

Automated Statistical and Machine Learning Platform

- ₂ for Biology Research
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Summary

This software provides a platform that combines machine learning and statistical analysis for chemical biology research, deployable as both a browser-based application and standalone desktop software. Researchers can upload CSV data, train Random Forest classification models with fully automated hyperparameter optimization, and perform comprehensive statistical tests through a unified interface requiring no programming expertise. The platform integrates data preprocessing, model training with version control, feature importance analysis, and interactive visualization (Figure 1), addressing the common workflow challenge of using multiple disconnected tools. Built with React 18.3 and TypeScript, it efficiently handles typical research datasets while allowing researchers to save and iteratively improve models through versioned training sessions. The complete implementation workflow from user interaction through model storage is illustrated in Figure 2.

Statement of Need

Biological and biomedical researchers routinely need to apply machine learning and statistics to experimental data, but existing tools create significant barriers. Powerful frameworks like scikit-learn (Pedregosa et al., 2011) and R (R Core Team, 2023) require programming expertise that many experimental scientists lack. Tools operate in isolation—researchers must manually transfer data between separate programs for statistical testing, machine learning, and visualization, reducing efficiency and introducing errors (Baker, 2016).

This software addresses these gaps by providing both web-based and desktop applications that combine Random Forest classification (Breiman, 2001) with standard statistical tests (t-tests, ANOVA, correlation) in one interface. The dual deployment model offers flexibility: researchers can use the browser version with no installation, or download the standalone desktop application for offline work and enhanced data privacy. Unlike Jupyter notebooks (Kluyver et al., 2016), it requires no coding knowledge. Unlike visual tools like Orange (Demšar et al., 2013), it includes comprehensive statistical testing alongside machine learning. The platform enables complete workflows—upload data, train models iteratively with version control, test hypotheses, generate visualizations—without switching applications or writing code.

Key Features and Implementation

The platform's modular interface organizes functionality into distinct tabs for data upload, model training, prediction, results visualization, and statistical analysis (Figure 1). This



- workflow-oriented design guides users through the complete analysis pipeline while maintaining
- access to all features.

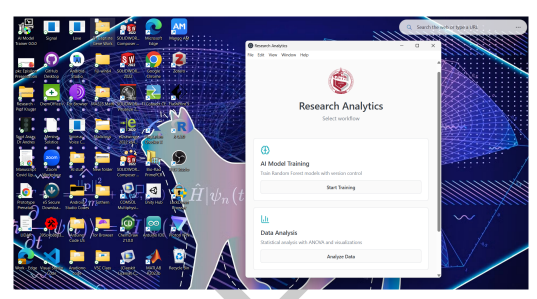


Figure 1: Interface dashboard showing the main analysis modules.

Architecture and Core Technologies

- The application is built with React 18.3 and TypeScript, leveraging Vite for optimized production
- builds and Electron for desktop packaging. The implementation follows a modular component 44
- architecture that separates concerns across data processing, model training, statistical analysis,
- and visualization layers. Core dependencies include ml-random-forest (v2.1) for machine
- learning algorithms, papaparse (v5.5) for robust CSV parsing, and recharts (v2.15) for 47
- SVG-based interactive visualizations. All computation occurs client-side, eliminating server
- dependencies and ensuring data privacy. The desktop application packages the same codebase
- for Windows, macOS, and Linux platforms.

Data Upload and Preprocessing

- The platform supports CSV file upload through drag-and-drop or file browser interfaces. Upon
- upload, the system performs automatic file structure detection and displays an interactive
- preview table showing the first 100 rows. Summary statistics (mean, median, standard deviation,
- quartiles, min/max) are computed for all numerical columns. Data validation identifies missing
- values, offering users options for row deletion or mean/median imputation. Preprocessing
- capabilities include z-score normalization, min-max scaling to [0,1], and automatic integer
- encoding of categorical variables. Column type detection distinguishes between numerical, 58
- categorical, and target variables, with manual override options.

Machine Learning Pipeline

- The platform implements Random Forest classification (Breiman, 2001), widely used for
- chemical property prediction and QSAR modeling (Svetnik et al., 2003). Figure 2 illustrates
- the complete implementation workflow from initial data upload through final model storage, 63
- showing how user interactions flow through data preprocessing, automated hyperparameter
- optimization, model training, evaluation, and version management.



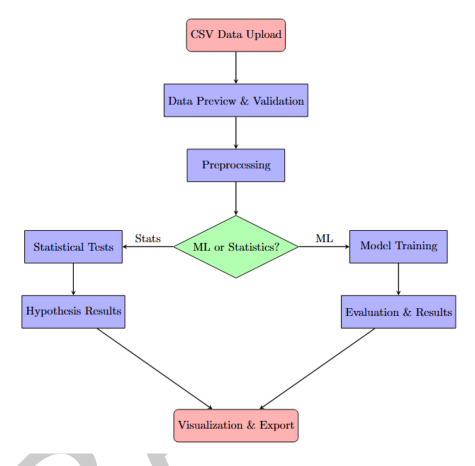


Figure 2: Implementation workflow from data upload through model storage.

- Recognizing that most researchers lack expertise in hyperparameter tuning, the system automat-
- 67 ically optimizes Random Forest parameters based on dataset characteristics. The optimization
- algorithm adjusts the number of trees (range: 10-500), maximum tree depth, and minimum
- samples per split according to dataset size and feature dimensionality, eliminating the need for
- manual configuration. Training executes asynchronously with real-time progress indicators to
- maintain interface responsiveness.
- $_{72}$ The system performs stratified 80/20 train-test splitting to preserve class distribution, crucial for
- imbalanced chemical datasets. Post-training, the interface displays comprehensive performance
- metrics including accuracy, precision, recall, F1-score, and interactive confusion matrices.
- 75 Feature importance scores computed via mean decrease in impurity reveal which molecular
- descriptors most influence classification, supporting interpretable model analysis.
- 77 Trained models persist in browser local storage or local file system (desktop version) with
- comprehensive version control. Researchers can save multiple model versions, each tagged
- 79 with training timestamp, dataset characteristics, and performance metrics. This versioning
- 80 system enables iterative model refinement—users can load previous versions, add new training
- 81 data, and create improved versions while maintaining the training history. Models export as
- ⁸² JSON files for deployment, sharing, or backup purposes.

Statistical Analysis Tools

- 84 The platform provides both parametric and non-parametric statistical tests for hypothesis
- testing and exploratory analysis. For comparing group means, Welch's t-test (Welch, 1947)



- $_{86}$ handles unequal variances, while the Mann-Whitney U test offers a distribution-free alternative
- 87 for non-normal data. One-way ANOVA enables multi-group comparisons. Correlation analysis
- includes Pearson's coefficient (Pearson, 1895) for linear relationships and Spearman's rank
- 89 correlation for monotonic associations.
- ₉₀ All statistical tests output comprehensive reports including p-values, effect sizes (Cohen's d. r).
- and 95% confidence intervals. The interface provides contextual guidance on assumption check-
- 92 ing (normality, homoscedasticity) and appropriate test selection based on data characteristics.
- Visual diagnostics include Q-Q plots and residual plots for assumption validation.

94 Interactive Visualization

The visualization module generates publication-quality SVG charts using Recharts, including scatter plots with regression lines, histograms with kernel density overlays, box plots with outlier detection, feature importance bar charts, confusion matrices with color-coded cells, and correlation heatmaps. All visualizations support interactive features: hover tooltips displaying precise values, zoom/pan controls for dense datasets, legend toggling for multi-series plots, and responsive sizing for different display resolutions. Charts export as high-resolution PNG images suitable for manuscript figures. The color schemes follow accessibility guidelines for colorblind users.

103 User Interface Design

As shown in Figure 1, the interface employs tab-based navigation mirroring typical analysis workflows: Data Upload \rightarrow Model Training \rightarrow Prediction \rightarrow Results \rightarrow Statistical Analysis. 105 Tabs remain disabled until prerequisite steps complete, preventing workflow errors. Form inputs 106 include real-time validation with error messages and tooltip hints. The responsive design 107 adapts to desktop and tablet viewports. Model management features include persistent storage (browser local storage with 5MB capacity or unlimited desktop file system), version control with 109 timestamp metadata and performance tracking, and JSON import/export for model sharing 110 and backup. The version history interface allows researchers to compare model performance 111 across iterations and load any previous version for continued training or deployment. 112

Research Applications

The platform supports prediction based on work data and trained AI, and exploratory data analysis in chemical biology. Typical applications include QSAR modeling, compound screening, and comparative analysis of experimental conditions. The integrated workflow reduces analysis time and technical barriers for laboratory researchers.

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References

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Abadi, M., Barham, P., Chen, J., Chen, Z., Davis, A., Dean, J., Devin, M., Ghemawat, S., Irving, G., Isard, M., & others. (2016). TensorFlow: A system for large-scale machine learning. *Proceedings of the 12th USENIX Symposium on Operating Systems Design and Implementation (OSDI)*, 265–283.



- Baker, M. (2016). 1,500 scientists lift the lid on reproducibility. *Nature*, *533*, 452–454. https://doi.org/10.1038/533452a
- Bergstra, J., & Bengio, Y. (2012). Random search for hyper-parameter optimization. *Journal* of Machine Learning Research, 13, 281–305.
- Breiman, L. (2001). Random forests. *Machine Learning*, 45(1), 5–32. https://doi.org/10. 1023/A:1010933404324
- ¹³⁴ Cortes, C., & Vapnik, V. (1995). Support-vector networks. *Machine Learning*, 20(3), 273–297. https://doi.org/10.1007/BF00994018
- Darnag, R., Minaoui, B., Glorennec, P. Y., Fakri, A., Zahrae, O., & Mourchid, M. (2010).

 QSAR studies of HEPT derivatives using support vector machines and neural networks.

 QSAR & Combinatorial Science, 29(5), 567–577. https://doi.org/10.1002/qsar.200960055
- Demšar, J., Curk, T., Erjavec, A., Gorup, Č., Hočevar, T., Milutinovič, M., Možina, M., Polajnar, M., Toplak, M., Starič, A., & others. (2013). Orange: Data mining toolbox in python. *Journal of Machine Learning Research*, *14*, 2349–2353.
- Friedman, J. H. (2001). Greedy function approximation: A gradient boosting machine. *Annals* of Statistics, 29(5), 1189–1232. https://doi.org/10.1214/aos/1013203451
- Hastie, T., Tibshirani, R., & Friedman, J. (2009). The elements of statistical learning:

 Data mining, inference, and prediction (2nd ed.). Springer. https://doi.org/10.1007/978-0-387-84858-7
- Kluyver, T., Ragan-Kelley, B., Pérez, F., Granger, B., Bussonnier, M., Frederic, J., Kelley, K., Hamrick, J., Grout, J., Corlay, S., & others. (2016). Jupyter notebooks—a publishing format for reproducible computational workflows. In F. Loizides & B. Schmidt (Eds.), Positioning and power in academic publishing: Players, agents and agendas (pp. 87–90). IOS Press. https://doi.org/10.3233/978-1-61499-649-1-87
- Mann, H. B., & Whitney, D. R. (1947). On a test of whether one of two random variables is stochastically larger than the other. *Annals of Mathematical Statistics*, 18(1), 50–60. https://doi.org/10.1214/aoms/1177730491
- Murphy, K. P. (2012). *Machine learning: A probabilistic perspective*. MIT Press. ISBN: 978-0262018029
- Pearson, K. (1895). Notes on regression and inheritance in the case of two parents. *Proceedings* of the Royal Society of London, 58, 240–242.
- Pedregosa, F., Varoquaux, G., Gramfort, A., Michel, V., Thirion, B., Grisel, O., Blondel, M.,
 Prettenhofer, P., Weiss, R., Dubourg, V., Vanderplas, J., Passos, A., Cournapeau, D.,
 Brucher, M., Perrot, M., & Duchesnay, E. (2011). Scikit-learn: Machine learning in python.
 Journal of Machine Learning Research, 12, 2825–2830.
- Perkel, J. M. (2021). Ten simple rules for writing and sharing computational analyses in jupyter notebooks. *PLOS Computational Biology*, *17*(7), e1008993. https://doi.org/10.1371/journal.pcbi.1008993
- R Core Team. (2023). *R: A language and environment for statistical computing*. R Foundation for Statistical Computing. https://www.R-project.org/
- Spearman, C. (1904). The proof and measurement of association between two things. *American Journal of Psychology*, 15(1), 72–101. https://doi.org/10.2307/1412159
- Svetnik, V., Liaw, A., Tong, C., Culberson, J. C., Sheridan, R. P., & Feuston, B. P. (2003).
 Random forest: A classification and regression tool for compound classification and QSAR modeling. *Journal of Chemical Information and Computer Sciences*, 43(6), 1947–1958. https://doi.org/10.1021/ci034160g



Van der Maaten, L., & Hinton, G. (2008). Visualizing data using t-SNE. *Journal of Machine Learning Research*, *9*, 2579–2605.

Virtanen, P., Gommers, R., Oliphant, T. E., Haberland, M., Reddy, T., Cournapeau, D.,
Burovski, E., Peterson, P., Weckesser, W., Bright, J., & others. (2020). SciPy 1.0:
Fundamental algorithms for scientific computing in python. *Nature Methods*, 17, 261–272.
https://doi.org/10.1038/s41592-019-0686-2

Welch, B. L. (1947). The generalization of student's problem when several different population variances are involved. *Biometrika*, *34*(1-2), 28–35. https://doi.org/10.1093/biomet/34. 1-2.28

