


HIGHLIGHTS FROM THE LITERATURE

- 425 What We're Reading

CANCER IMMUNOLOGY AT THE CROSS-ROADS

- 426  Novel "Elements" of Immune Suppression within the Tumor Microenvironment
Devikala Gurusamy, David Clever, Robert Eil, and Nicholas P. Restifo

MEETING REPORT

- 434 Cancer Immunology and Immunotherapy: Taking a Place in Mainstream Oncology
Keystone Symposia Meeting Summary
Matthew M. Gubin

CANCER IMMUNOLOGY MINIATURES

- 439 4-1BB-Enhanced Expansion of CD8⁺ TIL from Triple-Negative Breast Cancer Unveils Mutation-Specific CD8⁺ T Cells
Michiko Harao, Marie-Andrée Forget, Jason Roszik, Hui Gao, Gildy V. Babiera, Savitri Krishnamurthy, Jessica A. Chacon, Shumin Li, Elizabeth A. Mittendorf, Sarah M. DeSnyder, Korrene F. Rockwood, Chantale Bernatchez, Naoto T. Ueno, Laszlo G. Radvanyi, Luis Vence, Cara Haymaker, and James M. Reuben
Triple-negative breast cancers have low somatic mutational loads. By culturing tumor-infiltrating lymphocytes with agonistic 4-1BB mAb, tumor-specific T cells were expanded and the mutations to which they responded identified, providing a rationale for adoptive T cell therapy.

RESEARCH ARTICLES


- 446 Combined Anti-VEGF and Anti-CTLA-4 Therapy Elicits Humoral Immunity to Galectin-1 Which Is Associated with Favorable Clinical Outcomes
Xinqi Wu, Jingjing Li, Erin M. Connolly, Xiaoyun Liao, Jing Ouyang, Anita Giobbie-Hurder, Donald Lawrence, David McDermott, George Murphy, Jun Zhou, Matthias Piesche, Glenn Dranoff, Scott Rodig, Margaret Shipp, and F. Stephen Hodi
Galectin-1 is often produced by tumors and is protumoral, proangiogenic, and immunosuppressive. Ipilimumab plus bevacizumab induced production of neutralizing antibodies to galectin-1, which correlated with better clinical outcomes in metastatic melanoma patients, highlighting its utility as a therapeutic target.
- 455 RIG-I Resists Hypoxia-Induced Immunosuppression and Dedifferentiation
Christina Engel, Grethe Brüggmann, Silke Lambing, Larissa H. Mühlenbeck, Samira Marx, Christian Hagen, Dorottya Horváth, Marion Goldeck, Janos Ludwig, Anna-Maria Herzner, Jan W. Drijfhout, Daniela Wenzel, Christoph Coch, Thomas Tüting, Martin Schlee, Veit Hornung, Gunther Hartmann, and Jasper G. Van den Boom
Solid tumors are generally hypoxic. RIG-I, but not IFN α , still functioned under hypoxia. Activating RIG-I and using vitamin C to scavenge free radicals in a melanoma model augmented NK and CD8⁺ T cell antitumor functions and prolonged survival.
- 468 A STING Agonist Given with OX40 Receptor and PD-L1 Modulators Primes Immunity and Reduces Tumor Growth in Tolerized Mice
Jeremy B. Foote, Marlene Kok, James M. Leatherman, Todd D. Armstrong, Bridget C. Marcinkowski, Lauren S. Ojalvo, David B. Kanne, Elizabeth M. Jaffee, Thomas W. Dubensky Jr, and Leisha A. Emens
The efficacy and immune dynamics of STING modulation in the toleragenic tumor microenvironment were examined. Combining a STING agonist, PD-L1 blockade, and OX40R stimulation created an inflamed tumor microenvironment that recruited T cells and activated tumor-specific immunity.
- 480  Soluble PD-L1 as a Biomarker in Malignant Melanoma Treated with Checkpoint Blockade
Jun Zhou, Kathleen M. Mahoney, Anita Giobbie-Hurder, Fengmin Zhao, Sandra Lee, Xiaoyun Liao, Scott Rodig, Jingjing Li, Xinqi Wu, Lisa H. Butterfield, Matthias Piesche, Michael P. Manos, Lauren M. Eastman, Glenn Dranoff, Gordon J. Freeman, and F. Stephen Hodi
Melanoma cells could secrete several splice variants of PD-L1. Secretion differed among patients, and was affected by checkpoint therapy, with some changes associated with progressive disease, and others with favorable outcomes, suggesting circulating PD-L1 as a dynamic biomarker.

Table of Contents

493 Tumor-Derived α -Fetoprotein Directly Drives Human Natural Killer-Cell Activation and Subsequent Cell Death

Lazar Vujanovic, Elizabeth C. Stahl, Angela D. Pardee, David A. Geller, Allan Tsung, Simon C. Watkins, Gregory A. Gibson, Walter J. Storkus, and Lisa H. Butterfield

Low NK cell numbers, function, and infiltration into tumors predict poor outcomes for patients with hepatocellular carcinoma (HCC). Tumor-derived α -fetoprotein (AFP) from HCCs directly impacted NK cell function and viability, through both the AFP protein and AFP's low-molecular-mass cargo.

503 Combining DNA Vaccine and AIDA-1 in Attenuated *Salmonella* Activates Tumor-Specific CD4⁺ and CD8⁺ T-cell Responses

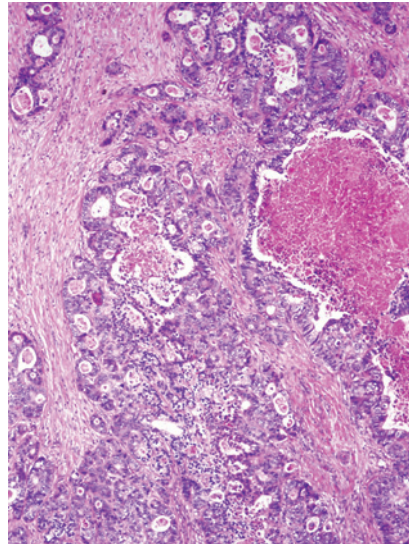
Yu Mei, Lixiang Zhao, Yonghao Liu, Huanle Gong, Yuan Song, Lei Lei, Ying Zhu, Ziqi Jin, Shoubao Ma, Bo Hu, Qing Sun, and Haiyan Liu
Effective tumor vaccines activate both CD4⁺ and CD8⁺ T cells. A melanoma DNA vaccine delivered by a bacterial system that ensured presentation of both class I and class II peptides activated both arms of the adaptive immune response.

 AC icon indicates AuthorChoice

For more information please visit www.aacrjournals.org

ABOUT THE COVER

This issue of *Cancer Immunology Research* includes a Cancer Immunology at the Crossroads article that highlights two of the many ways in which tumors can evade immune surveillance. Both of these “tricks” employ the repurposing of how two elements, oxygen and potassium, engage with the tumor microenvironment and the resultant suppression of antitumor immune responses. Read more in Gurusamy et al., beginning on page 426. The original image is of a resected metastatic colon tumor stained with hematoxylin and eosin. Photo from the Restifo laboratory. Artwork by Lewis Long.



Cancer Immunology Research

5 (6)

Cancer Immunol Res 2017;5:425-514.

Updated version Access the most recent version of this article at:
<http://cancerimmunolres.aacrjournals.org/content/5/6>

E-mail alerts [Sign up to receive free email-alerts](#) related to this article or journal.

Reprints and Subscriptions To order reprints of this article or to subscribe to the journal, contact the AACR Publications Department at pubs@aacr.org.

Permissions To request permission to re-use all or part of this article, use this link <http://cancerimmunolres.aacrjournals.org/content/5/6>. Click on "Request Permissions" which will take you to the Copyright Clearance Center's (CCC) Rightslink site.