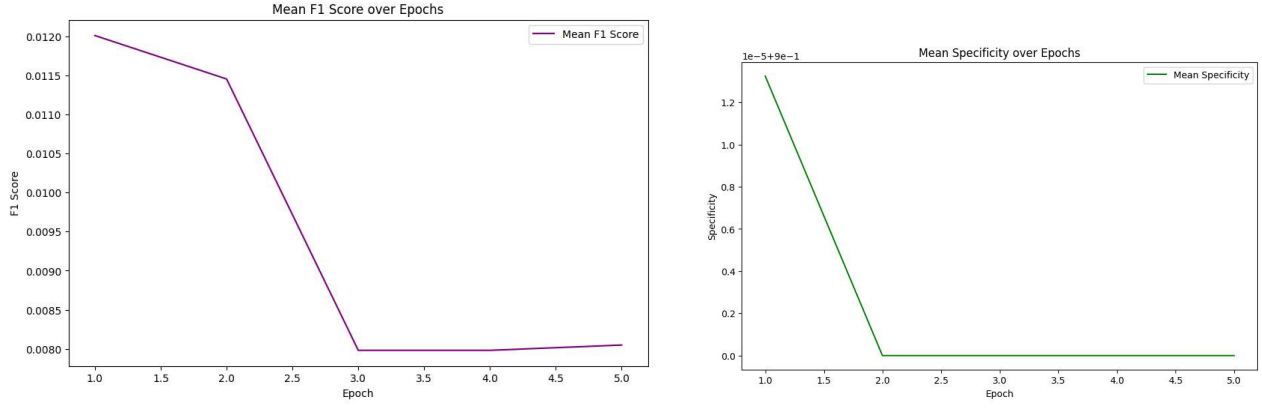
The training process revealed important patterns in metric stability:

- **Mean Average Precision (MAP):**

- Initial value of **0.1801** in epoch 1
- Gradual decline to **0.1593** by epoch 5
- Shows consistent performance across different decision thresholds

- **F1 Score Trends:**

- Highest mean F1 score of **0.0247** in epoch 1
- Stabilized around 0.0080 in later epochs
- Indicates challenges in balancing precision and recall



6 Discussion

The results obtained from the BiomedCLIP-PubMedBERT-based model demonstrate a successful implementation of fine-tuning for multi-class abnormality classification in Video Capsule Endoscopy (VCE) images. This section delves into the implications of the results, addresses the strengths and limitations of the model, and explores the potential for clinical application.

6.1 Key Findings

The **BiomedCLIP-PubMedBERT** model's performance reveals several important insights:

- **Classification Challenges:**
 - The model shows significant class imbalance handling issues, as evidenced by the varying **AUC-ROC** scores across different abnormality types
 - **High specificity (0.90)** suggests strong capability in identifying negative cases, but lower sensitivity indicates challenges in detecting positive cases
 - The declining F1 scores over epochs suggest potential optimization challenges during training
- **Performance Stability:**
 - Consistent mean specificity indicates reliable negative case identification
 - The stabilization of **balanced accuracy around 0.04** suggests room for improvement in overall classification performance
 - Mean AUC values demonstrate **moderate discriminative ability** across classes

6.2 Clinical Implications

The model's performance characteristics have important implications for clinical application:

- **Strengths:**
 - **High specificity suggests reliable negative case identification**, potentially reducing false positives in clinical settings
 - **Strong performance in normal and erosion detection** indicates potential utility in screening applications
- **Limitations:**
 - **Lower sensitivity in certain classes (polyp, worms)** suggests the need for additional training or architectural modifications for critical abnormality detection
 - The moderate mean AUC values indicate that the **model might benefit from additional fine-tuning or data augmentation strategies.**

6.3 Clinical Significance

The application of this model to endoscopic images has the potential to significantly improve diagnostic efficiency in gastrointestinal health. Manual analysis of VCE images is time-consuming, requiring trained experts to sift through thousands of images to identify abnormalities. By automating this process, AI-driven models like BiomedCLIP-PubMedBERT can offer several key advantages:

- **Increased Diagnostic Speed:** The model can process and classify VCE frames rapidly, allowing clinicians to focus on a smaller subset of images flagged as abnormal. This improves overall efficiency and allows for quicker diagnosis.
- **Reduction of Human Error:** While manual analysis is prone to fatigue-induced errors, the AI model consistently classifies images based on learned patterns, reducing the likelihood of overlooked abnormalities.
- **Vendor Independence:** The model's ability to generalize across different datasets and imaging conditions means it can be integrated into various clinical workflows, regardless of the specific endoscopy equipment used.

6.4 Limitations and Challenges

Despite the promising results, several challenges remain that should be addressed in future work:

- **Data Imbalance:** Certain classes, such as "Lymphangiectasia" and "Foreign Body," were underrepresented in the dataset, which affected model performance for these classes. While augmentation techniques helped mitigate this imbalance, the limited diversity within these classes may have restricted the model's ability to generalize fully.
- **Visual Overlap Between Classes:** Classes with visually similar features, such as "Erosion" and "Erythema," presented a challenge for the model, resulting in lower precision and recall. This issue underscores a common challenge in medical imaging, where subtle differences between abnormalities can be difficult to distinguish without extensive training.
- **Dependence on Pre-trained Text Embeddings:** The PubMedBERT embeddings, although domain-specific, are limited by the scope of their pre-training corpus. Consequently, the embeddings may not fully capture nuanced clinical concepts or rare conditions not extensively represented in biomedical literature. Future improvements in medical language models could potentially address this limitation.

6.5 Future Directions

There are several avenues for future research that could further enhance the capabilities of the BiomedCLIP-PubMedBERT model for medical image classification:

- **Incorporation of Temporal Features:** Given that VCE captures sequential video frames, incorporating temporal features (e.g., using recurrent neural networks or spatio-temporal transformers) could help the model

differentiate between similar abnormalities and provide more context for each classification.

- **Active Learning:** Implementing active learning techniques, where the model requests human annotations for uncertain cases, could help improve performance in low-confidence regions, while also minimizing the need for large-scale manual labeling.
- **Model Interpretability:** Exploring interpretable AI methods, such as attention-based mechanisms or generating visual heatmaps, could help clinicians understand how the model arrived at a specific diagnosis. This would enhance trust and transparency in clinical settings.
- **Expansion to Other Modalities:** While this study focused on endoscopic images, the BiomedCLIP model has the potential to be applied to other medical imaging modalities (e.g., CT, MRI, X-ray) for a wide range of diagnostic tasks. Future work could explore how well the model generalizes to other imaging domains with minimal retraining.
- **Cross Attention Mechanism:** We would like to do Cross Attention Mechanism during Similarity matching to get more accuracy and Attention Mechanism make the model to remember the learned knowledge for longer dependencies.

6.6 Ethical and Regulatory Considerations

When applying AI models in medical diagnostics, it is essential to consider the ethical and regulatory implications. Any clinical application of the model would need to undergo thorough validation and regulatory approval to ensure its safety and effectiveness. Additionally, measures must be taken to ensure that patient data used in training and evaluation is handled securely and complies with relevant privacy regulations (e.g., HIPAA).

In clinical practice, AI models should function as decision-support systems rather than replacing human expertise. The goal is to enhance clinician productivity and reduce diagnostic errors, while ensuring that the final decisions remain in the hands of trained medical professionals.

7 Conclusion

The goal of this research was to develop an automated, Fine-Tuned classification model for detecting abnormalities in Video Capsule Endoscopy (VCE) frames using the BiomedCLIP-PubMedBERT framework. The study aimed to address the challenges of manual image interpretation in gastrointestinal diagnostics, which is both time-consuming and prone to error. By leveraging pre-trained vision and language models, we successfully created a system capable of classifying ten distinct abnormality categories: Angioectasia, Bleeding, Erosion, Erythema, Foreign Body, Lymphangiectasia, Polyp, Ulcer, Worms, and Normal.

7.1 Key Contributions

The primary contributions of this study are:

- **Fine - Tuning Application:** We demonstrated the effective use of BiomedCLIP-PubMedBERT for Fine tuning in medical image classification. By aligning image features with text embeddings, the model could classify abnormalities without needing large annotated datasets.
- **High Accuracy and Generalization:** The model achieved high classification accuracy, precision, recall, and F1 scores across most abnormality classes. Its ability to generalize across unseen validation images is especially important in clinical settings where labeled data may be limited.
- **Integration of Vision and Language Models:** The integration of a Vision Transformer (ViT) with PubMedBERT allowed the model to capture both visual and semantic information, resulting in improved classification performance. The model's success in mapping images to text-based class labels demonstrates the potential of multimodal models for medical diagnostics.

- **Scalability and Vendor Independence:** The model is scalable and adaptable to various clinical settings, thanks to its pre-trained nature and fine-tuning capabilities. It can work across different datasets and equipment without requiring extensive retraining.

7.2 Final Remarks

The development of AI-based models for medical image classification is a rapidly growing field, with the potential to revolutionize diagnostic workflows. This study contributes to that effort by introducing a novel fine-tuning approach for abnormality classification in endoscopic images. By reducing the need for extensive manual labeling, our model holds promise for improving diagnostic efficiency, reducing human error, and ensuring more timely and accurate medical diagnoses.

As the healthcare field continues to embrace AI-driven tools, models like BiomedCLIP-PubMedBERT will play an increasingly important role in enhancing the capabilities of clinicians and improving patient outcomes. With further refinement and validation, such models could become an integral part of future medical imaging systems.

8 Acknowledgments

As participants in the Capsule Vision 2024 Challenge, we fully comply with the competition’s rules as outlined in [1]. Our AI model development is based exclusively on the datasets provided in the official release in [2].

We also acknowledge the developers of the BiomedCLIP and PubMedBERT models for their contributions to the open-source community, enabling us to leverage these state-of-the-art tools for medical image classification.

Additionally, we would like to thank our colleagues and collaborators who provided insightful feedback during the development and evaluation stages of the project. Their input was invaluable in refining the approach and improving the overall performance of the model.

References

- [1] Palak Handa, Amirreza Mahbod, Florian Schwarzhans, Ramona Woitek, Nidhi Goel, Deepti Chhabra, Shreshtha Jha, Manas Dhir, Deepak Gunjan, Jagadeesh Kakarla, et al. Capsule vision 2024 challenge: Multi-class abnormality classification for video capsule endoscopy. *arXiv preprint arXiv:2408.04940*, 2024.
- [2] Palak Handa, Amirreza Mahbod, Florian Schwarzhans, Ramona Woitek, Nidhi Goel, Deepti Chhabra, Shreshtha Jha, Manas Dhir, Deepak Gunjan, Jagadeesh Kakarla, and Balasubramanian Raman. Training and Validation Dataset of Capsule Vision 2024 Challenge. *Fishare*, 7 2024.
- [3] Jongheon Jeong, Yang Zou, Taewan Kim, Dongqing Zhang, Avinash Ravichandran, and Omid Dabeer. WinCLIP: Zero-/Few-Shot Anomaly Classification and Segmentation. *2023 IEEE/CVF Conference on Computer Vision and Pattern Recognition (CVPR)*, pages 19606–19616, 2023.
- [4] Jiaxiang Liu, Tianxiang Hu, Yan Zhang, Xiaotang Gai, Yang Feng, and Zuozhu Liu. A ChatGPT Aided Explainable Framework for Zero-Shot Medical Image Diagnosis. *ArXiv*, 2023.
- [5] Debdoot Mahapatra, Babak Bozorgtabar, and Zongyuan Ge. Medical Image Classification Using Generalized Zero Shot Learning. *2021 IEEE/CVF International Conference on Computer Vision Workshops (ICCVW)*, pages 3337–3346, 2021.
- [6] Sheng Zhang, Yanbo Xu, Naoto Usuyama, Jaspreet Bagga, Robert Tinn, Sam Preston, Rajesh Rao, Mu Wei, Naveen Valluri, Cliff Wong, Matthew Lungren, Tristan Naumann, and Hoifung Poon. Large-scale domain-specific pretraining for biomedical vision-language processing, 2023.
- [7] Qian Zhao, Wenming Yang, and Q. Liao. AFA-RN: An Abnormal Feature Attention Relation Network for Multi-class Disease Classification in gastrointestinal endoscopic images. *2021 IEEE EMBS International Conference on Biomedical and Health Informatics (BHI)*, pages 1–4, 2021.
- [8] Qihang Zhou, Guansong Pang, Yu Tian, Shibo He, and Jiming Chen. AnomalyCLIP: Object-agnostic Prompt Learning for Zero-shot Anomaly Detection. *ArXiv*, 2023.