Mitigating Hallucinations of Large Language Models in Medical Information Extraction via Contrastive Decoding

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Abstract

The impressive capabilities of large language models (LLMs) have attracted extensive interests of applying LLMs to medical field. However, the complex nature of clinical environments presents significant hallucination challenges for LLMs, hindering their widespread adoption. In this paper, we address these hallucination issues in the context of Medical Information Extraction (MIE) tasks by introducing ALternate Contrastive Decoding (ALCD). We begin by redefining MIE tasks as an *identify*and-classify process. We then separate the identification and classification functions of LLMs by selectively masking the optimization of tokens during fine-tuning. During the inference stage, we alternately contrast output distributions derived from sub-task models. This approach aims to selectively enhance the identification and classification capabilities while minimizing the influence of other inherent abilities in LLMs. Additionally, we propose an alternate adaptive constraint strategy to more effectively adjust the scale and scope of contrastive tokens. Through comprehensive experiments on two different backbones and six diverse medical information extraction tasks, ALCD demonstrates significant improvements in resolving hallucination issues compared to conventional decoding methods.

1 Introduction

Medical Information Extraction (MIE), including tasks such as medical entity recognition and relation extraction, is a fundamental component of medical NLP (Hahn and Oleynik, 2020; Xu et al., 2024b). It enables the derivation of structured knowledge from plain text, benefiting a wide array of applications (Wang et al., 2024; Liang et al., 2023; Qi et al., 2024), like medical knowledge graph construction (Wu et al., 2023; Xu et al., 2024a, 2023), medical dialogue (Gao et al., 2023;



Figure 1: An example demonstrating the hallucination generated by LLMs in MIE tasks. The green font in medical dialogue indicates a high correlation with ground truth. The blue font in the output represents correct token, while the red font represents tokens with hallucination problems. These problems mainly include the presence of nonexistent entities and reasoning errors.

Wu et al., 2024), and medical report generation (Liu et al., 2021). Previous MIE tasks (Yu et al., 2019; Guan et al., 2020) have been supervised, and their performance heavily depends on the quality and quantity of available training data. However, labeling medical documents requires specific knowledge which is both costly and time-consuming.

Recently, the remarkable zero-shot capabilities of large language models (LLMs) such as Chat-GPT and GPT-4 (OpenAI, 2023) have inspired researchers to transform MIE tasks into a generation paradigm (Zhu et al., 2023). However, the medical domain is less tolerant of errors compared to other domains. While there have been attempts to apply LLMs to the medical field (Singhal et al., 2022; Sharma et al., 2023; Liu et al., 2024a,b), there is a growing concern about the issue of hallucination (Huang et al., 2023). In the context of MIE, two types of hallucinations exist: (1) LLMs may

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identify medical entities that are not present in original texts, thereby fabricating facts and deviating from the original information. (2) LLMs may face reasoning errors when classifying medical entities, due to statistic biases in the pre-trained corpus. We show such a hallucination problem in Figure 1.

In this paper, we address the challenges of hallucination when applying LLMs to MIE tasks. We observe that LLMs for MIE can be conceptualized as an identify-and-classify process: initially identifying potential medical concept spans from the plain text, and then classifying these text spans into predefined categories (e.g., start token of a specific entity, subject of a specific relation), as shown in the 'Output' of Figure 1. The natural approach to applying LLMs is to prompt them to simultaneously complete both *identify* and *classify* steps in a unified decoding process (Lu et al., 2022; Wang et al., 2023b). We speculate that the hallucination problem may be linked to the joint next-word generation abilities of identification and classification, which could have inadvertently compromised each other's performance. Therefore, we believe that decoupling abilities of identification and classification, allowing LLMs to concentrate on specific sub-tasks, could simplify the complexity of the MIE task and potentially reduce hallucination issues (Khot et al., 2022; Bian et al., 2023).

Motivated by the aforementioned observation, we introduce ALternate Contrastive Decoding (ALCD), a straightforward decoding strategy designed to enhance the performance of LLMs on MIE tasks. In the training stage, we mask the optimization of tokens separately to decouple the identification and classification models. For instance, when fine-tuning the parameters of the identification model, classification tokens are masked to focus the model's attention solely on identification tokens, thereby ignoring its classification capability. During the inference stage, ALCD bolsters its classification/identification ability and contrasts logit predictions with another model. This contrastive decoding process alternates between classification and identification, depending on the type of the next token, which is determined by a simple rulebased judgment. Furthermore, we propose an adaptive constraint strategy to dynamically adjust the scale and scope of contrastive tokens. This allows individual samples to adapt to their unique characteristics by measuring the consistency among the three models and the level of confidence. Overall, this work makes three key contributions:

- To our knowledge, we are the first to employ contrastive decoding as a strategy to reduce hallucinations in LLMs for MIE tasks.
- We validate the broad applicability of our ALCD approach through experiments using two LLM backbones across six diverse medical tasks, such as determining causal relationships in medical concepts (Zhu et al., 2023).¹
- Our experimental results underscore the superiority of ALCD over eight established decoding methods.

2 Related Work

2.1 LLMs for Medical Domain

Rapid development has been seen in directly employing general LLMs (e.g., ChatGPT (OpenAI, 2023), ChatGLM (Du et al., 2022), and Qwen (Bai et al., 2023)) to the medical domain and training medical LLMs using medical data, such as Med-PaLM (Singhal et al., 2022), clinicalGPT (Wang et al., 2023a), and MedAlpaca (Han et al., 2023). Both general LLMs and medical LLMs may suffer from hallucinations, the undesired phenomenon of LLMs generating contents not based on training data or facts when applying them to complex medical tasks. Hallucinations could be caused by multiple factors, such as imperfect representation learning or erroneous decoding (Ji et al., 2023a). Due to the high demand for reliability in the medical domain, the hallucinations are thus less tolerated. Although previous works have explored the problem of hallucination in the medical domain (Umapathi et al., 2023; Ji et al., 2023b), there is a lack of exploration in MIE task, particularly regarding the efficiency of different decoding methods for mitigating hallucination.

2.2 Contrastive Decoding

The idea of contrastive decoding for LLM has been explored in various previous works, and different decoding strategies focus on different aspects of LLM improvements. Contrastive Decoding (CD) (Li et al., 2023) is proposed to contrast output probability of large-scale expert LLMs with small-scale amateur LLMs to diminish undesired amateur behavior and improve fluency and coherence in the generated contents. Context-aware Decoding (CAD) (Shi et al., 2023) focuses on the issue of LLMs' insufficient attention to context.

¹https://github.com/quqxui/quqxui-AlternateCD

CAD downweights output probability associated with LLMs' prior knowledge to promote LLMs' attention to context, thus improving the faithfulness of the generated contents. Chuang et al. (2024) introduced DoLa, where the output next-word probability is obtained from the difference in logits between a higher layer versus a lower layer, to reduce hallucinations and enhance truthfulness in the knowledge-based question-answering tasks. Visual Contrastive Decoding (VCD) is another decoding method to mitigate object hallucinations for large vision-language models by contrasting output distributions from original and distorted visual inputs (Leng et al., 2023). Sanchez et al. (2023) adapted Classifier-Free Guidance (CFG) (Ho and Salimans, 2022) from text-to-image generation to text-to-text generation and they showed CFG can increase the LLMs' performance and adherence to various prompts, including basic prompting, chainof-thought prompting, and chatbot prompting.

Although previous contrastive decoding strategies have been shown effective in addressing specific hallucinations in LLMs, their performance is inadequate for MIE tasks. In contrast, our ALCD effectively decouples the abilities to contrast and decode outputs, leading to notable enhancements.

3 Methodology

In this section, we introduce ALternate Contrastive Decoding (ALCD), a method specifically designed for medical information extraction tasks. Section 3.1 provides the foundational knowledge of Contrastive Decoding, while Section 3.2 delves into the details of our proposed ALCD method.

3.1 Preliminary

For generative LLMs, the common method for text generation is to predict next token in an autoregressive manner. Specifically, we denote the parameters of an LLM as θ . The model utilizes input text x and system instructions (prompts) i to generate a response y. For each time step t, we have:

$$y_t \sim \mathcal{P}_{\theta}(y_t | \boldsymbol{i}, \boldsymbol{x}, \boldsymbol{y}_{< t}),$$

$$\sim softmax(logit_{\theta}(y_t | \boldsymbol{i}, \boldsymbol{x}, \boldsymbol{y}_{< t})), \qquad (1)$$

where y_t represents the output token at a specific time step t, and $y_{<t}$ denotes the sequence of generated token sequence until the time step t - 1. The common ways of the next token selection include selecting the highest probability token (greedy search), exploring multiple high-probability paths simultaneously (beam search), or sampling according to the probability distribution (e.g., nucleus sampling (Holtzman et al., 2019)).

While, in contrastive decoding, there are typically two logits, which may be obtained from different LLMs using the same input source (Li et al., 2023) or the same LLM using different input sources (Shi et al., 2023). It should be noted that they need to share the same tokenizer to keep consistency between different logits. The probability for the next token is adjusted through subtraction:

$$logit_{\theta}(y_t|\boldsymbol{i}, \boldsymbol{x}, \boldsymbol{y}_{< t}) - logit_{\theta'}(y_t|\boldsymbol{i}, \boldsymbol{x}, \boldsymbol{y}_{< t}).$$
 (2)

The $logit_{\theta}$ and $logit_{\theta'}$ are usually generated from an LLM with high capabilities and low capabilities, respectively. For example, in CD (Li et al., 2023), $logit_{\theta}$ comes from a large expert LLM and $logit_{\theta'}$ comes from a small amateur LLM. Subtracting these two logits helps amplify the ground-truth tokens in $logit_{\theta}$ and downplay hallucinated tokens in $logit_{\theta'}$. Inspired by CD, we propose to alternately amplify or downplay the classification and identification capabilities of LLMs during the decoding process, to improve final generation results.

3.2 Alternate Contrastive Decoding

The process of our proposed ALCD is illustrated in Figure 2. We break down medical information extraction into two stages: identification and classification. In Section 3.2.1, we fine-tune LLMs separately for identification and classification. In Section 3.2.2, we utilize the decoders of three LLMs (identification, classification, and normal) together to perform MIE. As the two new LLMs are trained with Lora (Hu et al., 2021), they do not cause an excessive increase in training.

3.2.1 Decoupling with optimization masking

To effectively harness identification and classification capabilities of LLMs while minimizing interference from one another, we propose to decompose their respective abilities. Typically, it is natural to fine-tune two subtasks independently, resulting in a identification model \mathcal{M}_{id} and a classification model \mathcal{M}_{cl} . But this method has distinct instructions and input-output formats compared to normal model \mathcal{M}_{nl} . It poses an issue when these models are combined during the inference step, which can lead to inconsistent input with finetuning step, ultimately reducing the accuracy.

In this work, we propose to optimize two capabilities separately using optimization masking during the fine-tuning process, as shown in Figure



Figure 2: The overall pipeline of our proposed ALCD consists of two main steps. In Step #1, we aim to fine-tune submodels individually to decouple the abilities of identification and classification. In Step #2, we adaptively contrast the predictions at each time step by applying scale and scope constraints on tokens. The figure shows how LLMs generate token y_t at time step t based on previous tokens $y_{<t}$. The terms cls, ide, other represent classification, identification, and other tokens, respectively. The output logits of normal, classification and identification models are represented as l_{nl}^{θ} , l_{cl}^{θ} , and l_{id}^{θ} .

2(Step #1). We employ the same inputs as original task for fine-tuning both \mathcal{M}_{id} and \mathcal{M}_{cl} models. During fine-tuning, we selectively optimize tokens, and for instance, when optimizing parameter θ_{id} of identification model \mathcal{M}_{id} , we mask the tokens for classification task:

$$\max_{\theta_{id}} \sum_{(\boldsymbol{x}, \boldsymbol{y}) \in \mathcal{D}} \sum_{t=1, t \notin \mathcal{T}_{cl}}^{|\boldsymbol{y}|} log(\mathcal{P}_{\theta_{id}}(y_t | \boldsymbol{i}, \boldsymbol{x}, \boldsymbol{y}_{< t})), \quad (3)$$

where \mathcal{T}_{cl} represents the time step of classification tokens, which do not require optimization, and \mathcal{D} denotes training dataset. On the other hand, when optimizing parameter θ_{cl} of classification model \mathcal{M}_{cl} , we mask the tokens for identification task:

$$\max_{\theta_{cl}} \sum_{(\boldsymbol{x}, \boldsymbol{y}) \in \mathcal{D}} \sum_{t=1, t \notin \mathcal{T}_{id}}^{|\boldsymbol{y}|} log(\mathcal{P}_{\theta_{cl}}(y_t | \boldsymbol{i}, \boldsymbol{x}, \boldsymbol{y}_{< t})), \quad (4)$$

where \mathcal{T}_{id} represents time step of identification tokens. By employing masking optimization, we expect to develop LLMs that possess diverse capabilities. For fine-tuning normal model \mathcal{M}_{nl} , we also employ formulas similar to 3 and 4, but without any masking operations. Given the constraints of computational resources, we implemented parameterefficient fine-tuning techniques (e.g., LoRA (Hu et al., 2021)) to train these models.

3.2.2 Adaptively Contrasting the Predictions

After decoupling the capabilities, a significant challenge arises: how can we effectively harness the individual abilities of sub-models? To address this, ALCD is designed to alternate the enhancement of the classification ability of \mathcal{M}_{cl} and the identification ability of \mathcal{M}_{id} during LLM's inference stage, while excluding the influence of other capabilities originally present in normal model \mathcal{M}_{nl} . An illustration is shown in Figure 2(Step #2).

We denote $n_t \in \{cls, ide, other\}$ as the type of next token prediction, where cls, ide, other indicate classification, identification, and other tokens, respectively. Generally, in order to facilitate the evaluation of text generated from LLMs, it is typically to present the output of MIE in a structured format (Lu et al., 2022). Therefore, when LLMs generate token y_t at time t, we can determine the next token based on previous tokens $y_{< t}$ using a simple rule-based judgment: In our case, we require LLMs to utilize colon ':' to split ide and *cls* tokens, and each *ide-cls* pair is separated by a newline character '\n'. For instance, in this text: "Dizziness: positive $\ n$ fever: negative", the ide tokens (Dizziness or fever) are expected to be followed by a colon and then a *cls* token (*posi*tive or positive). We abbreviate the representation $logit_{\theta}(\cdot | \boldsymbol{i}, \boldsymbol{x}, \boldsymbol{y}_{< t})$ generated by $\mathcal{M}_{nl}, \mathcal{M}_{cl}$, and

 \mathcal{M}_{id} as l_{nl}^{θ} , l_{cl}^{θ} , and l_{id}^{θ} , respectively. The overall formula is as follows:

$$\begin{aligned} l_{nl}^{\theta} + \alpha (d_{cl} * l_{cl}^{\theta} - d_{id} * l_{id}^{\theta}), & \text{if } n_t = cls \\ l_{nl}^{\theta} + \alpha (d_{id} * l_{id}^{\theta} - d_{cl} * l_{cl}^{\theta}), & \text{if } n_t = ide \\ l_{nl}^{\theta}, & \text{if } n_t = other \end{aligned}$$
(5)

where α is a hyper-parameter and analyzed in Section 4.4. d_{id} and d_{cl} are adaptive scales proposed to measure the distance between two logit distributions: one between \mathcal{M}_{nl} and \mathcal{M}_{id} , and the other between \mathcal{M}_{nl} and \mathcal{M}_{cl} . We leverages Jensen-Shannon Divergence (JSD) to calculate them:

$$d_{id} = JSD(logit_{\theta_{nl}} || logit_{\theta_{id}}), d_{cl} = JSD(logit_{\theta_{nl}} || logit_{\theta_{cl}}).$$
(6)

Specifically, when predicting the next token in Formula (5), ALCD includes two extra components in addition to the logit l_{nl}^{θ} of the normal model. For example, if n_t is a cls token, The first component is enhancing l_{cl}^{θ} , with the motivation to utilize the classification ability of sub-model \mathcal{M}_{cl} . If the outputs of \mathcal{M}_{cl} is more different from \mathcal{M}_{nl} (e.g., larger d_{cl}), we will be more inclined towards the classification model. The second component involves contrasting the influence of sub-models \mathcal{M}_{id} , by decreasing logit values l_{id}^{θ} through adaptive scales (d_{id}) . The motivation behind this is that if the outputs of \mathcal{M}_{id} is more different from \mathcal{M}_{nl} (e.g., larger d_{id}), indicating a stronger contrast (denoted as $-d_{id} * l_{id}^{\theta}$), which makes sure that ALCD has the potential to mitigate the hallucinations arising from identification ability.

Conversely, when the next token is an *ide* token, the same rule is applicable. For the next token that do not belong to either *ide* or *cls*, we solely utilize logit output l_{nl}^{θ} of normal model. By employing this alternating contrast prediction, ALCD has the capability to modify the overall probability of tokens and then harness the abilities of sub-models.

3.2.3 Scope Constraints on Tokens

In addition, it is worth noting that certain tokens may exhibit a significant discrepancy when subjected to contrastive decoding, which makes the implausible tokens receive a high score after contrast, leading to what is referred to as the false positives (Li et al., 2023; Chuang et al., 2024). In light of this, we implement a constraint that is contingent upon the confidence level:

$$\mathcal{V}_{head}(\boldsymbol{y}_{< t}) = \{ v \in \mathcal{V} : \\ \mathcal{P}_{\theta}(v | \boldsymbol{i}, \boldsymbol{x}, \boldsymbol{y}_{< t}) \ge \beta \max_{v} \mathcal{P}_{\theta}(v | \boldsymbol{i}, \boldsymbol{x}, \boldsymbol{y}_{< t}) \},$$
(7)

Dataset	#Train	#Valid	#Test
CMeEE-V2	4,600	400	400
CMeIE-V2	4,600	400	400
IMCS-V2-NER	4,600	400	400
CMedCausal	2,600	400	400
IMCS-V2-SR	4,600	400	400
CHIP-MDCFNPC	4,600	400	400

Table 1: Dataset partitioning statistics.

where \mathcal{V} represents the output vocabulary of LLMs, v is the token of output vocabulary, and β is a hyper-parameter used to determine the max truncation rate of low-probability tokens. Instead of employing constraints with a single model in Li et al. (2023), our approach involves combining the intersection of confidence values $\mathcal{V}_{head}^{inter}$ obtained from three models (outputs of \mathcal{M}_{nl} , \mathcal{M}_{id} , and \mathcal{M}_{cl}). Tokens with confidence levels below a specific threshold are assigned a negative infinity value:

$$\begin{aligned}
\mathcal{V}_{head}^{inter} &= \mathcal{V}_{head}^{nl} \cap \mathcal{V}_{head}^{cl} \cap \mathcal{V}_{head}^{id}, \\
logit_{\theta}(v|\boldsymbol{i}, \boldsymbol{x}, \boldsymbol{y}_{< t}) &= -\infty, \text{ if } v \notin \mathcal{V}_{head}^{inter}(\boldsymbol{y}_{< t}).
\end{aligned} \tag{8}$$

By combining token constraints to enhance and contrast predictions, our proposed ALCD is able to effectively leverage capabilities of \mathcal{M}_{id} or \mathcal{M}_{cl} , while addressing the issue of hallucinations in \mathcal{M}_{nl} that arise from other capabilities in \mathcal{M}_{cl} or \mathcal{M}_{id} .

4 Experiments

4.1 Experimental Setup

Tasks and Datasets. We apply six MIE tasks from a Chinese medical dataset named PromptCBLUE (Zhu et al., 2023) for evaluation. CMeEE-V2 is a task of Chinese medical entity recognition. IMCS-V2-SR aims to normalize the patient-doctor dialogue by medical concepts. IMCS-V2-NER targets extracting medical concepts from dialogues. CMedCausal is a task of causal relation extraction for medical texts. CHIP-MDCFNPC refers to clinical concept finding and discrimination. CMeIE-V2 aims to recognize and categorize the entity relation contained in medical texts. The output forms of all tasks are built with the *identify-and-classify* pattern, as mentioned in Section 1. Due to space limitations, we leave more details about the tasks to Appendix A.1. Since the open-source test set was not available, we used the validation set as our test set. Subsequently, we partition the training set into a new training set and validation set and ensure the validation set contains the same number of samples as the test set. Table 1

Decoding Method	CMeEE-V2	CMeIE-V2	IMCS-V2-NER	CMedCausal	IMCS-V2-SR	CHIP-MDCFNPC
			ChatGLM-6B			
Greedy Search	66.48	45.60	88.37	41.01	71.55	42.58
Beam Search	66.77	45.80	88.60	41.41	71.84	42.77
Top K Sample	63.38	39.02	88.19	39.41	69.40	38.87
Nucleus Sample	64.93	41.13	88.26	40.58	69.88	41.92
CFG (Sanchez et al., 2023)	66.95	43.84	88.76	40.61	72.06	42.49
CAD (Shi et al., 2023)	66.88	44.04	88.77	40.57	72.06	42.49
CD (Li et al., 2023)	66.34	46.03	88.54	40.72	72.40	42.33
DoLa (Chuang et al., 2024)	66.46	43.78	88.96	40.47	38.68	42.92
ALCD (Ours)	67.45 [*]	46.83 [*]	89.49	42.28*	73.01 *	43.71 [*]
		ļ	Qwen-7B-Chat			
Greedy Search	65.49	42.87	88.65	30.10	71.28	40.61
Beam Search	66.61	43.40	89.46	30.21	71.35	40.94
Top K Sample	65.71	36.34	88.83	19.55	71.04	40.19
Nucleus Sample	66.04	33.87	89.08	25.81	70.09	39.40
CFG (Sanchez et al., 2023)	65.18	39.07	88.64	12.96	71.15	40.18
CAD (Shi et al., 2023)	66.09	36.67	88.00	14.40	71.72	39.49
CD (Li et al., 2023)	65.19	35.86	88.98	14.69	70.27	39.35
DoLa (Chuang et al., 2024)	65.16	35.51	88.49	16.52	71.29	39.37
ALCD (Ours)	67.91 [*]	44.19 [*]	90.88 [*]	31.57 [*]	72.14*	41.88

Table 2: Experiment results (micro F1 score \uparrow : higher is better) on six medical datasets with the best scores highlighted **in bold**. All baselines are based on the same fine-tuned normal model, and the model-agnostic parameters for fine-tuning and inference are kept consistent, with only the specific decoding method being changed. "*" indicates the statistically significant improvements (i.e., two-sided t-test with p < 0.05) over the best baseline.

presents the dataset partitioning statistics.

Models and Baselines. To improve the learning of data, we experimented with two widely-used multilingual LLMs, ChatGLM-6B v1 (Du et al., 2022) and Qwen-7B-Chat v1 (Bai et al., 2023). We compared our method for mitigating hallucinations with eight decoding baselines, which can be categorized as follows: Deterministic decoding: 1) greedy search decoding; 2) beam search decoding; Stochastic decoding: 3) Top K sample decoding; 4) nucleus sample decoding; Contrastive decoding: 5) CFG (Ho and Salimans, 2022); 6) CAD (Shi et al., 2023); 7) CD (Li et al., 2023); 8) DoLa (Chuang et al., 2024). For the validation of Deterministic and Stochastic methods, we utilized the implementation provided by the Huggingface toolkit (Wolf et al., 2020). However, for the contrastive decoding methods, adjustments were required when applying them to MIE tasks as they were not specifically designed to tackle the hallucination problem in MIE. For CFG, we simply use logits with normal input text and logits with the last token of input text as a comparison. For CAD, we employ both normal input text and input text

without classification labels to contrast the output in different contexts. For CD, we employ the normal model as the expert model and proceed with a model using only half the number of fine-tuning steps for the amateur model. DoLa is implemented following their published paper.

Implementation Details. We conducted all experiments using four NVIDIA V100 GPUs. As we finetuned LLMs using LoRA, the decoding process was performed using a single GPU. All experimental results were evaluated using the Micro-F1 score following Zhu et al. (2023). For ALCD, we conducted a search in the validation set to determine the appropriate values for the scale of contrasting prediction α , the maximum rate of constraint β , and the step of fine-tuning. For α , we limit the search scope to the values of [0.01, 0.1, 0.2, 0.3, 0.4, 0.5]. For β , we limit the search scope to the values of [0.4, 0.45, 0.5, 0.55, 0.6, 0.65]. The finetuning step of the normal model remains consistent across all baselines. We employ a batch size of 8 and perform 1,000 steps to fine-tune all datasets and LLMs, except for Qwen-7B-Chat where we use 3,000 steps in CMeIE-V2, CMedCausal, and



Figure 3: Ablation study on six medical datasets using ChatGLM-6B.

CHIP-MDCFNPC, due to that extra steps are required for convergence.

4.2 Main Results

In this section, we provide a comprehensive performance comparison of ALCD against other baselines on six medical datasets and two different backbone LLMs. As shown in Table 2, our proposed ALCD outperforms both contrastive decoding and non-contrastive decoding methods and the performance gap reaches the largest of 4.87% in Qwen-7B-chat on the CMedCausal dataset. Our proposed ALCD has been shown to improve performance on both ChatGLM-6B and Qwen-7B-Chat, which confirms its universality. Besides, ALCD particularly performs well on CMeEE-V2, IMCS-V2-NER, and CHIP-MDCFNPC datasets, and outperforms other baselines by a large margin. This finding aligns with our motivation as these datasets include more entity candidates, more classification labels, and thus higher difficulties for LLMs. Some contrastive decoding methods, such as DoLa, achieve much lower results on IMCS-V2-SR in the ChatGLM-6B, indicating the coupled difficulties for the medical identify-and-classify tasks. We find that the proposed adaptive method of DoLa predominantly selects the 2nd or 8th layer as the optimal premature layer, which suggests that DoLa's intended ability to amplify factual knowledge across different layers may not be fully aligned with the MIE tasks. We observed that the poor performance of



Figure 4: (a) Analysis of the scale of contrasting prediction α (in Formula 5); (b) Analysis of max rate of constraint β (in Formula 7).

sampling methods (Top K and Nucleus Sample) indicates that high diversity generation may not be essential for the MIE task.

4.3 Ablation Study

In this section, we analyze the effects of different components on ALCD. Specifically, we experiment with ALCD against three variants: 1) ALCD without Constraint: removing the dynamic constraints on tokens, 2) Alternate Sum: alternately summing the logits from three models instead of utilizing contrastive decoding, 3) Weighted Sum: directly summing the logits from three models with the same weight of ALCD. As depicted in Figure 3, the results confirm that incorporating token constraints enhances the performance of the normal model. Specifically, on the CMeIE-V2 dataset, the micro F1 score decreased from 47.02% to 46.19% when no constraints were utilized. Moreover, removing the alternate contrasting with either Alternate Sum or Weighted Sum resulted in performance declines, with Weighted Sum yielding the poorest overall performance. This finding highlights the effectiveness of applying alternate contrastive decoding and indicates that solely ensembling multiple LLMs for these tasks does not lead to performance improvement.

4.4 Scale of Contrasting Prediction

To investigate the effect of hyper-parameter α in Formula 5, we set different values from 0.01 to 0.5 and conduct experiments on CMeEE-V2 and IMCS-V2-SR datasets. A larger α means a larger

Dataset	Constraint in CD	Ours
CMeEE-V2	66.38	67.45
CMeIE-V2	46.11	46.83
IMCS-V2-NER	89.02	89.49
CMedCausal	41.73	42.28
IMCS-V2-SR	72.64	73.01
CHIP-MDCFNPC	42.88	43.71

Table 3: Comparison of token constraint method on all datasets using ChatGLM-6B.

scale of contrastive decoding. As shown in Figure 4(a), it can be observed that increasing the scale of contrastive decoding appropriately enhances the micro F1 score of both backbone LLMs, indicating the efficiency of our contrastive decoding method. While, excessively large values of α (e.g., exceeding 0.4), can lead to a decline in performance, which demonstrates that excessive utilization or weakening of the sub-models' ability may result in a decrease in the final effect.

4.5 Max Rate of Constraint

In this section, we examine the effect of β in Formula 7, which controls the max truncation rate of low-probability tokens for contrastive decoding. The results are shown in Figure 4(b). We observed that small β values (e.g., smaller than 0.45) have a minimal impact on the low-probability tokens, suggesting that these tokens are unlikely to significantly influence the model. We also found that the performance reaches its peak at around 0.5 and subsequently decreases with a further increase in β . This finding aligns with our analysis, as larger values of β tend to remove more false positive tokens. However, excessively large values of β can also result in the removal of true positive tokens, thereby reducing overall performance.

4.6 Comparison of Token Constraint

To further validate the effectiveness of our proposed constraint method for avoiding noisy tokens in contrastive decoding, we compare against the constraint method of CD. Specifically, we replace the token constraint related to scale and range in ALCD with a constraint employed in CD, while maintaining the alternative contrastive decoding technique unchanged. As shown in Table 3, our method consistently outperforms the 'constraint in CD' approach across all datasets. We attribute this improvement to the successful implementation of alternating adaptive token constraints on both scale



Figure 5: Analysis of varying decoupling steps during fine-tuning on IMCS-V2-SR dataset. 'Vanilla' refers to the performance of normal model using greedy search.

and scope in our ALCD, whereas CD relies solely on a maximum value judgment.

4.7 Affect of Decoupling Steps

To investigate how the capabilities of sub-models affect overall performance of ALCD, we conducted experiments by individually fine-tuning two subtask LLMs (i.e., \mathcal{M}_{id} and \mathcal{M}_{cl}) with varying steps while keeping normal model (i.e., \mathcal{M}_{nl}) unchanged. As illustrated in Figure 5, we observed that finetuning on sub-models effectively enhances performance, resulting in higher micro F1 scores compared to vanilla ones with 300 steps or larger. When the number of fine-tuning steps increases, the performance rises for both LLMs, while decreases after 600 steps for ChatGLM-6B and 400 steps for Qwen-7B-Chat, respectively. We believe the reason is that excessive fine-tuning steps can potentially improve the identification capabilities of \mathcal{M}_{cl} and the classification capabilities of \mathcal{M}_{id} , consequently compromising the desired decoupling effect between the two abilities. As a result, contrasting the predictions in ALCD fails to improve performance.

4.8 Case study

We also present a case study from the CHIP-MDCFNPC dataset in Table 4. It can be observed that the results generated by gready search identify entities (e.g., 'Swollen throat') that do not exist in the text or classify symptoms incorrectly (e.g., 'classifying Cough as Other'). However, our method can accurately predict *High fever: Positive*, *Cough: Positive, Fever: Positive, Common cold: Positive.* This further demonstrates the effectiveness of our approach in addressing hallucination issues.

Input	请根据下面医生与患者之间的对话片段判断临床发现实体的阴阳 性:\n患者:学校活动娃穿太少后感冒发烧,白天晚上反复发高烧 都39度多!并且发烧多次~有偶尔有点咳嗽!发高烧都是喝的美林
	退烧。{口服的小儿芬黄颗粒,利巴伟。阿莫西林,止咳冲剂}快两 天了这样高烧反复发作为什么还不好!求方法跪谢(男,8岁)\n医 生:你好,宝宝除了发烧还有其他症状\n患者:偶尔带点咳嗽。
(English Translation)	Please determine the positive and negative results of clinical findings based
	on the dialogue between the doctor and the patient below: Patient: After
	wearing too little clothing for school activities, the child caught a cold and
	had a high fever. The fever has been recurring during the day and night,
	reaching over 39 degrees Celsius each time. Occasionally, there is a slight
	cough. The fever is relieved by drinking Meilin. It has been almost two days,
	why is the high fever still not getting better? Please help (male, 8 years old).
	Doctor: Hello, besides the fever, does the baby have any other symptoms?
	Patient: Occasionally, there is a slight cough.
Gready search output	咳嗽: 其他\n高烧: 阳性\n喉咙肿大: 阳性
(English Translation)	Cough: Other High fever: Positive Swollen throat: Positive
ALCD output (ours)	高烧: 阳性\n咳嗽: 阳性\n发烧: 阳性\n感冒: 阳性
(English Translation)	High fever: Positive Cough: Positive Fever: Positive Common cold: Positive

Table 4: Case Study in the CHIP-MDCFNPC dataset.

5 Conclusion

In this paper, we propose ALCD to address hallucinations of LLMs in MIE tasks. ALCD utilizes decoupled fine-tuning process to separately learn LLM's identification and classification abilities. During inference, ALCD alternately enhances these abilities while excluding other capabilities that may result in hallucinations. We also introduce adaptive scales based on distribution similarities to enable the flexible use of identification or classification abilities. Extensive experiments conducted on two backbones have demonstrated substantial enhancement achieved by ALCD in MIE tasks.

6 Limitation

Our approach aims to decouple the identification and classification abilities of LLMs in the medical information extraction tasks and leverage their respective capabilities through alternate contrastive decoding. However, this strategy leads to an increase in both fine-tuning and inference costs. In this paper, ALCD switches between identification or classification capabilities based on simple rulebased judgment, but it is worth exploring more automatic and flexible judgment methods in future work. Furthermore, we have only investigated the effectiveness of our approach in medical information extraction tasks, and expanding our ALCD framework to other medical tasks, other domains, and other language settings is an avenue for future exploration. Exploring more robust decoupling methods and contrasting decoding techniques are also potential future research directions.

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A Appendix

A.1 Tasks and Datasets

In the experiments, we adopt a Chinese medical dataset, named PromptCBLUE (Zhu et al., 2023), including several common tasks. Due to limited resources, we select 6 tasks for validation. The statistics are in Table 1, and the dataset details are listed as follows:

- **CMeEE-V2**. Chinese medical name entity recognition. We consider "extracting entities from medical texts" as *identify* and "categorizing the entities" as *classify*.
- **CMeIE-V2**. Chinese medical entity relation extraction. We consider "recognizing the head and tail entities from medical texts" as *identify* and "categorizing the relation types between entities".
- **IMCS-V2-NER**. Medical entity recognition from the doctor-patient dialogue. We consider "identifying the medical entities from dialogues" as *identify* and "classifying the medical entity types" as *classify*.
- **CMedCausal**. Causal relation extraction for medical texts. We consider "recognizing the causal and effect words from medical texts" as *identify* and "categorizing the causal relation" as *classify*.
- **IMCS-V2-SR**. Medical normalization of the doctor-patient dialogue. We consider "extracting the normalized words from dialogues" as *identify* and "imputing the normalization labels" as *classify*.
- **CHIP-MDCFNPC**. Clinical concept finding and discrimination for the clinical report. We consider "extracting the clinical concepts from reports" as *identify* and "classifying the derived clinical concepts" as *classify*.

We show the number of prompt templates for each dataset 5. The large number of prompt templates in this PromptCBLUE allows us to better validate our ideas and the generalization ability of our model.

Furthermore, we provide some examples in Table 7 and 8.

Variations	prompt templates
CMeEE-V2	23
CMeIE-V2	37
CHIP-MDCFNPC	14
IMCS-V2-NER	25
IMCS-V2-SR	13
CMedCausal	12
Overall	124

Table 5: The number of Prompt Templates for eachdataset.

Variations	Beam search	ours
Identification_not_exit	0.0767	0.0593
Identification_pred_wrong	0.5241	0.5068
Classification_pred_wrong	0.0908	0.0775

Table 6: LLM's prediction accuracy in identificationand classification.

A.2 Ability to Mitigate Hallucinations

Furthermore, we conducted experiments to validate our method's ability to mitigate hallucinations. We evaluated the LLM's prediction accuracy in identification and classification. This includes three categories: **Identification_not_exist**: The entity or symptom identified by the LLM does not exist in the original text. **Identification_pred_wrong**: The entity or symptom identified by the LLM exists in the original text, but it is incorrect. **Classification_pred_wrong**: The LLM correctly identifies the entity or symptom, but the classification result is incorrect. Table 6 shows the results based on Qwen and CHIP-MDCFNPC datasets.

The table presents the results indicating that the effectiveness of ALCD is enhanced when evaluating classification and recognition capabilities separately. In all three categories, our ALCD method achieved lower error rates compared to the Beam Search method. This suggests that our method has some effectiveness in reducing LLM hallucinations. For the "Identification_not_exist" category, the ALCD method is more accurate in identifying entities or symptoms and generates less content that does not exist in the original text. For the "Identification_pred_wrong" category, which refers to cases where the model identifies entities or symptoms that exist in the original text but makes errors in identification, it indicates that the ALCD method has improved in correctly identifying entities or symptoms. For the "Classification_pred_wrong" category, it suggests that the ALCD method has

IMCS-V2- SR	prompt template
template1	找出当前对话中的症状,并判断阴阳性: \\n[INPUT_TEXT]\\n症状阴阳性选
	项: [LIST_LABELS]\\n答:
(English	Identify the symptoms in the current conversation and determine their positive or neg-
translation)	ative nature: [INPUT_TEXT] Symptom positive/negative options: [LIST_LABELS]
	Answer:
template2	[INPUT_TEXT]\\n根据上述对话历史,当前对话中症状有哪些?这些症状的阴
-	阳性是?\\n选项: [LIST_LABELS]\\n答:
(English	[INPUT_TEXT] Based on the previous conversation history, what are the symptoms in
translation)	the current conversation? What is the positive or negative nature of these symptoms?
	Options: [LIST_LABELS] Answer:
template3	依据之前对话内容,抽取当前对话中出现的症状实体,并明确它们的阴阳
•	性: \\n[INPUT_TEXT]\\n备选阴阳性标志: [LIST_LABELS]\\n答:
(English	Based on the previous conversation content, extract the symptom entities that appear in
translation)	the current conversation and specify their positive or negative nature: [INPUT_TEXT]
·	Available positive/negative indicators: [LIST_LABELS] Answer:

Table 7: Examples in the IMCS-V2-SR dataset.

also improved in classifying entities or symptoms. This demonstrates that ALCD has been proven to effectively minimize specific types of errors (identification and classification).

CMeIE-V2	prompt template
template1	找 出 指 定 的 三 元 组 : \\n[INPUT_TEXT]\\n实 体 间 关
	系:[LIST_LABELS]\\n答:
(English	Find the specified triplet: [INPUT_TEXT] Relationship between entities:
translation)	[LIST_LABELS] Answer:
template2	根据下述文本,提取出具有[LIST_LABELS]关系的实体
	对: \\n[INPUT_TEXT]\\n答:
(English	Based on the following text, extract entity pairs with the relationship [LIST_LABELS]:
translation)	[INPUT_TEXT] Answer:
template3	[INPUT_TEXT]\\n问题: 找出句子中描述的[LIST_LABELS]三元组的内
-	容。\\n答:
(English	[INPUT_TEXT] Question: Find the content of the triplet described in the sentence for
translation)	[LIST_LABELS]. Answer:

Table 8: Examples in the CMeIE-V2 dataset.