

Multi-stage Retrieve and Re-rank Model for Automatic Medical Coding Recommendation

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Abstract

The International Classification of Diseases (ICD) serves as a definitive medical classification system encompassing a wide range of diseases and conditions. The primary objective of ICD indexing is to allocate a subset of ICD codes to a medical record, which facilitates standardized documentation and management of various health conditions. Most existing approaches have suffered from selecting the proper label subsets from an extremely large ICD collection with a heavy long-tailed label distribution. In this paper, we leverage a multi-stage “retrieve and re-rank” framework as a novel solution to ICD indexing, via a hybrid discrete retrieval method, and re-rank retrieved candidates with contrastive learning that allows the model to make more accurate predictions from a simplified label space. The retrieval model is a hybrid of auxiliary knowledge of the electronic health records (EHR) and a discrete retrieval method (BM25), which efficiently collects high-quality candidates. In the last stage, we propose a label co-occurrence guided contrastive re-ranking model, which re-ranks the candidate labels by pulling together the clinical notes with positive ICD codes. Experimental results show the proposed method achieves state-of-the-art performance on a number of measures on the MIMIC-III benchmark.

1 Introduction

Electronic health records¹ (EHRs) contain a comprehensive repository of essential administrative and clinical data pertinent to a person’s care within a specific healthcare provider setting. In order to conduct meaningful statistical analysis, these EHR data are annotated with structured codes in a classification system known as *medical codes*. The International Classification of Diseases² (ICD) is

¹<https://www.cms.gov/Medicare/E-Health/EHealthRecords>

²<https://www.who.int/standards/classifications/classification-of-diseases>

one of the most widely-used coding systems, and it provides a taxonomy of classes, each uniquely identified by a code assigned to an episode of patient care.

The task of medical coding associates ICD codes with EHR documents. The *status quo* of assigning medical codes is a manual process, which is labour-intensive, time-consuming, and error-prone (Xie and Xing, 2018). To reduce coding errors and cost, the demand for automated medical coding has become imperative. Previous deep learning approaches regarded medical coding as an extreme multi-label text classification problem (Shi et al., 2017; Mullenbach et al., 2018; Baumel et al., 2018; Xie et al., 2019; Yuan et al., 2022), where an encoder is typically employed to learn the representations of the clinical notes and a label-specific binary classifier is subsequently constructed on top of the encoder for label predictions. However, some remaining difficulties have still posed immense challenges. First, clinical documents are lengthy (containing on average 1596 words in the MIMIC-III dataset) and noisy (including terse abbreviations, symbols, and misspellings). Second, the label set is extremely large and complex; for instance, in the 10th ICD edition, there are over 130,000 codes³. Third, the distribution of ICD codes is extremely long-tailed; while some ICD codes occur frequently, many others seldom appear, if at all, because of the rarity of the diseases. For instance, among the 942 unique 3-digit ICD codes in the MIMIC-III dataset (Johnson et al., 2016), the ten most common codes account for 26% of all code occurrences and the 437 least common codes account for only 1% of occurrences (Bai and Vucetic, 2019). To address the aforementioned challenges, we propose a novel multi-stage retrieve and re-rank framework, where the goal is to first generate a curated ICD list and then provide suggested ICD

³https://www.cdc.gov/nchs/icd/icd10cm_pcs.htm

ICD Codes:

- 401.9 Unspecified essential hypertension
- 151.9 Malignant neoplasm of stomach, unspecified site
- 285.1 Acute posthemorrhagic anemia
- 331.0 Alzheimer's disease
- 185 Malignant neoplasm of prostate
- 294.10 Dementia in conditions classified elsewhere without behavioral disturbance
- 45.16 Esophagogastroduodenoscopy [EGD] with closed biopsy
- 041.86 Helicobacter pylori [H. pylori]
- ...

Discharge Summary:

...Patient is a 83 year-old man with a history of hypertension, prostate ca (per son this has been stable, untreated for several months), and dementia who presented with an upper gastrointestinal bleed and was noted at his NSG home to have malaise, poor PO intake and low grade fevers (no note of fever in paperwork) for past 2d ... For his upper GI bleeding, the patient received IV fluids and was transfused with [**Year/Month/Day **]. He received intravenous pantoprazole therapy. Patient underwent an EGD that showed edematous mucosa and thickened folds concerning for malignancy with no evidence of active. H. pylori testing was positive and he was started on lansoprazole, amoxicillin, and clarithromycin. He had biopsies taken during endoscopy... He had dark maroon colored stool... Abnormal mucosa in the stomach (biopsy)...The patient did not have any active issues regarding his dementia. He was continued on his Namenda and Aricept during this admission.

Figure 1: An example of a medical record from the MIMIC-III dataset which includes the discharge summary, assigned ICD codes and auxiliary knowledge. We colour each code and its corresponding mentions in the discharge summary and auxiliary knowledge. We use the auxiliary knowledge of the notes to retrieve the candidate subset of the label space.

codes for a given medical record. In contrast to prior approaches, for instance, CAML (Mullenbach et al., 2018), MultiResCNN (Li and Yu, 2020) and KEPTLongformer (Yang et al., 2022), that primarily consider ICD indexing as a multi-label text classification task, we introduce a new perspective that conceptualizes the task as a recommendation problem. More precisely, we first conduct a two-stage retrieval process leveraging auxiliary knowledge and BM25 to obtain a small subset of candidate ICD codes from the large number of labels to alleviate issues caused by the label set and imbalanced label distribution. EHR auxiliary knowledge holds significant potential, but it has often been underutilized in prior studies. In addition to clinical texts, our focus centers on two code terminologies: Diagnosis-Related Group codes⁴ (DRG) and Current Procedural Terminology codes⁵ (CPT), as well as patient prescribed medications. These external sources can serve as robust indicators for predicting ICD codes. For instance, within a drug prescription, the presence of a medication like “Namenda” can

⁴<https://www.cms.gov/Medicare/Medicare-Fee-for-Service-Payment/AcuteInpatientPPS/MS-DRG-Classifications-and-Software>

⁵<https://www.ama-assn.org/amaone/cpt-current-procedural-terminology>

Auxiliary Knowledge:

CPT Codes:

- 99231 Hospital inpatient services

DRG Codes:

- 2402 Digestive Malignancy
- ...

Prescriptions:

- Bicalutamide
- Pantoprazole
- Midazolam
- Namenda
- Donepezil
- Atorvastatin
- Potassium Chloride
- ...

strongly imply a likelihood of Alzheimer’s disease, as depicted in Figure 1. Subsequently, we design a re-ranking model via co-occurrence guided contrastive learning to refine the candidate set, which can deal with lengthy clinical notes and generate semantically meaningful representations via the pre-trained language model and leverage code co-occurrence to generate co-occurrence-aware label representations. The co-occurrence of codes in clinical texts yields valuable insights into the interconnections among different diseases or conditions. As illustrated in Figure 1, the code for “Dementia in conditions classified elsewhere without behavioral disturbance” (294.10) can be easily found in the text; however, inferring the code “Alzheimer’s disease” (331.0) presents a more intricate challenge with less explicit clues. Fortunately, a robust association exists between these two diseases, with “Alzheimer’s disease” serving as a prevalent cause of “dementia”. This linkage can be effectively captured as these two diseases frequently co-occur within the clinical notes. This empowers us to gain a deeper understanding of the contexts, which could mitigate the limitation of long-tailed label distributions as rare labels might be suggested based on these relationships. We train the re-ranking model via contrastive learning as it has strong discriminative power that can extract features uniquely associated with each class, which empowers the model to make more accurate recommendations.

To summarize, the major contributions of this paper are:

- We formalize the medical coding task as a recommendation problem and present a novel multi-stage retrieve and re-rank framework to make more accurate predictions by ruling out the irrelevant codes before ranking, rather than making direct predictions on the entire large label set.
- To address the large label set and long-tailed distribution issues, in the two-stage retrieval process, we use external knowledge and BM25 to retrieve a subset of candidate labels from the large label space. We further leverage the code co-occurrence in the re-ranking stage to capture the internal connections among the codes.
- We apply contrastive learning in the re-ranking stage. It effectively pulls together the

150 representations of a clinical note and its cor-
151 responding golden truth labels, which allows
152 the model to make more accurate predictions.

153 2 Related Work

154 The automatic ICD indexing task is well estab-
155 lished in the healthcare domain. Extensive re-
156 search using deep learning has been dedicated to
157 ICD indexing, including recurrent-based neural
158 networks (RNNs), convolution-based neural net-
159 works (CNNs), and their variations. These archi-
160 tectures are able to extract and categorize semantic
161 features, reducing the need for medical domain
162 expertise during the traditional feature selection
163 stage seen in conventional algorithms (Teng et al.,
164 2023). The ICD indexing task is formulated as
165 a multi-label classification problem in these ap-
166 proaches. Mullenbach et al. (2018) introduced a
167 combination of CNN with an attention mechanism
168 to effectively capture pertinent information within
169 clinical texts for each ICD code. Building on this
170 foundation, Xie et al. (2019) enhanced the CNN
171 attention model by integrating a multi-scale fea-
172 ture attention technique. Many CNN variants were
173 subsequently introduced to address the challenges
174 posed by lengthy and noisy clinical texts, including
175 MultiResCNN (Li and Yu, 2020), DCAN (Ji et al.,
176 2020), and EffectiveCAN (Liu et al., 2021). RNN-
177 based models, renowned for their capacity to cap-
178 ture contextual information across input texts, have
179 also been widely used for ICD indexing. Shi et al.
180 (2017) proposed a character-aware Long Short-
181 Term Memory (LSTM) recurrent network to learn
182 the underlying representations of clinical texts. Xie
183 and Xing (2018) introduced a tree-of-sequences
184 LSTM architecture alongside adversarial learning
185 to capture hierarchical relationships among ICD
186 codes. Additionally, Baumel et al. (2018) presented
187 a Hierarchical Attention-Bidirectional Gated Re-
188 current Unit (HA-GRU) model, facilitating docu-
189 ment labeling by identifying sentences relevant to
190 each ICD code. LAAT (Vu et al., 2020) used a bidi-
191 rectional Long-Short Term Memory (BiLSTM) en-
192 coder and a customized label-wise attention mech-
193 anism to cultivate label-specific vectors across dis-
194 tinct clinical text fragments.

195 To address the hierarchical relationships intrin-
196 sic to ICD codes, Graph Convolutional Neural Net-
197 works (GCNNs) (Kipf and Welling, 2017) have
198 emerged as a powerful tool. Rios and Kavuluru
199 (2018) and Xie et al. (2019) used GCNNs to capture

200 both the hierarchical interplay among ICD codes
201 and the semantic information specific to each code.
202 HyperCore (Cao et al., 2020) took a comprehensive
203 approach by considering both code hierarchy and
204 code co-occurrence, employing GCNNs to learn
205 code representations within the co-graph.

206 Incorporating external knowledge beyond ICD
207 code information has also gained traction. Bai and
208 Vucetic (2019) introduced a Knowledge Source
209 Integration (KSI) model that integrates external
210 knowledge from Wikipedia. This integration calcu-
211 lated matching scores between clinical notes and
212 disease-related Wikipedia documents, in order to
213 enrich the available information for ICD predic-
214 tions. Additionally, Yuan et al. (2022) proposed a
215 Multiple Synonym Matching Network (MSMN) to
216 use synonyms of ICD codes, enhancing the qual-
217 ity of code representation learning. Expanding on
218 this, Yang et al. (2022) integrated a pre-trained lan-
219 guage model with three domain-specific knowledge
220 sources: code hierarchy, synonyms, and abbrevia-
221 tions. This fusion of knowledge sources contributes
222 significantly to the performance of ICD classifica-
223 tion.

224 3 Method

225 3.1 A Multi-stage Framework

226 We formulate the medical coding task as a rec-
227 ommendation task, given medical records $\mathcal{D} =$
228 $\{d_1, d_2, \dots, d_N\}$ and a set of ICD codes $\mathcal{Y} =$
229 $\{y_1, y_2, \dots, y_L\}$ with associated external auxiliary
230 knowledge \mathcal{K} . We construct the label information
231 as a graph structure \mathcal{G} , using code co-occurrence
232 relations, and we train a multi-stage recommender
233 system \mathcal{R} , based on the text information \mathcal{D} , con-
234 structed label information \mathcal{G} , and the external auxil-
235 iary knowledge \mathcal{K} . The system \mathcal{R} needs to predict
236 the relevant labels given a document $d \notin \mathcal{D}$.

237 In this section, we present a multi-stage retrieve
238 and re-rank framework for ICD indexing, which
239 is shown in Figure 2. Our model is composed of
240 a two-stage retrieval process that uses auxiliary
241 knowledge of the EHR and BM25 to obtain a short-
242 ened candidate list, and a re-ranking process that
243 conducts code co-occurrence guided contrastive
244 learning to further improve the recommended ICD
245 list.

246 3.2 The Retrieval Stage

247 **Using Auxiliary Knowledge** To retrieve the candi-
248 date list using auxiliary knowledge, we incorpo-

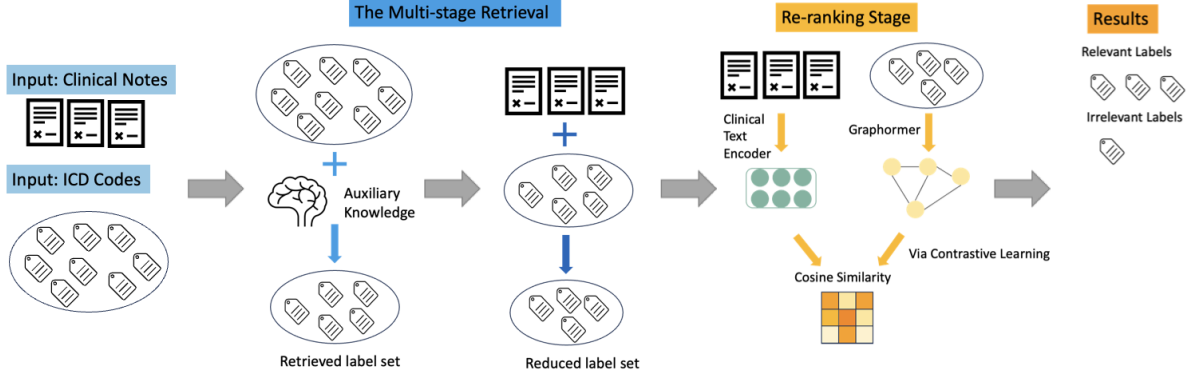


Figure 2: Overview of the proposed multi-stage retrieve and re-rank framework. The model first leverages auxiliary knowledge and BM25 to retrieve a candidate list from the full label space, then use a re-rank model that leverages the code co-occurrence guided contrastive learning to generate the final relevant labels.

rate insights from three external sources of knowledge: diagnosis-related group (DRG) codes, current procedural terminology (CPT) codes, and medications prescribed to patients. DRG codes are used by hospitals and healthcare providers to classify patients into groups based on their diagnosis, treatment, and length of stay. These codes are used for reimbursement purposes, and they help determine the amount that healthcare providers are remunerated for their services. DRG codes are further classified into medical DRGs (which exclude operating room procedures) and surgical DRGs. CPT codes are used to describe medical procedures and services provided by healthcare providers. They provide a standardized way of documenting and billing for medical services. CPT codes are used by insurance companies to determine reimbursement rates for healthcare providers. Such code terminologies significantly contribute to the refinement of ICD indexing. Moreover, the medications prescribed to patients offer a wealth of predictive information for ICD codes. These prescriptions often mark the conclusion of a patient’s care episode. As patients approach the conclusion of their treatment, the prescribed medications serve a critical role in managing their conditions. Consequently, these medications emerge as potent indicators of underlying health conditions or diagnoses. Their inclusion in the retrieval process greatly enhances the accuracy and relevance of the corresponding ICD code recommendations.

Given a clinical note d , we retrieve the candidate ICD list by calculating the auxiliary knowledge and label co-occurrence matrix using conditional probabilities, i.e., $P(y_i | k_j)$, which denote the prob-

abilities of occurrence of ICD y_i when auxiliary knowledge k_j appears.

$$P(y_i | k_j) = \frac{C_{y_i \cap k_j}}{C_{k_j}}, \quad (1)$$

where $C_{y_i \cap k_j}$ denotes the number of co-occurrences of y_i and k_j , and C_{k_j} is the number of occurrences of k_j in the training set. To avoid the noise of rare co-occurrences, a threshold η filters noisy correlations. \tilde{K}_j denotes the selected ICD set for auxiliary knowledge j .

$$\tilde{K}_j = \{y_i | P(y_i | k_j) > \eta, i = 1, \dots, L\}, \quad (2)$$

where L is total number of ICD codes in the label set. We then join the ICD codes retrieved from the auxiliary knowledge co-occurrences for the DRG codes, CPT codes and prescribed drugs to form the candidate ICD subset $\mathcal{C}_{\text{auxiliary}}$:

$$\mathcal{C}_{\text{auxiliary}}(d) = \tilde{K}_{\text{DRG}}(d) \cup \tilde{K}_{\text{CPT}}(d) \cup \tilde{K}_{\text{drug}}(d), \quad (3)$$

where $\mathcal{C}_{\text{auxiliary}} \subseteq \mathcal{Y}$.

Using BM25 The retrieval stage using auxiliary knowledge incorporates the co-relations between ICD codes and external knowledge, but ignores the relationship between clinical texts and labels. To increase the recall of the retrieval stage, we adopt BM25 (Robertson and Walker, 1994) to allow lexical matching between the medical documents and labels on the retrieved candidate list $\mathcal{C}_{\text{auxiliary}}$. Given a medical record d and an ICD code y , the score between d and y is calculated as:

$$\text{BM25}(d, y) = \sum_{w \in d \cap t_y} \text{IDF}(w) \frac{\text{TF}(w, t_y) \cdot (k_1 + 1)}{\text{TF}(w, t_y) \cdot k_1 (1 - b + b \frac{|Y|}{\text{avgdl}})}, \quad (4)$$

and

$$avgdl = \frac{1}{|\mathcal{Y}|} \sum_{y \in \mathcal{Y}} |t_y|, \quad (5)$$

where t_y represents the words in the label descriptors, $|\mathcal{Y}|$ is the length of the label descriptors in words, $avgdl$ is the average length of text information in the label.

When the BM25 score between d and y_i exceeds a certain threshold θ , we add y_i as a candidate of d :

$$\mathcal{C}_{\text{BM25}}(d) = \{y_i | \text{BM25}(d, y_i) > \theta, y_i \in \mathcal{C}_{\text{auxiliary}}\} \quad (6)$$

Given a clinical note d , its candidate ICD set is first generated by using the auxiliary knowledge in the retrieval stage and then reduced by using BM25, where $\mathcal{C}_{\text{BM25}} \subseteq \mathcal{C}_{\text{auxiliary}}$ and $\mathcal{C}_{\text{auxiliary}} \subseteq \mathcal{Y}$.

3.3 The Re-ranking Stage

Clinical Text Encoder Encouraged by the success of the pre-trained language model Longformer (Beltagy et al., 2020) in dealing with longer texts, we use a Clinical-Longformer (Li et al., 2023), specifically pre-trained in the medical domain, as a text encoder. Given a medical document d as input that consists of a sequence of tokens:

$$d = \{[\text{CLS}], x_1, x_2, \dots, x_{n-2}, [\text{SEP}]\}, \quad (7)$$

where [CLS] and [SEP] are two special tokens that indicate the beginning and end of the sequence, and n is the sequence length, the Clinical-Longformer encodes the tokens and outputs the hidden representations for each token:

$$H_{\text{hidden}} = \text{ClinicalLongformer}(d), \quad (8)$$

where $H_{\text{hidden}} \in \mathbb{R}^{n \times h_e}$, and h_e is the hidden size. Following previous work (Wang et al., 2022; Yang et al., 2022), we use the hidden state of the [CLS] token to represent the document, denoted as H_{T} .

Label Encoder The occurrence of two ICD codes together in clinical texts frequently indicates a simultaneous presence or a causal connection between specific diseases. This implies that the codes representing these interconnected diseases often manifest together within clinical notes. We employ a Graphormer (Ying et al., 2021) to incorporate the co-occurrence relationships among ICD codes. Unlike the original GNN, Graphormer models graphs using Transformer layers (Vaswani et al., 2017) with spatial encoding and edge encoding, which could effectively encode the structural information (i.e., code co-occurrence) of a

graph into the model. We create a directed code co-occurrence graph $\mathcal{G} = (\mathcal{Y}, \mathcal{E})$, where node set \mathcal{Y} is the labels and edge set \mathcal{E} denotes the co-occurrence relations. This graph is constructed using the code co-occurrence matrix, which has been used as the edge matrix for the graph. We create the code co-occurrence matrix by using the correlated relationship between labels based on conditional probabilities. This approach encapsulates the interdependence between various ICD codes in a quantifiable manner, offering valuable insights into the underlying connections among disease codes within the clinical texts. To be more specific, we calculate the probability of occurrence of label y_j when label y_i appears as follows:

$$P(y_j | y_i) = \frac{C_{y_i \cap y_j}}{C_{y_i}} \quad (9)$$

where $C_{y_j \cap y_i}$ denotes the number of co-occurrences of y_i and y_j , and C_{y_i} is the number of occurrences of y_i in the training set. To facilitate graph construction, we binarize the correlation probability $P(y_j | y_i)$. This entails converting the probability values into binary values, indicating whether a correlation exists (or not) between two labels. The operation can be written as:

$$\mathcal{E}_{ij} = \begin{cases} 0, & \text{if } P(y_j | y_i) < \lambda \\ 1, & \text{if } P(y_j | y_i) \geq \lambda, \end{cases} \quad (10)$$

where \mathcal{E} is the binary correlation matrix also known as the edge set, and λ is the hyper-parameter threshold to filter the noise edges. $\lambda = 1$ in our experiment, which means that an edge is formed when the two labels in each pair always appear together.

To encode the graph \mathcal{G} , we first generate the initial node features using the ICD full descriptors for each code y via Clinical-Longformer:

$$y = \{[\text{CLS}], x_1, x_2, \dots, x_{n-2}, [\text{SEP}]\}, \quad (11)$$

$$H_v = \text{ClinicalLongformer}(y),$$

where y represents a sequence of words in the label descriptors of label y , $H_v \in \mathbb{R}^{n \times h_e}$, and h_e is the hidden size. We use the hidden state of the first token ([CLS]) for representing the initial node feature denoted as H_{node}^i for the i^{th} label.

With all initial node features stacked as a matrix $V = \{H_{\text{node}}^1, H_{\text{node}}^2, \dots, H_{\text{node}}^L\}$, where $V \in \mathbb{R}^{h_e \times L}$, a standard self-attention layer is then used for feature migration. To leverage the structural information, a novel spatial encoding method is

used to modify the Query-Key product matrix $A^{\mathcal{G}}$ in the self-attention layer:

$$A_{ij}^{\mathcal{G}} = \frac{(H_{\text{node}}^i W_Q^{\mathcal{G}})(H_{\text{node}}^j W_K^{\mathcal{G}})^T}{\sqrt{h_e}} + b_{\phi(y_i, y_j)}, \quad (12)$$

where $W_Q^{\mathcal{G}}$ and $W_K^{\mathcal{G}}$ are layer-specific weight matrices, and $\phi(y_i, y_j)$ is the spatial relation between y_i and y_j in graph \mathcal{G} , and the function $\phi(\cdot)$ is defined as the connectivity between the nodes in the \mathcal{G} , which is the co-occurrence relation among labels. $b_{\phi(y_i, y_j)}$ is a learnable scalar indexed by $\phi(y_i, y_j)$, and shared across all layers. The attention score $A_{ij}^{\mathcal{G}}$ then has been used to aggregate the multi-head attention for the final output:

$$h^{l+1} = \text{MHA}(\text{LN}(h^l)) + h^l, \quad (13)$$

where LN denotes the layer normalization, MHA denotes the multi-head self-attention, h^l and $h^{l+1} \in \mathbb{R}^{L \times h_e}$ indicate the node representation of the l^{th} and $(l+1)^{\text{th}}$ layers. We use the last layer to represent the label feature denoted as H_L . For more details on the full structure of Graphormer, please refer to the original paper (Ying et al., 2021).

Contrastive Learning for Re-ranking Now we construct a code co-occurrence guided contrastive learning framework. Unlike supervised learning that aims to understand “what is what”, contrastive learning adopts a different perspective by learning “what is similar or dissimilar to what”. In our problem setting, we focus on the distances between a clinical document and its associated ICD codes, rather than solely between samples themselves. We consider the ground truth labels as positive samples, while the negative samples comprise all the other labels within the label space. Given H_T , the representation for a clinical note d and the set of representations of its corresponding ICD codes denoted as H_L^+ , we denote the representations of N negative ICD codes randomly chosen from the ICD codes of the documents in the batch (batch size is N) which are not ICD codes of document d as H_L^- . Contrastive learning aims to learn the effective representations by pulling d and H_L^+ together while pushing apart d and H_L^- , represented as S and D , respectively, in the equation below. The contrastive loss can be defined as:

$$\mathcal{L} = -\log \frac{S/\tau}{S/\tau + D/\tau}, \quad (14)$$

where $S = \exp(\sum_{c \in L_L^+} \cos(H_T, c)/|H_L^+|)$, $D = \exp(\sum_{c' \in L_L^-} \cos(H_T, c')/N)$, and τ is the temperature hyper-parameter.

4 Experiments

4.1 Dataset and Pre-processing

We conduct our experiments on the publicly available benchmark MIMIC-III (Johnson et al., 2016) dataset that contains a variety of patient data types, including discharge summaries, demographic details, interventions, laboratory results, physiologic measures, and medication information. Following previous work, we are interested in the de-identified discharge summaries with annotated ICD-9 codes. There are 52,722 discharge summaries and 8,922 unique ICD-9 codes in the dataset. We mainly use three major data resources from the dataset: (1) de-identified discharge summaries (from the NOTEVENTS table); (2) ICD-9 codes (from DIAGNOSES_ICD and PROCEDURES_ICD tables); and (3) auxiliary knowledge including DRG codes, CPT codes and drug prescriptions (from DRGCODES, CPTEVENTS, and PRESCRIPTIONS tables).

To preprocess the clinical notes, we first remove all de-identified information, then replace punctuation and atypical alphanumeric character combinations (e.g., ‘3a’, ‘4kg’) with white space, and lowercase every token. We truncate the discharge summaries at a maximum length of 4000 tokens. We follow Mullenbach et al. (2018) to form two settings: full codes (MIMIC-III-full) and top-50 frequent codes (MIMIC-III-top 50). In MIMIC-III-full, there are 47,719 discharge summaries for training, with 1,632 for validation, and with 3,372 for testing.

4.2 Implementation and Evaluation

We implement our model in PyTorch (Paszke et al., 2019) on a single NVIDIA A100 40G GPU. We use the Adam optimizer and early stopping strategies using Micro-F1 score over the validation set as stopping criterion to avoid over-fitting. We set the initial learning rate as $5e-5$ with batch size 16. We choose a learning rate scheduler which is warmed up with cosine decay, and the warm up ratio is set to 0.1.

For evaluating the performance of our proposed model, we employ three commonly used metrics: F1-score (Micro and Macro), AUC (Micro and Macro), and precision at K (P@K).

5 Results and Discussion

In order to assess the efficacy of our proposed framework, we compare with the existing state-of-the-art

Models	MIMIC-III-full						MIMIC-III-top 50				P@5
	AUC		F1		P@K		AUC		F1		
	Macro	Micro	Macro	Micro	P@8	P@15	Macro	Micro	Macro	Micro	
CAML (Mullenbach et al., 2018)	0.895	0.986	0.088	0.539	0.709	0.561	0.875	0.909	0.532	0.614	0.609
DR-CAML (Mullenbach et al., 2018)	0.897	0.985	0.086	0.529	0.690	0.548	0.884	0.916	0.576	0.633	0.618
MultiResCNN (Li and Yu, 2020)	0.910	0.986	0.085	0.552	0.734	0.584	0.899	0.928	0.606	0.670	0.641
LAAT (Vu et al., 2020)	0.919	0.988	0.099	0.575	0.738	0.591	0.925	0.946	0.666	0.715	0.675
Joint-LAAT (Vu et al., 2020)	0.921	0.988	0.107	0.575	0.735	0.590	0.925	0.946	0.661	0.716	0.671
EffectiveCAN (Liu et al., 2021)	0.915	0.988	0.106	0.589	0.758	0.606	0.915	0.938	0.644	0.702	0.656
MSMN (Yuan et al., 2022)	0.950	0.992	0.103	0.584	0.752	0.599	0.928	0.947	0.683	0.725	0.680
KEPTLongformer (Yang et al., 2022)	-	-	0.118	0.599	0.771	0.615	0.926	0.947	0.689	0.728	0.672
Ours	0.949	0.995	0.114	0.603	0.775	0.623	0.927	0.947	0.687	0.732	0.685

Table 1: Comparison to previous methods across three main evaluation metrics MIMIC-III dataset. Bold: the optimal values.

(SotA) models, as outlined in Table 1. The top score for each metric is denoted in bold. As shown, our model outperforms in the majority of evaluation metrics, with the exception of Macro-AUC and Macro-F1 on the MIMIC-III-full and MIMIC-III-top 50. Notably, our model achieves comparable performance on Micro-F1 and Micro-AUC, and improves precision at K on both MIMIC-III-full and MIMIC-III-top 50. These results provide solid evidence to validate the efficacy of integrating auxiliary knowledge in the retrieval stage and leveraging code co-occurrence guided contrastive learning in the re-ranking stage.

As the occurrence frequencies of each ICD codes are imbalanced, our focus lies in assessing the efficacy of our model specifically on infrequently appearing ICD codes. We categorize the ICD codes into four groups based on their occurrences in the training set: [0, 10), [10, 50), [50, 500), and [500,). Figure 3 illustrates the distribution of ICD codes and their occurrence percentages across the four categorized groups in the training set, which show that the distribution of ICD frequency is highly biased, conforming to a long-tail distribution. Figures 3b and 3c present the performance of our model on MIMIC-III-full in comparison to the CAML baseline (Mullenbach et al., 2018) across the four ICD groups on Macro-AUC and Micro-F1, respectively. Our model demonstrates significant improvements for both frequent and infrequent labels on both metrics.

To confirm the specific contributions of these modules in terms of enhancing both the effectiveness and robustness of the model, we conduct ablation studies with three different settings: (a) we examine the effectiveness of using auxiliary knowledge in the retrieval stage by removing the retrieval stage and rank the ICD codes on the whole label set;

Methods	F1		P@K	
	Macro	Micro	P@8	P@15
Full Model	0.114	0.603	0.775	0.623
w/o auxiliary knowledge	0.097	0.579	0.748	0.587
embedded w/ Clinical-BERT	0.083	0.548	0.711	0.546
w/o Graphormer	0.102	0.583	0.753	0.591

Table 2: Ablation experiment results on the MIMIC-III-full. Bold: the optimal values.

(b) we examine the influence of different embedding methods by replacing the Clinical-Longformer with Clinical-BERT; and (c) we test the effectiveness of label embedding by replacing the encoding of the label with the average of words embeddings in the label descriptors. The experimental results are shown in Table 2. We also conduct case studies to qualitatively understand the effects of incorporating the label co-occurrence and the auxiliary knowledge. Two have been presented in the Appendix A.

Effectiveness of Using Auxiliary Knowledge for Retrieval We employ three distinct types of auxiliary knowledge in the retrieval stage: DRG codes, CPT codes, and drug prescriptions. As shown in Table 2, removing auxiliary knowledge leads to a decline in performance, indicating the pivotal role of the retrieval stage. This outcome further provides evidence that external knowledge effectively addresses the challenge presented by a large pool of potential ICD codes. Through integrating external knowledge, the retrieval stage attains the capability to refine the candidate list using the co-occurrence relationships between ICD codes and the auxiliary knowledge, thereby amplifying both the efficiency and accuracy of the re-ranking stage. The selection of an appropriate candidate list for a given medical record hinges upon a hyperparameter, specifically the threshold ρ governing the co-occurrence between auxiliary knowledge

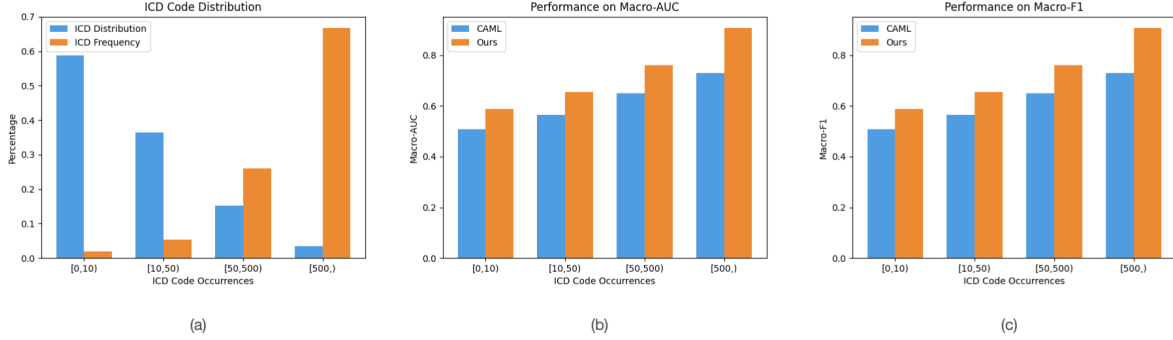


Figure 3: (a) ICD code distribution. (b) Macro-AUC performance comparison of our model and CAML on ICD codes at different frequency. (c) Micro-F1 performance comparison of our model and CAML on ICD codes at different frequency.

and ICD codes. The choice of ρ determines the candidate numbers that implicitly affect the overall performance of the model. Setting $\rho = 0.005$, the candidate list guarantees inclusion of 99.22% of the gold-standard ICD codes, resulting in an average of 1,460 codes within the subset. Notably, this accounts for approximately one-sixth of the complete code set. A further reduction using BM25 limits candidate list to 1299 on average.

Comparison of Clinical-Longformer and Clinical-BERT Increasing the maximum token limit is important in the context of clinical notes analysis as clinical texts are lengthy. Specially, in the MIMIC-III dataset, the average length of the discharge summaries is 1,596. Given the substantial token volume within the clinical notes, encoding a maximum number of tokens prior to downstream analysis becomes a pivotal requirement, which facilitates robust and meaningful subsequent analysis. To test the effectiveness of using longer sequences, we compare the model performance of Clinical-Longformer and a BERT-based pre-trained language model (i.e., Clinical-BERT) which can only encode a maximum of 512 tokens. As shown in Table 2, Clinical-Longformer substantially outperforms Clinical-BERT, indicating that the importance of maximum token limit over language models in automatic medical coding task.

Effectiveness of Learning Label Features Using Code Co-occurrence The graph structure has been shown to be effective in modeling code correlations and the Graphormer efficiently learns code representations. The findings presented in Table 2 highlight the affirmative impact of integrating code

co-occurrence into label representations. By using Graphormer, the model effectively captures and exploits the intricate connections and interdependencies among the labels, thereby improving the overall performance. This indicates that incorporating code co-occurrence information within the Graphormer empowers the model to gain insights from the collaborative behaviours of the labels, consequently facilitating a more holistic comprehension of the underlying label co-relations.

6 Conclusion

In this paper, we regard the medical coding task as a recommendation problem and present a novel multi-stage retrieve and re-rank framework. The primary objective of the proposed framework is twofold: to construct a curated list of ICD codes and subsequently to further refine the candidate list for a given medical record. Specifically, we first conduct a two-step retrieval process, incorporating auxiliary knowledge and the BM25 algorithm. This approach retrieves of a concise subset of the candidate list, mitigating the challenges of a very large and imbalanced label distribution. We then use a re-ranking model to refine the previously obtained candidate list, which employs code co-occurrence guided contrastive learning. Experimental results demonstrate that our proposed framework outperforms the previous SotA, which suggests that it provides more precise and contextually grounded ICD recommendations for the given medical records. In the future, our proposed framework may be extended with more external knowledge such as the Unified Medical Language System (UMLS) and code synonymy.

634 Limitations

635 Our usage of auxiliary knowledge is limited using
636 external knowledge including DRG codes, CPT
637 codes, and drug prescriptions only. Other knowl-
638 edge including disease-symptom, disease-lab rela-
639 tions, Unified Medical Language System (UMLS),
640 and others could also be potentially useful for the
641 auto ICD coding task.

642 Our study is constrained by its evaluation limited
643 to MIMIC-III-full and MIMIC-III-top 50 datasets,
644 primarily concentrated on common disease. To
645 comprehensively assess the model’s performance
646 on rare diseases, future work could benefit from a
647 curated list of rare diseases validated by domain
648 experts.

649 Ethics Statement

650 We are using the publicly available clinical dataset
651 MIMIC-III, which contains deidentified patient in-
652 formation. We do not see any ethics issue here in
653 this paper.

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A Case Studies

We conducted case studies to qualitatively explore the impacts of integrating label co-occurrence (illustrated in Figure 4) and auxiliary knowledge (depicted in Figure 5). We compared the full model with models that did not integrate the label co-occurrence and the external knowledge on the predictions of two patient records. For each patient, we present the discharge summary, ground truth ICD codes, label co-occurrence information, or auxiliary knowledge information, along with the predicted ICD codes from the full model and ablated models.

Case 1: Effectiveness of Incorporating Label Co-occurrence	
Discharge Summary	<p>Chief Complaint: heroin overdose</p> <p>Major Surgical or Invasive Procedure: s/p intubation, s/p cvc placement</p> <p>SINGLE SUPINE AP PORTABLE CHEST RADIOGRAPH: An endotracheal tube is in optimal position terminating 3.5 cm above the carina. A nasogastric tube coils within the stomach, with the tip terminating in the distal stomach. No pneumothorax or large pleural effusions are seen. There is diffuse opacity overlying the entire right lung and major portion of the left upper lung, which likely represent diffuse pulmonary edema, ARDS or hemorrhage. No acute osseous abnormality seen.</p> <p>IMPRESSION: Diffuse opacities in the right lung and left upper lung, likely represents pulmonary edema, ARDS or hemorrhage. ET tube in optimal position.</p> <p>Brief Hospital Course:</p> <p>TITLE: Medical ICU Resident/MERIT Admission Note Reason for transfer to the MICU: post-arrest management</p> <p>History of Present Illness and MICU Course: Mr. [Known last name 12303**] is a 19 year old male with a history of polysubstance abuse most significant for intravenous heroin, who presented to the [Hospital1 18**] ED for post-cardiac arrest care in the setting of an apparent heroin overdose. He was transferred from [Hospitals3 **].</p> <p>Briefly, he was discharged from a rehab center in [State 108**] one day prior to admission. Last night, at 3AM on [2145-4-10**], his mother found him down with needles around. She immediately called 911 and initiated CPR. He was intubated in the field per the [Location (un) 5700**] service ambulance record and dopamine and levofed were initiated; his pupils were reportedly fixed and dilated at that point. Patient cooling was also performed via ice packs. At [Hospital3 **], advanced cardiopulmonary life support was continued for over 40 minutes at which point the patient regained a pulse. His labs at [Hospital1 **] were notable for a WBC 14.2, Hct 45.2, Plt 210. Sodium 145, k 6.5, cl 107, co2 14, gluc 247, bun 19, cr 2.3, ca [43**]. He was then transferred to the [Hospital1 18**] ED. In the [Hospital1 18**] ED he was on three pressors (epinephrine, levophed, and vasopressin). His blood cases were checked twice and showed 6.79/86/61 -->6.87/67/82. He was transferred to the MICU. In the MICU, he did not have a femoral pulse. A cardiac monitor was placed and he was noted to have pulseless electrical activity. ACLS was initiated. He received sodium bicarbonate, calcium chloride, d50, NS, and boluses of epinephrine. His rhythm converted to ventricular fibrillation and he was shocked. He then converted to PEA and regained a pulse after another bolus of epinephrine. The family was present. The code lasted just under ten minutes. After discussion with the family, the decision was made not to escalate care (see Dr. [Last Name (STitle) **]????? note). He remained on three pressors with ventilatory support. Within one hour he became bradycardic and expired. See written death note in the chart. The organ bank declined the case for donation. The Medical examiner accepted the case. The family declined discretionary autopsy. Death report and other necessary documentation was filed.</p>
Ground Truth ICD Codes	427.5 Cardiac arrest; 96.71 Continuous invasive mechanical ventilation for less than 96 consecutive hours; V49.86 Do not resuscitate status; 518.81 Acute respiratory failure; 785.51 Cardiogenic shock; 99.60 Cardiopulmonary resuscitation, not otherwise specified; 304.71 Combinations of opioid type drug with any other drug dependence, continuous
Examples of Label Co-occurrence Information	<ol style="list-style-type: none"> 427.5 Cardiac arrest relates to 785.51 Cardiogenic shock 96.71 Continuous invasive mechanical ventilation for less than 96 consecutive hours relates to V49.86 Do not resuscitate status
Predictions of Full Model	427.5 Cardiac arrest; 96.04 Insertion of endotracheal tube; 96.71 Continuous invasive mechanical ventilation for less than 96 consecutive hours; 965.01 Poisoning by heroin; 99.6 Cardiopulmonary resuscitation, not otherwise specified; 785.51 Cardiogenic shock; V49.86 Do not resuscitate status
Predictions of No Label Co-occurrence	427.5 Cardiac arrest; 96.04 Insertion of endotracheal tube; 96.71 Continuous invasive mechanical ventilation for less than 96 consecutive hours; 965.01 Poisoning by heroin; 99.6 Cardiopulmonary resuscitation, not otherwise specified; 969.6 Poisoning by psychodysleptics (hallucinogens)

Figure 4: Case study on the effectiveness of incorporating label co-occurrence. Correctly predicted labels are marked in green and the incorrect ones are marked in red.

In Case 1, the ground truth ICD codes include ‘785.51 Cardiogenic shock’ and ‘V49.86 Do not resuscitate status’ are not explicitly mentioned in the discharge summary. The observed label co-occurrence between ‘427.5 Cardiac arrest’ and ‘785.51 Cardiogenic shock’, as well as co-relation between ‘96.71 Continuous invasive mechanical ventilation for less than 96 consecutive hours’ and ‘V49.86 Do not resuscitate status’ provide strong indicator suggesting the presence of the codes ‘785.51’ and ‘V49.86’. Without the label co-occurrence signals, the ablated model missed the predictions of codes ‘785.51’ and ‘V49.86’,

807 indicating a failure to leverage latent label informa-
 808 tion.

809 In Case 2, the patient has been diagnosed with
 810 ‘244.9 Unspecified acquired hypothyroidism’ with
 811 less explicit information in the discharge summary.
 812 Notably, the presence of the medication ‘Levothy-
 813 roxine’ in the drug prescription, an element of
 814 auxiliary knowledge, suggests that the patient is
 815 likely to have acquired hypothyroidism. The ab-
 816 lated model, lacking of the auxiliary knowledge,
 817 misses the prediction of code ‘244.9’. From the
 818 aforementioned cases 1 and 2 highlight the benefits
 819 of incorporating label co-occurrence and auxiliary
 knowledge, respectively.

Case 2: Effectiveness of Incorporating Auxiliary Knowledge	
Discharge Summary	<p>Chief Complaint: Subdural hematoma</p> <p>History of Present Illness: 78 year-old male with hypertension, ITP on Rituximab transferred from OSH for further management of SDH. Felt poorly yesterday. Woke up this morning with severe HA. Unresponsive in EMS. Went to [**Hospital1 **], found to have decerebrate posturing, fixed and dilated pupils. CT head with large left-sided SDH with 2mm shift, and transtorial herniation. Intubated (succ/etomidate), mannitol. Also received atropine for unknown reason. ... Discussed with neurosurgery, radiology; determined to benefit in intervention at this point. Per report from ED resident, patient converted to CMO, and awaiting arrival of family prior to extubation. Propofol restarted for comfort. On transfer to ICU, 67, 151/65, 10, 100% AC 10/500 PEEP 5, FIO2 100%. On the floor, patient is intubated and not responsive.</p> <p>Brief Hospital Course: 78M with hypertension, ITP with subdural hematoma complicated by mass effect. Expired shortly after admission.</p> <p>#. Subdural hematoma: In context of thrombocytopenia and known hypertension. Complicated by mass effect. Patient noted initially to be decorticate. Unresponsive with fixed/dilated pupils off of sedation. With down titrating ventilatory support, patient with rare breaths and with low tidal volumes. Discussed with family; plan for comfort.</p> <p>#. ITP: Thrombocytopenic. Held off on platelet transfusion as would not change outcome.</p> <p>#. Hypertension: Held anti-hypertensives.</p>
Ground Truth ICD Codes	V58.65 Long-term (current) use of steroids; 401.9 Unspecified essential hypertension; 432.1 Subdural hemorrhage; 96.71 Continuous invasive mechanical ventilation for less than 96 consecutive hours; 244.9 Unspecified acquired hypothyroidism; 348.4 Compression of brain; 287.31 Immune thrombocytopenic purpura
Examples of Using Auxiliary Knowledge	1. Levothyroxine relates to 244.9 Unspecified acquired hypothyroidism
Predictions of Full Model	287.31 Immune thrombocytopenic purpura; 348.4 Compression of brain; 348.5 Cerebral edema ; 401.9 Unspecified essential hypertension; 432.1 Subdural hemorrhage; 96.71 Continuous invasive mechanical ventilation for less than 96 consecutive hours; 244.9 Unspecified acquired hypothyroidism; E888.9 Unspecified fall ; V66.7 Encounter for palliative care
Predictions of No Auxiliary Knowledge	287.31 Immune thrombocytopenic purpura; 348.4 Compression of brain; 348.5 Cerebral edema ; 401.9 Unspecified essential hypertension; 432.1 Subdural hemorrhage; 96.71 Continuous invasive mechanical ventilation for less than 96 consecutive hours; E888.9 Unspecified fall ; V66.7 Encounter for palliative care ; 852.20 Subdural hemorrhage following injury without mention of open intracranial wound, unspecified state of consciousness

Figure 5: Case study on the effectiveness of incorporating auxiliary knowledge. Correctly predicted labels are marked in green and the incorrect ones are marked in red.

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