

Dear ARR Reviewer,

We write to kindly request you to consider the following where we address the issues raised in the previous round of reviews to this paper. Text in blue are the additions to the paper. We are also attaching a color-coded version of our paper. Text in blue are either new text or new sections.

Reviewer KnHk

1. We respectfully disagree. The innovation in the paper is the fact that we show that foundational models even those specifically trained in the domain do not work very well for some closed-domain question answering in the medical area. We then suggest how one can extract entities, use them to create prompts, generate synthetic text and retrain existing models in order to improve these models' performance. We demonstrate their efficacy and superiority on two closed-domain medical question-answering benchmarks, COVID-QA and RadQA. The method is not obvious and the experiments generate knowledge about our proposed method working better than any existing method. There lies the innovation.

With respect to generality, the point is taken that it would be good if we can run these on other datasets. However, that argument can be made against any paper. We have a limited number of resources both with respect to human time and computational time. We also have limitations on the space we are allowed to write a paper. In this space and with these resources we have addressed the issues to the best of our ability.

Is this the end of the story? No. More work is needed and our group is already working on extending this. Does this paper advance science? A resounding yes. We show that at least in these two domains using entity-based corpora generation works improve the state-of-the-art. We strongly believe the scientific community deserves to know this result. And, we are willing to make whatever specific edits, suggestions etc. your committee may have to improve the paper but a generic "not enough" as raised by the previous review is hard to address.

We highlight our contributions at the end of the Introduction section,

Line 111: In summary, our contributions are,

Proposing a pipeline for generating customized pre-training data for closed (in our case medical) domains.

Demonstrating the effectiveness of synthetic data for achieving sizable gains with reduced memory footprint.

Showing the benefits of creative prompting and dataset awareness.

Benchmarking numerous pre-trained biomedical FMs on COVID-QA and RadQA and contrasting our models with them by showing their superiority.

2. We have put in a dedicated Related Work section and addressed all known works, to the best of our knowledge, in the area directly related to our efforts.

Line 124: 2 Related Work

In this section, we discuss prior work on decoder-based FMs for QA, highlight their limitations and describe recent knowledge distillation pipelines.

3. Our paper does not require customized prompts. We can use the same basic template where we use the entity name to instruct a FM to generate synthetic texts related to that prompt. If we customize the prompt, we may get slightly better results. I asked practitioners whether they would be interested in manually creating a 2-3 line prompt template for the fancy template and

the work is so minimal that all practitioners were ready to do so. We can also automate this process with a heuristic solution. This should not be an impediment in the real-world to using our work nor would it take that much manual effort to set up a “fancy” prompt template (as defined in our paper).

4. We regret the unprofessional writing in the last draft. We have removed colloquial language, archaic phrasing, unnecessary unprofessional abbreviations and worked to the best of our ability given the time we had. We will do another full pass with a native speaker to edit the paper for the final camera-ready version should this paper be accepted.
5. We have tried to meticulously check every claim made in the paper in every line and either give citations or tone down the claims to make it clear that that is a probable conjecture. We still keep the conjecture in the hope that the intuition is useful for a follow-up work to verify them more rigorously. But, we agreed with the previous review and marked our conjectures more carefully.
6. We have tried to explain the methods and settings as much as we can given the space provided. We include the rest in the appendix due to space limitations. We will have a full-version of a technical report where we will report all parameters, etc. carefully to enable full reproduction.

Reviewer cmuw

1. Novelty – we claim that no prior work has done entity-based synthetic corpora generation and shown that it improves domain-specific foundation models on our closed-domain benchmarks. We would welcome counterexamples that disprove our claim; our best literature review did not find a paper that does that.
2. We have added real-life examples in the whole presentation.

Line 932: Appendix B – We refer readers to Figure 6 (top) to see an example of such kind of report.

3. We show improvements even in a non data-leak setting at the end. Note here that a “data-leak setting” is nuanced here. For example, if I have a student who does badly in a test I give, I can give the student a set of entities to read up on and then give another test. In this case, although I have given the student the list of entities to read up on, the second test is not really a seen test.

In fact, real practitioners may be fine giving us a set of diseases and entities of interest on which they want to ask questions and then we can generate our synthetic documents based on them and prepare for the unseen exam. Then, the practitioners can use the fine-tuned model to ask questions they seek to ask.

Nevertheless, as we should in our cross-validation experiments on COVID-QA, our method still performs better. Also, note that in RadQA, there is no data leak. So, the claim that our method works only in data-leak settings is flat out wrong.

Reviewer: rTgr

1. This may be an interesting solution but we do not claim that we have tried out all possible solutions. It would be another project or part of a new project. We do not know if that would work. Our claim is that our method of augmenting FMs work on our datasets that

we establish by our experiments. We have tried to make the claims we make in the paper more tight and accurate though.

2. We have tried to address the presentation issues.

TOP-Training: Target-Oriented Pretraining for Medical Extractive Question Answering

Anonymous ACL submission

Abstract

This work studies extractive question answering in the medical domain (Medical EQA). This problem casts two main challenges: i) domain specificity; most AI models lack the necessary knowledge; ii) extraction-based answering style, which restricts most autoregressive LLMs due to the potential hallucination concern. To handle those challenges, this work proposes TOP-Training, a target-oriented pretraining paradigm that stands out among all domain adaptation techniques with two desirable features: i) TOP-Training moves one step further than popular domain-oriented fine-tuning since it not only moves closer to the target domain, but also familiarizes itself with the target dataset, ii) it does not assume the existence of large set of unlabeled instances towards the target domain. Specifically, for a target medical EQA dataset, we extract its entities and leverage large language models (LLMs) to generate synthetic texts containing those entities; pretraining on this synthetic text data is shown better performance on the target medical EQA benchmarks. Overall, our contributions are threefold: i) TOP-Training, a new pretraining technique to effectively adapt LLMs to better solve a target problem; ii) TOP-Training has a wide application scope because it doesn't require the target problem to have a large set of unlabeled data; iii) Our experiments highlight the limitations of autoregressive LLMs, emphasizing TOP-Training as a means to unlock the true potential of bidirectional LLMs.

1 Introduction

EQA involves identifying a span of tokens in a passage to answer a given question. While language models such as BERT (Devlin et al., 2019), T5 (Raffel et al., 2020), and GPT-3 (Brown et al., 2020) have made remarkable leaps in this task, they suffer drastically when applied to domain-specific datasets, especially medical (Moradi et al., 2021).

We believe that the performance discrepancy is linked to the definition of a *domain*, i.e., the loose NLP equivalency of *domain = genre* or *thematic content*, which is quite restrictive, as argued by Plank (2016). Theoretically, a model trained on a specific theme should excel in tasks related to that subject matter. However, not all domain-specific models are equal as illustrated by the difference in performance of BioBERT (Lee et al., 2020) and PubMedBERT (Gu et al., 2021), even though both were trained on PubMed data (Tables 1,2). We suggest redefining *domain = [genre + dataset]*, and emphasizing the importance of tailoring the training data to not just the theme, but also on the target task dataset. This approach acknowledges that a *one-domain-model-to-rule-them-all* is not universally applicable, and that learning should also focus on concepts relevant to specific tasks.

Recent progress in generative FMs (Ye et al., 2023; Achiam et al., 2023) gives us reasons to believe that a solution to the closed-domain problem has been reached, particularly with ChatGPT, which has demonstrated strong performance, e.g., on the USMLE (Kung et al., 2023) exam. However, our expectations might be myopic as we (authors) demonstrate that decoder-only FMs struggle (Tables 1, 2) with specialized language, such as COVID-QA (Möller et al., 2020) and RadQA (Soni et al., 2022). Moreover, their autoregressive architecture is not well-suited for EQA as they are designed to *synthesize* new text rather than *extract* spans from given text (§ 5.1, 2.1). For sequences exceeding the model's maximum limit, they need to be divided into overlapping segments (pre-processing). Though this issue applies to both bi-directional and generative models, the former is more suitable due to its inherent capabilities (2.1). Furthermore, these models handle EQA as autoregressive language modelling rather than start/end token prediction, which can lead to either mismatch in generated/true answer tokens (§2.1) or even *hal-*

lucinated text (Ji et al., 2023).

To overcome these limitations, we propose distilling the knowledge from causal FMs into smaller MLM-style models, which are better suited for EQA. Our approach uses a causal FM to *generate synthetic corpora* tailored to a specific application and fine-tuning open-domain MLM models on this corpus. We adapt open-domain models instead of further training domain-centric ones (ex. Bio/SciBERT) (we did train the best performing biomedical model on our dataset, but saw no noticeable gains) to show the resource benefits of our pipeline. That is, without relying on gigabytes of domain-specific pre-training data, we can achieve respectable gains by honing our models on modest-sized corpora. Our results demonstrate the efficacy and the running time improvements with respect to existing domain-specific models.

We generate corpora instead of using existing text because: (a) it provides the *flexibility* to create content in a certain style, (b) some corpora can be *unavailable* for privacy reasons, ex. clinical diagnostic reports, or “internal” versions of corpora such as that used by Gururangan et al. (2020), and (c) our tests can be used to determine if the content produced by such FMs is *factually grounded* and able to teach student models specific writing styles.

In summary, our contributions are,

- Proposing a pipeline for generating customized pre-training data for closed (in our case medical) domains.
- Demonstrating the effectiveness of synthetic data for achieving sizable gains with reduced memory footprint.
- Showing the benefits of creative prompting and dataset awareness.
- Benchmarking numerous pre-trained biomedical FMs on COVID-QA and RadQA and contrasting our models with them by showing their superiority.

2 Related Work

In this section, we discuss prior work on decoder-based FMs for QA, highlight their limitations and describe recent knowledge distillation pipelines.

2.1 Autoregressive modeling for QA

On release, GPT-4 (Achiam et al., 2023) became the benchmark FM on almost all canonical NLP

tasks. Remarkably, Nori et al. (2023) proved that just by careful prompting, GPT-4 can achieve SOTA on various medical QA datasets. Initiatives such as Meditron-70B (Chen et al., 2023) became the next best thing to parallel GPT-4’s medical expertise. On closer inspection however, we see that the datasets on which these two models are applied are all multiple-choice style. Thus, their capability of handling EQA datasets, especially in the medical domain remains unexplored (cf. App. C).

Attempting to autoregressive FMs, Xu et al. (2021), through indirect supervision, train BART (Lewis et al., 2020) for span extraction by considering cross-attention weights as start/end token probabilities to align the generated and true spans. Noting that generating tokens is perhaps not the best strategy, Mallick et al. (2023) reframe the problem as generating *numeric indices* indicating either token or sentence-level spans. Innovative as these solutions are, we see two issues, i) marginal performance gains, and ii) only Mallick et al. (2023) use medical EQA dataset (Xu et al. (2021) focus only on open-domain) that too without much success - both points as compared to their encoder-based SOTA indicating that they are ineffective.

Luo et al. (2022) show that on average *encoders outperform decoders* in short-span detection with the added advantage of being better at *out-of-domain generalization* whereas decoder models can be comparably used in *long-context* EQA. Additionally, they show that the encoder alone from Seq2Seq models such as T5 follow a similar trend. While they do make the case for autoregressive FMs, later studies (Liu et al., 2023) demonstrate how newer instruction-tuned FMs such as MPT-30B-Instruct (Team et al., 2023) are *sensitive to the location of the gold span* Although they do not focus on encoders-alone, they do show how Seq2Seq models are relatively robust to such positional changes due to their encoders.

2.2 Knowledge Distillation

In the area of knowledge distillation, West et al. (2022) demonstrate how GPT3 can be utilized to create high-quality knowledge graphs via prompting. He et al. (2022) show how a GPT model can be used as a “teacher” to distil knowledge into a “student”. Similarly, Peris et al. (2022) use unlabelled task-relevant data and trained multilingual students with varying proportions of general/task-specific data and report the most gains using “only the downstream task’s unlabelled data”.

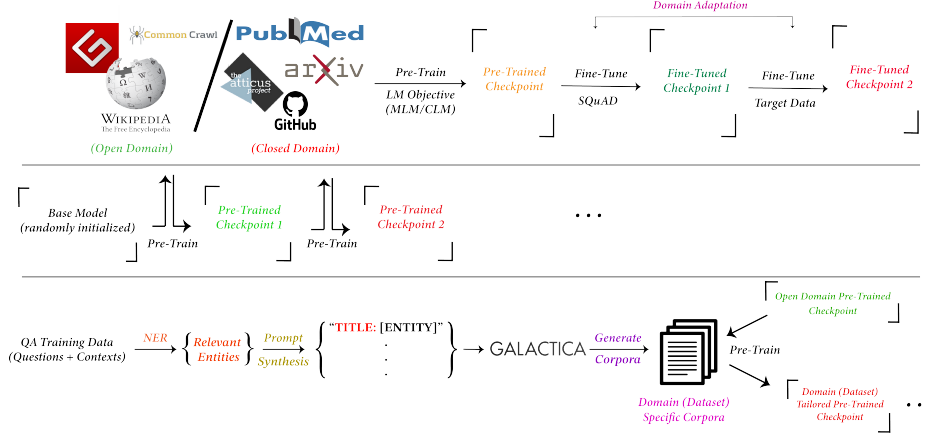


Figure 1: Pre-Training Pathways: From scratch (top); Extended (middle); **Targeted** (bottom; ours) | Note: We only show fine-tuning on EQA as it is the task of interest | The prompt handle is written in CAPITAL for emphasis.

Gururangan et al. (2020) introduced DAPT (Domain-Adaptive Pretraining) and TAPT (Task-Adaptive Pretraining), which share similarities with our work. DAPT involves extended pretraining on domain-specific corpora without labels, while TAPT focuses on pretraining on the unlabelled training set of the downstream task. Although they demonstrate the effectiveness of TAPT compared to DAPT, closed-domain datasets like COVID-QA typically lack a separate unlabelled training set and may not even have train/dev/test splits. Further, DAPT considers knowledge beyond what is necessary to the task data, whereas our approach confines training to only required concepts.

While the above techniques have achieved much success, they typically rely on high quantities of unlabelled corpora to yield useful results, thus raising the question: *What happens when we do not have enough “relevant” domain data, either in style or volume?* Inspired by the works of Gunasekar et al. (2023) and Zhou et al. (2023), who hearken the use of smaller yet *better quality* corpora, we introduce the notion of TOP-Training, which focuses on a specific subset of the domain, tailor-made for the ultimate downstream dataset.

3 Formulation of medical EQA

Each labeled data point consists of three elements:

- CONTEXT (\mathcal{C}): A piece of text in the medical domain that introduces the necessary information about a topic;
- QUESTION (\mathcal{Q}): A question sentence acquiring the information from CONTEXT. *It can be answerable or unanswerable.*

- ANSWER (\mathcal{A}): A consecutive span in CONTEXT that acts as the answer of QUESTION.

Typical datasets for medical EQA are split into *train*, *test* and optional *dev* sets.

Two main challenges for medical EQA problems: i) domain specificity; most AI models lack the necessary knowledge; ii) extraction-based answering style, which restricts most autoregressive LLMs due to the potential hallucination concern. The two challenges motivate us to propose TOP-Training accompanied with bi-directional LLMs for this particular problem.

4 TOP-Training

For the target closed-domain medical QA problem, our TOP-Training, depicted in Figure 1, first extract entities from the target dataset, then leverage an existing LLM to generate entity-related CONTEXT (Section 4.1), which later acts as the data for further tuning a bi-directional LLM for the target problem (Section 4.2).

4.1 LLM generates target-oriented CONTEXT

Entity collection from the target problem. In this work, we use entities in the target dataset as the connection between it with the newly generated synthetic data. To extract entities from the target EQA dataset,

First, we combine all the QUESTIONS and CONTEXTS from the *train* split of the medical EQA dataset. Next, we extract entities through Named Entity Recognition (NER) using spaCy¹. This step identifies roughly 47k and 11k entities in COVID-QA and RadQA, respectively.

¹With “en_core_sci_sm”.

Synthetic CONTEXT generation. Next, we create prompts for the identified entities to generate CONTEXTS mimicking the target datasets. This required studying the characteristics of the target datasets such as the text genre (full research articles in COVID-QA & radiology reports in RadQA), lengths, and relevant keywords.

Galactica, a decoder-based LLM pre-trained on a collection of text encompassing research articles, knowledge bases, code and even \LaTeX markup, is used to generate the synthetic data² with the following prompts.

- **Prompt for COVID-QA.** Since COVID-QA comprises research papers, based on this characteristic, we develop the following prompt for Galactica to generate pseudo research articles based on retrieved entities:

Title: [Entity]

where Title is the prompt handle/keyword and entity is the entity identified in the above stage.

- **Prompt for RadQA.** RadQA’s CONTEXTS are redacted radiology reports without any consistent format (Hartung et al., 2020). The *Findings* and *Impressions* sections are the most vital in a patient’s report (akin to the experiment and results section in a research paper). Inspired, we propose the following prompt:

Patient has Entity. FINDINGS
AND IMPRESSION

It is worth noting that Galactica had not been trained on radiology reports. Through this prompt, we synthesize *pseudo-reports* bypassing any privacy concerns. To maintain size parity between the two target datasets, five CONTEXTS are generated for each entity identified in RadQA, yielding around 55k (11k*5) total CONTEXTS, and one CONTEXT for each identified entity in COVID-QA, resulting a set of size 47K.

4.2 System pre-training on synthetic data

After generating CONTEXTS for each target data, we perform TOP-Training i.e., extended pre-training of BERT/RoBERTa on our generated corpus. In addition, TOP-Training is followed by two rounds of fine-tuning:

- The system will be first fine-tuned on the

SQUAD dataset to learn what EQA is in a generic domain;

- Subsequently, it is further fine-tuned on the *train* of each target data (either COVID-QA or RadQA), to solve the *test*.

5 Experiments

We focus on two datasets: COVID-QA, comprising 2,019 answerable QA pairs (no train/dev/test splits) sourced from CORD-19 (Wang et al., 2020), and RadQA, consisting of 6,148 QA pairs from radiology reports, with a train/dev/test split of 4,878/656/614. We experiment in two areas: baselines and TOP-Training, and provide mean and standard deviation scores over three random seeds for each.

5.1 Baselines & influencing factors

We consider 13 encoder models to each dataset taking into account which model made most sense to apply to a dataset. On COVID-QA, we applied models from checkpoints fine-tuned on SQuAD v1 while RadQA, containing unanswerable questions, was tackled with those fine-tuned on SQuAD v2 (Rajpurkar et al., 2018). For consistency, we utilized the *cased*, *base* version of each architecture when available. We use five-fold cross-validation to fine tune the models and report the results in Table 1. Results of models applied to the prescribed splits (RadQA) are presented in Table 2. The metrics used are exact match (EM), binary measure of whether the prediction and gold-standard spans are identical and F1, the harmonic mean of the number of shared words in the two spans with respect to the number of words in the prediction (precision) and with respect to the number of words in the gold-standard span (recall).

We selected Galactica-1.3B for consistency with our corpus generation experiments, MedL-LaMA(13 B) as a strong open-source medical checkpoint, and MedAlpaca(13 B) as a medical QA-specific LLaMA checkpoint. We measure the ability of the three decoder models to generate answers without fine-tuning, considering that decoders do not extract spans, but generate answers for comparison to gold-standard spans.

Following Yue et al. (2021), each sample is formatted as Question:<question_text> Context:<part_of_context> Answer:.. Due to the large size of COVID-QA contexts, they were segmented as they exceeded the maximum

²Other generative models such as BLOOM (Scao et al., 2022) and PubMedGPT performed worse in our experiments.

sequence length of each model (2,048 tokens). We report overall EM/F1 on each dataset and average best EM/F1 (parenthesis in Table 1) from each Q+C+A chunk for COVID-QA (N/A for RadQA since the context size was much smaller than the models’ maximum input length).

Corpus Size. We investigated the impact of synthetic dataset size on downstream performance in COVID-QA. We examined the effects of generating one and 10 contexts per entity.

Context Length. The average context length for COVID-QA is 6k tokens, and Galactica has a maximum context size of 2k, resulting in a misalignment between the synthetic corpus and the target dataset. We cannot increase the context size of Galactica. Training it from scratch with architectural changes is infeasible for us. Thus, we explore the impact of sequence length in the synthetic corpus by limiting the records to only 1k tokens. We cannot determine if *longer* sequences are *beneficial*; we can evaluate if *shorter* ones are *detrimental*.

Prompting Style. We explore the use of two different prompts when encouraging Galactica to generate *pseudo* radiology reports - “Patient has [entity]. FINDINGS AND IMPRESSION” (*fancy prompt*), and simply “[entity]” (*normal prompt*).

Human-Generated Contexts. We establish a *Wikipedia* baseline alongside our domain-specific models to assess the influence of content and text structure during domain adaptation. Additionally, Micallef et al. (2022) have shown that mBERTu Wiki, pre-trained on Maltese Wikipedia data, surpassed the performance of mBERT, thereby proving to be a competitive baseline. For each entity, we query Wikipedia and retrieve the complete page associated with the top search result. The number of entities available for this baseline is much smaller than that in our approach since most of the entities do not exist in Wikipedia due to either being extremely esoteric, e.g., *pulmonary parenchymal infiltrate* or improperly formed, e.g., *Bao &*.

6 Results & Discussion

We discuss the results of existing baselines (Tables 1 & 2) and TOP-Training (Tables 3 and 4). The last 3 rows of Tables 1 and 2 provide zero-

shot performance of our chosen decoder models on RadQA and COVID-QA respectively. We do not perform multiple trials here as the extremely poor performance would not benefit from additional runs. Granted they were not fine-tuned on our datasets, their size, pre-training data coverage and reported zero-shot performance on related datasets, should have allowed them to at least perform on par or better than open-domain BERT/RoBERTa. Overall, we see that MedAlpaca seems to be the “best” among the three for RadQA and only marginally poorer in terms of F1 for COVID-QA probably since MedAlpaca is an instruction-tuned version of MedLLaMA. On COVID-QA, none of the models generated text in line with the gold standard (~0 EM) and only showed positive F1.

6.1 COVID-QA

We now discuss the results benchmarking trials on COVID-QA contrasting the methods.

6.1.1 Baselines

Our experiments demonstrate that a one-size-fits-all approach does not work always for domain adaptation. BioBERT and PubMedBERT were trained on similar corpora and yet score in the same range, indicating no clear winner. PubMedBERT that is trained from scratch using a custom vocabulary covering a range of medical jargon performs better.

SciBERT (+CORD-19) checkpoint, trained on CORD-19 articles, performs worse than regular SciBERT, suggesting potential issues in training choices or noisiness in the data. Notably, LUKE, trained solely on Wikipedia data, emerges as the best baseline model, possibly due to its entity-recognition pre-training objective, which aids in identifying relevant entities for QA tasks (Van Aken et al., 2019) and highlighting the need for entity representations in closed-domains. Models capable of handling longer context i.e. Longformer, BigBird and XLNet do not show marked improvements. XLNet, degrades completely on COVID-QA, potentially due to token permutation hindering its reasoning across large and conceptually dense contexts.

6.1.2 TOP-Training

Wiki Baseline: Fine-Tuning BERT on Wikipedia yields marginal improvement of 0.9% EM and 1.2% F1; RoBERTa shows a 3.7% increase in EM and a 1.3% increase in F1. Our Wikipedia corpus, while small, contains relevant information per-

Table 1: Baseline Bio Models (COVID-QA). ¹(Peng et al., 2019); ²(Yuan et al., 2022); ³(Beltagy et al., 2020); ⁴(Zaheer et al., 2020); ⁵(Yamada et al., 2020); **Blue = best/red = worst** scores overall; **bold = best decoder**

Model	Pre-Training Corpus	Corpus Size	EM	F1
BioBERT	PubMed	4.5B words	37.62 ± 0.18	65.73 ± 0.35
SciBERT	Semantic Scholar	3.2B words	37.52 ± 0.23	65.58 ± 0.18
SciBERT(+CORD-19)	Semantic Scholar + CORD-19	3.2B words + 20GB	35.61 ± 0.30	63.60 ± 0.59
PubMedBERT	PubMed	3.1B words / 21GB	39.87 ± 0.74	68.47 ± 0.13
BlueBERT ¹	PubMed + MIMIC	4.5B words	27.35 ± 0.30	52.18 ± 0.40
CODER ²	Unified Medical Language System	NA	39.33 ± 0.47	67.01 ± 0.30
Longformer ³	Books + Wiki + RealNews + Stories	6.5B tokens	37.79 ± 0.39	66.58 ± 0.22
BigBird ⁴	Books + CC-News + Stories + Wiki	160GB (same as RoBERTa)	32.79 ± 0.13	60.06 ± 0.41
LUKE ⁵	Wikipedia	3.5B words	41.01 ± 0.30	68.23 ± 0.19
XLNET	BooksCorpus + Wikipedia + Giga5 + ClueWeb 2012-B + Common Crawl	32.89B words	2.45 ± 0.08	8.64 ± 0.19
Galactica	c.f. section 4	106B tokens	0 (0)	5.01 (11.11)
MedLLaMA	Medical Corpora	NA	0 (0)	5.81 (12.79)
MedAlpaca	Medical Meadow	NA	0.03 (0.2)	5.21 (12.73)

Table 2: Baseline Bio Models (RadQA). H(F1): HasAns_F1, H(EM): HasAns_EM; **Blue = best/red = worst** scores overall; **bold = best decoder**; ¹(Yan et al., 2022); ²(Alsentzer et al., 2019); ³(Gururangan et al., 2020)

Model	Corpus	Corpus Size	Dev				Test			
			EM	F1	H(EM)	H(F1)	EM	F1	H(EM)	H(F1)
BioBERT	PubMed	4.5B words	26.42 ± 0.49	44.26 ± 0.09	40.79 ± 0.76	68.31 ± 0.14	49.95 ± 1.08	63.32 ± 0.40	45.65 ± 1.21	63.50 ± 0.57
SciBERT	Semantic Scholar	3.2B words	27.03 ± 0.32	44.40 ± 0.06	41.65 ± 0.62	68.45 ± 0.22	53.04 ± 0.38	67.17 ± 0.73	48.62 ± 0.70	67.49 ± 1.02
PubMedBERT	PubMed	3.1B words / 21GB	31.45 ± 0.17	47.89 ± 0.46	48.40 ± 0.27	73.77 ± 0.62	54.07 ± 0.71	68.76 ± 0.22	49.49 ± 0.87	69.09 ± 0.80
BlueBERT	PubMed + MIMIC	4.5B words	30.08 ± 1.33	47.14 ± 0.81	46.12 ± 2.01	73.44 ± 2.78	54.99 ± 1.91	68.11 ± 1.41	48.55 ± 1.66	66.06 ± 1.40
CODER	UMLS	N/A†	40.50 ± 1.31	57.32 ± 1.74	47.37 ± 1.70	73.34 ± 1.29	53.74 ± 0.71	68.36 ± 0.36	49.86 ± 0.50	69.36 ± 1.00
LUKE	Wikipedia	3.5 billion words	27.44 ± 0.70	44.77 ± 0.40	42.35 ± 1.08	69.10 ± 0.62	50.92 ± 1.26	64.47 ± 1.75	46.16 ± 0.25	64.25 ± 1.28
RadBERT ¹	Radiology reports	2.6 GB	30.34 ± 1.50	48.00 ± 1.43	45.73 ± 0.68	73.00 ± 0.73	54.40 ± 2.84	67.34 ± 1.74	51.52 ± 0.87	68.80 ± 0.80
ClinicalBERT ²	MIMIC	0.5B words / 3.7GB	27.18 ± 1.89	44.69 ± 0.54	41.88 ± 2.86	68.90 ± 0.71	50.27 ± 1.63	63.40 ± 1.52	46.89 ± 0.13	64.41 ± 0.16
BioMed-RoBERTa ³	S2ORC	7.55B tokens / 47GB	27.44 ± 1.10	45.44 ± 0.64	42.35 ± 1.70	70.14 ± 0.99	52.82 ± 0.57	66.52 ± 0.32	48.62 ± 0.33	66.91 ± 0.82
Galactica	c.f. section 4	106B tokens	1.37	8.5	1.37	8.5	0.49	10.23	0.49	10.23
MedLLaMA	Medical Corpora	NA	0.3	10.63	0.3	10.63	0.16	12.14	0.16	12.14
MedAlpaca	Medical Meadow	NA	1.68	15.18	1.68	15.18	1.3	16.95	1.3	16.95

taining to COVID literature, which in turn aids in answering related questions.

47k corpus: With TOP-Training, BERT achieves a 4.01% increase in EM and a 3.5% increase in F1, while RoBERTa shows a 6.7% increase in EM and a 2.5% increase in F1, **setting a new SOTA on COVID-QA**. RoBERTa even outperforms the previous SOTA model (LUKE) by 1.2% in EM and 1.3% in F1, despite using a training corpus significantly smaller (67.4 MB/0.032B words) than LUKE’s 3.5B-word corpus (0.9% of the size). Moreover, any variation of our approach, ablation or otherwise, improves performance for both models over the Wikipedia baseline. These results indicate that our models benefit from additional training on corpora aligned with entity information from the downstream dataset.

470k [10x] corpus: Training with a 10x corpus (10 contexts per entity) led to the most improvements for BERT with EM increasing by 8.2% and F1 by 6.4%. This is consistent with Liu et al. (2019) who argue that BERT was *significantly un-*

dertrained. Though this improvement does not achieve RoBERTa performance, it demonstrates the scalability of our approach. RoBERTa improves too, but, not as much as when using the base 47k corpus. Undertrained BERT is more *malleable* to learning new concepts whereas RoBERTa seems to have hit its *ceiling for learning* in this domain.

Filtration - 50k corpus: Surprisingly, on removing ill-formed entities, Table 3 (Row 5) the performance declined with respect to the best BERT (10x corpus - row 4) and best RoBERTa (base 47k corpus - row 3) model. Our regex-based filtering rules mistakenly (as they cannot distinguish between true/false patterns) removed entities relevant to research articles such as author names or URLs, leading to the decline in performance.

Reduced context length - 47k (at most 1k context tokens) corpus: due to limitations in Galactica’s token generation i.e. at most 2048 tokens (last row of Table 3). Both models perform worse on both metrics as Galactica is unable to generate content matching the style of COVID-QA’s con-

Table 3: TOP-Training(COVID-QA). Time#: to generate corpus; ♣: entity filter; Gal = Galactica; max_length = Context Max Length. **Blue = best/red = worst** scores overall; **bold = best BERT/RoBERTa setup**.

Train Dataset	Time#	Corpus Size	Model	EM	F1
NA [Vanilla Fine-Tuning]	NA	NA	BERT	33.62 ± 0.59	60.01 ± 0.36
			RoBERTa	38.89 ± 0.52	67.44 ± 0.47
Wikipedia	≈ 2.5 hrs	139.6 MB	BERT	33.95 ± 0.13	60.76 ± 0.78
			RoBERTa	40.33 ± 0.60	68.30 ± 0.54
Gal(47k)	≈ 6.5 hrs	67.4 MB	BERT	34.97 ± 0.18	62.11 ± 0.32
			RoBERTa	41.51 ± 0.48	69.10 ± 0.27
Gal(470k) [10x]	≈ 2.5 days	558.2 MB	BERT	36.39 ± 0.27	63.84 ± 1.16
			RoBERTa	41.31 ± 0.22	68.84 ± 0.28
Gal(25k*2 = 50k)♣	≈ 6.5 hrs	64.0 MB	BERT	35.03 ± 0.38	62.14 ± 0.48
			RoBERTa	41.36 ± 0.35	69.00 ± 0.52
Gal(47k) [max_length = 1k] ≈ 2.5 hrs		44.8 MB	BERT	34.90 ± 0.14	62.02 ± 0.95
			RoBERTa	41.57 ± 0.33	68.98 ± 0.31

texts (research papers), underscoring the importance of domain-aware writing styles for adaptation pipelines.

6.2 RadQA

Results on RadQA are presented for both its validation and test splits. We consider various combinations of contexts (prompts) and entity filtration. Higher test scores on average are observed as compared to validation scores, which we attribute to fewer unanswerable questions in the test set (154 vs. 231) and slightly shorter contexts (73.82 vs. 78.1 tokens). We found no information leakage. We report scores for both, but mainly focus our analysis on the validation set.

6.2.1 Baselines

On the RadQA benchmark dev set, CODER, a PubMedBERT checkpoint, has the best EM & F1 but suffers slightly v/s PubMedBERT on only answerable questions presumably because CODER learned clinical embeddings from the UMLS knowledge graph with radiology terms.

Surprisingly, PubMed/Blue-BERT perform similarly on both the dev and test sets. Theoretically, BlueBERT should have performed better being pre-trained on MIMIC clinical notes. RadBERT, which is a superior RoBERTa architecture, and specifically trained on radiology reports did not perform well overall. Although it marginal improved over PubMed/Blue-BERT, it comes at the cost of a fraction of the training data. This again indicates the importance of proper domain alignment i.e. *what* data the models are trained on.

Unfortunately, LUKE performed poorly as compared to Bio/Sci-BERT, showing little, v/s both on dev, to no gain, v/s SciBERT on test, during evaluation. The impact of writing styles in the training corpora is clearly evident in the performance gap between Clinical/Rad-BERT. While the former was

trained on more clinical data, it was not the *right* type of data i.e. radiology reports, leading RadBERT to outperform it on both splits.

6.2.2 TOP-Training

Wiki Baseline: Training on the Wikipedia corpus, BERT shows an increase in EM only (2.8%) on the dev set, but an overall improvement in all measures on the test set compared to vanilla fine-tuning. RoBERTa, on the other hand, showed overall improvement on the dev set while test scores suffered compared to the the vanilla baseline.

Normal Prompting: Training on the unfiltered corpus with normal prompts (row 3) improved BERT and RoBERTa 3.7% EM, and 1.4% F1 for RoBERTa, over the vanilla baseline, which is greater than that of our Wiki baseline. However, **when the filter is applied (row 4)** BERT shows ~7% EM & 0.5% F1 increase while RoBERTa shows 6% EM & 3% F1 increase over regular fine-tuning. These scores indicate the necessity of using even a simple filter for selecting a subset of training corpora.

Best BERT setting: When both filtered entities and the corpus from the fancy prompt are used (row 6), improvements over basic fine-tuning (8.1% EM, 0.4% F1 on dev) and the Wikipedia baseline (5.2% EM, 1.1%F1 on dev) occur. Note: a) the benefit of studying RadQA and designing targeted prompts, and b) that BERT reaches these scores with a modest 34.3MB corpus, which is much smaller than the benchmarked models.

Best RoBERTa setting: RoBERTa demonstrates improvements across different combinations of filtration methods, prompt styles and using the Wikipedia corpus. However, the improvements are inconsistent with respect to a specific approach. Excluding the combined corpora settings, we see it achieve the best performance, on validation, using the corpus obtained from the filtered entities

Table 4: TOP-Training Results (RadQA). H(F1): HasAns_F1, H(EM): HasAns_EM; *: [Vanilla Fine-Tuning]. [†: normal, ‡: fancy] prompt, ♣: entity filter. **Blue** = **best**/red = **worst** overall; **bold** = **best BERT/RoBERTa setup**.

Train dataset / Corpus Size	Time#	Model	Dev				Test			
			EM	F1	H(EM)	H(F1)	EM	F1	H(EM)	H(F1)
NA*	NA	BERT	23.83 ± 0.49	42.91 ± 0.46	36.79 ± 0.76	66.23 ± 0.70	46.20 ± 1.96	59.42 ± 1.10	40.65 ± 1.15	58.30 ± 0.74
		RoBERTa	26.12 ± 0.69	43.83 ± 0.44	40.31 ± 1.06	67.65 ± 0.68	51.68 ± 0.73	64.94 ± 0.62	46.45 ± 0.66	64.14 ± 0.42
Wikipedia/18.4 MB	≈30 mins	BERT	24.49 ± 0.38	42.62 ± 0.08	37.80 ± 0.59	65.79 ± 0.13	47.40 ± 1.85	60.11 ± 1.59	41.95 ± 1.73	58.92 ± 1.56
		RoBERTa	27.19 ± 0.32	44.49 ± 0.31	41.96 ± 0.49	68.68 ± 0.47	50.54 ± 0.62	63.43 ± 0.75	45.72 ± 0.50	62.92 ± 0.67
Galactica(≈55k) †/ 81.6 MB	≈11 hrs	BERT	24.70 ± 0.46	42.88 ± 0.30	38.12 ± 0.71	66.19 ± 0.46	46.74 ± 1.84	59.69 ± 0.62	41.60 ± 2.31	58.88 ± 1.64
		RoBERTa	27.09 ± 0.70	44.42 ± 0.55	41.81 ± 1.09	68.56 ± 0.86	51.25 ± 0.41	64.41 ± 0.95	46.45 ± 0.66	64.01 ± 0.71
Galactica(≈55k) †♣/ 80.3 MB	≈11 hrs	BERT	25.51 ± 0.38	43.11 ± 0.23	40.38 ± 2.31	64.76 ± 2.75	46.36 ± 1.79	59.41 ± 1.67	46.59 ± 8.06	57.83 ± 1.81
		RoBERTa	27.69 ± 0.23	45.16 ± 0.35	42.74 ± 0.36	69.70 ± 0.54	51.14 ± 0.49	64.29 ± 0.38	46.31 ± 1.36	63.85 ± 0.76
Galactica(≈55k) ‡/ 120.8 MB	≈11 hrs	BERT	25.10 ± 0.32	42.78 ± 0.55	38.74 ± 0.49	66.03 ± 0.84	46.85 ± 1.74	59.54 ± 1.19	41.66 ± 0.45	58.60 ± 0.37
		RoBERTa	27.64 ± 0.49	44.99 ± 0.11	42.67 ± 0.76	69.45 ± 0.16	52.39 ± 0.80	65.57 ± 0.93	47.76 ± 1.85	65.34 ± 1.97
Galactica(≈55k) ‡♣/ 34.3 MB	≈11 hrs	BERT	25.76 ± 0.66	43.10 ± 0.27	39.68 ± 1.11	66.44 ± 0.55	46.52 ± 1.20	58.98 ± 1.09	40.44 ± 0.95	57.06 ± 0.94
		RoBERTa	27.08 ± 0.46	44.67 ± 0.23	41.81 ± 0.72	68.95 ± 0.35	51.30 ± 1.17	64.14 ± 0.41	47.39 ± 1.64	64.53 ± 0.85
Galactica(≈100k) †‡/ 115.6 MB	≈22 hrs	BERT	25.10 ± 0.54	42.74 ± 0.06	38.74 ± 0.83	65.97 ± 0.09	47.07 ± 1.23	60.04 ± 1.19	42.46 ± 0.13	59.78 ± 0.41
		RoBERTa	27.49 ± 0.49	44.82 ± 0.38	42.43 ± 0.75	69.17 ± 0.58	51.79 ± 1.98	64.99 ± 1.80	47.03 ± 1.45	64.45 ± 1.21
Galactica(≈100k) †‡♣/ 115.6 MB	≈22 hrs	BERT	24.75 ± 0.23	42.96 ± 0.10	38.20 ± 0.36	66.31 ± 0.16	47.50 ± 1.06	60.38 ± 0.48	41.96 ± 1.31	59.14 ± 1.97
		RoBERTa	27.85 ± 0.09	45.16 ± 0.11	42.98 ± 0.14	69.71 ± 0.18	51.79 ± 0.59	64.77 ± 0.40	46.88 ± 1.28	64.21 ± 0.75

and normal prompting (row 4) and on test, using filtered entities and fancy prompting (row 6).

RoBERTa v/s Benchmarks: In row 4, RoBERTa, outperforms Bio/Sci-BERT and LUKE on all metrics along with long-context models Big-Bird & Longformer. Interestingly, this RoBERTa even beats out BioClinicalBERT which was trained using much higher quality clinical notes scoring 1.9% EM & 1.1% F1 more than it.

Combined Prompting Styles: Merge the contexts from both prompt styles (rows 7 and 8) for the filtered and the unfiltered entities separately. BERT shows better performance in row 7 i.e. using the unfiltered corpus (1.7% EM increase over the filtered variant and roughly same F1) and RoBERTa in row 8 i.e. using the filtered corpus (1.3% EM and 0.8%F1 over the unfiltered variant). Overall, this version of RoBERTa resulted in our best overall model suggesting that incorporating a mixture of prompt styles will create more diverse corpora, enhancing domain alignment.

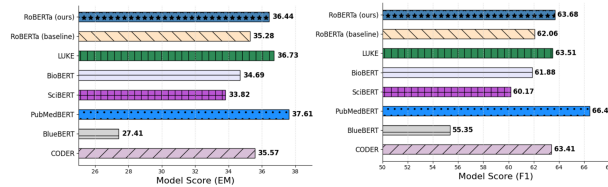


Figure 2: Information Leakage Validation Trials (Left - EM | Right - F1): RoBERTa (ours) was trained on a subset of the 47k corpus with entities only from the 80% train set. All of the models were fine-tuned in the usual manner i.e. SQuAD→COVID-QA (80% train set) and evaluated on the 20% test set.

6.3 Investigating Information Leakage

Given that the synthetic corpus generated for COVID-QA in §5 contains entities identified in the *entire* COVID-QA dataset - not from the *train* split within each *fold* - we explore if the performance gains from TOP-Training are a result of information leak. To this end, we construct a roughly 80%/20% train/test split (1,676/343 records), ensuring no context overlap, and apply a suite of models to this new split. When applying our TOP-Training, a synthetic corpus is generated *only* from entities identified in the train split.

RoBERTa subjected to TOP-Training still yields strong performance in this restricted scenario, only surpassed by PubMedBERT (and marginally by LUKE in EM) (Fig. 2) demonstrating that the improved performance on COVID-QA cannot be attributed to information leak from the test set. Although the scores are lower than those in Table 3, the relative scores produced by each model leads to a similar conclusion that TOP-Training yields optimal results.

7 Conclusion

In this study, we introduced TOP-Training, an innovative pretraining technique designed to enhance the alignment of a LLM with rare-domain target problems. The distinctive feature of TOP-Training lies in its automated synthesis of target-oriented data with the assistance of a LLM. Our experiments on medical EQA demonstrate the effectiveness of TOP-Training, shedding light on the limitations of widely used autoregressive LLMs.

Limitations

We identify two limitations of our work. First, we use a number of GPUs to generate our corpus. While we were fortunate to have access to powerful computing clusters, this could form a bottleneck when being deployed on low-end hardware. However, with cloud services being made more and more affordable, we feel that this point can only be a deal-breaker in severely budget-constrained settings. And second, in this study, we have only shown how to generate corpora for the biomedical domain. For an even wider applicability, we need to study generation techniques for other closed domains such as Finance, Law, Aviation, etc.

Ethics Statement

Our work relied on publicly available datasets. Ethical ramifications here are limited. To use RadQA, we had to acquire certifications to access it. Even though it is deidentified, we store the data in a protected fashion and have attempted not to quote even proxy identifiers that can be used to decode a person's identity.

Indirect hazards may be that our improved Q&A methodology can be used to ask questions that can be used to design bio-attacks on humans but such abusive uses are there with all technology and we strongly believe that the benefits world-wide of an improved closed-domain Q&A system is much bigger than the risks involved due to this technology.

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	A Token Filtering	916
	We performed entity filtering as a common abla-	917
	tion technique for both datasets. We used regular	918
	expressions to remove entities with special charac-	919
	ters such as *, !, etc., as well as specific text pat-	920
	terns like <code>https*</code> and <code>baby</code> . We implemented	921
	a length-based filter, retaining only entities longer	922
	than a certain number of characters. Additionally,	923
	for COVID-QA, we applied a second round of fil-	924
	tration using TF-IDF, considering the questions +	925
	context as the corpus and retaining the top 25k en-	926
	tities with the highest IDF scores. However, as	927
	this approach did not lead to substantial gains, we	928
	decided not to use it for RadQA. Due to the large	929
	number of possible combinations, we did not ex-	930
	tensively explore these settings in our experiments.	931
	B Synthetic Corpora Samples	932
	We provide samples from our generated corpus.	933
	In Figure 3 we show two positive results for our	934
	COVID-QA directed corpus. The top one is in	935
	the style of a research paper while the bottom one,	936
	though shorter in length, details useful information	937
	on the required entity. We show negative examples	938
	of the same in Figure 4. The top one completely	939

degrades into noise while the bottom one although coherent is talking about an unrelated topic.

For RadQA, we show similar examples. In Figure 5 we show positive (top) and negative (bottom) samples from normal prompting. Note that since programming languages were a part of Galactica’s training corpora, it sometimes generates the same during prompting. Finally, Figure 6 shows positive (top) and negative (bottom) examples of fancy prompting. While the positive sample shows a strong correlation with an actual radiology report, the bottom one completely breaks down.

C ChatGPT trials

We were curious to see if ChatGPT (with GPT3.5) would be able to answer a few questions from COVID-QA. Only the questions were provided to it without the context since they were much longer than its input capacity. We did not test RadQA since its questions were heavily linked to the patient report on hand and were not as *general* as the former’s. Figure 7 shows a few examples from our trials. As we can see, it seems to answer the questions correctly. However, as mentioned before, these answers need validation from a professional in the field. Also, we see that sometimes it **cannot answer** questions on topics not found in its training data, a problem not faced by our encoder models which rely only on the provided context.

D Hardware Details

To run our corpus generation code, we used a total of 8 40GB NVIDIA A100 or 8 32GB NVIDIA Tesla V100 cards, depending on our institutes job scheduler and availability. Although the former is a superior card, we preferred the latter since it gives relatively quick turnaround and is a general purpose GPU unlike the A100 which is required for even heavier compute. In fact we were requested by our system administrator to use the V100 over the A100, if it was possible, to avoid blocking off even more compute-heavy experiments, which we did not have an issue obeying. That being said, if the A100 cluster was free, we did take advantage of it to run our code. For pre-training and fine-tuning BERT/RobERTa, we used only 1 A100 80GB GPU as RobERTa was unable to fit on the 40/32 GB cards owing to the large batch size.

E Prompts Tried

As mentioned before, we knew the prompt to be used for COVID-QA since it was a feature of Galactica. However, we had to experiment with a few prompt styles for RadQA before landing on our final one. We describe the prompts we tried below for the entity small-bowel injury and the reasons for not using them. The text in *courier* is the prompt and in *italics* is the generated text.

- IMPRESSION: small-bowel injury (SBI) was defined as injury to the small bowel wall, or transmural disruption of the bowel wall, in the absence of penetrating abdominal trauma (The management of small-bowel injuries in the emergency room., Brown). Patients were excluded if the SBIs occurred after admission to the emergency department or were complicated by perforation. ... (truncated)

– Issue - Did not include the FINDINGS section.

- FINDINGS: small-bowel injury was observed in 11.1% and 21.9% of patients, respectively. Small-bowel injury associated with liver injury occurred in 10.1%. The mortality rate was significantly higher in the small-bowel injury group than in the non-small-bowel injury group (11.6% vs. 2.1%, $P = 0.005$). This study shows that small-bowel injury is associated with a high mortality rate, and that this complication is an important cause of morbidity and mortality in patients with severe acute pancreatitis.

– Issue - Did not include the IMPRESSIONS section.

- Patient has small-bowel injury. FINAL REPORT This patient was treated with conservative methods. The small-bowel injury was treated surgically with resection of the bowel, primary anastomosis, and end-to-end anastomosis between the ends of the resected bowel. She did not have recurrence of intestinal obstruction after 3 years.

– Issue - Did not include either IMPRESSIONS or FINDINGS section.

- John Doe has small-bowel injury. IMPRESSIONS : This

<p>Title: chest pain Title: an unusual cause of chest pain in an elderly patient Authors: Sushila Shrivastava, Vinay Kumar, Pramod Kumar Keywords: Chest pain, Eosinophilic pneumonia, Chest pain</p> <p># Abstract</p> <p># Background</p> <p>Pulmonary eosinophilic infiltration is one of the rare causes of non-specific chest pain in the elderly patients. We present a case of elderly patient who presented with a non-specific chest pain of 1 day duration with eosinophilia and chest CT scan showing ground glass opacities.</p> <p># Case presentation</p> <p>A 74-year-old lady presented with a 1-day history of chest pain, nausea and vomiting. The patient had a history of cough for 1 year, and she was on regular medication for cough. The patient had a significant history of hypertension, diabetes mellitus and COPD. On examination, the patient was afebrile with an oxygen saturation of 98% on room air. She had a heart rate of 90 beats per minute and blood pressure of 115/70 mmHg. Chest auscultation revealed a crackles in the lower lung fields. Her chest X-ray showed bilateral pulmonary infiltrates. A diagnosis of eosinophilic pneumonia was made and she was started on oral corticosteroids. She recovered without any further complications after two weeks of treatment.</p> <p># Conclusion</p> <p>Chest pain in the elderly patient is not always indicative of a life-threatening condition. A high index of suspicion for this rare entity should be kept in mind when a non-specific chest pain in elderly patients is seen.</p> <p># Background</p> <p>Pulmonary eosinophilic infiltration is a rare cause of non-specific chest pain in elderly patients. The clinical presentation of the disease is similar to other lung diseases with non-specific symptoms, and a high index of suspicion for this rare entity is required in elderly patients presenting with non-specific chest pain. We present a case of a 74-year-old lady with a history of cough who presented with a non-specific chest pain of 1 day duration with eosinophilia.</p> <p># Case presentation</p> <p>A 74-year-old lady was admitted in a state of mild depression. The patient had a significant history of hypertension, diabetes mellitus and COPD. She had a history of chronic cough for 1 year and was on regular medication for the cough. On examination, the patient was afebrile with an oxygen saturation of 98% on room air. She had a heart rate of 90 beats per minute and blood pressure of 115/70 mmHg. Chest auscultation revealed a crackles in the lower lung fields. She was investigated with a chest X-ray which showed bilateral pulmonary infiltrates. An ECG showed no sign of ST-T segment elevation. Her complete blood count (CBC) was as follows: haemoglobin 12.4 gm%, leucocytes 10,300/cu mm, platelets 253,000/cu mm, red blood cells (RBCs) 2.28 million/cu mm, and haemoglobin percentage (Hb%) 24.9%. Her electrolytes were normal. Her C-reactive protein (CRP) was 37.1 mg/L. D-dimers 0.24 µg/ml, and erythrocyte sedimentation rate (ESR) was 66 mm/1st hour. Her sputum was clear. The chest X-ray showed bilateral pulmonary infiltrates (Fig. 1). A diagnosis of eosinophilic pneumonia was made, and she was started on oral corticosteroids. She recovered without any further complications after two weeks of treatment. Fig. 1: Chest X-ray shows bilateral pulmonary infiltrates</p> <p># Discussion</p> <p>Eosinophilic pneumonia is an uncommon cause of non-specific chest pain. The incidence of this condition is approximately 0.1% of all pulmonary infiltrations [Idiopathic eosinophilic pneumonia: clinical features, radiological appearances and long-term outcome., Brown, Eosinophilic pneumonia: imaging findings., Bongiovanni]. The most common age group affected is between 40 and 60 years [Idiopathic eosinophilic pneumonia: clinical features, radiological appearances and long-term outcome., Brown]. However, there have been several case reports of eosinophilic pneumonia in the elderly, with the peak age between 60 and 70 years [Eosinophilic pneumonia in elderly patients: a case series and review of the literature, Kara, Eosinophilic pneumonia in the elderly, De Luca]. Most of the patients with eosinophilic pneumonia are asymptomatic at presentation, but it can also present with fever, dyspnea, cough, and chest pain [Idiopathic eosinophilic pneumonia: clinical features, radiological appearances and long-term outcome., Brown, Eosinophilic pneumonia in elderly patients: a case series and review of the literature, Kara, Eosinophilic pneumonia in the elderly, De Luca]. The chest pain in patients with eosinophilic pneumonia can be localized or generalized and can be of mild, moderate or severe intensity. It can be persistent or acute in nature [Idiopathic eosinophilic pneumonia: clinical features, radiological appearances and long-term outcome., Brown, Eosinophilic pneumonia in the elderly, De Luca]. The patient can be admitted to hospital with a cough, dyspnea, chest pain and fever. However, the patient may not show any of these symptoms at presentation [Idiopathic eosinophilic pneumonia: clinical features, radiological appearances and long-term outcome., Brown, Eosinophilic pneumonia in the elderly, De Luca].</p> <p>The most common clinical feature of eosinophilic pneumonia is cough, which is present in 85% of patients [Idiopathic eosinophilic pneumonia: clinical features, radiological appearances and long-term outcome., Brown]. Fever is present in 50% to 75% of patients [Idiopathic eosinophilic pneumonia: clinical features, radiological appearances and long-term outcome., Brown, Eosinophilic pneumonia in the elderly, De Luca]. Dyspnea and chest pain are seen in less than 10% of patients [Idiopathic eosinophilic pneumonia: clinical features, radiological appearances and long-term outcome., Brown, Eosinophilic pneumonia in the elderly, De Luca]. The most common respiratory symptoms are cough and dyspnea [Eosinophilic pneumonia in the elderly, De Luca]. The majority of the patients with eosinophilic pneumonia are asymptomatic, while a minority of patients present with non-specific symptoms of cough, fever, dyspnea and chest pain [Idiopathic eosinophilic pneumonia: clinical features, radiological appearances and long-term outcome., Brown, Eosinophilic pneumonia in the elderly, De Luca]. Patients with pulmonary eosinophilia and elevated ESR or CRP have been reported to have a poorer prognosis [Idiopathic eosinophilic pneumonia: clinical features, radiological appearances and long-term outcome., Brown]. Chest radiography can be nonspecific and shows bilateral or unilateral infiltrates [Idiopathic eosinophilic pneumonia: clinical features, radiological appearances and long-term outcome., Brown, Eosinophilic pneumonia in the elderly, De Luca]. A chest CT scan is often required to diagnose eosinophilic pneumonia and show a variety of appearances. These include ground glass opacities (GGO) and reticular opacities [Idiopathic eosinophilic pneumonia: clinical features, radiological appearances and long-term outcome., Brown, Eosinophilic pneumonia in the elderly, De Luca]. In our case, the patient had a history of chronic cough and had no fever on presentation. A chest X-ray showed bilateral infiltrates. A chest CT scan showed GGO and reticular opacities.</p> <p>The etiology of eosinophilic pneumonia is unknown, but it has been proposed that it is a T-helper 2-mediated inflammatory disorder [Idiopathic eosinophilic pneumonia: clinical features, radiological appearances and long-term outcome., Brown]. The diagnosis of eosinophilic pneumonia can be difficult because of the absence of typical radiological findings. It can be difficult to distinguish eosinophilic pneumonia from other inflammatory lung diseases. Treatment consists of corticosteroids, and the use of immunosuppressive drugs, such as azathioprine, hydroxychloroquine, and methotrexate, has been reported to improve the symptoms of eosinophilic pneumonia [Idiopathic eosinophilic pneumonia: clinical features, radiological appearances and long-term outcome., Brown, Eosinophilic pneumonia in the elderly, De Luca].</p> <p># Conclusion</p> <p>Eosinophilic pneumonia is a rare cause of non-specific chest pain in elderly patients. In these patients, the diagnosis can be difficult and a high index of suspicion should be kept in mind in such patients. Early diagnosis and prompt treatment can improve the prognosis.</p> <p># Consent</p> <p>Written informed consent was obtained from the patient for publication of this case report and any accompanying images. A copy of the written consent is available for review by the Editor-in-Chief of this journal.</p> <p>Footnote: Competing interests</p> <p>The authors declare that they have no competing interests.</p> <p>Footnote: Authors' contributions</p> <p>SS, KK and PK were involved in the management of the patient. SS drafted the manuscript. All authors read and approved the final manuscript.</p>	
<p>Title: Polymerase chain reaction #1: a new diagnostic tool in the management of tuberculosis.</p> <p>Abstract: Polymerase chain reaction (PCR) is an advanced molecular diagnostic technique. The potential of PCR in the diagnosis of tuberculosis (TB) has been evaluated by several investigators. PCR techniques have been used in the diagnosis of extra-pulmonary TB as well as the diagnosis of drug resistance. This article reviews the current status of PCR in the diagnosis of TB.</p>	

Figure 3: COVID-QA positive examples (underline = entity | **red** = prompt) | **olive** = generation



Figure 4: COVID-QA negative examples (underline = entity | **red** = prompt) | **olive** = generation

The role of FNA in the diagnosis of metastases to non-breast related lymph nodes has not been extensively studied. It has been reported that FNA was helpful in the diagnosis of metastases in patients with primary cancers other than breast cancer, such as non-small cell lung cancer [1]. The diagnostic value of fine-needle aspiration cytology for the diagnosis of nodal metastases in patients with non-small cell lung cancer, KIMJ, melanoma [1]. Cytologic diagnosis of metastases to nonbreast-related lymph nodes in breast cancer patients, ParkJ, esophageal cancer [2]. Fine needle aspiration cytology of lymph node metastases in esophageal carcinoma, YamagishiJ, and non-small cell lung cancer [3]. Fine-needle aspiration cytology of metastatic lung cancer to non-breast-related lymph nodes, KangJ. In our study, 44 lymph nodes were diagnosed as metastatic by FNA (Table 4). Table 4. The accuracy of FNA in staging of metastatic breast cancer to non-breast-related lymph nodes

Conclusions

In conclusion, FNA of the cervical lymph nodes was accurate in diagnosing metastatic breast cancer. In cases of lymph node enlargement in patients with breast cancer, FNA should be performed. In patients with negative cytomorphology on FNA, clinical, imaging, and biopsy findings should be considered. In these cases, additional procedures should be considered.

```

displacement(whatIs()) {
  EXPECT_EQ(1, dynamic_cast<const Shape*>(>shape_>shape()).dimensions());
  EXPECT_EQ(1, shape_>rank());
  EXPECT_TRUE(!shapeRank(shape_>shape(), shape_));
  EXPECT_EQ(1, shape_>dimensions());
}
}

TEST_P(ShapeOpTest, ShapeNotSameRank_1D) {
  Shape shape_;
  shape_.set_rank(1);
  shape_.set_dims({});
  Shape shape_copy = *shape_;
  shape_.set_rank(0);
  shape_copy.set_rank(1);
  shape_expected(shape_copy);
  TF_EXPECT_OK(
    shape_.AddScalar(2.0, /*add_shape=*/nullptr, /*is_dynamic=*/false));
  TF_EXPECT_OK(shape_.AddScalar(3.0, /*add_shape=*/nullptr, /*is_dynamic=*/false));
  TF_EXPECT_OK(shape_.AddScalar(4.0, /*add_shape=*/nullptr, /*is_dynamic=*/false));
  TF_EXPECT_OK(
    shape_.AddScalar(2.0, /*add_shape=*/nullptr, /*is_dynamic=*/true));
  TF_EXPECT_OK(shape_.AddScalar(3.0, /*add_shape=*/nullptr, /*is_dynamic=*/true));
  TF_EXPECT_OK(shape_.AddScalar(4.0, /*add_shape=*/nullptr, /*is_dynamic=*/true));
  TF_EXPECT_OK(shape_.AddScalar(2.0, /*add_shape=*/nullptr, /*is_dynamic=*/true));
  TF_EXPECT_OK(shape_.AddScalar(3.0, /*add_shape=*/nullptr, /*is_dynamic=*/true));
  TF_EXPECT_OK(shape_.AddScalar(4.0, /*add_shape=*/nullptr, /*is_dynamic=*/true));
  TF_EXPECT_OK(shape_.AddScalar(2.0, /*add_shape=*/nullptr, /*is_dynamic=*/true));
  TF_EXPECT_OK(shape_.AddScalar(3.0, /*add_shape=*/nullptr, /*is_dynamic=*/true));
  TF_EXPECT_OK(shape_.AddScalar(4.0, /*add_shape=*/nullptr, /*is_dynamic=*/true));
  TF_EXPECT_OK(shape_.AddScalar(2.0, /*add_shape=*/nullptr, /*is_dynamic=*/true));
  TF_EXPECT_OK(shape_.AddScalar(3.0, /*add_shape=*/nullptr, /*is_dynamic=*/true));
  TF_EXPECT_OK(shape_.AddScalar(4.0, /*add_shape=*/nullptr, /*is_dynamic=*/true));
  EXPECT_TRUE(ShapeIsSame(shape_, expected));
}

TEST_P(ShapeOpTest, ShapeNotSameRank_2D) {
  Shape shape_;
  shape_.set_rank(1);
  shape_.set_dims({1, 2});
  shape_.set_strides({1, 2});
  shape_.set_sizes({1, 2});
  Shape shape_copy = *shape_;
  shape_.set_rank(0);
  shape_copy.set_rank(1);
  shape_copy.set_dims({1, 2});
  shape_copy.set_strides({1, 2});
  shape_copy.set_sizes({1, 2});
  shape_expected(shape_copy);
  TF_EXPECT_OK(
    shape_.AddScalar(2.0, /*add_shape=*/nullptr, /*is_dynamic=*/false));
  TF_EXPECT_OK(shape_.AddScalar(3.0, /*add_shape=*/nullptr, /*is_dynamic=*/false));
  TF_EXPECT_OK(shape_.AddScalar(4.0, /*add_shape=*/nullptr, /*is_dynamic=*/false));
  TF_EXPECT_OK(
    shape_.AddScalar(2.0, /*add_shape=*/nullptr, /*is_dynamic=*/true));
  TF_EXPECT_OK(shape_.AddScalar(3.0, /*add_shape=*/nullptr, /*is_dynamic=*/true));
  TF_EXPECT_OK(shape_.AddScalar(4.0, /*add_shape=*/nullptr, /*is_dynamic=*/true));
  TF_EXPECT_OK(shape_.AddScalar(2.0, /*add_shape=*/nullptr, /*is_dynamic=*/true));
  TF_EXPECT_OK(shape_.AddScalar(3.0, /*add_shape=*/nullptr, /*is_dynamic=*/true));
  TF_EXPECT_OK(shape_.AddScalar(4.0, /*add_shape=*/nullptr, /*is_dynamic=*/true));
  EXPECT_TRUE(ShapeIsSame(shape_, expected));
}

TEST_P(ShapeOpTest, ShapeNotSameRank_3D) {
  Shape shape_;
  shape_.set_rank(1);
  shape_.set_dims({1, 2, 3});
  shape_.set_strides({1, 2, 3});
  shape_.set_sizes({1, 2, 3});
  Shape shape_copy = *shape_;
  shape_.set_rank(0);
  shape_copy.set_rank(1);
  shape_copy.set_dims({1, 2, 3});
  shape_copy.set_strides({1, 2, 3});
  shape_copy.set_sizes({1, 2, 3});
  shape_expected(shape_copy);
  TF_EXPECT_OK(
    shape_.AddScalar(2.0, /*add_shape=*/nullptr, /*is_dynamic=*/false));
  TF_EXPECT_OK(shape_.AddScalar(3.0, /*add_shape=*/nullptr, /*is_dynamic=*/false));
  TF_EXPECT_OK(shape_.AddScalar(4.0, /*add_shape=*/nullptr, /*is_dynamic=*/false));
  TF_EXPECT_OK(
    shape_.AddScalar(2.0, /*add_shape=*/nullptr, /*is_dynamic=*/true));
  TF_EXPECT_OK(shape_.AddScalar(3.0, /*add_shape=*/nullptr, /*is_dynamic=*/true));
  TF_EXPECT_OK(shape_.AddScalar(4.0, /*add_shape=*/nullptr, /*is_dynamic=*/true));
  TF_EXPECT_OK(shape_.AddScalar(2.0, /*add_shape=*/nullptr, /*is_dynamic=*/true));
  TF_EXPECT_OK(shape_.AddScalar(3.0, /*add_shape=*/nullptr, /*is_dynamic=*/true));
  TF_EXPECT_OK(shape_.AddScalar(4.0, /*add_shape=*/nullptr, /*is_dynamic=*/true));
}

```

Figure 5: RadQA - normal prompt (underline = entity | red = prompt) | olive = generation

[illegible]

Figure 6: RadQA - fancy prompt (underline = entity | red = prompt) | olive = generation

Question: What is DC-GENR and where is it expressed?

ChatGPT: As an AI language model, I do not have access to up-to-date information or research beyond my training data. Therefore, I am not aware of any current information on "DC-GENR." It is possible that this refers to a specific term or concept within a particular field or context. If you could provide me with more context or information about what you are referring to, I may be able to assist you further.

Question: What is the main cause of HIV-1 infection in children?

ChatGPT: *The main cause of HIV-1 infection in children is mother-to-child transmission of the virus during pregnancy, childbirth, or breastfeeding. This is also known as vertical transmission. If the mother is HIV-positive and not receiving treatment, there is a high risk of transmitting the virus to her baby. However, with proper medical care and treatment, the risk of mother-to-child transmission can be significantly reduced.*

Question: What plays the crucial role in the Mother to Child Transmission of HIV-1 and what increases the risk?

ChatGPT: *The level of the mother's HIV viral load plays a crucial role in mother-to-child transmission of HIV-1. A higher viral load in the mother's blood and genital secretions increases the risk of transmission to the baby. Factors that can increase the risk of mother-to-child transmission include lack of antiretroviral therapy during pregnancy, vaginal delivery, and breastfeeding. However, with proper medical care and treatment, the risk of mother-to-child transmission can be greatly reduced. Pregnant women living with HIV should seek medical care and treatment to help protect their babies*

Figure 7: Testing ChatGPT

1034	<i>case study was made possible by his support</i>	• Patient alpha has small-bowel	1081
1035	<i>by a research grant from the National</i>	injury requiring an open operation and	1082
1036	<i>Institutes of Health (CA 10644).</i>	was diagnosed with <i>C. difficile colitis</i> . The	1083
1037	– Issue - Had to make sure that the pro-	patient had a history of severe anemia and	1084
1038	nouns used were same throughout the	leukopenia, which were also observed in	1085
1039	report otherwise it would not be consis-	patient beta. Patient alpha was treated with	1086
1040	tent.	a single course of intravenous moxifloxacin	1087
1041	• Jane Doe has small-bowel	400 mg/day, which resulted in the resolution	1088
1042	injury. IMPRESSIONS : This case	of the diarrhea. He was followed closely after	1089
1043	study was made possible by the support of	discharge. In patient beta, he had a similar	1090
1044	a research grant from the Department of	episode of diarrhea after hospital admission,	1091
1045	Veterans Affairs.	but the diarrhea did not resolve. He was	1092
1046	– Issue - Same as above.	prescribed with an oral vancomycin 1000 mg	1093
1047	• Patient I has small-bowel	twice a day, and the diarrhea was resolved. ...	1094
1048	injury <i>He has no history of abdomi-</i>	(truncated)	1095
1049	<i>nal trauma or surgery. At first, he has only</i>	– Issue - This would have been a good	1096
1050	<i>mild abdominal pain. But with the progress</i>	choice. However, as we see above, pa-	1097
1051	<i>of disease, he has become distressed. At</i>	tient qualifiers such as A, I and alpha,	1098
1052	<i>the initial examination, the abdominal pain</i>	<i>might</i> bias the model towards more <i>male</i>	1099
1053	<i>was alleviated, but the patient still has slight</i>	patients (He). As such, we decided to	1100
1054	<i>distention of abdomen. He was admitted to</i>	drop the qualifier altogether.	1101
1055	<i>our hospital on the third day of admission.</i>		
1056	<i>His vital signs were: body temperature 37.5</i>	F Hyperparameters Used	1102
1057	<i>°C, blood pressure 110/70 mmHg, pulse 105</i>	Hyperparameters for each experiment is detailed	1103
1058	<i>beats/min, respiratory rate 26 breaths/min</i>	in Table 5. These were selected mostly from preex-	1104
1059	<i>and oxygen saturation 97% on room air.</i>	isting implementations or through minimal explo-	1105
1060	<i>He had mild abdominal distention. The</i>	ration of known settings.	1106
1061	<i>results of routine blood tests, urinalysis</i>	G Model Cards	1107
1062	<i>and abdominal ultrasound examination</i>	All models used in this study were downloaded	1108
1063	<i>were normal. X-ray showed free air in the</i>	from the HuggingFace library (Wolf et al., 2020).	1109
1064	<i>abdomen. ... (truncated)</i>	Each model, along with its model card (name as it	1110
1065	– Issue - Galactica could get confused	appears in the HuggingFace model hub) and URL	1111
1066	between I (alphabet) and I (roman nu-	is listed in Table 6.	1112
1067	meral).	H Note on Stability	1113
1068	• Patient A has small-bowel	All of our experiments were run using PyTorch	1114
1069	injury (Fig. 1). <i>He has no history</i>	1.13.1 and Huggingface 4.26.1. How-	1115
1070	<i>of abdominal trauma or surgery. At first,</i>	ever, we have noticed fluctuations in results when	1116
1071	<i>an abdominal CT scan was performed to</i>	training with other versions of these libraries. Thus,	1117
1072	<i>evaluate abdominal pain. Abdominal CT</i>	in order to replicate our scores to the best extent, we	1118
1073	<i>showed a partial small-bowel obstruction at</i>	recommend installing the aforementioned versions	1119
1074	<i>the splenic flexure of the colon (Fig. 2). An</i>	of the packages.	1120
1075	<i>oral contrast medium was then administered</i>		
1076	<i>via a nasogastric tube and an abdominal CT</i>		
1077	<i>scan was performed. ... (truncated)</i>		
1078	– Issue - Galactica could get confused mis-		
1079	interpret “A” for the beginning of a sen-		
1080	tence (it was observed for a few cases)		

Experiment	Hyperparameters
Corpus Generation	random seed: 42 renormalize_logits: True do_sample: True max_length (prompt + generated tokens): 2,048 top_p: 0.9 temperature: 0.9
Pre-Training	batch_size: 40 learning_rate: 5e-5 epochs: 3
Fine-Tuning (SQuAD)	batch_size: 16 max_input_length (question + context): 384 stride: 128 learning_rate: 2e-5 epochs: 3 n_best (top n answer spans): 20 max_answer_length: 30 optimizer_type: AdamW
Fine-Tuning (COVID-QA)	batch_size: 40 max_input_length (question + context): 384 stride: 128 learning_rate: 2e-5 epochs: 1 n_best (top n answer spans): 20 max_answer_length: 1000 optimizer_type: AdamW
Fine-Tuning (RadQA)	batch_size: 16 max_length: 384 stride: 128 learning_rate: 3e-5 epochs: 1 n_best (top n answer spans): 20 max_answer_length: 1000 optimizer_type: AdamW

Table 5: Hyperparameters for each experiment. We use three random seeds during pre-training and fine-tuning, 41, 42, 43 but only 42 when generating the corpus. This is done since otherwise to run the entire pipeline from generation to training across all ablations would take an infeasible amount of time.

Model	Model Card (URL)
BERT-Base, Cased	bert-base-cased
BERT-Base, Cased, SQuAD v1	batterydata/bert-base-cased-squad-v1
BERT-Base, Cased, SQuAD v2	deepset/bert-base-cased-squad2
RoBERTa-Base	roberta-base
RoBERTa-Base, SQuAD v1	csarron/roberta-base-squad-v1
RoBERTa-Base, SQuAD v2	deepset/roberta-base-squad2
BioBERT	dmis-lab/biobert-base-cased-v1.2
SciBERT	allenai/scibert_scivocab_uncased
SciBERT (+CORD-19)	lordtt13/COVID-SciBERT
PubMedBERT	microsoft/BiomedNLP-PubMedBERT-base-uncased-abstract-fulltext
BlueBERT	bionlp/bluebert_pubmed_mimic_uncased_L-12_H-768_A-12
CODER	GanjinZero/UMLSBert_ENG
LUKE	studio-ousia/luke-base
XLNet, SQuAD v1	arrafmousa/xlnet-base-cased-finetuned-squad
STonKGs *	stonkgs/stonkgs-150k
RadBERT	zzxslp/RadBERT-RoBERTa-4m
Clinical BERT	emilyalsentzer/Bio_ClinicalBERT
BioMed-RoBERTa	allenai/biomed_roberta_base
MedLLaMA	chaoyi-wu/MedLLaMA_13B
MedAlpaca	medalpaca/medalpaca-13b
Galactica	facebook/galactica-1.3b
Longformer, SQuAD v1	valhalla/longformer-base-4096-finetuned-squadv1
Longformer, SQuAD v2	mrm8488/longformer-base-4096-finetuned-squadv2
BigBird, SQuAD v1	FredNajjar/NF-bigbird-squad
BigBird, SQuAD v2	FredNajjar/bigbird-QA-squad_v2

Table 6: Model cards and URLs for all models used in our paper. * We wanted to use STonKGs (Balabin et al., 2022). However, there was no vocabulary file for the model which resulted in errors.