biopixR: Extracting Insights from Biological Images

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Summary

biopixR is an R package designed specifically for the analysis of bioimage data. The package comprises 15 functions that cater to a variety of tasks, including image import, segmentation, feature extraction, quantification, clustering, parameter optimization, and a distinctive algorithm for line gap mitigation. Among these is an optimized edge detection algorithm based on the Canny edge detection algorithm. Together with a noise-resilient threshold method, these two algorithms form the foundation of feature extraction within the biopixR package. Both edge detection and thresholding can be parameterized via an interactive Tcl/Tk user interface. For circular objects, the threshold adjustment factor and the smoothing factor for edge detection can be automatically calculated using Gaussian process regression models (Kriging models). Thereby making an automated feature extraction, which includes segmentation and labeling, available in R. Furthermore, all available methods for feature extraction and filtering can be accessed within a single pipeline function, not found in other packages. The package also includes a function for clustering objects within an image based on their shape features using Self-Organizing Maps (SOM).

Designed for medium-throughput analysis, the package is able to analyze whole directories, utilize multiple cores for parallel processing, and generate detailed log files to track the analytical process. Furthermore, the biopixR package offers distinctive data sets of microbead images and microbeads in water-oil emulsions. The fundamental functionality of biopixR was recently employed in the study by Geithe et al. (2024) to perform quality control on microbeads. As the biopixR package provides access to a range of fundamental data, including object size, quantity, shape, and intensity, it is a valuable tool for researchers in the field. The biopixR package leverages capabilities from the imager (Barthelmé and Tschumperlé 2019) and magick (Ooms 2024) packages to perform its own tasks related to biological image processing and analysis.

Statement of need

Imaging plays a pivotal role in data acquisition within biological laboratories and the broader life sciences. It offers crucial insights into cells (Rödiger et al. 2018; Schneider et al. 2019), biomarkers (Vafajoo et al. 2018), stress responses (Korkmaz, Debelec Butuner, and Roggenbuck 2018), and gene expression (Brenner et al. 2000), which are vital for diagnostics and disease prediction. Microbead technology, known for its rapid and flexible data collection, relies heavily on imaging techniques like microscopy (Rödiger et al. 2014; Ding et al. 2020). The challenge lies in the fast and accurate extraction of valuable information from complex biological images, particularly in the quantification of objects within these images.

The quantification of biological image data is a fundamental aspect of deriving insights into biological processes. This is exemplified by microbead-assays, which offer extensive multiplexing abilities, high-throughput capacity, reduced analysis time, and minimal sample requirements (Rödiger et al. 2014). Methods for detecting microbeads include flow cytometry, microfluidics, and image-based techniques (Choi et al. 2019). A user-friendly tool integrated into R could enhance analysis by combining statistical analysis and visualization techniques. The advancement of smart device imaging and chip development is facilitating the utilization of

microbeads in point-of-care testing (POCT) and disease diagnosis (Dinter et al. 2019; Zhang et al. 2019). This progress underscores the necessity for the development of efficient methods for extracting biological information from images, thereby eliminating the necessity for the use of complex laboratory equipment. Software that extracts attributes such as fluorescence intensity, size and shape allow for microbead encoding and differentiation between populations, thereby enhancing multiplexing capabilities (Zhang et al. 2019).

Microbead-based emulsion Polymerase Chain Reaction (ePCR) assays, which are typically analyzed using fluorescence-activated cell sorting (FACS) (Fraser et al. 2015), could potentially benefit from the use of imaging techniques. As this software is capable of analyzing microbead-based ePCR by preprocessing brightfield droplet images with the provided gap-filling algorithm. The aforementioned applications extend to the assessment of wastewater for the detection of microplastics (Ding et al. 2020), the real-time localization in microbead-based drug delivery systems (Bannerman and Wan 2016), and other fields of life science, such as cell biology.

Consequently, the biopixR package for R is a fundamental tool. The automated evaluation process enables medium-throughput analysis directly from images, simplifying analytical procedures and expanding experimental possibilities.

Software engineering

biopixR (1.0.0, LGLP-3.0 license) is an R (R Core Team 2023a) package (S3 class system). To ensure code quality, we employed various methodologies, including Continuous Integration (CI), unit testing (Wickham 2011), adherence to naming conventions (Bååth 2012), and the application of style guidelines (Wickham 2019). The choice of R as a programming language for image processing is based on its open-source framework, bindings to other languages (Reimert et al. 2024) to improve perfomance, advanced packages such as imager (Barthelmé and Tschumperlé 2019) and magick (Ooms 2024), and its strong support for reproducible research (Xie 2019). The importance of ensuring software reproducibility is widely recognized, not only for our own work but also for the broader scientific community (Gentleman and Temple Lang 2007; Rödiger et al. 2015). One measure to achieve this is by minimizing dependencies on other packages or libraries and single archives whenever possible. Therefore, biopixR depends on R ($\geq 4.2.0$), imager, magick and tcltk, imports data.table and cluster and suggests knitr, rmarkdown, doParallel, kohonen, imagerExtra, GPareto and foreach exclusively from the Comprehensive R Archive Network (CRAN). These features collectively provide a sophisticated environment for image analysis and editing, with the added advantage of community-driven improvements (Chambers 2008). R's integration of analysis with documentation ensures methodological precision and transparency in scientific research, making it a preferred choice for complex image processing tasks. The biopixR package underwent quality control through unit testing using the testthat package (Wickham 2011), ensuring its reliability.

Installation

The biopixR package can be installed from CRAN or GitHub, providing users with stable and developmental versions respectively.

To install the stable version of biopixR from CRAN, execute the following command in R:

install.packages("biopixR")

This command will download and install the latest stable release of the package, ensuring compatibility and reliability.

For users interested in the latest features and ongoing developments, the developmental version of biopixR is available on GitHub. To install this version, it is first necessary to install the devtools package if it is not already present in your R environment:

This command enables the installation of the development build from the GitHub repository. This provides access to the most recent features and updates that may not yet be available in the CRAN release.

Functions

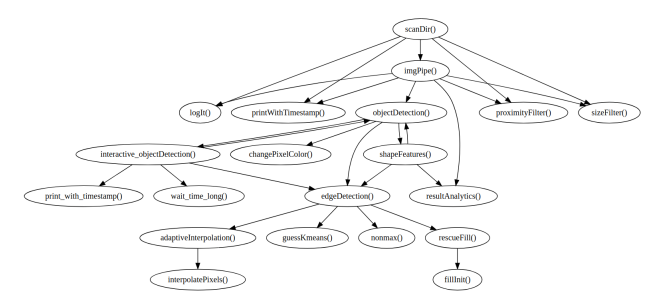


Figure 1: Dependency graph of the functions present in the biopixR package. Showing the levels of complexity by showing the descendants and ancestors of the imgPipe() function. The figure was created using the foodwebr package from Appleton-Fox (2022) (package version 0.1.1, RStudio 2023.09.0+463, R 4.3.2 on Linux, Ubuntu 22.04.3 LTS).

The biopixR package is intended for analyzing bioimage data in R, with a specific focus on the analysis and characterization of microbeads. The package provides tools for image preprocessing, segmentation, feature extraction, filtering, and visualization. It supports automation for medium-throughput analysis, utilizing algorithms to identify spherical objects, extract their features, and implement interactive tools for threshold and smoothing factor selection. Furthermore, it offers features for removing clumped or closely positioned particles to prevent inaccurate results, with the goal of improving the analysis of microparticles in diverse scientific disciplines.

The shapeFeatures() function is capable of extracting shape-related information from detected objects and grouping them using the SOM algorithm (Wehrens and Buydens 2007). The central function of biopixR is imgPipe(), which performs object detection and applies individual filters. This pipeline integrates all the fundamental procedures for comprehensive analysis into a single function. The user is required to provide the input image and select the appropriate detection methods to gain insights into the specific objectives present in the image. Furthermore, the user has the option to customize their workflow with individual filters. In addition, the function is capable of processing multiple color channels, such as the analysis of dual-color microbeads. The integration of this function within the dependency network and its interaction with other functions is illustrated in Figure 1. This function serves as a comprehensive pipeline for image analysis, offering a variety of selectable functions:

• importImage(), joins import functions of the imager (Barthelmé and Tschumperlé 2019) and magick (Ooms 2024) packages,

- edgeDetection(), a combination of a Canny edge detector and gap filling (Barthelmé and Tschumperlé 2019),
- objectDetection(), detects objects in an image by edge detection or thresholding,
- sizeFilter(), eliminates objects that exceed or fall below a certain size threshold,
- proximityFilter(), filters objects that are in proximity,
- resultAnalytics(), summarizes the extracted features in a clear and concise manner.

The biopixR package includes functions for analyzing entire directories, allowing for medium-throughput analysis. Making feature extraction and image clustering easily accessible:

- haralickCluster(), extracts Haralick features and clusters information using Partitioning Around Medoids (PAM) (Haralick, Shanmugam, and Dinstein 1973; Carlson 2018; Maechler et al. 2023),
- scanDir(), utilizing the pipeline for whole directory analysis.

The fillLineGaps() algorithm, along with helper functions:

- interpolatePixels(), calculates the coordinates required to connect two given points,
- adaptiveInterpolation(), searches a given radius surrounding a line end for contours and connects them,

addresses the issue of discontinuous edges by iteratively scanning for line ends within the image and reconnecting them to adjacent contours. Other approaches for dealing with discontinuous contours are not found in other R packages.

Examples demonstrating the use of biopixR for image analysis tasks can be found in the package's vignette (Brauckhoff, Rödiger, and Kieffer 2024).

Graphical User Interface

The biopixR package has broader applicability beyond microbead detection. The biopixR package is adaptable and can be utilized in any research field where the identification of distinct objects in images can be achieved through the use of a Canny edge detector or thresholding. This encompasses research areas such as foci detection, microplastics, and plant seeds. The automation methods employed in biopixR are predicated on the assumption of circular objects, rendering it particularly well-suited for the detection, quantification, and extraction of useful information from circular objects within images. Another integrated tool is an interactive function that assists the user in selecting the optimal input for their analysis. The function interactive_objectDetection() initiates a graphical user interface (GUI) (Figure 2) that utilizes the Tcl/Tk framework (R Core Team 2023b), enabling users to adjust the threshold and smoothing settings of the image.

Conclusion

In summary, biopixR represents a significant advancement in image analysis capabilities for R users. By providing both automation methods and interactive tools, the package empowers researchers to extract valuable insights from images with ease. Its adaptability across various research fields makes it a valuable tool for researchers seeking efficient solutions. Leveraging the power of R and its extensive library of packages, users can seamlessly integrate biopixR into their existing workflows, streamlining data analysis and visualization tasks. Moreover, the package's minimal dependencies ensure long-term stability and maintainability over time, making it an attractive choice for researchers seeking reproducible software. A comparative analysis between human and software-based object quantification from images underscores the importance of the biopixR package in terms of accuracy and reproducibility (see extended vignette: https://github.com/Brauckhoff/biopixR/tree/main/vignettes/detailed).

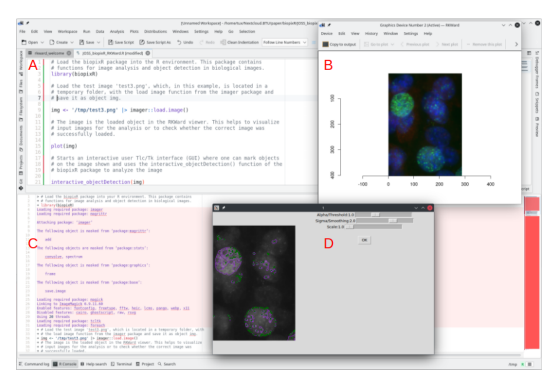


Figure 2: Graphical User Interface for interactive parameter selection. The function interactive_objectDetection() provides a simple interface with sliders to adjust threshold, smoothing, and scale. It highlights object contours in purple and centers in green for easy visualization. A) In this example, the GUI was used in RKWard (0.7.5z+0.7.6+devel3, Linux, TUXEDO OS 2, (Rödiger et al. 2012)). With a few commands, an image can be imported and analyzed. B) The plot() function displays the false-color image as a preview. In this figure, cells with DNA damage (similar to Rödiger et al. (2018)) are visible. C) Loading the biopixR package in the R console shows additional information such as loaded libraries and the number of CPU threads (n = 20, parallel processing). D) The rendering process is displayed on the console, including timestamps and the current state.

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