quicR: An R Library for Streamlined Data Handling of Real-Time Quaking Induced Conversion Assays

Gage R. Rowden^{a,b,c,*}, Peter A. Larsen^{a,b,c}

^aDepartment of Veterinary and Biomedical Sciences, University of Minnesota, USA. ^bMinnesota Center for Prion Research and Outreach, University of Minnesota, USA. ^cPriogen Corp., USA.

Abstract

Real-time quaking induced conversion (RT-QuIC) has become a valuable diagnostic tool for protein misfolding disorders such as Creutzfeldt-Jakob disease and Parkinson's disease. Given that the technology is relatively new, academic and industry standards for quality filtering data and high throughput analysis of results have yet to be fully established. The open source R library, quicR, was developed to provide a standardized approach to RT-QuIC data analysis. quicR provides functions, which can be easily integrated into existing R workflows, for data curation, analysis, and visualization.

Keywords: Chronic Wasting Disease, Creutzfeldt-Jakob Disease, Nano-QuIC, RT-QuIC, prion diagnostics, Parkinson's Disease, seed amplification assay

Metadata

Nr.	Code metadata description	Metadata
C1	Current code version	V2.1.0
C2	Permanent link to code/repository used for this code version	https://github.com/gage1145/quicR/releases/tag/v2.1.0
C3	Permanent link to Reproducible Capsule	https://cran.r-project.org/web/packages/quicR
C4	Legal Code License	GPL-3
C5	Code versioning system used	Git
C6	Software code languages, tools, and services used	R
C7	Compilation requirements, operating environments & dependencies	R (>=4.1.0)
C8	If available Link to developer documentation/manual	https://cran.r-project.org/web/ packages/quicR/quicR.pdf
C9	Support email for questions	rowde002@umn.edu

^{*}Corresponding author.

¹E-mail address: rowde002@umn.edu

1. Motivation and significance

Real-time quaking induced conversion (RT-QuIC) is within the family of seed amplification assays (SAAs), similiar to protein misfolding cyclic amplification (PMCA), and has garnered significant attention for its ultra-sensitive detection of misfolded protein aggregates [1, 2]. The assay works by converting a recombinant protein substrate into an amyloid aggregate in the presence of a misfolded seed [1, 3, 4, 5, 6, 7, 8, 9, 10]. The assay's sensitivity and specificity make RT-QuIC a promising tool for diagnosing diseases such as prion disorders and other protein misfolding pathologies [11, 12, 13, 14]. However, the relatively recent development and novelty of the assay have left a gap in widely accepted academic and industry standards for data analysis and interpretation [15].

To address this gap, we introduce **quicR**, an open-source library, developed in R [16], dedicated to the cleaning, analysis, and visualization of RT-QuIC data. By consolidating key metrics and providing robust analytical tools, **quicR** aims to standardize the analysis pipeline and foster reproducibility within the field of quaking induced assays including related assays such as Nano-QuIC [17] and Micro-QuIC [18]. **quicR** is designed with both researchers and diagnosticians in mind, providing a user-friendly interface that integrates seamlessly with existing R workflows.

While universal diagnostic criteria for RT-QuIC have yet to be established, certain analytical metrics have emerged as valuable tools for interpreting assay results and kinetics. These include:

- 1. Time-to-threshold (TtT): The time required for the fluorescence signal to exceed a predefined threshold (also known as lag time) [5].
- 2. Rate of amyloid formation (RAF): A measure of the kinetics of aggregate growth, which provides insight into the relative quantity of misfolded seed [19].
- 3. Maxpoint ratio (MPR): A ratio-based metric measuring peak normalized fluorescence intensities [15].
- 4. Maximum slope (MS): The steepest rate of fluorescence increase, reflecting the most rapid phase of aggregation [20].

Together, these metrics enable researchers to characterize the kinetics of RT-QuIC reactions comprehensively, enhancing the rigor and reliability of diagnostic decisions.

In addition to analytical tools, quicR provides flexible and customizable visualization capabilities. Leveraging the powerful ggplot2 library [21], quicR enables users to generate high-quality, publication-ready figures. These visualizations can be further customized using the intuitive '+' syntax of ggplot2, allowing for tailored presentations of RT-QuIC data.

By combining standardized metrics, advanced visualization tools, and a commitment to open source science, **quicR** serves as a foundational tool to empower researchers to analyze and present RT-QuIC data with clarity, consistency, and cohesion.

2. Software description

2.1. Software architecture

quicR was developed to address the growing need for efficient data conversion, analysis, and visualization of RT-QuIC data (Figure 1). The overall architecture revolves around integration with the proprietary MARS software (BMG Labtech, Ortenberg, Germany) which exports raw data as Excel workbooks. The data from these workbooks is then curated into usable objects in the R environment, and visualized.

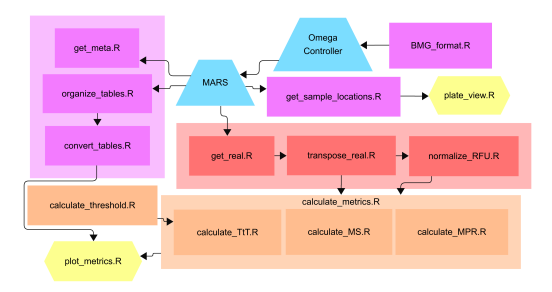


Figure 1: Workflow hierarchy of the **quicR** package. Blue nodes indicate steps where BMG software is needed. Purple nodes indicate functions dedicated to handling metadata. Red nodes are functions that acquire and manipulate raw data. Orange nodes are functions which calculate some metric. Finally, yellow nodes represent data analysis endpoints.

2.2. Software functionalities

The implementation of the quicR package encompasses several streamlined processes designed to facilitate data input, cleaning, transformation, and analysis of RT-QuIC data. This section provides a comprehensive guide to utilizing the package's key functionalities, detailing how to:

- 1. Format and input sample data into Omega control software (BMG Labtech, Ortenberg, Germany).
- 2. Extract, clean, and organize metadata and raw data and apply transformations/normalization for downstream analysis.
- Calculate critical analytical metrics, such as time-to-threshold (TtT), rate
 of amyloid formation (RAF), maxpoint ratio (MPR), and maximum slope
 (MS).
- 4. Visualize raw and analyzed data.

These steps are designed to enhance reproducibility, minimize manual data handling, and enable seamless integration with the MARS software workflow.

2.2.1. Input of Sample IDs into Omega Control Software

The Omega control software allows input of a TXT file containing sample IDs, dilution factors, and their well locations. This file is uniquely formatted, and not easily reproduced manually.

1. BMG_format(): This function allows for input of a CSV file containing the plate layout (see Table 1 for proper formatting), and exports the formatted TXT file. The file can then be imported into the Omega control software before running.

	1	2	3	4	5	6	7	8	9	10	11	12
Α	Р	S01	S02	S03	S04	S05	S06	S07	S08	S09	S10	S11
В	Р	S01	S02	S03	S04	S05	S06	S07	S08	S09	S10	S11
С	Р	S01	S02	S03	S04	S05	S06	S07	S08	S09	S10	S11
D	Р	S01	S02	S03	S04	S05	S06	S07	S08	S09	S10	S11
Ε	N	S01	S02	S03	S04	S05	S06	S07	S08	S09	S10	S11
F	N	S01	S02	S03	S04	S05	S06	S07	S08	S09	S10	S11
G	N	S01	S02	S03	S04	S05	S06	S07	S08	S09	S10	S11
Н	N	S01	S02	S03	S04	S05	S06	S07	S08	S09	S10	S11

Table 1: Example CSV file 96-well plate layout for input into the BMG_format() function. The top left corner is cell "A1" in the CSV file. The top numbered row and the left-most lettered column should never be altered.

2.2.2. Data Cleaning and Transformation

The MARS software exports real-time data as an Excel workbook. Typically, the first sheet in the workbook will include microplate views of both raw data and metadata; however, the metadata on this page is what is most useful for downstream processes. Those tables include the "Sample IDs" and "Dilutions" tables (if dilutions were included in the MARS export). For much of the downstream analysis, it is crucial that the "Sample IDs" table was exported from MARS. If there is no table, the user can simply add it manually.

- organize _tables(): returns a list of tables contained in the first sheet of the exported Excel sheet. These tables contain valuable metadata such as sample IDs, dilution factors, and microplate locations.
- 2. **convert_tables()**: accepts tables outputted from organize_tables() and converts them to columns in a data frame.
- 3. **get_sample_locations()**: extracts the well locations for each sample. Output of this function is used as an argument for visualizing a microplate-level view of real-time data.

2.2.3. Retrieving and Manipulating Raw Data

The raw, real-time data is typically found on the second sheet of the Excel work-book exported from MARS. There are three functions dedicated to the retrieval and cleaning of raw data.

- 1. **get_real()**: Retrieves the raw data from the Excel file, and outputs it as a data frame.
- 2. **transpose_real()**: Swaps the rows and columns which facilitates downstream analyses.
- 3. **normalize_RFU()**: normalizes the raw data by dividing each read by background fluorescence at a given cycle.

2.2.4. Calculations

Three analytical metrics have dedicated functions: TtT, MPR, and MS. RAF does not have a designated function since it is simply the reciprocal of the time-to-threshold (1/TtT). However, it can be calculated using the calculate_metrics() function if TtT is chosen as an optional parameter. Each function below accepts input from the transpose_real() or the normalize_RFU() functions. See Figure 2 for an example of the output of these functions.

 calculate_threhold(): returns a value which is a given number of standard deviations above the average background fluorescence of the entire microplate. This is a popular method of threshold calculation as reviewed in Rowden, et al.[15].

- calculate TtT(): takes the real-time data and calculates the time in hours needed to reach a given threshold value. This threshold can be supplied by calculate threshold() or determined separately by the user.
- 3. calculate MPR(): accepts raw or normalized data and returns the maximum value obtained during the run. If supplied with raw data, it will make a call to the normalize RFU() function.
- 4. calculate _MS(): computes the approximate derivative of the real-time data and returns the maximum value obtained during the run.
- 5. calculate _metrics(): makes a call to functions 2–4 above and generates a data frame with the sample-matched metrics. Also computes RAF if TtT is given as an argument.

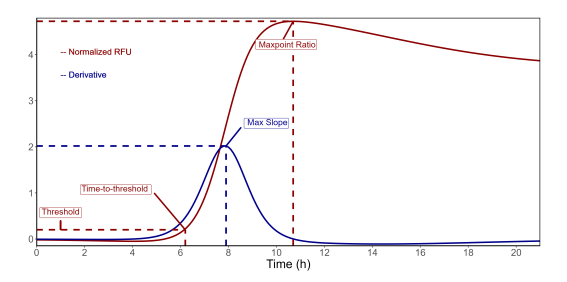


Figure 2: Example graph highlighting the calculated metrics described herein. The red curve represents a raw data curve that has been normalized against background. The maxpoint ratio is calculated as the maximum fluorescent value achieved in the normalized raw data. Time-to-threshold is determined as the time required to cross a given threshold (in this example, the threshold is set at 0.2). The blue curve represents the derivative of the raw data, and max slope is determined as the maximum of the derivative.

2.2.5. Visualization

The ensuing goal of this package is visualizing RT-QuIC data. While there are many ways to represent these data, **quicR** includes two functions which provide a rapid assessment of the RT-QuIC results.

1. plate view(): accepts the output from the get_real() function and get sample locations() function. Makes an 8×12 or 16×24 faceted plot

- depending on if the plate has 96 or 384 wells. Each facet shows the real-time data of each well.
- 2. **plot_metrics()**: requires the output of calculate_metrics() and generates a boxplot of each sample's MPR, MS, RAF, and TtT.

2.3. Comparable Methods

To contextualize the functionality and performance of quicR, we compared it with existing RT-QuIC analysis approaches from the published R packages, QuICSeedR [22] and rtquicR [23]. While all three offer overlapping core functionalities, their design philosophies, input handling, and computational trade-offs differ substantially.

Benchmarking was performed using the rbenchmark library [24] with 50 replications for each function. Comparisons are approximate as equivalent functions were not necessarily found between the three packages, and results are summarized in Table 2. In general, QuICSeedR out-performs quicR across the board. However, quicR addresses several practical limitations that the authors believe justifies the slower times. Notably, QuICSeedR requires an additional, manually formatted Excel file to input sample metadata, whereas quicR directly parses raw and meta data embedded within a single MARS export file. This design reduces user overhead and minimizes potential for transcription errors.

quicR also provides more granular control over metric calculations, allowing users to compute MPR, MS, TtT, and RAF independently rather than as a fixed batch. While QulCSeedR includes built-in support for bulk analysis and diagnostic outputs, these features can be approximated in quicR via standard R idioms (e.g., 'apply' family of functions), and are planned for future development. Overall, quicR favors flexibility, reproducibility, and usability with raw instrument output at the expense of execution speed—trade-offs that align with the package's intended use in diagnostic pipelines.

Feature / Benchmark	quicR	QuICSeedR	rtquicR
Available on CRAN	Yes	Yes	No
Number of input files required	1	2	2
Number of threshold methods	2	3	4
Number of normalization methods	1	1	2
Automated metadata entry	Yes	No	No
Automated sample locations	Yes	Yes	No
Allows empty data	Yes	Yes	No
Compatible with 384-well plates	Yes	Yes	No
Built-in bulk analysis support	No	Yes	Yes
Diagnostic analysis functionality	No	Yes	No
Incorporates dilution factors	Yes	No	Yes
Converts time units	No	Yes	Yes
Calculates metrics independently	Yes	No	Yes
Calculates metrics in bulk	Yes	Yes	No
Calcualates MPR	Yes	Yes	Yes
Calculates MS	Yes	Yes	No
Calculates TtT	Yes	Yes	Yes
Calculates RAF	Yes	Yes	No
Calculates area under the curve	No	No	Yes
Calculates SD50	No	No	Yes
Plate view plot execution time (s)	0.082	0.062	NA
Metric plot execution time (s)	0.009	0.005	0.007
Metric calculations (MPR, MS, etc.) (s)	0.355	0.018	NA
MPR calculation (s)	0.002	NA	0.028
TtT calculation (s)	0.001	NA	0.131
Metadata parsing time (s)	0.127	0.037	NA
Raw data parsing time (s)	0.069	0.050	0.191
Bulk analysis (s/file)	6.550	0.849	1.986

Table 2: Comparison of core features and execution times between **quicR** and **QuICSeedR**. While **QuICSeedR** demonstrates superior execution speed, **quicR** offers enhanced metadata handling, metric flexibility, and support for unprocessed instrument output files. All benchmarks were performed using the same dataset and machine (Windows 11, 16GB RAM).

2.4. Maintenance and Community Involvement

The quicR package is developed with long-term maintainability and community engagement in mind. The source code is hosted on GitHub (https://github.com/gagerowden/quicR), where users can report issues, request features, and contribute improvements via pull requests. A comprehensive CONTRIBUT-ING.md file and issue templates are provided to streamline collaboration and lower the barrier for external contributions. In addition to continuous integration testing, we plan to tag releases with semantic versioning and maintain a changelog to track updates and improvements. Feedback from diagnostic users has already led to enhancements in metadata parsing and output plotting, and we anticipate continued improvements through engagement with the broader scientific community. We encourage users to share their use cases, raise issues, and help shape future development directions.

3. Illustrative examples

3.1. Example Code

To demonstrate the utility of quicR, we used a typical RT-QuIC run performed on a 96-well plate. The reaction was performed on a FLUOstar Omega plate reader (BMG Labtech, Ortenberg, Germany). The file was exported as an Excel workbook where the metadata appears on the first sheet and the raw data on the second.

```
# Step 1: Install and load the package
install.packages("quicR")
library(quicR)
# Step 2: Identify the raw file.
file <- "example.xlsx"
# Step 3: Extract the raw data.
raw <- get real(file)[[1]]
# Step 4: Normalize the data against the background.
normal < - normalize RFU(raw, transposed = FALSE)
# Step 5: Extract the metadata.
meta <- organize tables(file) |> convert tables()
# Step 6: Get sample locations.
locations <- get sample locations(file)</pre>
# Step 7: Create the analyzed data frame.
analyzed <- calculate metrics(normal, meta)
# Step 8: Plot the analyzed data frame.
plot metrics(analyzed)
# Step 9: Plot the plate view.
```

plate view(raw, locations)

Once the file name is identified, the get_real() function is used to extract the raw data. This data is imported as a data frame with each sample designated as an individual column and the first column designated as time. Much of the downstream analysis works best with background-normalized data, so the raw data is then passed to the normalize_RFU() function [15]. This function accepts an integer as an argument for the desired cycle to be selected for background measurement.

Next, the metadata is extracted using the organize_tables() and convert_tables() functions. These take the metadata found on the first sheet of the Excel file and convert them into data frame columns. Finally, metrics are calculated and assigned to their given sample ID using calculate_metrics(). This generates a data frame with sample IDs, dilution factors (if chosen), MPR's, MS's, RAF's, and TtT's.

Once the proper variables have been defined, they can be used as input in the visualization functions. The analyzed data frame can be used as an argument in plot_metrics() to generate a plot such as Figure 3. Additionally, the raw data and sample locations are used to plot the plate view as in Figure 4.

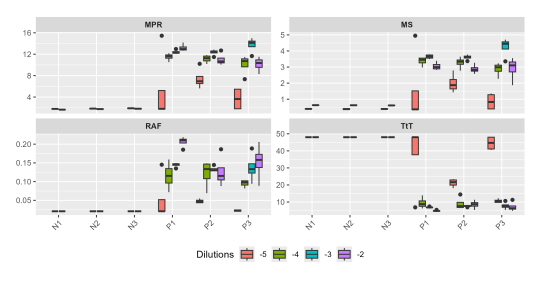


Figure 3: Boxplot of the four critical metrics calculated by the quicR package.

4. Impact

Since its development in 2010 [1, 2], the utility of RT-QuIC has expanded to a host of neurodegenerative diseases caused by prions and other misfolded proteins

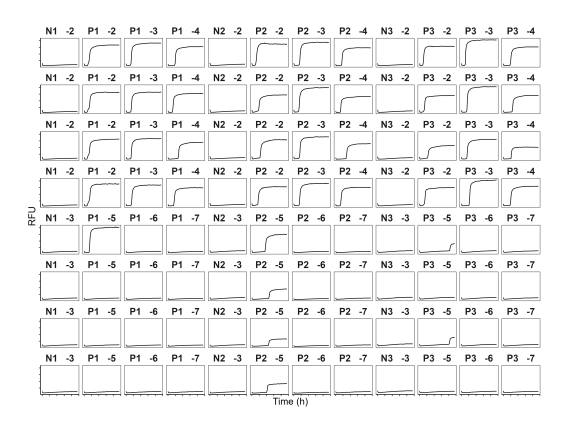


Figure 4: Plate view of RT-QuIC. Wells are identified with a sample ID and dilution factor separated by an optional delimiter (in this case, a whitespace).

[25, 26, 27]. The improved speed and accuracy of the assay has been a boon to human health [5, 28, 29, 30] and animal management efforts [31, 32, 33, 34]. Thus, it is imperative that standardized practices and analyses are adopted.

The quicR package represents a significant advancement in the standardization and reproducibility of RT-QuIC data analysis. By integrating key metrics such as TtT [5], RAF [19], MPR [15], and MS [20], quicR addresses critical gaps in the field, providing researchers and diagnosticians with a robust toolkit for interpreting RT-QuIC data.

One of the primary strengths of quicR lies in its streamlined workflow and user-centric design. The package leverages R's powerful ecosystem and the tidyverse [35] to create high-quality, customizable visualizations, ensuring accessibility for a wide range of users. Additionally, the incorporation of open-source principles allows the broader scientific community to contribute to its development, fostering innovation and adaptability.

Despite these strengths, there are limitations to consider. Currently, quicR is

tailored to data exported from the MARS software, which may limit its applicability to researchers using alternative microplate readers. Future iterations of the package could expand compatibility by incorporating functions to handle diverse data formats. Furthermore, while **quicR** includes robust visualization tools, users seeking highly specialized plots like those made by Li, et al.[22] may require additional customization beyond the package's default capabilities.

Another avenue for improvement lies in the standardization of RT-QuIC diagnostic criteria. **quicR** provides tools to calculate key metrics, but consensus on thresholds and diagnostic interpretations remains a challenge for the field [15]. Collaborative efforts among researchers and clinicians are necessary to define universal criteria, enabling **quicR** to fully realize its potential as a diagnostic tool. Diagnostic determinations could easily be built into the library, but a larger consensus within the research community will need to be reached to warrant inclusion.

5. Conclusions

quicR offers a powerful solution for the cleaning, analysis, and visualization of RT-QuIC data, addressing critical needs in a rapidly evolving field. By enabling consistent data handling and interpretation, quicR provides the foundation for improved diagnostic consistency and reproducibility. The package's open-source nature ensures that it will continue to evolve, integrating new insights and technologies as they emerge.

As RT-QuIC technology advances, tools like quicR will play a pivotal role in bridging the gap between assay development and practical application. By equipping researchers with reliable, standardized tools, quicR not only supports the study of prion and protein misfolding disorders but also serves as a model for the development of software solutions in other diagnostic fields.

Declaration of Conflicting Interests

Peter A. Larsen is a co-founder and stock owner, and Gage R. Rowden is a stock owner of Priogen Corp., a diagnostic company specializing in the ultra-sensitive detection of pathogenic proteins associated with prion and protein-misfolding diseases. The University of Minnesota licensed patent applications to Priogen.

Acknowledgements

Special thanks to Beni Altmann at The Comprehensive R Archive Network (CRAN) for help during the submission process to CRAN. We thank Tiffany Wolf and Marc Schwabenlander for their support through the Minnesota Center

for Prion Research and Outreach. We thank Miranda Huang and Kristin Bondo for testing the code. We would like to acknowledge Suzanne Stone and Sarah Gresch for maintaining lab operations. This work was funded by the Legislative-Citizen Commission on Minnesota Resources.

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