Editorial: The Role of the EMT Program in Regulating the Immune Response in Carcinoma

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Editorial on the Research Topic
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The Epithelial-to-Mesenchymal Transition (EMT) is a cell-biological process that operates during embryonic development, wound-healing, and carcinoma progression. During this process, cells typically shed the expression of epithelial markers, such as E-Cadherin, and express instead, mesenchymal markers, such as vimentin, fibronectin and certain master EMT-inducing transcription factors (EMT-TFs), notably Zeb1, Twist, Snail and Slug. This is a highly dynamic process that often gives rise to a series of intermediate phenotypic states arrayed along the epithelial-mesenchymal polarization spectrum (1, 2). While the EMT program has long been known to enhance metastatic potential and resistance to chemotherapies and targeted therapies, its ability to regulate the immune response and promote resistance to anti-tumor immunity has only recently been uncovered (3, 4).

Immunotherapies in general, and checkpoint blockade inhibitors in particular, have led to durable responses and prolonged survival in patients whose tumors failed to respond to conventional therapies. However, while immunotherapies have had unprecedented success in some patients with certain cancers such as melanoma and non-small cell lung cancer, a number of other malignancies have shown little or no response. This has generated the need to identify novel therapeutic vulnerabilities, which can be targeted in combination with known drivers of resistance to enhance the efficacy of immunotherapies. The ability of the EMT program to regulate the immune response brings to the forefront the possibility of targeting this tumor cell-intrinsic process to enhance anti-tumor immunity and subsequent responses to many forms of therapy. In this particular Research Topic, we present a collection of reviews and original research articles that document multiple mechanisms by which the EMT program regulates the immune response.

A few studies have demonstrated the ability of the EMT program to promote immune-evasive and immune-suppressive features in carcinomas. An unexplored avenue has been the universal applicability of the EMT program in majority of solid tumors to modulate the immune response in multiple other cancer types. Along these lines, Benboubker et al., discuss in their review how the phenotypic plasticity of melanomas promotes immune escape. They document several studies that demonstrate how de-differentiated melanoma cells with more-mesenchymal properties evade immune attack by the loss of melanoma antigens. Moreover, over-expression of Zeb1 in melanoma cells dampens their ability to release CXCL10, a potent chemoattractant for CD8+ T-cells. As a consequence, Zeb1 expressing melanoma cells exclude CD8+ T-cells and are resistant to anti-PD1 immune checkpoint blockade.
Meng et al., focus specifically on uveal melanoma and demonstrate that the transcription factor PRRX1 promotes uveal melanoma progression by activating the EMT program in these cells. Strikingly, in addition to promoting invasion and motility, the expression of PRRX1 was associated with the expression of multiple inhibitory immune checkpoint molecules and poor prognosis.

Further elaborating on the broad spectrum of EMT-induced immune-suppression, Gu et al., address the acquisition of immune-evasive and immune-suppressive properties by soft tissue sarcomas, which are also enriched in cancer stem-like cells. By using a combination of machine learning and transcriptomic approaches, the authors identified stemness signatures in patient samples. Importantly, a low stemness signature was associated with increased activation of various innate and adaptive immune cells, better prognosis and an increased likelihood to respond to immunotherapy.

As mentioned above, the EMT program is a highly dynamic process with multiple intermediate or hybrid states that can also exist along the polarization spectrum. Sahoo et al., focused on determining whether carcinoma cells residing in a hybrid epithelial-mesenchymal state also exhibit immune-evasive features. To this end, the authors constructed and simulated the dynamics of a minimalistic regulatory network that contained hallmark EMT markers and PD-L1 (Programmed death-ligand 1). The simulation network, which was also integrated with scRNA-Seq and bulk RNA-Seq, revealed that cancer cells residing in a hybrid EMT state are indeed likely to express higher levels of PD-L1. Moreover, in addition to expressing PD-L1, breast cancer cells that exhibit a hybrid EMT state showed enhanced resistance to anti-estrogen therapy.

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In terms of clinical utility, Zhao et al., review recent literature that is centered on targeting the hepatocyte growth factor/cellular-mesenchymal-epithelial transition (HGF/c-Met) signaling pathway in hepatocellular carcinomas. The authors summarize multiple clinical studies that demonstrate the potent anti-tumor activity of a MET inhibitor and suggest targeting MET in combination with other biomarkers as a therapeutic strategy for patients with advanced hepatocellular carcinoma. Additionally, Engelsen et al., review emerging evidence supporting the utility of targeting the receptor tyrosine kinase, AXL, to enhance the efficacy of immune checkpoint blockade inhibitors. While AXL is expressed by multiple cell types, in the context of cancer cells, it is closely associated with epithelial plasticity. Strikingly, its expression is induced on cells undergoing an EMT as well as on cells with high stem cell-like phenotype. Moreover, targeting AXL can induce profound changes within the tumor microenvironment, likely enhancing sensitization to immune checkpoint blockade inhibitors.

We hope that this special Research Topic collection contributes to further strengthening the association of the EMT program with immune-suppression in multiple cancer types, making it an attractive therapeutic target and predictive criteria to consider while implementing immunotherapeutic strategies.

**AUTHOR CONTRIBUTIONS**

All authors listed have made a substantial, direct, and intellectual contribution to the work and approved it for publication.

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