

# $R^3$ : End-to-End Reasoning-based Planning for Multi-step Retrosynthesis via Reinforcement Learning

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## Abstract

Multi-step retrosynthetic planning is a fundamental challenge in organic chemistry, traditionally modeled as a combinatorial search problem guided by single-step prediction models. However, this search-centric paradigm often disconnects from the explicit chemical reasoning processes employed by human experts. In this paper, we propose  $R^3$  (Reinforced Reasoning Retrosynthesis), a novel framework that reformulates this task as end-to-end generative reasoning. Instead of traversing a search tree,  $R^3$  simulates the problem-solving logic of chemists to directly generate complete synthetic pathways. To achieve this, we initialize the model with domain knowledge and employ end-to-end Reinforcement Learning (RL) to optimize the entire planning policy. Experimental results on Retrobench show that  $R^3$  achieves a state-of-the-art Top-1 accuracy of 43.7%, demonstrating that generative reasoning offers a superior alternative to traditional search algorithms in solving complex retrosynthetic problems.

## 1 Introduction

Retrosynthesis planning is a fundamental strategy in organic chemistry and drug discovery, aiming to systematically deconstruct complex target molecules into simpler, commercially available precursors (Corey, 1991; Zheng et al., 2022). This task is generally categorized into two levels: single-step retrosynthesis and multi-step retrosynthetic planning. Specifically, single-step retrosynthesis aims to identify a set of immediate reactants that can transform into a given target molecule via a one-step chemical reaction (Jiang et al., 2023). In contrast, multi-step planning seeks to discover

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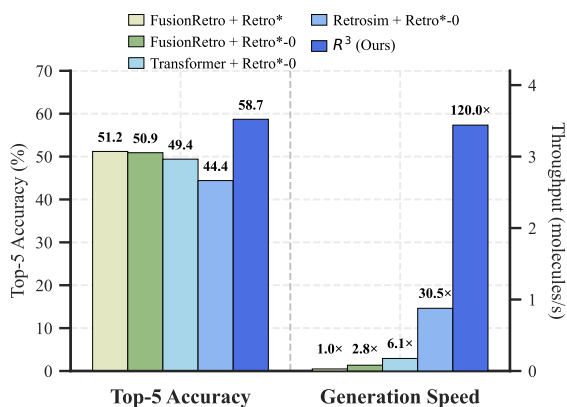


Figure 1: Performance comparison on Retrobench. The left chart displays the Top-5 accuracy (%), where  $R^3$  achieves 58.7%. The right chart illustrates the generation speed (throughput in molecules/s) annotated with relative speedup factors.  $R^3$  demonstrates superior efficiency, achieving a 120.0 $\times$  speedup compared to the FusionRetro + Retro\* baseline. Here, Retro\*-0 denotes the variant of Retro\* that does not employ a value model for search guidance; by contrast, Retro\* integrates a value model to optimize its search strategy, which leads to a slight improvement in performance but a trade-off of slower generation speed.

a complete, sequential reaction pathway that recursively transforms the target into a set of commercially available starting materials (building blocks) (Zheng et al., 2022; Zhong et al., 2023).

Traditionally, AI-driven methods have treated multi-step planning as a heuristic search problem. These approaches typically focus on either enhancing the accuracy of single-step models to prune the search tree (Kim et al., 2021; Liu et al., 2023a) or training value networks to guide algorithms like Monte Carlo Tree Search (MCTS) (Segler et al., 2018; Hong et al., 2023) or A\* search (Chen et al., 2020). While effective to some extent, these methods often suffer from high computational overhead and a “black-box” nature, lacking the explicit chemical reasoning that human experts employ when de-

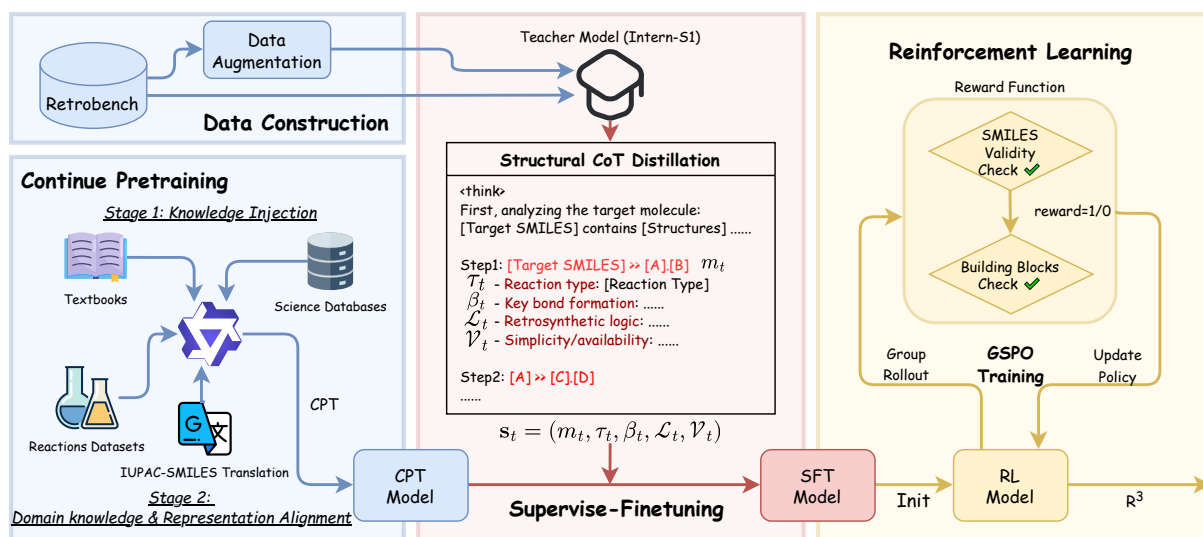


Figure 2: The overall framework of  $R^3$ . The training pipeline consists of three phases: (1) Continual Pretraining (CPT): Starting from Qwen3-8B-Base, the model undergoes a two-stage pretraining process on biomedical texts and reaction datasets to align chemical representations (SMILES/IUPAC) and inject domain knowledge. (2) Structural CoT Distillation (SFT): A teacher model (Intern-S1) distills reasoning capabilities into the student model using a structural reasoning protocol. The reasoning trajectory is structured into a 5-tuple sequence (reaction type  $\tau$ , key disconnection  $\beta$ , rationale  $L$ , etc.) to prevent logical drift. (3) Reinforcement Learning (RL): The model is further optimized using Group Sequence Policy Optimization (GSPO). The reward function  $r(x, y)$  strictly enforces both chemical validity of intermediate SMILES and the successful termination of the route at commercially available building blocks.

signing a route. Additionally, they are constrained by the performance ceiling of single-step models.

Recently, Large Language Models (LLMs) have shown remarkable capabilities in complex reasoning tasks across various domains (Guo et al., 2025; Jaech et al., 2024; Yu et al., 2025; Trinh et al., 2024; Li et al., 2022). Recent research has begun to explore the use of LLMs for retrosynthesis; however, these efforts are primarily limited to single-step retrosynthesis (Zhang et al., 2025a) or rely on integrating LLMs into search algorithms (Kang et al., 2025; Liu et al., 2025; Wang et al., 2025). The dependency on single-step models and search algorithms prevents these methods from fully exploiting the generative reasoning capabilities of LLMs for direct multi-step retrosynthetic route planning.

In this paper, we propose a novel framework to train an LLM (Qwen3-8B-Base) as a specialized agent for multi-step retrosynthetic planning. Our method shifts the paradigm from “searching” to “generative reasoning-based planning”, enabling the model to directly generate complete retrosynthetic routes in an end-to-end manner. This approach mimics the problem-solving logic of human chemists, who reason through each step of the retrosynthesis process rather than exhaustively

searching through all possibilities.

To achieve this, we first establish a robust chemical reasoning foundation by injecting domain knowledge and structured logic through continual pretraining and supervised fine-tuning. Building upon this, we employ end-to-end Reinforcement Learning (RL) to optimize the entire planning policy, ensuring that the model learns to generate valid retrosynthetic routes. Experimental results on Retrobench (Liu et al., 2023b) demonstrate that our model achieves state-of-the-art performance with a Top-1 accuracy of 43.7% and a Top-5 accuracy of 58.7%. Beyond accuracy,  $R^3$  demonstrates superior test-time scaling and inference efficiency, achieving a 120x speedup compared to traditional search-based baselines (Figure 1).

## 2 Related Work

**Search-based Multi-step Retrosynthesis** Traditionally, search-based multi-step retrosynthesis formulates the planning task as a search problem over a molecular graph or tree, aiming to identify a valid pathway of chemical reactions to transform a target molecule into commercially available starting materials. Foundationally, Monte Carlo Tree Search (MCTS) (Segler et al., 2018) and A\*-like

search strategies, such as Retro\* (Chen et al., 2020), formulate planning as a search problem guided by neural policy and value networks. Subsequent research has focused on enhancing search efficiency and route quality: Retro\*+ (Kim et al., 2021) and PDVN (Liu et al., 2023a) introduce self-improving mechanisms and dual-value networks to better estimate synthesis costs; EG-MCTS (Hong et al., 2023) and MEEA\* (Zhao et al., 2024) incorporate experience guidance and look-ahead exploration into tree search. Beyond single-target planning, methods like RetroGraph (Xie et al., 2022) and DreamRetroer (Zhang et al., 2025b) enable efficient group retrosynthesis. Additionally, recent frameworks like FusionRetro (Liu et al., 2023b) and CREBM (Liu et al., 2024a) integrate advanced molecular representations and energy-based reranking to further ensure the feasibility of the predicted routes.

**LLM-based Retrosynthesis** Recently, LLMs are increasingly being explored for retrosynthesis tasks due to their powerful reasoning capabilities. Some studies train foundation models on chemical data, however not specifically for retrosynthesis tasks (Frey et al., 2023; Bai et al., 2025). On single-step retrosynthesis tasks, LLM-based methods have been proven to possess state-of-the-art (SOTA) capabilities. (Zhang et al., 2025a; Yang et al., 2025b; Liu et al., 2024b). In multi-step retrosynthesis tasks, some studies have explored integrating LLMs into search algorithms to assist in refining retrosynthetic routes (Wang et al., 2025) or utilizing text descriptions as a modality to guide the search process (Kang et al., 2025). Specifically, Retro-R1 (Liu et al., 2025) provides the single-step model as a callable tool to the LLM, transforming multi-step retrosynthesis into an agent tool-use scenario. While the aforementioned methods leverage LLMs for retrosynthesis tasks, most are limited by the performance of the underlying single-step models and fail to utilize the LLMs’ superior reasoning capabilities to propose retrosynthetic routes directly.

## 3 Methodology

### 3.1 Problem Formulation

We formulate multi-step retrosynthetic planning as a sequential decomposition problem. Let  $\mathcal{M}$  denote the chemical space and  $\mathcal{B} \subset \mathcal{M}$  be the set of available building blocks. Given a target molecule  $x \in \mathcal{M} \setminus \mathcal{B}$ , the goal is to generate a

retrosynthesis plan  $\mathcal{T}$  — a directed acyclic graph where the root is  $x$  and all leaves belong to  $\mathcal{B}$ .

Traditionally, this problem is solved using search algorithms. The search process can be defined as finding a path in a state space graph where nodes represent molecules or sets of molecules, and edges represent chemical reactions. Let  $S_t$  be the set of precursor molecules at step  $t$ , with  $S_0 = \{x\}$ . At each step, a single-step retrosynthesis model  $f(m)$  predicts a set of possible reactants  $R$  for a molecule  $m \in S_t$ . The search algorithm (e.g. A\*) explores the tree of possible reactions to find a sequence such that the final set of leaves  $S_T \subseteq \mathcal{B}$ . The objective is often to minimize the cost of the route, defined as  $C(\mathcal{T}) = \sum_{r \in \mathcal{T}} c(r)$ , where  $c(r)$  is the cost of an individual reaction or material, or the depth of the route  $D(\mathcal{T})$ , which is the maximum number of reaction steps in any branch of the retrosynthesis tree.

Unlike searching algorithms, we propose a Generative Reasoning Paradigm. We model the retrosynthesis planner as a parameterized policy  $\pi_\theta$ . The model does not merely output the reaction pathway  $y$ ; instead, it generates a joint distribution of a reasoning trajectory  $z$  and the final plan  $y$ :

$$\pi_\theta(z, y|x) = \pi_\theta(y|x, z)\pi_\theta(z|x). \quad (1)$$

In this framework, the reasoning process  $z$  acts as a latent variable that bridges the gap between the target  $x$  and the solution  $y$ . The model first generates the chain-of-thought  $z$  to deduce the synthesis strategy step-by-step, and conditioned on this reasoning, it subsequently generates the structured retrosynthetic plan  $y$ .

### 3.2 Continual Pretraining

To enhance the model’s domain-specific foundation and specialized reasoning capabilities, we perform a two-stage continual pre-training process starting from Qwen3-8B-Base model (Yang et al., 2025a).

The first stage focuses on injecting general biomolecular and scientific knowledge. The training corpora for this stage are curated from specialized databases and literatures, including PubChem (Kim et al., 2025), PubMed (White, 2020), bioRxiv (Sever et al., 2019), and the biology and chemistry subsets of FineFineWeb (M-A-P et al., 2024). To ensure the model retains its general-purpose capabilities, we also incorporate a balanced portion of general data from FineWeb-Edu (Lozhkov et al., 2024). During this phase, we

follow standard unsupervised pre-training objectives, where the loss is calculated over the entire text sequence.

The second stage is specifically designed to enhance retrosynthesis-specific domain knowledge and representational alignment. We utilize a vast collection of chemical reactions sourced from the Open Reaction Database (ORD) (Kearnes et al., 2021a) and the USPTO (Lowe, 2012). We also integrate bidirectional translation tasks between molecular IUPAC names and SMILES. This representational alignment is essential because, during the subsequent reasoning process, molecular fragments and functional groups are frequently referenced by their IUPAC names in the textual chain-of-thought. Aligning these two modalities enables the model to bridge the gap between structural data and linguistic chemical concepts. This second stage shifts to a training paradigm closer to SFT, where the loss is calculated exclusively on the output tokens. Detailed statistics regarding dataset scales and specific data formats for each task are provided in Appendix Section A.

### 3.3 Structural Chain-of-Thought Distillation

A key challenge in end-to-end planning is that multi-step retrosynthesis inherently requires the model to maintain and navigate a complex search tree during its reasoning process. Although long CoT reasoning, as seen in models like DeepSeek-R1, enhances problem-solving, it often produces excessively long and unstructured text that is inefficient for representing tree-structured planning. This lack of structure makes it difficult for model to effectively learn the underlying chemical logic during the SFT process. To address this, we introduce Structural CoT Distillation.

**Reasoning Structure** Instead of free-form generation, we enforce a structured reasoning protocol. We define the reasoning trajectory  $z$  as a sequence of steps,  $z = (s_1, s_2, \dots, s_L)$ , where  $L$  represents the depth of the retrosynthesis tree. Each step  $s_t$  is modeled as a 5-tuple representing a comprehensive chemical decision unit:

$$s_t = (m_t, \tau_t, \beta_t, \mathcal{L}_t, \mathcal{V}_t) \quad (2)$$

$m_t$  (**Transformation**): The specific reaction mapping (e.g., SMILES string  $P \gg R_1.R_2$ ).

$\tau_t$  (**Reaction Type**): The categorical classification (e.g., 1,3-Dipolar cycloaddition), providing mechanistic context.

$\beta_t$  (**Key Disconnection**): The identification of the strategic bond to break (e.g., N-O bond of isoxazole ring), ensuring topological awareness.

$\mathcal{L}_t$  (**Strategic Rationale**): A natural language justification explaining why this step is favorable (e.g., thermodynamic stability, functional group tolerance).

$\mathcal{V}_t$  (**Availability Check**): A verification step assessing if the generated precursors are commercially available building blocks.

**Reasoning Data Distillation** We utilize the training dataset of Retrobench (Liu et al., 2023b) as the initial data source. To capture the multimodal nature of retrosynthesis, we augment the dataset using retro\* (Chen et al., 2020) to discover diverse valid pathways.

We employ Intern-S1 (Bai et al., 2025) as the teacher model. During the distillation phase, prompts provided to the teacher model includes the ground-truth route to facilitate the generation of high-quality reasoning traces. During the SFT phase, prompts for the student model contain only target molecules, ensuring that the model learns to plan without access to the answer.

**Supervised Fine-Tuning** We fine-tune the student model  $\pi_\theta$  to minimize the negative log-likelihood of this structured sequence:

$$\mathcal{L}_{SFT} = - \sum_{(x,z,y) \in \mathcal{D}_{SFT}} \log P_\theta(z, y|x) \quad (3)$$

### 3.4 Reinforcement Learning

To further improve the model’s performance on retrosynthesis tasks, we employ GSPO (Zheng et al., 2025), a variant of GRPO (Shao et al., 2024), as our RL algorithm.

Given a target product  $x$ , the policy  $\pi_\theta$  generates a retrosynthetic route  $y$ . We utilize a binary outcome-supervised reward function  $r(x, y)$ , defined to enforce both chemical validity and precise termination at the desired starting materials. Specifically,  $r(x, y) = 1$  if and only if all intermediate products in the route  $y$  are chemically valid SMILES, and the set of leaf nodes (starting materials) matches the ground-truth building blocks. Following the standard evaluation protocol on Retrobench (Liu et al., 2023b), this matching is

Table 1: Main Results on Retrobench. We compare  $R^3$  with SOTA baselines across three categories.  $R^3$  achieves new SOTA performance on all metrics, from Top-1 to Top-5 accuracy.

Category	Method	Top-1	Top-2	Top-3	Top-4	Top-5
		(%)	(%)	(%)	(%)	(%)
Template-based	Retrosim (Coley et al., 2017)	35.1	40.5	42.9	44.0	44.4
	Neuralsym (Segler and Waller, 2017)	42.0	49.3	52.0	53.6	54.3
	GLN (Dai et al., 2019)	39.6	48.9	52.7	54.6	55.7
Template-free	Transformer (Karpov et al., 2019)	31.3	40.4	44.7	47.2	49.4
	Megan (Sacha et al., 2021)	19.5	29.7	37.2	42.6	45.9
	FusionRetro (Liu et al., 2023b)	37.5	45.0	48.3	50.2	51.2
	FusionRetro+CREBM (Liu et al., 2024a)	39.6	46.7	49.5	51.0	51.7
	RetroInText (Kang et al., 2025)	42.1	49.9	53.0	54.7	55.7
Generative LLMs	Intern-S1 (Bai et al., 2025)	2.0	3.5	5.7	8.1	11.5
	GPT-5 (OpenAI, 2025)	2.4	6.3	10.7	12.0	14.9
	Gemini-3-pro (Google, 2025)	6.4	8.5	12.5	14.5	20.0
	$R^3$ (Ours)	<b>43.7</b>	<b>50.4</b>	<b>55.0</b>	<b>57.3</b>	<b>58.7</b>

performed by comparing the canonical InChiKeys of the molecules.

The GSPO objective is written as:

$$\mathcal{J}_{\text{GSPO}}(\theta) = \mathbb{E}_{\substack{x \sim \mathcal{D} \\ \{y_i\} \sim \pi_{\theta_{\text{old}}}}} \left[ \frac{1}{G} \sum_{i=1}^G \mathcal{L}_i(\theta) \right], \quad (4)$$

where  $\mathcal{L}_i$  represents the clipped objective for the  $i$ -th sample:

$$\mathcal{L}_i = \min \left( s_i \hat{A}_i, \text{clip}(s_i, 1 - \varepsilon, 1 + \varepsilon) \hat{A}_i \right). \quad (5)$$

GSPO utilize the group-based standardized advantage  $\hat{A}_i = (r(x, y_i) - \mu_r) / \sigma_r$  and the importance ratio  $s_i = (\pi_{\theta}(y_i|x) / \pi_{\theta_{\text{old}}}(y_i|x))^{1/|y_i|}$ . The importance ratio of GSPO is based on sequence-level, making it a more stable RL algorithm than GRPO (Zheng et al., 2025; Shao et al., 2024).

## 4 Experiments

### 4.1 Experimental Setup

**Dataset** We evaluate our method on Retrobench, a widely used benchmark for retrosynthesis prediction proposed by FusionRetro (Liu et al., 2023b). This benchmark is constructed from the USPTO reaction dataset and a set of commercially available building blocks. Detailed statistics regarding the dataset are provided in Appendix B. To ensure fair comparison and data consistency, we canonicalize all molecules in the dataset following the

methodology established in FusionRetro (Liu et al., 2023b).

**Baseline** We compare our method against two categories of baselines: (1) Search-based methods, including Retrosim (Chen et al., 2020), Neuralsym (Kim et al., 2021), GLN (Dai et al., 2019), Transformer (Karpov et al., 2019), Megan (Sacha et al., 2021), FusionRetro (Liu et al., 2023b), CREBM (Liu et al., 2024a), and RetroInText (Kang et al., 2025); (2) Generative LLMs, including Intern-S1 (Bai et al., 2025), GPT-5 (OpenAI, 2025), and Gemini-3-pro (Google, 2025). For search-based methods, we report the results using the search algorithm (Retro\* or Retro\*-0) that achieves the best performance. For LLM baselines, we use the same prompt as our method during inference to ensure a fair comparison, and evaluate them on a subset of the test set due to computational constraints.

**Metric** We evaluate the performance using the Top-1 exact match accuracy. The match is based on the comparison of the InChiKey of the predicted building blocks and the ground-truth building blocks, following the evaluation protocol established in FusionRetro (Liu et al., 2023b).

For Generative LLMs, we adopt a specific protocol for Top-K evaluation. We oversample 8 responses for each test instance and remove duplicates. To compute Top-K accuracy, we randomly

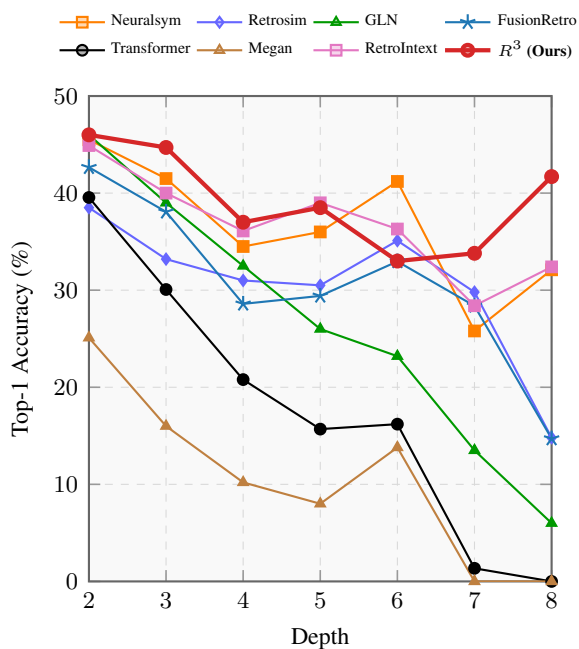


Figure 3: Comparison of Top-1 Accuracy across varying synthetic route depths.  $R^3$  demonstrates superior performance, particularly at greater depths, compared to baselines including Neursym, Retrosim, and FusionRetro.

select  $k$  unique samples, extract their corresponding retrosynthetic route trees, and merge them into a unified graph. A test case is considered solved if the set of ground-truth starting materials is contained within the nodes of this merged graph. We conduct sampling across various temperatures and top-p settings and report the best performance for Top-K metrics.

## 4.2 Results

**Comparison with Baselines** The main results are summarized in Table 1. Our method achieves a Top-1 accuracy of 43.7% on Retrobench, establishing a new state-of-the-art. This performance significantly surpasses both the strongest search-based baseline, RetroInText (42.1%), and the fusion-based approach, FusionRetro (37.5%).

Beyond Top-1 accuracy,  $R^3$  demonstrates superior capability in generating diverse valid pathways. In terms of Top-k performance, our model consistently outperforms all baselines, achieving 50.4% at Top-2 and 58.7% at Top-5. Crucially, this significant performance gain at higher  $k$  values evidences the model’s **test-time scaling** capability. It demonstrates that by increasing the computational budget at inference time,  $R^3$  can effectively tackle a wider array of challenging retrosynthesis problems that

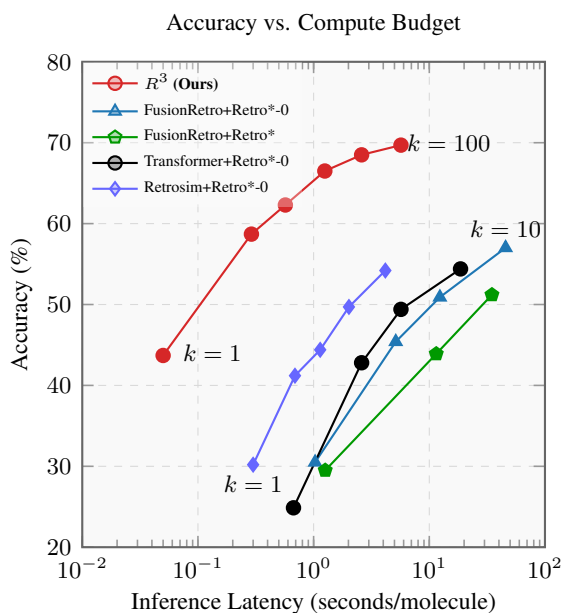


Figure 4: Inference Scaling Efficiency. We plot the cumulative Top-K accuracy against the average inference latency per molecule on a logarithmic scale. For FusionRetro, distinct points correspond to increasing beam sizes (from  $k = 1$  to 10), illustrating the significant latency cost required to expand the search space. In contrast,  $R^3$  demonstrates a superior Pareto frontier, efficiently converting test-time compute (sampling budget  $k$ ) into performance gains without the exponential overhead of planning algorithms.

remain unsolved by a single greedy decoding pass.

Furthermore, the results highlight the necessity of domain-specific adaptation. General-purpose LLMs, such as GPT-5 and Gemini-3-pro, achieve negligible accuracy ( $< 7\%$ ), confirming that the complex logic of retrosynthesis cannot be solved by reasoning capabilities alone without specialized chemical knowledge injection and alignment.

**Performance at Different Depths** To analyze the model’s robustness across varying levels of retrosynthesis complexity, we evaluate  $R^3$  against baselines across route depths ranging from 2 to 8. As illustrated in Figure 3,  $R^3$  consistently outperforms all baselines at nearly every depth level.

Notably, traditional search-based methods often suffer from the “curse of dimensionality” as the search tree deepens. In contrast,  $R^3$  exhibits a substantial advantage at greater depths (e.g., maintaining  $> 41\%$  accuracy at depth 8), whereas methods like FusionRetro degrade significantly (dropping to  $\sim 14\%$ ). This suggests that our generative reasoning paradigm effectively captures global structural dependencies, preventing the error propagation typ-

Model	Top-1 Acc (%)
Qwen3-8B-Base	0.0
+ CPT	0.0
+ SFT (w/o Structural CoT)	4.0
+ SFT	28.3
+ SFT + RL	29.2
+ CPT + SFT	33.1
+ CPT + SFT + RL	<b>43.7</b>

Table 2: Ablation study on Retrobench Top-1 Accuracy. We illustrate the impact of CPT, SFT, and RL (200 steps) on the Qwen3-8B-Base model.

ical in greedy or heuristic search algorithms.

### 4.3 Computing Efficiency

In high-throughput scenarios, inference latency is a critical bottleneck. As shown in Table 1,  $R^3$  achieves an average throughput of **3.44** molecules per second to generate 5 samples, representing a **120 $\times$  speedup** over FusionRetro+Retro\* (0.028mol/s). This dramatic acceleration stems from our shift to a generative paradigm: unlike search algorithms that iteratively navigate a combinatorial tree (often stalling GPU utilization below 35% due to CPU-GPU overhead),  $R^3$  generates the complete route in a single autoregressive pass. This allows  $R^3$  to leverage optimizations like FlashAttention to maintain  $> 90\%$  GPU utilization, while search-based methods only achieve  $\sim 35\%$  GPU utilization.

Figure 4 further visualizes the trade-off between compute budget and performance.  $R^3$  establishes a superior **Pareto frontier**, delivering competitive accuracy at a fraction of the latency. Crucially, our model demonstrates effective **test-time scaling**: increasing the sampling budget to  $k = 100$  boosts coverage to 69.7% while maintaining a total latency ( $\sim 5.7$ s per molecule) that is still orders of magnitude faster than the search-based baseline. In contrast, search algorithms like FusionRetro face a scaling wall: expanding the search width (e.g., beam size) from  $k = 1$  to  $k = 10$  incurs a massive latency penalty (from  $\sim 1$ s to  $\sim 45$ s) due to the combinatorial explosion of the search space, yielding only marginal accuracy gains due to the diminishing returns of exploring lower-probability branches.

### 4.4 Ablation Study

To decouple the contributions of our three-stage training pipeline, we analyze the functional role of each phase as shown in Table 2. The results highlight a synergistic dependency among the stages:

**SFT as the Cold-Start Foundation** SFT is the prerequisite for the model to function. The base model (0.0%) completely fails to solve any retrosynthesis tasks. SFT is essential to “cold start” the policy; without the structural constraints learned during this phase, the model cannot generate valid reasoning traces, rendering subsequent RL impossible to launch.

**CPT as the Knowledge Reservoir** While SFT adapts the format, the model does not acquire plentiful new domain knowledge. The necessity of CPT is evident when comparing *Base+SFT+RL* (29.2%) with our full method (43.7%). Without the domain-specific representation space constructed during CPT, the model hits an early performance ceiling. CPT provides the fundamental chemical logic that allows the model to respond to RL optimization effectively.

**RL as the Performance Catalyst** Finally, RL acts as the catalyst to unlock the model’s latent potential. On top of the CPT-enhanced backbone, RL drives a massive **10.6% improvement** (33.1%  $\rightarrow$  43.7%). This confirms that while CPT provides the knowledge and SFT provides the form, only RL can align these capabilities with the long-horizon goal of successful multi-step planning.

**Ablation on Structural CoT** To assess the impact of our proposed Structural Chain-of-Thought distillation, we conduct an ablation where we directly fine-tune the model using free-form CoT traces generated by the teacher model, without enforcing the structured 5-tuple schema. This variant achieves only 4.0% accuracy. By analyzing the SFT data, we observe that the teacher model often produces excessively long and unstructured reasoning paths, and tends to reveal ground-truth answers in the prompt. This leads to logical drift and poor generalization during inference. In contrast, our structured approach compels the model to explicitly reason about each chemical decision, significantly enhancing both interpretability and performance.

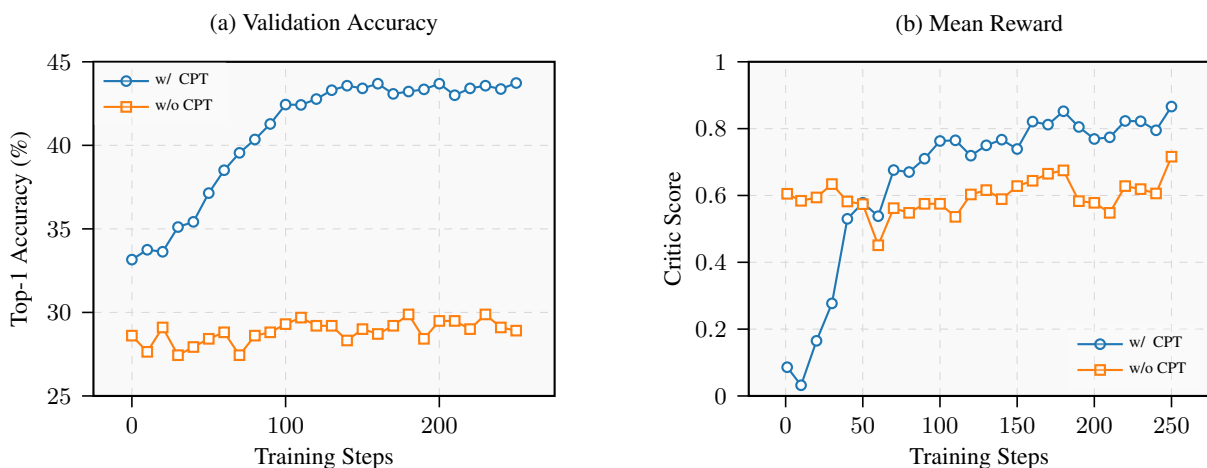


Figure 5: Training dynamics of CPT model vs. non-CPT model during RL stage.

#### 4.5 Building Block Awareness Analysis

A critical concern for generative planners is the potential hallucination of unavailable starting materials. Unlike search-based methods that query a database at every step,  $R^3$  must internalize the definition of available building blocks ( $\mathcal{B}$ ).

To quantify this capability, we constructed a classification probe dataset from Retrobench. We labeled ground-truth starting materials as *positive* and intermediate products as *negative*. We then queried Qwen3-8B to determine if a given molecule belongs to  $\mathcal{B}$ . As shown in Table 3, Qwen3-8B achieves a Pass@1 accuracy of 82.0% and Pass@3 of 94.3%. This high accuracy demonstrates that the base model already learned the chemical space of commercially available materials, which is crucial for generating valid retrosynthetic routes without external database lookups.

Table 3: **Building Block Identification Accuracy.** The model is asked to classify whether a molecule is a commercially available building block.

Model	Pass@1	Pass@3
Qwen3-8B	82.0%	94.3%

#### 4.6 RL Training Dynamics

We present the training dynamics during the RL stage in Figure 5. The model enhanced with CPT exhibits steady improvements in both validation accuracy and mean reward over 250 training steps. In contrast, the model without CPT fails to improve, stagnating at its initial SFT performance level. We observe that the CPT model starts with

a lower initial reward but quickly surpasses the non-CPT model within the first 50 steps. This may be attributed to the injection of non-conversational corpora during the CPT phase, which temporarily misaligns the response style but ultimately provides a richer knowledge base for effective RL optimization.

## 5 Conclusion

In this work, we present  $R^3$ , a novel framework that leverages Large Language Models for multi-step retrosynthetic planning. By reformulating the task as a generative reasoning problem and employing end-to-end reinforcement learning, we enable the model to generate interpretable and chemically valid retrosynthesis routes. Our extensive experiments on Retrobench demonstrate that  $R^3$  achieves state-of-the-art performance, significantly outperforming existing search-based and LLM-based methods. This work highlights the potential of LLMs in complex scientific reasoning tasks and opens new avenues for future research in AI-driven retrosynthetic planning.

### Limitations

Despite the promising results, this work relies primarily on the exact match accuracy of starting materials to evaluate performance, which may not fully capture the holistic quality of the generated retrosynthetic routes, such as reaction conditions, yield, or atom economy. Furthermore, the evaluation is constrained by the Retrobench dataset, which contains a limited number of ground-truth pathways for each target molecule; consequently, chemically valid and efficient routes generated by

our model that deviate from these annotated references may be overlooked, potentially underestimating the model's true planning capabilities due to the incomplete coverage of the vast chemical search space.

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## A Details of Continual Pretraining

This section provides a comprehensive breakdown of the data sources, formatting strategies, and training objectives employed during the two-stage continual pre-training process. The detailed statistics for each dataset are summarized in Table 4, and the training hyperparameters are summarized in Table 5.

### A.1 Stage 1: Biomolecular and Scientific Knowledge Injection

The primary objective of Stage 1 is to imbue the model with a broad foundation in biomolecular science and chemical structures. We categorize the data used in this stage into three distinct types based on their structural format:

**Molecule+Text:** This data type consists of molecular structural information paired with descriptive natural language. Specifically, we extract SMILES sequences from PubChem (Kim et al., 2025) and pair them with their corresponding textual descriptions (e.g., chemical properties, taxonomy, and pharmacological action). This allows the model to associate structural representations with scientific concepts.

**Interleaved Text:** To enhance the model’s ability to handle multi-modal scientific literature, following Pei et al. (2024), we process corpora from bioRxiv (Sever et al., 2019) and PubMed (White, 2020) Abstracts using BERN2 (Sung et al., 2022), a high-performance biomedical entity recognition tool. We identify biological and chemical entities within the text and either insert or replace them with their corresponding SMILES sequences or IUPAC names. This creates a dense, interleaved representation that enables the model to establish a direct associative link between natural language mentions and their corresponding chemical structures.

**Pure Sequence Data:** This includes large-scale collections of SMILES and IUPAC names from PubChem (Kim et al., 2025), as well as scientific text from FineWeb-biology/chemistry (M-A-P et al., 2024) and PubMed Central (White, 2020). We also include FineWeb-Edu (Lozhkov et al., 2024) to maintain capabilities in the general domain. These datasets are treated as pure sequences.

During this stage, we employ a standard unsupervised pre-training objective, calculating the cross-entropy loss over the entire sequence to maximize the model’s predictive accuracy across both textual and chemical modalities.

### A.2 Stage 2: Retrosynthesis Specialization and Representational Alignment

The second stage transitions toward retrosynthesis reaction and the alignment of different molecular notations. The data types in this stage are formulated as follows:

**Retrosynthesis Task:** Utilizing data from the Open Reaction Database (ORD) (Kearnes et al., 2021b) and USPTO (Lowe, 2012), we formulate retrosynthesis as a sequence-to-sequence task. The input is the SMILES of product molecule, and the model is trained to predict the corresponding reactants.

**Cross-Modal Alignment (IUPAC  $\leftrightarrow$  SMILES):** To bridge the gap between nomenclature and structure, we perform bidirectional translation tasks. For the IUPAC to SMILES task, the input is a molecule’s IUPAC name and the output is its SMILES sequence; the SMILES to IUPAC task follows the reverse logic. This alignment is critical for ensuring the model can accurately reference molecular fragments by name during complex chain-of-thought reasoning.

**USPTO-Application:** A unique case in Stage 2 is the USPTO-Application dataset. We use its interleaved version provided by PRESTO (Cao et al., 2024). Specifically, the USPTO-Application dataset comprises chemical reaction equations and experimental procedures, where molecular entities identified by BERN2 (Sung et al., 2022) are replaced with their corresponding SMILES. To accommodate the SFT loss format, its input is set as an empty sequence, while the entire interleaved text serves as the output.

Unlike Stage 1, the training objective in Stage 2 shifts to an SFT-style loss, where the gradient is calculated exclusively on the output tokens (e.g., the reactants, the translated notation, or the interleaved content in USPTO-Application), thereby refining the model’s precision in specialized chemical generation.

## B Dataset Details

The Retrobench dataset consists of 46,458 training routes, 5,803 validation routes, and 5,838 test routes for evaluation. We summarize the data statistics of Retrobench in Table 6. The dataset is categorized by the depth of the ground-truth retrosynthetic routes.

Table 4: Detailed data configuration of the two-stage continual pre-training.

Data Source	Data Type	Samples	Tokens
<i>Stage 1</i>			
PubChem (Kim et al., 2025)	Molecule+Text	387.4K	70.7M
bioRxiv (Sever et al., 2019)	Interleaved Text	2.3M	495.3M
PubMed Abstract (White, 2020)	Interleaved Text	33.4M	10.4B
PubChem (Kim et al., 2025)	SMILES	114.8M	4.1B
PubChem (Kim et al., 2025)	IUPAC	114.8M	5.6B
FineFineWeb-biology (M-A-P et al., 2024)	Text	7.7M	8.1B
FineFineWeb-chemistry (M-A-P et al., 2024)	Text	9.0M	8.0B
PubMed Central (White, 2020)	Text	70.6M	13.0B
FineWeb-Edu (Lozhkov et al., 2024)	Text	19.7M	20.2B
<i>Stage 2</i>			
Open Reaction Database (Kearnes et al., 2021b)	Retrosynthesis	1.8M	135.5M
USPTO-Full (Dai et al., 2019)	Retrosynthesis	771.9K	50.1M
USPTO-MIT (Zhong et al., 2022)	Retrosynthesis	406.9K	31.2M
USPTO-Application (Cao et al., 2024)	Interleaved Text	1.6M	532.1M
PubChem (Kim et al., 2025)	IUPAC to SMILES	15M	1.2B
PubChem (Kim et al., 2025)	SMILES to IUPAC	15M	1.2B

Table 5: Detailed training configuration of the two-stage continual pre-training.

	Continual Pretraining Stage 1	Continual Pretraining Stage 2
Batch Size	1,024	256
Max Length	4,096	4,096
Training Tokens	70.1B	3.2B
Learning Rate Schedule	Cosine with min lr	Cosine
Peak Learning Rate	$6.0 \times 10^{-5}$	$1.0 \times 10^{-4}$
GPU Devices	4 × A100	4 × A100

## C Hyperparameters

We utilize the Qwen3-8B-Base model (Yang et al., 2025a) as base model, LlamaFactory (Zheng et al., 2024) as SFT framework, and VeRL (Sheng et al., 2024) as RL framework. During SFT, we set the batch size to 64, learning rate to  $3e - 5$ , and train for 3 epochs. During RL, we set the batch size to 64, group size to 16, and learning rate to  $1e - 6$ . The training curves are shown in Figure 5.

## D Case Study

We provide a concrete example of a retrosynthetic route generated by  $R^3$  in Figure 6. This visualization demonstrates the model’s ability to decompose a complex target into available building blocks. For brevity, the input prompts and the output reasoning traces are truncated.

Table 6: Data statistics of Retrobench. The table reports the number of molecules for each route depth across Training, Validation, and Test sets.

Dataset	Depth											
	2	3	4	5	6	7	8	9	10	11	12	13
Training	22,903	12,004	5,849	3,268	1,432	594	276	107	25	0	0	0
Validation	2,862	1,500	731	408	179	74	34	13	2	0	0	0
Test	2,862	1,500	731	408	179	74	34	13	2	32	2	1

### MODEL INPUT (ABBREVIATED)

You are an expert in retrosynthesis planning. Your task is to propose a sequence of chemical reactions that can synthesize a target molecule from available starting materials.

#### ## Task Description

Given a target molecule (in SMILES format), work backwards to find a sequence of reactions that breaks down the target into simpler molecules that are reasonably available as starting materials.

#### ## Output Format

Provide the final answer as a JSON list of reaction strings in the format "PRODUCT>>REACTANTS".

Please propose a retrosynthesis route for the following target molecule:

Target molecule: Cn1ncc(C(=O)N2CCCC2)c1C(=O)Nc1ccc2nc(-c3ccnc3)nn2c1

### MODEL OUTPUT (PARTIAL DISPLAY)

<think>

Step 1: Cn1ncc(C(=O)N2CCCC2)c1C(=O)Nc1ccc2nc(-c3ccnc3)nn2c1>>Cn1ncc(C(=O)O)c1C(=O)Nc1ccc2nc(-c3ccnc3)nn2c1.C1CCNC1

- Reaction type: Amide Bond Disconnection (Aliphatic)
- Key bond formed: Amide bond between a pyrazole-carboxylic acid and pyrrolidine.
- Retrosynthetic Logic: The aliphatic amide is disconnected first. This is strategic because pyrrolidine is a simple, commercially available secondary amine. Forming this bond late-stage avoids potential side reactions involving the amine during the earlier construction of the heteroaromatic core.
- Reactant Availability: Pyrrolidine (C1CCNC1) is a common commodity chemical.

Step 2: Cn1ncc(C(=O)O)c1C(=O)Nc1ccc2nc(-c3ccnc3)nn2c1>>CCOC(=O)c1cnn(C)c1C(=O)Nc1ccc2nc(-c3ccnc3)nn2c1

- Reaction type: Functional Group Interconversion (Ester Hydrolysis)
- Key bond formed: Transformation of carboxylic acid back to ethyl ester.
- Retrosynthetic Logic: The free carboxylic acid is traced back to its ethyl ester precursor. This is necessary because the ester group is typically carried through the ring-forming steps (e.g., Knorr synthesis or condensation) used to build the pyrazole skeleton. Reverting to the ester prepares the molecule for the \*next\* disconnection: breaking down the complex core into simpler fragments.
- Reactant Availability: The ester is a key synthetic intermediate (not yet a starting material) that facilitates further structural decomposition.

Step 3: ... [Remaining steps omitted for brevity] ...

</think>

<answer> ["Cn1ncc(C(=O)N2CCCC2)c1C(=O)Nc1ccc2nc(-c3ccnc3)nn2c1>>Cn1ncc(C(=O)O)c1C(=O)Nc1ccc2nc(-c3ccnc3)nn2c1.C1CCNC1", ... ] </answer>

Figure 6: A representative case study generated by  $R^3$ .