Defects in epithelial tissue organisation – A question of life or death

April 13 2017

A. Monolayer epithelial tissue

B. Emergence of a topological defect

C. Cell extrusion from the tissue
Schematics showing the arrangement of cells in normal epithelial tissue (a), following the introduction of a topological defect (b) and the site of cell death and extrusion following the appearance of a topological defect in the epithelial layer (c). The blue gradient region represents a concentration of compressive force. Credit: Mechanobiology Institute, Singapore

Researchers from the Mechanobiology Institute, Singapore (MBI) at the National University of Singapore have discovered the primary mechanism driving the extrusion of dying cells from epithelial monolayers. This work was published in Nature on 13 April 2017.

**Misalignment of cells predicts cell death and removal in epithelial tissue**

The removal of cells from a tissue occurs regularly. Not only are damaged or dying cells removed, but the process of cell extrusion can prevent regions from becoming overcrowded. This is particularly important not only during developmental processes when tissues and organs are being formed, but also in diseases such as cancer, when tumors grow uncontrollably. Despite the importance of cell extrusion in development and aging, as well as the pathological importance in cancer progression, the cues that flag a cell for removal were poorly understood.

Now, by studying single-layers of epithelial cells grown in the lab, scientists from MBI and the Institut Jacques Monod, Centre national de la recherche scientifique (CNRS) and University Paris Diderot in collaboration with researchers from Oxford University and Institut Curie have found that the major factor driving cell death and removal relies on the physical arrangement of cells in the surrounding cell layer. In particular, the appearance of defects in the cellular patterns of epithelial
layers promotes cell death and elimination from the tissues.

There are several examples in nature where a molecule or cell type aligns in a defined manner. Bacteria colonies, fat molecules, and even internal components of the cell, are just a few examples. Another well-known example corresponds to liquid crystals, a state of matter between a solid and a liquid, which can consist of rod-shaped molecules. Under certain conditions, these molecules can align along a preferential orientation when altered by electric, magnetic fields or temperature. This phenomenon is particularly well-known since it is exploited in technologies such as liquid crystal displays. In this case the optical properties of liquid crystals are determined by their alignment. Shifts in their alignment determine what we see on the display. As in any crystal, a perfect arrangement does not exist in liquid crystals and defects emerge in their arrangement that strongly modify their physical properties. The starting point of this study was to show how the behaviour of the cell sheet is analogous to liquid crystals.

Here, PhD candidate Mr Thuan Beng Saw, together with Professor Chwee Teck Lim of MBI, and Professor Benoit Ladoux of Institut Jacques Monod (IJM, CNRS) and MBI, in collaboration with Dr Amin Doostmohammadi and Professor Julia M. Yeomans (Oxford), Professor Philippe Marcq (Curie Institute) and Assistant Professor Yusuke Toyama (MBI), found that like the liquid crystals in a phone or laptop monitor, epithelial cells were arranged parallel to each other with their 'long' sides all facing the same direction. Following this analogy, they also observed the emergence of topological defects', which caused the cells to realign so that they resembled a comet. In this case, cells at the head of the comet pattern had shifted so that they now aligned perpendicular to the cells that made up the tail. Some cells turned up to 90 degrees. In a liquid crystal display, such realignments of the molecules merely alter the optical properties of the material. However, in an epithelial sheet, such changes in the pattern can mean life or death
for the cells involved. Remarkably, it was after this cell realignment that cells near the head of the 'comet pattern' died and were removed from the surrounding tissue.

To further investigate the relationship between cell death and topological defects, they examined the forces being generated around these particular areas of cell misalignment. They found that compressive force concentrated at the head of the comet pattern. This force generated over an hour prior to cell extrusion, and was sufficient to trigger cell death at topological defects. As cells are connected to each other by protein cables and adhesions structures, any movement of a cell causes tension to be propagated to its neighbours. The misalignment of cells causes significant bending of cells and this leads to high compressive stresses in these regions. These stresses are sufficient to trigger apoptosis and cell extrusion of a nearby cell.

Tissue engineering and regenerative medicine requires scientists to carefully control cell growth and tissue development in a lab. The findings presented in this work are of fundamental importance towards achieving this control. Indeed, the researchers successfully controlled how cells aligned by introducing shapes in areas where the cells grew that mimicked the topological defects associated with cell death and extrusion. This allowed them to pinpoint where in the cell sheet extrusion would occur. These discoveries provide a significant step forward in our understanding of how the physical microenvironment plays a role in tissue development, and provides new approaches with which researchers can control, analyse and study cell growth and death.
