

# FINDR: A Fast Influential Data Selector for NL2Code Pretraining

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## Abstract

Pretraining on massive corpora has given rise to large language models (LLMs) with multi-task capabilities. However, real-world applications often require more specialized training, as is the case of NL2Code. We approach this specialization through the lens of *data selection*, i.e., identifying a subset of a large corpus that aligns with a desired target distribution—a challenge that remains under-explored within NL2Code. Existing methods are typically designed for selecting instruction-tuning data, and might not easily scale to large-scale code repositories; while methods for NL2Code do exist, they primarily rely on coarse heuristics—such as repo stars—for filtering. To bridge this gap, we propose FINDR, an efficient data selection method that extends logistic regression with feature-wise importance reweighting—marking it, to our knowledge, the first fine-grained solution to NL2Code pretraining. Our method uses hashed n-grams and *code-aware features* to capture code-specific patterns, and then apply informative priors to *reweight* feature importance when computing *influence scores*. Extensive experiments on NL2Python and NL2SQL, with two model families, show that FINDR consistently outperforms strong baselines in both execution accuracy and token efficiency. Notably, pre-training on only 2% of FINDR-selected data boosts Gemma by over 29% in both domains, even surpassing CodeGemma (pretrained on 300x more examples) by 10% in Python.<sup>1</sup>

## 1 Introduction

Large language models (LLMs), such as GPT-4 (OpenAI, 2023), LLaMA3 (AI@Meta, 2024), Gemma (Mesnard et al., 2024), and Mistral (Jiang et al., 2023) have demonstrated remarkable capabilities across a wide range of natural language (NL)

<sup>1</sup>Due to budget constraints, we study the pretraining of *small LLMs* ( $< 3B$ ) in this work (see *Limitation* section).

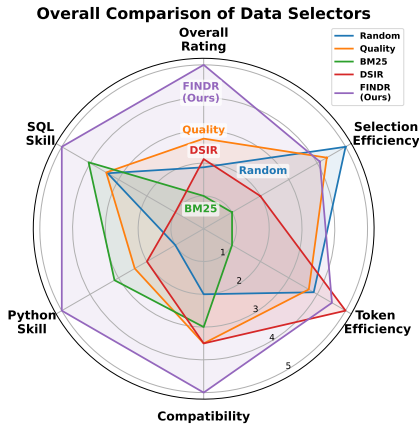


Figure 1: Overall comparison of our proposed FINDR with strong baseline data selectors. FINDR yields the best rating by balancing effectiveness and efficiency. See §A for detailed rubric behind this chart.

tasks, often surpassing expert-engineered NLP systems. Their success can be largely attributed to training on massive corpora (Gao et al., 2021), which capture diverse linguistic patterns and world knowledge (Chowdhery et al., 2023). However, as these models are adapted to more specialized downstream tasks, a significant fraction of pretraining data can become extraneous and counterproductive to such target tasks. In NL2Code (translating natural language into code; Dehaerne et al., 2022; Xu and Zhu, 2022), the choice of relevant and quality training data is even more crucial, since irrelevant data could introduce noisy, misleading examples that hinder the model’s code generation quality (Wang et al., 2023a; Zan et al., 2023). Thus, selecting a subset of code-specific data is essential for adapting a general-purpose LLM to NL2Code tasks. However, manually curated code datasets are time-consuming, error-prone, and limit both scale and diversity (Yu et al., 2018; Hendrycks et al., 2021). Therefore, there presents a pressing need for *automated data selection framework for NL2Code* to mitigate these drawbacks and dynamically tailor to users’ needs.

A growing body of work highlights the importance of *data selection* for efficient pretraining or fine-tuning of LLMs (Feng et al., 2022; Chowdhery et al., 2023; Albalak et al., 2024). Two complementary perspectives are typically involved in this space: *data composition* (Soldaini et al., 2024)—deciding the ratio and types of data to include (e.g., NL vs. code)—and *coreset selection* (Phillips, 2016), where the goal is to identify the most important subset of the training data. In this work, we focus on *coreset selection*, where we seek a small subset of pretraining data that leads to performance on par with or better than full-dataset training. Recent advances, such as targeted instruction tuning (Xia et al., 2024), further highlight the impact of selecting the “right” data to cut computational costs and improve target skills.

Despite growing interest in data selection (Xie et al., 2023; Han et al., 2023; Xia et al., 2024), most existing methods remain computationally expensive because they were originally designed for instruction-tuning data (Wang et al., 2024), which might not scale easily to large pretraining corpora. While data selection techniques do exist for NL2Code (e.g., filtering by doc length or repo stars; Cao et al., 2023; Zan et al., 2023), they primarily rely on overly coarse heuristics that overlook subtle differences in examples. Consequently, *scalable and fine-grained data selection in NL2Code are left relatively under-explored*. To address these limitations, we propose an **efficient** data selector, FINDR (Fast INfluential Data Ranker), an extension of logistic regression with *feature-wise importance reweighting*. In addition, to capture code-specific constructs (e.g., vectorized function calls; Nasrabadi et al., 2023), we augment FINDR’s hashed n-gram feature extractor with *code-aware feature*.<sup>2</sup> With these fast yet accurate designs, FINDR manages to strike a balance between data selection *effectiveness* and *efficiency*, making it particularly well-suited for large-scale settings.

We perform extensive experiments to evaluate FINDR on NL2Python (Lai et al., 2022) and NL2SQL (Li et al., 2024b) using two distinct small LLM (*sLLM*) families, demonstrating both its *effectiveness* and *efficiency*. Our results reveal that FINDR selects *on-target* data that consistently boosts base models across domains and generally outperforms strong baselines by non-trivial mar-

<sup>2</sup>Although we introduce *code feature* tailored to NL2Code, FINDR can be seamlessly adapted to large-scale unlabeled data by adjusting the custom feature space for other domains.

gins. For instance, on Python, FINDR improves the base model by 16% to 36%, and surpasses the SOTA selector, DSIR (Xie et al., 2023), by 10%. While BM25 remains a strong baseline (Xia et al., 2024), FINDR is substantially more efficient, processing 47 million Python files in 3.5 GPU hours compared to BM25’s 760 CPU hours. Notably, training Gemma on 2% of data selected by FINDR outperforms CodeGemma by 10%, which consumes 300 times more examples. Moreover, the proposed FINDR exhibits robust generalization as verified in two model families including both general-domain and code-specific sLLMs.

We summarize major contributions as follows:

- We are the *first* to systematically study a suite of data selection methods for NL2Code continued pretraining, and perform comprehensive comparison among them (Figure 1).
- We propose FINDR, an *efficient* data selector to capture nuanced data influence at scale, which integrates code-aware features into hashed n-gram representations, and augments logistic regression with informative priors for feature-wise importance reweighting.
- Experiments on downstream NL2Python and NL2SQL tasks showcase that FINDR boosts base models substantially while efficiently identifying *on-target* data from large-scale pretraining corpus.
- We validate that FINDR robustly outperforms four baselines, including the SOTA selector (Xie et al., 2023), across two sLLMs (DeepSeek-Coder and Gemma) and two languages (Python and SQL) by large margins.

## 2 Related Work

### 2.1 Data Selection

Data selection has recently emerged as a fundamental research topic for LLMs (Coleman et al., 2020; Xia et al., 2020; Paul et al., 2021; Sachdeva et al., 2024). Two primary directions are *data composition* (Soldaini et al., 2024), which optimizes the mix of different data sources (e.g., natural language vs. code), and *coreset selection* (Phillips, 2016), which identifies a small, representative subset (the “coreset”) that captures the dataset’s essential features. While data composition can improve transparency and help inform decision-making process (Gebru et al., 2021; Elazar et al., 2024), ever higher compute spending has attracted increasing attention to coreset selection (Killamsetty et al., 2021; Xia

et al., 2023; Griffin et al., 2024). By focusing on the most *influential* data, coresnet selection significantly cuts training costs without degrading performance, benefiting both pretraining (Xie et al., 2023; Han et al., 2023) and instruction tuning (Xia et al., 2024; Wang et al., 2024). In this work, we frame data selection as a **coreset selection** problem, developing **efficient** methods for identifying training subsets that match, or even exceed, the performance of training on manual selections (Feng et al., 2022) or even the full corpus.

Existing data selection approaches mainly fall into three broad categories. The first is *random sampling*, which, despite its simplicity and unbiased nature, results in uninformative selections that fail to represent the target domain (Devlin et al., 2019; Gururangan et al., 2020; Guo et al., 2022). The second relies on *surface-level matching* to make informed but efficient selections, including BM25 (Robertson and Zaragoza, 2009), DSIR (Xie et al., 2023), among others (Jiang and Zhai, 2007; Moore and Lewis, 2010; Du et al., 2022). DSIR, for instance, uses hashed n-gram features to represent documents and applies a Naïve Bayes model for data selection. Yet, they struggle with document-length variations: BM25 favors lengthy documents while DSIR selects overly short ones. Quality Classifier (Brown et al., 2020) is a more robust method, which leverages logistic regression and has become a standard for pretraining data selection (Gao et al., 2021; Chowdhery et al., 2023). However, this approach overlooks per-feature importance. FINDR is a *surface-level matching* method which, inspired by importance weighting for domain adaptation (Shimodaira, 2000; Sugiyama et al., 2007), introduces feature-wise importance reweighting to capture nuances among features.

The third category relies on fine-grained feature representations—such as embeddings (Chen et al., 2023; Wu et al., 2023; Xiao et al., 2024), gradients (Han et al., 2023; Xia et al., 2024), perplexities (Li et al., 2024c) or entropies (Kousar et al., 2025)—often combined with pairwise comparisons. While these methods can capture more subtleties in the data, they typically incur quadratic compute complexity, thus restricted to instruction data selection only (Wang et al., 2024). Recent approaches use ChatGPT (Zheng et al., 2023; Liu et al., 2024) to assess data relevance via prompting, but high API costs limit their scalability, especially when re-runs are needed. In contrast, FINDR offers an informed solution that is *tractable* at the scale

needed for unlabeled pretraining data selection.

## 2.2 Natural Language to Code (NL2Code)

Translating natural language problem description into code (NL2Code) has attracted substantial attention for its potential to enhance developer productivity and democratize software development (Allamanis et al., 2018; Dehaerne et al., 2022; Zan et al., 2023). Early studies approached NL2Code through RNN (Iyer et al., 2016), LSTM (Eriguchi et al., 2016) and CodeBERT models (Feng et al., 2020), often incorporating syntax-aware architectures to capture the structural nature of code (Yin and Neubig, 2017). While these methods mark significant progress over rule-based baselines (Allamanis and Sutton, 2014), they rely on large amounts of labeled language-code pairs, limiting coverage and incurring considerable implementation costs.

Most recent progress stems from LLMs (Chen et al., 2021a; Fried et al., 2023; Guo et al., 2024) (continuously) pretrained on massive *unlabeled code* from GitHub and StackOverflow. These models exhibit strong *zero-* and *few-shot* learning capabilities, often requiring minimal tuning or just prompt engineering to excel at coding tasks (Barke et al., 2023; Zheng et al., 2024a; Zhang et al., 2024b). As model sizes grow, LLMs demonstrate emergent capabilities such as debugging (Kang et al., 2025). Despite showing promise as coding assistants, LLMs can still introduce bugs (Nguyen and Nadi, 2022), indicating a need for further refinement before reaching human-level competence.

While data selection has been increasingly studied for NL generation, it remains *under-explored* for NL2Code. In contrast to NL domains, where selecting instruction data drives sophisticated algorithms, NL2Code datasets are predominantly **unlabeled** (HuggingFace, 2021; Kocetkov et al., 2022). Consequently, current practice for NL2Code is limited to basic filtering techniques to ensure code files are deduplicated, complete, and clean (Chen et al., 2021a; Li et al., 2022; Fried et al., 2023; Nijkamp et al., 2023): remove incomplete or auto-generated files and discard rarely used repos. While these heuristics offer decent opportunities for *filtering out* undesired code, they are not meant for *filtering in* (“finding”) the relevant code for a target domain.

Move beyond coarse heuristics, we introduce FINDR, a more fine-grained data selection algorithm *tailored* to NL2Code, while extensible to other *large-scale unlabeled scenarios*. To our knowledge, this is the first systematic study of data

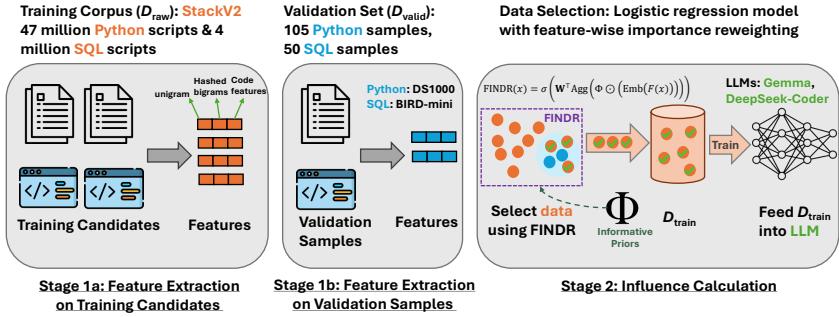


Figure 2: Overview of FINDR. We first extract code-feature-augmented representations (§3.1), and then leverage informative priors to apply feature importance reweighting to compute influence scores (§3.2). The light blue bubble (stage 2) denotes decision boundary.

selection for NL2Code, enabling more efficient and targeted pretraining of sLLMs for code generation.

### 3 Method: FINDR

Figure 2 provides an overview of the two stages behind FINDR, **feature extraction** and **influence calculation**, specifically instantiated in the NL2Code setting. Here, we aim to select the most influential subset of data,  $\mathcal{D}_{train}$ , from a large corpus,  $\mathcal{D}_{raw}$ , for continued pretraining of LLMs.

To begin, we obtain **code-tailored features** by running our feature extractor (§3.1) over the entire training corpus ( $\mathcal{D}_{raw}$ ). Next, we utilize a small set of validation examples,  $\mathcal{D}_{val}$ , which mirrors the target test data,  $\mathcal{D}_{test}$ . Indeed, small-scale validation sets have proven effective for model tuning and domain adaptation (Kirstain et al., 2022; Zhou et al., 2023; Zhang et al., 2024c). The stage 1 process is highly flexible: whenever we have a new target dataset ( $\mathcal{D}_{test}$ ) or a shift in domain, we can easily gather a reasonably small validation set to guide the data selection. Hence, FINDR becomes a plug-and-play solution that can seamlessly adapt to evolving requirements.

Stage 2 is where we measure the **influence** of candidate training points with respect to the small validation set ( $\mathcal{D}_{val}$ ), detailed in §3.2. By focusing on the most influential data, we can reduce training time and resource consumption, which is especially critical for LLMs. After computing the influence scores, we feed the selected data into an LLM for **continued pretraining**. Notably, this can be done either *stochastically*—sampling data points based on normalized FINDR scores (with high-scoring samples possibly repeated in  $\mathcal{D}_{train}$ )—or *deterministically* by selecting the top  $k\%$ <sup>3</sup> of  $\mathcal{D}_{raw}$  to

construct  $\mathcal{D}_{train}$ , ensuring stable coverage of top-scoring examples. We adopt the deterministic approach because, as seen in preliminary experiments, it yields more consistent improvements and simplifies hyperparameter tuning, leaving the stochastic approach for future exploration.

#### 3.1 Feature Extraction (Stage 1)

Considering our goal is to extract features for a massive number of data points in an efficient manner, we choose to trade off some representational expressiveness for higher efficiency. Instead of relying on semantically rich embedding methods (e.g., static (Pennington et al., 2014) and generative (Devlin et al., 2019) embeddings), we adopt  $n$ -gram bag-of-words as a practical solution. Inspired by existing work on feature extraction for NL data (Joulin et al., 2017; Xie et al., 2023), we note that using unigrams alone fails to capture subtle surface-level signals, such as word-pair interactions and ordering cues, but enumerating all bigrams is intractable. Hashed bigrams, instead, strike a practical balance by reducing computational overhead while retaining valuable contextual information.

Beyond textual features, we additionally introduce a code-specific representation—**code feature**—that model programming language-specific functions/patterns. For instance, one feature bucket tracks the frequency of NumPy array creation functions (e.g., `numpy.array()`, `numpy.zeros()`). This helps highlight operations that are particularly relevant to coding. Refer to Table A6 for more examples of code-feature buckets. To summarize, our feature extractor encapsulates both lexical and semantic properties of code (i.e., surface-level unigram, hashed bigrams, and abstracted function descriptions), enabling richer representation for subsequent selection steps.

<sup>3</sup>In this paper, we set a fixed  $k = 2$  unless otherwise noted, similarly to Xia et al. (2024).

### 3.2 Influence Calculation (Stage 2)

In order to efficiently compute influence scores from the feature representations introduced in §3.1, we build upon a logistic regression (LR) model. LR has long been favored in large-scale natural language understanding tasks for its simplicity, ease of interpretation, and robust performance (Brown et al., 2020; Chowdhery et al., 2023; Gao et al., 2021). Equation (1) depicts our FINDR score calculation for each data point  $x$ , extending prior LR-based models with **feature-wise importance reweighting** and incorporating a novel  $\Phi[\cdot]$  function that implements informative priors.

$$\text{FINDR}(x) = \sigma \left( \mathbf{W}^\top \text{Agg}(\Phi \odot (\text{Emb}(F(x)))) \right) \quad (1)$$

where the parameters  $\mathbf{W}$  and  $\text{Emb}[\cdot]$  are trainable, while  $\text{Agg}[\cdot]$  denotes the aggregation function (i.e., generating document-level representations), and  $F(x) \in \mathbb{R}^N$  is the extracted feature for an input data  $x$  (described in §3.1).

The **feature-wise importance** score is computed using  $\Phi[\cdot] \in \mathbb{R}^N$ <sup>4</sup>, formulated as in eq. (2).

$$\Phi[\mathcal{D}_{\text{val}}, \mathcal{D}_{\text{raw}}] = \min(\text{REG}[\frac{\Phi'_{\mathcal{D}_{\text{val}}}[\mathbf{f}]}{\Phi'_{\mathcal{D}_{\text{raw}}}[\mathbf{f}]}], M) \quad (2)$$

where  $\Phi'_{\mathcal{D}}[\mathbf{f}]$  is the frequency-based raw importance score (eq. (3)),  $\text{REG}$  is a regularization factor balancing priors vs. uniform weights (eq. (4)), and  $M$  caps scores to prevent feature-wise shortcuts.

**Raw Importance Score Calculation.** We begin with computing *raw importance scores*,<sup>5</sup> by counting the relative frequency of each individual feature for two sets of data:  $\mathcal{D}_{\text{val}}$  as the positive set and  $\mathcal{D}_{\text{raw}}^6$  as the negative set. Formally,

$$\Phi'_{\mathcal{D}}[\mathbf{f}] = \frac{\sum_{j=1}^n \mathbf{f}_j}{\sum_{i=1}^n \mathbf{1}^\top \mathbf{f}_i} \quad (3)$$

where  $\mathbf{f}_i$  represents the feature vector for the code file  $i$ , extracted as in §3.1, and  $n$  denotes the size  $|\mathcal{D}|$ . This frequency-based score calculation strikes a balance between simplicity and scalability, making it especially appropriate for large collections of unlabeled code.

**Regularization Component (REG).** The REG component serves as a regularization mechanism controlled by the hyperparameter  $\gamma \in [0, 1]$ . The

<sup>4</sup>For simplicity, we use  $\Phi[\cdot]$  to represent  $\Phi[\mathcal{D}_{\text{val}}, \mathcal{D}_{\text{raw}}]$ .

<sup>5</sup>We use “influence” to denote data point-level FINDR score, while “importance” means feature-wise weights.

<sup>6</sup>The construction of  $\mathcal{D}_{\text{raw}}^*$  is detailed in §E.1.

$\gamma$  modulates the reliance on *priors* versus *uniform feature weighting* as follows:

- $\gamma = 1$ : The model reduces to a standard LR model, assigning equal weights to features.
- $\gamma = 0$ : The model fully leverages priors, allowing nuanced feature distinctions.

$$\text{REG}[\phi] = \gamma(1 - \phi) + \phi \quad (4)$$

where  $\phi$  is instantiated as the difference of raw importance scores between positive and negative sets, as shown in eq. (2).

Furthermore, to address the size imbalance between  $\mathcal{D}_{\text{val}}$  and  $\mathcal{D}_{\text{raw}}^*$ , we introduce a rescaling factor  $C$ , and thus replace  $\phi$  with  $\frac{\phi}{C}$  in Equation (4). In fact, such imbalances can lead to *skewed feature importance scores*, particularly when smaller sets disproportionately influence the learning process (Henning et al., 2023). In contrast, the factor  $C$  ensures comparability between sets of varying sizes. Specifically, we implement two types of rescaling factors: *accumulated feature count difference (AFC)* and *document count difference (DC)*. That is, AFC addresses the size difference based on the total number of feature occurrences, while DC is only concerned about the number of code files.

**Capping Scores.** In our preliminary studies, we find that certain features, e.g., project-specific variable names, can show up in short bursts, yielding tremendously large importance scores. Those *rare yet inconsequential* tokens will, however, disrupt the training process of FINDR. Thereby, as indicated in eq. (2), we cap each feature’s importance score at  $M$  to resolve such anomalies and prevent FINDR from picking up unexpected artifacts.

## 4 Experiments

### 4.1 Datasets and Evaluation Metrics

**Pretraining corpus.** We consider **StackV2** (Lozhkov et al., 2024) as  $\mathcal{D}_{\text{raw}}$  for selecting  $\mathcal{D}_{\text{train}}$  for our target tasks. StackV2 is a large-scale code corpus of more than 3 billion files in 600+ programming languages, primarily sourced from public GitHub repos. Our focus is on two subsets: Python and SQL. Combined, these subsets comprise approximately 50 million scripts with an average length of 3,412 characters. Specifically, there are 46.64M Python scripts (totaling 300GB) and 3.63M SQL scripts (totaling 40GB).

**Evaluation benchmarks.** To assess performance and generalizability, we evaluate baselines and

	DeepSeek-Coder						Gemma					
	Origin	Surface	Semantic	Difficult	Perturbation	Overall	Origin	Surface	Semantic	Difficult	Perturbation	Overall
Base Model	19.9	9.2	17.5	6.8	12.0	15.1	13.8	7.2	9.8	4.9	7.6	10.1
Random Selection	20.7	8.6	16.7	4.9	11.0	14.7	15.3	6.6	10.7	6.2	8.2	10.9
Quality Classifier	21.2	9.3	15.8	6.2	11.2	15.1	17.0	10.5	11.5	6.2	9.7	12.5
BM25	22.2	6.6	14.1	5.6	9.5	14.4	21.0	7.9	14.5	4.9	9.8	14.2
DSIR	22.2	9.9	17.5	5.9	12.0	15.9	15.3	5.3	12.4	5.6	8.4	11.1
FINDR (Ours)	24.2	12.2	18.4	7.1	13.3	17.5	19.0	9.2	14.1	6.2	10.4	13.7

Table 1: Comparison of FINDR with data selection baselines in the Python domain, measured by Pass@1, when training with 2% of selected data. Base model denotes out-of-the-box evaluation without additional training. Per Lai et al. (2022), we conduct 0-shot evaluation, and we report individual results on 4 problem types and the aggregated perturbation set. Best results are **bold**, and informed data selectors that outperform the base model are highlighted on a scale of 5 red shades (color schemes in §B). Overall, FINDR improves base Coder and Gemma by 16% and 36%. Notably, FINDR attains the highest score on perturbed items, showcasing the robustness of FINDR. Efficiency comparison, i.e., data selection efficiency, is provided in Table A4 (Selection Time column).

FINDR on two target tasks/domains ( $\mathcal{D}_{test}$ ).<sup>7</sup> In the Python domain, we focus on the still largely unresolved **DS-1000** (Lai et al., 2022), instead of widely studied benchmarks like HumanEval (Chen et al., 2021b) and MBPP (Austin et al., 2021), which have approached saturated performance (Table A8). DS-1000 comprises 1,000 data science-oriented code generation problems spanning seven scientific computing libraries, e.g., NumPy. For SQL, we adopt the challenging **BIRD** (Li et al., 2024b), which comprises 95 databases across 37 professional areas, and narrows the gap between experimental and real-world settings seen in other benchmarks (Zhong et al., 2017; Yu et al., 2018). In this work, we use its recent derivative, BIRD-miniDev,<sup>8</sup> released in June 2024, which supports diverse database management systems.

Following the literature (Xia et al., 2024), we also hold out a subset of examples as  $\mathcal{D}_{val}$  for guiding data selection. Statistics are shown in Table A5.

**Evaluation metrics.** For Python DS1000 (Lai et al., 2022), we use the **pass@1 accuracy**, which evaluates functional correctness based on test case success and adherence to surface-form constraints (e.g., mandatory use of vectorized operations). For SQL (Li et al., 2024b), we report **Execution (EX)**, which checks if predicted and ground-truth queries produce identical results, and **Soft F1-score**, which measures the similarity between the tables produced by generated and reference SQL queries.

## 4.2 Experiment Setup

We include recent small LLMs that excel at reasoning and code completion tasks: coding-specialized sLLM, **DeepSeek-Coder-1.3B-base** (hereafter,

<sup>7</sup>We rigorously compared nearly 20 benchmarks to determine the most suitable ones for our evaluation (Table A7).

<sup>8</sup>[https://github.com/bird-bench/mini\\_dev](https://github.com/bird-bench/mini_dev)

Coder; Guo et al., 2024), and generalist sLLM, **Gemma-2B** (Mesnard et al., 2024).

As our goal is to study if a selected subset of influential data can boost sLLM performance on the target task, we only perform **continued pre-training** on  $\mathcal{D}_{train}$  without additional fine-tuning. We use base versions of the models and employ Llama-Factory (Zheng et al., 2024b) for parallel training with 8 40GB GPUs. We set the context length to 4,096, gradient accumulation to 32 and per-device batch size to 1. Models are trained for 2 epochs for Python and 3 epochs for SQL. For all other hyperparameters, we keep the default values.

For evaluation, we follow the official evaluation protocols (Lai et al., 2022; Li et al., 2024b), and use greedy decoding with few-shot demonstrations if needed (0-shot for Python and 1-shot for SQL).

**FINDR Setup and Training:** We build  $\mathcal{D}_{FINDR}$  for training FINDR, using  $\mathcal{D}_{val}$  and a sample of  $\mathcal{D}_{raw}$  as positive and negative sets. In the feature extractor, we use all unigrams in  $\mathcal{D}_{FINDR}$  and, for bigrams, apply the FNV-1a algorithm (Fowler et al., 2012) to obtain hashed features using 100k buckets. The code feature is enabled only for Python, and we semi-automatically define 618 classes (buckets), covering 8,721 Python functions.<sup>9</sup> The training process of FINDR consists of two stages: first, learning  $\Phi[\cdot]$  (§E.2), and then supervised learning on  $\mathcal{D}_{FINDR}$ ,<sup>10</sup> which updates the randomly initialized parameters  $\mathbf{W}$  and  $\text{Emb}[\cdot]$  for 10 epochs.

## 4.3 Data Selector Baselines

We include major *efficient* pretraining data selection baselines. The simplest baseline is **random selection**, where we randomly sample data from the training corpus. For informed data selection

<sup>9</sup>See §D for semi-automatic construction process.

<sup>10</sup> $|\mathcal{D}_{FINDR}^{\text{Python}}| = 1,000$  and  $|\mathcal{D}_{FINDR}^{\text{SQL}}| = 500$ .

	DeeoSeek-Coder				Gemma			
	Easy	Med.	Hard	Overall	Easy	Med.	Hard	Overall
Base	<b>26.5</b>	8.8	<b>2.0</b>	11.1	12.2	2.8	<b>3.9</b>	5.1
Random	18.4	5.7	<b>2.0</b>	7.6	11.2	2.8	2.9	4.7
Quality	19.4	3.8	1.0	6.6	<b>18.9</b>	2.0	2.9	<b>5.9</b>
BM25	22.5	6.4	<b>2.0</b>	8.9	<b>17.9</b>	<b>3.2</b>	2.0	6.1
DSIR	4.1	0.0	0.0	0.9	6.6	0.6	0.0	1.8
FINDR	25.5	<b>11.2</b>	<b>2.0</b>	<b>12.2</b>	<b>18.9</b>	2.8	<b>3.9</b>	<b>6.6</b>

Table 2: EM performance in the SQL domain when training with 2% of selected data (F1 in Table A2). Base model denotes out-of-the-box evaluation. Following Li et al. (2024b), we conduct 1-shot evaluation, and we report individual results on 3 problem types. Best results are **bold**, and data selectors superior to Base are highlighted on a scale of 5 red shades. In general, FINDR leads to the best performance across the board.

baselines, we compare with **BM25** (Robertson and Zaragoza, 2009), which is based on word frequency statistics to rank examples to determine how relevant a training document is. Another baseline, LR-based **Quality Classifier**, is widely used for filtering and selecting data from large-scale pretraining corpora (Gao et al., 2021; Chowdhery et al., 2023). We also compare to prior art for selecting unlabelled data, **DSIR** (Xie et al., 2023). It applies n-gram features to weight candidate training data through Naive Bayes formulation, and sample data points accordingly.

We do not compare to instruction data selection approaches, due to extremely *slow* pace and *high* compute cost (Wang et al., 2024), such as representation-(Xiao et al., 2023) and gradient-based (Xia et al., 2024) methods, often exceeding 10k GPU hours to process StackV2 Python subset.

## 5 Results

### 5.1 Main Results and Analyses

We present our main results in Table 1 and Table 2 for Python and SQL domains, respectively, compared against baseline approaches. More results are in §C. We summarize five key findings.

1) *Random selection often degrades performance.* It can be tempting to assume that adding more data—no matter how it is sampled—will improve model performance. But in line with pre-LLM findings (Moore and Lewis, 2010; van der Wees et al., 2017), our experiments confirm that randomly chosen data can degrade performance, even compared with using the base model out-of-the-box. Thus, in the absence of an informed data selector, defaulting to the base model is preferred.

2) *FINDR selects on-target data that consistently boosts base models across domains.* As

has been shown, training on FINDR-selected data consistently enhances code generation capabilities across all evaluated sLLMs. In particular, overall performance gains range from 16% to 36% on Python and 9% to 29% on SQL. These results suggest that FINDR can indeed select the most influential examples, which works robustly across experimental settings.

3) *FINDR generally outperforms strong baselines by non-trivial margins.* Apart from Gemma model on Python, FINDR performs the best across the board. Interestingly, BM25 sometimes surpasses FINDR, but at a *significant computational cost*: it has the slowest selection pace (Figure 1), and the Python scripts it selects are five times longer (Table A4). As a result, token counts rise substantially, thereby increasing LLM training time. In contrast, FINDR offers a more *balanced* solution for *token-efficiently* capturing the most influential data. Notably, on “Perturbed” Python examples—which mitigate potential data leakage from LLM pretraining—FINDR achieves the highest scores, underscoring its robust performance in settings that rely less on memorized knowledge. For further evidence, see Table A1 and Table A2.

4) *FINDR demonstrates superior robustness on “Difficult” examples.* Beyond the overall performance leap, a key advantage of FINDR is its robustness in handling “difficult” scenarios. On Python’s “Difficult” split, FINDR consistently yields the largest improvement relative to other baselines, indicating that its selected examples effectively target the reasoning skills needed for complex code generation. Likewise in SQL, FINDR preserves, and sometimes improves (Table A2), the ability to generate challenging SQL queries, whereas several baselines worsen performance in these tough cases.

5) *NL-targeted selectors do not necessarily excel at NL2Code.* Finally, we note that DSIR (Xie et al., 2023), SOTA method for selecting pure NL data, proves much less effective when adapting to code, especially on the SQL domain. Indeed, as discussed in Xia et al. (2024), we have also observed that DSIR-selected examples are extremely short, thus weakening the code generation capacity of trained models. Furthermore, our finding highlights a notable gap between NL-only and NL2Code data, while our method, FINDR, offers—to the best of our knowledge—the first solution to help bridge the gap.

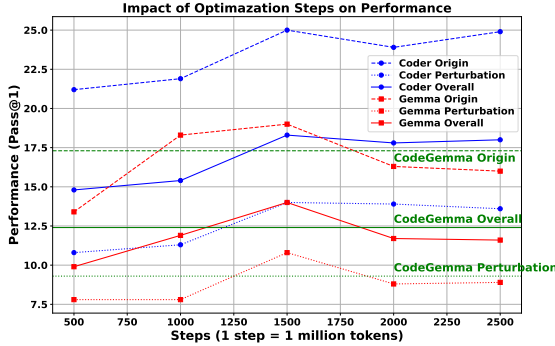


Figure 3: Pass@1 scores of continuously trained sLLMs in the Python domain, in relation to the optimization steps. We identify the performance peaking at around 1,500 steps, trained with 1.5B *on-target* tokens, and Gemma even outperforming CodeGemma, despite the latter being trained on 500B *general* tokens.

## 5.2 Further Study on Data Selection

Data, the essential component in this study, has been the driving force behind ever capable LLMs. Here, we are particularly interested in how LLMs respond to varying *training data*. We address this question under two conditions: (1) varying the **optimization steps** (which translates to the number of training tokens), and (2) varying the **selection ratio** (i.e., the proportion of data selected by FINDR).

**Impact of Optimization Steps.** We first analyze how increasing the training budget—in terms of optimization steps—impacts final performance. Figure 3 shows performance trends for Coder and Gemma up to 2,500 steps.<sup>11</sup> Both models improve until about 1,500 steps (1.5B tokens), after which Coder plateaus and Gemma slightly declines. Thus, 1,500 steps provides a clear balance of performance gains and training efficiency under the default 2% selection ratio. We also compare Gemma to CodeGemma (Zhao et al., 2024), which benefits from extra 500B tokens of continued pre-training. Despite CodeGemma being a stronger LLM, Gemma continuously trained on 1.5B *on-target* tokens chosen by FINDR effectively closes the gap. This highlights the advantage of informed data selection like FINDR—it consumes just **0.3%** of the CodeGemma training tokens yet outperforms massive-scale training at random.

**Impact of Selection Ratio.** We next examine how different selection ratios (1%, 2%, 5%, 10%) affect performance, as depicted in Figure 4 and Fig-

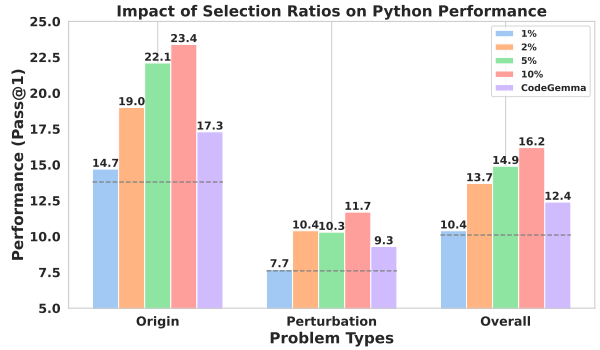


Figure 4: Gemma results in Python domain with varying selection ratios. Dashed line denotes the off-the-shelf Gemma’s result. Performance improves steadily with higher selection ratios, with a notable cut-off at 2%, where additional training yields benefits. Complete results refer to Figure A1. We also observe similar trend for the SQL domain, shown in Figure A2.

ure A2. Dashed lines denote the out-of-the-box Gemma’s performance. On both tasks, continuously trained models exceed the base model once the ratio reaches 2%. Beyond that threshold, performance generally rises further, albeit with marginal gains from 2% to 5% in certain splits (e.g., Python Perturbation, SQL Hard). Moreover, for Python, starting at 2%, Gemma trained on FINDR-selected data surpasses CodeGemma despite its extensive pretraining. However, no such leap is observed for SQL, even at 10%. This is likely due to the fact that Python’s full set (47M scripts) far exceeds SQL’s (4M), so 10% of SQL data still translates to fewer samples than 1% of Python.

In summary, our findings reveal that both the size of the training budget (i.e., the number of steps) and FINDR’s selection ratio play significant roles in reshaping downstream NL2Code capabilities. More importantly, *informed data selection* can substantially improve performance with only a small fraction of the entire corpus. In the future, we will explore optimal training steps for each selection ratio and further investigate the scaling law for informed data selection.

## 5.3 Ablation Study of FINDR

We conduct ablation experiments to analyze contributions of design elements in FINDR.

## 5.4 $\eta$ Ratio for $\Phi$ Estimation

We vary  $\eta$ <sup>12</sup> from 1 to 100 to assess how the sampling size of negatives for *informative priors* es-

<sup>11</sup>Each optimization step processes 1M tokens, so 500–2,500 steps correspond to training on 0.5B–2.5B tokens.

<sup>12</sup>a hyperparameter introduced in the learning process of  $\Phi[\cdot]$  (§E.2).

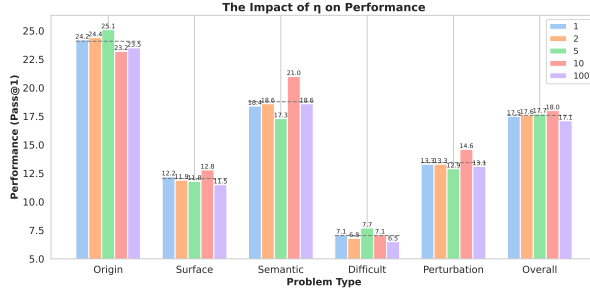


Figure 5: Impact of  $\eta$  (sampling size of negatives for informative priors estimation) on performance in Python domain. We have witnessed a steady performance boost from  $\eta = 1$  to 10, but a sharp drop afterwards, likely due to diminishing returns from overly large sampling of negatives.

timation influences downstream performance. Intuitively, a larger set should yield more accurate feature-weight estimates, thus improving downstream selection. Figure 5 confirms this trend for most problem types, recording a steady performance boost from  $\eta = 1$  to 10. However, a noticeable drop is observed beyond that point, as evidenced by the lowest overall result when  $\eta = 100$ , likely due to diminishing returns from overly large sampling of negatives.

Balancing efficiency and accuracy, we adopt  $\eta = 1$  in our main experiments. This setting requires minimal computational overhead yet maintains near-peak performance. Future work can explore adaptive strategies for tuning  $\eta$  to improve the *informative priors* estimation.

**Code Features and Rescaling Approach.** We ablate the *code feature* in feature extractor (§3.1), and test DC-based versus AFC-based *rescaling* approaches (§3.2). As displayed in Table 3, removing code features consistently degrades results across all splits, showing the importance of code-specific features in code-related tasks (Nasrabadi et al., 2023; Jiang et al., 2024). While DC rescaling sometimes performs competitively, AFC generally yields superior results. This reflects the benefit of fine-grained rescaling for data imbalance issues (Henning et al., 2023). Altogether, the ablation studies validate the design choices in FINDR.

## 6 Conclusion

In this work, we introduce FINDR, an efficient pretraining data selection method based on logistic regression but enhanced with feature importance reweighting. Concretely, we augment hashed n-

	Gemma					
	Origin	Surface	Semantic	Difficult	Perturbation	Overall
FINDR	<b>19.0</b>	<b>9.2</b>	<b>14.1</b>	<b>6.2</b>	<b>10.4</b>	<b>13.7</b>
- Code feature	17.4	8.6	11.1	4.6	8.5	12.0
FINDR (DC rescaling)	18.3	6.9	11.5	<b>6.2</b>	8.7	12.4
- Code feature	<b>19.0</b>	<b>9.2</b>	12.8	3.7	9.1	12.9

Table 3: Ablation study of FINDR. We find that removing code features consistently degrades results across all splits. Complete results (incl. DS-Coder) see Table A3.

gram features with *code features* to capture code-specific constructs, then apply informative priors to *reweight* feature importance when computing *influence* scores. Notably, our FINDR is the first data selection algorithm tailored to NL2Code pretraining. Experiments on Python and SQL demonstrate FINDR’s superiority over strong baselines and its compatibility across diverse sLLMs. Our further study confirms that a small, influential subset of data can yield significant performance improvements, even outperforming an LLM trained on 300 times more examples.

## Limitation

**GPU resources.** The base sLLMs used for continued-pretraining in this work are of 1.3 to 2.5 billions parameters. It is thus more time-consuming than training smaller previous-generation models like BART (Lewis et al., 2020), which in turn results in a significantly higher carbon footprint. Specifically, we train each model on 8 NVIDIA A100 (40GB VRAM) with significant CPU and memory resources. The training time for each model ranges from several hours to 2 days, depending on the configurations.

Meanwhile, due to the limited GPU resources at hand (8 NVIDIA A100-40GB), this work serves as a *pilot study* to rigorously assess the effectiveness of various data selection methods for continued pretraining of small LLMs on the NL2Code task. We will study the scaling effect (i.e., increasing model sizes) in future work, as additional compute becomes available.

**Evaluation Domains.** In this work, we have included two challenging evaluation benchmarks, aiming to cover a diverse array of code styles and domains. Yet, these two benchmarks cannot comprehensively represent the entire spectrum of the NL2Code space. Indeed, evaluation remains an ongoing challenge in data selection—existing studies typically rely on only 3–4 benchmarks as well (Xia et al., 2024; Li et al., 2024a). In future research, we plan to extend FINDR to more program-

ming languages, e.g., Java and C++, and examine its robustness as new large-scale benchmarks are constructed.

**Generalizability of FINDR.** In this work, we focus primarily on developing and validating FINDR for NL2Code. As is common in this area (Wang et al., 2024), evaluations are typically performed on the motivating target domains only, leaving the question of generalizability to a broader range of domains for future work. For instance, DSIR (Xie et al., 2023), a SOTA data selector in the natural language (NL) space, performs poorly in the coding space (Table 1, Table 2). Therefore, we plan subsequent work focusing on extending the evaluation of FINDR to non-NL and non-NL2Code domains, while expecting others to also investigate FINDR beyond NL2Code as we have done with the strong baselines (e.g., DSIR) in this work.

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## A Rubrics for Radar Plot

In order to produce the overall comparison radar plot (Figure 1), we consider five distinct metrics.

- **Selection Efficiency:** This metric is derived from the *Selection Time* column in Table A4. Specifically, we first apply add-one smoothing to all raw values, followed by a logarithmic transformation, and is finally rescaled to a range of 1 to 5 using min-max normalization. Note, for all raw values (unit: hours), they represent the entire time cost of selecting 2% of data from the Python set of StackV2, including parameter learning and inference (i.e., data selection) stages. For the random baseline, the time cost is trivial, so we assign 0.
- **Token Efficiency:** This metric is directly derived from the *Character Count* column in Table A4, and is rescaled to a range of 1 to 5 using min-max normalization. The raw value indicates the average number of characters in a Python script within the selected data.
- **Compatibility:** The purpose of this metric is to assess whether a data selection approach can boost performance for both evaluated models, and whether it exhibits any model preference or bias. This metric is relatively subjective, and the scores we assign to each approach are displayed in Table A4. For example, both Quality Classifier and BM25 yield improvements when the underlying model is Gemma, but no improvement is observed for Coder. Therefore, we rate their compatibility as moderate (3). However, since Quality classifier does not hurt the performance in the Python domain, we slightly increase its score to 3.5.
- **Python Skill:** This metric is directly derived from Table 1, and is rescaled to a range of 1 to 5 using min-max normalization.
- **SQL Skill:** This metric is directly derived from Table 2, and is rescaled to a range of 1 to 5 using min-max normalization.

Finally, we derive the **overall rating** by placing equal emphasis on both efficiency and effective/performance dimensions. The efficiency dimension encompasses the selection and token efficiencies, while the other three metrics are grouped under the effective/performance dimension. At the authors' discretion, we assign the five metrics weights of [1, 1, 0.5, 0.75, 0.75], considering that

compatibility is a relatively subjective metric. We then compute a weighted sum of per-metric ranks for each data selection approach, which is subsequently rescaled to a range of 1 to 5 using min-max normalization.

## B Color Scheme

For all data selectors (including both FINDR and baseline approaches), we highlight them on a scale of 5 red shades based on the relative improvements over the off-the-shelf base models. We design the following scheme to color Table 1, Table 2, Table A1 and Table A2:

- if the relative gain is in the range of (0%, 5%], the value is highlighted in pale pink .
- if the relative gain is in the range of (5%, 15%], the value is highlighted in pink .
- if the relative gain is in the range of (15%, 30%], the value is highlighted in rose-pink .
- if the relative gain is in the range of (30%, 50%], the value is highlighted in rose-red .
- if the relative gain is over 50%, the value is highlighted in dark red .

## C Supplementary Main Results

Due to space limitation in the main text, this section supplements §5.1.

**Python.** Table A1 presents results where features for validation data are extracted from the *complete* script, including both *context* and *answer*, whereas for Table 1, features are extracted solely from the *context*. Overall, the performance difference between the *context-only* and *complete* script settings is minimal. Therefore, for all the other experiments performed in the Python domain, we extract features using only the context for two reasons: (1) Context is shorter than the complete script, making feature extraction more efficient, and (2) incorporating solutions requires extensive human annotation, which limits scalability.

**SQL.** As discussed in §4.1, we adopt two evaluation metrics for the SQL domain: EM and F1. Table A2 complements Table 2 by presenting performance in terms of F1. Note, for all SQL domain experiments, features for validation data are only extracted from the *answer*. This choice is based on our observation that using *context-only* or *complete*

	DeepSeek-Coder						Gemma					
	Origin	Surface	Semantic	Difficult	Perturbation	Overall	Origin	Surface	Semantic	Difficult	Perturbation	Overall
Base Model	19.9	9.2	17.5	6.8	12.0	15.1	13.8	7.2	9.8	4.9	7.6	10.1
Random Selection	20.7	8.6	16.7	4.9	11.0	14.7	15.3	6.6	10.7	6.2	8.2	10.9
Quality Classifier	<b>23.6</b>	<b>9.9</b>	17.9	6.8	12.4	16.8	<b>19.9</b>	<b>7.9</b>	12.4	<b>7.4</b>	<b>9.7</b>	<b>13.6</b>
BM25	22.8	6.6	15.4	5.6	10.1	15.0	<b>23.1</b>	5.3	15.0	6.2	9.7	<b>14.9</b>
DSIR	21.0	9.2	<b>18.8</b>	6.8	12.6	15.9	16.4	7.9	12.0	4.9	8.8	11.7
FINDR (Ours)	<b>23.3</b>	<b>12.5</b>	<b>18.8</b>	<b>7.4</b>	<b>13.7</b>	<b>17.4</b>	20.1	<b>9.2</b>	<b>16.2</b>	3.7	<b>10.6</b>	14.3

Table A1: Comparison of FINDR with strong data selection baseline approaches in the Python domain, measured by Pass@1, when training with 2% of selected data. Base model denotes out-of-the-box evaluation without additional training. In contrast to Table 1, features of validation examples are extracted from the complete script (i.e., context and answer). Following Lai et al. (2022), we conduct 0-shot evaluation, and we report individual results on 4 problem types and the aggregated perturbation set. Best results are **bold**, and informed data selectors that outperform the base model are highlighted on a scale of 5 red shades (see color schemes in §B). Overall, FINDR improves over base Coder and Gemma by 15% and 39%, respectively. Notably, FINDR achieves the highest score on perturbed items, showcasing the robustness of FINDR.

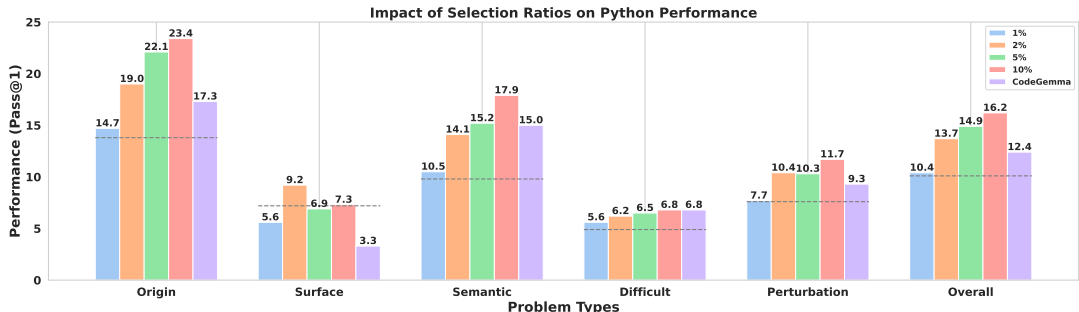


Figure A1: Gemma results in Python domain with varying selection ratios. Dashed line denotes the off-the-shelf Gemma’s result. Performance improves steadily with higher selection ratios, with a notable cut-off at 2%, where additional training leads to better performance than base Gemma and CodeGemma.

	DeepSeek-Coder				Gemma			
	Simple	Moderate	Challenging	Overall	Simple	Moderate	Challenging	Overall
Base	<b>28.4</b>	10.7	3.3	12.8	13.3	3.6	4.6	6.0
Random	19.1	6.5	<b>4.1</b>	8.7	12.8	3.4	4.2	5.6
Quality	19.8	4.4	1.5	7.1	<b>18.5</b>	2.7	3.3	6.3
BM25	24.0	7.9	3.3	10.3	<b>19.6</b>	<b>4.3</b>	4.1	<b>7.6</b>
DSIR	4.1	0.1	0.0	1.0	7.0	0.7	0.8	1.8
FINDR	27.5	<b>11.7</b>	2.7	<b>13.1</b>	<b>20.0</b>	4.1	<b>4.9</b>	<b>7.7</b>

Table A2: F1 performance comparison in the SQL domain when training with 2% of selected data. Base model denotes out-of-the-box evaluation. Following Li et al. (2024b), we conduct 1-shot evaluation, and we report individual results on 3 problem types. Best results are **bold**, and data selectors superior to base are highlighted on a scale of 5 red shades. In general, FINDR leads to the best performance across the board. EM results refer to Table 2.

scripts results in inferior performance and higher results variance.

## D Construction of Code Feature Buckets

To construct a comprehensive set of code feature buckets, we leverage powerful LLMs such as GPT-3.5.<sup>13</sup> Using the Numpy library as an example, we prompt GPT-3.5 to generate major function cate-

gories and their corresponding expressions. This process is performed three times with varying output sizes: 10, 50, and 70 classes. We merge the results, removing only duplicated expressions while retaining all unique classes. This procedure is repeated for each of the seven libraries. In total, we compile 618 feature classes encompassing 8,721 distinct Python expressions. Examples of these feature buckets are presented in Table A6.

## E More details of Informative Priors ( $\Phi$ )

The informative priors ( $\Phi$ ) are intended to capture global beliefs about the relative importance of different features. In an ideal scenario,  $\Phi$  would be independent of any specific experience. However, in practice,  $\Phi$  inevitably depends on the target domain. Moreover, generating an exhaustive list of importance scores for all features in the training set is impractical—especially since the training set is dynamic, and even if it were fixed, manually assigning these scores would be prohibitively labor-intensive. To this end, we decide to estimate  $\Phi$  in an on-demand fashion using Equation (2).

<sup>13</sup><https://chat.openai.com/>; gpt-35-turbo-16k-0613, training data up to Sept. 2021

	DeepSeek-Coder						Gemma					
	Origin	Surface	Semantic	Difficult	Perturbation	Overall	Origin	Surface	Semantic	Difficult	Perturbation	Overall
FINDR	<b>24.2</b>	<b>12.2</b>	18.4	<b>7.1</b>	13.3	<b>17.5</b>	<b>19.0</b>	<b>9.2</b>	<b>14.1</b>	<b>6.2</b>	<b>10.4</b>	<b>13.7</b>
- Code feature	22.5	11.2	18.8	6.8	13.1	16.8	17.4	8.6	11.1	4.6	8.5	12.0
FINDR (DC rescaling)	23.4	10.6	18.6	6.8	12.9	17.0	18.3	6.9	11.5	<b>6.2</b>	8.7	12.4
- Code feature	22.2	11.9	<b>20.5</b>	6.5	<b>14.0</b>	17.2	<b>19.0</b>	<b>9.2</b>	12.8	3.7	9.1	12.9

Table A3: Ablation study of FINDR. We find that removing code features consistently degrades results across all splits. Meanwhile, using DC rescaling approach generally hurts the performance, in comparison with the default AFC.

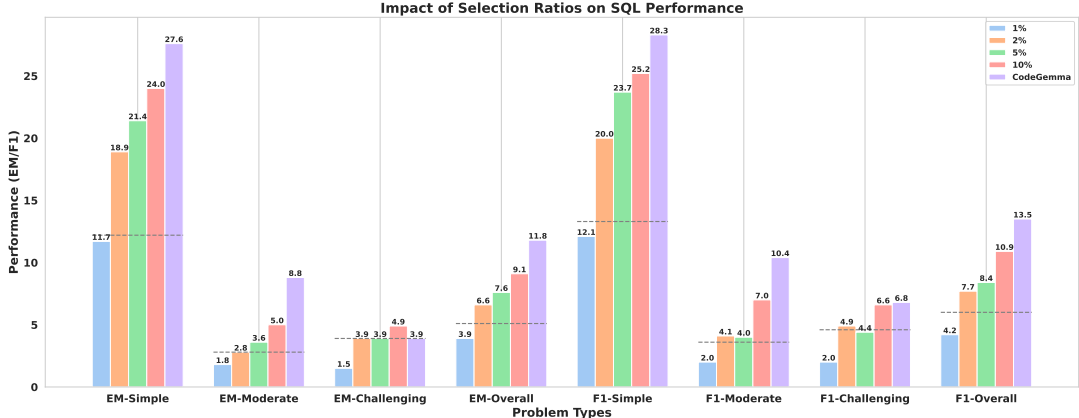


Figure A2: Gemma results (EM and FI) in SQL domain with varying selection ratios. Dashed line denotes the off-the-shelf Gemma’s result. Performance improves steadily with higher selection ratios, with a notable cut-off at 2%, where additional training leads to better performance than base Gemma.

	Selection Time	Character Count	Compatibility
Random	0	3,410	2.0
Quality	2	3,855	3.5
BM25	760	10,788	3.0
DSIR	145	533	3.5
FINDR (Ours)	3.5	1,762	5.0

Table A4: Raw statistics for radar plot (Figure 1), and efficiency comparison (i.e., selection time) among select baselines and FINDR. The unit for selection time is *hours*, including parameter learning and inference stages. The detailed usage of each column is documented in §A.

### E.1 Construction of $\mathcal{D}'_{raw}$

To address the large size disparity between  $\mathcal{D}_{val}$  and  $\mathcal{D}_{raw}$ , and mitigate shortcut learning (see §3.2), we introduce a reduced negative set,  $\mathcal{D}'_{raw}$ . Specifically,  $\mathcal{D}'_{raw}$  is a small subset of  $\mathcal{D}_{raw}$  that serves as a proxy for negative examples. To stabilize subsequent supervised learning stage of FINDR, we ensure that all negative instances in  $\mathcal{D}_{FINDR}$  are contained within  $\mathcal{D}'_{raw}$ . We further introduce a hyperparameter  $\eta$ , which defines the size of  $\mathcal{D}'_{raw}$  as a multiple of the negative set in  $\mathcal{D}_{FINDR}$ . That is,  $|\mathcal{D}'_{raw}| = \eta |\mathcal{D}_{FINDR}^-|$ . By tuning  $\eta$ , we can balance data coverage against computational overhead.

### E.2 Learning Process of $\Phi$

After obtaining  $\mathcal{D}'_{raw}$ , we combine it with  $\mathcal{D}'_{val}$  (serving as the positive set) to learn  $\Phi$ . The learning process of  $\Phi$  is detailed in §3.2. Once  $\Phi$  is learned, the corresponding parameters are frozen in the subsequent supervised learning stage.

### E.3 Default Setting of $\Phi$

Our FINDR method introduces three core hyperparameters in  $\Phi$ :

- $\gamma$ : Balances the contribution of the priors vs. uniform weighting ( $\gamma = 0.75$  by default).
- $M$ : Caps the maximum importance score for each feature ( $M = 3$  by default).
- $\eta$ : Controls the ratio of negative samples between the training set and the Prior estimation set ( $\eta = 1$  by default, due to efficiency and representativeness).

Benchmarks	$ \mathcal{D}_{\text{val}} $	$ \mathcal{D}_{\text{test}} $		Splits	$ C $	$ A $	#Shot	#Tasks	Domain
DS1000 (Lai et al., 2022)	105	895	452 (105)/152/234/162	2,857	141	0	7	7	Python
BIRD-miniDev (Li et al., 2024b)	50	450	148 (50)/250/102	4,270	201	1	12	12	SQL

Table A5: Statistics of evaluation benchmarks.  $|\mathcal{D}_{\text{val}}|$  and  $|\mathcal{D}_{\text{test}}|$  denote the size of validation and test sets. Splits represent the fine-grained data splits by problem types, as seen in Table A1 and Table 2. That is, there are *Origin*, *Surface*, *Semantic* and *Difficult* in the Python domain, and *Simple*, *Moderate* and *Challenging* in the SQL domain. We also ensure that all validation data are sampled from the simplest category, as indicated by parentheses, allowing for the study of LLM generalizability and true intelligence.  $|P|$  and  $|A|$  denote the average length of *context* (C) and *answer* (A). For #Shot, we follow the official practice in respective benchmarks (Lai et al., 2022; Li et al., 2024b). #Tasks represent the number of libraries (Python) and subjects (SQL) included in each benchmark.

Library	Feature Bracket	Expressions/Functions
Matplot	Plotting Functions	<code>pyplot.plot, plot, matplotlib.pyplot.hist, plt.hist, boxplot, plt.scatter, bar, matplotlib.pyplot.boxplot, matplotlib.pyplot.plot, scatter, matplotlib.pyplot.bar, plt.bar, pyplot.bar, plt.plot, pyplot.scatter, pyplot.hist, hist, pyplot.boxplot, plt.boxplot, matplotlib.pyplot.scatter</code>
Numpy	Array Creation	<code>np.ones, numpy.eye, array, numpy.zeros, numpy.array, np.zeros, np.array, numpy.ones, empty, zeros, np.eye, ones, eye, np.empty, numpy.empty</code>
Pandas	Input/Output	<code>to_csv, pd.read_json, pandas.read_csv, pd.read_sql, pandas.to_csv, pandas.read_html, pandas.read_sql, read_csv, read_html, read_json, pandas.read_json, pd.to_csv, read_sql, pd.read_html, pd.read_csv, pandas.read_excel, pd.read_excel, read_excel</code>
PyTorch	Math Operations	<code>torch.log, torch.cos, add, torch.sub, pow, sub, torch.sqrt, exp, sin, cos, sqrt, mul, div, torch.sin, torch.exp, torch.mul, log, torch.pow, torch.add, torch.div</code>
SciPy	Data Structures	<code>scipy.sparse.dok_matrix, coo_matrix, scipy.sparse.coo_matrix, scipy.sparse.bsr_matrix, scipy.sparse.lil_matrix, sparse.lil_matrix, sparse.dok_matrix, sparse.bsr_matrix, scipy.sparse.csc_matrix, sparse.csc_matrix, sparse.coo_matrix, csc_matrix, sparse.csc_matrix, sparse.bsr_matrix, dok_matrix</code>
Sklearn	Model Selection	<code>sklearn.model_selection.KFold, model_selection.GridSearchCV, StratifiedKFold, sklearn.model_selection.GridSearchCV, cross_val_score, model_selection.train_test_split, KFold, sklearn.model_selection.cross_val_score, train_test_split, model_selection.StratifiedKFold, sklearn.model_selection.train_test_split, GridSearchCV, sklearn.model_selection.StratifiedKFold, model_selection.KFold, model_selection.cross_val_score</code>
TensorFlow	Tensor Manipulation	<code>tf.constant, tf.Variable, concat, tf.concat, Variable, constant, tf.reshape, reshape, transpose, tf.transpose</code>

Table A6: Example feature brackets. For each library in DS1000, we show one bracket with associated expressions.

Benchmark	Size	#PL
HumanEval (Chen et al., 2021b)	164	Python
HumanEval+ (Liu et al., 2023)	164	Python
MBPP (Austin et al., 2021)	974	Python
MBPP+ (Liu et al., 2023)	378	Python
Spider (Yu et al., 2018)	8,034	SQL
BIRD-Dev (Li et al., 2024b)	500	SQL
ODEX (Wang et al., 2023b)	945	Python
CoderEval (Yu et al., 2024)	460	Python, Java
ReCode (Wang et al., 2022)	1,138	Python
StudentEval (Babe et al., 2024)	1,749	Python
BigCodeBench (Zhao et al., 2024)	1,140	Python
ClassEval (Du et al., 2023)	100	Python
NaturalCodeBench (Zhang et al., 2024a)	140	Python, Java
LiveCodeBench (Jain et al., 2025)	713	Python
DSP (Chand et al., 2022)	1,119	Python
ExeDS (Huang et al., 2022)	534	Python
DS-1000 (Lai et al., 2022)	1,000	Python

Table A7: Overview of major NL2Code benchmarks. We pick BIRD-DEV and DS-1000 based on the following rationales: 1) we chose Python for it being widely used in NL2Code benchmarks, and we adopted DS-1000 due to the unsaturated performance (also see Table A8); 2) we included SQL, particularly BIRD-Dev, because of the introduced challenges due to its complexity and recency (2024), helping mitigate data contamination issues; and 3) we excluded Java, as the Java partitions of NaturalCodeBench only contains 70 problems, which seemed insufficient to reveal models’ true performance. For each benchmark, we report its size and programming languages covered.

Model	Human Eval (%)	MBPP (%)
DeepSeek-Coder-1.3B (Guo et al., 2024)	65.9	65.3
DeepSeek-Coder-6.7B (Guo et al., 2024)	74.4	74.9
GPT-4 Turbo (OpenAI, 2023)	85.4	85.7

Table A8: Model performance on two widely adopted NL2Code benchmarks, Human Eval and MBPP. As explained in §4.1, we do not use these two since they have approached saturated performance. Results are reported as of 04/03/2025.