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DOI: 10.21105/joss.04080

Software
- **Review**
- **Repository**
- **Archive**

**Summary**

Rainbow is a free, open-source and cross-platform Python-based tool for Air-Liquid Interface (ALI) cell culture time-lapse image analysis. Rainbow accepts input time-lapse images in standard image formats, such as TIFF, or microscopy file formats, such as ND2, and then computes the optical flow; that is, the apparent motion of individual pixels in an image ([Beauchemin & Barron, 1995; Turaga et al., 2010; Zhai et al., 2021]), across multiple frames using a deep learning based pretrained optical flow model ([Jiang et al., 2021]). Rainbow then applies circular data analysis to the pixel-level optical flow information to calculate the average magnitude [0, ∞] (μm) and direction [0, 360) (°) of motion between adjacent frames to quantify cell motility. Additionally, the variance of the magnitude [0, ∞] (μm) and direction [0, ∞] (°) of motion between adjacent frames is calculated to quantitatively capture the degree of heterogeneity in cell motility.

For each experiment, a CSV file with the minimum, maximum, mean, standard deviation, and variance of the magnitude and direction of cell movement between adjacent frames in an image sequence is produced. Multiple CSV files from different experiments can be combined into one file that can be analyzed for differences in cell motility across multiple experiments. Rainbow also includes a high-resolution and easily readable unified hue/saturation-based visualization scheme for the instantaneous vector field of motion between adjacent frames of an image sequence to qualitatively show cell motion. Rainbow can be used through a graphical user interface or command line interface and can generate a HTML report containing output images, videos, publication ready figures, and CSV files detailing cell dynamics (refer to Examples folder on GitHub). Importantly, our software is not limited to ALI culture image analysis, and developers can extend the software’s existing pipeline to other use cases. For example, the optical flow model can be readily substituted with different models, as we utilized the Factory Method creational software design pattern. The data analysis and report generated can be adjusted through interactive Jupyter Notebooks, allowing for a flexible and versatile system.

Some of Rainbow's visualizations are shown in Figure 1.

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Figure 1: Rainbow optical flow visualization. **A:** The direction of motion at any position within Rainbow-generated optical flow images is measured clockwise from the initial horizontal position of a unit circle (left) and is shown using hue values (right). **B:** The magnitude of motion at any position within optical flow images is shown using saturation values. High saturation (100%) corresponds to high magnitude of motion and low saturation (25%) corresponds to low magnitude of motion. **C, G:** Still frames taken from two separate ALI culture image sequences. **D, H:** Unified visualization of optical flow magnitude and direction between adjacent frames of two ALI culture image sequences using Rainbow. The arrow indicates the average direction of motion across the image sequence. The circles indicate three localized vortexes that the cells move around in a swirl-like motion as they change direction. **E, I:** Traditional visualization of optical flow between adjacent frames of two ALI culture image sequences using quiver plots containing vector arrows at every 70 px. **F, J:** Polar plots visualizing motion magnitude (concentric circles; µm) and direction (azimuthal angle; degrees) in the same frame of two ALI culture image sequences. Colour scale indicates the number of points migrating towards given direction. All left positioned subfigures from row 2 onwards (C-F) correspond to the same ALI culture image sequence while right positioned subfigures correspond to a different image sequence (G-J). For complete insight, refer to Examples folder on GitHub containing videos.

Statement of need

Differentiated primary airway epithelial cells cultured at air-liquid interface (ALI) are commonly used to assess airway epithelial function in vitro in health and disease, such as asthma (Chen & Schoen, 2019; Looi et al., 2018; Martinovich et al., 2017). The integration of image analysis and ALI cell cultures has provided novel insights into cell dynamics, such as the recently identified unjammed-to-jammed transition of AEC characterized by changes in cell motility (Mitchel et al., 2020; Park et al., 2015). In chronic respiratory diseases like asthma, increased cell motility and AEC unjamming have been linked to airway remodeling and disease development (Mitchel et al., 2020; Park et al., 2015). However, the image analyses performed in these studies are limited. For example, handcrafted methods from the MATLAB Computer Vision Toolbox that compute optical flow have been used to extract cell motion information from ALI culture images (Mitchel et al., 2020). This approach requires licenced software, and handcrafted optical flow estimation methods have been outperformed in terms of accuracy by deep learning methods (Savian et al., 2020). Furthermore, commonly used cell motion metrics, such as average cell speed, do not capture all unique aspects of cell motion, such as the heterogeneity of cell migration patterns across time. Cell motion is commonly visualized using vector fields, which are useful but bound by an inverse relationship between resolution and readability (Henkes et al., 2020; Nnetu et al., 2012; O’Sullivan et al., 2020).

To increase understanding of lung disease mechanisms and development of new treatment options for patients, there is a need for open-source solutions for ALI culture image analyses that can be broadly implemented across cell biology laboratories. To the best of our knowledge, Rainbow is the first easy-to-use software tool that performs all the above analyses automatically for efficient utilization by non-programmers. Rainbow produces automatic cell motion quantifications, figures, and reports that are all easily transferrable into publications. We anticipate that Rainbow will provide cell motion characterization for each experiment and allow for easy comparisons among multiple experiments to uncover cell migration mechanisms previously undetermined in health and disease.

Acknowledgements

We would like to thank the contribution and assistance of all the respiratory fellows, anaesthetists, nurses, hospital staff at St John of God Hospital, Subiaco (Human Research Ethics Committee study approval #901), and Western Australian Epithelial Research Program (WAERP) members. We would also like to thank the families and children participating in this project. This work was supported by the Wal-Yan Respiratory Research Centre Inspiration Award, Cystic Fibrosis Charitable Endowment Charles Bateman Charitable Trust, Western Australian Department of Health Merit Award, and BHP-Telethon Kids Blue Sky Award. Furthermore, this project relies on high quality open source Python packages: NumPy (Harris et al., 2020), matplotlib (Hunter, 2007), pandas (Wes McKinney, 2010), PyYaml, ND2Reader, Gooey, Physt, imutils, MoviePy, natsort, PIMS, tqdm (Costa-Luis et al., 2021), pytests (Krekel et al., 2004), FFmpeg (Tomar, 2006), OpenCV (Bradski, 2000), Jupyter Notebook (Kluyver et al., 2016), Astropy (Astropy Collaboration et al., 2018, 2013), PyTorch (Paszke et al., 2019), and scikit-image (Walt et al., 2014).

References


