

AtomSQL: Interactive Disambiguation of NL-to-SQL via User-Guided Atom-Level Alignment

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ABSTRACT

Natural language to SQL (NL-to-SQL) systems are widely used for querying databases in natural language, achieving high accuracy on benchmark datasets. In practice, however, these systems face a fundamental challenge: a single natural language question can have multiple reasonable SQL interpretations, differing in predicates, aggregation functions, join paths, or set modifiers such as `DISTINCT`. This ambiguity is typically localized to specific parts of a query rather than the query as a whole. Existing approaches either assign a single confidence score to the entire query, which does not reveal where models disagree, or engage users in multi-turn clarification dialogues that resolve ambiguity without ever showing the candidate queries that motivated the questions, leaving users unable to make informed decisions about their intent.

We demonstrate `ATOMSQL`, a system that helps users resolve ambiguity in NL-to-SQL at the granularity of individual query components, and learns from their choices to produce better results over time. `ATOMSQL` runs multiple NL-to-SQL models on a user’s question, decomposes each generated query into *atoms* (clause-level units such as join conditions, predicates, and aggregation functions), and computes a confidence score for each atom based on cross-model agreement. Atoms are highlighted in the interface alongside the competing alternatives, so the user can pinpoint exactly where interpretations differ and select the one that matches their intent. Each selection is recorded and used to personalize future query results toward that user’s preferences.

1 INTRODUCTION

Natural language to SQL (NL-to-SQL) systems are increasingly used to help users query databases without writing SQL. Recent systems can generate executable SQL queries for complex questions and achieve high accuracy on benchmark datasets like Spider [11]. As a result, NL-to-SQL is often treated as a solved or nearly solved problem from an accuracy perspective.

However, in practice, users face a fundamental challenge: a natural language question may have multiple reasonable SQL interpretations, even when all generated queries are valid and executable. Real user questions are often underspecified: the same question can map to several distinct SQL queries that differ in predicates, aggregation functions, join paths, or set modifiers like `DISTINCT`. Recent work [3, 9] has formally characterized this phenomenon, showing that a single question can have multiple valid SQL answers and that such ambiguity is prevalent in real-world queries. This ambiguity is often localized to specific parts of the SQL query rather

than the entire query, so two systems may agree on most of a query but disagree sharply on one or two components.

One approach to handle this ambiguity is to attach a confidence score to each generated query. If the system is uncertain about its output, a low score can warn the user to double-check. A more interactive alternative is to detect ambiguity and engage the user in a clarification dialogue, asking targeted questions until the intended SQL is pinned down. Both directions capture something important: confidence signals help users decide when to trust the output, and clarification dialogues can resolve ambiguity before a query is executed. Yet, as shown in Example 1, each also leaves a significant gap when confronted with the localized, component-level nature of SQL ambiguity.

Example 1.1 (Atom-level ambiguity in NL-to-SQL). Consider an administrator querying a student-pet database. She wants to find students meeting certain age and pet ownership criteria for a housing allocation report, so she asks:

Q: “Show Two Older Students Who Have Two Pets.”

When multiple NL-to-SQL models process this question, they generate substantially different queries:

T5-LM generates:

```
SELECT has_pet.StuID
FROM Has_pet
JOIN Student ON has_pet.StuID = student.StuID
GROUP BY has_pet.StuID
HAVING COUNT(*) = 2
```

Llama 3 (8B parameters) generates:

```
SELECT *
FROM Student
WHERE Age > (SELECT AVG(Age) FROM Student)
AND LName IN (
  SELECT student.LName
  FROM Student
  JOIN Has_Pet ON student.StuID = has_pet.StuID
  JOIN Pets ON has_pet.PetID = pets.PetID
  GROUP BY student.LName
  HAVING COUNT(pets.PetType) = 2
)
```

SQLCoder generates:

```
SELECT s.fname, s.lname, COUNT(h.petid) AS
  pet_count
FROM student s
JOIN has_pet h ON s.stuid = h.stuid
GROUP BY s.fname, s.lname
HAVING COUNT(h.petid) >= 2
```

*Joint first author

```
ORDER BY s.age DESC
LIMIT 2;
```

All three queries are schema-valid and executable, but they encode different interpretations. Some parts are stable across models (all agree on joining Student and Has_Pet), while others diverge sharply: whether “two pets” means exactly two ($= 2$) or at least two (≥ 2), whether “older” requires an age predicate, and whether to apply ORDER BY and LIMIT.

Related Work. The example above reflects a recurring pattern in practice. Most NL-to-SQL systems [5, 6, 8] produce a single SQL query without modeling ambiguity or exposing uncertainty to the user, and existing efforts to address this fall into two categories, each with a distinct shortcoming.

Query-level confidence methods [1, 10] assign a single confidence score to each generated SQL query, treating the entire query as an atomic unit. Returning to our example: a confidence score might flag that the output is uncertain, but it cannot tell the user whether the disagreement is about the HAVING threshold, the age predicate, or both. Ambiguity in NL-to-SQL is typically localized to specific components. A whole-query score does not reveal where the models disagree, so the user cannot tell which parts of the query to trust and which to question.

Interactive clarification systems [2, 4, 7] use multi-turn conversations to produce a single SQL query. These systems ask the user targeted questions to resolve ambiguity, but the interaction is one-sided: the user answers questions without ever seeing the candidate queries that motivated them. In our example, such a system might ask “Do you mean exactly two pets or at least two?” without revealing that there are also open questions about the age predicate and whether to apply an ORDER BY clause. The user resolves one ambiguity while remaining unaware of the others, and the final query may still not match their intent.

Recent datasets and benchmarks [3, 9] demonstrate that ambiguity is prevalent in real-world NL-to-SQL queries. AMBROSIA [9] and PRACTIQ [3] formally characterize how a single question can yield multiple valid SQL answers.

ATOMSQL. We address these limitations by treating each SQL query as a composition of *atoms*: indivisible semantic units such as tables, join conditions, predicates, aggregation functions, and modifiers like DISTINCT. ATOMSQL runs multiple NL-to-SQL models, decomposes each output into atoms, and compares atoms across models. Cross-model agreement yields per-atom confidence scores that show exactly where interpretations diverge, providing the per-atom granularity that whole-query confidence scores lack. Instead of asking the user abstract questions, the interface shows the generated query with uncertain atoms highlighted and the available alternatives for each atom shown alongside. When a user selects a preferred interpretation, ATOMSQL learns this preference and applies it to future queries.

Demonstration. In our demonstration, participants will interact with ATOMSQL and observe how it exposes ambiguity in NL-to-SQL queries. Using real-world datasets, we will showcase how ATOMSQL identifies which query atoms are stable versus uncertain, presents alternative interpretations for ambiguous atoms, and adapts to user preferences over time. Participants will experience firsthand how

the interface enables informed decision-making without requiring multi-turn conversations.

2 SYSTEM OVERVIEW

We model NL-to-SQL ambiguity as a probabilistic alignment between natural language (NL) atoms and SQL atoms. ATOMSQL combines cross-model agreement and user preference to produce a final alignment. It is composed of five components: (1) a **model execution layer** that runs all active NL-to-SQL models in parallel; (2) an **atom decomposer** that parses each SQL output and the user query into clause-level atoms; (3) an **agreement and confidence engine** that measures cross-model consensus per atom; (4) a **user preference store** that records and applies past user choices; and (5) a **query assembler** that scores candidates and selects the best one. We now describe components (2)–(5) in detail, using the motivating query “Show Two Older Students Who Have Two Pets” as a running example. The full atom matrix for this example is larger and sparse, so for clarity we focus on the subset of atoms that capture the main ambiguities.

2.1 Atoms and Setup

A user query is decomposed into n NL atoms x_1, \dots, x_n , and the outputs of all active models are parsed into a set of m SQL atoms a_1, \dots, a_m . Each NL atom maps to one or more SQL atoms; the goal is to find the most likely mapping.

Using the motivating query “Show Two Older Students Who Have Two Pets”, we identify the following NL atoms:

$x_1 = \text{students}$, $x_2 = \text{older}$, $x_3 = \text{two pets}$, $x_4 = \text{show two}$

The full union of SQL atoms across all model outputs is larger than what is convenient to show inline. In practice, most of these atoms are unambiguous and yield sparse rows in the alignment matrix. For clarity, we focus on the atoms corresponding to the main ambiguities in this example:

$a_1 = \text{WHERE Age} > \text{AVG(Age)}$, $a_2 = \text{ORDER BY Age DESC}$,
 $a_3 = \text{HAVING COUNT(*)} = 2$, $a_4 = \text{HAVING COUNT(petid)} \geq 2$,
 $a_5 = \text{LIMIT } 2$

These focused atoms correspond to the ambiguous NL atoms:

$x_2 \rightarrow \{a_1, a_2\}$, $x_3 \rightarrow \{a_3, a_4\}$, $x_4 \rightarrow \{a_5\}$

Here, x_2 (“older”) admits multiple interpretations, such as a filtering condition (WHERE Age > AVG(Age)) or a ranking signal (ORDER BY Age DESC). Likewise, x_3 (“two pets”) can be interpreted as exactly two pets or at least two pets. The atom x_4 (“show two”) corresponds to the LIMIT 2 constraint.

2.2 Model Agreement

The agreement matrix $M \in \mathbb{R}^{n \times m}$ encodes cross-model consensus, where each row corresponds to an NL atom x_i and each column corresponds to a SQL atom a_j . The entry M_{ij} represents the proportion of models that map x_i to a_j , with $M_{ij} = 1$ indicating full agreement and lower values indicating disagreement.

In the full example, the matrix includes many atoms and is highly sparse: most NL atoms map consistently to a single SQL atom across models, producing rows with a single dominant nonzero entry. For clarity, we focus on the ambiguous rows corresponding to x_2

(“older”), x_3 (“two pets”), and x_4 (“show two”), and the focused atoms a_1, \dots, a_5 above.

With three models in our running example, the focused agreement matrix is:

$$M = \begin{bmatrix} \frac{1}{2} & \frac{1}{2} & 0 & 0 & 0 \\ 0 & 0 & \frac{2}{3} & \frac{1}{3} & 0 \\ 0 & 0 & 0 & 0 & \frac{1}{3} \end{bmatrix}$$

This means that for the atom “older,” the models split between a filtering interpretation and an ordering interpretation; for “two pets,” two models prefer the exact-count interpretation while one prefers the at-least-two interpretation; and for “show two,” only one model explicitly produces a LIMIT 2 atom.

2.3 User Preference and Alignment

With no prior user history, we initialize $U = M$:

$$U = \begin{bmatrix} \frac{1}{2} & \frac{1}{2} & 0 & 0 & 0 \\ 0 & 0 & \frac{2}{3} & \frac{1}{3} & 0 \\ 0 & 0 & 0 & 0 & \frac{1}{3} \end{bmatrix}$$

We combine model agreement and user preference using a logarithmic formulation:

$$P_i = \text{softmax}(\log M_i + \beta \log U_i)$$

where β controls the influence of user preference.

When no prior user history is available, we set $\beta = 0$, yielding:

$$P_i = \text{softmax}(\log M_i) = M_i$$

Thus, the initial alignment follows model agreement exactly:

$$P = \begin{bmatrix} \frac{1}{2} & \frac{1}{2} & 0 & 0 & 0 \\ 0 & 0 & \frac{2}{3} & \frac{1}{3} & 0 \\ 0 & 0 & 0 & 0 & \frac{1}{3} \end{bmatrix}$$

As user feedback becomes available ($U \neq M$), we increase β , allowing user preferences to influence the alignment.

2.4 Preference Update

When a user selects atom a_j for NL atom x_i , we increment U_{ij} by a step size $\alpha > 0$ and renormalize row i , shifting P toward the user-selected atom in future queries. Suppose the user selects a_3 (exactly two pets) over a_4 for x_3 (“two pets”), with step size $\alpha = 0.3$. The corresponding row of U is updated as:

$$[0, 0, \frac{2}{3}, \frac{1}{3}, 0] \rightarrow [0, 0, \frac{2}{3} + 0.3, \frac{1}{3}, 0] = [0, 0, 0.967, 0.333, 0]$$

We then renormalize the row to obtain a probability distribution:

$$[0, 0, 0.744, 0.256, 0]$$

The updated U shifts the alignment P toward the user-selected atom in subsequent queries.

If none of the candidate SQL atoms matches the user’s intent, the user can reformulate the query or enable additional models to generate a richer set of candidate atoms.

In addition to atom-level feedback, we maintain a model preference score $\text{Pref}(m)$ for each model m , which reflects how often a user selects atoms originating from that model. This captures user trust in different models over time.

2.5 Confidence and Query Scoring

For each atom, we define a confidence score $C_{ij} = M_{ij}^{raw} \cdot P_{ij}$, where M_{ij}^{raw} is the unnormalized agreement count (i.e., the fraction of models that produced SQL atom a_j for NL atom x_i before row normalization), and P_{ij} is the final alignment probability. After the user interaction above:

$$C = \begin{bmatrix} \frac{1}{2} & \frac{1}{2} & 0 & 0 & 0 \\ 0 & 0 & 0.496 & 0.085 & 0 \\ 0 & 0 & 0 & 0 & \frac{1}{3} \end{bmatrix}$$

To select the best SQL output, we compute scores at the query level. While confidence is estimated at the atom level, the final decision is made at the query level by aggregating these signals. The atom-level confidence C_{ij} captures the reliability of individual NL-to-SQL mappings, and these are combined to score each candidate query.

The atom score $S_{\text{atom}}(q)$ measures how well a SQL output q aligns with the user’s NL query by averaging alignment probabilities P_{ij} over all atoms in q , effectively summarizing per-atom confidence into a single query-level score. The final score $S_{\text{final}}(q)$ further incorporates model-level preference:

$$S_{\text{atom}}(q) = \frac{1}{|A_q|} \sum_{j \in A_q} \sum_{i=1}^n P_{ij}, \quad S_{\text{final}}(q) = S_{\text{atom}}(q) \cdot (1 + \lambda \cdot \text{Pref}(m))$$

where λ controls the influence of model preference and $\text{Pref}(m)$ is the cumulative preference score for model m . The SQL output with the highest S_{final} is shown to the user.

2.6 End-to-End Flow

On each query, the atom decomposer extracts x_i and a_j ; the agreement engine builds M from model outputs; the preference store contributes U from past user choices; the two are combined into the final alignment P ; and the query assembler picks the SQL output with the highest S_{final} . With no prior history, ATOMSQL relies entirely on model agreement. As the user provides feedback, U shifts and future results increasingly reflect their preferences.

3 DEMONSTRATION SCENARIO

We will demonstrate ATOMSQL using the *pets* database from the Spider benchmark, containing three relations (Student, Has_Pet, and Pets) with attributes such as StuID, Age, FName, and PetType. We guide the users through nine steps (annotated in Figure 1) impersonating Nicole, a university housing coordinator preparing a report on pet-friendly housing assignments. She needs to identify older students with exactly two pets to prioritize them for a new pet-friendly dorm wing, and asks ATOMSQL “*Show two older students who have two pets.*” demonstrating how atom-level confidence exposes ambiguity and how user feedback adapts future results.

Steps (A) & (B) (Selecting models and reviewing schema).

In (A), Nicole selects the NL-to-SQL models from the *Admin Panel*. The panel lists both commercial models (GPT-4o, Claude 3.5 Sonnet, Gemini 1.5 Pro) and open-source models (Llama 3 8B, DeepSeek-Coder 7B); in this scenario, SQLCoder 34B, T5SQL, and Llama 3 8B are toggled active. The panel also exposes two sliders that control how ATOMSQL balances its scoring: the *User Preference Value* slider sets β , which controls how strongly past user atom selections influence the alignment P , and the *Historical Preference Value* slider sets

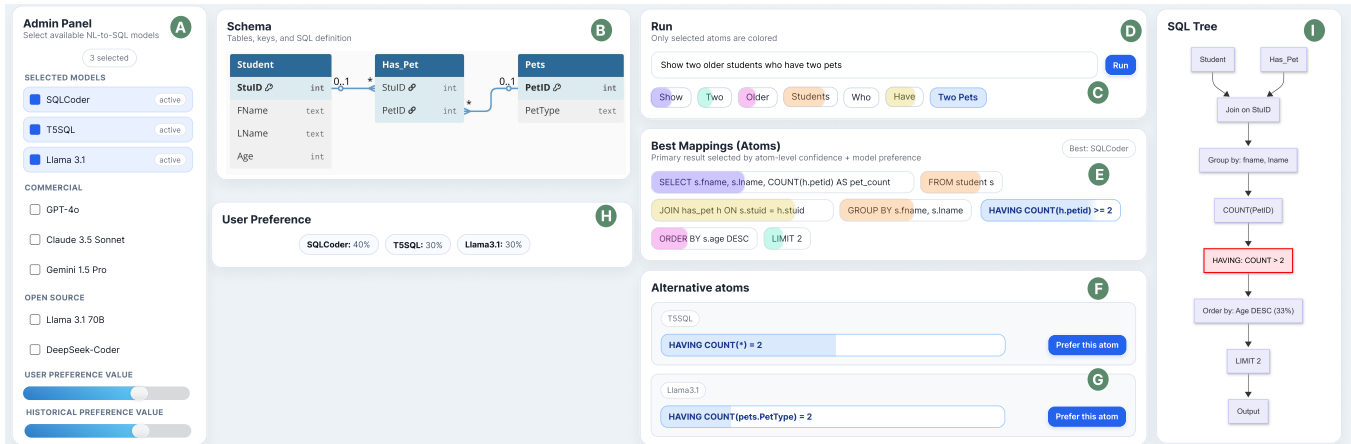


Figure 1: The demo interface: (A) select models, (B) review the schema, (C) enter the natural-language query, (D) run the selected models, (E) inspect the best atom mappings, (F) view alternative atoms, (G) choose a preferred atom, (H) update user preferences, and (I) view the SQL tree.

λ , which controls the weight given to historically preferred models when computing S_{final} . In (B), she reviews the *Schema* panel, which renders the database as an entity-relationship diagram showing *Student*, *Has_Pet*, and *Pets* with their columns and foreign-key join paths, making table structure and join relationships visible before issuing a query.

Steps (C) & (D) (Entering and running the query). In (C), Nicole types her natural-language question into the *Run* panel. In (D), she clicks *Run*, dispatching the query to all three active models simultaneously. The interface tokenizes the input into NL atom chips (*Show, Two, Older, Students, Who, Have, Two Pets*), highlighting the semantically significant phrases *ATOMSQL* will align to SQL. *ATOMSQL* collects each model’s generated SQL query, decomposes each query into clause-level atoms, and computes cross-model agreement scores.

Step (E) (Inspecting best mappings). In (E), *ATOMSQL* presents the primary SQL interpretation in the *Best Mappings (Atoms)* panel, with the banner indicating “Best: SQLCoder” since SQLCoder’s output achieved the highest alignment score S_{final} for this query. The assembled query is shown as a sequence of color-coded atom chips, where each color corresponds to a distinct NL atom: atoms mapped from the same NL phrase share a color, making it easy to trace which part of the question each SQL clause originates from.

Steps (F) & (G) (Viewing alternatives and selecting a preference). In (F), the *Alternative atoms* panel surfaces competing formulations for the *HAVING* atom: T5SQL proposes `HAVING COUNT(*) = 2` and Llama 3 8b proposes `HAVING COUNT(pets.PetType) = 2`, each with a *Prefer this atom* button that signals to *ATOMSQL* what Nicole meant by “two pets.” In (G), Nicole clicks *Prefer this atom* to select T5SQL’s formulation. This targeted correction replaces only the disputed atom without requiring her to rewrite the full SQL.

Step (H) (Updating model weights). The effect of Nicole’s selection is immediately visible in the *User Preference* panel: the initial weights (SQLCoder 34b 40%, T5SQL 30%, Llama 3 8b 30%) are updated so that T5SQL’s share grows and SQLCoder’s decreases, ensuring future rankings better reflect her demonstrated preference and allowing *ATOMSQL* to learn her intent over time.

Step (I) (Viewing the SQL tree). In (I), the *SQL Tree* panel renders the active query as a top-down execution tree, exposing the full hierarchical structure of the SQL rather than just its flat atom decomposition: *Student* joined with *Has_Pet* on *StuID*, grouped by *fname* and *lname*, aggregated with `COUNT(*)`, filtered by `HAVING COUNT > 2` (highlighted in red as the atom Nicole selected), ordered by `Age DESC`, and capped with `LIMIT 2`. This lets Nicole verify that join paths, aggregation scopes, and nesting are logically sound, going beyond what the flat atom chips in (E) convey.

Beyond the student-and-pets scenario, we will demonstrate *ATOMSQL* on other NL-to-SQL queries with different schemas and ambiguous questions, showcasing its ability to expose multiple reasonable interpretations at the atom level. The key takeaway from our demo is that users can inspect confidence, compare alternatives, and provide feedback on specific SQL atoms, while *ATOMSQL* adapts to incorporate their preferences in future queries.

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