Abstract

In recent years, the mammary gland epithelium has been shown to be a mixture of differentiated cell populations in a hierarchical relationship with their stem and progenitor cells. However, the mechanisms that regulate their cellular differentiation processes are still unclear. The identification of genes that govern stem and progenitor cell expansion, or that determine daughter cell fate, will be of crucial interest for understanding breast cancer diversity and, ultimately, improving treatment. Two recent analyses have identified some of the key genes that regulate these processes, lighting up the highway to normal mammary gland development.

In the mammary epithelium, recent successes in identifying stem and progenitor cells and lineage-specific transcription factors have raised the possibility of a much deeper understanding of what controls lineage commitment and cell fate. We have some way to go before we can approach the current level of understanding of haematopoietic lineage commitment. Nevertheless, every new report adds another piece to the puzzle and two recent publications have done just that.

Pietersen and co-workers have described the expression of Bmi1 in mammary cells and its crucial role in mouse mammary gland biology [1]. Bmi1 is a member of the polycomb group of proteins and a regulator of stem/progenitor cell function in other systems [2,3]. However, Pietersen and co-workers found that Bmi1 was expressed in all mammary epithelial cells and reported the profound inhibition of mammary gland development in a Bmi1 knockout mouse model, as well as a reduction of the mammary stem cell activity and premature lobuloalveolar differentiation. Interestingly, the co-deletion of Ink4a/Arf (repressors of the cell cycle down-regulated by Bmi1) partially rescued normal mammary gland development, whereas pregnancy did so completely. Thus, Bmi1 regulates stem and progenitor cell proliferation not only through the inhibition of Ink4a/Arf, but also through the inhibition of other genes related to differentiation processes that are significantly up-regulated by pregnancy.

While Bmi1 is a gene ubiquitously expressed in mammary cells, other genes are differentially expressed between cell populations and are important for lineage commitment and cell-fate. Raouf and co-workers [4] have recently compared the gene expression profiles of four human mammary cell populations isolated using cell-surface marker expression. Genes from the NOTCH signalling pathway, known to regulate cell fate decisions in other systems [5-7], were described as differentially expressed between two cell populations with different colony forming capacities: undifferentiated or bipotent colony forming cells (CFCs) and luminal-restricted CFCs. NOTCH4 was highly expressed in bipotent CFCs and markedly down-regulated in luminal-restricted CFCs, whereas an opposite pattern was found for NOTCH3 and HES6, as well as for NOTCH1 and NOTCH2 (although the changes were less dramatic for the latter two genes). Functional studies in which NOTCH3 signalling was blocked in bipotent CFCs showed a substantial decrease in formation of luminal-type colonies and an increase in myoepithelial-type colonies, leading the authors to propose NOTCH3 as one of the key genes for the luminal cell commitment. Although it was not directly demonstrated, it seems likely that the bipotent CFCs correspond to stem and/or multilineage progenitor cells whereas the luminal CFCs correspond to a lineage-restricted progenitor population.

These studies have highlighted two classes of genes driving mammary epithelial cell differentiation: genes accelerating or putting brakes on the development and differentiation processes (‘traffic lights’); and genes defining directions in these processes (‘road signs’) (Figure 1). Bmi1 is a promoter of stem and progenitor cell expansion but blocks differentiation, preventing its premature initiation [1]. Interestingly, both studies agreed in describing Bmi1 as commonly expressed in all mammary cells [1,4]. In addition, pregnancy favours both cell expansion and differentiation, co-operating with Bmi1 in regulating stem cell kinetics but opposing its...
effects in differentiation [1]. On the other hand, the NOTCH signalling pathway determines mammary cell fate, as in other tissues [6,7]. In previous studies on human mammospheres, NOTCH4 was shown to be essential for maintaining stem/progenitor cell activity and to direct progenitor cells to myoepithelial cell commitment [8], indicating a dual role for NOTCH4 as a ‘traffic light’ gene in stem cells but a ‘road sign’ gene in progenitor cells. Raouf and colleagues have confirmed the importance of NOTCH signalling pathways in mammary gland biology, showing that the luminal cell lineage is defined by down-regulation of NOTCH4 and the increase of NOTCH3. This NOTCH signalling role in cell-fate determination is also supported by conditional knockout mouse models of Notch effectors [9]. Although an interaction between Notch and Bmi1 has not been shown in these studies, a complex interacting model may exist with the involvement of Hedgehog and Wnt signalling pathways for the self-renewal and differentiation in mammary stem cells, as previously proposed [10].

Given that both traffic lights and road signs must be phased correctly for smooth fate determination, it is important to be able to integrate studies that address stem/progenitor cell kinetics and lineage commitment in both human and rodent systems. However, the different cell-sorting methods currently in use to isolate mammary cell populations, and the differing terminology used to describe them, confound our ability to create coherent models. Once the cell markers we use have been fully validated, the mammary stem cell community should be encouraged to develop common isolation protocols and nomenclature to facilitate the integration of data.

The mammary stem cell field has changed almost beyond recognition in the last five years. We are now in a position to go further than simply identifying stem cells and to truly begin to understand the molecular control of cell fate in the stem-progenitor-differentiated cell hierarchy. This is not simply an academic exercise. We feel (as no doubt do many others working in the field) that the different gene expression profiles of different breast cancer subtypes reflect real underlying differences in the cells of origin of these tumours and in the differentiation potential of these cells [11-13]. Therefore, we feel that molecular pathology data can only be fully interpreted and understood by reference to normal mammary epithelial cell subtypes and the regulation of their differentiation. The more we understand lineage selection and fate determination in the normal mammary epithelium, the closer we get to a comprehensive understanding of the biology of breast cancer subtypes.

**Competing interests**

The authors declare that they have no competing interests.
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