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Why does epithelia display heterogeneity? Bridging physical and biological concepts

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Abstract

Technological and computational advances in the past few decades have allowed biophysicists to describe behaviour of epithelia, using mesoscale physical principles. In such description, similar to a glassy solid or dense particulate matter, epithelial cells are shown to transit from fluid-like motion to a jammed and kinetically arrested state upon crowding. This jamming transition is characterized by dynamic heterogeneity, which is revealed by fluctuations in intercellular stresses arising from multicellular cooperation. Even though recent studies are suggestive of the role of dynamic heterogeneity in tissue homeostasis and wound healing, very little is known about its physiological meaning. Meanwhile, recent studies in epithelial cell biology reveal an intrinsic cellular heterogeneity arising from variations in genome and protein expression patterns. Interestingly, such heterogeneity is also shown to be relevant in regulating tissue homeostasis. In the light of such observations, it becomes intuitive to ask how the inherent biological heterogeneity of epithelial tissues influences its physical behaviour. In this review, we attempt to bridge this gap by connecting studies describing dynamic heterogeneity in epithelial tissues with those that describe intrinsic biological heterogeneity and discuss how these might be linked in order to potentially regulate epithelial functionality.

Key words: heterogeneity, epithelial tissue, jamming, collective behaviour, extra-cellular matrix

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Epithelial cells construct inner and outer linings of our organs and function as physical barriers, thus, protecting the underlying tissue from infections, dehydration, and also aiding in efficient absorption of nutrients and gases (Alberts 2008). Cells within the epithelia perform these tasks, being jammed at their place while also making sure that epithelial homeostasis is maintained, failing in which can be potentially fatal for the tissue (Macara et al. 2014). Interestingly, the same cells can unjam and flow almost like a fluid during physiological and pathological situations such as organ development, wound healing and cancer metastasis (Friedl and Gilmour 2009; Mongera et al. 2018; Park et al. 2016; Sadati et al. 2013; Scarpa and Mayor 2016). In such situations, cells, rather than moving individually, migrate as a group in various patterns (Haeger et al. 2015; Petitjean et al. 2010; Poujade et al. 2007; Rorth 2012; Tarle et al. 2015). Reductionist view holds that such cooperative cellular events are mediated at the level of cell-cell interactions where local signals are translated into physical forces (such as those generated in the cellular cytoskeleton and those exerted across cell-cell junctions), which are then translated into cell motility (Das et al. 2015; Keller 2012; Ladoux and Mège 2017; Trepap et al. 2009). Such physical forces are believed to be fundamental to biological form and function but have remained hidden until recently when experimental methods are finally making them visible (Angelini et al. 2010; Angelini et al. 2011; Edwards and Schwarz 2011; Malinverno et al. 2017; Sabass et al. 2008; Schwarz and Soine 2015; Sunyer et al. 2016; Tambe et al. 2011; Trepap and Fredberg 2011). Furthermore, recent advances in mathematical biology have also led to the development of models that can predict various parameters of epithelial behaviour in both jammed and unjammed states (Edwards and Schwarz 2011; Garcia et al. 2015; Henkes et al. 2011; Mark et al. 2010; Mehes and Vicsek 2014; Sepulveda et al. 2013; Steinberg 2007). Together, these studies have revealed unpredicted behaviour of epithelial tissues and are beginning to explain why cells jam and unjam, and how collective cell behaviour is orchestrated. Since excellent reviews have been written on the topic (Friedl and Gilmour 2009; Haeger et al. 2015; Merkel and Manning 2017; Park et al. 2016; Park and Fredberg 2016; Pegoraro et al. 2016; Sadati et al. 2013), we will only briefly describe the heterogeneous nature of the jamming transition from the physical perspective and then discuss its implications in regulating epithelial functionality while also taking into account the inherent biological heterogeneity present within the epithelium.

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Jamming transition and dynamic heterogeneity. Ongoing cell divisions, apoptosis and cell mingling make the epithelia a highly dynamic place (Al-Hussaini et al. 2016; Christ et al. 1990; Gardner 1986; Macara et al. 2014). Interestingly, monolayer stress profiles of such epithelial layers reveal dynamic heterogeneity, with intercellular stress displaying stochasticity in space and time, meaning that stress is tied neither to any particular position nor to any particular cell within the monolayer (Angelini et al. 2010; Angelini et al. 2011; Garrahan 2011; Tambe et al. 2011). Topography of these intercellular forces, at any given instant, can be compared with a rugged landscape, similar to that of a mountain range, where peaks arise from cooperation between tens of cells pulling together (Tambe et al. 2011) (Fig. 1a). Interestingly, cell density also plays a key role in regulating dynamic heterogeneity i.e. when cells start to crowd, their movement becomes arrested and zones of cooperativity grow bigger (Angelini et al. 2011). Such a scenario is intriguingly analogous to glass transition within a supercooled fluid or dense particulate matter in which a non-equilibrium jammed state is reached by cooling, crowding or by decreasing applied load (Debenedetti and Stillinger 2001; Mattsson et al. 2009; Mayer et al. 2008; Nagel 1998; Trappe et al. 2001). Hallmarks similar to glass transition (spontaneous intermittent fluctuations, dynamic heterogeneity, cooperativity, and kinetic arrest) are observed by epithelial cell monolayer, wherein, the dynamical arrest is caused

1 upon crowding and depends upon parameters such as active motility, cellular forces, cell shape and applied stress.
2 When these parameters are comprehended in a jamming phase diagram (Nagel 1998; Sadati et al. 2013; Trappe
3 et al. 2001), predictions on epithelial physical behaviour can be made. For instance, as intercellular adhesion or
4 crowding progressively increases, cell motility and rearrangement would become rare and therefore, cooperativity
5 would increase, leading to a topologically frozen epithelium (Sadati et al. 2013). Subsequently, then, the question
6 is what the extension of jamming at homeostasis should be that allows the epithelia to achieve their vital
7 physiological functions such as regulating homeostasis and orchestrating collective cell migration.
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11 **Physiological relevance of heterogeneity.** The ability of epithelial cells to dynamically remodel their
12 surroundings as well as their own cytoskeleton in response to external cues such as damage or mechanical stresses
13 is known to provide a mechanical resilience to epithelial tissues (Khalilgharibi et al. 2019; Trepap and Sahai 2018).
14 Recent studies are suggestive of the hypothesis that, by maintaining a striking balance between jammed and
15 unjammed phases, the epithelial monolayer might have evolved to attain such resilience, by virtue of which, it
16 can efficiently undergo switch-like changes required for physiological functions (Park et al. 2015; Sadati et al.
17 2014; Saw et al. 2017; Vishwakarma et al. 2018). For instance, a recent study demonstrates that cooperative forces
18 owing to dynamic heterogeneity control the selection as well as frequency of leader cells which guide collective
19 migration during wound healing (Vishwakarma et al. 2018). Another study demonstrates that hot spots of
20 compressive stresses within the epithelial monolayer induce topological defects that subsequently lead to local
21 cell extrusion (Saw et al. 2017). Since hot spots of compressive stresses build up regions of multicellular
22 cooperation (Tambe et al. 2011) which show density dependence (Angelini et al. 2011), efficient cell extrusion
23 for regulating tissue homeostasis would intuitively require the right extent of cell packing. Such extrusion events
24 are important, not only due to their relevance in regulating cell density during epithelial homeostasis (Fadul and
25 Rosenblatt 2018; Gudipaty et al. 2018) but also in removing aberrant or tumour cells via a mechanism described
26 as cell competition, by virtue of which, epithelia gains the ability to defend itself against cancer (Kajita and Fujita
27 2015; Wagstaff et al. 2013). Understanding physiological relevance and extent of jamming in epithelia becomes
28 even more important in tissues that are naturally subjected to elevated levels of stress, such as lung epithelium
29 which goes through cyclic breathing stress and, therefore, tissue plasticity plays an important role in maintaining
30 its integrity, especially during lung injury (Frank and Matthay 2003).
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43 The physical heterogeneity described above is most likely to be influenced by the existing innate biological
44 heterogeneities in the epithelia which are associated with variations in genome or protein expression patterns
45 (Fig. 1b). A known outcome of this genetic variability is somatic mosaicism that leads to the presence of multiple
46 cell clones within an adult tissue. Somatic mosaicism can originate from epigenetics events (Rakyan et al. 2002;
47 Sutherland et al. 2000) such as, for instance, the inactivation of one of the X-chromosomes in females (Rakyan et
48 al. 2002) or from mobile DNA elements such as retrotransposons (Beck et al. 2011; De 2011). A classic example
49 of somatic mosaicism can be observed in the skin with the presence of mosaicisms in the pigmentation known as
50 *café-au-lait* spots (De 2011; Rawles 1947). In addition to these genetic differences, differential regulation of
51 proteins expression induced by external cues, such as extracellular matrix (ECM), can also create cellular
52 heterogeneity within the epithelial layer (Fig. 1b). The importance of such heterogeneity in regulating tissue
53 homeostasis has been shown in the basal layer of esophageal epithelium containing stem cells responsible for
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1 tissue renewal (DeWard et al. 2014). It has been shown that, in this layer, population of stem cells has
2 heterogenous proliferation rates which are distinguishable by the expression of specific cell-surface markers such
3 as the laminin receptor integrin $\alpha6\beta4$. Here, the involvement of laminins, major components of extra-cellular
4 matrix, suggests the importance of cell-ECM adhesion in maintaining cellular heterogeneity, and subsequently in
5 regulating tissue homeostasis (DeWard et al. 2014). Interestingly, cellular heterogeneity dictated by differential
6 laminin expression has also been shown to be involved in regulating functionality of endothelial cells. For
7 example, the extracellular matrix of endothelium in postcapillary venules consists of areas of high and low
8 expression of the laminin 511 isoform as compared to the capillaries where the expression of laminin 511 is
9 homogeneous (Di Russo et al. 2017; Sixt et al. 2001). Such differential distribution of laminin controls endothelial
10 cell junction tightness, thereby dictating the location of leucocytes extravasation through the blood-brain barrier
11 which occurs only in low laminin 511 regions (Sixt et al. 2001; Song et al. 2017). In addition to the biochemical
12 composition of ECM, its topography has also been shown to control the heterogeneity of epithelial cells. Recently,
13 an elegant experimental setting using undulated elastomer surfaces revealed the effect of ECM topography on
14 heterogeneity of keratinocytes (Mobasserri et al. 2019). After seeding primary keratinocyte on the surfaces, the
15 monolayer assembled within a range of cellular stiffness, cell-cell adhesion forces and acto-myosin contractility
16 levels. The results provided new insights into the possible heterogenous control of keratinocytes proliferation
17 rates by the topography of the dermal ECM during ageing and inflammation (Mobasserri et al. 2019). Differential
18 ECM expression also impacts on the aetiology of retinal degenerative disease, i.e. age-related macular
19 degeneration. The early stage of the disease is characterized by high level of ECM accumulation known as *drusen*
20 that occurs between the retinal pigment epithelium and the underlying Bruch's membrane (Coleman et al. 2008).
21 Drusen formation is a common age effect, but only the accumulation of high number of large drusen ($> 63\mu\text{m}$ in
22 diameter) correlates with epithelium degeneration and photoreceptor detachment (Coleman et al. 2008). Since the
23 retina pigment epithelium presents a very high heterogeneity in cell shape (Fig. 2), protein synthesis and granules
24 accumulation, it is tempting to speculate that this diversity of cell shape might also correspond to high
25 heterogeneity in monolayer tensions and, therefore, might control drusen formation and their growth.
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39 **Conclusion.** Even though physical and biological heterogeneities are currently known to be distinct, they are
40 likely to be interactive and interdependent. Local cellular heterogeneity might influence the mechanical properties
41 of epithelia, its ability to transduce forces and, hence, the nature of physical heterogeneity. Recent technological
42 advancements in biophysics, cell biology and mathematical biology has now made it possible to analyze the
43 physics and biology of the epithelia within the same framework. Such approach allows us to attain a more
44 comprehensive understanding on epithelial physiology and would subsequently require devising new treatment
45 strategies for epithelia degenerative diseases
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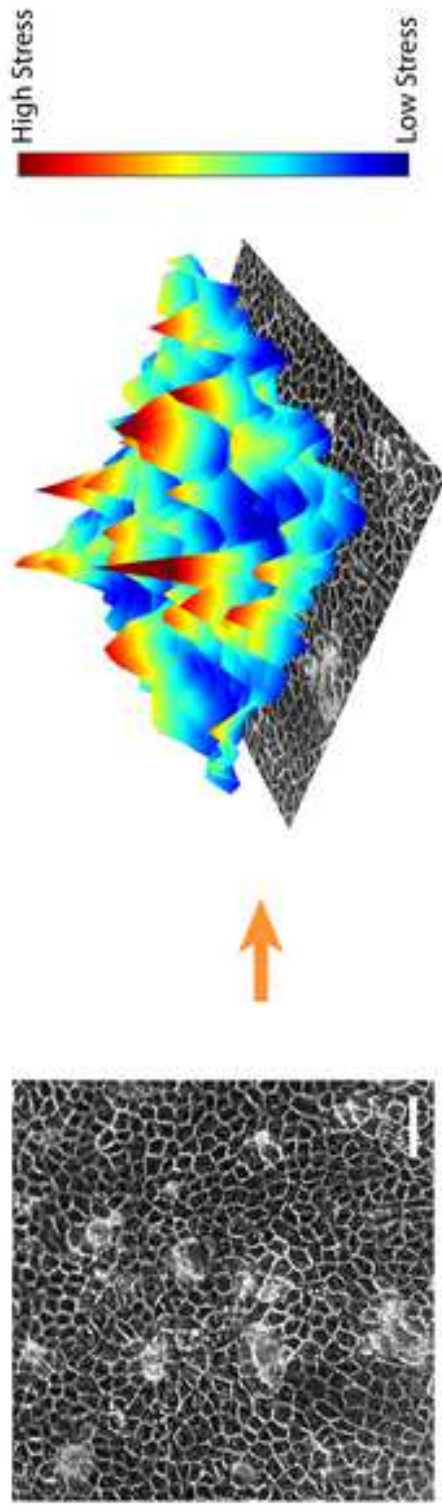
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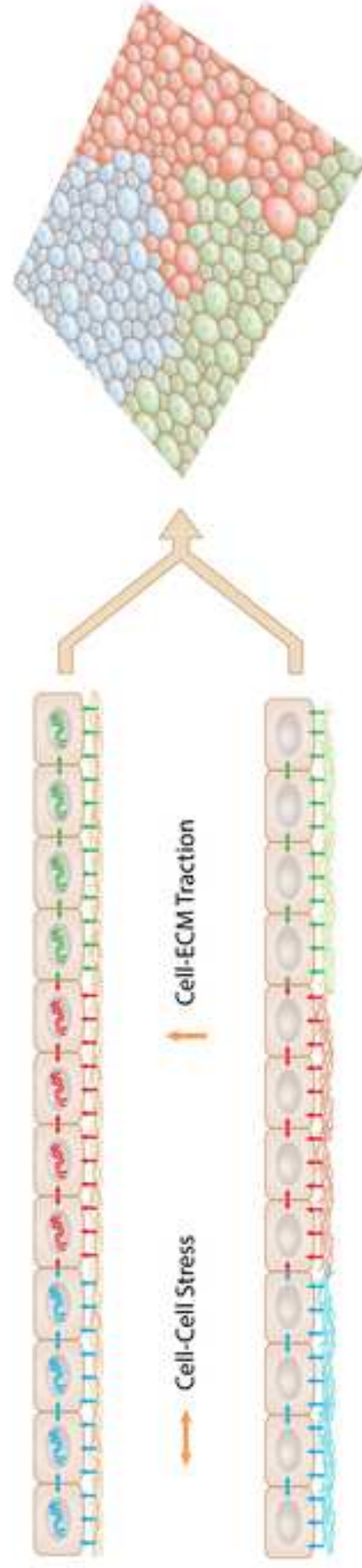
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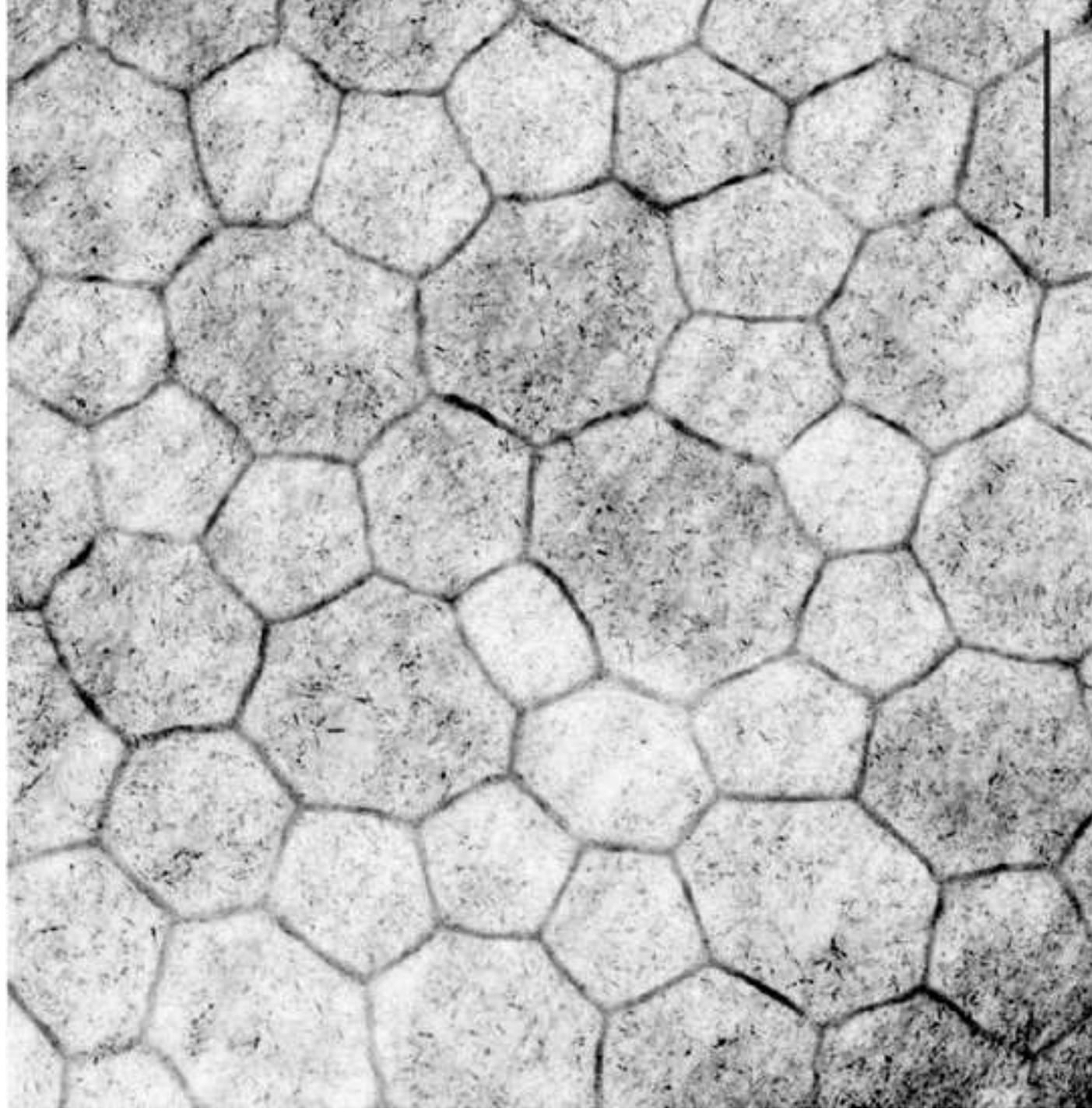
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1 **Figure legends.**

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4 **Figure 1:** a) The intercellular stress profile in a confluent epithelial monolayer of canine kidney epithelial cells
5 (MDCK) reveal a rugged stress profile at a given time point. Scale bar is 50µm. b) Cellular heterogeneities can
6 arise from genetic differences or differential regulation of protein expression which are also influenced by external
7 cues such as ECM components. In addition, heterogenous clones in epithelia might differ in their mechanical
8 properties, having different levels of adhesion forces (cell-cell stresses and cell-ECM tractions), thus impacting
9 on physical nature of epithelia.
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14 **Figure 2:** Immunofluorescent staining of *en-face* preparation of murine retinal pigment epithelium for F-actin
15 reveals highly heterogenous character of this epithelium. To be noted, is the postmitotic nature of these epithelial
16 cells, that excludes correlation of cell size with the cell cycle. Scale bar is 20 µm.
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