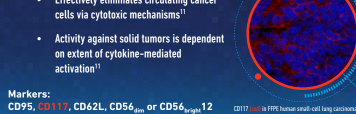
# STARS OF THE SHOW

The immune system plays a pivotal role in tumor formation, development, and metastasis. Cancer cells are inherently antigenic, which normally allows immune cells to identify and eliminate them prior to tumor formation. Tumor formation occurs when cancer cells develop methods to evade or outpace immune-mediated killing. Understanding this relationship between immune and cancer cells is therefore integral to restoring immune system potency for cancer therapeutics.

# NATURAL KILLER (NK) CELLS

# Mechanism:

- Effectively eliminates circulating cancer cells via cytotoxic mechanisms<sup>11</sup>
- Activity against solid tumors is dependent on extent of cytokine-mediated







• Dendritic cells (DCs) and macrophages are professional antigen-presenting cells (APCs) pivotal for activating T cells13

DENDRITIC CELLS AND MACROPHAGES:

- Macrophages also kill cells via phagocytosis or cytotoxic mechanisms; phenotypes range from pro-inflammatory to anti-inflammatory/pro-repair14 • Cancer cell-secreted cytokines cause tumor-infiltrating DCs to switch to an immuno-suppressive
- phenotype, while tumor-associated macrophages (TAMs) present anti-inflammatory phenotypes, inhibit T cell activity, and promote angiogenesis, tumor growth, and metastasis 13,14

Checkpoint proteins and the pathways they activate are critical for immune self-regulation<sup>15</sup>

• The ability to inhibit immune responses is key for limiting collateral damage and maintaining

• Cancer cells have co-opted the activation of these pathways to deactivate immune-mediated

 Checkpoint inhibition – using exogenous agents to prevent cancer cell-mediated checkpoint pathway activation – is a popular anti-cancer therapeutic strategy undergoing intensive

tumoricidal mechanisms, thereby facilitating tumor immune evasion<sup>19</sup>



Macrophage-marker CD68 expression (magenta) in FFPE human tonsil.

The primary effectors of immune-mediated cell death, T cells exert their tumoricidal functions by recognizing antigens presented on tumor cells' surfaces.<sup>2</sup> Tumor cells evade T cells through nutrient deprivation,<sup>3</sup> promoting cell inactivation, and activating immunosuppression mechanisms.<sup>2</sup> Augmenting T cell activity to counteract these effects is a primary focal point of immuno-oncology research.

T CELLS



# CYTOTOXIC T CELLS (CTLs)

# Mechanism:

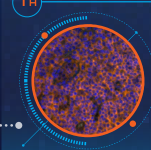
(TCR)-major histocompatibility complex (MHC)-antigen presentation

Markers:

- Primed and activated through T cell receptor
- Releases cytotoxins to kill cells expressing

CD8, CD44, CD62, 4

# HELPER T CELLS (TH CELLS)



# Mechanism:

- Regulates immune system function through cytokine secretion and activation of macrophages, B cells, and CTLs
- Vital for anti-tumor protection<sup>5</sup> Markers: CD4; distinguished from T<sub>reg</sub> cells (also CD4+) by secretion profile (Th1 cells secrete IFNy, The interleukins (ILs) 4, 13, and 5, and T<sub>h17</sub> ILs 17 and 21)6

REGULATORY T CELLS (TREG CELLS)

- Suppresses immune system activity to prevent deleterious inflammation and autoimmune
- Tumor cells promote T<sub>req</sub> recruitment, resulting in immunosuppression and evasion8

# Markers:

FoxP3, CD258

# microenvironment," Nat Immunol 14(10):1014-1022, 2013.

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# **ENDOTHELIAL CELLS**

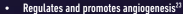
Checkpoint Pathway Proteins: PD-1, PD-L1; CTLA-4, CD80/CD86<sup>19,20</sup>

' IMMUNE CHECKPOINTS

Mechanism:

self-tolerance19

# Mechanism:



 Controls tumor cell intra/extravasation, metastasis, and immune cell infiltration<sup>23</sup>



PD-L1 (orange) in FEPE human lung carcinoma.

Markers: CD31, von Willebrand Factor<sup>24</sup>

# CD31 expression (red) in FFPE human breast carcinoma

 Produces antibodies that promote anti-tumor T cell, macrophage, and NK cell activity9

B CELLS

Mechanism:

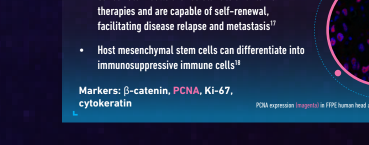
 Can encourage tumor development by producing growth factors and autoantibodies9

CD19, CD20, CD21, CD40, CD80, CD86, & CD69<sup>10</sup>

# **CANCER CELL MARKERS**

## Mechanism:

 Cancer stem cells are resistant to anti-tumor therapies and are capable of self-renewal. facilitating disease relapse and metastasis1



# **FIBROBLASTS**

# Mechanism:

- Creates a favorable environment for tumor growth by secreting growth factors and extracellular matrix21
- Promotes angiogenesis as well as recruitment of

Markers: α-smooth muscle actin, vimentin, desmin, platelet derived growth factor receptor<sup>22</sup>









