NF-κB, an Active Player in Human Cancers
Yifeng Xia, Shen Shen, and Inder M. Verma

Combining Radiation and Immunotherapy: A New Systemic Therapy for Solid Tumors?
Chad Tang, Xiaohong Wang, Hendrick Soh, Steven Seyedin, Maria Angelica Cortez, Sunil Krishnan, Erminia Massarelli, David Hong, Aung Naing, Adi Diab, Daniel Gomez, Huiping Ye, John Heymach, Ristuko Komaki, James P. Allison, Padmanee Sharma, and James W. Welsh

Antigen-Specific Culture of Memory-like CD8 T Cells for Adoptive Immunotherapy
Adam J. Litterman, David M. Zellmer, Rebecca S. LaRue, Stephen C. Jameson, and David A. Largaespada

In Vitro Characterization of the Anti-PD-1 Antibody Nivolumab, BMS-936558, and In Vivo Toxicology in Non-Human Primates
Changyu Wang, Kent B. Thudium, Minhua Han, Xi-Tao Wang, Haichun Huang, Diane Feingersh, Candy Garcia, Yi Wu, Michelle Kuhne, Mohan Srinivasan, Sujata Singh, Susan Wong, Neya Garner, Heidi Leblanc, R. Todd Bunch, Diann Blanet, Mark J. Selby, and Alan J. Korman

Mechanisms That Can Promote Peripheral B-cell Lymphoma in ATM-Deficient Mice
Suprawee Tepsuporn, Jiazhi Hu, Monica Gostissa, and Frederick W. Alt

Synopsis: Tepsuporn, Hu, and colleagues generated the first mouse models for B-cell lymphoma in the context of ATM deficiency, and they provide a detailed characterization of the mature B-cell lymphomas that arise, revealing an unanticipated mechanism for the developmental propagation of V(D)J recombination-initiated DNA double-strand breaks.

Targeting 4-1BB Costimulation to the Tumor Stroma with Bispecific Aptamer Conjugates Enhances the Therapeutic Index of Tumor Immunotherapy
Brett Schrand, Alexey Berezhnoy, Randall Brennesen, Anthony Williams, Agata Levay, Ling-Yuan Kong, Ganesh Rao, Shouhao Zhou, Amy B. Heimberger, and Eli Gilboa

Synopsis: Schrand and colleagues report the efficacy in five murine tumor models of an immunotherapeutic approach whereby systemic administration of tumor stroma-targeted 4-1BB aptamer conjugates, which target disseminated tumor lesions, elicits potent antitumor immunity with minimal dose-limiting toxicity.

CALGB 150905 (Alliance): Rituximab Broadens the Antilymphoma Response by Activating Unlicensed NK Cells
Juan Du, Sandra Lopez-Verges, Brandelyn N. Pitcher, Jeffrey Johnson, Sin-Ho Jung, Lili Zhou, Katharine Hsu, Myron S. Czuczman, Bruce Cheson, Lawrence Kaplan, Lewis L. Lanier, and Jeffrey M. Venstrom

Synopsis: Du and colleagues report that a “missing ligand” genotype predictive of unlicensed NK cells was associated with higher progression-free survival in 101 follicular lymphoma patients treated with rituximab-containing mAb combinations, and that rituximab triggered responses in vitro from healthy-donor unlicensed NK cells.

IL32 Is Progressively Expressed in Mycosis Fungoides Independent of Helper T-cell 2 and Helper T-cell 9 Polarization
Hanako Ohmatsu, Daniel Humme, Nicholas Culati, Juana Gonzalez, Markus Möbs, Mayte Suárez-Fariñas, Irmia Cardinale, Hiroshi Mitsui, Emma Guttmann-Yassky, Wolfram Sterry, and James G. Krueger

Synopsis: Ohmatsu and colleagues report the consistently high and increasing expression of IL32 in cutaneous T-cell lymphoma mycosis fungoides (MF) compared with benign inflammatory skin diseases, and these findings correlate with increases in IFNγ mRNA suggesting that IL32 may be an autocrine cytokine in MF progression.
<table>
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<tr>
<th>Page</th>
<th>Title</th>
<th>Authors</th>
<th>Synopsis</th>
</tr>
</thead>
<tbody>
<tr>
<td>901</td>
<td>STING Ligand c-di-GMP Improves Cancer Vaccination against Metastatic Breast Cancer</td>
<td>Dinesh Chandra, Wilber Quispe-Tintaya, Arthee Jahangir, Denise Asafu-Adjei, Ilyssa Ramos, Herman O. Sintim, Jie Zhou, Yoshihiro Hayakawa, David K.R. Karaolis, and Claudia Gravekamp</td>
<td>Chandra, Quispe-Tintaya, and colleagues show that stimulator of IFN genes (STING) ligand c-di-GMP activated caspase-3, stimulated T cells, and nearly completely eliminated all metastases in mouse breast cancer model 4T1, when combined with Listeria monocytogenes–based Mage-b vaccine in a therapeutic setting.</td>
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<td>911</td>
<td>Episomal Expression of Truncated Listeriolysin O in LmddA-LLO–E7 Vaccine Enhances Antitumor Efficacy by Preferentially Inducing Expansions of CD4⁺FoxP3⁻ and CD8⁺ T Cells</td>
<td>Zhisong Chen, Laurent Ozbun, Namju Chong, Anu Wallecha, Jay A. Berzofsky, and Samir N. Khleif</td>
<td>Chen and colleagues developed an improved, attenuated L. monocytogenes–based vaccine that induced the regression of established mouse TC-1 tumors; they show that listeriolysin O serves as a vaccine adjuvant that decreases Treg frequency by inducing the expansion of non-Treg T cells.</td>
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<td>923</td>
<td>Correction: Bevacizumab plus Ipilimumab in Patients with Metastatic Melanoma</td>
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ABOUT THE COVER

Nuclear factor kappa-light-chain-enhancer of activated B cells (NF-κB) comprises a family of five transcription factors (NF-κB1/p105/p50, NF-κB2/p100/p52, RelA/p65, RelB, and c-Rel) that form distinct heterodimer or homodimer protein complexes, which bind to consensus DNA sequences at promoter regions of responsive genes. Members of the NF-κB family are involved in signaling pathways controlling vital biologic processes, and stringent regulation of NF-κB activity is indispensable for the integrity of cellular functions. Oncogenic mutations can lead to constitutive and/or elevated NF-κB activity. NF-κB is also activated by cytokines, growth factors, cellular and environmental stresses, and DNA damage. In premalignant lesions with elevated NF-κB activity, the accumulation of proinflammatory cytokines contributes to the protumorigenic microenvironment. NF-κB is the master regulator mediating a cross-talk between inflammation and cancer at multiple levels. NF-κB activity promotes proliferation and angiogenesis, suppresses apoptosis, and induces epithelial–mesenchymal transition, which can lead to distant metastasis. At tumor sites, NF-κB activation may remodel local metabolism and anergize the immune system to favor tumor growth. Suppression of NF-κB in myeloid cells or tumor cells could lead to tumor regression, making the NF-κB pathway a promising therapeutic target. For more details of NF-κB activities in human cancers, see the Masters of Immunology primer by Inder M. Verma and colleagues starting on page 823 of this issue.

ABOUT THE MASTER

Inder Mohan Verma, PhD, is the Irwin and Joan Jacobs Chair Professor in Exemplary Life Science, Professor and Director of the Laboratory of Genetics at The Salk Institute for Biological Sciences (La Jolla, CA), and an American Cancer Society Professor of Molecular Biology. He is also an adjunct professor in the department of biology at the University of California, San Diego, and a distinguished adjunct professor in the Biotechnology Research Group at the King Abdulaziz University (Jeddah, Saudi Arabia). Dr. Verma was educated at the Lucknow University in India, and received his PhD in Biochemistry from the Weizmann Institute of Science (Rehovot, Israel), characterizing mitochondrial ribosomal RNA from the fungus Aspergillus nidulans, under the mentorship of Professor Uri Littauer. In 1971 he joined the laboratory of Nobel laureate David Baltimore at the Massachusetts Institute of Technology (Cambridge, MA), where he studied reverse transcriptase from RNA tumor viruses, including avian myeloblastosis virus, mouse leukemia virus, and hamster leukemia virus. Dr. Verma established his independent laboratory at The Salk Institute in 1974, and at the age of 26, he was one of the youngest faculty members.

Dr. Verma’s work on RNA tumor viruses and reverse transcriptase led to his identification and/or characterization of several oncogenes, including c-fos, c-rel, and the breast cancer genes (BRCA1, BRCA2). The Verma laboratory has contributed significantly to the delineation of the regulation and expression of c-fos during prenatal and postnatal development, growth, and differentiation. These investigators have characterized the mechanism of BRCA1 tumor suppression. C-rel is a member of the nuclear factor-κB (NF-κB) family of master transcription factors, the subject of this Masters primer. With expertise in molecular biology and retrovirology, Dr. Verma developed viral expression vectors for gene transfer to replace missing or defective cellular proteins, and this work has become the foundation for cell and gene therapy. The Verma laboratory has used the gene therapy technology to generate a mouse model of glioblastoma, from which they have identified neural cancer stem cells, as few as ten of which can induce tumors in immunodeficient mice. They are using the same technique to investigate the initiation and treatment of lung and prostate cancers.

Dr. Verma was born in Sangrur, Punjab, India. He has received many honors, including an Outstanding Investigator Award from the NIH (1988), and he was elected as a member of the Third World Academy of Sciences (1995), the National Academy of Sciences, India (1997), the U.S. National Academy of Sciences (NAS; 1997), the U.S. NAS Institute of Medicine (1999), the American Academy of Arts and Sciences (2000), the European Molecular Biology Organization (1998), and the American Philosophical Society (2006). Dr. Verma was elected as a Foreign Fellow of the Indian National Science Academy (2005). He was awarded the Vilcek Foundation Prize (2008), the ASGT Outstanding Achievement Award (2009), the Spector Prize (2010), and the Pasarow Award in Cancer Research (2010). Dr. Verma now serves as the Editor-in-Chief of the Proceedings of the National Academy of Science of the United States.