

Quantitative analysis of surface wave patterns of Min proteins

Meindlhumer, S.; Kerssemakers, J.W.J.; Dekker, C.

DOI

[10.3389/fphy.2022.930811](https://doi.org/10.3389/fphy.2022.930811)

Publication date

2022

Document Version

Final published version

Published in

Frontiers in Physics

Citation (APA)

Meindlhumer, S., Kerssemakers, J. W. J., & Dekker, C. (2022). Quantitative analysis of surface wave patterns of Min proteins. *Frontiers in Physics*, 10, Article 930811. <https://doi.org/10.3389/fphy.2022.930811>

Important note

To cite this publication, please use the final published version (if applicable). Please check the document version above.

Copyright

Other than for strictly personal use, it is not permitted to download, forward or distribute the text or part of it, without the consent of the author(s) and/or copyright holder(s), unless the work is under an open content license such as Creative Commons.

Takedown policy

Please contact us and provide details if you believe this document breaches copyrights. We will remove access to the work immediately and investigate your claim.



OPEN ACCESS

EDITED BY

David M Holloway,
British Columbia Institute of
Technology, Canada

REVIEWED BY

Petra Schwille,
Max Planck Society, Germany
Eldon Emberly,
Simon Fraser University, Canada

*CORRESPONDENCE

Cees Dekker,
c.dekker@tudelft.nl

SPECIALTY SECTION

This article was submitted to Biophysics,
a section of the journal
Frontiers in Physics

RECEIVED 28 April 2022

ACCEPTED 30 June 2022

PUBLISHED 03 August 2022

CITATION

Meindlhumer S, Kerssemakers J and
Dekker C (2022), Quantitative analysis of
surface wave patterns of Min proteins.
Front. Phys. 10:930811.
doi: 10.3389/fphy.2022.930811

COPYRIGHT

© 2022 Meindlhumer, Kerssemakers
and Dekker. This is an open-access
article distributed under the terms of the
[Creative Commons Attribution License
\(CC BY\)](https://creativecommons.org/licenses/by/4.0/). The use, distribution or
reproduction in other forums is
permitted, provided the original
author(s) and the copyright owner(s) are
credited and that the original
publication in this journal is cited, in
accordance with accepted academic
practice. No use, distribution or
reproduction is permitted which does
not comply with these terms.

Quantitative analysis of surface wave patterns of Min proteins

Sabrina Meindlhumer, Jacob Kerssemakers and Cees Dekker*

Department of Bionanoscience, Kavli Institute of Nanoscience Delft, Delft University of Technology, Delft, Netherlands

The Min protein system is arguably the best-studied model system for biological pattern formation. It exhibits pole-to-pole oscillations in *E. coli* bacteria as well as a variety of surface wave patterns in *in vitro* reconstitutions. Such Min surface wave patterns pose particular challenges to quantification as they are typically only semi-periodic and non-stationary. Here, we present a methodology for quantitatively analysing such Min patterns, aiming for reproducibility, user-independence, and easy usage. After introducing pattern-feature definitions and image-processing concepts, we present an analysis pipeline where we use autocorrelation analysis to extract global parameters such as the average spatial wavelength and oscillation period. Subsequently, we describe a method that uses flow-field analysis to extract local properties such as the wave propagation velocity. We provide descriptions on how to practically implement these quantification tools and provide Python code that can directly be used to perform analysis of Min patterns.

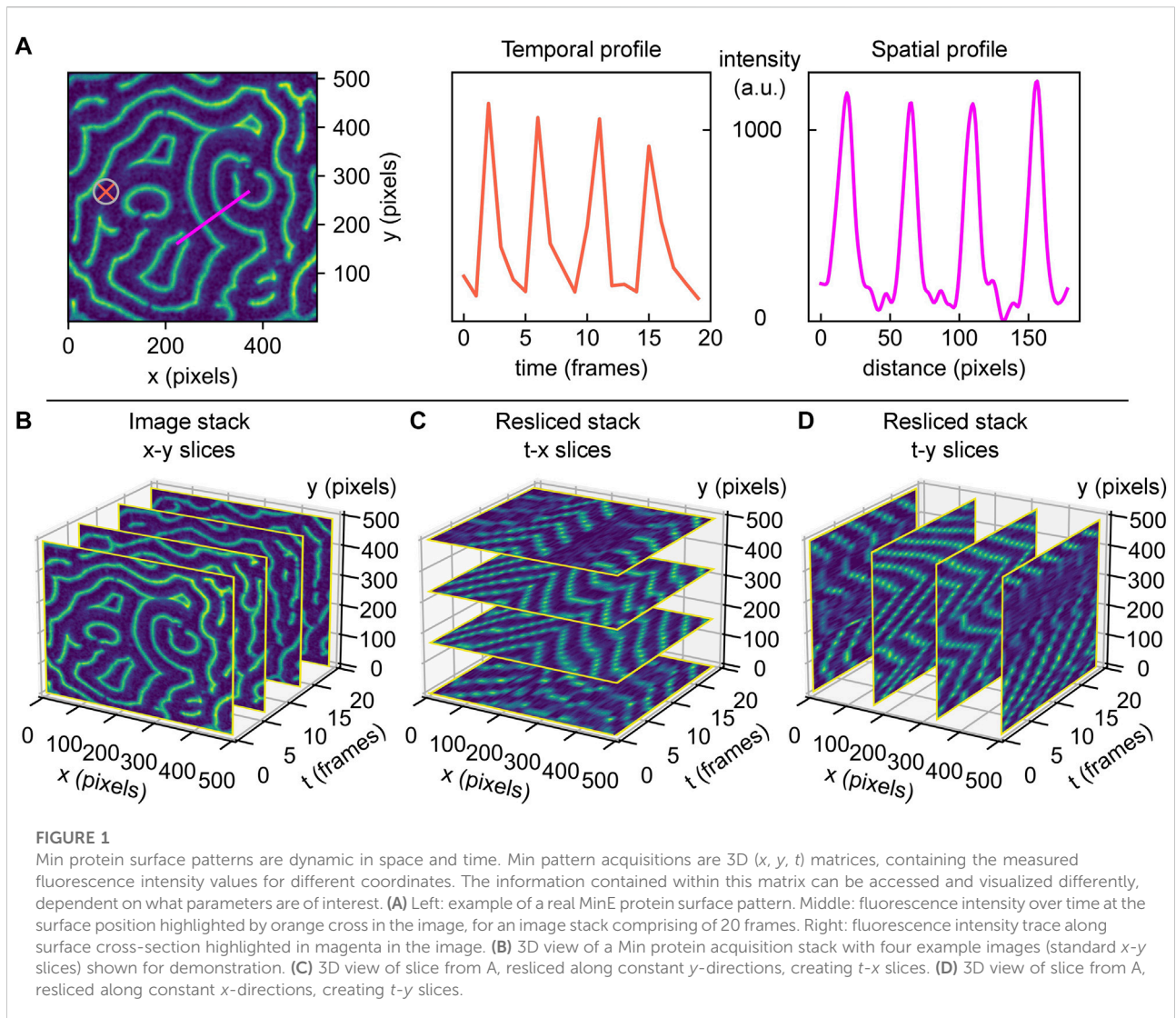
KEYWORDS

min proteins, pattern formation, image analysis, surface protein waves, quantification, python

1 Introduction

Pattern formation is a fascinating basic phenomenon that occurs across many scales, from galaxy formation all the way down to embryology and beyond. In chemistry and biology, self-organizing patterns emerge from the combination of specific intermolecular interactions and molecular transport processes [1]. Acting together, reaction and diffusion can result in inhomogeneous concentrations of molecules that constitute spatiotemporal patterns. Such patterns provide useful functions as they impose directional or positional preferences on processes in cells and tissues [2–5]. Indeed, pattern formation is of vital importance for the description of a multitude of biological phenomena, ranging from bacterial and eukaryotic cell division [4], to embryonic development of multicellular organisms [3, 5], up to entire ecosystems [6].

For example, in *E. coli* bacteria, a pattern-forming mechanism acts to determine the central position of the rod-shaped cell, ensuring that the required protein-machinery is guided to the correct location to start a fully symmetric division into daughter cells. This particular pattern-forming system, formed by the Min proteins, relies on a reaction-diffusion mechanism and is widely considered to be the best-studied model system for intracellular pattern formation [4, 7]. As the moment of cell division approaches, Min proteins will periodically bind and unbind the inner membrane at the poles of the



bacterium. When visualized by fluorescent labelling, the Min proteins are observed to oscillate from pole to pole with a period of approximately 1 minute. As a result, their temporally averaged concentration is minimized at the mid-cell location. One component of the Min system, MinC, exerts an inhibitory effect on a component of the cell division machinery. As the lowest concentration of MinC is found at mid-cell, the proteins that facilitate cell division will preferably bind there, and the cell gets divided evenly [8].

While MinC is important for this downstream process, the pattern as such is formed by only two proteins, MinD and MinE. MinD is an ATPase that upon binding ATP can attach to the membrane, while MinE is its ATPase activator. MinE can bind to the membrane upon recruitment by membrane-bound MinD and subsequently facilitate MinD's ATP-consuming membrane detachment [7, 9–11]. The Min proteins thus constitute a pattern-forming model system with only 2 essential

components, which is appealingly simple for both theoretical and experimental studies. A wide range of experiments have been reported that explore particular features of the Min system [9, 12, 13]. A multitude of Min protein models [14, 15] have been proposed, and continue to be developed as new molecular details are discovered [10, 16–18].

Arguably, the most iconic images of Min protein patterns present themselves in *in vitro* studies of these proteins. Such experiments typically reconstitute Min proteins on supported lipid bilayers on a glass slide, with a fraction of Min proteins carrying a fluorescent label [9]. Min proteins exhibit mesmerizing dynamic membrane patterns in such an artificial *in vitro* environment: Over a wide range of concentrations, one encounters characteristic patterns such as rotating spirals or travelling planar wave fronts, with typical wavelengths in the order of tens of micrometers [9, 12]. An example is given in Figure 1A, showing a snapshot image for labelled MinE.

Examination of this image reveals various features that are typical for Min patterns. For example, Min proteins exhibit planar waves, but these occur only locally and become less correlated on longer length scales. Min patterns thus organize in surface domains with one dominating type of pattern, such as spiral or planar wave fronts travelling in a certain direction. Boundaries between domains are recognizable by phase shifts.

Thorough quantitative analysis of Min patterns is nontrivial. While visual inspection is often sufficient to deduce the basic Min pattern, quantitative information is required to extract trends in the patterns' characteristic parameters, such as the average or local wavelength, oscillation period, or propagation velocity. For perfectly periodic Min patterns with only one type of pattern with a set orientation over the entire imaged surface, analysis can easily be achieved by Fourier analysis. However, this is most often not possible due to the fact that real Min patterns are usually a patchwork of domains separated by phase-shift boundaries. The semi-periodic nature of Min patterns makes image analysis challenging, and so far there are no clear guidelines or standards on how to realise this without user bias. Extraction of quantitative parameters is typically performed by manual selection of individual surface positions [19], line traces [20], or rectangular regions-of-interest [12, 21]. Plotting of fluorescence intensity along a defined axis allows to determine the pattern's local wavelength. Similarly, the temporal characteristics such as its oscillation period and wave propagation velocity can be measured by monitoring the intensity versus time at one spot [19], see [Figure 1A](#) for an illustration of these possible practices.

While these approaches are widely used, such manual selection is cumbersome, prone to user bias, and it underuses the vast amount of data in videos of Min patterns that could improve data accuracy. In the example given in [Figure 1A](#), the intensity profile of denoted line trace (magenta) shows approximately equidistant peaks that indicate the wavelength. However, selection of another spot or a slightly different trace that is not perpendicular to the wave fronts would have led to a deviating result. Furthermore, note how extending the trace over the region of the spiral domain would erroneously lead to the inclusion of shorter or longer wavelengths. Notably, these domain boundaries do not always remain stationary over the time of acquisition, and hence line traces would have to be carefully adapted for every single frame so as to adapt to possible reorientations of wave fronts (compare for example the domain variation over time in [Supplementary Figure S1A](#)). In view of all these limitations, we conclude that an automated user-independent analysis of large regions is preferable, as it allows to obtain solid statistics on the patterns' characteristic features.

In this paper, we present a methodology for analysing Min patterns that aims for thorough quantitation, easy usage, reproducibility, and user independence. We start by briefly proposing strategies for image cleaning. After that, we proceed to presenting strategies for global and local analysis of Min

protein surface patterns. For extracting global parameters, we rely on calculating autocorrelation maps to examine the average periodic features of Min patterns, following other groups [16] as well as our own [18]. We treat a Min pattern acquisition stack as a three-dimensional matrix containing information in space and time (x, y, t) , and we propose strategies on how to efficiently access this information. Subsequently, we introduce an analysis pipeline which allows to quantify local properties of Min patterns, such as the wave propagation speed and direction of propagation. The approach we present here relies on the identification of individual wave crest points and their movement from one frame to the next. While other tools have been used to extract the directional preferences of Min surface wave propagation [22], our approach allows not only for obtaining large distributions of parameters, but also for accessing multiple parameters at the same time. For both global and local analysis, we offer guidance to researchers who would like to implement similar strategies for their own applications. Code is openly available and provided in Python 3 [23].

2 Image processing methodology

Min patterns are phenomena that occur at lipid membranes. Accordingly, the most suitable forms of microscopy for *in vitro* experiments are those that acquire an image along the membrane-coated surface. Examples are total internal reflection fluorescence (TIRF) microscopy [12, 16, 19, 24], laser scanning confocal microscopy [20–22, 25] or spinning disc confocal microscopy [18, 26]. Epifluorescence microscopy [27] is generally not suitable as it leads to high background signal from the fluorescent molecules in the bulk, significantly reducing the pattern's quality or even making it unrecognizable.

“Image cleaning” includes procedures and operations performed on microscope data that improve the overall quality of the image by making the features of interest better recognizable, while not distorting or removing essential components. Good image cleaning, where background and spurious contributions are removed, is important for successful quantitative analysis of Min protein patterns, as insufficient cleaning can lead the algorithms to fail to recognize the pattern as such. There are multiple imaging artefacts that need to be corrected. For example, local fluorescent impurities such as protein aggregates may give a static signal that is not part of the Min pattern. Depending on the microscope, illumination is typically inhomogeneous across the field-of-view. In many cases, researchers are interested in the time-dependent behavior of Min patterns, and for this, they acquire images at the same spatial positions at regular time intervals ranging from seconds to minutes. Consequently, Min proteins may bleach due to prolonged imaging [18]. Flow-cell setups with constant bulk flow may pose additional challenges in the form of objects entering and passing through the field-of-view or adhering to and spontaneously detaching from the surface [26]. We empirically found that the

occurrence of imaging artefacts can vary strongly, depending on the experimental setup (specifically, the type of sample chamber and the surface preparation), protein concentration (higher concentrations may result in more aggregates), and experiment duration (the longer, the more aggregates occur).

With image cleaning, we aim to tackle all of these issues. Here, we propose a sequence of standard actions that we found to yield overall good results for a time series (several frames of the same sample position saved sequentially in an image stack) of images I_{movie} of *in vitro* Min surface patterns. Our protocol involves the following steps:

- Correct for fluorescence bleaching by normalizing each frame to its mean intensity value.
- Create an ‘illumination correction map’ I_{illum} by smoothing and averaging all movie images, and normalizing the images to their maximum intensity.
- Create a static background image I_{stat} by averaging over all the moving (surface pattern) features of all images in the stack. This background image then only contains static fluorescent features such as local specks, holes, and scratches.
- Correct each image via the following image operation: $I_{cor} = (I_{movie} - I_{stat})/I_{illum}$.
- At this point, there may still be some artefacts left, for example, those that appeared during the acquisition time and might thus not have been included in the static background image. Therefore, remaining bright or dark artefacts can be removed manually or by thresholding.
- Images can finally be slightly smoothed to diminish effect of sharp edges and artefacts from the latter cropping.

In cases where it is required to explore Min patterns across very large areas, it may furthermore be of interest to stitch multiple fields-of-view of Min patterns into larger images [26, 28]. If the microscope’s software does not offer an automatized solution for this, stitching can be achieved by good bookkeeping and a few lines of code. When planning to stitch images, it can be helpful to choose individual field-of-views in such a way that there is a bit of an overlap (e.g. 5% of the width) between adjacent areas, as this makes it easier to correctly reassemble the full image afterwards. Note that in general, adjoining areas may not have exactly the same median intensity. Therefore, differences in median intensity levels between individual field-of-views need to be corrected for when assembling the stitched image. This way, the resulting large image will appear more homogeneous in brightness and stitching borders will be less visible.

3 Pattern analysis strategies

3.1 Global parameters

In many cases, researchers are interested in quantifying parameters such as the average spatial wavelength and

oscillation period of a Min pattern. For this purpose, they acquire image stacks of Min patterns at a certain surface region (x, y in pixel or distance units) and at regular time-intervals (t in time units or frames). However, the existence of domains within and dynamics of patterns make manual extraction of these parameters a laborious task, and the results of such analyses (cf. Figure 1A) may suffer from poor statistics and user bias. Here, it is important to realize that what we are interested in are essentially global, image-averaged parameters. In this section, we describe how quantification of such global parameters can be achieved by performing autocorrelation analysis [29] for different slices along time or space coordinates (x - y frames such as in Figure 1B, t - x slices such as in Figure 1C or t - y slices as in Figure 1D). We start with spatial autocorrelation, aiming to quantify the wavelength, and then continue to present how temporal autocorrelation can be used to quantify the oscillation period of a pattern.

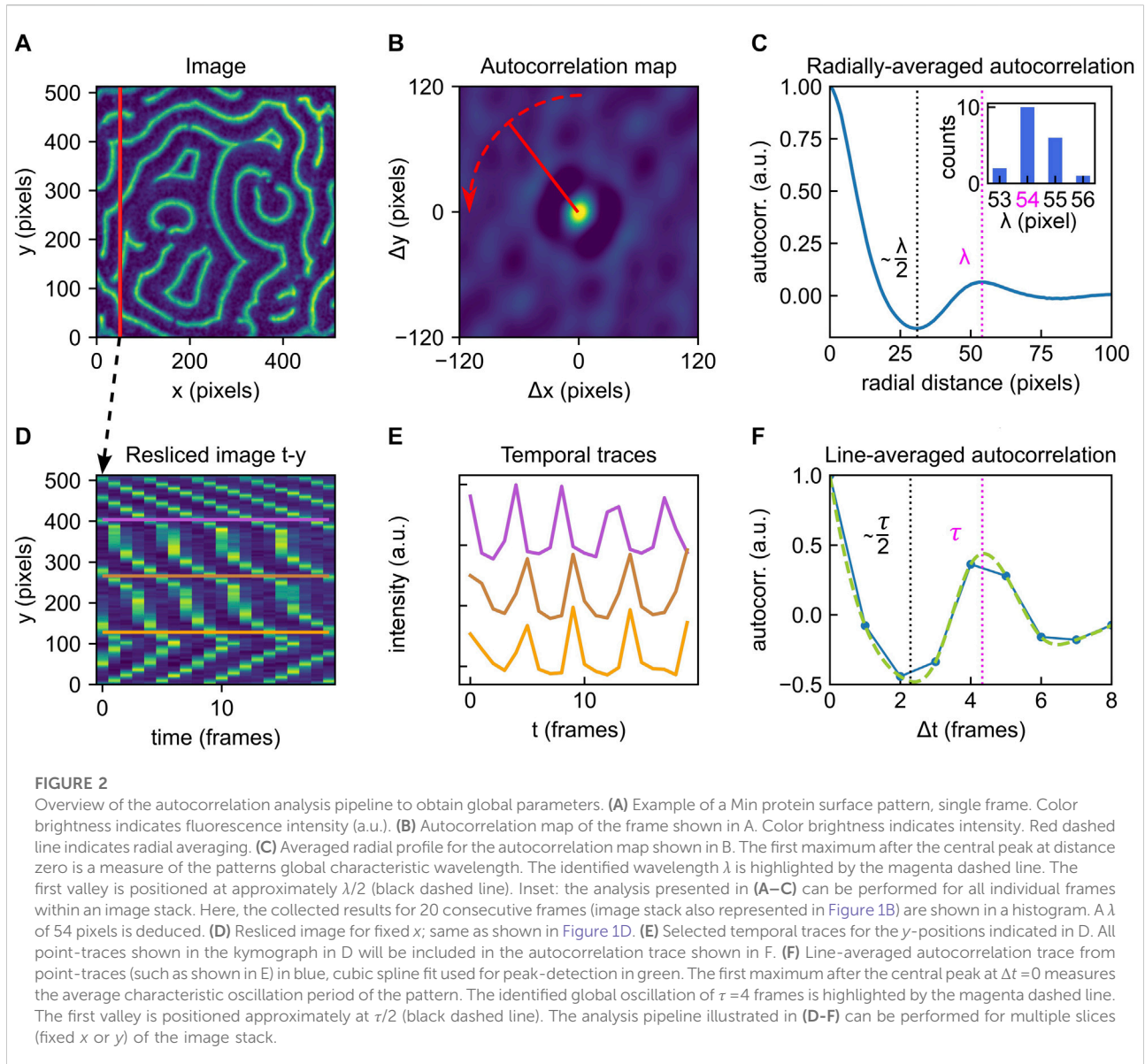
3.1.1 Global wavelength

Spatial autocorrelation analysis of an image essentially compares each pixel to other pixels around it, quantifying their (dis)similarity as a function of distance [29]. Starting from a single image frame as shown in Figure 2A, an autocorrelation map can be calculated using routines from scientific libraries provided for most programming languages. Before performing these operations, the image should be normalized by a series of actions (subtraction of minimum intensity, division by summed-up total intensity and subtraction of mean intensity). Using standard functions supplied within most environments, a twodimensional autocorrelation map $crmx$ can be calculated for an image frame $image$ by the transformation

$$crmx = \text{real}(\text{ifft2}(\text{fft2}(image) \cdot \text{conj}(\text{fft2}(image)))) \quad (1)$$

with fft2 calculating a two-dimensional discrete Fourier Transformation, ifft2 calculating its inverse and real and conj returning the real part and complex conjugate of the input, respectively. The four quadrants of the autocorrelation map can be re-arranged so as to place the position ($\Delta x = 0, \Delta y = 0$) in the center of the image, rather than having four partial peaks in the corners.

Close examination of the autocorrelation map in Figure 2B reveals a general speckled pattern as well as a ring-shaped feature that is present around the center, which is characterized by a peak in intensity. As this central peak corresponds to a distance of zero, its high intensity is a consequence of the self-correlation of each pixel with itself. Moving radially outward from the center, one observes a decrease in intensity (negative autocorrelation) followed by a peak (positive autocorrelation). An easy way to extract the pattern’s dominant wavelength is to perform an angular averaging of the profile around the central peak, where the radial line profiles are averaged over all angles, as illustrated by the red arrow in panel Figure 2B. This leads to a



profile as shown in **Figure 2C**. Here, the first peak after the central peak identifies the pattern's global wavelength λ (magenta dashed line). The first minimum corresponds to the dark value from the autocorrelation map. The valley in fact corresponds to $\sim \lambda/2$ (black dashed line). Note that this value is not exactly half the wavelength, since Min profiles are typically somewhat asymmetrical (cf. **Figure 1A**).

If, as often is the case, Min patterns display steady-state dynamics, then multiple time frames can be used to provide better statistics in estimating the global wavelength. Min patterns are routinely acquired at constant time intervals, which provides a series of sequential image frames, as represented in **Figure 1B**. Calculating and analysing autocorrelation maps for several

frames within such an image stack allows to collect a distribution of wavelengths, such as the histogram shown in the inset in **Figure 2C**, summarising results for 20 consecutive frames. Indeed, for dynamic patterns, we recommend performing this analysis strategy for multiple (e.g. 10) frames and averaging the results to obtain a reliable value for the pattern's spatial wavelength. As Min patterns are typically dynamic, it is also possible that their wavelengths change over time, e.g. due to changes in external parameters. The spatial autocorrelation analysis presented here is however equally applicable to static patterns.

Identifying the peaks shown in **Figures 2C,F**, see following section) is trivial to the human eye, but not necessarily

straightforward to automate. For our data, we achieved this via a simple algorithm that identifies the second local maximum of the trace. In doing so, we implicitly assume that the trace is fine enough to achieve sufficient accuracy and upsampling is not required. While a curve such as shown in [Figure 2C](#) looks smooth to the eye, the original image data had a finite resolution, and accordingly, this curve contains small rags which an algorithm may erroneously detect as a local maximum. In many cases, this issue can be solved by smoothing the curve slightly before subjecting it to peak analysis. This can be achieved by applying a smoothing kernel stretching over a few pixels, for example a small fraction of the image dimension.

To avoid erroneous results, it is recommendable to plot at least one example trace (for one frame) and inspect whether the detected (averaged) wavelength position correctly co-localizes with the valley position. If no clear valley can be identified (characterized by a small or even vanishing intensity difference between the local minimum and maximum), the pattern may lack a recognizable periodicity, which could be an intrinsic property of the given pattern. Alternatively, this issue can also be encountered if insufficient image cleaning has been applied to the stack before analysis and the pattern is superimposed with too many artefacts, resulting in low autocorrelation. However, we find that this method is generally quite robust towards noise, as shown in [Supplementary Figure S2B](#).

Further, the ratio between image size and spatial wavelength is of importance. In [Supplementary Figure S3](#), a large-scale image stack is sequentially cropped to smaller sizes and global autocorrelation analysis as presented here is performed on randomly selected areas. Based on these results, we find that the standard deviation increases as the image sizes gets smaller, and estimate that the image dimension should be large enough to contain at least 5–10 full wavelengths for the presented algorithm to provide reliable results.

3.1.2 Global oscillation period

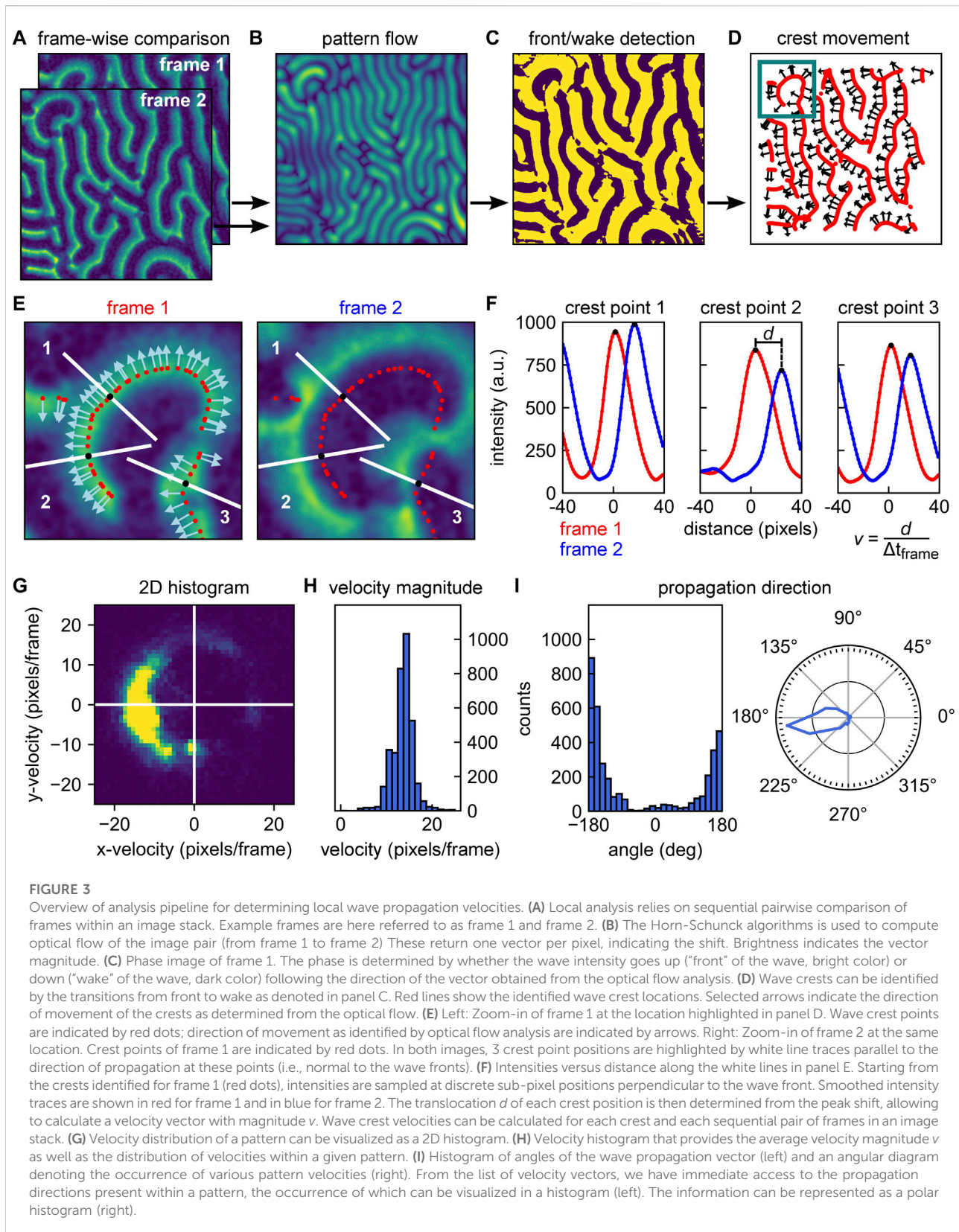
Another global property of interest is the average oscillation period. This parameter describes how at any given point within the imaged region, the membrane protein density (which is proportional to the fluorescence intensity) can be expected to change over time. We quantify the global oscillation period of a Min pattern following a strategy that closely resembles the one presented for obtaining the global wavelength in the preceding section.

To obtain quantitative information on the temporal dynamics of Min patterns, they are typically acquired as time series. A time series such as the one shown in [Figure 1B](#) is essentially a three-dimensional (x, y, t) matrix, containing information on fluorescence intensity as a function of space and time. Notably, for temporal analysis to be reliable, image data has to be acquired at a rate above the Nyquist rate. Min patterns are characterized by point-wise temporal periodicity (compare

the time evolution at a random sample position in [Figure 1A](#)). Hence, when acquiring a time series, intervals have to be chosen short enough to ensure the acquisition of enough data to adequately depict the pattern's temporal dynamics. If consecutive acquisition times in a series are chosen very far apart, the pattern may even appear to travel towards the opposite direction. According to Nyquist theorem, for patterns of an oscillation period τ , consecutive images in a series have to be acquired at intervals spaced no longer than $\tau/2$ apart. In principle, the shorter these intervals are, the better the temporal dynamics can be characterized. Min protein patterns in lipid bilayer reconstitutions will typically exhibit wave propagation velocities of several 100 nm s^{-1} [12] and wavelengths around $50 \mu\text{m}$ [9]. This leads to an expected oscillation period in the order of minutes. Acquiring sufficient images to fulfill Nyquist theorem should therefore not be an issue in most microscope setups. In practice, finding the optimal imaging rate is a trade-off between achieving a high temporal resolution and minimizing fluorescence bleaching.

Upon analysing a given Min pattern stack with respect to its temporal characteristics, we found that reslicing the stack along fixed spatial coordinates simplifies data processing and makes it more straightforward to access the temporal information contained within the stack. [Figure 2D](#) shows a (t, y) slice through our example image stack at fixed pixel position x . These t - x or t - y slices are collections of kymographs for all points along the cross-section at the fixed spatial coordinate. In [Figure 2E](#), we show example traces of these kymographs along selected y positions, which then correspond to individual surface locations (x, y) on the field-of-view shown in [Figure 2A](#). Analogous to the strategy presented for spatial global analysis, [Eqn. 1](#) can be used to obtain an autocorrelation map for this slice (cf. [Supplementary Appendix DEMO_MinDE_global_analysis](#)). To extract the pattern's global oscillation period, we consider the autocorrelation map's trace at $\Delta y = 0$ for $\Delta t \geq 0$, plotted in [Figure 2F](#). Including the $\Delta t \leq 0$ trace (going back in time) of the curve would not yield additional resolution, as it contains the same information as the positive trace. This profile then immediately provides the averaged temporal autocorrelation trace for all points along one fixed spatial coordinate (as indicated for fixed x by the vertical line in [Figure 2A](#)). If the line-averaged autocorrelation trace does not provide sufficiently high resolution to identify a minimum indicative of the oscillation period, imaging over a longer time and/or at shorter intervals could be necessary.

The first maximum of this curve beyond the central peak again identifies the predominant global oscillation period τ (magenta dashed line), while the first minimum is located at approximately $\tau/2$ (black dashed line). Statistics on τ can be collected by performing the analysis pipeline illustrated in [Figures 2D–F](#) for multiple cross-sections at fixed y or x , that is, for multiple resliced frames as shown in [Figures 1C,D](#). Compared to the traces obtained from spatial autocorrelation,



which are calculated via radial averaging, these temporal autocorrelation traces may appear rather coarse. A more advanced peak detection can be performed with cubic-spline fitting and subsequent extreme point analysis. Further, the presented method for temporal autocorrelation analysis is less robust towards noise than spatial autocorrelation, as shown in [Supplementary Figure S2C](#). Adequate image preprocessing and cleaning is again of high importance. Image smoothing was found to slightly increase the detected oscillation period, albeit not above the error margins obtained for unsmoothed images, as shown in [Supplementary Figure S4](#).

Note furthermore that the propagation speed of a pattern may depend on the concentrations of all components, and may change over time. Therefore, for very long time series, it can make sense to split the stack up into multiple stacks of shorter total duration and perform analysis for subsequent sub-stacks (compare [Supplementary Figure S8](#)). However, we estimate that an image stack needs to span the duration of a minimum length of 3 oscillation periods for the presented method to yield reliable results (cf. [Supplementary Figure S4](#)).

3.2 Local parameters

In this section, we present an analysis strategy that aims to quantify local parameters of Min patterns. These parameters include the local wave propagation velocity or characteristic local distances such as the distance between MinD and MinE wave crests. As we will describe in more detail, local parameters can be quantified by implementing two steps: 1) identification of wave crests for each frame within an image stack, and 2) comparison of intensities at and around the identified crest points from one frame to the next.

3.2.1 Local wave propagation velocity

The local wave propagation velocity is determined as the shift of individual crest points from one frame to the next. To calculate the local velocity, we first need to identify these crest points, and then determine their positional difference. We achieve this by applying optical flow analysis. Optical flow is defined by how brightness patterns move from one image to another [30, 31]. For our Min patterns, we estimate an optical flow vector field using Horn-Schunck algorithm [30, 32]. This algorithm works on a pair of sequential image frames (see [Figure 3A](#) for example data, recorded for MinE) and returns a vector field, describing the estimated optical flow for each pixel from the first to the second frame. [Figure 3B](#) shows the magnitude (indicated by image brightness) of the optical flow vectors obtained for our example image pair from [Figure 3A](#). Note that the collected optical flow vectors do not necessarily represent the local wave propagation velocities. The relation between the two measures will depend on the wave shape, making it hard to generalize.

As a first step to identifying the positions of the wave crests within the first frame, we use the information obtained from

optical flow analysis to create a phase image. This phase image is essentially a binary map over the whole image, which indicates whether at any given point we are at the “front” (‘ahead of the wave’) or “wake” region of the wave (‘behind the wave’). These regions are characterized by a decrease or increase in fluorescence intensity, respectively, along the direction of movement. If we consider any specific surface location and its corresponding vector obtained from optical flow, we can follow the direction of that vector over a short distance (say 3 pixels) within the same frame, and determine whether the fluorescence intensity is decreasing or increasing. The outcome of this procedure for our example frames is shown in [Figure 3C](#). Using this phase image, we can identify those positions in the image that make up the wave crests – which form the backbone structure of the pattern. Wave crests are identified by the positions at which a transition from the front to a wake region occurs. [Figure 3D](#) shows the wave crests identified for the first frame in red. Subsequently, we need to determine the direction of wave propagation for these individual crest points. While we have a set of vectors available for distinct (x, y) positions from optical flow analysis, their directions will in general not be exactly normal to the wave crests due to noise. Therefore, we calculate vectors which are perpendicular to the lines of wave crests, and determine their correct directionality (pointing towards the front of the wave) using the information already present in the phase image. In [Figure 3D](#) the direction of propagation is indicated by vectors at distinct crest points throughout the image for illustration. Following the procedure outlined here and in [Figures 3A–D](#), we thus effectively obtained 1) a list of positions, identifying the wave crest point positions, and 2) a list of vectors, describing their direction of movement from the current to the next frame.

The next step is to quantify how far these crest points moved during the transition from the current to the next frame. Upon quantifying this positional shift, the velocity magnitude is given by this distance divided by the time that passed between the acquisition of subsequent frames. The procedure is illustrated in [Figures 3E,F](#) for three example crest points. For each crest point, we considered the fluorescence intensity traces along the direction of movement at the given position (i.e., perpendicular to the wave front). Starting from the crest position, we plotted this trace up to a distance corresponding to roughly half the (global) wavelength of the pattern. Choosing this distance should ensure that the next frame’s peak is still within the sampling range (provided that the time resolution of the acquisition stack is sufficiently high, cf. [Section 3.1.2](#)). In [Figure 3E](#), the example traces along which intensities are plotted are highlighted in white. For sampling positions along this trace that are located off the pixel grid, intensities can be estimated by interpolation. This yields an intensity trace at sub-pixel resolution. This intensity trace will have a maximum located very close to the center of the originally determined crest position, with a small offset that can be attributed to the

additional smoothing step performed before optical flow analysis, and is generally not significant (cf. [Supplementary Figure S5](#) and adjoined table).

Next, we performed exactly the same procedure at the same positions on the next frame. As shown in [Figure 3E](#) and [Figure 3F](#), these positions are now no longer centered around the maximum of the intensity trace, but offset from them by a certain distance. This distance is the positional shift that we need to obtain for quantification of the local wave propagation velocity. Performing peak fitting allows to identify the two traces' peak values and measure the shift d in crest peak position at sub-pixel resolution.

Performing this analysis for all identified crest points along all sequential pairs of frames in a stack eventually yielded a list of velocity vectors with components (v_x, v_y) at frame n for a wave crest at position (x, y) . Image smoothing is an important technical aspect of the local analysis presented here. We first perform a light image smoothing (with a limited initial smoothing kernel) to de-noise the image and to simplify peak detection such as shown in [Figure 3F](#). For optical flow analysis ([Figure 3B](#)), we perform an additional smoothing step with a more extensive secondary smoothing kernel. The effect of different initial smoothing kernels is illustrated in [Supplementary Figure S6](#), the effect of increasing image noise in [Supplementary Figure S2D,E](#). Further, applying Horn-Schunck algorithm [30] requires setting a few parameters, particularly the regularization constant and the number of iterations. Here, we determined their values by trial-and-error from evaluating simulated as well as real Min protein pattern data and kept both constant from then on.

Depending on what kind of information is of interest, the obtained distribution of velocity vectors can be represented in different ways. A representation that combines information on the magnitude and directional preference is a 2D histogram such as the one shown in [Figure 3G](#) for an example image stack of MinE data. If the magnitude of the wave propagation velocity is most important, the velocity magnitudes v can be plotted in a histogram to visualize their distribution and peak value, as shown in [Figure 3H](#). In other cases, the directional information may be of prime interest, for example, when studying the influence of an external bulk flow [26] or physical environmental parameters on the pattern [22]. For each vector component, an angle α can be calculated that corresponds to the wave crest point's direction of movement with respect to the direction of applied flow. The occurrence of different directions can then be plotted in a regular or polar histogram, as shown in [Figure 3I](#). As shown [Supplementary Figure S7](#), the method can be applied to a wide range of image sizes. Decreasing the area on which analysis is performed generally leads to smaller standard deviation in velocity magnitude distribution, which is linked to a loss in directional information. The full local velocity pipeline is also presented in the [Supplementary Appendix DEMO_MinDE_local_analysis](#).

3.2.2 Local MinD/MinE crest shift

When studying Min protein patterns, researchers routinely acquire data in two channels, for fluorescently-labelled MinD and MinE, respectively. Both these proteins exhibit a peak in their surface density, but these two peaks do not coincide in space and time (an example is given in [Figure 4A](#)).

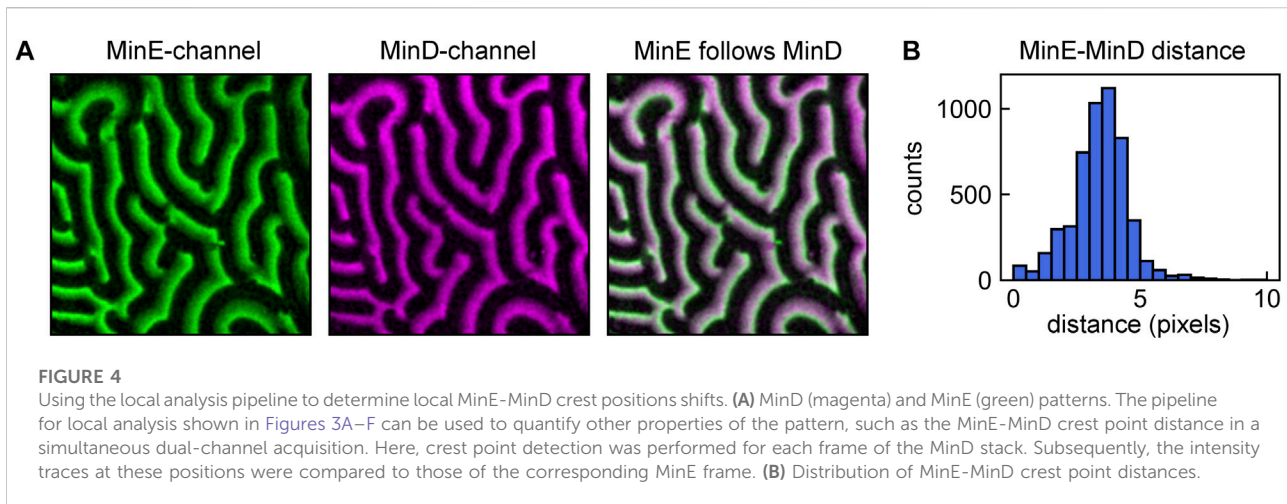
To determine these differences, the analysis pipeline presented in [Section 3.2.1](#) for obtaining wave propagation velocity between sequential frames can be slightly adapted for measuring the difference in crest position between MinD and MinE waves. This can be achieved by first performing crest detection (analysis pipeline shown in [Figures 3A–D](#)) for one stack, e.g. the MinD stack. The crest points attributed to the MinD stack can then be compared to the corresponding frame of the MinE stack, analogous to the comparison shown in [Figure 3E](#). The collected peak shifts d can be visualized in a histogram, such as shown in [Figure 4B](#) for our example data.

4 Discussion

In this paper, we proposed protocols that can be applied to analyse data on *E. coli* Min protein surface patterns with respect to quantitative parameters such as the spatial wavelength, oscillation period, or wave crest propagation velocities. Our primary goal was to develop user-independent and largely automatized strategies for pattern analysis in order to 1) improve comparability of results obtained from different studies and research groups, and 2) provide tools for quantification of experimental data for comparing results to theoretical studies. We presented global and local analysis methods which both can be valuable depending on what kind of pattern information is of interest. All of the routines that we suggested and described can be further modified to better fit the requirements posed by a particular research question – thus providing a framework that others can build upon.

The methodology we describe can be combined with many of the aspects and practices of Min pattern analysis which we did not cover here. For example, in studies investigating absolute or relative local concentrations or concentration gradients of MinD and MinE on the surface, fluorescence calibration can be important. Calibration allows to relate fluorescence intensity to the local membrane protein density [12, 16, 19, 21]. However, in many cases the patterns as such – their shapes, domains, wavelength, or dynamics – can be studied without this step, using the acquired fluorescence intensity data in arbitrary units, and this is what we focused on in this paper.

The global analysis that was presented here provides parameters averaged over entire image frames (spatial analysis) or slices along fixed spatial coordinates (temporal analysis). We found that for well-cleaned images, spatial autocorrelation robustly quantifies the main dominant wavelength within an image frame,



as identified by the distance at which maximum self-correlation is present. This yielded results that are very comparable to those obtained by manual quantification, albeit with higher statistical reliability (see [Supplementary Figure S1](#) for an example comparison of manual and automatic analysis). For temporal autocorrelation, we recommend re-orienting the image stack into slices of fixed x - or y -coordinates and calculating autocorrelation-maps along sets of kymographs, as described. We found this to be an efficient way to access large parts of the information provided within the image stack. Pixelwise analysis of intensity traces at distinct surface location, autocorrelation thereof and subsequent averaging is a more straightforward approach. However, we found that taking this detour via reslicing as presented in this paper reduces computational time by about two orders of magnitude, and hence, allows for easier processing of large amounts of data (cf. SI).

In many cases, parameters obtained from spatial and temporal analysis can be averaged for several frames/slices. However, whether this is justified depends on the time evolution of the pattern over the course of the acquisition. [Supplementary Figure S8](#) shows the temporal autocorrelation results for an example image stack which shows a decline in oscillation period over time. This is visible from the spreading of peaks in the resliced image (cf. [Supplementary Figure S8A](#)) as well as in the variation in the shape of the $\Delta x = 0$ traces (cf. [Supplementary Figure S8B](#)). In such a case, averaging is only justified over the part of the stack that shows a constant temporal periodicity. If required, the routines can be expanded to include peak analysis of single traces in the kymograph set and automatically determine up to which point averaging is permitted.

Local analysis, in contrast to global analysis, directly provides large-number distributions of parameters even when analysing only two consecutive images. Our approach to local analysis relies on the identification of wave crests and quantification of the positional shift thereof from one frame to the next. The first step can in principle be achieved by various strategies. For example, one could implement an algorithm that relies on thresholds of fluorescence intensity to

identify peaks. Here, we presented a strategy that relies on established algorithms for optical flow analysis, as it offers certain advantages, in particular access to directional information. We have already successfully used this local analysis methodology to examine and quantify the response of Min protein patterns to hydrodynamic bulk flow [26]. Numerous other applications can be imagined, in which a surface pattern is either exposed to an external stimulus and responds to it, or in which the time evolution of its characteristic properties is relevant. Using the local analysis pipeline presented here (or modifications thereof) allows to systematically quantify properties of interest with proper statistics rather than relying on qualitative description alone.

As our local analysis routine essentially offers a way to acquire a high number of intensity traces that are locally perpendicular to the wave fronts, it opens up possibilities for further in-depth analysis. Our brief illustration of a MinD-MinE crest distance detection is only one example of a parameter that can be extracted using an optical-flow-based routine. For example, it is imaginable to modify our local analysis pipeline such that it facilitates fast visualization of local fluorescence intensity profiles for both MinD and MinE waves. Upon fluorescence calibration, the fluorescence intensity can be converted to membrane protein density, and hence the relative peak heights can be connected to the protein ratio at the surface. Further, information obtained from local and global analysis can be combined, and local velocity analysis can provide an alternative to global methods if their requirements are not met. For example, while we found that a certain minimum number of entire oscillations is required for global temporal analysis to yield results (cf. [Supplementary Figure S4](#)), local analysis can already be performed on only two consecutive frames, and can be combined with global spatial analysis to serve as an alternative route to determining the predominant oscillation period.

Finally, we note that the general strategies presented within this paper are not restricted to application to the analysis of *E. coli* Min protein surface patterns. Min protein patterns have some properties that are very specific to them – such as their quasi-

periodic nature, dynamic oscillatory behavior, and their organization within domains – and our routines are designed for these specific properties. However, the routines presented here may very well be modifiable and applicable to quantify other types of semi-periodic surface patterns, biological or not, for example to the fascinating reaction-diffusion patterns of chemical systems like the BZ-AOT system [33]. We hope that our analysis toolbox will help to further disentangle the intriguing mechanisms that underlie pattern formation.

Data availability statement

The code created for this study as well as example data are openly available and can be found on Zenodo at zenodo.org/record/6724666 (second release). Please refer to the [Supplementary Material](#) for further information on how to access and use the analysis tools.

Author contributions

SM performed experiments with Min proteins, contributed to the provided image analysis tools and their representation, created the figures and co-wrote the paper. JK identified suitable image analysis methods, developed the majority of the provided tools and co-wrote the paper. CD co-developed the underlying methodology and co-wrote the paper.

Funding

Financial support was provided by the Netherlands Organisation for Scientific Research (NWO/OCW) as part of

References

1. Turing A. The chemical basis of morphogenesis. *Philos Trans R Soc Lond Ser B, Biol Sci* (1952) 237:37–72. doi:10.1098/rstb.1952.0012
2. Bois JS, Jülicher F, Grill SW. Pattern formation in active fluids. *Phys Rev Lett* (2011) 106:028103. doi:10.1103/PhysRevLett.106.028103
3. Kondo S, Miura T. Reaction-diffusion model as a framework for understanding biological pattern formation. *Science* (2010) 329:1616–20. doi:10.1126/science.1179047
4. Halatek J, Brauns F, Frey E. Self-organization principles of intracellular pattern formation. *Phil Trans R Soc B* (2018) 373:20170107. doi:10.1098/rstb.2017.0107
5. Schweisguth F, Corson F. Self-organization in pattern formation. *Develop Cell* (2019) 49:659–77. doi:10.1016/j.devcel.2019.05.019
6. Rietkerk M, van de Koppel J. Regular pattern formation in real ecosystems. *Trends Ecol Evol* (2008) 23:169–75. doi:10.1016/j.tree.2007.10.013
7. Ramm B, Heermann T, Schwille P. The *E. coli* MinCDE system in the regulation of protein patterns and gradients. *Cell Mol Life Sci* (2019) 76:4245–73. doi:10.1007/s00018-019-03218-x
8. Kretschmer S, Schwille P. Pattern formation on membranes and its role in bacterial cell division. *Curr Opin Cell Biol* (2016) 38:52–9. doi:10.1016/j.cob.2016.02.005

the NanoFront and BaSyC programs, as well as by ERC Advanced Grant no. 883684.

Acknowledgments

We thank Grzegorz Pawlik for providing example Min data, Julian Hofer for support with Python coding, and Alberto Blanch Jover, Angeliki Goutou, Ashmiani van den Berg, Eli van der Sluis, Erwin Frey, Fridtjof Brauns, Jaco van der Torre, Jernej Rudi Finžgar and Nicola De Franceschi for helpful discussions.

Conflict of interest

The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

Publisher's note

All claims expressed in this article are solely those of the authors and do not necessarily represent those of their affiliated organizations, or those of the publisher, the editors and the reviewers. Any product that may be evaluated in this article, or claim that may be made by its manufacturer, is not guaranteed or endorsed by the publisher.

Supplementary material

The Supplementary Material for this article can be found online at: <https://www.frontiersin.org/articles/10.3389/fphy.2022.930811/full#supplementary-material>

9. Mizuuchi K, Vecchiarelli AG. Mechanistic insights of the Min oscillator via cell-free reconstitution and imaging. *Phys Biol* (2018) 15:031001. doi:10.1088/1478-3975/aa9e5e
10. Wettmann L, Kruse K. The min-protein oscillations in *Escherichia coli*: An example of self-organized cellular protein waves. *Phil Trans R Soc B* (2018) 373:20170111. doi:10.1098/rstb.2017.0111
11. Frey E, Halatek J, Kretschmer S, Schwille P. Protein pattern formation. In: Bassereau P, Sens P, editors. *Physics of biological membranes*. Cham: Springer International Publishing (2018). p. 229–60. doi:10.1007/978-3-030-00630-3_10
12. Vecchiarelli AG, Li M, Mizuuchi M, Mizuuchi K. Differential affinities of MinD and MinE to anionic phospholipid influence Min patterning dynamics *in vitro*: Flow and lipid composition effects on Min patterning. *Mol Microbiol* (2014) 93:453–63. doi:10.1111/mmi.12669
13. Heermann T, Steiert F, Ramm B, Hundt N, Schwille P. Mass-sensitive particle tracking to elucidate the membrane-associated MinDE reaction cycle. *Nat Methods* (2021) 18:1239–46. doi:10.1038/s41592-021-01260-x
14. Meinhardt H, de Boer PAJ. Pattern formation in *Escherichia coli*: A model for the pole-to-pole oscillations of min proteins and the localization of the division site. *Proc Natl Acad Sci U S A* (2001) 98:14202–7. doi:10.1073/pnas.251216598

15. Kruse K, Howard M, Margolin W. An experimentalist's guide to computational modelling of the Min system. *Mol Microbiol* (2007) 63:1279–84. doi:10.1111/j.1365-2958.2007.05607.x
16. Vecchiarelli AG, Li M, Mizuuchi M, Hwang LC, Seol Y, Neuman KC, et al. Membrane-bound MinDE complex acts as a toggle switch that drives Min oscillation coupled to cytoplasmic depletion of MinD. *Proc Natl Acad Sci U S A* (2016) 113:E1479–E1488. doi:10.1073/pnas.1600644113
17. Denk J, Kretschmer S, Halatek J, Hartl C, Schwille P, Frey E, et al. MinE conformational switching confers robustness on self-organized Min protein patterns. *Proc Natl Acad Sci U S A* (2018) 115:4553–8. doi:10.1073/pnas.1719801115
18. Brauns F, Pawlik G, Halatek J, Kerssemakers J, Frey E, Dekker C, et al. Bulk-surface coupling identifies the mechanistic connection between Min-protein patterns *in vivo* and *in vitro*. *Nat Commun* (2021) 12:3312. doi:10.1038/s41467-021-23412-5
19. Ivanov V, Mizuuchi K. Multiple modes of interconverting dynamic pattern formation by bacterial cell division proteins. *Proc Natl Acad Sci U S A* (2010) 107:8071–8. doi:10.1073/pnas.0911036107
20. Ramm B, Glock P, Mücksch J, Blumhardt P, García-Soriano DA, Heymann M, et al. The MinDE system is a generic spatial cue for membrane protein distribution *in vitro*. *Nat Commun* (2018) 9:3942. doi:10.1038/s41467-018-06310-1
21. Loose M, Fischer-Friedrich E, Herold C, Kruse K, Schwille P. Min protein patterns emerge from rapid rebinding and membrane interaction of MinE. *Nat Struct Mol Biol* (2011) 18:577–83. doi:10.1038/nsmb.2037
22. Zieske K, Schweizer J, Schwille P. Surface topology assisted alignment of Min protein waves. *FEBS Lett* (2014) 588:2545–9. doi:10.1016/j.febslet.2014.06.026
23. Van Rossum G, Drake FL. *Python 3 reference manual*. Scotts Valley, CA: CreateSpace (2009).
24. Vecchiarelli AG, Li M, Mizuuchi M, Ivanov V, Mizuuchi K. MinE recruits, stabilizes, releases, and inhibits MinD interactions with membrane to drive oscillation. *Microbiology* (2017). doi:10.1101/109637
25. Ramm B, Goychuk A, Khmelinskaia A, Blumhardt P, Eto H, Ganzinger KA, et al. A diffusio-phoretic mechanism for ATP-driven transport without motor proteins. *Nat Phys* (2021) 17:850–8. doi:10.1038/s41567-021-01213-3
26. Meindlhumer S, Brauns F, Finžgar J, Kerssemakers J, Dekker C, Frey E. Directing min protein patterns with advective bulk flow. *BioRxiv* (2021). doi:10.1101/2021.12.23.474007
27. Caspi Y, Dekker C. Mapping out Min protein patterns in fully confined fluidic chambers. *eLife* (2016) 5:e19271. doi:10.7554/eLife.19271
28. Würthner L, Brauns F, Pawlik G, Halatek J, Kerssemakers J, Dekker C, et al. Bridging scales in a multiscale pattern-forming system. *arXiv:2111.12043 [nlin, physics:physics]* (2021). ArXiv: 2111.12043.
29. Robertson C. Theory and practical recommendations for autocorrelation-based image correlation spectroscopy. *J Biomed Opt* (2012) 17:080801. doi:10.1117/1.JBO.17.8.080801
30. Horn BKP, Schunck BG. Determining optical flow. *Artif Intelligence* (1981) 17:185–203. doi:10.1016/0004-3702(81)90024-2
31. Mesbah M. Gradient-based optical flow: A critical review. In: *Isspa '99. Proceedings of the fifth international symposium on signal processing and its applications (IEEE cat. No.99EX359)*, 1. Brisbane: Queensland Univ. Technol (1999). p. 467–70. doi:10.1109/ISSPA.1999.818213
32. [Dataset] Michael I. *scivision/pyoptflow: update implementation, compare 3 implementations* (2021). doi:10.5281/zenodo.5574844
33. Vanag VK, Epstein IR. Pattern formation mechanisms in reaction-diffusion systems. *Int J Dev Biol* (2009) 53:673–81. doi:10.1387/ijdb.072484v