

- 1 MotilA A Python pipeline for the analysis of
- ² microglial fine process motility in 3D time-lapse
- multiphoton microscopy data
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DOI: 10.xxxxx/draft

Software

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Editor: Open Journals ♂ Reviewers:

Copenjournals

Submitted: 01 January 1970 Published: unpublished

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Summary

MotilA is a Python-based image analysis pipeline for quantifying fine process motility of microglia from 3D time-lapse two-channel fluorescence microscopy data. Developed for high-resolution multiphoton in vivo imaging datasets, MotilA enables both single-file and batch processing across multiple experimental conditions. It performs image preprocessing, segmentation, and motility quantification over time, using a pixel-based change detection strategy that yields biologically interpretable metrics such as the turnover rate (TOR) of microglial fine processes. While originally designed for microglial imaging, the pipeline can be extended to other cell types and imaging applications that require analysis of dynamic morphological changes. MotilA is openly available, platform-independent, and includes extensive documentation, tutorials, and example data to facilitate adoption by the broader scientific community. It is released under the GPL-3.0 open-source license.

Statement of need

Microglia are innate immune cells of the central nervous system and exhibit highly dynamic, motile processes that continuously scan their environment (M. Fuhrmann et al., 2010; Nebeling et al., 2023; Nimmerjahn et al., 2005; Prinz et al., 2019; Tremblay et al., 2010). Quantifying microglial motility at the level of fine processes is crucial for studying their function in health and disease, including neurodegeneration, inflammation, and synaptic remodeling. However, despite the biological importance of this analysis, there is currently no dedicated open-source tool tailored for this task.

To date, researchers typically quantify microglial motility manually (see., e.g., Nebeling et al. (2023)) using general-purpose image processing software such as Fiji/ImageJ (Schindelin et al., 2012) or ZEISS ZEN (Carl Zeiss Microscopy GmbH, Accessed 2025). While these approaches are well established in the field, they are time-consuming, lack reproducibility, and are not well suited for batch processing or cohort-level comparisons. They often focus on individual microglia, whereas *MotilA* enables analysis of the full field of view, allowing for more comprehensive and scalable quantification. Manual workflows are also more susceptible to human bias, limiting their scalability and objectivity (Brown, 2017; Lee et al., 2024; Misra et al., 2015; Wall et al., 2018).

MotilA addresses these limitations by providing an end-to-end, user-friendly, and batchcapable pipeline specifically designed for 3D time-lapse two-channel microscopy data. It
supports standardized workflows for single- and multi-channel datasets, integrates essential
preprocessing steps (registration, spectral unmixing, histogram normalization), and derives
biologically meaningful motility metrics from binarized pixel dynamics. The method builds on



- 41 strategies used in prior studies but automates the workflow in a reproducible, scalable, and
- open-source manner. MotilA thus fills a critical gap in neuroimaging analysis pipelines and is
- particularly valuable for labs working with multiphoton in vivo imaging.

What does *MotilA* do?

45 MotilA is a modular and customizable image analysis pipeline written in Python that quantifies

microglial fine process motility from time-lapse fluorescence microscopy data, typically acquired

with two-photon (Denk et al., 1990; Helmchen & Denk, 2005) or three-photon in vivo imaging

(F. Fuhrmann et al., 2024; Horton et al., 2013). Although it was originally developed for

microglial analysis, the pipeline is adaptable to other cell types and imaging contexts involving

50 dynamic morphological changes over time.

51 At its core, MotilA extracts sub-volumes from 3D time-stacks, performs 2D maximum intensity

projections, and segments the resulting images to classify pixel-wise changes in microglial

morphology. These changes are quantified frame-by-frame and used to calculate biologically

interpretable metrics, including the turnover rate (TOR). The design is tailored to biological

55 imaging data, with particular attention to typical issues such as z-axis projection loss, channel

bleed-through, motion artifacts, photobleaching, and signal heterogeneity.

57 The pipeline supports both single-file processing and large-scale batch analysis. Parameters are

highly customizable either programmatically or via metadata files, and all results are automati-

cally logged, saved, and summarized for downstream statistical analysis. The outputs include

segmented image series, intermediate diagnostics (e.g. histograms, projections, brightness

traces), and Excel spreadsheets with motility metrics.

To accommodate large-scale, high-resolution imaging data, MotilA supports memory-efficient

file handling via the Zarr format (Miles et al., 2025), enabling processing of large TIFF files

using memory mapping to avoid RAM overload.

MotilA can be run via Python scripts or Jupyter notebooks, and it includes extensive docu-

mentation, examples, and a tutorial dataset to make onboarding straightforward.

67 We welcome community contributions and issue reports via the GitHub repository: https://orunners.com/https://or

//github.com/FabrizioMusacchio/motila.

How is "motility" determined?

70 MotilA quantifies motility by analyzing pixel-wise changes in microglial fine processes' morphol-

71 ogy over time. The pipeline first extracts a sub-volume around a user-defined z-axis center from

each 3D image stack and applies a 2D maximum intensity projection to reduce dimensionality.

Although this sacrifices some z-axis information, it enables efficient segmentation and pixel-level

tracking while maintaining biological interpretability.

At each time point t_i , the projected and binarized image $B(t_i)$ is compared to the next time

point $B(t_{i+1})$. A temporal variation map $\Delta B(t_i)$ is computed as:

$$\Delta B(t_i) = 2 \times B(t_i) - B(t_{i+1})$$

Based on this difference image, each pixel is classified as:

- Stable (S) if $\Delta B=1$
- Gained (G) if $\Delta B = -1$
 - Lost (L) if $\Delta B = 2$

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From these categories, *MotilA* calculates the microglial fine process turnover rate (TOR), a

82 central metric representing the fraction of pixels that changed:



$$TOR = \frac{G+L}{S+G+L}$$

- This approach allows for a quantitative assessment of microglial process dynamics at each
- 84 time point and across the full recording session. The same principle can be extended to other
- motile cell types or dynamic cellular structures where morphological changes manifest as gain
- 86 or loss of segmented pixels over time.
- 87 The implementation is based on analytical strategies described in previous studies such as M.
- Fuhrmann et al. (2010) and Nebeling et al. (2023), with added flexibility for batch processing,
- 89 filtering, and parameter tuning.

Mey features

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MotilA offers a combination of modularity, reproducibility, and scalability specifically tailored to motility analysis in multiphoton *in vivo* imaging. Its key features include:

Automated preprocessing pipeline

Includes optional steps for image registration (2D and 3D), spectral unmixing, histogram equalization for contrast enhancement within time points, and histogram matching for brightness normalization across time points (e.g. to correct for photobleaching), as well as noise reduction via median and Gaussian filtering.

Flexible segmentation and thresholding

Supports multiple adaptive thresholding methods (e.g. Otsu, Li, Triangle) and customizable blob detection settings to isolate fine microglial processes or similar structures.

Pixel-based motility quantification

Tracks pixel-wise changes between time points to classify stable, gained, and lost pixels, allowing biologically interpretable metrics like the turnover rate (TOR).

Batch processing capabilities

Enables large-scale processing of multiple datasets with a standardized folder structure and parameter metadata sheets, suitable for cohort-level studies.

User-defined projection settings

Allows flexible extraction of sub-volumes and z-projection around multiple centers to avoid overlapping cells and vascular artifacts.

Memory-efficient file handling

Supports memory mapping of large TIFF files via the Zarr format, enabling efficient processing of high-resolution time-lapse datasets without exhausting system RAM.

Metadata integration and parameter logging

Automatically reads per-dataset settings from metadata files (e.g. Excel sheets), and stores processing parameters and outputs in structured result folders.

Cross-platform compatibility

Runs on Windows, macOS, and Linux, tested with Python 3.9 and compatible with common scientific computing environments via Conda.

Tutorials and example data included

Comes with Jupyter notebooks, example datasets, and clear documentation to help new users get started quickly.

122 Pipeline steps

The *MotilA* pipeline follows a modular sequence of image processing and analysis steps designed for robust and reproducible quantification of motility from multi-dimensional imaging data. It supports both single-file and batch workflows and includes options for fine-grained customization at each step.



27 Core pipeline steps

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For single datasets, *MotilA* executes the following sequence (Figure 1a)):

1. Load image data

Supports TIFF files (Gohlke, 2025) in TZCYX (multi-channel) or TZYX (single-channel) format, following ImageJ/Fiji conventions (T: time, Z: z-axis, C: channel, Y: height, X: width).

2. Extract sub-volumes

Selects a z-stack around a projection center for each time point, allowing focused analysis and optional multiple projections per stack.

3. (Optional) Register sub-volumes

Applies 3D motion correction (Anuta, 1970; Guizar-Sicairos et al., 2008; Kuglin & Hines, 1975) to each time series stack using user-defined template strategies (e.g. mean, median).

4. (Optional) Perform spectral unmixing

Removes signal bleed-through between channels, especially relevant for two-channel imaging setups.

5. **Z**-projection

Projects each 3D sub-volume into a 2D image via maximum intensity projection to simplify segmentation and speed up processing.

6. (Optional) Register projections

Aligns the 2D projections across time to correct for lateral motion artifacts.

7. (Optional) Apply histogram equalization

Enhances local contrast within each projection using contrast-limited adaptive histogram equalization (CLAHE) (Pizer et al., 1987; Walt et al., 2014).

8. (Optional) Apply histogram matching

Normalizes brightness across time points to mitigate bleaching effects or intensity drift (Walt et al., 2014).

9. (Optional) Apply filtering

Reduces noise with optional median filtering (square or circular kernel) and/or Gaussian smoothing (Harris et al., 2020; Virtanen et al., 2020).

10. Segment microglial processes

Applies adaptive thresholding (Glasbey, 1993; Li & Tam, 1998; Otsu, 1979; Prewitt & Mendelsohn, 1966; Ridler et al., 1978; Yen et al., 1995; Zack et al., 1977) and blob filtering (Fiorio & Gustedt, 1996; Walt et al., 2014; Wu et al., 2005) to identify and isolate morphologically relevant structures.

11. Analyze motility

Quantifies pixel-level changes over time to classify stable, gained, and lost regions, from which motility metrics are derived.

All intermediate outputs and metrics are saved for validation and further analysis.

Batch processing steps

MotilA supports fully automated batch processing using a standardized folder structure and Excel-based metadata configuration. This enables reproducible cohort-level analysis across many animals or experimental conditions.

1. Define a project folder

Each dataset is placed in an ID-specific subdirectory, containing imaging files, metadata, and optional result directories.

2. Run the batch process

The core pipeline is executed for each dataset using shared or per-dataset parameters defined in metadata.xls.

3. Save results

Segmentation outputs, projections, and motility metrics are stored in structured result



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folders for each dataset.

4. Batch-collect metrics

Aggregates metrics across datasets into cohort-level Excel files for downstream statistical analysis.

This design enables large-scale, reproducible quantification of microglial motility with minimal manual intervention.

184 Assessing results and analyzing outputs

MotilA provides rich output in the form of diagnostic plots, intermediate image files, and structured Excel tables to support both per-dataset assessment and cohort-level statistical analysis.

188 Per-dataset assessment

For each processed image stack, MotilA generates:

- Segmented images and overlays showing gained, lost, and stable regions across time points.
 - Histogram plots for brightness, pixel area, and thresholding diagnostics.
 - Motility metrics table (motility.xlsx) containing:
 - Gained pixels (G)
- Lost pixels (L)
 - Stable pixels (S)
 - turnover rate (TOR) per time point
- Brightness metrics (brightness.xlsx) tracking average pixel intensity over time.
- Cell area metrics (cell_pixel_area.xlsx) reporting the segmented microglial pixel area per time point.
- These outputs help assess segmentation quality, evaluate photobleaching or signal loss, and refine preprocessing parameters as needed.
- 203 Cohort-level batch analysis
- During batch processing, *MotilA* can aggregate key metrics from all datasets into shared summary files, including:
 - all_motility.xlsx All G/L/S/TOR metrics across datasets.
 - all_brightness.xlsx Mean brightness per dataset and time point.
 - all_cell_pixel_area.xlsx Segmented area per dataset and time point.
 - average_motility.xlsx Dataset-wise average motility metrics across the full recording.
- These results allow for statistical comparison of motility dynamics across experimental conditions and facilitate downstream visualization and modeling in tools like Python, R, or Excel.
- This multi-level output strategy ensures both technical validation and biological insight, making *MotilA* suitable for both exploratory and hypothesis-driven studies.

215 Main functions

216 The three main entry points for the pipeline are:

- process_stack Processes a single image stack, performing the full pipeline.
- batch_process_stacks Executes the pipeline across multiple datasets in a project folder.
- batch_collect Gathers motility metrics from all datasets for cohort-level analysis.



- Each function supports extensive parameterization via arguments or metadata files.
- A complete overview of configurable parameters for single-file processing, batch workflows, and image enhancement is provided in the MotilA README.

224 Useful helper functions

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225 Several additional functions assist with data preparation and quality control, including:

- tiff_axes_check_and_correct Automatically adjusts TIFF axis order to TZCYX/ TZYX if needed.
- hello_world Verifies a successful import of the MotilA module.
- logger_object Initializes logging for the current analysis session.

230 Applications and limitations

MotilA was designed with a primary focus on the analysis of microglial fine process motility in vivo, using high-resolution 3D time-lapse two-channel fluorescence microscopy data. Its modular design and general image processing framework, however, make it applicable to a broader range of dynamic imaging contexts.

235 Applications

Microglial dynamics

Quantification of process turnover during surveillance, neuroinflammation, or disease models such as neurodegeneration and injury.

Neuronal structural plasticity

While *MotilA* is optimized for microglial processes, its pixel-based change detection framework can in principle be adapted to analyze dynamic changes in dendrites or axons — such as growth, retraction, or remodeling — provided the structures can be reliably segmented across time.

■ Two-channel in vivo imaging

Effective for experiments involving simultaneous imaging of microglia and neurons (e.g., Cx3Cr1-GFP with Thy1-YFP), with spectral unmixing to reduce bleed-through from overlapping channels or fluorophores.

Cohort-level studies

Designed to analyze and compare motility metrics across large experimental groups, enabling high-throughput, statistically robust results.

Teaching and prototyping

The example datasets and tutorials make *MotilA* a useful tool for training purposes or prototyping new analysis approaches.

254 Limitations

Loss of z-resolution

The use of 2D maximum intensity projections simplifies processing but sacrifices z-axis information. This may lead to overlapping structures and limits spatial specificity.

Segmentation-dependent

Accuracy depends on appropriate thresholding and image quality. Overlapping processes, blood vessels, or low signal-to-noise ratios can reduce segmentation performance.

Limited spectral unmixing

The current unmixing approach is a simple channel subtraction. More advanced unmixing strategies may be required for some experimental setups.

Not a general-purpose tracking tool

MotilA is optimized for pixel-level process motility, not for full cell tracking or object-based morphological quantification over time.



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Assumes TIFF input with standardized axis order
 Input images must conform to TZCYX or TZYX structure; other formats require conversion.

Despite these limitations, *MotilA* provides a powerful, reproducible framework for analyzing microglial motility and similar biological processes, especially in experimental setups where manual analysis would be impractical.

273 Real-world example

To demonstrate its practical utility, *MotilA* includes a fully compatible example dataset of *in vivo* two-photon time-lapse imaging stacks from the mouse frontal cortex (Gockel et al., 2025).

These data were acquired to assess microglial fine process motility under control conditions and during complement C4 overexpression, a genetic risk factor for schizophrenia.

The dataset contains two 5D TIFF stacks with the following structure:

- T: 8 time points (5-minute intervals over 35 minutes)
- C: 2 imaging channels (Cx3cr1-GFP for microglia, tdTomato for neurons)
- **Z**: ~60 optical sections (1 µm step size)
- Y, X: ~1200 \times 1200 px (~125 \times 125 μ m² field of view)

The files are formatted for direct use with *MotilA*, requiring no manual reorganization or preprocessing.

Figure 1 summarizes the full MotilA workflow as applied to the example dataset. For visualization purposes, the original full-field dataset was cropped around a single microglial cell 286 to reduce background clutter and allow detailed inspection of each processing step. Panel a) 287 outlines the core and optional steps in the processing pipeline. Panel b) shows z-projections of the microglial cell at time points t_0 and t_1 , including raw data, contrast enhancement, and 289 filtering prior to segmentation. Panel c) displays the delta image used for motility quantification 290 and the corresponding pixel-wise classification into stable (S, blue), gained (G, green), and lost 291 (L, red) pixels. Panel d) tracks the average brightness of the segmented cell relative to the first time point, which helps assess signal stability and potential bleaching. Panel e) presents the turnover rate (TOR) across all time points, capturing the dynamics of microglial process remodeling.



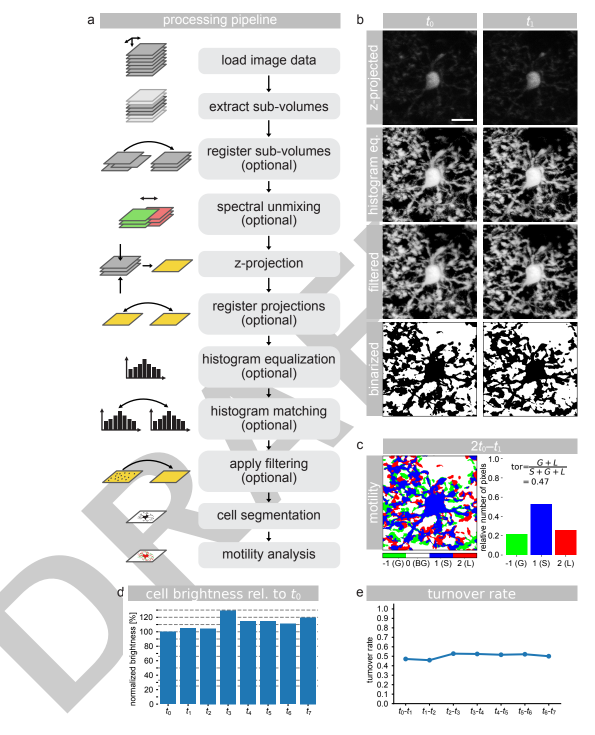


Figure 1: Step-by-step illustration of the MotilA pipeline using the included test dataset. a) Overview of the image processing pipeline, showing core and optional steps. b) Example projections of a cropped microglial cell at time points t_0 and t_1 , including raw, histogram-equalized, filtered (median and Gaussian), and binarized versions. c) Binarized pixel-wise comparison between t_0 and t_1 , with classification into stable (S, blue), gained (G, green), lost (L, red), and background (BG, white) pixels, along with the corresponding pixel statistics. d) Normalized cell brightness over time, relative to t_0 , used to assess bleaching and signal stability. e) Turnover rate (TOR) plotted across all time points for the same cell, representing process-level motility dynamics. All microglial image panels in b and c are shown at the same scale. Scale bar in the top-left image of panel b represents $10\,\mu m$.



Past and ongoing projects

MotilA has already been successfully applied in multiple neuroscience studies involving in vivo imaging of microglia and neurons in the mouse brain.

The following published and preprint works used *MotilA* to analyze fine process motility in physiological and pathological contexts:

Crux et al. (2024)

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Investigated the role of actin depolymerizing factors ADF/Cofilin1 in microglial motility and memory formation. *MotilA* was used to quantify reduced motility in knockout mice. \rightarrow https://doi.org/10.1101/2024.09.27.615114

■ F. Fuhrmann et al. (2024)

Employed deep three-photon imaging of microglia in the medial prefrontal cortex to measure sub-cellular process dynamics in awake mice. *MotilA* was used to quantify microglial turnover at depths beyond 1 mm.

→ https://doi.org/10.1101/2024.08.28.610026

Gockel et al. (2025)

Generated and published the example dataset accompanying this pipeline, which was used to demonstrate microglial motility changes in response to complement C4 overexpression.

→ https://doi.org/10.5281/zenodo.15061566

These studies showcase the pipeline's suitability for both targeted microglial investigations and large-scale, high-resolution imaging projects. Ongoing work continues to extend *MotilA*'s application to additional brain regions, genetic perturbations, and imaging modalities, including multi-channel and high-speed two-photon datasets.

Acknowledgements

We gratefully acknowledge the Light Microscopy Facility (LMF) and Animal Research Facility (ARF) at the German Center for Neurodegenerative Diseases (DZNE), Bonn, for their essential support in data acquisition and technical infrastructure.

This work was supported by the DZNE and by grants to MF from the European Union ERC-CoG (MicroSynCom 865618) and the German Research Foundation DFG (SFB1089 C01, B06; SPP2395). MF is a member of the DFG Excellence Cluster ImmunoSensation2. This work was also supported by the iBehave network to MF and the CANTAR (CANcerTARgeting) network to FN, both funded by the Ministry of Culture and Science of the State of North Rhine-Westphalia. The funders had no role in study design, data collection and interpretation, or the decision to submit the work for publication. FN received additional funding from the Mildred-Scheel School of Oncology Cologne-Bonn.

All animal procedures related to the example dataset were conducted in compliance with institutional, national, and international regulations. Experiments were approved by the relevant animal care and use committees at DZNE (Germany), following guidelines equivalent to the ARRIVE 2.0 framework. All efforts were made to reduce the number of animals used and to refine experimental conditions in accordance with the 3Rs (Replacement, Reduction, and Refinement) principles.

We also acknowledge the open-source community whose tools and contributions made the development of *MotilA* possible.

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