insideOutside: an accessible algorithm for classifying interior and exterior points, with applications in embryology

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Abstract

A crucial aspect of embryology is relating the position of individual cells to the broader geometry of the embryo. A classic example of this is the first cell-fate decision of the mouse embryo, where interior cells become inner cell mass and exterior cells become trophoblast. Fluorescent labelling, imaging, and quantification of tissue-specific proteins have advanced our understanding of this dynamic process. However, instances arise where these markers are either not available, or not reliable, and we are left only with the cells’ spatial locations. Therefore, a simple, robust method for classifying interior and exterior cells of an embryo using spatial information is required. Here, we describe a simple mathematical framework and an unsupervised machine learning approach, termed insideOutside, for classifying interior and exterior points of a three-dimensional point-cloud, a common output from imaged cells within the early mouse embryo. We benchmark our method against other published methods to demonstrate that it yields greater accuracy in classification of nuclei from the pre-implantation mouse embryos and greater accuracy when challenged with local surface concavities. The code is freely available as a well-documented, easy-to-use MATLAB script. This method should prove useful within and beyond embryology, with broader applications to similar data from the biological and life sciences.
Introduction

The mouse embryo undergoes three major morphogenetic events between fertilization and implantation. These are termed compaction, cavitation, and hatching (Figure 1A,B) (Tarkowski and Wróblewska, 1967; Smith and McLaren, 1977; Yoshinaga et al., 1976). Compaction coincides with the first binary cell-fate decision, which is ultimately driven by cellular position within the embryo (Figure 1C) (Tarkowski and Wróblewska, 1967). Exterior cells polarize to become the extraembryonic trophoderm (TE), precursors of the placental (Lawson et al., 1999), while the interior cells become the inner cell mass (ICM) (Ziomek and Johnson, 1980; Johnson and Ziomek, 1981). The ICM cells undergo a second binary cell-fate decision to become either the embryonic epiblast, source of the foetus (Gardner and Rossant, 1979) and embryonic stem cells (Evans and Kaufman, 1981; Martin, 1981), or the primitive endoderm (PrE), founder of the yolk sac (Gardner and Johnson, 1972). This second cell-fate decision coincides with cavitation, where a fluid-filled cavity, called the blastocoel, forms between the TE and one side of the ICM (Smith and McLaren, 1977). Finally, prior to implantation, the embryo must hatch. This is the process where the embryo will break out of, and dissociate from, the zona pellucida (Malter and Cohen, 1989; Suzuki et al., 1995; Yoshinaga et al., 1976).

Molecular profiling of these tissues has been performed at the levels of RNA-sequencing (Boroviak et al., 2018; Guo et al., 2010, 2017; Kurimoto et al., 2006) and immunohistochemistry (Chazaud et al., 2006; Niwa et al., 2005; Palmieri et al., 1994; Plusa et al., 2008) to reveal key lineage-markers. These lineage-markers have been used to study the dynamic emergence and plasticity of distinct cell-identities throughout the stages of pre-implantation development by employing fluorescent reporter knock-ins (Arnold et al., 2011; Grabarek et al., 2012; Hamilton et al., 2003; Kalkan et al., 2017; McDole and Zheng, 2012; Plusa et al., 2008). However, there remain instances where reliable lineage-markers do not exist. For example, in unspecified cells of the compacting morula, where a single cell can co-express lineage-markers for TE (CDX2), PrE (GATA6), and epiblast (SOX2) (Plusa et al., 2008). There are also instances in genetic knock-outs (KOs) where lineage-markers no longer faithfully mark their lineage. In the example of the pluripotency factor OCT4 (Pou5f1), expression levels of early ICM lineage-markers KLF4 and TFCP2L1 are significantly lower in OCT4-KO mice than that of wild-type mice at the mid-blastocyst stage (Stirparo et al., 2021). Additionally, OCT4-KO embryos exhibit lower levels of expression in early PrE marker GATA6 and fail to initiate expression of late PrE marker SOX17 (Le Bin et al., 2014). A more severe case can be seen in the KO of the early PrE marker GATA6, where embryos not only failed to activate the expression of other PrE markers, PDGFRA, SOX17 and GATA4, but also expressed the ICM marker NANOG in the TE (Bessonard et al., 2014; Schrodé et al., 2014). Finally, moving beyond the mouse, definitive sets of lineage-markers have yet to be firmly established for other mammalian systems, such as human and non-human primates (Boroviak et al., 2018; Guo et al., 2021; Stirparo et al., 2018). In such instances, we must find alternative methods to classify the tissues under investigation.

Spatial information is a classical method for classifying cell populations in the pre-implantation mouse embryo (Fleming, 1987; Nichols and Gardner, 1984). Advances in both image acquisition
A. Bright-field images from pre-implantation mouse development, from morula to late blastocyst. B. Three major morphogenetic events occur during pre-implantation mouse development: compaction, cavitation, and hatching. C. Cells in the pre-implantation embryo make two sets of binary cell-fate decisions: first, blastomeres become inner cell mass (ICM) (interior) or trophectoderm (exterior); second, ICM become epiblast or primitive endoderm. These decisions coincide with compaction and cavitation, respectively, and are completed by hatching. D. A shape is convex if for any pair of points, $P_0$ and $P_1$, the resulting line segment is entirely contained within the shape; biological examples of convex shapes include the compacted morula and blastocyst. E. A shape is concave if there exists at least one pair of points whose resulting line segment passes to the exterior of the shape; biological examples of concave shapes include trophectoderm cells and the blastocoel cavity.

and image processing technologies have improved the accuracy of this spatial information. A common analysis method in mouse embryology involves quantitative immunofluorescence (qIF) of cell nuclei. This is usually the process in which three-dimensional (3D) confocal fluorescence microscopy images of nuclei are segmented and quantified using software such as Fiji [Schindelin et al. 2012], MINS [Lou et al. 2014], or Nessys [Blin et al. 2019]. Output parameters from qIF may include total nuclear fluorescence, nuclear volume, and the geometric center (centroid) of the nucleus. The point-cloud of centroids can then be used to classify individual nuclei by their relative position to all other nuclei. Classification of interior and exterior nuclei is of particular interest when investigating the relationship between the cells of the ICM (interior) and the TE (exterior).

To date, three methods have been used to classify interior and exterior nuclei of the mouse embryo from qIF. We refer to these as the random sample consensus (RANSAC) Ellipsoidal, Convex Hull, and insideOutside methods. The RANSAC Ellipsoidal method, employed by the nuclear segmentation software MINS [Lou et al. 2014], robustly fits an ellipsoid to the
point-cloud generated by segmented nuclear centroids through the RANSAC iterative method \cite{Fischler1981}. A nucleus is then classified as exterior if the distance from the ellipsoid’s centre to the nuclear centroid exceeds 0.95 times the distance from the ellipsoid’s centre to the point on the ellipsoid that is closest to the nucleus’s centroid; otherwise, the cell is classified as interior. The MINS software package has been widely used for qIF and has been cited in nearly 100 manuscripts. The Convex Hull method, employed by the spatial analysis software IVEN \cite{Forsyth2021}, constructs a convex hull, the smallest convex set that contains all centroids, from all nuclear centroids of the embryo and then classifies a nucleus as exterior if it belongs to the boundary of the convex hull. IVEN does allow for manual correction of the classification, however, this requires user input which may introduce bias. The insideOutside method, employed by \cite{Stirparo2021} and the focus of the present work, is an accessible position-based approach to the classification of interior and exterior nuclei. Here we use unsupervised classification over two parameters that relate a nuclear centroid to the boundary of the convex hull generated from all nuclear centroids of the embryo.

An additional fourth method, which we will refer to as the Naïve Ellipsoidal method, was introduced by Forsyth et al. \cite{Forsyth2021} for their benchmarking. This approach accepts the first fit of an ellipsoid to the nuclear centroid point-cloud, as opposed to generating a consensus ellipsoid through iterative random sampling, as in the RANSAC approach. The same distance rule as the RANSAC Ellipsoidal method is then applied to classify cell position.

These four methods share the common assumption that the embryo is convex. A shape is said to be convex if the line segment between any pair of points within the shape is entirely contained within the shape (Figure 1D); otherwise the shape is said to be concave (Figure 1E). The RANSAC Ellipsoidal method models the embryo as an ellipsoid, which is a convex shape. Similarly, the Convex Hull method explicitly defines the exterior points as being a member of a convex shape. Therefore, these methods underperform if the surface of the embryo exhibits small local concavities or if the embryo incurs indentations through fixation and mounting. The insideOutside method softens the assumption of convexity by classifying points using a two-dimensional parameter space instead of requiring strict membership of a convex shape.

We finish by noting that it has become increasingly important for experimentalists to perform rigorous quantification of the data they generate. Thus it is necessary to develop easy-to-deploy software that does not require high levels of programming expertise. Furthermore, benchmarking of the three mentioned classification methods has yet to be performed. Therefore, we present the accessible insideOutside algorithm for the classification of interior and exterior points of a relatively convex shape. We detail the mathematical framework underpinning the two-dimensional parameter space used for unsupervised classification, along with accuracy testing. We then proceed to benchmark current classification method using pre-implantation mouse blastocysts, showing that the Convex Hull and insideOutside methods outperform the Ellipsoidal methods. We conclude by demonstrating that the insideOutside method outperforms the Convex Hull method when challenged with local surface concavities, similar to what would be found in empirical data sets.
Results

The minimum distance and variance in distances from a point to the surface of a convex shape are inversely related

Here we will establish minimum distance to the surface and the variance in all distances to the surface of a convex shape as the parameters used for the insideOutside algorithm. An intuitive understanding for the selection of these parameters follows by considering what happens to these two parameters for two different points within a sphere: a point at the centre and a point at the surface.

![Diagram showing minimum distance and variance in distances for points at the origin and on the surface of a sphere.]

**Figure 2**: A. For a point $P$ (orange dot) coincident with the centre of a sphere ($O$, black dot) of radius $r$, the minimum distance from $P$ to the sphere is $r$ and the variance in distances is zero. B. For a point $P$ located on the sphere, the minimum distance to the sphere is zero, and the variance in distances is greatest. C. Analytic expressions for the minimum distance, $m$, and variance in distances, $v$, from a point $P$ located on/inside a sphere of radius $r$ centred at the origin to the sphere. D. This relationship

\[ v = r^2 + x^2 - \left( \frac{3r^2 + x^2}{3r} \right)^2 \]
is tested for spheres that are discretized using equidistant points. 100 equidistant points (blue dots) are plotted on the unit sphere (rainbow surface). \( m \) and \( v \) are calculated for 50 test points (red dots) along the vector from the origin (black dot) to the surface point \( P = [0, 0, 1] \). Shown are the three-quarters view (left) and the three orthogonal views (right). E. The inverse relationship between \( m \) and \( v \) are shown for the continuous case of the unit sphere (see (5)) (red line) and discrete cases of 100 equidistant points on the unit sphere (Figure 2D, black dotted line) and 100 uniform random points on the unit sphere (Figure S1, translucent black lines, 1000 realizations).

First, let us consider what happens to these parameters for the point at the exact centre of the sphere (Figure 2A). The minimum distance to the sphere is exactly the radius of the sphere. In fact, the distance from the centre to all other points of the sphere are identically the radius, meaning that the variance in distance to the surface of the sphere is exactly zero. Thus, minimum distance to the surface is maximized and the variance in distances is minimized for the point at the centre of a sphere.

On the other hand, consider an arbitrary point on the surface of that same sphere (Figure 2B). For that surface point, the minimum distance to the surface of the sphere is exactly zero. If we then draw line segments from that point to all other points on the surface of the sphere, we see that we are drawing line segments of every length between zero and the diameter of the sphere. Meaning that the original point on the surface of the sphere achieves the most diversity of line segment lengths possible for the sphere. In other words, a point on the surface of the sphere has the maximum variance in distances to the surface. Thus, minimum distance to the surface is minimized and the variance in distances to the surface has been maximized for any point on the surface of a sphere. We therefore arrive at an inverse relationship between the minimum distance and the variance in distances to the surface of a sphere as we move from the centre of the sphere to the surface of the sphere.

We now formalize this relationship. First, we derive the expression for the minimum distance from any point on/inside the sphere of radius \( r \), centred at the origin, to the sphere. Intuitively, a point \( P \) on/inside the sphere, its closest point on the sphere, and the origin all lie on a straight line (Figure 2C, top). Hence, if \( P \) is located a distance \( x \in [0, r] \) from the origin, then since the distance from any point on the sphere to the origin is \( r \), the minimum distance from \( P \) to the sphere is given by \( m = r - x \).

Next, we derive the expression for the variance in distances from a given point on/inside the sphere to the sphere. Let \( P \) be located at distance \( x \in [0, r] \) from the origin, and without loss of generality let \( P \) lie on the z-axis. Consider a point \( Q \) on the sphere, whose angle to the z-axis is given by \( \phi \) (Figure 2C, bottom). Since the distance from \( Q \) to the origin is \( r \), by the law of cosines the distance from \( P \) to \( Q \) is given by \( d(\phi) = \sqrt{r^2 + x^2 - 2rx \cos \phi} \). The variance in distances from \( P \) to the sphere is thus given by

\[
v = E[d^2] - E[d]^2. \tag{1}
\]
Computing surface integrals, we find that the mean distance $E[d]$ is given by

$$E[d] = \frac{1}{4\pi r^2} \int_0^\pi \int_0^{2\pi} d(\phi) \ r^2 \sin \phi \ d\theta \ d\phi = \frac{3r^2 + x^2}{3r},$$

(2)

while the second moment $E[d^2]$ is given by

$$E[d^2] = \frac{1}{4\pi r^2} \int_0^\pi \int_0^{2\pi} (d(\phi))^2 \ r^2 \sin \phi \ d\theta \ d\phi = r^2 + x^2,$$

(3)

hence the variance in distances is given by

$$v = r^2 + x^2 - \left( \frac{3r^2 + x^2}{3r} \right)^2.$$

(4)

We next show that the minimum distance and variance in distances to the sphere are inversely related. Substituting $x = r - m$ into our expression for $v$, after some algebra we obtain

$$v = -\frac{1}{9r^2} (m - r)^2 \left( m^2 - 2mr - 2r^2 \right),$$

(5)

from which we obtain

$$\frac{dv}{dm} = -\frac{2}{9r^2} (m - r)(2m^2 - 4mr - r^2).$$

(6)

Since $m - r \leq 0$ and $2m^2 - 4mr - r^2 < 0$ for $m \in [0, r]$, we have $dv/dm \leq 0$ for $m \in [0, r]$, hence $v$ is a decreasing function of $m$ for $m \in [0, r]$. Thus, for points on/inside the sphere, the variance in distances is inversely related to the minimum distance to the sphere. Finally this relationship can be seen by plotting (5) for the unit sphere ($r = 1$) (Figure 2E).

We conclude this section by showing that this inverse relationship holds for a convex surface generated by a set of discrete points. For this we simulate 100 points on the surface of a sphere that are spaced either equidistant (Figure 2D) or uniformly at random (1000 realizations) (Figure S1, Deserno 2004). For each discrete surface, the minimum distance and variance in distances to the surface points are calculated for 50 test points equally spaced between the centre of the sphere and a surface point. Both sets of simulations closely match the analytical solution (Figure 2E). While this has been demonstrated for the case of the sphere and discretized derivatives of the sphere, it provides a theoretical foundation for the use of $m$ and $v$ in the classification of convex point-clouds. Importantly, the demonstration using discretized surfaces indicates that this relation is directly applicable to empirical data which consist of discrete points, e.g. the centroids of nuclei within an embryo.
insideOutside: a two-dimensional decision space for classifying interior and exterior positions

Motivated by the theoretical result of the previous section, we proceed to describe an algorithm for the classification of interior and exterior points of a 3D point-cloud. The insideOutside algorithm (Algorithm 1) takes in a set of 3D Cartesian coordinates, \( S \in \mathbb{R}^{m \times 3} \) (Figure 3Ai), and returns indexing vector \( I \in \mathbb{B}^m \) with 0 indexing the inside points and 1 indexing the outside points. For pre-implantation embryos, the input data can be generated through manual nuclear segmentation in Fiji (Schindelin et al., 2012) or MATLAB’s volumeSegmenter App (Copyright 2020 The MathWorks, Inc) or through automated 3D nuclear segmentation pipelines like MINS (Lou et al., 2014) or Nessys (Blin et al., 2019). The algorithm begins by computing the Delaunay triangulation, \( \mathcal{D} \), over \( S \) (Figure 3Aii). From \( \mathcal{D} \), we generate a convex hull \( \mathcal{H} \) (Barber et al., 1996) (Figure 3Aiii). Now, using \( \mathcal{H} \) we can calculate the distance function, \( d(P, \mathcal{H}) \), \( \forall P \in S \) (Figure 3Aiv). We calculate the minimum and variance in \( d(P, \mathcal{H}) \), \( \forall P \in S \), and then scale each parameter such that the maximum is 1 and the minimum is 0 (Figure 3Av). Finally, hierarchical clustering by ward linkage is performed on the parameters to classify the points into two groups (Figure 3Avi, vii).

Algorithm 1: insideOutside takes in an \( n \times 3 \) matrix of Cartesian points and returns a bit vector that classifies each point as either inside, 0, or outside, 1.

Data: Set of points \( S \in \mathbb{R}^{m \times 3} \)  
Result: Classification vector \( I \in \mathbb{B}^m \)  
Delaunay triangulation \( \mathcal{D} \) over \( S \)  
Generate convex hull \( \mathcal{H} \) from \( \mathcal{D} \)  
for each point \( P \in S \) do  
  for each face \( f \in \mathcal{H} \) do  
    Calculate \( \min d(P, f) \)  
  end  
  Calculate \( m = \min d(P, \mathcal{H}) \)  
  Calculate \( v = \text{Var}(d(P, \mathcal{H})) \)  
end  
Perform unsupervised classification for two groups

The accuracy of the insideOutside method classification was performed on test shapes designed to resemble the late mouse blastocyst, whose cell number ranges between 100-150 (Plusa et al., 2008), where approximately 60-70% of the cells belong to the TE (outside cells) (Fleming, 1987; Saiz et al., 2016, 2020; Morgani et al., 2018). Therefore we constructed shapes with 100 uniform random points on the unit sphere (outside) and 50 uniform random points within balls of radii between 0.01 and 1 (inside), both centered at the origin (Figure 3B and Figure S2). 1000 shapes were simulated and classified for each of 100 inner ball radii.

Initial tests revealed near-perfect classification rates for True Outside points at all inner ball radii (Figure 3C). There was, however, a significant drop in the True Inside classification rate at an inner ball radius of 0.82 where the True Inside classification rate dropped below 0.99 with minimum rate of 0.48 ± 0.17 (mean ± standard deviation) at an inner ball radius of 1. To
Figure 3: A. Outline of the `insideOutside` algorithm. B. Accuracy testing was performed on a shape.
constructed of 100 outside points, uniform random points on the unit sphere, and 50 inside points, uniform random points in a ball centered at $O$ of radii ranging from 0.01 to 1 (see Figure S2). Bi-iv. The steps of the algorithm performed on an example shape with inner ball radius of 1. Bi. Ground truth of inside points, blue dots enclosed by blue surface, and outside points, orange points on orange surface. Bii. The triangulated hull generated from making a convex hull over the Delaunay triangulation. Biii. The classification of points using hierarchical clustering over the calculated parameter space. Shown are True Inside points (blue), True Outside points (orange), and Misclassified Outside points (green). Biv. The classification mapped onto the original shape. C-D. Accuracy testing was performed by classifying the points of 1000 shapes for each of 100 different inner ball radii. The mean True Inside rate (blue dotted line) is shown with standard deviation (blue filled region) and the mean True Outside rate (orange solid line) is shown with standard deviation (orange filled region). C. Accuracy test for the parameters $m$ and $v$. D. Accuracy test for the parameters $\log_{10}(m + 0.01)$ and $v$.

To improve the True Inside classification rate, we modified the parameter space by taking the log of $m$, which results in a greater separation of inside and outside points along the minimum distance axis (Figure 3D). Simulations bear out marked improvements in True Inside classification rates with no detriment to True Outside classification rates. The resulting True Inside classification rates do not drop below 0.99 until an inner ball radius of 0.94 and achieve a minimum of only 0.82 ±0.09 at an inner ball radius of 1. Thus we have established our algorithm using the parameters of $[\log_{10}(m + 0.01), v]$ and we now proceed to challenge it with empirical data.

**insideOutside and Convex Hull methods outperform the Ellipsoidal methods when classifying cells of the mouse blastocyst**

In this section we set out to show that the insideOutside method can successfully classify the nuclei of real world embryos. In doing so, we also benchmark our method against the three other methods (Naïve Ellipsoidal, RANSAC Ellipsoidal (Lou et al., 2014), and Convex Hull (Forsyth et al., 2021)) using previously quantified mouse mid-blastocysts (Stirparo et al., 2021) (Figure 4A). SOX2 staining, which can be used to mark all nuclei of the early ICM (Wicklow et al., 2014), was used as the ground truth for benchmarking, where SOX2 positive nuclei indicate inside nuclei and SOX2 negative nuclei indicate outside nuclei. SOX2 positive/negative status was determined through statistical inference (Gaussian mixture modelling) with 758 cells from 14 embryos (Figure 4B,C). We then used the SOX2 ground truth (Figure 4D) to calculate the True Inside and True Outside rates for the four classification methods (Figure 4E).

For inside nuclei classification (Figure 4F, left), we find that both the Convex Hull (rate = 0.92 ± 0.06, mean ± standard deviation) and insideOutside (rate = 0.91 ± 0.06) methods outperform the Naïve Ellipsoidal method (rate = 0.59 ± 0.29; p-values of $6.8 \times 10^{-4}$ and $1.0 \times 10^{-3}$, respectively, Kruskal-Wallis) and RANSAC Ellipsoidal method (rate = $0.62 \pm 0.12$; p-values of $8.1 \times 10^{-5}$ and $1.3 \times 10^{-4}$, respectively). The Naïve and RANSAC Ellipsoidal methods show no difference in ability to classify inside nuclei (p-value = 0.96). Similarly, the Convex Hull and insideOutside methods show no difference in ability to classify inside nuclei (p-value = 0.99, Kruskal-Wallis).

For outside nuclei classification (Figure 4F, right), we see that the Naïve Ellipsoidal method
(rate = 0.73 ± 0.13) underperforms compared to the RANSAC Ellipsoidal method (rate = 0.95 ± 0.06, p-value = 4.9 \times 10^{-4}) , Convex Hull method (rate = 0.89 ± 0.09, p-value = 0.031), and insideOutside method (rate = 0.90 ± 0.08, p-value = 0.019). However, no difference is seen between the RANSAC Ellipsoidal, Convex Hull, and insideOutside methods (all pair-wise p-values ≥ 0.63).

Thus, the Naïve Ellipsoidal method shows the lowest rates of classification for the both inside and outside points. While the RANSAC Ellipsoidal method only underperforms in classifying interior points. These data together show that ellipsoid fitting has the least satisfactory classification accuracy.

insideOutside has higher accuracy than the Convex Hull method when challenged with local surface concavities

While both the insideOutside and Convex Hull methods perform comparably on the empirical blastocysts, the Convex Hull method holds a systematic error of misclassifying outside points as inside points when small local surface concavities are introduced. We highlight this issue by emulating increasing levels of local surface concavities via the introduction of increasing levels of normally distributed random noise to the surface points of the test shapes (Figure 4G). We then compute the classification rates over the parameter space of inner ball radius (100 radii between 0.01 and 1) and noise factor (100 levels between 0 and 0.25) for 100 shapes (Figure 4H and Figure S3A). The Convex Hull method shows a uniform decrease of True Outside classification rates across all inner ball radii for increasing levels of noise, eventually dropping below a rate of 0.4 around a noise factor greater than 0.2. The insideOutside method does not display this uniform decrease of True Outside classification rates, instead maintaining a rate of greater than 0.9 for the majority of the parameter sets tested. The insideOutside method only begins to lose accuracy when both the inner ball radius and noise factor become large. Surface concavities have negligible effects on the classification rates of inside points for both methods (Figure S3B,C).

Discussion

Motivated by the need to accurately classify cells of mouse embryos by position alone, we present insideOutside, an accessible algorithm for the classification of interior and exterior points of a three-dimensional point-cloud. We established, both analytically and computationally, that for a convex shape, there exists an inverse relationship between the minimum distance to the shape’s surface and variance in distances to the shape’s surface. We then harnessed this inverse relationship to build an algorithm which allows for faithful classification of interior and exterior points by hierarchical clustering. We then proceeded to benchmark our method against three other methods, Naïve Ellipsoidal, RANSAC Ellipsoidal [Lou et al. 2014] and Convex Hull [Forsyth et al. 2021], showing that the insideOutside method was as reliable, or better, at classifying nuclei of the pre-implantation mouse embryo. We demonstrated that the insideOutside method has greater accuracy than the Convex Hull method in classifying exterior points when challenged...
Figure 4: **A.** Confocal images of a mid-blastocyst stained for DNA and early ICM marker SOX2. A single slice is shown for transmitted light and maximum intensity projections are shown for fluorescence images. **B.** Gaussian mixture modelling (GMM) was performed on the SOX2 nuclear signal of 758 cells from 14
embryos to classify SOX2 positive (blue) and negative (orange) nuclei. Nuclear signal was normalized by nuclear volume, log$_{10}$ transformation, and re-scaling to the interval [0,1]. C, GMM classification of cells applied to the embryo from A. D, SOX2 GMM classification of embryo from A shown in three-quarters view. SOX2 GMM classification was used as the ground truth for methods benchmarking. Shown are True Inside (blue) and True Outside (orange) cell classification. E, Classification of embryo from A by the Naïve Ellipsoidal, RANSAC Ellipsoidal [Lou et al., 2014], Convex Hull (Forsyth et al., 2021), and insideOutside [Stirparo et al., 2021] methods shown in three-quarter view. Shown are True Inside (blue), True Outside (orange), Misclassified Outside (green; inside classified as outside), and Misclassified Inside (gray; outside classified as inside) nuclei. F, Classification rates for True Inside (left, blue) and True Outside (right, orange). Kruskal-Wallis Test, p-values: *, 0.05 ≥ p > 0.01; **, 0.01 ≥ p > 0.001; ***, 0.001 ≥ p > 0.0001; ****, 0.0001 ≥ p. All other pairwise relationships were not significant with p-values ≥ 0.63. G, Increasing levels of noise were added to the surface points of the test shape to simulate increasing local surface concavities. H, The mean True Outside rate (orange scale) is shown over the parameter space of inner ball radius (100 radii between 0.01 and 1) versus noise factor (100 levels between 0 and 0.25) for the Convex Hull (left) and insideOutside (right) methods. 100 test shapes were classified for each parameter pair. Additional contour lines are shown to delineate drops in classification rate.

with local surface concavities. Finally, we have packaged the algorithm as a stand-alone MATLAB function that takes in a set of points as an $n \times 3$ matrix that returns two outputs: an $n \times 1$ bit vector indexing inside, 0, and outside, 1, points and an $n \times 2$ matrix of the calculated parameter values of minimum distance and variance in distances to the surface generated from the input set of points. This is freely available at https://github.com/stanleystrawbridge/insideOutside.

We have shown here that the Convex Hull and insideOutside methods both outperform the Naïve and RANSAC Ellipsoidal methods in the classification of interior nuclei of pre-implantation mouse embryos. While, the Naïve Ellipsoidal method underperforms the other three methods when classifying exterior nuclei. In all four methods we see that instances of misclassification are highest where the ICM is in contact with the TE (Figure 4E). This show that ellipsoid fitting methods (Naïve and RANSAC) are not as suitable in classifying interior and exterior points as convex hull based methods (Convex Hull and insideOutside).

Moreover, we have shown, through simulation, that the insideOutside method has greater accuracy when challenged with surface concavities. This is of particular importance for classifying model systems whose exterior points exhibit high levels of noise. This noise can either be biological, an example being columnar epithelium whose nuclei exhibit differential apicobasal polarity. Or the noise can be technical, such as that resulting from inherent segmentation inaccuracies, which will be true of any method used.

These findings speak to the appropriateness of each method. There may be instances when the user has a large number of points in a low noise situation where exterior points should be strictly classified as belonging to the surface. In such a case the Convex Hull method is most appropriate. Alternatively, the user may want to soften this condition in the case of a small number of points in a high noise situation, e.g. the pre-implantation mouse embryo. Here, the insideOutside method would be most useful, as it performs the best in such high noise situations. While the Ellipsoidal method has proved useful in identifying unique embryos from
images with many embryos ([Lou et al., 2014], we would not recommend the Ellipsoidal method for classifying the nuclei of those embryos. Finally, we note that the insideOutside method is slightly slower than the Ellipsoidal and Convex Hull methods by virtue of it performing more calculations, i.e. a step in the insideOutside algorithm is the generation of a convex hull. However, this does not affect practical application and only becomes apparent when classifying numbers of shapes on the order of \( > 10^6 \), as presented here in simulations.

There is scope for the refinement of the insideOutside algorithm. This could come by way of incorporating more information about the segmented nuclei, e.g. making use of nuclear aspect ratio and not just nuclear centroid. Additional parameters could also be introduced to the parameter decision space. IVEN has made use of number-of-neighbors, calculated from the Delaunay triangulation, in downstream spatial analysis. The addition of number-of-neighbors to the classification space may aid in better discrimination of interior and exterior points, especially in the problem case where the ICM meets the TE. Also the method of unsupervised clustering could be further explored. Here we present the insideOutside algorithm using hierarchical clustering with ward linkage. Initially, k-means clustering was attempted but proved to have lower levels of accuracy due to the irregular cluster shapes in the decision parameter space. Many other unsupervised clustering methods exist (DBSCAN, spectral clustering, Gaussian mixture modelling, etc.) which could also be deployed.

We find it of particular importance to make this method, and future methods, easy-to-deploy for biological and life scientists, as there is increasing need for them to perform rigorous quantification of their data. Development of the insideOutside method in Stirparo et al. (2021) was born of a need to refine the original MINS classification method and was driven by collaboration between experimentalists and theoreticians. While there is no expectation for experimentalists to do methods development, there is expectation that they should be able to use these methods, thus empowering future work. Both MINS and IVEN share in this ethos of empowering experimentalists in the journey of data analysis. However, the insideOutside method stands apart from its counterpart methods in that it is a stand alone function, whereas the classification methods in MINS and IVEN are members of a larger software package. This means the insideOutside method has greater flexibility in use and migration to other programming languages. For example, implementations in open source languages like Python and R, which are widely used in the biological and life sciences, will make the insideOutside method more accessible as it would remove the dependency on a MATLAB licence. Critically, the stand-alone nature of the insideOutside method lends itself to incorporation into other software pipelines. Indeed, the insideOutside method could be incorporated as an additional classification method into either MINS or IVEN, as both packages have MATLAB implementations.

Finally, other use cases for the insideOutside method include other mammalian organisms that undergo the process of blastocyst formation (humans, non-human primates, other rodents, ungulates, etc.). It also has use for certain organoid systems, such as quantifying the level of cell sorting in ICM organoids ([Mathew et al., 2019]). And while the insideOutside method was motivated by the need to discriminate between the ICM and the TE in the pre-implantation
blastocyst, it remains a general method for classifying the interior and exterior points of a point-cloud. This means it has extensibility to any data of this description. This includes, but is not limited to, the organization of transcription factor clusters from single-molecule localization microscopy [Liu et al., 2014], the pattern of RNA transcripts acquired through seq-FISH [Lohoff et al., 2021], and the relationship of genomic loci within the nucleus as determined by single-cell Hi-C structures [Stevens et al., 2017].

Materials and Methods

Embryo collection and bright-field imaging

Embryos were obtained from natural mating, detection of a copulation plug in the morning was used as confirmation of successful mating and indicated embryonic day (E) 0.5. 8-cell and compacted morula embryos were flushed from the oviduct at E2.5 and E3.0, respectively, and mid and late blastocysts were flushed from the uterine horns at E3.5 and E4.5, respectively, using M2 medium (Sigma-Aldrich, M7167). This research has been regulated under the Animals (Scientific Procedures) Act 1986 Amendment Regulations 2012 following ethical review by the University of Cambridge Animal Welfare and Ethical Review Body. Use of animals in this project was approved by the ethical review committee for the University of Cambridge, and relevant Home Office licences (Project licence No. 80/2597 and No. P76777883) are in place. Bright-field images were taken on a Leica DMI4000B microscope.

Quantitative immunofluorescence of embryos

Quantitative immunofluorescence data was originally published in Stirparo et al. (2021). In brief, embryos were fixed in paraformaldehyde, stained for DNA and SOX2, and imaged using confocal microscopy. Embryo nuclei were segmented and quantified using MINS [Lou et al., 2014] to calculate the nuclear parameters of total fluorescence (sum of pixel values), volume, and centroid.

Accuracy testing designations and rates

For accuracy testing, a point is assigned one of four designations: true inside, misclassified inside, true outside, or misclassified outside. A designation of "true inside" means the ground truth of the point and the classification of the point were both "inside". A designation of "misclassified inside" means the ground truth of the point was "inside" and the classification of the point was "outside". The same logic applies to the "outside" designations.

Two rates are calculated in the accuracy testing: true inside rate and true outside rate. The "true inside rate" is calculated as the number "true inside" points divided by the number of "total inside" points determined by the ground truth,

\[
\text{True Inside Rate} = \frac{\text{True Inside}}{\text{Total Inside}}.
\]
The same logic applies to the "true outside rate".

**Code availability**

The **insideOut** algorithm and all code used in this manuscript to perform simulations, analysis, and benchmarking are written in MATLAB (2021a) and are freely available at [https://github.com/stanleystrawbridge/insideOutside](https://github.com/stanleystrawbridge/insideOutside) under the GNU General Public License v3.0.

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**Author contributions**

SES conceived and developed the **insideOutside** algorithm. ECS collected bright field images of pre-implantation mouse embryos. SES and AGF performed mathematical analyses. AK performed embryo immunostainings and segmentation used for methods benchmarking. SES performed simulations and computational analyses. SES, AGF, and JN wrote the manuscript.

**References**


Supplemental material

Figure S1: Example of 100 random points (blue dots) drawn uniformly from the unit sphere (rainbow surface). The minimum distance to the surface, $m$, and variance in distances to the surface, $v$, are calculated for 50 test points (red dots) along the vector from the origin (black dot) to a random point on the surface. Shown are the three-quarters view (left) and three orthogonal views (right). Figure 2E.
Figure S2: Example shapes with different inner ball radii used for accuracy testing. **A.** Ground truth of inside points (blue dots enclosed by blue surface) and outside points (orange points on orange surface). **B.** The convex hull generated the Delaunay triangulation. **C.** Classification of points using hierarchical clustering over the calculated parameter space. Shown are True Inside points (blue), True Outside points (orange), and Misclassified Outside points (green). **D.** The classification mapped onto the original shape.
Figure S3: Classification rates shown over the parameter space of inner ball radius (100 radii be-
between 0.01 and 1) versus noise factor (100 levels between 0 and 0.25) for the Convex Hull (left) and insideOutside (right) methods. **A.** The standard deviation of the True Outside rate. **B.** The mean True Outside rate. Additional contour lines are shown to delineate drops in classification rate. insideOutside rate is > 0.9 everywhere except when the inner ball radius is close to 1 and the noise factor is small. **C.** The standard deviation of the True Inside rate.