Liver cancer was ranked the sixth most commonly diagnosed cancer with the fourth leading cause of cancer-related deaths worldwide in 2018 (1). Notably, hepatocellular carcinoma (HCC) constitutes 75–85% among all the liver cancer cases, and ranks first morbidity and mortality of malignancies in Chinese males under 60. Due to hypervascularity in liver and high heterogeneity, HCC was regarded as an aggressive disease. Recent clinical trials focusing on resection and transplantation after targeted therapy for advanced HCC show encouraging results. However, the adjuvant therapy for patients with high risk of recurrence after surgery still raises attention. Currently, emerging evidence suggested that the crosstalk between tumor cells and the microenvironment plays a pivotal role in tumorigenesis and metastasis. The tumor microenvironment of HCC provides tumor cells with necessary conditions for their sustained growth, invasion and metastasis. It is typically characterized by dysregulated immune network, sustained injury and regeneration, angiogenesis, inflammation and metabolic reprogramming. These aforementioned characteristics raise attention to the important role of infiltrating immune cell profile and tumor stroma in microenvironmental regulation. According to the immune cell subset and proportion, HCC could be classified into different immune subtypes, for example, Sia’s “immune class/exclusion” and Kurebayashi’s “immune high/mid/low”. Zhang et al. further identified three novel distinctive HCC subtypes with immunocompetent, immunodeficient, and immunosuppressive features (2). Immunocompetent and immunosuppressive subtype were characterized by robust immune cell infiltration and upregulated immune checkpoint molecules. These two subtypes, as well as the “immune class” and “immune-high subtype”, displayed significant increase in immune cells infiltration and better responsiveness to immunotherapy. Tumor stroma and vasculature could also cause considerable impact on tumor progression. Cancer-associated fibroblasts (CAFs) can result in a robust stromal reaction characterized by fibrotic extracellular matrix and make tumor cells convert to aggressive and immune-excluded phenotype via transforming growth factor-β (TGF-β) signaling pathway (3). Tumor-associated vasculature is another main hallmark of cancers. Endothelial cells (ECs) of the tumor vasculature can not only inhibit anti-tumor immunity by immune cells regulation and elimination, but also help to create a proangiogenic microenvironment via vascular endothelial growth factor (VEGF) signaling pathway (4). The disorganized tumor vasculature could remodel a special microenvironment characterized by hypoxia, acidosis, and high interstitial pressure, which both support tumor cells proliferation and structure a physical barrier for infiltrating immune cells (5). Furthermore, the interaction between different components relies on the regulation of chemokines and cytokines, which play essential roles in recruiting immune cells and maintaining cancer and stromal cell biology (6). In another word, these microenvironmental populations profoundly influence the tumor immune and metabolic homeostasis, which suggests immunometabolism as a novel perspective of HCC microenvironment.
The concept of immunometabolism has highlighted the significance of crosstalk between immune microenvironment and metabolic reprogramming rather than isolated immune or metabolic alteration. For instance, macrophage plays a leading role in the immunometabolic microenvironment of HCC due to its quantity. On the one hand, it could be activated by metabolic signals and subsequently stimulated hepatocyte lipogenesis and cell death. On the other, the metabolic alteration in macrophages could further lead to immune reaction suppression, tumor proliferation and angiogenesis (7). Hence, interruption of those immunometabolism-associated cells or molecules would be a new direction of targeting tumor microenvironment for the treatment of HCC.

The possibility of targeting immunometabolism for tumor therapy is extensively studied in T cells (8). Selective alteration of metabolic process such as glycolysis or oxidative phosphorylation could help rescue T cell exhaustion within microenvironment and in turn improve the efficacy of checkpoint inhibitors (9). As for adoptive cell transfer, T cells could be metabolically manipulated in vitro to promote mitochondrial integrity and oxidative metabolism, which would better support their in vivo anti-tumor function and longevity (9). Metabolic reprogramming of innate immune cells also provides options to restore normal regional immune surveillance. Recent work has shown that glycolytic enzyme PFKFB3 was a key driver of PD-L1 in macrophages, and thus conducted T cell exhaustion in HCC, which could be reversed by inhibition of PFKFB3 (10). And intracellular accumulation of cholesterol was known to have resulted in activated effector function of NK cells against hepatoma cells. Strategies to increase cholesterol uptake by NK cells might therefore be developed for treatment of HCC (11). These findings suggest that key metabolic molecules including enzymes and metabolites have the potential to be "immunometabolic checkpoints" that reprogram immune cells metabolically into an anti-tumor phenotype and support traditional immunotherapies like PD-1/L1 inhibitor or CAR-T.

However, targeting the primary tumor and its surrounding microenvironment is just the “first round” of tumor treatment. There is yet a distance to tumor eradication, due to the inevitable invasion and metastasis. Hence, we carefully proposed the attention to preventing secondary tumor microenvironment formation as well, that is, pre-metastatic niche. The pre-metastatic niche can be defined as a supportive “soil” which is undergoing the microenvironment profile changes to form the specific sites in preparation for metastasis, which can also be regarded as the extension of primary microenvironment. Targeting the pre-metastatic niche-promoting components to suppress pre-metastatic niche formation can consequently result in the “imprisonment” of HCC. Disabling its invasiveness provides valuable treatment time window for patients who are waiting for transplantation or those who are unable to sustain surgical damage. In these settings, targeting the molecular events evolved in multiple steps of metastasis provides options for us to interfere these processes. Several signaling axes including stromal-derived factor-1 (SDF-1)/C-X-C motif chemokine receptor 4 (CXCR4) and interleukin-6 (IL-6)/signal transducer and activator of transcription 3 (STAT3) have been reported to be crucial for pre-metastatic niche formation in pre-clinical animal models (12,13). Blocking the delivery of metastasis-promoting exosomes to recipient cells may be another effective strategy for preventing HCC metastasis due to their roles in educating the pre-metastatic niche (14). Given the complicated composition within exosomes, deeper understanding of exosomes’ role in metastasis and a more specific targeting tool are needed.

Besides pre-metastatic niche, some researchers have also bestowed their attention to field cancerization, which is defined as the cancer-primed cell population that underlies the development of many cancers, albeit presented without morphological changes (15). The surrounding microenvironment may be altered by the cancerized field and cancerization would progress in an appropriate microenvironmental context. Nowadays, the clinical issue presented by field cancerization mainly focused on risk prediction and surgical margin decision. Understanding the evolution of cells and surrounding microenvironment within these cancerized fields may make it possible to intervene ahead of tumor progression.

To date, the fundamental researches have already revealed the important role of tumor microenvironment in tumorigenesis and metastasis. A better understanding of microenvironmental regulation will facilitate a breakthrough in precision medicine and provide a rationale for clinical study. Besides that, targeting pre-metastatic niche as an adjuvant therapy will besiege tumor cells locally by disabling their invasions. This approach hopefully provides time window for patients to receive curative surgical treatment. It is promising that further investigation on the mechanism of pre-metastatic niche formation, even the cancerized field would provide a new insight into the
prevention and treatment of HCC.

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Footnote

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