EC-RAFT: Automated Generation of Clinical Trial Eligibility Criteria through Retrieval-Augmented Fine-Tuning

Anonymous ACL submission

Abstract

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Eligibility criteria (EC) are critical components of clinical trial design, defining the parameters for participant inclusion and exclusion. However, designing EC remains a complex, expertise-intensive process. Traditional approaches to EC generation often rely on user-prompted predefined categories, which limit adaptability and may fail to produce comprehensive, contextually appropriate criteria. To address these challenges, we introduce EC-RAFT, a method that utilizes Retrieval-Augmented Fine-Tuning (RAFT) to generate structured and cohesive EC directly from clinical trial titles and descriptions. EC-RAFT integrates contextual retrieval, synthesized intermediate reasoning, and fine-tuned language models to produce comprehensive EC sets. To enhance clinical alignment evaluation with referenced criteria, we also propose an LLM-guided evaluation pipeline. Our results demonstrate that our solution, which uses Llama-3.1-8B-Instruct as a base model, achieves a BERTScore of 86.23 and an EC-matched LLM-as-a-Judge score of 1.66 out of 3, outperforming zero-shot Llama-3.1 and Gemini-1.5 by 0.41 and 0.11 points, respectively. EC-RAFT was trained in a low-cost setup and, therefore, can be used as a practical solution for EC generation while ensuring quality and relevance in clinical trial design. We release our code on GitHub at

1 Introduction

Eligibility Criteria (EC) are essential components of clinical trial design, specifying the parameters for participant inclusion and exclusion (Su et al., 2023). These criteria ensure trials are scientifically valid, ethically sound, and capable of meeting their objectives. However, designing EC remains a laborintensive and expertise-driven process (Su et al., 2023). Tools that can suggest or generate relevant EC have the potential to significantly facilitate researchers' work in trial design (Kim et al., 2024). Generating these criteria is inherently complex be-044 cause consistency and clinical validity are needed 045 throughout the criteria set. Despite advances in 046 using large language models (LLMs) for summa-047 rization or specialized tasks in the biomedical domain, several barriers remain to creating a fully automated, contextually accurate system that can generate comprehensive sets of EC directly from 051 trial descriptions. Recent developments in instruction fine-tuning for LLMs have shown promise in generating logical reasoning outputs through techniques like chain-of-thought prompting and 055 rationale generation (Wei et al., 2022). Retrievalaugmented generation (RAG) has also emerged as an effective mechanism for grounding model 058 outputs with external domain knowledge, thereby improving factual correctness (Ram et al., 2023). 060 Retrieval-augmented fine-tuning (RAFT) extends 061 RAG by incorporating instruction fine-tuning to 062 improve both domain adaptation and retrieval ro-063 bustness (Zhang et al., 2024). These developments 064 allow the development of an end-to-end system to 065 generate a complete set of EC while preserving 066 essential clinical context and domain relevance. To 067 address these gaps, we propose EC-RAFT, a novel 068 approach that leverages Retrieval-Augmented Fine-069 Tuning (RAFT) (Zhang et al., 2024) for automated 070 EC generation. EC-RAFT aims to produce com-071 plete EC sets directly from trial titles and descrip-072 tions without requiring user-input EC categories 073 or a recommendation system. Our key features 074 include: 075

1. RAFT (Zhang et al., 2024) incorporates relevant external clinical trial information (existing trial details and eligibility criteria) and generates intermediate reasoning steps to finetune LLM. 076

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2. Generating a complete set of eligibility criteria results in a fully structured set of inclusion and exclusion criteria. We demonstrate that

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084synthesized intermediate reasoning steps pro-085duced by LLM, enhance the performance of086the base models during fine-tuning for EC087generation. Our results show that EC-RAFT088exceeds zero-shot baseline approaches across089multiple evaluation metrics, including seman-090tic similarity and LLM-as-a-judge scoring.

Our training setup was also optimized for cost efficiency using the Parameter-Efficient Fine-Tuning technique (PEFT) (Xu et al., 2023; Hu et al., 2021). Specifically, training our best model required 380 GPU hours on NVIDIA A100 costing approximately 452.20 USD while achieving superior performance compared to the baseline.

2 Related Work

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2.1 Eligibility Criteria Generation and Recommendation.

Over the past decade, various methods have been proposed to facilitate EC design. Trial2Vec (Wang and Sun, 2022) introduced a trial-level representation using contrastive learning to recommend relevant clinical trials to researchers, providing a foundation for trial similarity assessment. Based on trial representation approaches, CReSE (Kim et al., 2024) applied contrastive learning and rephrasing strategies to recommend relevant EC for a given trial context, focusing on high semantic similarity. AutoTrial (Wang et al., 2023) generates EC using LLM, offering interpretability through explicit reasoning chains. However, it uses predefined categories, which can restrict adaptability in complex clinical trials and potentially omit key criteria. Autocriteria (Datta et al., 2024) uses prompting on GPT4 to extract granular EC from clinical trial documents.

2.2 LoRA and Supervised Fine-Tuning (SFT).

Adapting LLMs to specialized tasks such as clin-120 ical trial EC generation often requires fine-tuning 121 on domain-specific datasets. Low-rank adaptation 122 (LoRA) (XTuner Contributors, 2023; Hu et al., 123 2021) has been applied in similar biomedical tasks 124 by efficiently integrating domain knowledge into 125 pre-trained models (Liao et al., 2024). Similarly, 126 127 supervised fine-tuning (SFT) has been employed in applications such as automated medical report gen-128 eration (Guo et al., 2024). However, while LoRA 129 and SFT have demonstrated significant efficacy in 130 these specialized tasks, they typically lack retrieval 131

strategies and do not generate domain-specific outputs, such as a complete set of EC.

2.3 Retrieval-Augmented Fine-Tuning (RAFT).

RAFT (Zhang et al., 2024) techniques have shown promise across various domains, including biomedical tasks, by simulating an "open-book" scenario in which a model can consult relevant external documents during both training and inference. Traditionally, RAFT involves providing the model with a mixture of "golden" and "distractor" retrieved texts, enabling it to learn when and how to utilize external information. However, RAFT methods often focus on short-form QA tasks rather than producing outputs such as fully articulated sets of EC.

2.4 Contributions of EC-RAFT.

While approaches such as AutoTrial (Wang et al., 2023), CReSE (Kim et al., 2024), or RAG-based pipelines have advanced the field, they each exhibit drawbacks. AutoTrial's category-based system may miss nuanced criteria critical for complex or adaptive trial designs. CReSE's strong clustering and recommendation focus lacks a mechanism for generating complete sets of EC. Standard RAFT-based pipelines (Zhang et al., 2024) often emphasize classification or short-form QA tasks, leaving the generation and evaluation of elaborate clinical EC largely unexplored. EC-RAFT integrates retrieval-augmented fine-tuning with synthesized chain-of-thought reasoning to generate a single structured set of inclusion and exclusion criteria to address these limitations. EC-RAFT provides a flexible and comprehensive solution for automated EC generation in complex trial contexts by bypassing the need for category-dependent generation and leveraging domain-specific retrieval as a backbone.

3 Methods

In this section, we introduce our approach, which leverages clinical trial data from ClinicalTrials.gov and integrates state-of-the-art techniques in embedding, retrieval, and fine-tuning to automate the generation of EC (Figure 1). We then describe the experiments designed to evaluate our system.

3.1 ClinicalTrials.gov Dataset

We collected 267,347 clinical trials from Clinical-Trials.gov, covering 2000 to 2024. To facilitate analysis, we split these trials into three datasets:



Figure 1: **Overview of the EC-RAFT pipeline. A.** Retrieve relevant trials and their EC (**D**) for the trial of interest (**X**) using SciNCL embeddings, then combine them with the desired EC (**y**) to generate intermediate reasoning steps (**R**). **B.** Fine-tune the model to generate a single response that includes both reasoning and final eligibility. **C.** Evaluate using two approaches: (1) **BERTScore** (Zhang et al., 2020) for semantic similarity, and (2) **LLM-Guided Evaluation** for clinical relevance of matched EC pairs.

213,877 trials for training, 26,735 trials for validation, and 26,735 trials for testing (Table 1). The training, validation, and test set contains around 168.4k, 20.9k, 21.1k interventional and 45.4k, 5.8k, 5.6k observational trials respectively. The training data contain 1.25M interventional trials with an average of 4.98 ± 5.11 inclusions and 7.46 ± 7.05 exclusions per trial and 137k observational trials with an average of 3.02 ± 2.85 inclusions and 3.44 ± 3.68 exclusions per trial.

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Our dataset consists of three primary sections: title, description, and EC. The description section includes a brief summary, a detailed description of the trial, and intervention details, including the type, name, description, and alternative names of the interventions involved. The EC section, which extracts from eligibilityModule within the protocolSection, contains key participant criteria, including both structured fields and free-text criteria. eligibilityCriteria within eligibilityModule section provides key eligibility details, including inclusion and exclusion criteria. While most trials specify age and gender requirements within the eligibilityCriteria section, some studies omit explicit references to these factors. Instead, these details are provided in dedicated fields within the same module: sex for gender information, minimumAge and maximumAge for age ranges, and healthyVolunteers for whether healthy volunteers are accepted. We extracted and processed these fields from both structured metadata and free-text EC to ensure that all EC are included.

3.2 Data Embedding and Retrieval

The first step involves obtaining comprehensive clinical trial data, including titles, descriptions, and eligibility criteria, from ClinicalTrials.gov (Figure 1A). We employ the SciNCL embedding model (Ostendorff et al., 2022) to embed clinical trials, which are subsequently retrieved to generate intermediate steps (\mathbf{R}) . The rationale for selecting SciNCL is its ability to embed semantics in domainspecific text. After embedding, we retrieve relevant trials and their EC (D) and Trial Information (X) using Euclidean distance. Importantly, only the training split was embedded. During testing and evaluation, we retrieved trials exclusively from the embedded training split. Our experiments vary the relevant trials (top-N) from N = 1 to 5 for generating the intermediate step (\mathbf{R}) (Section 5.3).

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3.3 Intermediate Steps Generation

In EC-RAFT, the generation of intermediate reasoning steps (**R**) plays a pivotal role in creating a structured pathway for training models. This process begins by integrating the retrieved trial information (**D**) which includes the title, description and ECs, the trial-of-interest information (**X**), consisting of its title and description, and the desired eligibility criteria (**y**) for the target study (Figure 1A). The **D** is retrieved from the vector database using **X**'s title and description while filtering out **X** out of retrieved documents, with different top-*N* values applied based on the experimental configuration. The desired eligibility criteria (**y**) serve as a *hint* that guides the LLM in breaking down

Statistic	Train (N = 213,877)		Validation (N = 26,735)		Test (N = 26,735)	
	Interventional	Observational	Interventional	Observational	Interventional	Observational
Number of Clinical Trials	168,429	45,448	20,928	5,807	21,129	5,606
Total Inclusion Criteria	838,948	137,234	103,910	17,531	103,982	16,990
Total Exclusion Criteria	1,256,242	156,298	154,896	20,470	156,212	19,000
Mean Inclusion Criteria per Trial (± SD)	4.98 ± 5.11	3.02 ± 2.85	4.97 ± 5.04	3.02 ± 2.77	4.92 ± 5.01	3.03 ± 2.99
Mean Exclusion Criteria per Trial (± SD)	7.46 ± 7.05	3.44 ± 3.68	7.40 ± 7.00	3.53 ± 3.64	7.39 ± 7.05	3.39 ± 3.61

Table 1: Statistics of clinical trials and EC. We calculate an average and a standard deviation of the number of EC of interventional and observational trials as these study types differ in their structure, particularly in the number of exclusion criteria.

each criterion, connecting them to evidence derived
from retrieved studies (D) and the study information (X).

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The primary objective is to generate intermediate reasoning steps (\mathbf{R}) that justify how each eligibility criterion (\mathbf{y}) is logically constructed and justified based on the retrieved trials (\mathbf{D}) and the target trial information (\mathbf{X}). The process can be written as:

$$\mathbf{D} + \mathbf{X} + [\text{Hint} : \mathbf{y}] \to \mathbf{R}$$
(1)

These intermediate steps will later be used in the fine-tuning steps formulated in (2). (see 3.4 for more details). These intermediate steps allow the model to learn how to derive eligibility criteria (y) from trial information (X) and retrieved studies information (D).

Including retrieved trials as part of the input provides the LLM with domain-specific examples, offering insights into established clinical practices. These examples enable the model to identify patterns and infer appropriate criteria for the target study. However, discrepancies may arise when the desired EC conflict with information from the retrieved trials. For instance, a retrieved trial might exclude patients with mild hypertension, whereas the target study explicitly includes them. In such cases, the LLM is tasked with identifying and articulating these conflicts, justifying deviations from established norms.

This conflict-resolution mechanism aims to ensure that the generated eligibility criteria (\hat{y}) are likely to be both contextually relevant and aligned with the specific goals of the target study, even when they may diverge from traditional practices. Our experiments explore the use of models including Gemini-1.5-flash-002 (Gemini Team, 2024) and Llama-3.1-8b-instruct (Grattafiori et al., 2024) to synthesize intermediate steps (**R**).

3.4 RAFT for Generating EC

RAFT in EC-RAFT enhances the model's ability to generate eligibility criteria (y) by leveraging relevant context retrieved from clinical trial data (D). Unlike traditional RAFT methods (Zhang et al., 2024) that classify documents as golden or distractors, EC-RAFT utilizes all retrieved trials holistically to account for varying levels of relevance. This ensures that the model is informed by diverse clinical contexts during fine-tuning. In this step, we utilized Llama-3.1-8b-instruct as a base model for supervised fine-tuning. We utilize Low-Rank Adaptation (LoRA) training techniques for cost efficiency. This fine-tuning process is structured as follows:

$$\mathbf{D} + \mathbf{X} \to \mathbf{R} + \mathbf{y} \tag{2}$$

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This approach aligns the model's training process with real-world scenarios, allowing it to learn directly from domain-specific documents in an open-book setting (Zhang et al., 2024). By integrating reasoning steps (\mathbf{R}), the model is encouraged to generate both eligibility criteria (\mathbf{y}) and output logical intermediate steps generated in the section above.

3.5 Generation of Eligibility Criteria

During inference, the fine-tuned model inputs the target trial's title and description (Figure 1B). It retrieves relevant trials from the vector database and uses the combined information to generate a complete set of EC. The output includes how eligibility criteria are derived ($\hat{\mathbf{R}}$) and the whole set of predicted eligibility criteria ($\hat{\mathbf{y}}$). Similar to the fine-tuning process, we can write this as:

$$\mathbf{D} + \mathbf{X} \to \hat{\mathbf{R}} + \hat{\mathbf{y}}$$
 (3)

We generate both the reasoning path and the predicted criteria. This allows the model to produce a reasoning process before predicting EC, which

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may improve results compared to direct inference(Wu et al., 2024).

To evaluate the effectiveness of our approach, we compare the performance of EC-RAFT with zeroshot inference from Llama-3.1-8b-instruct and Gemini-1.5-flash. We also vary the number of top-N during the generation of $\hat{\mathbf{R}}$ to evaluate its performance across different numbers of retrieved documents (**D**).

4 Evaluation

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Due to the challenging nature of semi-structured Eligibility Criteria, we employ three metrics to compare our predicted output (\hat{y}) with the ground truth (y) to measure: 1) **BERTScore** for overall semantic similarity, and 2) **LLM-Guided evaluation** which only evaluate the matched pair, **Pair-BERTScore**, identified by LLMs and utilize **LLM-as-a-Judge** to judge capability to assess clinical relevance for each matched pair.

4.1 BERTScore

We utilize BERTScore (Zhang et al., 2020) with the DistilBERT (uncased) (Sanh et al., 2020) model to assess the semantic similarity between the desired and predicted EC. BERTScore evaluates alignment based on token-level matches between the reference and predicted criteria, weighting these matches by their contextual embeddings to produce a similarity score. However, BERTScore may overestimate similarity due to the semi-structured nature of EC and may fail to distinguish logical inversions between inclusion and exclusion criteria.

4.2 LLM-Guided Evaluation

We propose an LLM-guided evaluation pipeline to assess how well-generated EC aligns with their corresponding reference criteria. This pipeline combines (1) **Pairing-and-scoring step** matching EC and calculating Pair-BERTScore (Section 4.2.1) and (2) **An additional match score** using an LLMas-a-Judge (Section 4.2.2). Below, we provide a general overview of the pipeline, followed by the unique details of each metric.

 Initial Evaluation We use Gemini-1.5-flash-002 to identify the most semantically and clinically relevant predicted criterion for each reference criterion. The model matches each reference criterion with the most pertinent predicted criterion, regardless of order, ensuring that all potential matches are considered. This process captures nuanced relationships between reference and predicted EC by explicitly accounting for inclusion-exclusion inversions, clinical parameters, and eligibility thresholds. The evaluation is generated in free-text format, prioritizing matching accuracy and judgment without enforcing a structured response, which could hinder accuracy (Tam et al., 2024). The evaluation prompt is provided in Figure A.

2. **Structured Output** We use watt-tool-8B's (watt-ai, 2023) structured response functionality to convert free-text evaluations into a JSON schema, ensuring consistency for accuracy calculations (Figure B). We utilized watt-tool-8B due to its state-of-the-art performance in tool-calling despite its size (Yan et al., 2024).

4.2.1 Pair-BERTScore

After getting the structured pairs of inclusion and exclusion, we calculate semantic similarity using BERTScore (Fig 2). This process enhances evaluation accuracy by removing any inflated scores that may arise from structural similarities. Note that Pair-BERTScore only accounts for the paired EC but not the excess generation of predicted criteria.

4.2.2 LLM-as-a-Judge

While Pair-BERTS core measures semantic similarity, it may fail to capture clinically significant distinctions between desired and predicted eligibility criteria $(\mathbf{y}, \hat{\mathbf{y}})$. To address this, we introduce LLMas-a-Judge, which evaluates the clinical and logical alignment between predicted and reference EC. For each matched EC pair, Gemini-1.5-flash also assigns a clinical relevance score (0-3) based on the degree of alignment, where higher scores indicate more substantial clinical similarity (Figure A). We calculate the mean of the judge's score to measure how well the generated EC $(\hat{\mathbf{y}})$ align with the desired EC (\mathbf{y}) .

4.2.3 Precision-Recall

Similar to Pair-BERTScore, the judge's score does not account for the excess EC generated. Thus, we also computed precision and recall to quantitatively measure the agreement between predicted and reference EC as follows



Figure 2: **LLM-guided Evaluation Metrics.** We align only generated EC, with corresponding reference EC and compute precision, recall, pair-BERTScore, and judge score. **Note** that in the actual pipeline, we also instruct the model to reason before evaluating each judge score (Fig. A, B)

(4)

$$Precision = \frac{N_M}{N_P}, Recall = \frac{N_M}{N_R},$$

where N_M represents the number of matched reference criteria with a positive match score (match_score > 0), N_R denotes the total number of reference criteria, and N_P is the total number of predicted criteria after de-duplication and filtering. De-duplication removes the exact predicted EC or a part of the same EC.

5 Results

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5.1 Comparison with zero-shot baselines

Using the ClinicalTrials.gov test split, we com-423 pare EC-RAFT performance against two zero-424 shot baselines: Llama-3.1-8B-Instruct and 425 Gemini-1.5-flash. As shown in Table 2, 426 EC-RAFT achieves a BERTScore of 86.35, 4.93 427 higher than base model Llama-3.1-8B-Instruct 428 and 4.17 higher than Gemini-1.5-flash which 429 is a larger model (Table 2). This indicates im-430 proved overall semantic similarity between the gen-431 erated and reference eligibility criteria. Regarding 432 clinical relevance, EC-RAFT with Gemini's R ob-433 tains the highest precision and mean judge score, 434 along with a superior mean Pair-BERTScore. This 435 means that EC-RAFT can generate precise EC to 436 437 the referenced EC. Although Gemini-1.5-flash registers a slightly higher recall, this advantage 438 comes at the expense of precision-likely due 439 to its tendency to generate excess criteria. On 440 top of that, our model was self-improved by 441

using base model Llama-3.1-8B-Instruct to generate **R** that could match the performance of Gemini-1.5-flash in some areas. Our results underscore the effectiveness of incorporating retrieval-augmented fine-tuning with intermediate reasoning steps, as it enables the model to generate eligibility criteria that are both semantically and clinically relevant.

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5.2 Effect of Larger model Intermediate steps

we want to see if reasoning steps Here. can affect the fine-tuned performance of EC-RAFT. We compare two variations differing in the model used to generate intermediate reasoning steps (R): Llama-3.1-8B-Instruct and Gemini-1.5-flash. As shown in Table 2, both approaches significantly improve BERTScore over the baselines, with EC-RAFT using Llama-3.1-8B-Instruct achieving a slightly higher BERTScore than the Gemini-based variant. However, EC-RAFT with Gemini-1.5-flash exhibits superior overall performance across LLMguided evaluations, achieving the highest precision, recall, mean Pair-BERTScore, and mean judge score, suggesting that its generated criteria are more clinically aligned. These results highlight the impact of selecting a strong LLM for generating intermediate reasoning steps, reinforcing that larger models like Gemini-1.5-flash can improve the accuracy and clinical relevance of EC generation.

5.3 Effect of LoRA hyper-parameters

LoRA (Low-Rank Adaptation) enables efficient fine-tuning by introducing trainable low-rank up-

Model	BERTScore †	LLM-guided Evaluations			
		Precision ↑	Recall †	Mean Pair-BERTScore \uparrow	Mean Judge Score ↑
Llama-3.1-8B-Instruct	81.42	77.16	67.63	51.95	1.3097
Gemini-1.5-flash	82.18	72.47	78.34	63.66	1.6004
EC-RAFT (R from Llama-3.1-8B-Instruct)	86.35	72.55	66.92	61.20	1.5932
EC-RAFT (R from Gemini-1.5-flash)	86.23	78.84	75.89	67.76	1.7150

Table 2: Comparison between EC-RAFT and baselines (Zero-shot)

dates. We evaluate the impact of Rank (r) and Scaling Factor (α) on Eligibility Criteria generation using BERTScore and LLM-guided evaluations. Results in Table 3 show slightly better in BERTScore, precision, and judge score when increasing r from 64 to 128 and α from 16 to 64, while recall remains stable, indicating that increasing LoRA's rank does not significantly enhance EC generation \hat{y} .

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5.4 Effect of top-N retrieval

We evaluate EC-RAFT with different top-N settings to examine the impact of retrieved documents on eligibility criteria generation. We generate **R** using L1ama-3.1-8B-Instruct by varying N retrieved documents. Table 4 shows that increasing N initially improves performance. BERTScore peaks at top-N of 4 before stabilizing, and precision follows a similar trend, suggesting excess documents may introduce noise. Recall remains stable with minor fluctuations, while Mean Pair-BERTScore and Mean Judge Score show slight variations. Overall, retrieving around four relevant documents provides modest benefits, but the overall impact remains limited.

5.5 Qualitative and Error Analysis

We sample a clinical trial on stroke and generate EC using EC-RAFT and Gemini-1.5-flash (Table 5). We found that EC from EC-RAFT are closely matches the reference in age and thrombectomy eligibility but omits intracranial vertebral artery involvement. Meanwhile, Gemini-1.5-flash are more restrictive, requiring prior endovascular therapy and a strict 90-day follow-up. It also excludes patients with a history of stroke/TIA and severe co-morbidities, further reducing eligibility.

Overall, EC-RAFT tracks the reference more closely, while Gemini-1.5-Flash generates a more lengthy EC, having higher recall but lower precision. This highlights the trade-off between precision and recall in automated EC generation.

6 Conclusion

In this work, we introduced EC-RAFT, a framework that leverages retrieval-augmented fine-tuning and synthesized intermediate reasoning to automate the generation of clinical trial eligibility criteria. EC-RAFT generates structured, robust, and clinically relevant eligibility criteria directly from trial descriptions. Our experiments on a largescale ClinicalTrials.gov dataset demonstrate that EC-RAFT outperforms zero-shot baselines despite being much smaller in model size, achieving higher BERTScores and clinical alignment as evidenced by LLM-guided evaluations. Notably, incorporating intermediate reasoning-proves instrumental in enhancing both the precision and overall quality of the output. While challenges remain, EC-RAFT represents a significant step towards automating the complex process of clinical trial design. Future work will refine the intermediate steps generation process and scale up model size and compute for better performance.

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Limitations

While EC-RAFT demonstrates promising results in automated EC generation, several limitations should be acknowledged. First, our approach relies on LLMs, which can produce plausible but inaccurate or inconsistent criteria that require human expert validation. Second, training data comes primarily from public clinical trial registries, which may not fully represent the diversity of trial designs or specialized medical domains. Third, our evaluation metrics (BERTScore and LLM-as-a-Judge) provide computational approximations of generated EC but may not fully capture clinical relevance or practical applicability. Specifically, the latter may suffer from inconsistent responses and bias. The performance may vary between different medical specialties and trial types, particularly for rare diseases or novel therapeutic approaches where training data are limited. Future work should address these limitations through expanded train-

Model	BERTScore ↑	LLM-guided Evaluations			
		Precision \uparrow	Recall ↑	Mean Judge Score \uparrow	Mean Pair-BERTScore ↑
EC-RAFT ($r = 64, \alpha = 16$)	86.1712	70.69	67.76	1.6039	61.73
EC-RAFT ($r = 128, \alpha = 64$)	86.2426	71.08	67.70	1.6046	61.76

Table 3: Comparison between different LoRA configuration (Rank r and Alpha α) with top-N = 2

Top- N	BERTScore ↑	LLM-guided Evaluations			
		Precision \uparrow	Recall ↑	Mean Pair-BERTScore ↑	Mean Judge Score ↑
1	0.861739	70.73	67.70	61.69	1.6003
2	0.861712	70.67	67.76	61.76	1.6039
3	0.863107	72.05	66.82	61.10	1.5897
4	0.863507	72.55	66.92	61.21	1.5932
5	0.863454	72.47	67.12	61.37	1.5981

Table 4: Comparison of different top-N configurations for EC-RAFT

Multicentre ObseRvational Study in Posterior mecHanical thrombEctomy: conscioUs Sedation vs General aneSthesia: MORPHEUS STROKE REGISTRY (NCT05082896)

Deference Criterie		Comini 1 5 flach
Reference Criteria Inclusion Criteria: Subject aged ≥ 18 years Evidence of posterior arterial occlusion Patients eligible for mechanical thrombectomy Exclusion Criteria: Patient's refusal to participate Arterial recanalization prior to endovascular thrombectomy Sex: ALL Ages: Minimum Age: 18 Years Age Group (Child: birth-17, Adult: 18-64, Older Adult: 65+): OLDER_ADULT, ADULT Accepts Healthy Volunteers: No 	 EC-RAFT Inclusion Criteria: Age ≥ 18 years Acute ischemic stroke with occlusion of the basilar artery or P1 segment of the posterior cerebral artery Treatment with endovascular therapy Treatment initiation within 24 hours after stroke onset Informed consent from patient or legal representative Exclusion Criteria: Pre-existing conditions with significant impact on functional outcome (pre-stroke mRS > 1) Intracranial hemorrhage Pregnancy Participation in other clinical trials Sex: ALL Ages: Age Group: OLDER_ADULT, ADULT Accepts Healthy Volunteers: No 	Gemini-1.5-flash Inclusion Criteria: - Confirmed diagnosis of ischemic stroke due to posterior circulation large vessel occlusion (PC LVO) demonstrated by imaging (CTA or MRA). - Underwent endovascular therapy (EVT) for the PC LVO. - Received either conscious sedation OR general anesthesia during EVT. - Able to complete at least 90 days of follow-up. - Provided informed consent to partici- pate in the study. Exclusion Criteria: - Contraindications to endovascular therapy (EVT). - Ischemic stroke not due to PC LVO (e.g., anterior circulation stroke). - Hemorrhagic stroke. - Incomplete follow-up data at 90 days post-EVT. - History of previous stroke or Tran- sient Ischemic Attack (TIA). - Severe co-morbidities that may con- found outcome assessment. Sex: ALL Ages: - Minimum Age: 18 Years - Maximum Age: 90 Years - Age Group: ADULT,
		 Maximum Age: 90 Years Age Group: ADULT, OLDER_ADULT Accepts Healthy Volunteers: No

Table 5: Comparison of NCT05082896's Eligibility Criteria: Reference, EC-RAFT, and Gemini-1.5-flash

ing datasets, domain-specific fine-tuning, and more rigorous clinical validation processes.

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A Appendix

This appendix provides documentation of the prompts used in EC-RAFT for EC generation. The following sections detail the exact prompts, implementation notes, and practices developed during our research.

A.1 LLM-guided Evaluation Prompt

To ensure a standardized and clinically grounded evaluation, we adapt the scoring methodology from the Evaluation Guideline for Assessing Clinical Relevance between an EC Pair (Su et al., 2023).

666	The adapted framework categorizes EC similarity
667	into four levels:

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- Clinical relevance $3 \rightarrow$ Clinically identical ECs.
 - Clinical relevance 2 → Strongly relevant due to factors like disease progression or epidemiology.
 - Clinical relevance 1 → Loosely relevant due to general treatment plans, disease progression, or epidemiological factors.
 - Clinical relevance 0 → Irrelevant from a clinical perspective.

The actual prompt can be found in figure A. The matched EC pairs and their scores can be found in figure B.

A.2 LLM-guided Evaluation JSON Schema

After the initial evaluation, we utilize watt-tool-8B to convert the free-text evaluation into a structured JSON format for quantitative analysis in Section 4.2. Since each reference criterion can match multiple predicted criteria, the predicted values are stored as a list of strings to accommodate the one-to-many relationship.

A.3 Implementation Details & Computational Cost

Our default LoRA configuration includes a Rank of 64, α of 16, and dropout of 0.1, except in section 5.3. We train on four NVIDIA A100 GPUs, requiring 192 to 470 GPU-hours per model, depending on the top-N value, totaling around 2,200 GPU-hours across this paper. Our best-performing model is trained in 380 hours, costing approximately 452.20 USD at a market rate of 1.19 USD per GPU-hour.

LLM-guided Evaluation Prompt

Please evaluate the clinical relevance of the following two eligibility criteria on a 4-point scale (0–3). Below is an example of a clinical situation by clinical relevance score and the corresponding EC pair.

Clinical relevance 3: The two eligibility criteria are essentially identical clinically. *Examples*:

- EC1: "[exclusion] serum albumin is 2.4 g/dL or less"
- EC2: "[inclusion] serum albumin is 2.4 g/dL or more"
- EC1: "Minimum Age : 18 Years" EC2: "Minimum Age : 18 Years"

Clinical relevance 2: The two eligibility criteria have strong relevance due to factors such as disease progression or epidemiology.

Example: ...omitted for brevity... ...omitted for brevity... Evaluation Process

For each reference criterion, compare it to the relevant predicted criteria. If no relevant predicted criterion exists, state this explicitly. The evaluation process is as follows:

- 1. Recite the reference exact criterion and state explicitly if it is from [inclusion] or [exclusion].
- 2. Search the predicted criteria list to identify the relevant matches, regardless of order (commaseparated), and explicitly state which part of the predicted criteria each match comes from ([inclusion], [exclusion], [age], [sex], [accepts healthy volunteers]).
- 3. Recite the reference **Sex**, **Ages**, and **Accepts Healthy Volunteers** one at a time and compare them with the relevant predicted values.
- 4. Provide a reason explaining how the criteria match or differ.
- 5. Assign a match score (0–3) based on the clinical relevance of the predicted criterion to the reference criterion.
- 6. If no predicted criterion matches the reference, state that explicitly and assign a score of 0.

At the end of the evaluation, please provide:

- Unmatched Predicted Criteria:
 - Unmatched Predicted Inclusion Criteria: List all predicted inclusion criteria that were not matched to any reference criteria (relevance score = 0). No explanation is needed—just list them (comma-separated).
 - Unmatched Predicted Exclusion Criteria: List all predicted exclusion criteria ...Same as before, omitted for brevity...

Figure A: LLM-guided Evaluation Prompt

```
LLM-guided Evaluation JSON Schema
{
  "inclusion_criteria": [
    {
    "reference": "criteria",
    "predicted": ["match"],
    " "cyplanation",
       "reason": "explanation",
       "match_score": 3
    }
  ],
"exclusion_criteria": [
    "predicted": ["match"],
"reason": "explanation",
       "match_score": 2
    }
  ],
  "sex": {
    "reference": "value",
    "predicted": [""],
"reason": "explanation",
    "match_score": 0
  },
"age": {
    "reference": "value",
"predicted": ["match"],
    "reason": "explanation",
    "match_score": 2
  "reason": "explanation",
     "match_score": 1
  },
"unmatched_predicted_criteria": {
    disted_inclusion
     "unmatched_predicted_inclusion
    _criteria": ["unmatched"],
"unmatched_predicted_exclusion
     _criteria": ["unmatched"]
  }
}
```

Figure B: JSON Schema parsed from free-text judge response: