Immunological fortification at our barrier organs: Protecting us as we age

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The barriers of the human body represent our first line of defence against the outside world. Our barrier organs therefore face the formidable immunological challenge of defending us against pathogenic insults and promoting a peaceful co-existence with the local microbiota that inhabit them. Most antigens faced by our immune system initially enter through breaches in the physical barriers of the skin, or across one of the mucosal surfaces of either the respiratory, gastrointestinal or urogenital tracts. Consequently, these tissues are ‘immunologically fortified’ with common and distinct features, and their preservation is an essential component of host survival.

This is beautifully illustrated in the female reproductive tract (FRT). In their review of immune responses in the FRT, Monin et al.1 highlight its dual identities. The lower regions of the FRT including the vagina are characterized by a multi-layered stratified squamous epithelium, a mucus layer containing antimicrobial peptides, antibodies and immune mediators, and a substantial Lactobacillus-rich microbial community, forming a substantial physical and chemical barrier against invasive pathogens. This is complemented by a full arsenal of immune cells, including substantial populations of tissue-resident memory T-cells. The upper parts of the FRT, the uterus and ectocervix, meanwhile contain only a single layer of epithelium and appear devoid of commensals. Instead, the uterus possesses an immune system that is adapted to balance mucosal immunity against microbial exposure and immune tolerance, allowing and promoting the growth of a fetus. Indeed recent use of single-cell RNA sequencing (scRNA-seq) has revealed distinct populations of decidual natural killer (NK) cells2 which, as Male and colleagues discuss, unlike their cytotoxic contemporaries in the circulation are poor killers but produce pro-
angiogenic and trophoblast chemoattractant factors, key to fetal development, confirming the importance of barrier immunity from the moment of conception.

With this in mind, Bottling and Hanifa reveal how the most overt physical barrier, the skin, begins to develop in utero (REF – Bottling et al.). Harnessing recent advances in scRNA-seq with knowledge generated through traditional approaches such as histology allows the authors to take us on a molecular and cellular exploration of the human skin from the earliest points of life. They discuss how the immune response of the skin in utero is drastically different from that seen after birth, and rapidly changing during each trimester. The review shows that while fetal skin is hypo-responsive to inflammatory stimuli, it is primed to drive remodelling and repair; containing populations of type-2-like macrophages and dendritic cells, ILCs and regulatory T-cells (Tregs) that all promote wound healing without scarring in the 1st and 2nd trimesters.

The UK’s Office for National Statistics estimates that over the next 50 years an additional 8.2 million people will be living over the age of 65 years – that’s an increase in the size of current-day London. In their review, Chambers and Vukmanovic-Stejic take an elegantly simplistic approach discussing the individual components of the stromal, innate and adaptive effector cells compartmentalized within the three layers of the skin in the steady-state, and how these change as we age. Of interest to many of us in an ever-aging society, the authors describe specific age-related ‘defects’ specific to the skin barrier. For example, with increasing lifespan comes a host of complications resulting from chronic low-grade inflammation termed ‘inflamm-aging’. Alongside a loss of the structural integrity of the skin itself, a thinning of the epidermis and fragmentation of the extracellular matrix occurs, driven by increased matrix metalloproteinases and a reduced production of pro-collagen. In addition, aging results in a significant reduction in local Langerhans cells and functionality of the local antigen-specific T-cell population, resulting in a greater incidence of bacterial or viral infections and cancer.

The immunological defences of the human skin provide protection to a 1.5–2 m² area. In comparison, the respiratory tract covers an area of 70 m² that for the most part is made up of a single layer of epithelial cells for gas exchange. Maintaining efficient uptake of O₂ and removal of CO₂ means limiting the potential of immune cells to infiltrate this epithelium, even in response to infection. Invernizzi, Lloyd and Molyneaux focus their review on the ability of the respiratory epithelium itself to regulate immunity (REF – Invernizzi et al.). Invernizzi et al. also highlight how recent 16S sequencing data of the lower airways has changed scientific dogma: what was once believed to be a largely sterile microenvironment is now considered an ideal niche for specific species of commensal microbiota. They also discuss how disruption of microbial homeostasis at the respiratory epithelium drives the pathogenesis of a number of lung diseases ranging from asthma to idiopathic pulmonary fibrosis. The lungs also play host to unique resident immune cells, most notably alveolar macrophages that populate the luminal side of the alveoli and airways. So distinctive are alveolar macrophages that they cannot currently be generated in vitro, and Willinger and colleagues use their review to examine emerging evidence from model systems including the use of humanized mice engineered to express human macrophage colony-stimulating factor (M-CSF) and granulocyte-macrophage colony-stimulating factor (GM-CSF) to highlight additional developmental cues vital to their generation in vivo. Further to this, the authors also discuss a number of dynamic changes that occur to lung-specific macrophages occupying different niches throughout life, that start out as hypo-responsive sentinels in early life, but can become hyper-reactive as we age – potentially contributing to age-related inflammatory diseases.

From an immunological perspective, perhaps the most extensively researched, discussed and reviewed barrier site has been the gastro-intestinal (GI) tract. The GI tract is the barrier through which the majority of our nutrients flow, and is home to the canonical microbiome, containing over 10^{12} microbiota containing a wide range of parasitic, bacterial and viral pathogens (recently the focus of another review series10). In their review on integrin-mediated activation of transforming growth factor (TGFB), Travis and colleagues highlight the importance of the αv integrin family in providing contextual signals, facilitating these diverse activities. They highlight the distinct roles played by the integrin-TGFB pathway at discreet barriers, driving tolerance, and limiting T-cell responses. Perhaps of particular interest in the current setting of COIVD-19, the authors also discuss the implications of over-active TGFB in limiting antiviral immunity and promoting specific disease pathogenesis within the lung.

Finally, our largest internal organ, the liver, is discussed by authors Swadling and Stamatakis. The liver co-ordinates many physiological processes, including the filtration of blood, metabolism and storage of macronutrients, and detoxification. Blood flowing to the liver transits via the GI tract and is therefore rich in antigens. As such, the liver is tolerized to avoid immune responses against innocuous antigens (similar to the upper FRT), whilst maintaining the ability to elicit immune responses to blood-borne pathogenic insult. The authors discuss how emerging evidence from state-of-the-art single-cell technologies contributes to our understanding of this barrier (REF – Swadling et al.). Until recently, accurate mapping of the immune compartment in the liver was considered challenging as it required access to difficult/precious
samples and high-level experimental resolution. Describing the latest tools used to examine liver immunity in unprecedented detail, the diversity in the function and phenotype of resident immune cells, and heterogeneity of the parenchyma, is explained. Notably, the authors draw parallels between the murine and human liver, revealing how single-cell analyses have advanced or redefined our understanding of immune responses at this barrier.

As so starkly highlighted by the on-going global SARS-Cov2 pandemic, we rely on the dynamic immunological homeostasis that occurs at our immunological fortified barriers every day. This collection of reviews highlight the use of novel technologies (microbial sequencing, metabolomics and single-cell transcriptomics) and model systems to reveal the complex tapestry of cellular and molecular interactions occurring at each of our barriers in health and disease. With the advent of collaborative, open-source initiatives such as the Human Cell Atlas, the next decade promises to be an exciting era in our understanding of how local immune cell ‘experts’ contribute to barrier immunity in both health and disease.

References

1 Monin, L; Whetlock, EM; Male, V. Immune responses in the human female reproductive tract. Immunology 2020; 160:106–15.

2 Vento-Tormo, R; Efremova, M; Botting, RA; Turco, MY; Vento-Tormo, M; Meyer, KB et al. Single-cell reconstruction of the early maternal-fetal interface in humans. Nature 2018; 563:347–53.

3 Coolen, NA; Schouten, KC; Middelkoop, E; Ulrich, MM. Comparison between human fetal and adult skin. Arch Dermatol Res 2010; 302:47–55.

4 Popescu, DM; Botting, RA; Stephenson, I; Green, K; Webb, S; Jardine, L et al. Decoding human fetal liver haematopoiesis. Nature 2019; 574:365–71.

5 Chambers, ES; Vukmanovic-Stejic, M. Skin barrier immunity and ageing. Immunology 2020; 160:116–25.

6 Invernizzi R; Lloyd CM; Molyneaux PL. Respiratory microbiome and epithelial interactions shape immunity in the lungs. Immunology 2020; 160:171–82.

7 Molyneaux, PL; Cox, MJ; Willis-Owen, SA; Malla, P; Russell, KE; Russell, AM et al. The role of bacteria in the pathogenesis and progression of idiopathic pulmonary fibrosis. Am J Respir Crit Care Med 2014; 190:906–13.

8 Eren, E; Ringqvist, E; Willinger, T. Origin and ontogeny of lung macrophages: from mice to humans. Immunology 2020; 160:126–38.

9 Rongvuo, A; Willinger, T; Martinek, J; Strowig, T; Gearty, SV; Teichmann, LL et al. Development and function of human innate immune cells in a humanized mouse model. Nat Biotechnol 2014; 32:364–72.

10 Bain, CC; Cerovic, V. Interactions of the microbiota with the mucosal immune system. Clin Exp Immunol 2020; 199:9–11.

11 McEntee, CP; Gunathilaj, S; Travis, MA. Regulation of barrier immunity and homeostasis by integrin-mediated transforming growth factor beta activation. Immunology 2020; 160:139–48.

12 Kubes, P; Jenne, C. Immune responses in the liver. Annu Rev Immunol 2018; 36:247–77.

13 Reger, A; Teichmann, SA; Lander, ES; Amit, I; Benoist, C; Barney, E et al. The human cell atlas. eLife 2017; 6:e3–30.