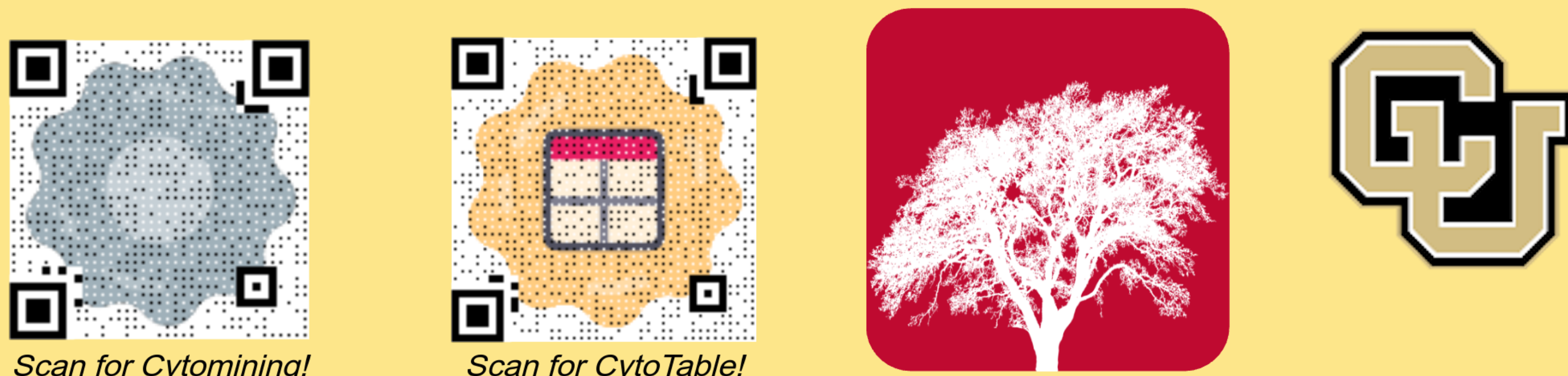


CytoTable: High performance and scalable single-cell morphology feature engineering

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Introduction

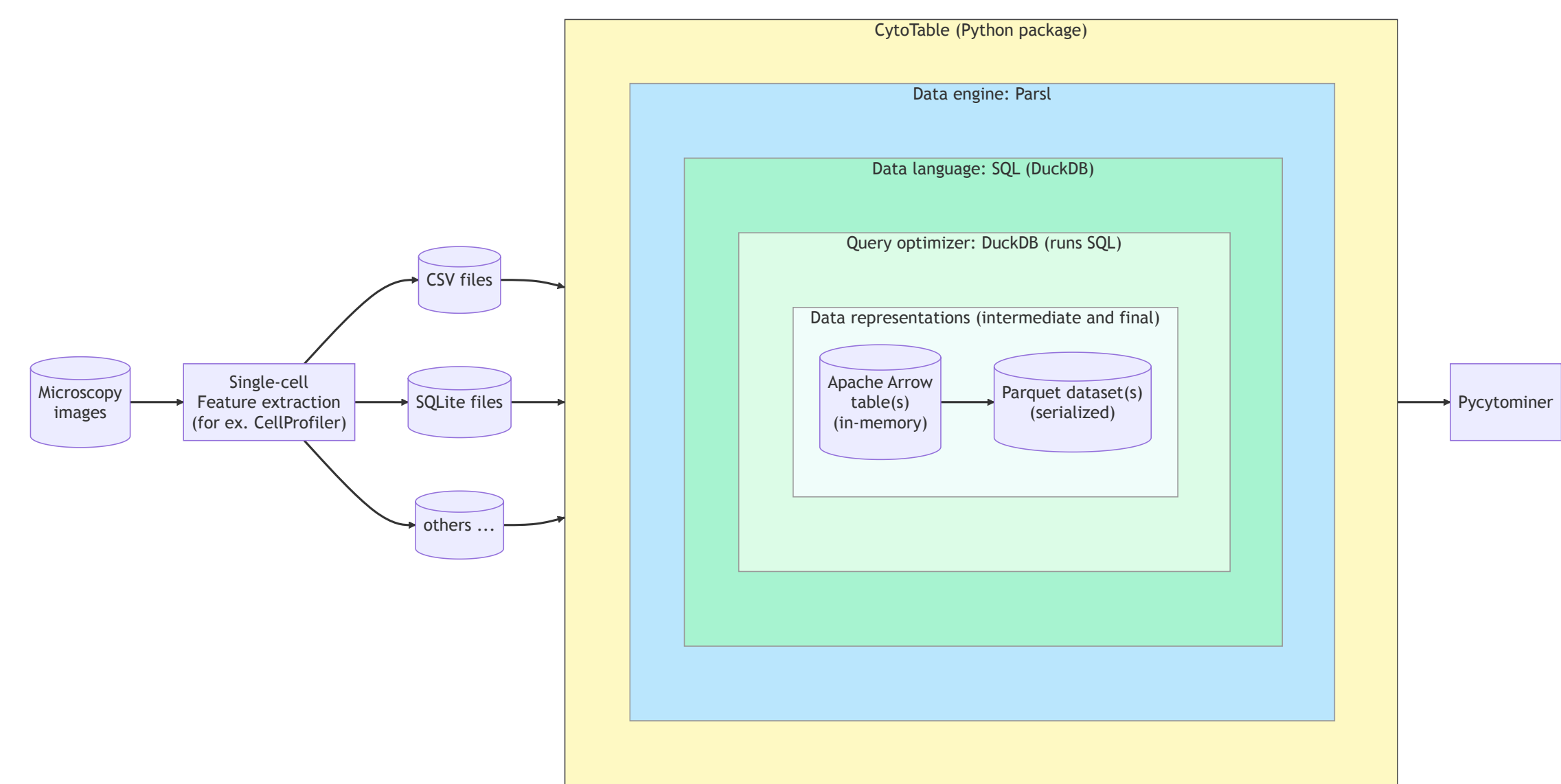


Figure 1. The flow of high-dimensional single-cell morphology data with relationship to CytoTable modular data stack components.

We are solving significant scalability and replicability challenges with high-dimensional single-cell morphology data (such as those extracted from CellProfiler[1]) by implementing novel and effective capabilities as a modular, portable, and cross-language single-cell data stack[2]: (a) language frontend: SQL (DuckDB[3]), (b) intermediate representation: Apache Arrow[4] and Apache Parquet[5], (c) query optimizer: DuckDB[3], (d) execution engine: Parsl[6] with Pythonic MapReduce design patterns[7], (e) execution runtime, Python package (PyPI, source)(Figure 1).

Microscopy feature data scalability

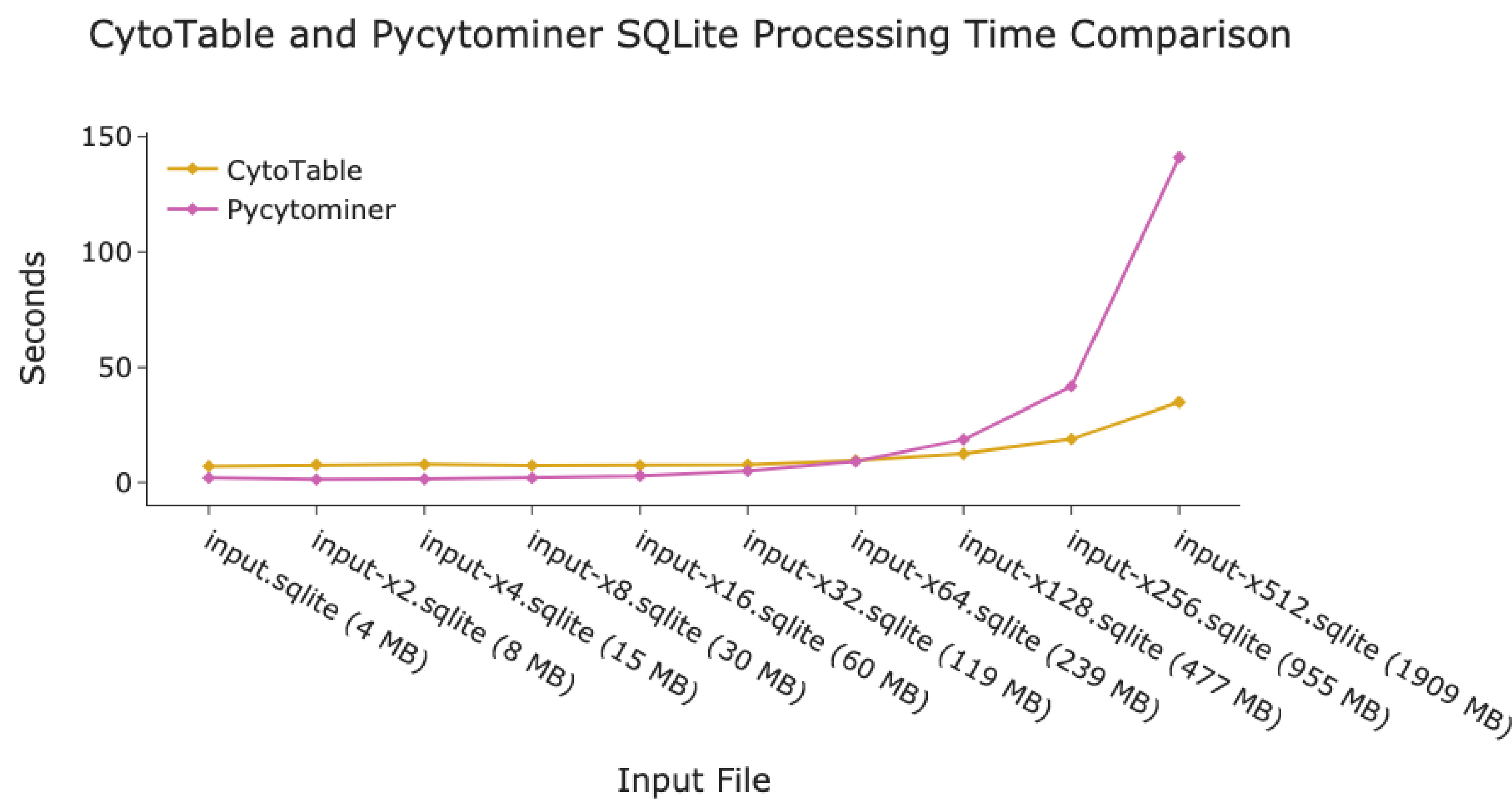


Figure 2. Comparing processing time duration for CytoTable and Pycytominer for various datasets of increasing size.

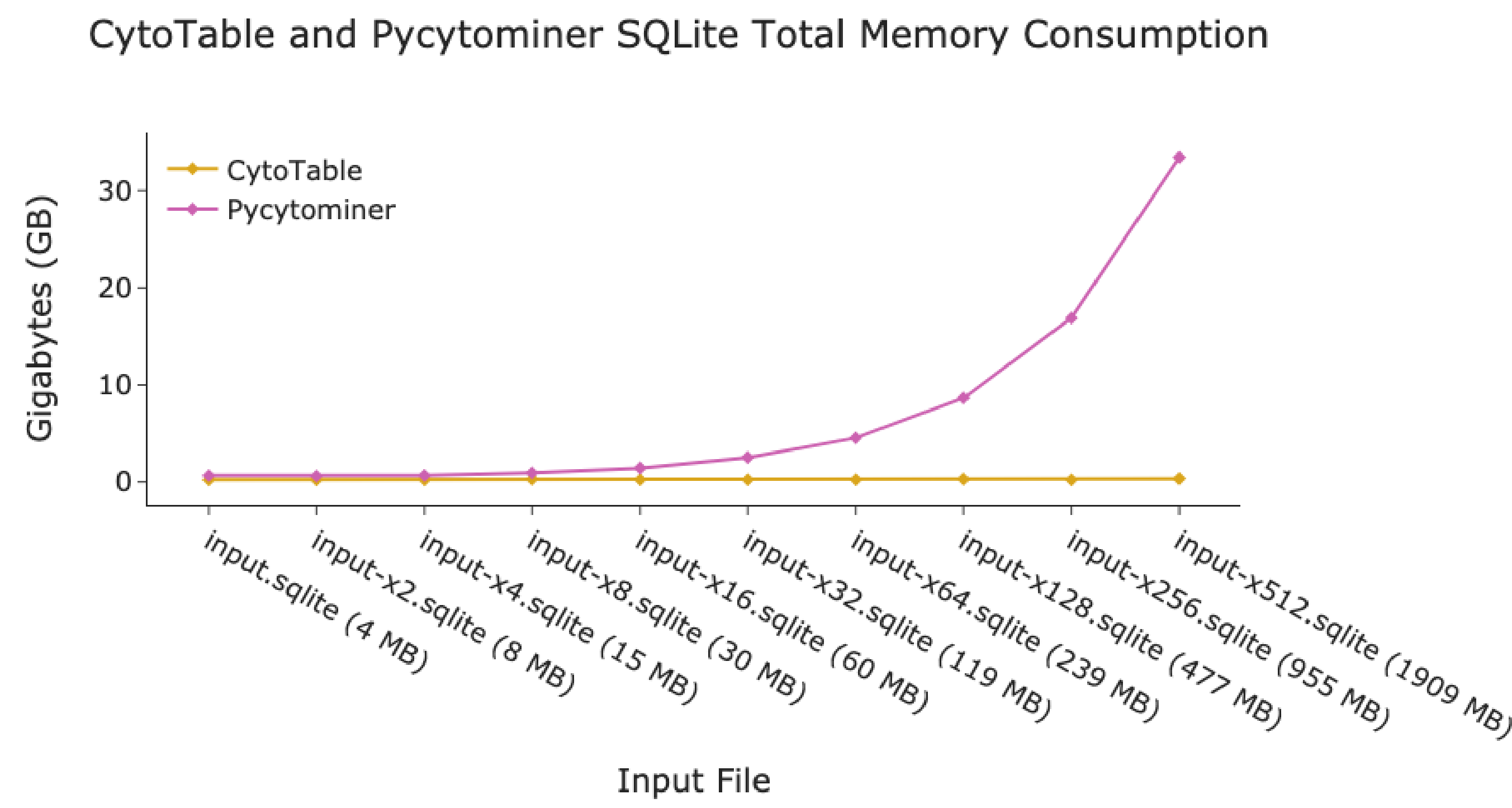


Figure 3. Comparing processing total memory consumption for CytoTable and Pycytominer for various datasets of increasing size.

CytoTable builds upon the shoulders of Pycytominer, helping to streamline the `SingleCells.merge_single_cells(...)` method. We decrease overall processing completion time (Figure 2) and memory consumption (Figure 3) for large amounts of data by leveraging composable data stack elements.

Empowering the Cytomining Ecosystem

Orchestration: CytoSnake

Authors: Erik Serrano, Jenna Tomkinson, Roshan Kern, Vince Rubinetti, Dave Bunten, Gregory P. Way

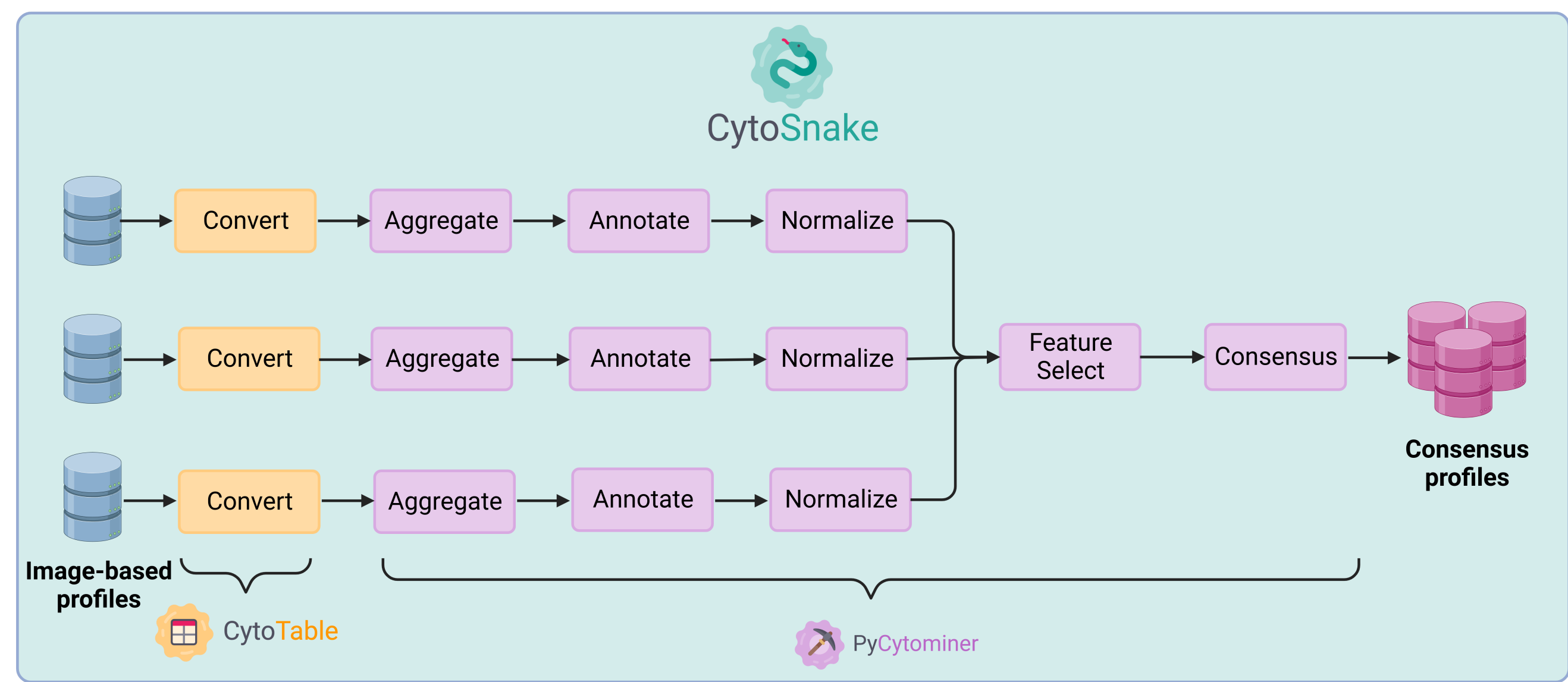


Figure 4. We orchestrate image-based profiling pipelines using CytoSnake. CytoTable is a critical tool for scalable orchestration.

CytoSnake is an innovative tool for orchestrating high-dimensional cell morphology data processing pipelines, including those which leverage CytoTable and other applied usecases.

Applied research: NF1 Schwann cell project

Authors: Jenna Tomkinson, Michelle Mattson-Hoss, Cameron Mattson, Herb Sarnoff, Gregory P. Way

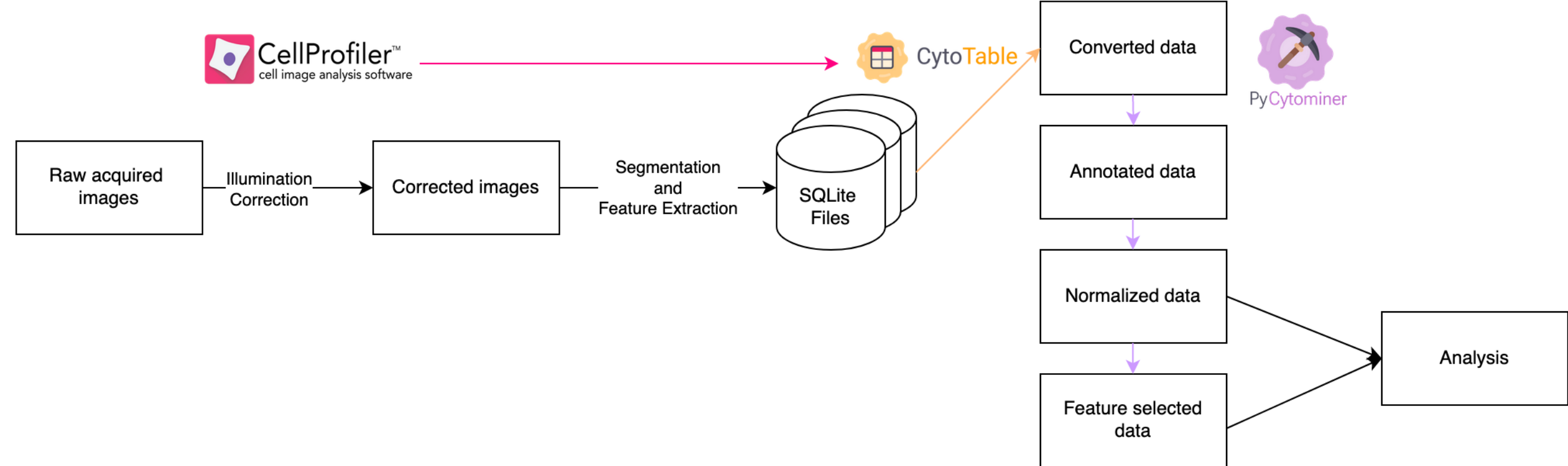


Figure 5. NF1 image-based profiling pipeline implementation details including CytoTable and Pycytominer.

We process Cell Painting images of Schwann cells with different NF1 *genotypes*. This application will increase understanding of cell morphology impacts of NF1 in Schwann cells and help prioritize rare disease treatments.

Applied research: Pyroptosis signature project

Authors: Michael J. Lippincott, Jenna Tomkinson, Interstellar Collaborators*, Masafumi Tsuboi, Carla Basualto-Alarcon, Gregory P. Way

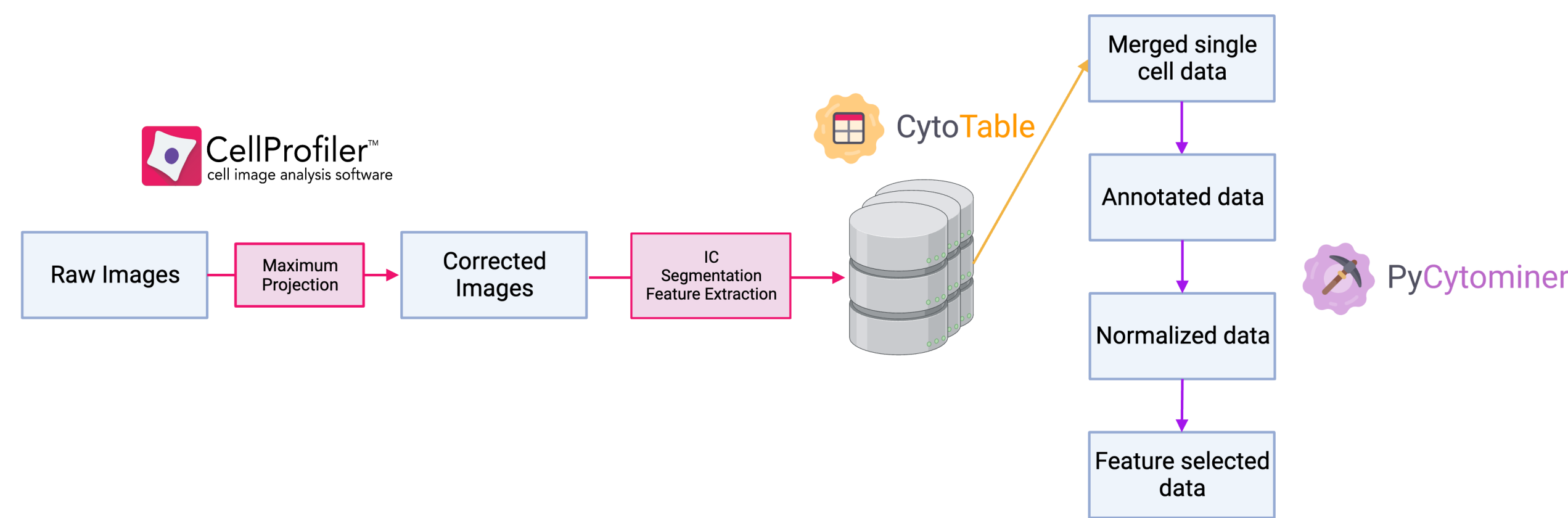


Figure 6. Image-based profiling pipeline of analyzing cells treated with various chemical agents to induce cell death. We are specifically focused on characterizing a cell morphology signature of pyroptosis that is distinct from apoptosis.

Identifying and characterizing pyroptosis signatures in cellular systems, aiding in the study of inflammatory cell death pathways as part of the Interstellar collaboration.

Applied research: CFReT project

Authors: Jenna Tomkinson, Erik Serrano, Gregory P. Way

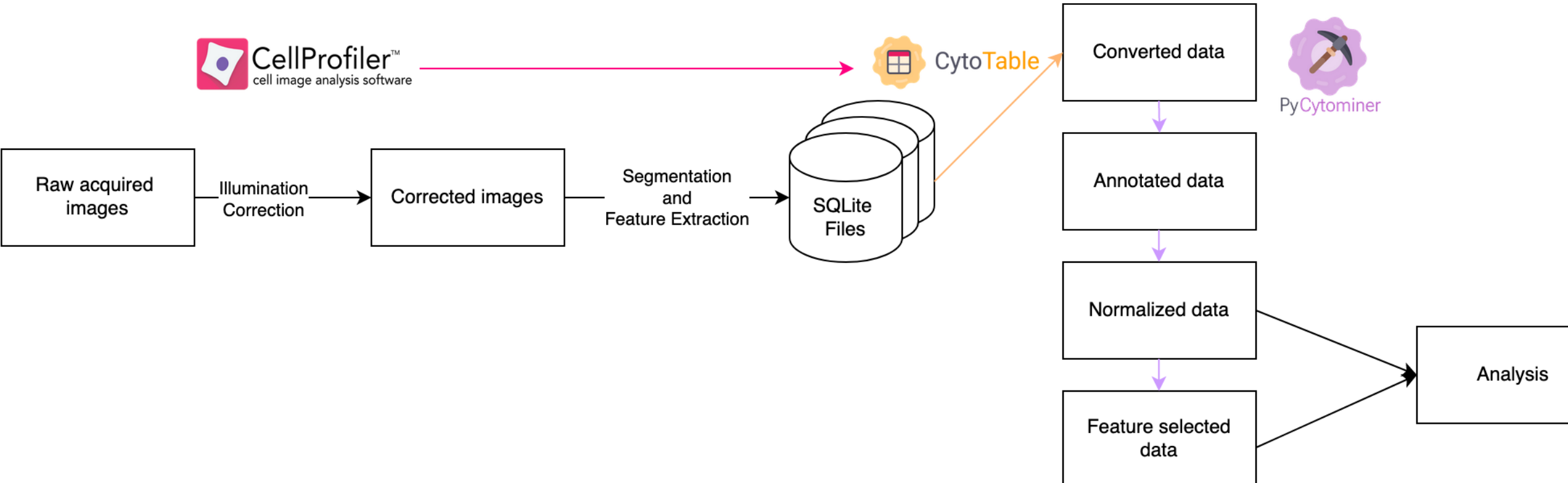


Figure 7. Image based profiling pipeline for characterizing cell morphology of failing cardiac fibroblasts through the Consortium for Fibrosis Research Translation (CFReT) project.

Image-based analysis of cardiac fibroblast datasets to uncover proprietary drug impact on reversing fibrosis.

Using CytoTable

```
import cytotable
result_file = cytotable.convert(
    source_path="path/to/feature-data",
    dest_path="destination/path.parquet",
    dest_datatype="parquet",
    preset="cellprofiler_csv",
)
```

Figure 8. Pythonic syntax for using CytoTable.

CytoTable may be installed from PyPI (`pip install cytotable`) and includes a Pythonic API which can be customized as needed or leverage existing presets (Figure 8). See the CytoTable documentation for more detail: <https://cytomining.github.io/CytoTable/>

Shape the future with us!



Figure 9. Cytomining Ecosystem software logos.

The Cytomining Ecosystem cultivates image-based profiling research through state-of-the-art software engineering and a vibrant, open-source community. We aim to provide the necessary infrastructure for a new era of bioinformatic innovation with high-throughput microscopy.

Interested in collaborating?
We welcome your input, contributions, and guidance!

Find us at <https://github.com/cytomining>.

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