

Regulation and potential drug targets of tumor-associated Tregs

January 16 2019



Regulatory T cells (Tregs) negatively regulate the antitumour immune responses by inhibiting the function of effector T cells (CTL) and antigen presenting cells



(APC). Credit: Science China Press

According to estimates from the World Health Organization (WHO), there will be 18.1 million new cases and 9.6 million cancer deaths worldwide in 2018. Cancer is a serious disease that affects people all over the world. And studies on the treatment of tumors have far-reaching significance.

In recent years, <u>tumor</u> immunotherapy has become the fourth-ranking cancer treatment measure in line with traditional surgical treatment, radiotherapy and chemotherapy, and targeted therapy. The 2018 Nobel Prize in Physiology or Medicine was awarded to this field because of the amazing effects of <u>cancer</u> immunotherapies.

Nobel laureate James Allison studied a known protein called CTLA-4, which functions as a brake when antigen-presenting cells (APCs) stimulate T cells. CTLA-4 antibodies for the treatment of melanoma were approved by the FDA in 2011. Meanwhile, Tasuku Honjo discovered PD-1, a protein also operates as a brake, but with a different mechanism to mediate immune escape of tumor cells. Therapies based on this discovery are strikingly effective, especially for <u>solid tumors</u>. Five antibody drugs targeting PD-1/PD-L1 have been used for more than a dozen indications in the past four years.

However, the objective response rate of patients is only 20 to 30 percent, on average, indicating that additional factors contributing to resistance to PD-1/PD- L1 monoclonal antibody therapies need to be identified. Evidence indicates that regulatory T cells (Tregs) might play a key role in regulating immune response to tumors.

Tregs, formerly known as suppressor T cells, play an important role in



maintaining self-tolerance and immune homeostasis, but an inhibitory role in evoking effective antitumour immune responses. Studies in both humans and animal models indicate that Tregs accumulate in the tumor sites of various cancers. Notably, the increased numbers of Tregs are positively correlated with poor prognosis, but negatively correlated with patient survival in the majority of tumours.

There is ample evidence indicating that the depletion of Tregs or the inhibition of Treg function will enhance antitumour effects. Based on these findings, targeting tumour-associated Tregs might be an attractive strategy for alleviating tumour immunesuppression and improving immune responses against cancers. However, suitable candidates with minimal toxic side effects have not been found so far.

In a new systematic review in *Science China Life Sciences*, researchers have summarized the characteristics of tumour-associated Tregs, the molecular basis of the regulation of Tregs, and the potential drug targets that have been discovered. Moreover, they discuss the scientific issues about the regulation of tumor-associated Tregs that require further research and the possible directions for exploring potential therapeutic targets.

Regarding the regulation of tumour-associated Tregs, both Treg-intrinsic and extrinsic factors contribute to the modulation. Endogenous factors are mainly derived from key signaling pathways in Treg cells, and exogenous factors are derived from other types of <u>cells</u> at the tumor site as well as the particularly complex tumor microenvironment. According to the researchers, the interaction between Tregs and other cell types in the TME and the effects of the abnormal TME metabolism on Tregs require deeper study.

"In addition to paying attention to highly expressed markers of tumourassociated Tregs, focusing on the instability of Tregs may also be a good



choice for developing Treg-targeted immunotherapies," addressed by the researchers, when talking about how to explore suitable targets.

More information: Shanshan Yan et al, The function and potential drug targets of tumour-associated Tregs for cancer immunotherapy, *Science China Life Sciences* (2019). DOI: 10.1007/s11427-018-9428-9

Provided by Science China Press

Citation: Regulation and potential drug targets of tumor-associated Tregs (2019, January 16) retrieved 26 April 2023 from <u>https://medicalxpress.com/news/2019-01-potential-drug-tumor-associated-tregs.html</u>

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