Morphometrics of complex cell shapes: Lobe Contribution Elliptic Fourier Analysis (LOCO-EFA)

Yara E. Sánchez-Corrales\textsuperscript{1,3}, Matthew Hartley\textsuperscript{1}, Jop van Rooij\textsuperscript{1,2}, Athanasius F. M. Marée\textsuperscript{1,4}, Verônica A. Grieneisen\textsuperscript{1,4}

\textsuperscript{1} Computational and Systems Biology, John Innes Centre, Norwich NR4 7UH, UK.
\textsuperscript{2} Theoretical Biology/Bioinformatics, Dept. of Biology, Utrecht University, Padualaan 8, 3584 CH Utrecht, The Netherlands.
\textsuperscript{3} Current address: MRC-Laboratory of Molecular Biology, Cambridge Biomedical Campus, Francis Crick Avenue, Cambridge CB2 0QH, UK
\textsuperscript{4} To whom correspondence should be addressed. E-mail: Stan.Maree@jic.ac.uk, Veronica.Grieneisen@jic.ac.uk

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Abstract

Quantifying cell morphology is fundamental to the statistical study of cell populations, and can help unravel mechanisms underlying cell and tissue morphogenesis. Current methods, however, require extensive human intervention, are highly parameter sensitive, or produce metrics difficult to interpret biologically. We therefore developed a method, Lobe Contribution Elliptical Fourier Analysis (LOCO-EFA), which generates from digitalised 2D cell outlines meaningful descriptors that can be directly matched to morphological features. This is shown by studying well-defined geometric shapes as well as actual biological cells from plant and animal tissues. LOCO-EFA provides a tool to phenotype efficiently and objectively populations of cells, here demonstrated by applying it to the complex shaped pavement cells of Arabidopsis thaliana wild type and speechless leaves, and Drosophila amnioserosa cells. To validate our method’s applicability on large populations, we analysed computer-generated tissues. By controlling in silico cell shape, we explored the potential impact of cell packing on individual cell shape, quantifying through LOCO-EFA deviations between the specified shape of single cells in isolation and the resultant shape when they interact within a confluent tissue.
**Summary statement**

Novel method (LOCO-EFA) quantifies complex cell shapes, extracting meaningful biological features such as protrusion number and amplitude; here shown for complex plant, animal and *in silico* cells.

**Introduction**

Cell geometry has long fascinated biologists (Thompson, 1917). This interest is driven by a wide range of underlying scientific questions. For instance, cell shape changes can be linked to physiological responses of cells, such as membrane protrusions during apoptosis and migration (Charras and Paluch, 2008), and can underlie cell behaviour, such as chemotaxis (Driscoll et al., 2012; Keren et al., 2008). It plays a key role in tissue morphogenesis during development (Lecuit and Lenne, 2007; Sherrard et al., 2010) and in homeostasis (Marinari et al., 2012; Veeman and Smith, 2013). Cell shape influences intracellular processes such as microtubule organisation (Ambrose et al., 2011; Gomez et al., 2016) and stress patterns in plant epithelia (Sampathkumar et al., 2014); it indirectly positions the plane of cell division (Bessen and Dumaîs, 2011; Minc et al., 2011) and can even determine how a flower attracts pollinators (Noda et al., 1994). Given the rich diversity of processes in which cell shape plays a decisive role, either actively or passively, cell morphometrics, the qualitative and quantitative study of cell shape characteristics, is becoming very important for developmental biology. In parallel, advances in imaging technology and software allow us to collect remarkable amounts of cell morphological data, which in turn calls for analytic tools to enable extracting meaningful cell shape information (Zhong et al., 2012). In stark contrast to the technological advances in imaging, there are relatively few automatic and quantitative tools available to analyse complex cell shapes (Ivakov and Persson, 2013; Ljosa et al., 2012; Rajaram et al., 2012). This gap reflects the non-trivial nature of this task: cell shape is often irregular and variable, making it very difficult to establish universal criteria encompassing cell geometry.

To illustrate the issues involved in quantitatively capturing complex cell shapes, we consider pavement cells (PCs) in the plant epidermis (Figure 1A, B) and amnioserosa cells in the Drosophila embryo (Figure 1C). PCs present a striking development, requiring multiple locally divergent growth fronts within each cell that are coordinated amongst neighbouring cells. Amnioserosa cells dynamically change their complex cell shape within a confluent tissue. Both cell types present challenges for quantifying cell shape: 1) their complex, non-holomorphic geometries cannot be captured in a meaningful way with traditional shape metrics; and 2) lack of recognisable landmarks excludes a myriad of shape analysis methods, such as Procrustes analysis (Klingenberg, 2010).

Traditional metrics for cell morphology include area, perimeter, aspect ratio and form factor. Useful as general descriptors, they deliver limited shape information. Very different shapes may yield a similar aspect ratio or form factor (Figure 1D–H). Besides not being unique, such descriptors tend to omit information regarding
biologically relevant shape features (Fu et al., 2005). Several approaches to quantify complex cell shapes are summarised in Table 1. Some of those methods, such as the skeleton method, are highly sensitive to image noise as well as to the precise choice of parameters (for an example, see Le et al., 2006). Other metrics, such as lobe length and neck width (Fu et al., 2005), require humans to judge what a lobe is, which strongly impacts the quantitative results (Figure 1 and Figure S1). It renders these metrics highly variable from cell to cell, from phenotype to phenotype and from human to human. To avoid such dependencies, an automatic method, LobeFinder, was developed to count lobes and indentations (Wu et al., 2016). This method, however, is less adapted to irregular cell shapes and estimates lobe numbers which do not closely correspond to those defined by human inspection (Figure 1). Moreover, it finds its limitations when the characteristics of a shape reside in the distribution and amplitude of the lobes, rather than in their number. For instance, some Arabidopsis mutants present PCs that are more elongated or have shallower lobes, but which occur at a similar spatial frequency (Lin et al., 2013). Recognising the need of automatic and non-biased quantification for PCs, Möller et al. (2017) developed PaCeQuant, a software to define lobes and necks in a systematic way based on local curvature. Alike LobeFinder, it is highly sensitive to small variations in the shape contour, with the sampling density of the contour biasing the local curvature estimation.

Promising alternatives are methods that consider the full cell outline, reducing it into a series of coefficients that can be employed as shape descriptors in a multivariate study (Ivakov and Persson, 2013; Pincus and Theriot, 2007). Elliptical Fourier Analysis (EFA) is such a method, used to quantify two-dimensional complex shapes (Diaz et al., 1989; Kuhl and Giardina, 1982; Schmittbuhl et al., 2003). In this method, the contour’s coordinates are decomposed into a series of related ellipses (described by EFA coefficients), which can be combined to reconstitute the original shape. Despite its wide usage in morphometric studies, EFA cannot retrieve information that directly relates to morphological features of a cell, obstructing biological interpretation. This is because the same outline can be represented by infinitely many different sets of EFA coefficients, depending on how the cell outline is approximated, and because there is no one-to-one relationship between EFA modes and the number of morphological features (see supplementary Materials and Methods for further details).

Here we present a new method based on EFA, termed Lobe-Contribution Elliptic Fourier Analysis (LOCO-EFA), that overcomes the common obstacles described above. Our method also uses the whole 2D cell contour but, unlike EFA, provides a set of metrics that directly relate to morphological features, permitting the assessment of cell shape complexity in an objective and automatic manner. Importantly, it is not sensitive to cell orientation or imaging resolution, and robustly yields similar coefficients for similar shapes, allowing shape comparisons to be drawn.

To validate the usage of our method on larger cellular datasets, we analyse confocal images of Arabidopsis thaliana PCs. We then complement this study with the analysis of synthetic tissues generated using the Cellular Potts model (Glazier and Graner, 1993; Graner and Glazier, 1992), in which complex-shaped cells have a parametrised
specified shape, allowing us to ask to what degree the resultant cell shape within a confluent tissue context is shaped by cell-to-cell interactions and to what degree it can be explained by intracellular shape control mechanisms. Applying LOCO-EFA to these abstract, in silico tissues (which rather mimic animal cells, with details regarding cell wall mechanics or chemical signalling not being considered), allows us to quantify the divergence of their specified cell shape when isolated to the shape taken up when immersed within a tissue. Finally, by applying LOCO-EFA to Drosophila data, we confirm its applicability to a wide range of biological systems.

Results

Quantitative characterisation of cell shape using LOCO-EFA

Applying EFA to quantify cell shapes, we came across a number of specific shortcomings. We first explain those issues to highlight our motivation and choices that led to the development of LOCO-EFA. See supplementary Materials and Methods for further details, such as mathematical implementation. Here, we focus on explaining the analysis in terms of its biological relevance, how it can be applied and interpreted.

The shape analysis proposed here is linked to frequency decompositions of digitalised 2D shape outlines. We find it useful to compare the decomposition of a complex cell shape, such as that of a PC, to the way the sounds of musical instruments can be decomposed. When listening to a musical note, a quantifiable observable is the pitch. Within the context of PC shape this corresponds to the observed number of lobes or, as we will explain in detail, to the dominant spatial frequency of the cell’s outline. Another quantifiable property of a musical note is its volume, or amplitude. For cell shape this corresponds to the extent to which lobes protrude and indentations retract, for which we also apply the term ‘amplitude’. Finally, the timbre of musical instruments is what essentially distinguishes, for example, a clarinet from an oboe playing the same note (pitch) at the same volume/amplitude. An analogous notion for cell shape studies is the ability to capture additional aspects of shape morphology that enable differences between cells to be quantified, even when the number of lobes (pitch) and their level of protrusion (amplitude) is the same.

As a starting point, Elliptical Fourier Analysis (EFA, Kuhl and Giardina, 1982) can describe the contour of any complex two-dimensional shape, including non-holomorphic shapes such as PCs, which most other methods are unable to handle (see Figure S2, and supplementary Materials and Methods for further details). Using the coordinates of the 2D outline (Figure S3A), EFA decomposes the shape into an infinite series of ellipses (also referred to as “modes” or “harmonics”, Figure 2A). This series of ellipses, $n = 1 \cdots \infty$, can then be combined to exactly retrieve the original shape: each $n$th elliptic harmonic traces $n$ revolutions around the first ellipse while orbiting around the previous $(n - 1)$ harmonic ellipse, which in turn orbits around its previous one $(n - 2)$, and so forth (Figure S3B). This summation results in an outline being ‘drawn’, shown in Movie 1. A cut-off sets the number of modes that are actually taken into account. In general, the value is determined for which the
reconstituted cell contour is sufficiently close to the original outline (see further below).

The fact that each ellipse represents a harmonic suggests that it captures dominant spatial frequencies within the original shape. EFA harmonics have therefore been considered to be reasonable descriptors for shape (Schmittbuhl et al., 2003). However, the pitch, i.e., the most basic cellular feature to quantify, is actually not directly retrieved by EFA, not even for simple shapes. For instance, a six-sided shape is expected to present a strong contribution from the 6th mode. Instead, EFA represents such a shape as a mixed contribution from the two adjacent modes, the 5th and 7th (Figure 2E and Figure S3C). This mismatch arises from how individual EFA modes contribute to the outline. When an outline is approximated, each elliptical mode rotates either clockwise or counterclockwise. The direction of this rotation in respect to the rotation direction of the first mode causes either an increase or a decrease in the number of features drawn, one off from the actual mode (Figure S3D, E, Movie 2 and Movie 3). As a consequence, the ‘pitch’ obtained using EFA does not correspond to actual cell features, hindering interpretation. Moreover, EFA coefficients are redundant, i.e., there are more parameters than needed to specify the same specific shape (Haines and Crampton, 2000). Consequently, comparison of cell shapes on the basis of their EFA coefficients (for example, by means of principle component analysis) is nonsensical. Together, these traits make the EFA method unsuitable for cell morphology quantification and renders prohibitive meaningful comparisons between multiple cell shapes.

Diaz et al. (1990) proposed a solution for the mismatch between actual shape features and EFA’s results, using the fact that the relative direction of rotation is a main determinant of the reconstructed dominant harmonic or ‘pitch’ (see supplementary Materials and Methods for further details). It turns out, however, that each ellipse simultaneously contributes to two different spatial frequencies, something their heuristic solution cannot solve (see Figure S3F and Movie 4). As a consequence, although their method is often (but not always) able to correctly recapitulate the ‘pitch’, it is never able to correctly capture the amplitude or timbre of the cell shape.

To overcome these limitations, we propose a new basis for the outline reconstruction, which we coined $L_n$, after lobe number. Similar to EFA, modes can be summed to recreate the original shape, and each mode is represented by a set of four parameters. There are also two important distinctions. Firstly, a cell outline is now decomposed into a unique series of $L_n$ coefficients. See supplementary Materials and Methods for further details regarding elimination of coefficient redundancy. Secondly, shape features such as the protrusion number (“pitch”), their amplitude and the characteristic lobe distributions (“timbre”) are now directly mapped to the $L_n$ coefficients. They are obtained by decomposing each EFA harmonic into its exact, specific contributions to two separate $L_n$ modes (Figure 2B–D). In general, EFA modes $n-1$ and $n+1$ both partly contribute to mode $L_n$, with some specific exceptions (see Figure S4). The resulting method, which we coin Lobe Contribution-EFA or LOCO-EFA, thus consists of: eliminating multiple representations of a given outline; decomposing each $n$th elliptical EFA mode into two separate lobe
contributions; and integrating those separate modes into single LOCO-EFA modes. Every $L_n$ mode can be regarded to represent two oppositely rotating circles, each with its own starting point for the rotation, the four $L_n$ coefficients providing the radii and starting angles of both circles. We next assign a scalar $L_n$ value to capture the amplitude of each mode. Quantifying the amplitude requires both the radii of and the angular distance between the starting points of the two contributing circles, as well as the starting point of the main circle, $L_1$, to be taken into account (see Figure 2D, and supplementary Materials and Methods for further details). The $L_n$ spectrum represents the relative contribution of each individual mode to the cell shape (Figure S3C). Indeed, the spectrum of the six-lobed test shape used for Figure S3C contains a pronounced peak at mode six, as well as a peak at mode one that represents the overall circular shape. To visually appreciate the contribution of specific modes, the original shape can be reconstructed using consecutive modes up to a given mode number (compare Figure 2E with Figure 2F).
To illustrate how LOCO-EFA quantifies different shapes, we first apply it to geometrical shapes with variable numbers of protrusions (Figure 3A–I). LOCO-EFA robustly determines the main $L_n$-mode of each shape, correctly estimating lobe number (Figure 3J). We next tested if LOCO-EFA also correctly captures the amplitude, by applying the method to shapes of the same “pitch”, but with variable amplitudes (Figure 3N–Q). Indeed, the $L_n$ magnitude changes accordingly (Figure 3T), its absolute value correctly measuring the size of the extensions.

Following the analogy of sound decomposition, a more nuanced quantification is timbre. Timbre resides in the entirety of the amplitude spectrum. It is determined by which overtones are emphasised in relation to one another. For cell shape studies, we consider “timbre” analysis the ability to capture additional aspects of shape complexity, besides the main number and amplitude of protrusions/lobes. This additional information should enable distinction between different cellular phenotypes, such as between wild type and mutants (Lin et al., 2013). To illustrate, Figure 3R, S show two additional six-sided shapes which differ in “timbre” from Figure 3Q, with their accompanying $L_n$ spectra (Figure 3T). For both shapes, a clear $L_6$ peak reflects their six-lobedness, while an additional peak at $L_2$ captures the elongated nature of these shapes, and so forth. Thus, LOCO-EFA retrieves not only the main number of morphological features of an hypothetical cell, but also important fine-grained characteristics.

From the full $L_n$ spectrum, additional objective metrics can be derived to help quantify different aspects of “cell shape complexity”. We here define four metrics: XOR difference; marginal difference; cumulative difference; and entropy.

Firstly, cell shape complexity can be estimated from the approximation of the original shape by the first $N$ LOCO-EFA modes only. It addresses how relevant each subsequent $L_n$ mode is for explaining that specific shape. Figure 2F illustrated the importance of a specific mode for reconstructing the original shape (in that case, mode six). One can straightforwardly quantify the relative contribution of each mode for explaining the shape by the total areal difference (either in number of grid points or $\mu m^2$) between the original and the reconstructed shape when the first $N$ LOCO-EFA modes are used. To do so, we take the XOR (exclusive or) between the original and reconstructed cell shapes (see Figure 2F–H, and supplementary Materials and Methods for further details). A more “complex” shape requires more LOCO-EFA modes to obtain a good match. Note that a circular cell can be reconstituted using only the contribution of the first LOCO-EFA mode ($N = 1$). On the other hand, cells presenting a high lobe number require a high number of modes for XOR to approach zero (Figure 3K, U).

Quantifying cellular complexity can be further compressed by integrating from $n=2$ onwards the area under the XOR curve. We coin the resultant scalar cumulative difference (cd), with higher values corresponding to more complex-shaped cells. Figure 3M, W shows the cd values for the series of test shapes, indicating that cd becomes high when morphological protrusions increase in number or become larger in amplitude.
**XOR** profiles are typically not smooth. Instead, some modes peak as they strongly contribute in capturing the main shape features. Hence, the marginal decrease in the **XOR** value when an extra mode is added, coined *marginal difference*, further highlights the shape’s dominant modes (Figure 3L, V). This profile is comparable to the $L_n$ spectrum, also determining something akin to “pitch” and “amplitude”. We found, however, that it bears a higher discriminatory power for more complex and irregular cell shapes. Moreover, when a cell’s shape has significant contributions from multiple modes, then high marginal difference levels can be directly linked to specific cellular features (see Figure 2G, H). Thus, marginal difference helps identifying which modes are most relevant for specific shape aspects.

Finally, shape complexity is not solely about protrusion number and amplitude, but can also arise from the irregularity of these protrusions. With the previous measures, a highly regular star-shaped cell with five outspoken lobes is quantified as being as complex as a highly distorted cell with different amplitudes and distributions of five lobes, albeit less pronounced than the star-shaped case. One might therefore prefer to define cell shape complexity as a cell’s deviation from well-defined periodic outlines. A useful measure for this alternative definition of “cell shape complexity” is the Shannon entropy $E$ of the $L_n$ spectrum. The entropy measure is based upon the information content within the whole $L_n$ spectrum (Equation 1). For many shapes, entropy yields very similar results to cumulative difference. However, they give distinct results for cell outlines that have a strong contribution from the lower modes. In such cases, entropy delivers more meaningful values regarding “complexity”. This is due to lower modes being able to strongly impact cumulative difference. For example, for a highly elongated cell there will be a high contribution from $L_2$. Simply being elongated, however, does not so much represent shape complexity in the way defined above. For such a simple but elongated shape, the cumulative difference can be very similar to a shape with contributions distributed among many modes. The latter outline, however, is typically considered to be more ‘complex’. Entropy correctly captures this form of complexity. In summary, we propose LOCO-EFA and derived metrics as a new method to quantify cell shape complexity. For fully unsupervised analysis without *a priori* knowledge of cell shape features, we recommend employing all the metrics discussed in this section.

LOCO-EFA generates an infinite series of modes, without a pre-specified cut-off. Besides the measures discussed above, the **XOR** analysis also provides an algorithmic and meaningful cut-off for LOCO-EFA data analysis. Cell shapes that will be analysed with this method will in general be derived from segmentation of microscopy images. The natural choice for the grid on which to calculate the **XOR** should therefore be equivalent to the microscopy image, at its acquired resolution. **XOR** analysis (properly performed, see details in Supplementary Materials and Methods) yields values which become zero when a sufficiently large, but finite number of modes are taken into account. Additional terms then only alter the reconstructed outline at a sub-pixel resolution, i.e., at a higher resolution than the microscopy image itself. Obviously, the latter cannot be meaningful in any possible way. The mode at which **XOR** reaches zero therefore provides a natural cut-off to truncate $L_n$ series.
LOCO-EFA applied to plant pavement cells

To validate our method, we analysed Arabidopsis thaliana leaf epidermal PCs. Actual biological cells, such as PCs, can be highly asymmetrical, with multiple peaks in their $L_n$ landscape (Figure 3S, T). The outline of an asymmetrical cell with a certain number of protrusions placed quasi-periodically along its edge results in multiple superimposed protrusion frequencies. In general, the total number of hand-counted lobes matches to a peak at the corresponding $L_n$ value (but note that hand-counting is subjective). For instance, for nine lobes a peak will be observed at $L_9$. However, if these lobes are clustered in a pentagonal fashion, an additional peak at $L_5$ appears, whilst superimposed on a triangular shaped cell basis a $L_3$ contribution would be found, and so forth.

PCs acquire their characteristic jigsaw puzzle-like shape through multipolar growth patterns, such that relative simple shaped PCs become highly complex during development (Figure 4A–G). Notably, the smooth shape changes are clearly reflected in the $L_n$ spectra over time (Figure 4I). Its initial squarish shape and later nine- and thirteen-lobedness are well captured by LOCO-EFA, through peaks at modes $L_4$, $L_9$, and $L_{13}$, and corresponding peaks in the marginal difference profile. In contrast, when EFA is used, the third and fifth mode are erroneously indicated to represent shape features, besides a number of other mismatches (Figure 4H). Importantly, the smooth cell shape development over time leads to smooth changes in the LOCO-EFA $L_n$ profile over the different time points (another example is shown in Figure S5), in contrast to highly irregular changes in the EFA profile. This illustrates that comparably shaped cells can have very different EFA profiles, making EFA unsuitable to analyse real PC populations.

To visualise the shape characteristics of populations of PCs, we analysed leaves of the speechless mutant (MacAlister et al., 2007), which does not generate during the leaf development any other cell types such as meristemoids or stomata (Figure 1B, Figure 5A), as well wild type leaf epidermis, consisting of PCs, stomata and other cells from the stomatal lineage (Figure 1A, Figure 5B).

Using LOCO-EFA, it is straightforward to dissect the precise contribution of each mode for each cell in the population. Figure 5A, B show the spatial distribution of cells within a tissue that are predominantly 4-, 5-, 6-, or 7-lobed, by colour-coding cells by their $L_n$ values. Very few cells are captured by a single $L_n$ peak. Instead the majority of shapes have significant contributions stemming from multiple modes. Consequently, simply counting the number of lobes, either manually or through automatic algorithms, would lead to incomplete information regarding the shape of such cells, making it for example difficult to compare mutant phenotypes. Moreover, our data shows that PCs lack a population-wide preferential $L_n$ (Figure 5A, B).

The heterogeneity in modes that composes real populations of PCs suggests that their resultant cell shapes cannot be easily explained solely by intracellular molecular mechanisms underlying lobe and indentation patterning. Currently proposed mechanisms, based on two counteracting pathways (one for lobe formation and another for indentation formation, see details in Xu et al., 2010) give rise to Turing-
like instabilities which tend to generate symmetrical shapes (Vanag and Epstein, 2009). Moreover, these patterning models would predict that equally sized cells exhibit equal lobe numbers. However, the cell shape patterning takes place within a confluent tissue, which complicates how individual cells generate their shape. In the experimental setting, it is very hard to distinguish between the preferred shape of a cell due to its intracellular patterning, and the acquired shape due to constraints imposed by the tissue. It is well-known that if cells prefer to be round, they will take up an hexagonal shape within a tissue context (Thompson, 1917), but it is unclear what to expect for multi-lobed shapes. Therefore, to explore what shapes arise when a population of cells with complex shape preferences form a confluent tissue, and to further validate LOCO-EFA on cell populations, we simulated interacting cells with prespecified shape preferences, and employed LOCO-EFA on the resulting in silico tissue.

Applying LOCO-EFA to in silico populations and the effect of interactions between preferred cell shapes

We create in silico cells using the Cellular Potts Model (CPM), an energy-based framework that describes cells and their dynamics through small membrane extensions and retractions (see Methods section). In its basic form, CPM cell shapes emerge due to the interaction between interfacial tension, internal cellular pressure and cortical tension (Magno et al., 2015). Here we used an extension of the CPM which predefines intrinsic forces causing elongation and lobedness, resulting in more complex cell shapes. This extension consists in applying additional, cell-specific forces to subcellular update events, resulting in elongated and/or multilobed preferred cell shapes (Equation 4). Three additional forces are used that capture (i) an intrinsic tendency to elongate; (ii) a tendency to form a specified number of lobes; and (iii) an additional force for the cell to round up (Figure 5E–G, van Rooij et al. (2017b) and Movie 5). The latter term robustly prevents cells from falling apart, which becomes important within a confluent tissue with conflicting preferred cell shapes. In the simulations, a population of cells, individually having the same preferred shape, interact with each other to form a tissue. In this way, we can compare the shape of a single cell in isolation with the shape cells attain within a tissue.

We here present the analysis for two distinct specified shapes (Figure 5C, D; see Table S1 for the specific parameters used). Both preferred shapes have six lobes, but the cells shown in Figure 5D also tend to be elongated. Although the same cell shape is specified for all cells within the population (above the panels, we show the acquired cell shape in isolation), local interactions within the tissue both change and diversify the cell shapes. We quantified this divergence using LOCO-EFA. For both specified shapes, the amplitudes of the main specified modes ($L_6$ in Figure 5C and $L_2, L_4, L_6$ in Figure 5D) strongly decrease within the population, while other modes that were not prominent in isolated in silico cells became relevant within the multi-cellular context (Figure 5H, I). Marginal difference portrays a comparable picture, through a broadening of the set of modes involved. XOR analysis presents a more nuanced picture: for the elongated cells depicted in Figure 5D a structural reduction in shape
complexity is observed, i.e., the tissue context prevents cells to take up their preferred shape complexity (Figure 5I). For the rounded cells in Figure 5C, however, the relative XOR level is smaller than one for $n \geq 6$, indicating additional high-mode shape complexity triggered by the cell-cell interactions. All measures indicate large cell-to-cell variations, reflecting a high shape diversity within the tissue. We further illustrate the changes in contributions and their spatial heterogeneity by colour-coding $L_4$–$L_7$ (as indicated for each panel), for both the isolated cells and the resultant shapes of all cells within the simulated tissues. The isolated cells present a very high contribution from $L_6$, with marginal contributions from the other modes. In contrast, due to the cellular interactions, other modes become prominent within the tissue, highly varying from cell to cell, even though all cells have identical specified shapes. Thus, although a cell in isolation would generate regular protrusions with specific amplitudes, periodical lobe formation becomes inhibited and gets modified within a packed tissue, with symmetry and shape distortions being directly linked to tissue packing (Figure 5C, D). Such dynamics were observed irrespective of specified cell shape, i.e., irrespective of the number of lobes, their amplitude, and the level of overall cell elongation, and robust over a wide CPM parameter range (Figure S6, Figure S7). Given that radially symmetric, periodically-spaced lobed cell shapes are highly unlikely to be space filling, resolving conflict between preferred shape and confluency could be a relevant driving force for complex cell shape morphogenesis.

**LOCO-EFA applied to *Drosophila* amnioserosa cells**

To demonstrate LOCO-EFA’s applicability to other (non-plant) developing tissues in which cells present a high level of shape complexity, we analysed *Drosophila* during dorsal closure (Knust, 1997). Amnioserosa, the squamous epithelial that covers the dorsal side of the embryo, undergoes dramatic cell shape changes during this morphodynamic event. Simple cuboidal to columnar epithelium covers the remainder of the embryo, with both captured in our image (Figure 6A, B). At the imaged interface, cells present a broad distribution in size (Figure 6C) and shape complexity. Analysing the spatial distribution in the magnitude of the different $L_n$ modes (Figure 6D–F) reveals how the surrounding epidermis can be well described by cell elongation alone (very high and dominant $L_2$ values), while the amnioserosa cells are characterised by higher $L_n$ modes (with Figure 6E and F showing $L_5$ and $L_8$, respectively). Differences in their patterning represent cell-to-cell variations in lobe numbers. These shape characteristics are consistent with classical studies (Young et al., 1993), which proposed that observed elongation of epidermal cells perpendicular to the long axis of the embryo could explain the change in surface area required to cover the amnioserosa. The cell shapes can further be analysed by depicting the mode corresponding to the maximum marginal difference for each cell. It indicates the dominant number of extensions best describing that shape, “counting” their major morphological feature (Figure 6G). It shows how elongation dominates in the epidermis while higher modes dominate in the amnioserosa. The cumulative difference is a measure of lobe richness, its value increasing as number and amplitude of lobes increases. It yields highest values for the multi-polar cells within the amnioserosa,
presenting low levels for the epidermal cells (Figure 6H). Entropy provides an alternative quantification of shape complexity, by measuring shape irregularity. Highly asymmetric cells require a broad range of modes to capture their shape, leading to high entropy values. The spatial distribution in entropy (Figure 6I) is similar to the spatial distribution in cumulative difference, with differences between the two being particularly interesting, entropy directly highlighting the most irregular cells. In short, LOCO-EFA and its derived quantifications retrieve both the level and type of shape complexity of the cells of both tissues.

**Discussion**

The progress in microscopy and imaging techniques generates a need for adequate analytic tools to capture relevant information efficiently and objectively (Zhong et al., 2012). Image acquisition through high-throughput microscopy generates large datasets beyond the human ability (or patience) to be analysed manually, demanding computational tools. We have developed a new analytic tool which takes as the input the contour of a 2D cell projection, extracting from it, in an efficient and parameter-independent manner, quantitative meaningful shape information. Importantly, the pipeline can be integrated within segmentation procedures (Fernandez et al., 2010; van Rooij et al., 2017a), to fully automate shape analysis of a series of images.

Our method can be intuitively grasped through the analogy of music perception. To quantify an instrument playing a certain note, say a violin playing the note A, one firstly wishes to have a device that determines the note played. We have shown here that LOCO-EFA, unlike EFA, correctly determines the analogous feature for shapes, which is the number of protrusions. Moreover, LOCO-EFA, in contrast to EFA, quantitatively measures the amplitude of that particular feature; alike determining the volume of a given note, when multiple notes are played concomitantly. In all examples presented here we have normalised to cell area. Hence an L-value of 0.15 indicates a peak-to-trough distance of 15% of the cell diameter (amplitude equal to 15% of the cell radius).

LobeFinder, the recent method developed by Wu et al.(2016) can also be employed to assess protrusion number. When the biological question asked requires not only the “pitch” to be measured, but also the “volume” and “timbre”, corresponding to lobe amplitude and other irregularities, such alternative methods will be insufficient. Indeed, LOCO-EFA provides a holistic set of measurements that allows complex morphologies to be quantified in a reproducible manner.

We illustrated how the measurements obtained via LOCO-EFA can be interpreted, first using simple shapes (geometrical or symmetrical forms), followed by using confocal images of Arabidopsis PCs and Drosophila amnioserosa, to better assess the performance of our method on actual, highly complex and asymmetric biological shapes. When analysing complex shapes through the $L_n$ spectrum only, it is non-trivial to ‘visualise’ the corresponding shape in the same manner as can be done for geometric shapes. In such cases it is useful to plot the XOR and marginal difference profiles, to gain a better notion of the major shape properties. PC shape analysis is directly biologically relevant, because many of the players accounting for the lobe and
indentation patterning are known (Jones et al., 2002; Xu et al., 2010), enabling one to now extend the study of cell shape control to mutants and experimental interferences. We found that few cells have a symmetrical shape, i.e., most cannot be represented well by a single high $L_n$ value. It is unlikely that such composition of real cell shapes in several $L_n$ values can be fully explained by the existence of two counteracting pathways specifying lobe and indentation identity. Our in silico approach rather suggests that the interactions between space-filling shapes can dramatically increase the overall irregularity: even when the CPM cells are specifically programmed to develop well-defined regular shapes, the interactions between them trigger dramatic cell shape deviations and variations. Within the tissue the main, specified mode decreases in strength while other modes become relevant. Thus, tissue confluency leads to asymmetric and variable resultant shapes.

Although our synthetic data is but a phenomenological description of real shapes, our results suggest that the local influence of neighbours during PC development could be important for shape acquisition. To assess this hypothesis further, it will be crucial to perform quantitative shape analysis on in vivo cell populations over time, combined with growth tensor analysis (i.e., anisotropy and spatial patterning in the growth rate). Such studies, combined with genetic or physical perturbations in cell growth and deformation and in silico cell growth models, could help untangle how cell shape specified at the cellular level is linked to the resultant shape arising at the tissue level.

Applying LOCO-EFA to cell tracking data, we observed that the LOCO-EFA profiles of those changing cells varied smoothly over time. Such trajectories are cell-specific and provide unique fingerprints of each individual developing cell. This opens the possibility of using the $L_n$ modes as cell identifiers within a temporal sequence of images, to help track populations of cells automatically.

To illustrate how this powerful tool can be used to measure complex undulating cells, we have here applied LOCO-EFA to Arabidopsis PCs and Drosophila amnioserosa. Although we focussed on discussing overall shape distributions throughout the tissue, LOCO-EFA shape descriptors could also be used to investigate correlations in shape between neighbouring cells, in a similar manner as done for topological traits in the same tissue (Carter et al., 2017). Moreover, LOCO-EFA analysis on shape dynamics and shape correlations between neighbouring cells can be easily extended to other cell types and other species, including less complex shapes. Furthermore, LOCO-EFA could also be relevant for understanding phenotypic morphology of subcellular structures, such as mitochondria, which can present different levels of shape complexity (Dimmer et al., 2002), and sperm cell nuclei, already analysed using EFA (Mashiko et al., 2017). Our method is also well-suited for studying organ shape development, specifically when landmarks are difficult to assign. It can thereby enhance quantification and biological meaningfulness of current EFA-based studies that, for example, decompose entire leaf shapes (Liao et al., 2017), insect wings (Yang et al., 2015), jaw shape and sizes (Rose et al., 2015), and pinniped whisker morphologies (Ginter et al., 2012). LOCO-EFA can even be employed at different levels within the same organism, for example, to quantify leaf shape,
serrations as well as root morphology (Li et al., 2017). Lastly, LOCO-EFA could constitute a powerful tool for whole organism analysis, especially within paleobiology, where it could enrich current elegant studies initiated using EFA, to, for example, analyse bivalves (Crampton, 1995), trilobite-like arthropod evolution (Jackson and Budd, 2017), and Triatominae eggs (Santillán-Guayasamín et al., 2017). For all such studies, when possible, we recommend that our method be integrated within recent image analysis pipelines, allowing extraction and analysis of shape information in a high-throughput manner (Heller et al., 2016; Stegmaier et al., 2016; van Rooij et al., 2017a).

In short, LOCO-EFA can be used to quantify morphologies described as closed 2D contours, across scales, from subcellular to organ and beyond.

**Material and Methods**

**Confocal images and image processing**

Columbia wild type or *speechless* mutant (MacAlister et al., 2007) leaves expressing pmCherry-Aquaporin (Nelson et al., 2007) were imaged using a confocal microscope Leica SP5 at comparable stages and in comparable regions. Cells changing over time were imaged using a custom-made perfusion chamber (Kuchen et al., 2012; Robinson et al., 2011; Sauret-Güeto et al., 2012). Further image processing to flatten the images was done using ImageJ. *Drosophila melanogaster* embryos expressing ubi-DE-Cadherin-GFP (Oda and Tsukita, 2001) were dechorionated in bleach, rinsed in water and attached to a coverslip with the dorsal side up using heptane glue and covered with Halocarbon Oil 27 for live imaging on a Zeiss 780 confocal. Both the *Arabidopsis* and *Drosophila* images were segmented using in-house software (Segmentation Potts Model, van Rooij et al., 2017a). In this study we present a single, typical example of a wild type and of a *spch* leaf, as well as five typical examples of static pavement cell outlines and two typical examples of developing pavement cells, all within a *spch* leaf. These images were selected from a study in which one WT and seven *spch* leaves were imaged at in total 15 time-points for the WT leaf and 121 time-points for the *spch* leaf (Carter et al., 2017). The amnioserosa image represents a typical example selected from four live-imaged embryos.

**Shape descriptors**

Average lobe lengths and neck widths were calculated using ImageJ (Analyse → Measure). The skeleton was calculated using Better Skeletonization by Nicholas Howe, available through MATLAB File Exchange.

**Geometric shapes**

All geometric shapes were generated by the “superformula” described in Gielis (2003), and were analysed in the same manner as the confocal images.
**XOR**

All the grid points belonging to each individual real or synthetic pavement cell were compared to all the grid points captured by the subsequent series of LOCO-EFA reconstructions. A reconstruction of level $N$ takes into account the first $N$ $L_n$ modes. The *in silico* cells were generated using the Cellular Potts Model, which is a grid-based formalism, while for the experimental data the grid points were directly defined by the imaging resolution. The scripts used to calculate the XOR and to colour-code the real and synthetic cells were written in the coding language C. See supplementary Materials and Methods for further details.

**Entropy and other measurements**

The entropy measure is defined as

$$E = -\sum_{l=1}^{L_l} f_l \ln f_l ,$$

where $f_l$ refers to the relative proportion of each $L_l$ for a given $L$ number of modes analysed, i.e., $f_l = \frac{L_l}{\sum_{l=1}^{L} L_l}$.

Shape approximations, cumulative difference and entropy were calculated using the first 50 $L_n$ modes. To capture cell shape complexity linked to protrusions rather than mere anisotropy, cumulative difference is calculated from the second $L_n$ mode onwards. This value turned out to be more than sufficient to capture any cell shape given the grid point resolution used for all cases here. Note, however, that very high-resolution images might require additional modes to fully capture the shape.

**Cellular Potts Model generating complex cell shapes**

The Cellular Potts Model (CPM) is an energy-based model formalism used to model cellular dynamics in terms of Cell Surface Mechanics (Magno et al., 2015). Individual cells are described by a set of grid points on a lattice. In this manuscript, we used the CPM to generate *in silico* cells with relatively complex shape preferences that are allowed to interact within a confluent setting. During each simulation step a grid point is chosen in a random fashion to evaluate whether its state changes into one of its neighbouring states, effectively corresponding to a small cell shape modification at that point. To evaluate if such state change will occur, the energy change is calculated that such a copy would cause. This is done by calculating the change in the configurational energy as defined by the following Hamiltonian, which sums up the energy contribution of each pixel within the entire field as well as of all cells:

$$H = \sum_{ij} \sum_{i'j'} J \left(1 - \delta_{c_{ij},c_{ij'}}\right) + \sum_c \lambda_a \left(a_c - A\right)^2 + \sum_c \lambda_p \left(p_c - P\right)^2 .$$

$J$ refers to the coupling energy, summed over all grid points $(i,j)$ and their eight (2nd order) neighbours $(i',j')$. The Kronecker delta term $\left(1 - \delta_{c_{ij},c_{ij'}}\right)$ simply assures that neighbouring lattice sites of the same state (i.e., belonging to the same cell) do not
contribute to the total energy of the system. The variables $a_c$ and $p_c$ denote respectively the actual cell area and the actual cell perimeter for each cell ($c$); the parameters $A$ and $P$ denote the target cell area and perimeter. The parameters $\lambda_a$ and $\lambda_p$ describe the resistance to deviate from the target area and perimeter, respectively. The probability a copying event is accepted depends on the change in the Hamiltonian,

$$\Delta \mathcal{H} = \mathcal{H}_{after} - \mathcal{H}_{before},$$

in the following way:

$$P = \begin{cases} 1 & \text{if } \Delta \mathcal{H} < -Y, \\ e^{(\frac{\Delta \mathcal{H} + Y}{T})} & \text{if } \Delta \mathcal{H} \geq -Y, \end{cases}$$

where $Y$ corresponds to the yield or ability of a membrane to resist a force and $T$ (simulation temperature) captures additional stochastic fluctuations. Copying events which decrease $\mathcal{H}$ by at least $Y$ will always be accepted, otherwise acceptance follows a Boltzmann probability distribution (Equation 3).

To generate cells with a particular number of preferred protrusions, we modify the change in the Hamiltonian as is calculated for every evaluated copying event, effectively shortcutting intracellular biochemistry and biophysics, in the following way. Simulated cells are attributed with a specified preferred number of lobes, amplitude of lobes, overall elongation and roundness, implemented by modifying the change in the Hamiltonian for every evaluated copy event as follows (van Rooij et al., 2017b):

$$\Delta \mathcal{H}' = \Delta \mathcal{H} - \nu \cos(n\theta) - \chi \cos(2\alpha) - \mu\left(\frac{\sqrt{A}}{\pi} - r\right).$$

Those three additional terms are evaluated for both cells involved in the copying event, so there are effectively six additional terms. The first term captures the tendency to form $n$ lobes, with $\nu$ capturing the propensity to extend to form a lobe or to retract to form an indentation, thus giving rise to the amplitude or pointedness of the lobes. $\theta$ describes the angle between any of the $n$ equally spread out target directions for outgrowth and the vector determined by the coordinates of the grid point under evaluation and the centre-of-mass of the cell (hereafter called the copy vector) (Figure 5E). To clarify, when a cell extension is considered right on top of one of the target directions, then $n\theta = 0$, $\cos(n\theta) = 1$, and tendency to extend is maximally increased, while halfway two target directions, $n\theta = \pi$, $\cos(n\theta) = -1$, and the tendency to extend is maximally suppressed.

The second term in Equation 4 captures an overall elongation, implemented in a similar fashion. The parameter $\chi$ corresponds to the propensity to elongate and $\alpha$ is the angle between the elongation vector and the copy vector.

If only these two terms are used, cells within tissue simulations can easily lose coherence, i.e., fall apart. Therefore, a third term was added, capturing a propensity to roundness. The parameter $\mu$ captures the resistance of a cell to deviate from a circle, with $r$ being the length of the copy vector, and $\sqrt{A}/\pi$ being the preferred radius of cell, given its target area. For further details, see van Rooij et al. (2017b).

Importantly, the target lobe and elongation vectors are not fixed during the simulation. At intervals of 100 simulation time steps they are dynamically updated, in order to attain the most favourable position, effectively “accommodating” its lobe positions with respect to its neighbours. During a vector update step, the preferred
directions of extension are matched to the set of directions the current shape of cells presents the strongest level of extension.

The initial cell positions within the field were randomly chosen. Simulations were run for 10000 time steps (see an example in Movie 5 Parameters used for each used specified cell shape are given in Table S1.

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Figures

Figure 1: Complex cell shapes and the shortcomings of traditional shape quantifiers. (A–C) Complex cell shapes in both plant (A, B) and animal (C) tissues. (A, B) Pavement cells (PCs) of Arabidopsis thaliana wild type (A) and speechless mutant (B), characterised by jigsaw-like shapes. Scale bars 50 μm. (C) Amnioserosa cells in the Drosophila embryo present cell shapes with similar complexity. Scale bar 20 μm. (D–G) Individual cells from the imaged tissues (upper panel), and the corresponding segmented cell outlines (lower panel). Scale bars 10 μm. (H) Traditional metrics to quantify cell shape lead to similar values for very different shapes and are image-resolution and parameter sensitive. Here we compare the cells D–G. See also Figure S1.
Figure 2: To get LOCO. (A) EFA decomposes a 2D cell outline into an infinite summation of related ellipses or modes that can also be used to approximate the cell outline. (B) Each EFA elliptic harmonic is decomposed into two counter-rotating circles. (C) Mode $L_n$ is composed of the counter-clockwise rotating $n+1$th EFA harmonic circle and the clockwise rotating $n-1$th circle. (D) The combined amplitude contribution to $L_n$ (yellow line) of the two counter-rotating circles with radii $\lambda_{n+1}$ and $\lambda_{n-1}$ is also depends on the offset in their starting points and the offset of the overall (mode 1) starting point, which together determine the initial phase shift (green dots) in the amplitude contribution of each rotor. (E, F) Comparison of closed contour reconstruction through either EFA (E) or LOCO-EFA (F). Although both approximations converge to the original 6-lobed star shape (labelled ‘Original’), the reconstruction using EFA harmonics (E) generates a spurious shape after addition of the 5th harmonic and only recovers the original shape after the 7th harmonic, whereas the LOCO-EFA (F) reconstitutes the original shape precisely at the 6th mode, matching the protrusion number. The number of modes used for each sequential reconstruction is indicated below each shape. (G) LOCO-EFA reconstruction of a real cell taking the first $n L_n$ modes into account, as indicated below the panels. (H) Determination of the level of mismatch between the original cell shape and its $n$th order truncated LOCO-EFA approximation, by applying the XOR (Exclusive OR) function (see Supplementary Materials and Methods).
Figure 3: Interpreting LOCO-EFA-derived measures for geometrical and asymmetric shapes. (A–I) Symmetrical and well-defined geometrical shapes with normalised area. (J–L) $L_n$, $XOR$, and marginal difference profiles for shapes (A–I). (J, L) For each geometric shape, a clear peak appears in the profiles, this main contributor to the shape always coinciding with the number of protrusions. (M) Cumulative difference (cd) and entropy for shapes (A–I). (N–Q) Symmetrical shapes with increasing protrusion amplitude. (R, S) Asymmetrical shapes. (T–V) $L_n$, $XOR$, and marginal difference profiles for shapes (N–S). Increasing protrusion amplitude leads to increasing peak levels in the profiles. Asymmetric shapes present multiple peaks, indicating that multiple modes are needed to recapitulate the original shape. (W) Cumulative difference (cd) and entropy for shapes (N–S).
Figure 4: LOCO-EFA metrics on a cell changing its shape over time. (A–G) Sequence of a tracked PC growing over time with normalised area. (H) $P_n$ and marginal difference profiles using EFA. Applying EFA modes to approximate the cell shapes leads to erratic profiles that fail to recover the biological sequence of development, observed in the $P_n$ profile and as spurious peaks at the third and fifth harmonics in the marginal difference profile. (I) $L_n$ and marginal difference profiles using LOCO-EFA. The LOCO-EFA measurements recover the smooth transitions during the cell morphogenesis. The overall square symmetry of the cell is captured by a peak at $L_4$, the formation of lobes by a smooth increase in $L_9$, and later $L_{13}$.
Figure 5: LOCO-EFA analysis on *in vivo* and *in silico* pavement cells. (A, B) LOCO-EFA applied to *speechless* mutant (A) and wild type (B) leaf tissue. Colour coding depicts the $L_n$ values for 4 different $L_n$ modes, as indicated above each panel, with the scale shown below. Very few cell shapes can be reasonably captured through a single $L_n$ mode, revealing cell shape complexity. (C, D) LOCO-EFA applied to *in silico* PCs reveals the degree of divergence from their specified shape that interacting cells within a tissue experience. Two different specified cell shape populations are shown (SCS1 and SCS3, each with six lobes, see Table S1). The specified shapes are depicted above each panel. Colour coding within the panels and of the specified shapes above each panel again depicts the $L_n$ values, with the scale shown below. Within the tissue, strong deviations in $L_n$ contributions are observed. (E–G) Modelling...
framework to generate the *in silico* tissues. (E) Standard Cellular Potts Model is modified to allow for a specified number of lobes (here, \( n = 5 \)) to form at regular radial spacings (\( \alpha \)). (F) This gives rise to a symmetric, multi-lobed specified cell shape, shown in red. (G) Within the tissue, however, cells with the same specified shape deform while interacting with neighbouring cells. (H, I) Distribution of the ratios, for three different LOCO-EFA metrics, between the cell in isolation and each of the cells within the tissue population, for SCS1 (H) and SCS3 (I), respectively. The central mark of the box plot indicates the median and the edges refer to the 25th and 75th percentiles. \( n = 66 \) (H) and 44 (I) *in silico* cells, respectively. The red lines highlight where the ratio is unity. \( L_n \) and XOR are plotted on a log scale.
Figure 6: Cell shape analysis during dorsal closure of the Drosophila embryo. (A) Confocal image of amnioserosa cells. (B) Segmentation identifies each cellular domain by a unique ID, represented through a distinct colour. (C–I) Several cell shape characteristics, quantified and depicted by the heat map shown below C. (C) Cell area. Amnioserosa cells are larger (red) than surrounding epithelia (blue and purple cells). (D) $L_2$ for each cell. Levels are high in surrounding epithelia, corresponding to predominant cell elongation. (E) $L_5$ for each cell. Amnioserosa cell shapes carry larger representations of higher mode numbers. The cell with the highest $L_5$ contribution indeed displays five distinct protrusions. (F) $L_8$ for each cell. Higher modes substantially contribute to the amnioserosa cells, with $L_8$ strikingly high for the cell with eight visibly prominent protrusions, and high for other multi-polar shapes. (G) Mode at which highest marginal difference occurs, depicted for each cell. Colours represent mode numbers, as indicated. For example, many cells can be described predominantly having an elongated axis (purple cells, with highest mode 2), while one cell is best described as being triangular (blue cell, with highest mode 3), etc. (H) Cumulative Difference for each cell, a measure of lobe richness (both number and amplitude). (I) Entropy for each cell, a measure of shape irregularity. Colour map between 0 and max for D–F and between min and max for C, H, I.
## Table

Table 1: Distinct shape descriptors have been used to quantify pavement cells.

| Measure                        | Description                                                                                                                                                                                                 | Reference                  |
|-------------------------------|-----------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|-----------------------------|
| Average lobe length and neck width | Length of each lobe and the distance between opposite indentations within a cell (called necks) are shown in Figure S1. The final measure for a cell is the average of all lobe lengths and the average of all the neck widths. These measurements depend on human criteria to identify lobes and necks, and are given in absolute length units (thus, incomparable throughout growth stages). | (Fu et al., 2005)           |
| Form factor (or circularity)  | Defined as $\frac{p^2}{4\pi A}$, where $p$ is the perimeter and $A$ is cell area. A circle corresponds to a form factor 1, the lowest value possible.                                                                 | (Andriankaja et al., 2012; Bai et al., 2010; Russ, 2011) |
| Skeleton                      | This metric relies on the number of end points of a skeleton representation of the cell shape. The skeleton is formed by iteratively removing pixels from a grid-based cell shape representation, such that eventually a branched one-dimensional graph remains. There are different variants of this algorithm to skeletonise shapes; the resulting branch patterns and length of branches highly depends on the parameters used and are very sensitive to the image resolution. | (Le et al., 2006; Russ, 2011) |
| Average polarity score        | Defined as $\frac{c + s}{2}$, where $c$ is the circularity and $s$ the number of skeleton end points.                                                                                                  | (Sorek et al., 2011)        |
| LobeFinder                    | This method calculates the convex hull or minimal polygon enclosing a cell. After applying certain thresholds, the number of lobes corresponds to the number of local minima.                               | (Wu et al., 2016)           |
Supplementary Materials and Methods

Decomposing shape: Lobe Contribution Elliptical Fourier Analysis (LOCO-EFA)

In this section, we first summarise previous efforts to make EFA coefficients interpretable within a morphometrics perspective and explain why matching EFA coefficients with shape features generally does not hold. We describe in detail our new method, Lobe Contribution Elliptical Fourier Analysis (LOCO-EFA). We show how it provides quantitative and biologically interpretable measurements that are unique for a given shape, overcoming the shortfalls of the previous methods.

Standard Fourier Analysis cannot be used to quantify complex cell shapes

Standard Fourier Analysis has been widely used to analyse cell morphology. It can, however, only be applied when cells present simple holomorphic shapes, i.e., when the radii emanating from the centroid of a cell intersect its outline only once (Figure S2A and Pincus and Theriot (2007)). When the geometry of a cell is more complex, as in the case of pavement cells, and radii emanating from the centroid can intersect the outline more than once, the shape cannot be decomposed using a Fourier expansion based on polar coordinates (Figure S2B and Schmittbuhl et al. (2003)).

Elliptical Fourier Analysis fails to align mode frequency with morphological features

In 1982 Kuhl and Giardina proposed the Elliptical Fourier Analysis (EFA) to describe the contour of any two-dimensional shape (both holomorphic and non-holomorphic), derived from the coordinates of all the points along its outline. In short, EFA takes the $x$ and $y$ coordinates of a closed contour and decomposes it into an infinite summation of related ellipses:

$$x(t) = \alpha_0 + \sum_{n=1}^{\infty} \left( \alpha_n \cos \left( \frac{2n\pi t}{T} \right) + \beta_n \sin \left( \frac{2n\pi t}{T} \right) \right), \quad (5a)$$

$$y(t) = \gamma_0 + \sum_{n=1}^{\infty} \left( \gamma_n \cos \left( \frac{2n\pi t}{T} \right) + \delta_n \sin \left( \frac{2n\pi t}{T} \right) \right), \quad (5b)$$

where $\alpha_n$, $\beta_n$, $\gamma_n$ and $\delta_n$ are the so-called EFA coefficients and $\alpha_0$ and $\gamma_0$ are the $x$- and $y$-offset of the initial contour. The detailed derivation of the formulae for $\alpha_0$, $\gamma_0$, $\alpha_n$, $\beta_n$, $\gamma_n$ and $\delta_n$ can be found in Kuhl and Giardina (1982). They are calculated from a discrete chain of contour points $(x_i, y_i)$ with $i = 1, \ldots, K$ (see Figure S3A), $K$ being the total number of points along the closed contour. We define $(x_0, y_0) \equiv (x_K, y_K)$, given that cell contours are closed. Now imagine drawing the contour of the cell, then $\Delta t_i$ is the time spent drawing the line segment of the contour that links $(x_{i-1}, y_{i-1})$ to $(x_i, y_i)$, i.e., $\Delta t_i = \sqrt{(x_i - x_{i-1})^2 + (y_i - y_{i-1})^2} = \sqrt{\Delta x_i^2 + \Delta y_i^2}$. Note that $\Delta t_i$ is not fixed but can vary for each interval. Define $T$ as the total time spent to draw the whole contour, i.e., $T=\sum_{i=1}^{K} \Delta t_i$. The “time” passed while drawing the contour, starting from contour point $(x_0, y_0)$, or, equivalently, the distance passed along the contour to reach each contour...
point \((x_i, y_i)\), is referred to as \(t_i\), i.e., \(t_i = \sum_{p=1}^{i} \Delta t_p\), with \(t_0 = 0\), and \(t_K = T\), the “total drawing time” or total perimeter length (see Figure 1 in the main text). Given that no equal spacing between the points is required, it is straightforward to define \(K\) observation points from any kind of cell contour. The only requirements are that the contour is closed and the coordinates form an ordered list that follows the contour. The EFA coefficients are then given by:

\[
\begin{align*}
\alpha_n &= \frac{T}{2n^2 \pi^2} \sum_{i=1}^{K} \frac{\Delta x_i}{\Delta t_i} \left( \cos \frac{2n\pi t_i}{T} - \cos \frac{2n\pi t_{i-1}}{T} \right) , \\
\beta_n &= \frac{T}{2n^2 \pi^2} \sum_{i=1}^{K} \frac{\Delta x_i}{\Delta t_i} \left( \sin \frac{2n\pi t_i}{T} - \sin \frac{2n\pi t_{i-1}}{T} \right) , \\
\gamma_n &= \frac{T}{2n^2 \pi^2} \sum_{i=1}^{K} \frac{\Delta y_i}{\Delta t_i} \left( \cos \frac{2n\pi t_i}{T} - \cos \frac{2n\pi t_{i-1}}{T} \right) , \\
\delta_n &= \frac{T}{2n^2 \pi^2} \sum_{i=1}^{K} \frac{\Delta y_i}{\Delta t_i} \left( \sin \frac{2n\pi t_i}{T} - \sin \frac{2n\pi t_{i-1}}{T} \right) .
\end{align*}
\]

The offset to the contour is given by:

\[
\begin{align*}
\alpha_0 &= \frac{1}{T} \sum_{i=1}^{K} \left( \frac{\Delta x_i}{2\Delta t_i} \left( t_i^2 - t_{i-1}^2 \right) + \xi_i (t_i - t_{i-1}) \right) + x_0 , \\
\gamma_0 &= \frac{1}{T} \sum_{i=1}^{K} \left( \frac{\Delta y_i}{2\Delta t_i} \left( t_i^2 - t_{i-1}^2 \right) + \varepsilon_i (t_i - t_{i-1}) \right) + y_0 .
\end{align*}
\]

where \(\xi_i = \sum_{j=1}^{i-1} \Delta x_j - \frac{\Delta x_i}{\Delta t_i} \sum_{j=1}^{i-1} \Delta t_j\), \(\varepsilon_i = \sum_{j=1}^{i-1} \Delta y_j - \frac{\Delta y_i}{\Delta t_i} \sum_{j=1}^{i-1} \Delta t_j\), and \(\xi_1 = \varepsilon_1 = 0\). Further details and full derivation can be found in Kuhl and Giardina (1982).

Each set of four coefficients yields an ellipse (also referred to as the “nth mode” or “nth elliptic harmonic”), with a certain orientation and a certain starting point. The original cell outline can thus be expressed as an infinite summation of ellipses. Note that \(x(t)\) and \(y(t)\) are periodic functions with period equal to \(T\).

A visual way to understand how the set of ellipses gives rise to the final shape is as follows: the second elliptic harmonic traces two clockwise or counter-clockwise revolutions around the first harmonic; the third harmonic traces three revolutions around the path drawn by the second harmonic; and the \(n\)th harmonic traces \(n\) revolutions around the path drawn by the previous harmonic (see Figure S3 and Movie 1).

Diaz et al. (1990) proposed a heuristic measure regarding the contribution of each harmonic to the shape through an approximation of the perimeter of each ellipse multiplied by its harmonic number \(n\):

\[
P_n = 2\pi n \sqrt{\frac{\lambda_{1n}^2 + \lambda_{2n}^2}{2}} ,
\]

where \(\lambda_{1n}\) and \(\lambda_{2n}\) are the major and minor axis of the \(n\)th ellipse. Moreover, Diaz et al. (1990) introduced an additional correction to capture the complex relationship between EFA modes and shape feature periodicity.
The direction of rotation of the \( n \)th harmonic ellipse is given by the determinant of the EFA coefficients matrix, \( \det \begin{bmatrix} \alpha_n & \beta_n \\ \gamma_n & \delta_n \end{bmatrix} \), i.e., the direction of rotation is given by the sign of

\[
r_n = \alpha_n \delta_n - \beta_n \gamma_n. \tag{9}
\]

If \( r_n < 0 \), the elliptic harmonic is rotating clockwise; if \( r_n > 0 \) the elliptic harmonic is rotating counter-clockwise. When EFA is used for shape approximation, mode \( n \) contributes to shape features with an \( n + 1 \) or \( n - 1 \) periodicity. This is in contrast to standard Fourier Analysis, in which mode \( n \) contributes to shape features with an \( n \) periodicity. Standard Fourier Analysis, however, is only possible for holomorphic shapes, and hence cannot be applied to, for example, pavement cells. Diaz et al. (1990) observed that whether mode \( n \) predominantly contributes to shape features with an \( n + 1 \) or with an \( n - 1 \) periodicity strongly depends on whether the \( n \)th harmonic rotates together with or against the direction of the first harmonic (see Movie2 and Movie3). This effect of presenting contributions to the \( n + 1 \)’th and \( n - 1 \)’th mode depending on the rotation direction of the first and \( n \)th harmonic is a common phenomenon observed for objects orbiting around others (hereafter referred to as the relative direction effect). A well-known example of the relative direction effect is the rotation of the Earth and its movement around the sun. The actual number of rotations our planet makes per year (as observed from “star-rise to star-rise”, the so-called sidereal days) is one off from the number of days we perceive in a year (from “sunrise to sunrise”, the so-called solar days). Because our planet rotates around its axis in the same direction as it moves around the Sun, the number of solar days per year is 365, one less than the number of sidereal days per year, which is 366. If the rotation of Earth would have been in the opposite direction as its movement around the sun, the number of solar days per year would instead have been 367. In light of exactly the same principle, Diaz et al. (1990) introduced that when the \( n \)th elliptic harmonic is moving in the same direction as the first harmonic, its shape contribution \( P_n \) should be assigned to \( n - 1 \); inversely, when the direction of a given mode is opposite to the first harmonic, its shape contribution \( P_n \) should be assigned to \( n + 1 \).

We will show below that this simple heuristic is reasonable as long as the ellipse marginally deviates from a circle, but is is not valid in general. When the aspect ratio of the ellipse (\( \lambda_{1n}/\lambda_{2n} \)) is large (i.e., the elliptical harmonic is very flat, deviating significantly from a circular shape), the proposed rule fails to apply. Figure S3F illustrates a situation when the rotation direction of the first and third harmonic are opposite (and no other modes are used), yet instead of generating a contour with \( n - 1 = 2 \) protrusions, as expected from the heuristic rule, a four-sided outline is generated, clearly illustrating that this method of \( P_n \) shifting does not work in general (see also Movie 4). Moreover, it is not possible to reconstruct the original shape using the \( P_n \) values, and therefore cannot be used for additional analysis based on shape reconstruction as presented in the main paper. This strongly limits usage of EFA for biological shape interpretations and statistical population analysis. Surprisingly, although EFA has been used to quantify morphology at the organ level, relative direction effect has typically been ignored altogether (Chitwood et al., 2013; Frieß and Baylac, 2003; Iwata et al., 2010, 1998; Neto et al., 2006; Yoshioka et al., 2005).

Realising that the source of the problem is linked to the eccentricity of the ellipses, it became clear to us that we could overcome this issue by essentially decomposing each ellipse into two counter-rotating circles. All circles can then be redistributed, forming a
new base. The details of how to do so are discussed below. We call the new base $L_n$, which, when summed up, can also reconstitute the original shape.

**Contouring the limitations: Lobe Contribution Elliptical Fourier Analysis (LOCO-EFA)**

To capture the biologically relevant cell shape features, overcoming the limitations of using $P_n$ and rotation-dependent $n + 1, n - 1$ adjustments, we have developed an alternative method coined Lobe Contribution Elliptical Fourier Analysis (LOCO-EFA). As the name indicates, it correctly maps the contribution of each mode/harmonic to the corresponding morphological features. This is done by separating each elliptic harmonic into two circular harmonics, each rotating in an opposite direction.

First we rewrite the EFA (Equation 5) in matrix form:

$$
\begin{bmatrix}
x(t) \\
y(t)
\end{bmatrix} = \begin{bmatrix}
\alpha_0 \\
\gamma_0
\end{bmatrix} + \sum_{n=1}^{N} \begin{bmatrix}
\alpha_n & \beta_n \\
\gamma_n & \delta_n
\end{bmatrix} \begin{bmatrix}
\cos\left(\frac{2n\pi t}{T}\right) \\
\sin\left(\frac{2n\pi t}{T}\right)
\end{bmatrix},
$$

with the infinite sum being truncated at the $N$th order harmonic. Equation 10 can concisely be expressed as

$$
[X(t)] = [A_0] + \sum_{n=1}^{N} [A_n] [M_n(t)],
$$

in which $[X(t)]$ corresponds to the drawn cell outline $\begin{bmatrix}
x(t) \\
y(t)
\end{bmatrix}$; $[A_0]$ represents the spatial offset $\begin{bmatrix}
\alpha_0 \\
\gamma_0
\end{bmatrix}$; $[A_n]$ corresponds to the EFA coefficients matrix $\begin{bmatrix}
\alpha_n & \beta_n \\
\gamma_n & \delta_n
\end{bmatrix}$; and $[M_n(t)]$ refers to the rotor $\begin{bmatrix}
\cos\left(\frac{2n\pi t}{T}\right) \\
\sin\left(\frac{2n\pi t}{T}\right)
\end{bmatrix}$. (For clarity, we will use the notation $[..]$ throughout to emphasise we are dealing with matrices; not to be confused with $|..|$ that represents determinant, which we here only refer to as $\text{det} [..]$.)

The LOCO-EFA method consists of three steps: 1) eliminate multiple representations of the same outline; 2) decompose each $n$th elliptic harmonic into two circular harmonics, each rotating in an opposite direction; and 3) determine $L_n$ and $L_n$ for all $N$ modes. Below we describe these steps in detail.

(1) **Eliminate multiple representations of the same outline**

It had already been noted that EFA coefficients are redundant and therefore compromise statistical analysis and shape comparisons (Haines and Crampton, 2000). We found that there are three sources of degeneracy in the EFA coefficients that therefore have to be eliminated. First, a contour can be drawn starting from any arbitrary initial point along the contour. While exactly the same outline is drawn, each starting point is represented by a completely different set of EFA coefficients for all modes $[A_n]$. Basically, whenever the starting point is changed, all elliptic harmonics take a different orientation (Kuhl and Giardina, 1982). The first step is therefore to transform the EFA coefficients such that the starting point of the first harmonic is always positioned at, for standardisation, the extreme of the semi-major axis (see further below). The second source of degeneracy, however, is that such a normalisation still allows for two possible representations of the
outline, since each of the two extremes along the semi-major axis can be chosen as the starting point. Moreover, a third source of degeneracy is due to the fact that the outline can be drawn clockwise or counter-clockwise. Clearly, all three sources of degeneracy have to be removed to make any comparison between cells sensible.

The first step is to determine where the new starting point should be positioned, as well as the scaled amount of time or temporal angle ($\tau_1 = \frac{2\pi t}{T}$) required to reach the starting point (see Kuhl and Giardina, 1982). As stated above, we wish the starting point to coincide with one of the extremes of the semi-major axis of the first harmonic. Points along the first harmonic ($x_1, y_1$) can be described as:

\begin{align}
x_1(\tau) &= \alpha_1 \cos \tau + \beta_1 \sin \tau, \\
y_1(\tau) &= \gamma_1 \cos \tau + \delta_1 \sin \tau,
\end{align}

where $\tau = \frac{2\pi t}{T}$ being the scaled time or temporal angle. By differentiating the magnitude of the first harmonic ellipse $E(\tau) = \sqrt{x_1(\tau)^2 + y_1(\tau)^2}$ and setting its derivative to zero ($\frac{dE(\tau)}{d\tau} = 0$), the temporal angles can be found at which the extremes along the semi-major and semi-minor axes of the first harmonic are reached (Kuhl and Giardina, 1982):

$$\tau_1(\nu) = \frac{1}{2} \arctan \left( \frac{2(\alpha_1 \beta_1 + \gamma_1 \delta_1)}{\alpha_1^2 + \gamma_1^2 - \beta_1^2 - \delta_1^2} \right) + \frac{\nu}{2} \pi.$$  

The values $\nu = 0, 1, 2, 3$ give the four possible solutions along both the axes, after which the same points get repeated. For LOCO-EFA, it is required (see further below) to limit the starting point to the semi-major axis only. To satisfy this condition, the second derivative of $E(\tau)$, evaluated at the temporal angle, should be negative, i.e., $\frac{d^2E(\tau)}{d\tau^2} |_{\tau_1} < 0$. Substituting the found solutions into the second derivative results in $\nu = 0$ and $\nu = 2$ belonging to the points along the semi-major axis whenever the denominator of the arctan term is positive, and the solutions $\nu = 1$ and $\nu = 3$ belonging to the points along the semi-major axis whenever the denominator of the arctan term is negative. A very straightforward computational implementation of this result is to make use of the four-quadrant inverse tangent function (atan2) as provided by most programming languages (i.e., such that $\text{atan2}(1,1) = \pi/4$ is different from $\text{atan2}(-1,-1) = -3\pi/4$).

Then, using

$$\tau_1 = 0.5\text{atan2} \left( \frac{2(\alpha_1 \beta_1 + \gamma_1 \delta_1)}{\alpha_1^2 + \gamma_1^2 - \beta_1^2 - \delta_1^2} \right)$$

automatically and unambiguously ensures that the temporal angle $\tau_1$ is located at one of the extremes of the semi-major axis.

This still leaves two ways to position the starting point (one for each of the extremes of the semi-major axis) and thereby two distinct representations of a same outline. We therefore further restrict $\tau_1$ to always lie within the first or second quadrant (I or II in Figure S8A). This is achieved by testing if the obtained $\tau_1$ that shifts the starting point of the first harmonic to the semi-major axis indeed positions it within quadrant I or II. To shift the starting point, we first introduce a time shift $\tau' = \tau - \tau_1$ such that at $\tau' = 0$ the first harmonic is positioned along its semi-major axis:
\[ [A_1] [M_1(t)] = [A_1] [M_1(t' + \tau_1)] , \]
\[ = [A_1] \begin{bmatrix} \cos (t' + \tau_1) \\
\sin (t' + \tau_1) \end{bmatrix} , \]  
(15a)
\[ = [A_1] \begin{bmatrix} \cos (\tau_1) \cos (t') - \sin (\tau_1) \sin (t') \\
\sin (\tau_1) \cos (t') + \cos (\tau_1) \sin (t') \end{bmatrix} , \]  
(15b)
\[ = [A_1] \begin{bmatrix} \cos (\tau_1) - \sin (\tau_1) \\
\sin (\tau_1) + \cos (\tau_1) \end{bmatrix} \begin{bmatrix} \cos (t') \\
\sin (t') \end{bmatrix} , \]  
(15c)
\[ = [A_1] \begin{bmatrix} \psi_{\tau_1} \end{bmatrix} [M_1(t')] , \]  
(15d)
\[ = [A_1] [M_1(t')] , \]  
(15e)

were \([\psi_{\tau_1}]\) is the rotation operator, rotating by an angle \(\tau_1\) and \([A_1'] = [A_1][\psi_{\tau_1}]\). The spatial angle at the shifted starting point \(\varrho\) is given by

\[ \varrho(\nu) = \arctan \left( \frac{\gamma_1'}{\alpha_1'} \right) + \nu \pi . \]  
(16)

Again, a single, unique and correct solution for \(\varrho\) can be obtained by using \(\varrho = \arctan2(\gamma_1', \alpha_1')\) instead. The starting point lies in quadrant III or IV when \(\varrho < 0\). In that case, \(\tau_1\) is modified as follows:

\[ \tau_1^* = (\tau_1 + \pi) . \]  
(17)

Otherwise (when the starting point is already in quadrant I or II), \(\tau_1^* = \tau_1\).

The new EFA coefficients corrected for the starting point then become:

\[ \begin{bmatrix} \alpha_n^* & \beta_n^* \\
\gamma_n^* & \delta_n^* \end{bmatrix} = \begin{bmatrix} \alpha_n & \beta_n \\
\gamma_n & \delta_n \end{bmatrix} \begin{bmatrix} \cos (n\tau_1^*) - \sin (n\tau_1^*) \\
\sin (n\tau_1^*) \cos (n\tau_1^*) \end{bmatrix} . \]  
(18)

Finally, we ensure that the direction of contour approximation of the first harmonic is always counter-clockwise (i.e., that \(r_1 \geq 0\), Equation 9). Besides removing redundancy by restricting the freedom of choice regarding the overall direction of contour approximation, this transformation also guarantees a unique correspondence between the properties of each subsequent harmonic and its contribution to the morphological features. When the direction of the first harmonic is clockwise \((r_1 < 0)\), we therefore invert the direction of motion of all ellipses, maintaining thereby their inter-relationships. This can be done by running “time” backwards:

\[ \begin{bmatrix} x(-t) \\
y(-t) \end{bmatrix} = \begin{bmatrix} \alpha_n^* & \beta_n^* \\
\gamma_n^* & \delta_n^* \end{bmatrix} \begin{bmatrix} \cos \left( \frac{2n\pi t}{T} \right) \\
\sin \left( \frac{2n\pi t}{T} \right) \end{bmatrix} \]  
(19)
\[ = \begin{bmatrix} \alpha_n^* & -\beta_n^* \\
\gamma_n^* & -\delta_n^* \end{bmatrix} \begin{bmatrix} \cos \left( \frac{2n\pi t}{T} \right) \\
\sin \left( \frac{2n\pi t}{T} \right) \end{bmatrix} . \]

In short, whenever \(r_1 < 0\), all indices \(\beta_n^*\) and \(\delta_n^*\) should be negated. After these steps, each unique cell contour is represented by a unique set of EFA coefficients. Note that the steps above do not alter the layout, nor do they rotate the shape. In certain study contexts, however, it might be desirable to rotate the contour itself, positioning
the semi-major axis, for example, to be parallel to the x-axis (or in any other preferred orientation). The details on how to perform those rotations can be found in Kuhl and Giardina (1982). Please note that unlike in their study, our subsequent analysis does not require such a cell contour realignment.

For simplicity of notation in the rest of the Supplementary Materials and Methods we refer to the \( [A_n] \) matrix, which elements have been normalised regarding the starting point and direction of reconstruction of the first harmonic:

\[
[A_n] \equiv \begin{bmatrix} a_n & b_n \\ c_n & d_n \end{bmatrix} \equiv \begin{bmatrix} \alpha_n^* & \beta_n^* \\ \gamma_n^* & \delta_n^* \end{bmatrix}.
\] (20)

After all possible sources of redundancy have been removed, the next step of the LOCO-EFA method is to split each elliptic harmonic into two counter-rotating circles.

(2) Decompose each \( n \)th elliptic harmonic into two circles with opposite direction of rotation

In order to find the contribution of \( n \)th harmonic to a given morphological feature, we rewrite the \( [A_n] \) matrices in Equation 10 such as to explicitly introduce the length of the semi-major and semi-minor axis of the \( n \)th ellipse (\( \lambda_{1n} \) and \( \lambda_{2n} \)).

For this purpose, it is necessary to introduce both the temporal and the spatial rotation operator of each elliptic harmonic, given by \( [\psi_{T_n}] \) and \( [\psi_{S_n}] \), respectively. The temporal operator is defined as

\[
[\psi_{T_n}] = \begin{bmatrix} \cos \phi_n & -\sin \phi_n \\ \sin \phi_n & \cos \phi_n \end{bmatrix},
\] (21)

and the spatial operator is defined as

\[
[\psi_{S_n}] = \begin{bmatrix} \cos \theta_n & -\sin \theta_n \\ \sin \theta_n & \cos \theta_n \end{bmatrix},
\] (22)

where \( \phi_n \) is the temporal angle (i.e., the time \( \tau_n \) required to rotate to the semi-major axis) and \( \theta_n \) the spatial angle (i.e., the angle of this position along the semi-major axis with the positive x-axis) (Figure S8A).

Equation 11 can be written as:

\[
[X(t)] = [A_0] + \sum_{n=1}^{N} [\psi_{S_n}] [\psi_{S_n}]^{-1} [A_n] [\psi_{T_n}] [\psi_{T_n}]^{-1} [M_n(t)],
\] (23)

given that \( [\psi_{S_n}] [\psi_{S_n}]^{-1} \) and \( [\psi_{T_n}] [\psi_{T_n}]^{-1} \) correspond to the identity matrix \([I]\).

Equation 11 can also be written in a form which directly highlights the contribution of the semi-major and semi-minor axis:

\[
[X(t)] = [A_0] + \sum_{n=1}^{N} [\psi_{S_n}] [A_n] [\psi_{T_n}]^{-1} [M_n(t)].
\] (24)

This equation can be understood as follows: for each mode, correctly position the starting point relative to the semi-major axis, transform the original circle into an ellipse, its semi-major axis along the x-axis and semi-minor axis along the y-axis, and finally rotate the ellipse to its correct position (see Figure S8B–D). Since both descriptions should
be equivalent, by combining Equation 23 and Equation 24, the $[\Lambda_n]$ matrix can be identified, with $\lambda_{1n}$ corresponding to the length of the semi-major axis, and the modulus of $\lambda_{2n}$ to the length of the semi-minor axis of the $n$th ellipse:

$$[\Lambda_n] = [\psi_{S_n}]^{-1} [A_n] [\psi_{T_n}] = \begin{bmatrix} \lambda_{1n} & 0 \\ 0 & \lambda_{2n} \end{bmatrix}. \quad (25)$$

Note that this process is similar to a singular value decomposition of $[A_n]$, with the difference that here $\lambda_{2n}$ (but not $\lambda_{1n}$) can be negative. The temporal angle $\phi_n$ corresponds, for each elliptic harmonic, to the scaled time $\tau_n$ to reach an extreme along the semi-major axis (Figure S8). Similarly to Equation 13, this corresponds to

$$\phi_n(\nu) = \frac{1}{2} \arctan \left( \frac{2(a_n b_n + c_n d_n)}{a_n^2 + c_n^2 - b_n^2 - d_n^2} \right) + \nu \frac{\pi}{2}. \quad (26)$$

Once again, using $\arctan$ ensures that $\phi_n$ corresponds to the semi-major axis (see above). For the next steps of the analysis (see below) it is essential that $\phi_n$ corresponds to the temporal angle to reach the semi-major axis, not the semi-minor axis, hence usage of $\arctan$ or any equivalent function which determines the quadrant of the return value is essential.

The spatial angle $\theta_n$ (see Figure S8) can be calculated after applying the temporal modification:

$$\theta_n(\nu) = \arctan \frac{c_n'}{a_n'} + \nu \pi, \quad (27)$$

where $c_n'$ and $a_n'$, are the new coefficients after the temporal transformation (i.e., after applying $[A_n] [\psi_{T_n}]$). Again, it is essential to use $\arctan$ to ensure that $\phi_n$ and $\theta_n$ are both relative to the same extreme of the semi-major axis. It does not matter, however, which of the two extremes is being used, hence for this step no check is required regarding the quadrants. Applying Equation 25 then provides $\lambda_{1n}$ and $\lambda_{2n}$. Deriving a temporal and spatial angle relative to the semi-major axis and both being related to the same extreme guarantees that $\lambda_{1n} \geq 0$ (while $\lambda_{2n}$ can be positive or negative, depending on the rotation direction of the rotor) and $|\lambda_{1n}| \geq |\lambda_{2n}|$.

Using the above, the EFA (Equation 11) can be rewritten as:

$$\begin{bmatrix} x(t) \\ y(t) \end{bmatrix} = [A_0] + \sum_{n=1}^{N} \begin{bmatrix} \cos \theta_n & -\sin \theta_n \\ \sin \theta_n & \cos \theta_n \end{bmatrix} \begin{bmatrix} \lambda_{1n} & 0 \\ 0 & \lambda_{2n} \end{bmatrix} \begin{bmatrix} \cos \phi_n & \sin \phi_n \\ -\sin \phi_n & \cos \phi_n \end{bmatrix} \begin{bmatrix} \cos \left( \frac{2n\pi t}{T} \right) \\ \sin \left( \frac{2n\pi t}{T} \right) \end{bmatrix}. \quad (28)$$

Written in this form, the contribution of each harmonic can easily be separated to correctly map to morphological feature number. The diagonal matrix containing the length of the semi-major and semi-minor axis of each $n$th mode can be decomposed into two diagonal matrices, each corresponding to circular orbits moving in opposite directions:

$$\begin{bmatrix} x(t) \\ y(t) \end{bmatrix} = [A_0] + \sum_{n=1}^{N} \begin{bmatrix} \cos \theta_n & -\sin \theta_n \\ \sin \theta_n & \cos \theta_n \end{bmatrix} \begin{bmatrix} \lambda_{+n} & 0 \\ 0 & \lambda_{+n} \end{bmatrix}$$

$$+ \begin{bmatrix} \lambda_{-n} & 0 \\ 0 & -\lambda_{-n} \end{bmatrix} \begin{bmatrix} \cos \phi_n & \sin \phi_n \\ -\sin \phi_n & \cos \phi_n \end{bmatrix} \begin{bmatrix} \cos \left( \frac{2n\pi t}{T} \right) \\ \sin \left( \frac{2n\pi t}{T} \right) \end{bmatrix}, \quad (29)$$
where $\lambda_{+n}$ and $\lambda_{-n}$ are the radii of those circles (Figure 2). Summing up the diagonal matrices in Equation 29 yields

$$
\begin{bmatrix}
x(t) \\
y(t)
\end{bmatrix}
= [A_0] + \sum_{n=1}^{N} \begin{bmatrix}
cos \theta_n & -\sin \theta_n \\
\sin \theta_n & \cos \theta_n
\end{bmatrix}
\begin{bmatrix}
\lambda_{+n} + \lambda_{-n} & 0 \\
0 & \lambda_{+n} - \lambda_{-n}
\end{bmatrix}
\begin{bmatrix}
\cos \phi_n & \sin \phi_n \\
-\sin \phi_n & \cos \phi_n
\end{bmatrix}
\begin{bmatrix}
cos \left(\frac{2\pi nt}{\lambda_{+n}}\right) \\
sin \left(\frac{2\pi nt}{\lambda_{+n}}\right)
\end{bmatrix},
$$

(30)
in which the major and minor axes of each elliptic harmonic are

$$
\begin{align*}
\lambda_{1n} &= \lambda_{+n} + \lambda_{-n}, \\
\lambda_{2n} &= \lambda_{+n} - \lambda_{-n},
\end{align*}
$$

(31a, b)

and hence the radii of each oppositely-rotating circle is given by

$$
\begin{align*}
\lambda_{+n} &= \frac{\lambda_{1n} + \lambda_{2n}}{2}, \\
\lambda_{-n} &= \frac{\lambda_{1n} - \lambda_{2n}}{2}.
\end{align*}
$$

(32a, b)

Given that $\lambda_{1n} \geq 0$ and $|\lambda_{1n}| \geq |\lambda_{2n}|$, $\lambda_{+n}$ and $\lambda_{-n}$ are always positive. To approximate the cell contour $(x(t), y(t))$ using the circles $\lambda_{+n}$ and $\lambda_{-n}$ requires completing the transformations using the spatial ($\theta_n$) and temporal angle ($\phi_n$) as calculated before, most clearly seen through the expression

$$
[X(t)] = [A_0] + \sum_{n=1}^{N} [\psi_{Sn}] [\Lambda_{+n} + \Lambda_{-n}] [\psi_{Tn}]^{-1} [M_n(t)].
$$

(33)
The term $\Lambda_{+n}$ presents the subset of the $n$th elliptic harmonic which is moving in the same direction as the first harmonic, therefore purely contributing to $n - 1$ “lobes”, i.e., shape features with periodicity $n - 1$. In contrast, $\Lambda_{-n}$ presents the subset moving in the opposite direction, purely contributing to $n + 1$ “lobes” (shape features) only. Their contributions can be separated by writing:

$$
[X(t)] = [A_0] + \sum_{n=1}^{N} [A_{+n}] [M_n(t)] + \sum_{n=1}^{N} [A_{-n}] [M_n(t)],
$$

(34)

where

$$
[A_{jn}] = [\psi_{Sn}] [\Lambda_{jn}] [\psi_{Tn}]^{-1}, \text{ for } j = +, -.
$$

(35)

This can be further simplified. Straightforwardly, $[\psi_{Sn}] [\Lambda_{+n}] = [\psi_{Sn}] \lambda_{+n} [I] = [\Lambda_{+n}] [\psi_{Sn}]$. Regarding $[\Lambda_{+n}]$,

$$
[\psi_{Sn}] [\Lambda_{-n}] = \begin{bmatrix}
\cos \theta_n & -\sin \theta_n \\
\sin \theta_n & \cos \theta_n
\end{bmatrix}
\begin{bmatrix}
\lambda_{-n} & 0 \\
0 & -\lambda_{-n}
\end{bmatrix},
$$

(36a)

$$
= \begin{bmatrix}
\lambda_{-n} \cos \theta_n & +\lambda_{-n} \sin \theta_n \\
\lambda_{-n} \sin \theta_n & -\lambda_{-n} \cos \theta_n
\end{bmatrix},
$$

(36b)

$$
= \begin{bmatrix}
\lambda_{-n} & 0 \\
0 & -\lambda_{-n}
\end{bmatrix}
\begin{bmatrix}
\cos \theta_n & \sin \theta_n \\
-\sin \theta_n & \cos \theta_n
\end{bmatrix},
$$

(36c)

$$
= \begin{bmatrix}
\lambda_{-n} & 0 \\
0 & -\lambda_{-n}
\end{bmatrix}
\begin{bmatrix}
\cos (-\theta_n) & -\sin (-\theta_n) \\
\sin (-\theta_n) & \cos (-\theta_n)
\end{bmatrix},
$$

(36d)

$$
= [\Lambda_{-n}] [\psi_{Sn}]^{-1}.
$$

(36e)
Thus,

\[ [A_{+n}] = [A_{+n}] \left[ \psi_{s_{n}} \right] [\psi_{r_{n}}]^{-1}, \]
\[ [A_{-n}] = [A_{-n}] \left[ \psi_{s_{n}} \right] [\psi_{r_{n}}]^{-1}. \]

Hence, introducing \( \zeta_{+n} = \theta_{n} - \phi_{n} \) and \( \zeta_{-n} = -\theta_{n} - \phi_{n} \), Equation 35 can be written as

\[ [A_{jn}] = [A_{jn}] \left[ \psi_{\zeta_{jn}} \right], \text{ for } j = +, - , \]

which corresponds to

\[ [A_{jn}] = \begin{bmatrix} a_{jn} & b_{jn} \\ c_{jn} & d_{jn} \end{bmatrix} \]
\[ = \begin{bmatrix} \lambda_{jn} & 0 \\ 0 & j\lambda_{jn} \end{bmatrix} \begin{bmatrix} \cos \zeta_{jn} & -\sin \zeta_{jn} \\ \sin \zeta_{jn} & \cos \zeta_{jn} \end{bmatrix}, \text{ for } j = +, - . \]

We next label those matrices with respect to their lobe contribution instead of their EFA mode. To make the distinction, we here use subscript \( n \) to indicate the EFA mode, and subscript \( l \) to indicate the LOCO-EFA mode. In general, \([A_{+}] = [A_{+n+1}] \) and \([A_{-}] = [A_{-n-1}] \). There are, however, a few exceptions, see Figure S4:

1) The overall offset of the contour is not solely given by \( [A_{0}] \). An additional contribution to the offset is coming from \([A_{n=0}] \). Note, however, that the contribution from \([A_{n=2}] \) is in fact not a perfect offset to the contour, but also causes a kidney bean-shaped distortion to the contour, its deviation from a pure offset becoming more pronounced when the contribution of this mode relative to the overall contour size is larger (see Figure S4B).

2) The overall circular shape of the contour (i.e., LOCO-mode 1) receives solely a contribution from \([A_{+n=1}] \) itself, i.e., \([A_{+n=1}] = [A_{+n=1}] \).

3) When \( N \) EFA modes are taken into account, then LOCO-mode \( N \) only receives a contribution from \([A_{-n=N-1}] \).

4) Likewise, when taking \( N \) EFA modes into account, there is still a contribution to LOCO-mode \( L = N + 1 \), solely coming from \([A_{-n=N}] \).

Defining \([M_{+}] = [M_{n+1}] \) and \([M_{-}] = [M_{n-1}] \), with the exceptions \([M_{+t=0}] = [M_{n=2}] \) and \([M_{-t=0}] = [M_{n=1}] \), then the same shape approximation can be achieved through LOCO-EFA as through EFA:

\[ [X(t)] = [A_{0}] + \sum_{L-2}^{L} [A_{+}] [M_{+t=0}(t)] + \sum_{L}^{L} [A_{+}] [M_{+t=1}(t)] + \sum_{L=2}^{L} [A_{+}] [M_{+t=l}(t)] + \sum_{l=2}^{L} [A_{-}] [M_{-t=l}(t)]. \]

Equation 40 can be used to reconstruct the shape up to a certain \( L \) number, requiring EFA coefficients up to mode \( L + 1 \):

\[ [X_{L=1}(t)] = [A_{0}] + [A_{+t=0}] [M_{+t=0}(t)] + [A_{+t=1}] [M_{+t=1}(t)], \]
\[ [X_{L=2}(t)] = [A_{0}] + [A_{+t=0}] [M_{+t=0}(t)] + [A_{+t=1}] [M_{+t=1}(t)] + \sum_{l=2}^{L} [A_{+}] [M_{+t=l}(t)] + \sum_{l=2}^{L} [A_{-}] [M_{-t=l}(t)]. \]
Each LOCO-EFA mode can be fully described by the combination of the radii and the starting positions of both circles. The radii are, as previously defined, $\lambda_+ = \lambda_{n+1}$ and $\lambda_- = \lambda_{n-1}$, with the exceptions $\lambda_{i=0} = \lambda_{n+2}$ and $\lambda_{i=0} = 0$, and $\lambda_{i=1} = \lambda_{n+1}$ and $\lambda_{i=1} = 0$. The starting points of both circles are defined as $\psi_{i=0} = \psi_{n+2}$ and $\psi_{i=0} = \psi_{n+2}$. Together this gives a set of four coefficients $L_1 = (\lambda_+, \lambda_-, \psi_+, \psi_-)$ that fully capture each LOCO-EFA mode and allow for a full reconstitution of the original shape:

$$[X(t)] = [A_0] + \sum_{l=2}^{L-2} [\lambda_{i=0}] [\psi_{i=0}] [M_{i=0}(t)] + [\lambda_{i=1}] [\psi_{i=1}] [M_{i=1}(t)] + \sum_{l=2}^{L} [\lambda_{i=0}] [\psi_{i=1}] [M_{i=1}(t)] .$$

(42)

The exceptions for the lowest modes, as depicted in Figure S4A might appear counterintuitive. Their particular repairing is however a direct consequence of how we have chosen to define mode 0 and mode 1. Our choice was driven by seeking to stay more in line with classical EFA. As in EFA, we have defined mode 0 as the offset of the contour with respect to the underlying coordinate system, and therefore mode 1 as the radius of the circle, the most basic capturing of the shape itself. LOCO-EFA modes, in contrast, should be viewed as corresponding to a specific number of perturbations along that circle (for its mathematical derivation see further details in the next session). Hence, an alternative choice of definitions could have been to denote the radius of the circle mode 0. The unidirectional perturbation of the circle as provided by EFA modes 0 and 2 could be interpreted as a ‘single lobe’, and could therefore be denominated mode 1, etc. Although such a defendable renaming of LOCO-EFA modes would make Figure S4A free of any apparent exceptions in mode reassignment, we preferred to maintain the definitions as here presented, for sake of clarity. The reason being that, although it introduces exceptions that carefully have to be taken care of, the chosen denomination captures, alike EFA, within mode 0 the non-oscillatory “DC” contour offset. Renaming that to mode 1 would in our eyes be more unnatural than to simply reassign modes as is currently done.

In the next section we derive how from those values a single amplitude value can be found, the $L_i$ contribution.

(3) $L_i$ amplitude contributions

In this section, we first show that the contribution to the shape of the two counter-rotating circles of a specific LOCO-EFA mode is not simply determined by the radii of those circles, but also depends on the relative position of the starting points. We then derive a heuristic which optimally captures its contribution, determining as well the limits of this approximation. To illustrate how a difference between the starting points $\psi_{i=0}$ and $\psi_{i=0}$ of mode $l$ can affect the generated amplitude, we first plot the contribution of $L_3 = (\lambda_{i=0}, \lambda_{i=0}, \psi_{i=0}, \psi_{i=0})$ superimposed on $L_1 = (1, 0, 0, 0)$, i.e., superimposed on the unit circle starting at zero degrees. As long as $\lambda_{i=0}$, $\lambda_{i=0}$ are not too large, a single amplitude value $a$ (which is the amplitude of deviation from the unit circle) can be observed for any phase $\omega$; peak amplitude values $A$ occur for specific phases $\Omega$ (as illustrated in Figure S9F). Figure S9A–C depict the shape contribution of the negative rotor (green), positive rotor (red) and their summed contribution (orange), as a function of the phase $\omega$, for $L_3 = (0.15, 0.15, 0, 0)$. This scenario illustrates that even though the negative rotor makes two sidereal rotations while the positive rotor makes four sidereal rotations, the amplitude
pattern that they generate not only has a period three in both cases, as argued throughout
the paper, but also perfectly match one another regarding the phases at which the peaks
and troughs in the amplitude pattern are reached. The summed contribution (Figure S9C)
therefore indeed yields a peak amplitude exactly equal to \( \lambda_{+3} + \lambda_{-3} \). In contrast, when
both rotors are exactly out-of-phase (\( L_3 = (0.15, 0.15, 0, \pi) \), Figure S9D–F), the peaks
and troughs are exactly out-of-phase as well, and the patterns almost (but not totally)
cancel each other out (Figure S9F). To determine the effective contribution of a LOCO-
EFA mode we therefore have to determine the phase at which each rotor reaches its peak
amplitude.

The angles \( \Omega_+ \), \( \Omega_- \) at which the positive and negative rotor reach their peak amplitude
can be calculated straightforwardly. They occur when the phase of the rotor itself is
equivalent to the overall phase generated by \( L_1 \) (as illustrated in Figure S9G). The phase
of \( L_1 \) starts at \( \zeta_+ + 1 \), while the phase of the positive rotor starts at \( \zeta_+ \) (illustrated in
Figure S9K). When the pattern is laid down, the phase of the positive rotor changes
\((l + 1)\) times faster than the phase of \( L_1 \) (Equation 42). Thus, regarding the phase at
peak amplitude it holds that

\[
\begin{align*}
\Omega_+ &= (l + 1)(\Omega_+ - \zeta_+) + \zeta_+ + 2\pi \nu, \\
-l\Omega_+ &= \zeta_+ - (l + 1)\zeta_+ + 2\pi \nu, \\
\Omega_+ &= \zeta_+ + \frac{\zeta_+ - \zeta_-}{l} + \frac{2\pi}{l} \nu.
\end{align*}
\]

where \( \nu \) is any integer, and the values \( \nu = 0, 1, \ldots, l - 1 \) provide the complete set of
phases at which peak amplitude is reached. Likewise, the negative rotor rotates \((l - 1)\)
times faster than \( L_1 \) and in the opposite direction, hence starting at \(-\zeta_-\) (see Figure S9J).
Regarding the phase at peak amplitude it therefore holds that

\[
\begin{align*}
\Omega_- &= -(l - 1)(\Omega_- - \zeta_-) - \zeta_- + 2\pi \nu, \\
l\Omega_- &= -\zeta_- + (l - 1)\zeta_+ + 2\pi \nu, \\
\Omega_- &= \zeta_+ - \frac{\zeta_+ + \zeta_-}{l} + \frac{2\pi}{l} \nu.
\end{align*}
\]

To assess the summed amplitude contribution, we next fit the amplitude pattern laid
down by each rotor to a sine wave, exactly matching both the peak amplitude and the
phase at which the peak amplitude is reached, and then sum those two contributions:

\[
\begin{align*}
a_+ &= \lambda_+ \cos(lw + \zeta_+ - (l + 1)\zeta_+) , \\
a_- &= \lambda_- \cos(lw + \zeta_- - (l - 1)\zeta_+) , \\
a_+ + a_- &= \lambda_+ \cos(lw + \zeta_+ - (l + 1)\zeta_+) + \lambda_- \cos(lw + \zeta_- - (l - 1)\zeta_+). 
\end{align*}
\]

Equation 45a exactly matches the peak amplitude and its phase of the positive rotor, as
derived in Equation 43, while Equation 45b exactly matches the peak amplitude and its
phase of the negative rotor, as derived in Equation 44. While peak amplitude and phase
match perfectly, Figure S9G, H illustrate that for not too large values of \( \lambda_{+1}, \lambda_{-1} \) also the
rest of the pattern presents a close match.

Using standard trigonometry, the summed amplitude can be written as
\[ a_{+l} + a_{-l} = a_l = \sqrt{\lambda_{+l}^2 + \lambda_{-l}^2 + 2\lambda_{+l}\lambda_{-l}\cos(\zeta_{+l} - \zeta_{-l} - 2\zeta_{+l})\cos(l\omega + \zeta_l)}, \]  

where

\[ \zeta_l = \text{atan2}(\lambda_{+l} \sin((\zeta_{+l} - (l + 1)\zeta_{+l})) + \lambda_{-l} \sin((\zeta_{-l} - (l - 1)\zeta_{+l}) + \lambda_{-l} \cos((\zeta_{-l} - (l - 1)\zeta_{+l})). \]  

Figure S9I illustrates that for not too large values of \( \lambda_{+l}, \lambda_{-l} \) this expression provides a close match to the pattern generated by the \( L_l \) mode, here illustrated using both a different phase and a different amplitude for both rotors. Using the equation above, the amplitude or contribution of mode \( l \) can hence be defined as

\[ L_l = \sqrt{\lambda_{+l}^2 + \lambda_{-l}^2 + 2\lambda_{+l}\lambda_{-l}\cos(\zeta_{+l} - \zeta_{-l} - 2\zeta_{+l})}. \]  

While above we have performed a more formal derivation, a more intuitive way to understand the amplitude contribution of the positive and negative rotor combined, is in terms of how much the amplitude contributions of both rotors are out-of-phase with each other. The most straightforward moment to determine their phase shift is at the initial point of drawing the cell’s outline. As illustrated in Figure S9J, K, the phase shift regarding the amplitude contribution of the negative rotor is given by \( \zeta_{+l} - \zeta_{+l} \), and of the positive rotor by \( \zeta_{+l} - \zeta_{+l} \). To understand the graphical argument for \( \zeta_{+l} - \zeta_{+l} \), one has to realise that the negative rotor rotates clockwise, and hence its starting angle is \( -\zeta_{+l} \) (for the positive rotor it is simply \( \zeta_{+l} \)), while for the same reason the phase shift between initial amplitude and maximum amplitude has to be calculated clockwise (instead of standard counterclockwise, as done for the positive rotor), again as shown in Figure S9J, K. Plotting the phase shift and strength of both amplitude contributions as vectors and then adding them up then provides the above equation for \( L_l \) (as is depicted in Figure 2C).

Note that the pattern can never be perfectly described by a sine wave, which is why four parameters are needed to describe each mode, rather than the two required by Equation 46. Even when following Equation 46 the two waves cancel each other out, this is not exactly the case, as seen in Figure S9F. There is therefore no additional level of redundancy, as the usage of the \( L_l \) numbers might suggest, even when the value of \( L_l = 0 \). When the amplitude of a rotor becomes larger, the generated pattern starts to deviate from a sine wave (Figure S9J, K), and their summed pattern from Equation 46 (Figure S9L). For simple closed contours (which cell outlines should always be), the deviations cannot be very large, since otherwise the contour becomes non-simple (i.e., crosses itself). This confinement in deviations for cell outlines allows us to base a significant part of our analysis on \( L_l \) values.

Moreover, we observe that the positive rotor generates protrusions that are flatter than a sine wave (Figure S9B, J), whereas the negative rotor generates lobes that are pointier (Figure S9A, K). The extent of “flattening” or “pointedness” of each mode can be determined by the proportion \( w_{+l} = \lambda_{+l}/(\lambda_{+l} + \lambda_{-l}) \) and \( w_{-l} = 1 - w_{+l} \), respectively; a useful additional measure if needed.

The method presented here cannot be straightforwardly extended to 3D. The essential issue is that this whole analysis hinges on a parametrisation which moves along a directed curve. As no directed curve can be constructed to represent a cell outline in 3D, there is no simple extension from our 2D LOCO-EFA to 3D.
**XOR measurement**

LOCO-EFA can be applied to any closed-curve cell outline defined by a series of connected points. There are no requirements regarding the number of points or the distance between them, nor do their positions have to be integer or positive values. Nevertheless, the cell outlines that will be analysed with our method are generally derived from segmentation of microscopy images. Many segmentation methods are available, with most of them assigning each pixel within a microscopy image to a specific cell domain.

In the paper we show that, besides directly looking at the $L_n$ values, a highly informative analysis is to capture the quality of cell shape reconstruction when only a certain number of modes are taken into account. This is done by applying an EXCLUSIVE OR (XOR) between the segmented cell and reconstructed cell. The natural choice for the grid on which the XOR is performed is therefore equivalent to the microscopy image itself, at the resolution the image has been acquired and segmentation performed, as is done for all the experimental images analysed in this study. In such a case, the cell contour, i.e., the edge of the segmented cell, takes up integer values (when expressed in grid points), with the distance between points being either 1 (when the cell edge moves horizontally or vertically) or $\sqrt{2}$ (when the cell edge moves diagonally). While the input of the LOCO-EFA analysis is a set of discrete points, the output that it provides is a continuous curve. The LOCO-EFA provides an infinite series of modes, passing increasingly closer to the original set of discrete points $(x_i, y_i)$ at parametrised time $t_i$ when more modes are taken into account. (For a derivation of the predicted bound on error as a function of mode number for the classical EFA, see (Kuhl and Giardina, 1982).) To reconstruct from that curve a cell shape on a grid therefore requires two steps: 1) Determine the grid points at which the curve passes at each time $t_i$. Please note that the time series $t_i$ has to correspond to the “time” passed while drawing the discrete chain of original contour points, as discussed below Equation 5, and hence will be a time series containing irregular intervals (of either 1 or $\sqrt{2}$). The reason is that by only evaluating the continuous curve at those time points, for sufficiently many modes the curve is guaranteed to lie within the grid point defining the original cell outline. In between those time points, however, the curve could very well cross through grid points that do not belong to the cell outline, especially when the cell outline involves a diagonal step. 2) Only in the case that the next reconstructed contour point lies beyond the Moore neighbourhood of the previous contour point, iteratively reduce the time steps between these two reconstructions (deviating from the specific time points as defined above), in order to acquire on the grid a contiguous chain of pixels defining the cell outline. Doing so allows for flood filling at the end to obtain the complete cell shape rather than just the cell outline, and to then apply an XOR with the original segmented image.

LOCO-EFA provides infinitely many modes, for which it is at forehand not defined where a required cut-off should be placed, since the approximation becomes asymptotically more precise. The XOR analysis, however, provides an algorithmic and objective cut-off for any LOCO-EFA data analysis, not only the XOR analysis. When the reconstruction is performed as explained above, then for a sufficiently large, but finite number of modes each individual contour grid point will be correctly reconstructed, which leads to the XOR value becoming zero. Adding additional terms will only change the reconstructed outline at a sub-pixel resolution, i.e., at a resolution that is higher than the original microscopy image. Obviously, the latter cannot be meaningful in any possible way. Using the XOR reaching zero therefore provides a natural cut-off to truncate the
series, as higher-mode contributions can only be informative provided the XOR is still non-zero, but cannot be informative beyond that point. In our experience, this point is usually reached between 20 and 200 modes, but will be cell-complexity and image-resolution dependent. Note that additional information acquired outside of the method could justify earlier cut-offs, for example when there is information available regarding the quality of segmentation.

Final remarks

To summarise, the LOCO-EFA method consists in: 1) eliminating degeneracies in the EFA coefficients; 2) decomposing each elliptic harmonic into two circles rotating in opposite directions ($\lambda_+n$ and $\lambda_-n$) and therefore contributing to $n - 1$ and $n + 1$ number of lobes (more generally, morphological features); and 3) calculating the offset between starting points of these two circles derived from each ellipse to estimate the amplitude of the $L_i$th lobe contribution.

To eliminate cell area effects (for example, when looking at shape diversity within a cell populations), it might also be desirable to normalise for cell size by dividing each $L_n$ value by the square root of the cell area. This then provides a complete description of the number of lobes and their amplitude, which can be used to characterise and quantify intrinsic cell shape properties, irrespective of cell area, spacing between sampling points, and rotations or inversions of the cell (Figure S10A, B). Changes in the image resolution, however can of course affect the fine-grained information retrieval, but only if the resolution becomes very low (Figure S10C, D).
Supporting Figures

Figure S1: **Subjective choices involved in neck width and lobe length determination and Skeletonisation.** (A) Neck width and lobe length depend on human criteria for both identifying and quantifying such structures, as indicated by the question-marks. (B) Both the number of skeleton end-points and the length of the branches strongly depend on the parameter settings used for the skeletonisation algorithm. Here, parameter $p$ (see material and Methods) was set to 45 (left), 65 (middle), or 85 (right). For those values, the number of predicted lobes varies between 6 and 8.
Figure S2: **Holomorphic and non-holomorphic shapes.** (A) In holomorphic shapes, all radii starting from the centroid intersect the outline only once. (B) In non-holomorphic shapes, some radii intersect more than once. Very few pavement cells have a holomorphic shape, the majority presenting highly complex non-holomorphic outlines. The outline of non-holomorphic shapes cannot be represented as a function of the angle, precluding, for example, standard Fourier analysis.
Figure S3
Figure S3: **Cell outline reconstruction using EFA and LOCO-EFA.** (A) Calculating the EFA coefficients from a cell outline. The discrete chain of contour points can be positioned arbitrarily (for example, the points do not have to be associated to an underlying grid). Also, the distances between points can be arbitrarily long. (B) Sequential approximation of the cell’s contour. The first harmonic forms an elliptic shape (1, blue). The second harmonic describes an elliptic orbit (2–6, red), orbiting twice while moving around the first harmonic (2, 3). Their summed trajectories are shown in green. The third elliptic harmonic (4–6, orange) orbits thrice while moving around this summed trajectory (4–6). The summation of the first three elliptic harmonics is shown in blue. See Movie 1 for this dynamical reconstitution of the cell contour. (C) The EFA coefficients cannot be directly linked to shape features, here shown through the power contribution of each harmonic ($P_n$) (upper panel): the main $P_n$ contributions of a six-pointed star-like shape (shown as an inset) come from the 5th and 7th harmonic. To align the coefficients to actual shape features, Diaz et al. (1990) proposed to shift the contribution of each harmonic to either $n + 1$ or $n - 1$, depending on the rotation direction of each individual harmonic (middle panel). This brings the main shape contributor and the actual number of shape features in alignment to each other. This method, however, does not always hold (see F), and moreover generates a range of spurious contributions from a large set of different modes, which is a notorious issue that hampers analysis of complex shapes using standard EFA (Haines and Crampton, 2000). The LOCO-EFA method correctly aligns the shape assessment and the real shape features (lower panel), without generating any additional spurious contributions. Moreover, unlike the other methods, the values correctly represent the amplitude of the shape features. (D–F) Contours (shown in blue), generated from the first (green) and third EFA harmonic (orange) only. The number of morphological protrusions (lobes) specified by the $n$th EFA mode is affected, but not fully determined, by its rotation direction. The heuristic rule proposed by Diaz et al. (1990) states that if the first harmonic and the $n$th harmonic rotate in the same direction, a contour is generated with $n - 1$ protrusions, while if their rotation direction is opposite, the contour will contain $n + 1$ lobes. This rule, however, is only correct when the elliptic harmonic has a circular shape (D, E), in which case indeed overall shapes are generated with either 2 (D) or 4 lobes (E), simply dependent on the rotation direction with respect to the first mode. When, however, the elliptic harmonic has a higher eccentricity (F), the final shape can have $n + 1$ protrusions (4 lobes) even though the rotation direction of the first and third harmonic are the same. In D–F the determinant of the first harmonic is $-1.0002$ (i.e., rotating clockwise), while the determinants of the third harmonics are $-6.554 \cdot 10^{-2}$ (clockwise), $6.446 \cdot 10^{-2}$ (counter-clockwise), and $-1.19 \cdot 10^{-4}$ (clockwise), respectively. The heuristic fails as a consequence of each single EFA mode actually contributing to two different spatial modes. See also Movie 2–4.
Figure S4: Schematic mapping between EFA modes and LOCO-EFA modes. (A) Each $n$th EFA mode contributes to both $n+1$ and $n-1$ morphological periodicities. The red arrows represent the contributions of the $n$th EFA mode to $n+1$ protrusions, due to the clockwise rotations of the circular harmonics $\lambda_{-n+1}$. The blue arrows indicate the contributions to $n-1$ protrusions, due to the counter-clockwise rotations of the circular harmonics $\lambda_{n-1}$. A few exceptions apply: The second EFA mode contributes to a shift in the positioning of the layout, i.e., to LOCO-EFA mode 0 ($\lambda_0$, blue and dashed line), rather than to the overall size of the layout, as might have been expected. The first EFA mode contributes to the overall circular size of the layout ($\lambda_1$, blue and dashed line), rather than to a shift in the positioning of the layout, as might have been expected. The zeroth EFA mode only contributes to a shift in the positioning of the layout (yellow line). Finally, the two highest LOCO-EFA modes have incomplete contributions, given any cutoff in the number of EFA modes. (B–E) The contribution $\lambda_0$ is not simply an offset of the contour, but also involves a kidney bean-shaped distortion, more pronounced for larger contributions. Mode $\lambda_1$ (the circular mode) is shown in blue; mode $\lambda_0$ in red; their summation in green; and the predicted shape if the contribution were purely an offset in magenta. (B) $\lambda_0$ being 10% of $\lambda_1$, effectively resulting in just an offset to the shape outline (C). (D, E) $\lambda_0$ being 30% of $\lambda_1$, resulting in a clear kidney bean-shaped distortion of the shape.
Figure S5: Additional example of LOCO-EFA metrics on a cell changing its shape over time. (A–G) Sequence of a tracked pavement cell growing over time with normalised area. (H) $P_n$, XOR difference and marginal difference profiles, cumulative difference and entropy using EFA. (I) $L_n$, XOR difference and marginal difference profiles, cumulative difference and entropy using LOCO-EFA. See also the first example presented in Figure 4 in the main text.
Figure S6
Figure S6: Confluent in silico cell populations, simulated with three types of specified cell shapes (SCSs) and different number of specified lobes. Parameters for the different specified cell shapes are given in Table S1. (A–H) Cells with large protrusions (Specified Cell Shapes 1, SCS1), with specified lobe number increasing from 3 (A) to 10 (H); (I–P) Cells with small protrusions (SCS2), with specified lobe number increasing from 3 (I) to 10 (P); (Q–X) Elongated cells (SCS3), with specified lobe number increasing from 3 (Q) to 10 (X). Specified cell shapes, as resulting from single-cell simulations, are shown above each panel. Cells are randomly coloured. See also Figure 5 and Figure S7.
Figure S7: Divergence between specified cell shapes and resultant population-level cell shape diversity within confluent in silico cell populations. The mismatch is visualised as the difference in LOCO-EFA-derived measures between the average for confluent population simulations and its value for single cell simulations. Three types of specified cell shapes (SCSs) were simulated, as indicated for each column, and parametrised in Table S1. Lobe numbers vary from 3 to 10, as indicated for each row and depicted in Figure S6. (A–C) Difference between the average $L_n (\langle L_n \rangle)$ values of the cells of the simulated populations and that of a single simulated cell. (D–F) Difference between average XOR ($\langle\text{XOR}\rangle$) of the cells of the simulated population and that of a single simulated cell. (G–I) Same as in (A–C, D–F), but now for marginal difference (Md). In all cases the specified cell shape becomes less pronounced in the confluent population simulations (indicated by negative values that all measures yield at the given specified lobe number), while the cells present an increased shape diversity and complexity (seen by the broad flanking regions with positive values in the profiles, indicating a large range of modes that contribute to the shapes). These effects are present regardless of the number of specified lobes or the combination of parameters used, but becomes less pronounced at higher lobe numbers. See also Figure 5. Each depicted line is based upon averaging all cells within three independent tissue simulations using the same shape descriptors; each simulated tissue contains in between 42 and 70 in silico cells.
Figure S8
Figure S8: **Temporal and spatial transformations required to calculate the precise contribution of each EFA mode.** (A) The blue ellipse depicts an elliptic harmonic, given by $[X(t)] = [A][M(t)]$, while the grey circle depicts the unit circle, given by $[X(t)] = [M(t)]$. The temporal angle $\phi$ is the scaled time required to move along an elliptic harmonic from the starting point at $\tau = 0$ to one extreme along the semi-major axis (green filled circles). The angle $\phi$ cannot be trivially derived from the spatial position at $\tau = 0$, requiring first an effective projection upon the unit circle (green open circles). The spatial angle $\theta$ is the inclination of the elliptic harmonic. After applying the appropriate spatial and temporal transformations to the EFA coefficients using this geometrical interpretation, the semi-major and semi-minor axis, $\lambda_1$ and $\lambda_2$, can be straightforwardly obtained. To eliminate multiple representations of the same outline, the starting point of the first harmonic is specifically positioned at the extreme along the semi-major axis which lies in either quadrant I or II, given by temporal angle $\tau^*_1$ (Equation 18). In contrast, all other temporal angles $\phi_n$ used, while also positioned along the semi-major axis, are not confined to quadrant I or II. (B–E) Visual interpretation of Equation 24 as a step-wise construction of the elliptic harmonic. The matrix $[M(t)]$ corresponds to the unit circle (B). For each mode, first the starting point relative to the semi-major axis is correctly positioned (C); then the original circle is transformed into an ellipse, its semi-major axis along the x-axis and semi-minor axis along the y-axis (D); and finally, the ellipse is rotated to its correct position (E). See further details in the Supplementary Materials and Methods.
Figure S9
Figure S9: Impact of LOCO-EFA starting points $\zeta_+$ and $\zeta_-$ on the amplitude of the reconstructed shape. All panels show $L_3 = (\lambda_{i3}, \lambda_{-3}, \zeta_{+3}, \zeta_{-3})$ superimposed on $L_1 = (1, 0, \zeta_{+1}, 0)$, i.e., the unit circle. In (A–F), $L_1 = (1, 0, 0, 0)$. (A) Amplitude $a$ as a function of phase $\omega$ (as depicted in I) for the negative rotor only ($L_3 = (0, 0.15, 0, 0)$), shown in green. For several phases also the rotor (light blue) itself is drawn. Note that by plotting the rotor in the $(\omega, a)$ plane, the circles are slightly deformed. (B) Same for the positive rotor only ($L_3 = (0.15, 0, 0, 0)$), in red. (C) Same for both rotors superimposed ($L_3 = (0.15, 0.15, 0, 0)$), in orange. (D–F) Alike (A–C), but for an out-of-phase starting angle of the negative rotor. (D) $L_3 = (0, 0.15, 0, \pi)$, in green; (E) $L_3 = (0.15, 0, 0, 0)$, in red; (F) $L_3 = (0.15, 0.15, 0, \pi)$, in orange. In (G–L), $L_1 = (1, 0, 2/3\pi, 0)$. (G) Pattern generated by the negative rotor only ($L_3 = (0, 0.1, 0, -5/12\pi)$), shown in green, compared to the equation $a_{-l} = \lambda_{-l} \cos(l\omega + \zeta_{-l} - (l - 1)\zeta_{+l})$ (Equation 45b), in dark blue. They present a close match. (H) Same for the positive rotor only ($L_3 = (0.075, 0, 3/4\pi, 0)$), shown in red, compared to $a_{+l} = \lambda_{+l} \cos(l\omega + \zeta_{+l} - (l + 1)\zeta_{+l})$ (Equation 45a), show in dark blue, again presenting a close match. (I) Same for both rotors superimposed ($L_3 = (0.075, 0.1, 3/4\pi, -5/12\pi)$), shown in orange, and $a_l = \sqrt{\lambda_{+l}^2 + \lambda_{-l}^2} + 2\lambda_{+l}\lambda_{-l} \cos(\zeta_{+l} - \zeta_{-l} - 2\zeta_{+l}) \cos(l\omega + \zeta_l)$ (Equation 46), show in dark blue, again presenting a close match. Panel (G) also illustrates that at peak amplitude the phase of the main and subrotor are equal; panel (I) also illustrates the concepts amplitude ($a$); peak amplitude ($A$); phase ($\omega$); and phase at peak amplitude ($\Omega$). (J–L) Alike (G–I), but for larger amplitude. (J) $L_3 = (0, 0.2, 0, -5/12\pi)$, in green; (K) $L_3 = (0.15, 0, 3/4\pi, 0)$, in red; (L) $L_3 = (0.15, 0.2, 3/4\pi, -5/12\pi)$, in orange. Panel (J) and (K) also illustrate how the initial phase shifts with respect to the amplitude contribution of the negative and positive rotor depend on $\zeta_{+l}$, and on $\zeta_{-l}$ and $\zeta_{+l}$, respectively. They are given by $\zeta_{+l} + \zeta_{-l}$ for the negative rotor and by $\zeta_{+l} - \zeta_{+l}$ for the positive rotor.
Figure S10: LOCO-EFA is invariant to image rotation or mirroring, nor sensitive to image resolution. (A) The cell outline of an experimentally observed cell was mirrored along the y-axis and/or rotated over different angles, after which LOCO-EFA was applied to each image separately. (B) The \( L_n \) values and other derived metrics were invariant to those transformations. (C) The resolution of the original image was reduced, such that the number of contour points decreased from 1104 to 253, 108 and 27 (from left to right, respectively). (D) The \( L_n \) numbers and the associated metrics only deviated from the high-resolution values when the resolution was very low and the cell outline was clearly deviating from original cell outline. This in contrast to the skeletonisation method, for which even marginal resolution reductions can cause large deviations in the outcome.
Supporting Table

Table S1: Three types of specified cell shapes (SCS) as used in Figure S6 and Figure S7.

|            | SCS1 | SCS2 | SCS3 |
|------------|------|------|------|
| Target Area| 858  | 1167 | 1197 |
| Pointedness| 6912 | 5328 | 5207 |
| Number of lobes | 3–10 | 3–10 | 3–10 |
| Roundness   | 382  | 518  | 434  |
| Elongation  | 4    | 28   | 5927 |

C code of LOCO-EFA method

Click here to Download C code of LOCO-EFA method

Outline example for code

Click here to Download Outline example for code
Supporting Movies

Movie 1: **Approximation of a closed contour using Elliptic Fourier Analysis.** A given two-dimensional shape can be approximated using EFA by summing $n$ elliptic harmonics as follows: each $n$th elliptic harmonic traces $n$ clockwise or counter-clockwise revolutions while moving around the previous elliptic harmonic.

Movie 2: **Direction of rotation opposite to the first harmonic ellipse.** EFA mode 3 generating a shape with 4 features, its rotation direction opposite to the rotation direction of the first harmonic.
Movie 3: **Direction of rotation same as the first elliptic harmonic.** EFA mode 3 generating a shape with 2 features, its rotation direction the same as the rotation direction of the first harmonic.

Movie 4: **Exception of the rule regarding rotation direction.** If the eccentricity of an elliptical harmonic is very high, the number of generated lobes does not follow the rule-of-thumb proposed by Diaz et al. (1990). Here, EFA mode 3 generates a shape with 4 features, although its rotation direction is the same as the rotation direction of the first harmonic.
Movie 5: Example of an in silico simulation of a population of cells with more complex specified shapes. CPM simulation of multi-lobed cells take up pavement-like cell shapes when they are allowed to interact with their neighbours within a confluent tissue.
Supporting Code

loco-efa_example.c Code written in C which applies the full procedure to calculate the $L_1$ and $L_4$ values to any file containing contour coordinates. For each computational step, it is indicated where the relevant mathematical details can be found in the Supplementary Materials and Methods. We have intentionally kept the code as bare as possible, without for example any graphical interface, to allow it to be trivially compiled and run on any platform. Details regarding compilation and execution can be found in the header of the file.

cell_outline.csv Contour data of the cell presented in Figure 3S, Figure S3A and Figure S10. This file, or any other file containing contour data, can be used as an input for the program.
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