
scBench: Evaluating AI Agents on Single-Cell RNA-seq Analysis

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Abstract

1 Introduction

Single-cell RNA sequencing (scRNA-seq) is a workhorse assay in research biology, providing transcriptional measurements at single-cell resolution to interrogate molecular state of tissues. As datasets grow in size and experimental usage broadens, drawing scientific conclusions increasingly depends on multi-step and resource-intensive computational methods that bridge techniques in statistics, high-dimensional data analysis, and programming. For many research groups, analysis—not sequencing—has become a rate-limiting step.

Agents—large language models (LLMs) that write code, invoke tools, and iterate toward a goal—have emerged with rapidly growing capabilities in software engineering and data analysis. However, agents for scRNA-seq remain both unreliable and underpowered, prone to scientific inaccuracies and hallucinations, and frequently fail to complete domain-specific analysis steps that depend on messy, real-world datasets.

Existing biology benchmarks emphasize recall, interpretation, or literature-style reasoning, and do not require empirical interaction with data or faithfully represent real-world analysis tasks. As a result, we lack a standard, deterministic yardstick for data-grounded scRNA-seq analysis.

We introduce **scBench**, a benchmark of 394 verifiable problems distilled from routine scRNA-seq workflows spanning six sequencing platforms and seven task categories. Each evaluation consists of a data snapshot, a natural-language task, and a deterministic grader. Across eight frontier models evaluated under a common harness, the best model reaches 52.8% accuracy, with large task- and platform-dependent performance swings. Together with SpatialBench for spatial transcriptomics, scBench provides a complementary diagnostic for measuring and improving agent competence on the two dominant transcriptional assays.

2 Results

2.1 scBench: Verifiable Problems from Real Workflows

scBench comprises 394 evaluations spanning six sequencing platforms and seven task categories (Table 1). Each evaluation pairs a data snapshot (often an AnnData .h5ad file) with a natural-language task prompt and a deterministic grader that scores the agent’s structured JSON output as a pass or fail. The benchmark mass concentrates in the analysis stages with the greatest dataset-specific variation: cell typing (118 evaluations, 30%) and differential expression (71, 18%) together account for nearly half the benchmark. Normalization (44) and QC (36) are smaller because these procedural steps admit fewer distinct problem formulations per dataset. Platform representation ranges from Illumina (85 evaluations) and MissionBio (81) to CSGenetics (42). ParseBio lacks QC evaluations because its vendor workflow omits explicit quality filtering, limiting cross-platform QC comparisons to five platforms.

The two axes, platform and task, enable stratified analysis of model performance. Task categories induce a gradient of accuracy: normalization applies standard transformations often with well-understood implementations; an agent need only identify the correct function call. Cell typing and differential expression require multi-step reasoning and contextual scientific reasoning: selecting marker genes, interpreting cluster identity, subsetting cells, choosing statistical tests, and ranking results. Platform diversity tests generalization beyond training-data familiarity. Chromium and Illumina dominate public repositories and tool documentation; MissionBio and ParseBio appear less frequently, use non-standard data structures and sport lesser known technical footguns. Models overfit on Scanpy tutorials without learning transferable analysis techniques should collapse on underrepresented platforms. Sections 2.3 and 2.4 quantify these effects.

2.2 Aggregate Model Performance

We evaluated eight frontier models from four providers (Table 2). Claude Opus 4.6 achieves the highest accuracy at 52.8% (95% CI: 48.3–57.2%), followed by Claude Opus 4.5 at 49.9% and GPT-5.2 at 45.2%. Claude Sonnet 4.5 reaches 44.2%, placing fourth despite being a smaller model. The bottom tier comprises GPT-5.1 (37.9%), Grok-4.1 (35.6%), Grok-4 (33.9%), and Gemini 2.5 Pro (29.2%).

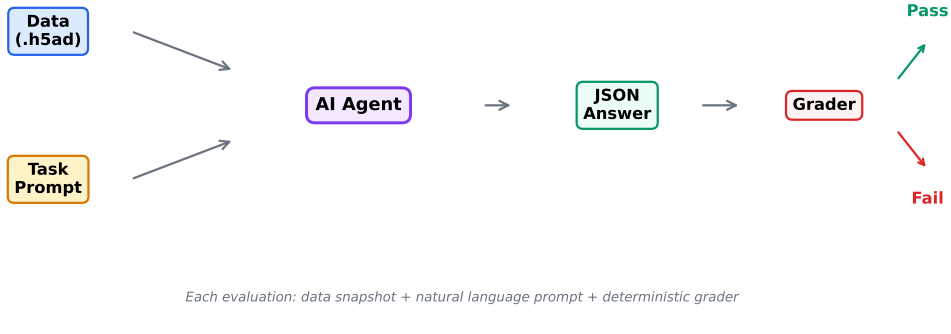


Figure 1: scBench evaluation structure.

Table 1: Number of evaluations by platform and task category.

	QC	Norm.	Dim. Red.	Clust.	Cell Typ.	Diff. Exp.	Traj.	Total
BD Rhapsody	6	11	14	7	13	10	—	61
Chromium	10	11	15	8	5	11	—	60
CSGenetics	4	5	7	5	20	1	—	42
Illumina	8	7	10	12	33	8	7	85
MissionBio	8	3	5	12	34	19	—	81
ParseBio	—	7	18	5	13	22	—	65
Total	36	44	69	49	118	71	7	394

The 23.6 percentage point spread between best and worst models exceeds SpatialBench’s 18.3 pp spread, indicating that scBench discriminates model capability despite the higher overall accuracy. Anthropic models occupy the top four positions, with both Opus variants and Sonnet outperforming all competitors. The confidence intervals for the top three models overlap, suggesting statistical equivalence at the aggregate level; stratified analysis (Sections 2.3–2.4) reveals where models diverge.

2.3 Task Category Analysis

Task categories exhibit a consistent difficulty gradient (Table 3, Figure 5). Normalization is easiest, with best-model accuracy of 83.8% and cross-model mean of 70.4%. QC follows at 55.3% mean. These procedural tasks—applying well-documented transformations—are approaching reliability. At the other extreme, differential expression is hardest: best-model accuracy of 41.4% (Opus 4.6) and cross-model mean of 27.0%. Cell typing (mean 34.9%) and clustering (mean 38.3%) occupy the middle. This gradient is nearly universal: seven of eight models follow the same difficulty ordering.

The hardest tasks are also the most discriminative. Differential expression shows a 27.7 pp spread between best and worst models (Opus 4.6 at 41.4% vs Gemini at 13.7%). Normalization shows a larger absolute spread (32.4 pp) but a lower coefficient of variation due to its high mean. Model capability differences concentrate in judgment-heavy stages—DE and cell typing—rather than procedural ones.

Rank inversions reveal task-specific strengths. Claude Sonnet 4.5 achieves near-parity with Opus 4.5 on normalization (82.9% vs 83.8%) but falls behind by 6 pp on differential expression. Opus 4.6 dominates judgment-heavy tasks while Opus 4.5 leads on procedural tasks. Grok-4 beats Grok-4.1 on clustering (34.8% vs 31.2%), demonstrating that version improvements do not translate uniformly across task types.

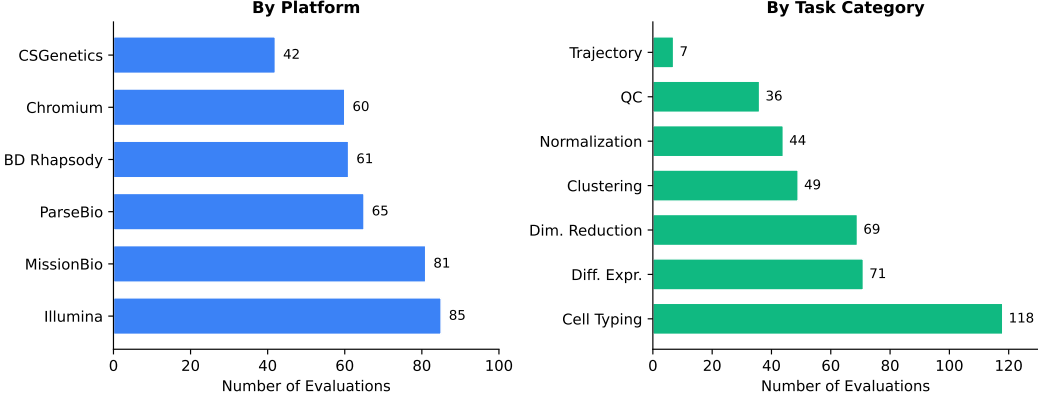


Figure 2: Distribution of 394 evaluations across platforms and task categories. Cell typing and differential expression dominate; ParseBio lacks QC evaluations.

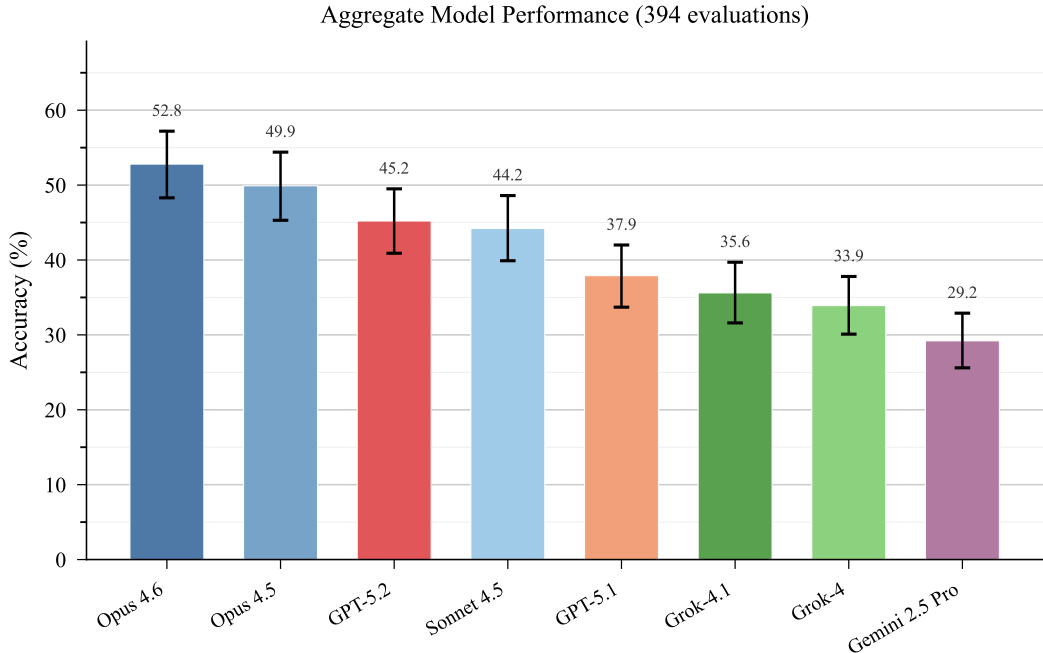


Figure 3: Aggregate accuracy of 8 frontier models on scBench (394 evaluations, 3 replicates each). Error bars show 95% confidence intervals computed via two-stage aggregation with the t -distribution.

2.4 Platform-Dependent Performance

Sequencing platform imposes a difficulty gradient comparable in magnitude to the model performance spread (Table 4, Figure 6). Per-platform mean accuracy ranges from 59.1% on CSGenetics to 26.4% on MissionBio—a 32.7 percentage-point gap that exceeds the 23.6-point spread between the best and worst models. Unlike the task gradient, where all models follow a consistent difficulty ordering, only two models (Opus 4.5 and GPT-5.2) follow the platform ordering exactly. The middle tier of platforms (BD Rhapsody, Illumina, Chromium) clusters within 3 percentage points, and models routinely swap rankings within this band. The extremes are more stable: CSGenetics is easiest for six of eight models, and MissionBio is hardest for all eight.

MissionBio is where the overall leaderboard breaks down. Cross-model coefficient of variation on MissionBio is 36.8%, versus 10.2% on Chromium—model choice matters 3.6 times more on

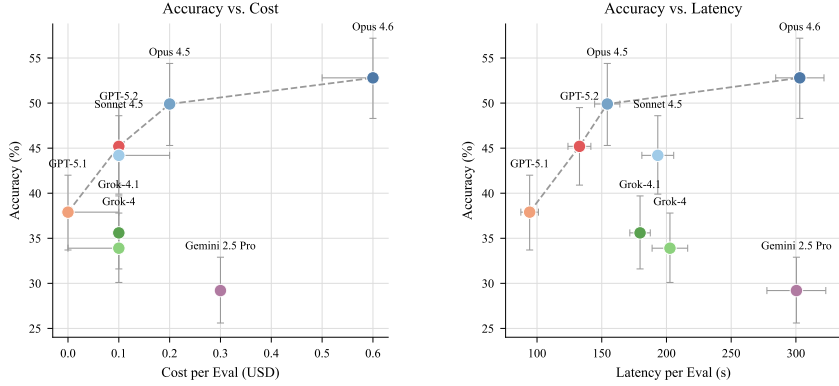


Figure 4: Accuracy–cost–latency tradeoffs across models. Marker size indicates mean latency per evaluation. GPT-5.2 achieves near-top accuracy at substantially lower cost than the Opus models.

Table 2: Overall model performance on scBench (394 evaluations, 3 replicates, mini-SWE-agent harness).

Model	Provider	Accuracy (%)	95% CI	Latency (s)
Claude Opus 4.6	Anthropic	52.8	(48.3, 57.2)	303
Claude Opus 4.5	Anthropic	49.9	(45.3, 54.4)	154
GPT-5.2	OpenAI	45.2	(40.9, 49.5)	133
Claude Sonnet 4.5	Anthropic	44.2	(39.9, 48.6)	193
GPT-5.1	OpenAI	37.9	(33.7, 42.0)	94
Grok-4.1	xAI	35.6	(31.6, 39.7)	180
Grok-4	xAI	33.9	(30.1, 37.8)	203
Gemini 2.5 Pro	Google	29.2	(25.6, 32.9)	300

the hardest platform. MissionBio does not merely lower scores; it inverts rankings. Grok-4 (sixth overall) beats GPT-5.2 (third overall) on MissionBio (24.7% vs 23.0%), and Sonnet 4.5 surpasses GPT-5.2 by 11.2 percentage points (34.2% vs 23.0%). The Anthropic models hold up on MissionBio (Opus 4.6 at 42.0%, Sonnet at 34.2%) while most competitors collapse. In contrast, Chromium shows the most compressed rankings, with all models falling between 33.0% and 51.7%.

Platform-specific leadership further complicates model selection. Opus 4.5 leads on four platforms (CSGenetics, BD Rhapsody) but GPT-5.2 leads on Illumina (54.5% vs 50.6%) and Opus 4.6 takes Chromium (51.7% vs 47.1%). On Chromium—the most common 10x Genomics platform—GPT-5.1 also beats Opus 4.5 (46.0% vs 47.1%), suggesting that OpenAI models have a specific advantage on the most documented platform. Gemini, ranked last overall, rises to fourth place on CSGenetics (52.4%), beating GPT-5.1, Grok-4.1, and Grok-4; easier platforms compress the bottom of the ranking.

Every model exhibits substantial platform brittleness. Gemini shows the largest swing: 42.1 percentage points between CSGenetics (52.4%) and MissionBio (10.3%). GPT-5.2 is nearly as variable (42.1 points, CSGenetics 65.1% to MissionBio 23.0%). Even Opus 4.5, the most consistent model, loses 39.1 points between its best and worst platforms (77.0% on CSGenetics, 37.9% on MissionBio). Grok-4 appears most stable with only a 15.8-point swing, but this stability reflects uniform mediocrity rather than genuine generalization. These platform effects likely reflect uneven coverage in training data: MissionBio’s multi-modal protein+RNA workflows and non-standard data structures are less represented in public documentation than 10x Chromium pipelines.

2.5 Comparison to Spatial Transcriptomics

scBench and SpatialBench together cover the two dominant single-cell modalities, enabling direct comparison of agent capabilities (Table 5). The top model reaches 52.8% on scBench versus 38.4% on SpatialBench—a 14.4 percentage-point advantage for scRNA-seq. This gap is consistent across the performance range: the bottom model scores 29.2% on scBench versus 20.1% on SpatialBench

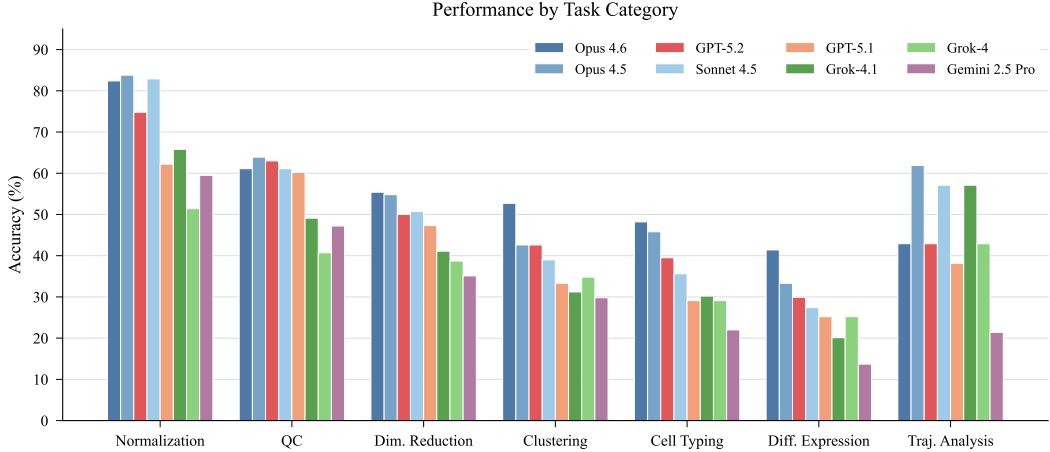


Figure 5: Accuracy (%) by model and task category. Tasks ordered by difficulty (normalization easiest, differential expression hardest). Error bars show 95% confidence intervals. The difficulty gradient is consistent across models.

Table 3: Accuracy (%) by task category with 95% CI. Best result per task in **bold**.

Model	QC	Norm.	Dim. Red.	Clust.	Cell Typ.	Diff. Expr.
Opus 4.6	61.1 (45.3, 76.9)	82.4 (71.8, 93.0)	55.4 (43.7, 67.1)	52.7 (40.6, 64.9)	48.2 (40.1, 56.2)	41.4 (31.7, 51.0)
Opus 4.5	63.9 (48.1, 79.7)	83.8 (73.1, 94.5)	54.8 (42.8, 66.7)	42.6 (29.5, 55.6)	45.8 (37.6, 53.9)	33.3 (23.8, 42.9)
GPT-5.2	63.0 (47.5, 78.4)	74.8 (62.1, 87.4)	50.0 (38.7, 61.3)	42.6 (30.0, 55.1)	39.5 (32.3, 46.8)	29.9 (21.0, 38.8)
Sonnet 4.5	61.1 (45.7, 76.5)	82.9 (71.6, 94.2)	50.7 (39.8, 61.7)	39.0 (26.9, 51.1)	35.6 (28.2, 43.0)	27.4 (18.5, 36.2)
GPT-5.1	60.2 (44.7, 75.6)	62.2 (48.3, 76.1)	47.3 (35.4, 59.2)	33.3 (21.3, 45.4)	29.1 (22.8, 35.4)	25.2 (17.1, 33.4)
Grok-4.1	49.1 (34.2, 64.0)	65.8 (52.3, 79.2)	41.1 (30.0, 52.1)	31.2 (20.5, 41.9)	30.2 (23.6, 36.8)	20.1 (12.3, 27.9)
Grok-4	40.7 (27.5, 54.0)	51.4 (39.5, 63.2)	38.7 (27.3, 50.1)	34.8 (24.6, 44.9)	29.1 (22.6, 35.6)	25.2 (17.0, 33.4)
Gemini	47.2 (34.2, 60.3)	59.5 (45.3, 73.6)	35.1 (24.8, 45.4)	29.8 (20.4, 39.2)	22.0 (16.3, 27.8)	13.7 (7.9, 19.4)

(+9.1 points). The performance spread is similar in both benchmarks (23.6 points for scBench, 18.3 for SpatialBench), indicating that both discriminate models comparably despite scRNA-seq sitting at a higher baseline. Model rankings are preserved at the extremes: Claude Opus leads both benchmarks and Gemini ranks last in both, suggesting a shared underlying “agent competence” axis that transfers across modalities.

The two benchmarks share structural regularities but diverge in failure modes. Normalization is the easiest category in both: best-model accuracy of 84% on scBench versus 76% on SpatialBench. Procedural transformations with well-documented implementations are tractable regardless of modality. The hardest categories differ: differential expression (41%) for scBench versus QC (22%) for SpatialBench, though the task taxonomies are not identical and cross-benchmark category comparisons are approximate. Both benchmarks exhibit strong platform effects, with 30–40 percentage-point swings depending on sequencing technology. The performance gap between benchmarks likely reflects training data availability: scRNA-seq has an order of magnitude more public datasets than spatial transcriptomics, and tools like Scanpy and Seurat dominate the ecosystem with extensive documentation. Spatial transcriptomics additionally requires coordinate-aware reasoning—neighborhood analysis, tissue architecture, spatial autocorrelation—that adds complexity beyond gene expression analysis.

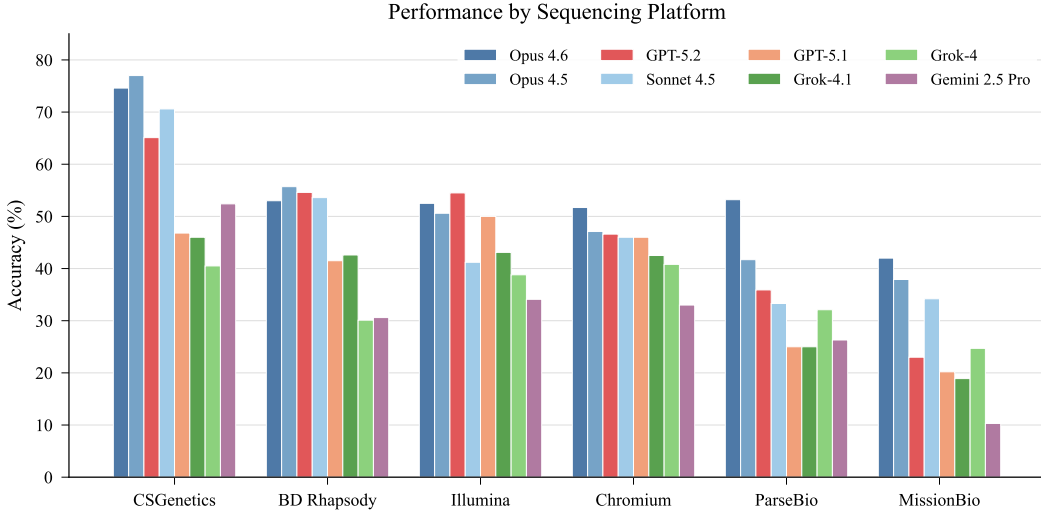


Figure 6: Accuracy (%) by sequencing platform. Platforms ordered by decreasing cross-model mean accuracy. Error bars show 95% confidence intervals.

Table 4: Accuracy (%) by sequencing platform with 95% CI. Best result per platform in **bold**.

Model	CSGenetics	BD Rhapsody	Illumina	Chromium	ParseBio	MissionBio
Opus 4.6	74.6 (62.5, 86.7)	53.0 (41.8, 64.2)	52.5 (42.8, 62.3)	51.7 (40.4, 62.9)	53.2 (41.3, 65.1)	42.0 (32.4, 51.6)
Opus 4.5	77.0 (65.9, 88.1)	55.7 (44.7, 66.7)	50.6 (40.4, 60.7)	47.1 (35.5, 58.7)	41.7 (28.2, 55.2)	37.9 (28.3, 47.4)
GPT-5.2	65.1 (53.2, 77.0)	54.6 (43.5, 65.8)	54.5 (45.9, 63.1)	46.6 (35.2, 57.9)	35.9 (23.3, 48.5)	23.0 (15.2, 30.9)
Sonnet 4.5	70.6 (57.8, 83.5)	53.6 (42.1, 65.0)	41.2 (31.9, 50.4)	46.0 (35.4, 56.6)	33.3 (22.3, 44.4)	34.2 (25.0, 43.3)
GPT-5.1	46.8 (33.6, 60.0)	41.5 (30.9, 52.2)	50.0 (41.6, 58.4)	46.0 (34.3, 57.7)	25.0 (13.5, 36.5)	20.2 (13.0, 27.3)
Grok-4.1	46.0 (33.5, 58.6)	42.6 (31.8, 53.4)	43.1 (34.1, 52.2)	42.5 (32.0, 53.0)	25.0 (13.4, 36.6)	18.9 (13.0, 24.9)
Grok-4	40.5 (27.6, 53.4)	30.1 (21.3, 38.8)	38.8 (31.0, 46.7)	40.8 (30.5, 51.1)	32.1 (19.6, 44.5)	24.7 (17.7, 31.7)
Gemini	52.4 (40.2, 64.6)	30.6 (21.2, 40.0)	34.1 (26.8, 41.4)	33.0 (22.8, 43.2)	26.3 (15.5, 37.1)	10.3 (5.5, 15.0)

Table 5: Comparison of scBench (scRNA-seq) and SpatialBench (spatial transcriptomics) under the mini-SWE-agent harness.

	scBench	SpatialBench
Number of evaluations	394	146
Number of platforms	6	5
Number of task categories	7	7
Top model accuracy	52.8%	38.4%
Bottom model accuracy	29.2%	20.1%
Top–bottom spread	23.6 pp	18.3 pp
Easiest task (best model)	Norm. 84%	Norm. 76%
Hardest task (best model)	DE 41%	QC 22%

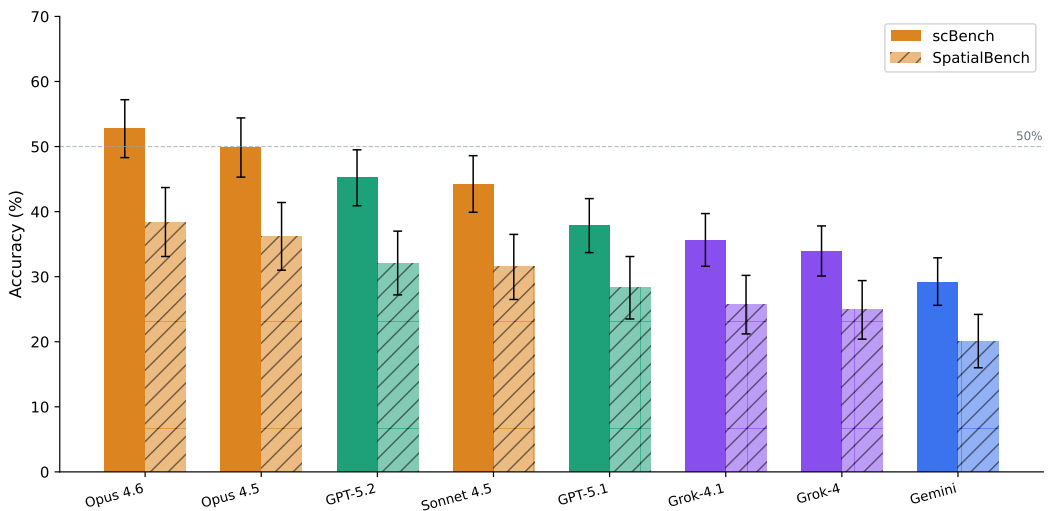


Figure 7: Model accuracy on scBench (solid bars) versus SpatialBench (hatched bars). scRNA-seq yields consistently higher accuracy across all models, but rankings are preserved: Claude Opus leads both benchmarks, Gemini ranks last. Error bars show 95% CIs.

3 Methods

3.1 Problem Construction

scBench is constructed from real scRNA-seq analysis workflows across six sequencing platforms: Chromium, BD Rhapsody, CSGenetics, Illumina, MissionBio, and ParseBio. Following Spatial-Bench, we identify analysis steps that satisfy three criteria: (1) the task arises in routine practice—a step that a working bioinformatician would perform as part of a standard pipeline (QC, normalization, HVG selection, clustering, annotation, or differential expression); (2) the answer requires empirical data interaction—it depends on the provided dataset and cannot be produced from textbook knowledge or memorized gene lists alone; and (3) the result is a verifiable quantitative artifact—a structured JSON output that can be graded deterministically by one of five grader families (Section 3.5).

Each candidate problem follows a five-stage construction pipeline. We first reproduce the target analysis step on the provided data using the published workflow. We then define the output artifact as an exact JSON schema with named fields and value types. Next, we select the grader family matching the output shape (e.g., `NumericTolerance` for cell counts, `DistributionComparison` for cell type proportions). We calibrate tolerances by running the analysis with multiple valid methods and parameter choices to establish the range of acceptable answers. Finally, we harden against shortcuts by removing precomputed embeddings, cached labels, and any fields that would allow the agent to read the answer without performing the intended computation.

Ground truth values are derived by re-running published pipelines from raw counts using author-specified parameters where available, then verified against domain understanding of expected biological results. When papers do not uniquely specify parameters (e.g., QC threshold not reported), we use standard defaults and widen tolerances to accept the resulting variation. Each problem is assigned an evaluation type—scientific, procedural, or observational (Section 3.3)—governing how aggressively tolerances must accommodate methodological variation. As a final quality-control step, we attempt to solve each problem by (a) reading `.obs` and `.uns` fields directly without computation, (b) answering from prior biological knowledge, and (c) running with alternative valid methods to verify tolerance coverage. Problems failing any of these checks are revised or removed.

3.2 Anatomy of a Problem

Each evaluation is a JSON specification with four agent-visible components and one internal component. The *data node* points to one or more `AnnData .h5ad` files containing the expression matrix, cell metadata (`.obs`), and gene annotations (`.var`); at runtime the harness downloads these files into an isolated workspace. The *task prompt* describes the analysis goal in natural language and specifies the exact JSON output format, including field names and value types. The *deterministic grader* defines the grader family and its configuration—ground truth values, tolerance parameters, and pass thresholds—that map the agent’s structured answer to pass/fail (Section 3.5). *Metadata* tags each problem by task category, evaluation type, sequencing platform, and computational complexity. A fifth component, *notes*, documents the solution approach, tolerance rationale, and known edge cases; notes are excluded from the agent’s context at the harness level and never appear at runtime.

Eval definitions are validated by a deterministic *linter* before entering the benchmark. The linter performs static schema validation, checking that required fields are present, grader configurations are well-formed (e.g., tolerance types are valid, thresholds are in range), and that the answer fields specified in the task prompt match what the grader expects. Evals that fail linting are blocked; evals with ambiguous tolerances or shortcut-prone structure are revised or removed during manual review.

3.3 Evaluation Types and Durability

Every evaluation is classified into one of three types that govern how aggressively tolerances must accommodate methodological variation.

Scientific. The prompt specifies a biological goal but leaves both the method and its parameters to the agent (e.g., “filter low-quality cells”). Because multiple QC thresholds, HVG selection methods, or clustering resolutions could defensibly be applied to the same data, tolerances must be wide

enough to accept all reasonable choices. Tight tolerances are used only when clean data causes valid methods to converge.

Procedural. The prompt names a specific method and leaves only parameter choices to the agent (e.g., “normalize using scran pooling”). Tolerances can be tighter than for scientific evaluations because the method is constrained.

Observational. The prompt asks the agent to interpret or report a property of the data (e.g., “which cell populations separate along PC1?”). Durability requirements are relaxed, and grading focuses on verifiability and anti-shortcut structure.

The distribution of evaluation types affects aggregate interpretation: a benchmark weighted toward procedural evaluations would yield higher scores because the method is specified, while a scientific-heavy benchmark tests judgment under ambiguity. Scientific evaluations carry wider tolerances on average than procedural evaluations, reflecting the greater methodological freedom.

3.4 Design Principles

Following SpatialBench, we apply three design criteria to every evaluation. The overarching rule is *specify what, not how*: tasks define the scientific goal and the exact output format, but do not prescribe a step-by-step method or parameters (with the exception of procedural evaluations, which name the method). The linter enforces structural compliance; manual review validates each criterion.

Verifiability. Each task specifies an exact JSON output format with named fields and value types, and is paired with a deterministic grader whose output shape matches the task (e.g., `NumericTolerance` for cell counts, `DistributionComparison` for cell type proportions). Success is automatically checkable with no subjective interpretation. Tasks that rely on subjective language (“interesting”, “meaningful”) without an operational definition are rejected. Importantly, omitting thresholds or algorithm names is acceptable and often desirable—it preserves anti-shortcut structure by forcing the agent to make data-driven choices.

Scientific durability. The intended answer must be stable across reasonable methodological choices, or tolerances must be wide enough to accept the resulting variation. Durability requirements scale with evaluation type (Section 3.3): scientific evaluations demand the widest tolerances, procedural evaluations can be tighter, and observational evaluations are the most relaxed. We specifically avoid two failure modes common in scRNA-seq benchmarking: random seed sensitivity (Leiden clustering is stochastic, so we do not test exact cluster counts) and library version artifacts (UMAP coordinates are arbitrary across versions, so we test biological interpretation rather than coordinates). Imprecision that a domain expert would resolve unambiguously (e.g., “filter low-quality cells” without specifying which metric) is not considered a durability failure; only ambiguity where reasonable interpretations yield materially different answers is flagged.

Anti-shortcut. The agent must load and analyze the provided dataset; prior knowledge alone is insufficient. During problem construction, we remove precomputed embeddings (`adata.obs["X_pca"]`, `adata.obs["X_umap"]`) when the evaluation tests computation, strip cached labels and summary statistics that would allow the agent to read the answer directly, and ensure that multiple-choice distractors are biologically plausible to prevent label leakage. That an answer is “just a number” does not make it guessable—dataset-specific quantities are not shortcuttable once precomputed fields are removed.

3.5 Graders

Each evaluation is paired with a deterministic grader that maps the agent’s structured JSON answer to pass/fail. We use five grader families; formal specifications are in Appendix C.

NumericTolerance. Validates numeric values such as cell counts, expression levels, and QC metrics. Supports four tolerance modes—absolute ($|x - x^*| \leq \epsilon$), relative ($|x - x^*| / |x^*| \leq \epsilon$), minimum ($x \geq x_{\min}$), and maximum ($x \leq x_{\max}$)—as well as asymmetric bounds. Multiple fields are checked

independently; all must pass. String values are coerced to floats; coercion failure counts as a field failure.

MultipleChoice. Validates discrete answers against one or more correct options. The agent’s response is trimmed and uppercased before comparison, making matching case-insensitive.

MarkerGenePrecisionRecall. Validates gene lists against canonical marker sets using recall@ K (fraction of canonical markers recovered) and precision@ K (fraction of returned genes that are canonical). Gene names are lowercased before comparison. Recall thresholds are set per evaluation (typically ≥ 0.50); precision thresholds default to ≥ 0.60 but are set to zero when the evaluation tests recall without penalizing novel DE genes. A per-cell-type mode supports multi-population differential expression by requiring a minimum recall for each cell type.

LabelSetJaccard. Validates unordered set predictions (e.g., predicted cell type labels) via the Jaccard index $J(A, B) = |A \cap B| / |A \cup B|$, with a default pass threshold of 0.90. Both missing and extra labels penalize the score equally. Labels are compared as-is without case normalization.

DistributionComparison. Validates multi-category proportions such as cell type distributions. Each ground-truth category is checked independently against an absolute tolerance (e.g., ± 5 percentage points); all categories must pass. Categories missing from the agent’s output fail automatically, while extra categories are ignored. Category names are lowercased before comparison. The all-must-pass rule ensures that agents cannot ignore rare cell types; tolerances are set wide enough to absorb reasonable per-category variation.

3.6 Agent Harness

We evaluate all models under mini-SWE-agent, an open-source harness that implements a simple action loop: the LLM generates a free-form response, the harness extracts the first fenced code block (delimited by markdown triple-backtick syntax), executes it in a local bash shell, and returns stdout/stderr to the model as the next observation. Each evaluation is capped at 100 action steps (LLM turn \rightarrow code extraction \rightarrow execution \rightarrow observation); if the agent exhausts the step budget without writing `eval_answer.json`, the evaluation scores zero.

The runtime environment provides scanpy, anndata, numpy, pandas, scipy, and matplotlib; all models share the same package versions. Network access is enabled, allowing agents to install additional packages if needed. Each evaluation runs in an isolated workspace: data files are symlinked from a local cache and the agent has read/write access only within that workspace. No GUI or interactive tools (Jupyter, plot display) are available.

Two timeout layers bound execution. An operation timeout of 300 seconds caps any individual bash command; an evaluation timeout of 600 seconds (configurable per evaluation) caps total wall-clock time via SIGALRM. On timeout or runtime crash, the harness grades whatever is in `eval_answer.json` at that point; if no answer file exists, the evaluation scores zero. There is no retry logic—each replicate is a single attempt. The harness records a complete trajectory for every run (conversation history, tool calls, and outputs), enabling post-hoc analysis of agent behavior.

3.7 Statistical Design

We follow the same two-stage aggregation used in SpatialBench. Each model–evaluation pair is run $K=3$ times. Replicates share the same prompt, data, and harness; the only source of variation is the model’s sampling nondeterminism (no explicit seed or temperature control). Each run receives a binary outcome from the grader, $s_{i,r} \in \{0, 1\}$.

In the first stage we compute the per-evaluation mean $\bar{s}_i = \frac{1}{K} \sum_r s_{i,r}$, yielding a value in $\{0, \frac{1}{3}, \frac{2}{3}, 1\}$. In the second stage we treat the $\{\bar{s}_i\}_{i=1}^n$ as independent observations and compute the aggregate accuracy $\hat{\mu} = \frac{1}{n} \sum_i \bar{s}_i$ with 95% confidence intervals via the t -distribution on $n-1$ degrees of freedom. All 392 evaluations are equally weighted; there is no upweighting by task category or platform. Per-evaluation means are approximately independent because evaluations use different datasets and different prompts; the shared model is the only common factor and is con-

stant within a model’s column. For stratified breakdowns (by task or platform), we apply the same procedure to the relevant subset, recomputing n , $\hat{\mu}$, and the corresponding t critical value.

4 Discussion

Agents for scRNA-seq occupy the same middle-capability regime that SpatialBench exposed for spatial transcriptomics: between knowing biology and writing code lies the harder skill of extracting biological insight from messy, real-world datasets. Across 394 verifiable problems with deterministic grading, the best model reaches 52.8% accuracy—leaving substantial room for progress. The 23.6-point spread across models shows that scBench discriminates capability, with significant task- and platform-dependent behavioral shifts. In practice, these results suggest that today’s agents can accelerate routine analysis but cannot yet be trusted to autonomously answer scientific questions without stringent verification of intermediate results and human oversight.

As with SpatialBench, the path forward appears to be a long tail of tractable engineering. Some runs fail for mundane reasons—instruction-following drift, output-format errors, or wasted steps chasing compliance rather than analysis—yet these account for a nontrivial fraction of zeros in a deterministic benchmark. scBench surfaces the same pattern: steps that demand contextual, often tacit judgment remain the least reliable. Normalization and QC are approaching reliability, while cell typing and differential expression require contextual decision-making and scientific reasoning currently outside the capabilities of frontier models. General-purpose coding skill is not sufficient; models need exposure to representative scRNA-seq workflows across diverse tissue and disease contexts, in addition to thorough understanding of technology-specific analysis techniques.

Platform-dependent performance swings often exceed task-dependent ones, suggesting that reliable agents will require platform-aware context, assay-specific tooling, and self-calibration heuristics rather than one-size-fits-all workflows. The MissionBio collapse and the Illumina-specific strength of certain models are not random noise—they reflect gaps in training data and the fragility of memorized pipelines when confronted with unfamiliar data structures.

scBench shares SpatialBench’s limitations: deterministic graders enable verifiable evaluation but necessarily discretize scientific judgment into automatically checkable outputs, and each evaluation snapshots a single workflow step rather than capturing long-horizon iteration where errors compound and thresholds are revisited. We hope scBench serves both as a measurement tool and a diagnostic lens—an evolving specification of scRNA-seq competence that supports test-driven development of agent systems whose behavior can improve through both model training and harness engineering.

5 Data and Code Availability

The benchmark framework—graders, linter, and agent harness—is available at <https://github.com/latchbio/latch-eval-tools>. Seven canonical evaluations (one per task category) are publicly released to demonstrate the benchmark format; the full 392-evaluation suite is withheld to prevent training contamination. Aggregate results (per-model, per-task, per-platform breakdowns) and complete run trajectories are available from the authors upon request. The evaluation framework supports custom agents via a pluggable `agent_function` interface, enabling direct comparison of new models against the published results.

A. Benchmark Inventory

scBench comprises 394 evaluations drawn from published analyses on six sequencing platforms. Each platform uses a distinct library preparation and capture technology, ensuring that the benchmark tests generalization across the scRNA-seq ecosystem rather than proficiency on a single data format.

- **BD Rhapsody**: microwell-based capture with targeted or whole-transcriptome panels. 61 evaluations.
- **Chromium** (10x Genomics): droplet-based capture. The most widely used scRNA-seq platform and the best represented in public datasets and documentation. 58 evaluations.
- **CSGenetics**: droplet-based capture with a proprietary barcoding chemistry. 42 evaluations.
- **Illumina**: plate-based single-nucleus RNA-seq (DRG tissue). 85 evaluations.
- **MissionBio** (Tapestri): targeted panel sequencing of DNA, RNA, and surface protein. Non-standard data structures and less common analysis tooling make this the hardest platform in the benchmark. 81 evaluations.
- **ParseBio**: split-pool combinatorial barcoding (no microfluidics). 65 evaluations.

Tissue types span PBMCs, tumor microenvironments (4T1 mammary carcinoma, CDX models), dorsal root ganglia (DRG), and hematopoietic samples. Table 1 shows the distribution of evaluations across platforms and task categories.

Table 6: Summary of scBench evaluations.

By Platform	Evals	By Task Category	Evals
BD Rhapsody	61	QC	36
Chromium	60	Normalization	44
CSGenetics	42	Dimensionality Reduction	69
Illumina	85	Clustering	49
MissionBio	81	Cell Typing	118
ParseBio	65	Differential Expression	71
		Trajectory Analysis	7
Total	394	Total	394

Grader Distribution

Evaluations use five grader families to assess agent outputs:

- **NumericTolerance**: QC metrics, cell counts, expression values, fold changes (most common)
- **MultipleChoice**: Biological interpretation, pattern identification
- **MarkerGenePrecisionRecall**: Marker discovery, differential expression gene lists
- **LabelSetJaccard**: Cell type prediction, cluster composition
- **DistributionComparison**: Cell type proportions, population distributions

Tissue Coverage

The benchmark covers four primary tissue/sample types:

- **PBMC** (BD Rhapsody, CSGenetics, ParseBio): 168 evaluations — T cell subtypes, monocyte populations, rare cell detection
- **Tumor microenvironment** (Chromium): 58 evaluations — 4T1 mammary carcinoma, CDX small-cell lung cancer, CAF subtypes
- **Dorsal root ganglia** (Illumina): 85 evaluations — neuron subclasses, satellite glial cells, age-related changes

- **Hematopoietic** (MissionBio): 81 evaluations — CCUS samples, clonal hierarchy, mutation burden

Complete Evaluation Inventory

Table 7 provides the complete list of all 392 evaluations organized by platform.

Table 7: Complete inventory of scBench evaluations.

Description	Platform	Task	Grader
Naive T cell marker comparison	BD Rhapsody	Cell Typ.	MCQ
Treg marker gene recall	BD Rhapsody	Cell Typ.	P@K
CD8 TEM vs naive classification	BD Rhapsody	Cell Typ.	MCQ
Effective subtype count	BD Rhapsody	Cell Typ.	Numeric
Baseline iNKT fraction	BD Rhapsody	Cell Typ.	Numeric
CD8 TEM trend contrast	BD Rhapsody	Cell Typ.	MCQ
Classical monocyte pattern	BD Rhapsody	Cell Typ.	MCQ
CD14 score separation	BD Rhapsody	Cell Typ.	Numeric
Proliferative lymphocyte rarity	BD Rhapsody	Cell Typ.	Numeric
Subtype stability under reclustering	BD Rhapsody	Cell Typ.	Numeric
Patient composition divergence	BD Rhapsody	Cell Typ.	Numeric
21-subtype distribution (v1)	BD Rhapsody	Cell Typ.	Dist
21-subtype distribution (v2)	BD Rhapsody	Cell Typ.	Dist
Marker program coverage	BD Rhapsody	Clust.	P@K
Cytotoxic program cluster	BD Rhapsody	Clust.	MCQ
Cluster count	BD Rhapsody	Clust.	Numeric
Program separation overlap	BD Rhapsody	Clust.	Numeric
Subtype expression shift	BD Rhapsody	Clust.	Numeric
S100A vs MHC enrichment	BD Rhapsody	Clust.	Numeric
Louvain resolution sweep	BD Rhapsody	Clust.	Numeric
Day 3 stress gene fraction	BD Rhapsody	DE	Numeric
CD4 TEM EGR1 log fold change	BD Rhapsody	DE	Numeric
CD4 TEM RGS1 log fold change	BD Rhapsody	DE	Numeric
CD14 monocyte TNFa log fold change	BD Rhapsody	DE	Numeric
IFITM3 temporal pattern	BD Rhapsody	DE	MCQ
IL1B temporal pattern	BD Rhapsody	DE	MCQ
FCER1G day 3 expression	BD Rhapsody	DE	MCQ
Adhesion gene return to baseline	BD Rhapsody	DE	MCQ
DE temporal pattern (09)	BD Rhapsody	DE	MCQ
DE temporal pattern (10)	BD Rhapsody	DE	MCQ
Dimensionality reduction (14 evals)	BD Rhapsody	Dim.Red.	Mixed
Normalization (11 evals)	BD Rhapsody	Norm.	Numeric
Quality control (6 evals)	BD Rhapsody	QC	Numeric
CAF subcluster cell typing (5 evals)	Chromium	Cell Typ.	Mixed
Tumor clustering (6 evals)	Chromium	Clust.	Mixed
Contractile CAF marker recovery	Chromium	DE	P@K
Differential expression (10 evals)	Chromium	DE	Mixed
Dimensionality reduction (15 evals)	Chromium	Dim.Red.	Mixed
Normalization (11 evals)	Chromium	Norm.	Numeric

Continued on next page

Table 7 – continued

Description	Platform	Task	Grader
Quality control (10 evals)	Chromium	QC	Numeric
PBMC cell type proportions	CSGenetics	Cell Typ.	Dist
T cell marker recovery	CSGenetics	Cell Typ.	P@K
T cell activation/exhaustion state	CSGenetics	Cell Typ.	MCQ
Cell typing (17 additional evals)	CSGenetics	Cell Typ.	Mixed
Clustering (5 evals)	CSGenetics	Clust.	Mixed
Differential expression (1 eval)	CSGenetics	DE	Numeric
Dimensionality reduction (7 evals)	CSGenetics	Dim.Red.	Mixed
Normalization (5 evals)	CSGenetics	Norm.	Numeric
Quality control (4 evals)	CSGenetics	QC	Numeric
Neuron subclass assignment	Illumina	Cell Typ.	Jaccard
Brain signature in DRG (adversarial)	Illumina	Cell Typ.	MCQ
Cell typing (31 additional evals)	Illumina	Cell Typ.	Mixed
Leiden cluster count	Illumina	Clust.	Numeric
Clustering (11 additional evals)	Illumina	Clust.	Mixed
Differential expression (15 evals)	Illumina	DE	Mixed
Dimensionality reduction (10 evals)	Illumina	Dim.Red.	Mixed
Normalization (7 evals)	Illumina	Norm.	Numeric
Quality control (8 evals)	Illumina	QC	Numeric
Cell type label set	MissionBio	Cell Typ.	Jaccard
Other cell fraction	MissionBio	Cell Typ.	Numeric
NK marker recovery (top 5)	MissionBio	Cell Typ.	P@K
Cell typing (19 additional evals)	MissionBio	Cell Typ.	Mixed
CCUS clonal typing (10 evals)	MissionBio	Cell Typ.	MCQ
Louvain cluster count	MissionBio	Clust.	Numeric
Clustering (11 additional evals)	MissionBio	Clust.	Mixed
Differential expression (19 evals)	MissionBio	DE	Mixed
Dimensionality reduction (5 evals)	MissionBio	Dim.Red.	Mixed
Normalization (3 evals)	MissionBio	Norm.	Mixed
Quality control (8 evals)	MissionBio	QC	Numeric
cDC2 annotation confusion	ParseBio	Cell Typ.	MCQ
Cell typing (12 additional evals)	ParseBio	Cell Typ.	Mixed
Clustering (5 evals)	ParseBio	Clust.	Mixed
IL-4 monocyte response	ParseBio	DE	Numeric
Differential expression (21 evals)	ParseBio	DE	Mixed
Dimensionality reduction (18 evals)	ParseBio	Dim.Red.	Mixed
Normalization (7 evals)	ParseBio	Norm.	Numeric

Grader abbreviations: MCQ = MultipleChoice, P@K = MarkerGenePrecisionRecall, Numeric = NumericTolerance, Jaccard = LabelSetJaccard, Dist = DistributionComparison. Full evaluation specifications are available in the benchmark repository.

B. Canonical Examples

We provide 2–3 representative evaluations from each platform to illustrate the benchmark format and grader diversity, with emphasis on downstream analysis tasks (cell typing, clustering, differential expression). For each we list the task category, grader type, and tolerance rationale.

BD Rhapsody

Cell Typing (MarkerGenePrecisionRecall). `bd_rhapsody_celltyping_02_treg....` The agent identifies marker genes for regulatory T cells from PBMC data. Canonical markers: *FOXP3*, *IL2RA*, *CTLA4*, *DUSP4*, *RGS1* (5 total). Pass: $\text{recall}@10 \geq 0.60$, $\text{precision} \geq 0$.

Clustering (NumericTolerance). `bd_rhapsody_clustering_03_count`. The agent clusters PBMC cells and reports the number of clusters. Ground truth: 12 clusters; tolerance ± 2 (absolute). The tolerance accommodates variation across resolution parameters and clustering algorithms.

Chromium

Cell Typing (LabelSetJaccard). `chromium_celltyping_03_caf_subcluster....` The agent subclusters cancer-associated fibroblasts (CAFs) from a 4T1 tumor dataset and identifies marker programs for each subcluster. Ground truth: 12 canonical markers including *Acta2*, *Col1a1*, *Mki67*, *Ly6c1*. Pass: $\text{Jaccard} \geq 0.60$.

Differential Expression (MarkerGenePrecisionRecall). `chromium_de_01....` The agent sub-clusters CAFs, identifies the contractile subcluster, and reports top 50 DE genes. Canonical markers include *Tpm1*, *Myl9*, *Tagln* (9 total). Pass: $\text{recall} \geq 0.67$.

Clustering (NumericTolerance). `chromium_cdx_sclc_heterogeneity....` The agent computes intra-cluster heterogeneity before and after cisplatin treatment in a CDX small-cell lung cancer model, reporting the fold change. Pass: $\text{fold change} \geq 1.0$ (minimum threshold).

CSGenetics

Cell Typing (DistributionComparison). `pbmc_cell_type_annotation_v1`. The agent annotates PBMC cells into five compartments (T cells, B cells, NK cells, Monocytes, Dendritic cells) and reports percentages. Ground truth: 59.0%/20.3%/5.0%/14.8%/0.7%; tolerance ± 5 pp per category.

Cell Typing (MarkerGenePrecisionRecall). `pbmc_t_cells_marker_recovery`. The agent identifies the top 20 marker genes for T cells. Canonical markers include *CD3D*, *CD3E*, *IL7R*, *TRAC*, *TRBC2* (10 total). Pass: $\text{recall} \geq 0.50$, $\text{precision} \geq 0$.

Cell Typing (MultipleChoice). `pbmc_tcell_dual_activation_exhaustion`. The agent interprets T cell states to determine which subpopulation shows both activation and exhaustion signatures. Correct answer: H. Requires understanding of T cell biology beyond marker lookup.

Illumina

Cell Typing (LabelSetJaccard). `snrna_anno_03_assign_neuron_subclasses....` The agent assigns neuron subclasses (NF, NP, PEP, TH) in DRG snRNA-seq data based on marker expression. Ground truth: 4 neuron subclasses. Pass: $\text{Jaccard} \geq 0.80$.

Clustering (NumericTolerance). `snrna_ic_11_leiden_cluster_and_report_n_clusters`. The agent applies Leiden clustering to DRG snRNA-seq data and reports the cluster count. Ground truth: 16 clusters; tolerance ± 3 (absolute).

Cell Typing (MultipleChoice). `snrna_anno_adv_forced_03....brain_region_bait`. An adversarial forced-choice evaluation: the agent is given DRG (peripheral nervous system) data but asked which brain-region-specific neuronal signature scores highest. The correct answer (F, Microglia_homeostatic) appears because of shared macrophage-like markers, not because microglia are present in DRG. Tests whether the agent can detect that brain signatures are biologically implausible in this tissue context.

MissionBio

Cell Typing (MarkerGenePrecisionRecall). `annotation_03_nk_marker_recovery_top5`. Using the Tapestri protein panel, the agent identifies the top 5 markers for NK cells. Canonical markers: *CD16*, *CD56*. Pass: recall ≥ 0.50 , precision ≥ 0.40 .

Cell Typing (MultipleChoice). `ccus_ct_09_highest_mutation_burden.....` Using the Tapestri multi-omic panel, the agent identifies which cell population carries the highest per-cell mutation burden. Correct answer: A. Requires integrating DNA variant calls with cell labels.

ParseBio

Cell Typing (MultipleChoice). `pbmc_cdc2_annotation_confusion`. The agent resolves an annotation ambiguity in a split-pool PBMC dataset by interpreting marker overlap between cDC2 and other myeloid populations. Correct answer: A.

Differential Expression (NumericTolerance). `parsebio_il4_monocyte_response`. The agent computes the log2 fold change of a target gene in monocytes under IL-4 stimulation. Ground truth: -1.25 ; pass if $\log_{2}FC \leq -1.1$ (maximum threshold, directional).

C. Grader Specification

The graders, linter, and harness are implemented in the open-source `latchbio/latch-eval-tools` repository.¹ Below we give formal specifications for each grader family.

NumericTolerance

Input: JSON object with one or more numeric fields. String values are coerced via `float()`; coercion failure counts as a field failure.

Tolerance modes (configured per field):

- Absolute: pass if $|x - x^*| \leq \epsilon$
- Relative: pass if $|x - x^*| / |x^*| \leq \epsilon$
- Minimum: pass if $x \geq x_{\min}$
- Maximum: pass if $x \leq x_{\max}$
- Asymmetric: pass if $x^* - \epsilon_{\text{lower}} \leq x \leq x^* + \epsilon_{\text{upper}}$

Multiple fields are checked independently; all must pass. Missing required fields fail. Extra keys are ignored.

MultipleChoice

Input: `{"answer": "A"}`.

Normalization: agent answer is trimmed and uppercased (`.strip().upper()`).

Pass criterion: agent answer is a member of the configured `correct_answers` list.

¹<https://github.com/latchbio/latch-eval-tools>

MarkerGenePrecisionRecall

Input: a gene list (flat mode) or a dictionary mapping cell types to gene lists (per-cell-type mode). Gene names are lowercased before comparison.

Flat mode. Let P be the agent's gene set and G the canonical marker set.

$$\text{precision}@K = \frac{|P \cap G|}{|P|}, \quad \text{recall}@K = \frac{|P \cap G|}{|G|}$$

Pass if precision $\geq \tau_p$ **and** recall $\geq \tau_r$. Defaults: $\tau_r = 0.50$, $\tau_p = 0.60$ (overridable; $\tau_p = 0$ disables precision penalty).

Per-cell-type mode. Recall is computed per cell type; a cell type passes if recall $\geq \text{min_recall_per_celltype}$. The evaluation passes if the count of passing cell types $\geq \text{min_celltypes_passing}$.

LabelSetJaccard

Input: a list of predicted labels. Labels are compared as-is (no case normalization).

Pass criterion: $J(A, B) = |A \cap B| / |A \cup B| \geq \tau$ (default $\tau = 0.90$). Missing and extra labels both reduce the Jaccard index.

DistributionComparison

Input: a dictionary mapping category names to percentages. Category names are lowercased.

Pass criterion: for each ground-truth category c , $|p_c^{\text{agent}} - p_c^{\text{gt}}| \leq \epsilon$ (default $\epsilon = 3.0$ pp). All ground-truth categories must pass. Missing categories fail; extra categories are ignored.

Failure Modes

All graders classify failures into four modes: (1) format error (missing or unparsable JSON), (2) missing field, (3) type error (coercion failure), (4) wrong value (out of tolerance). All yield score zero; the grader's reasoning field records which mode.