

Highlights of This Issue 2827

SPECIAL FEATURES


CCR Translations

- 2829** Precision Medicine for Diffuse Large B-cell Lymphoma
Megan S. Lim and Kojo S.J. Elenitoba-Johnson
See related article p. 2919
- 2832** MET Exon 14 Alterations in Lung Cancer: Exon Skipping Extends Half-Life
Alexander Drilon
See related article, p. 3048

CCR Drug Updates

- 2835** TAS-102 for Treatment of Advanced Colorectal Cancers That Are No Longer Responding to Other Therapies
Daphne L. van der Velden, Frans L. Opdam, and Emile E. Voest

Molecular Pathways

- 2840** Molecular Pathways: Emergence of Protein Kinase CK2 (CSNK2) as a Potential Target to Inhibit Survival and DNA Damage Response and Repair Pathways in Cancer Cells
 Adam J. Rabalski, Laszlo Gyenis, and David W. Litchfield

CLINICAL TRIAL BRIEF REPORT

- 2848** Phase Ib Study of PEGylated Recombinant Human Hyaluronidase and Gemcitabine in Patients with Advanced Pancreatic Cancer
Sunil R. Hingorani, William P. Harris, J. Thaddeus Beck, Boris A. Berdov, Stephanie A. Wagner, Eduard M. Pshevtolsky, Sergei A. Tjulandin, Oleg A. Gladkov, Randall F. Holcombe, Ronald Korn, Natarajan Raghunand, Samuel Dychter, Ping Jiang, H. Michael Shepard, and Craig E. Devoe

CANCER THERAPY: CLINICAL

- 2855** Phase I Study of Veliparib (ABT-888) Combined with Cisplatin and Vinorelbine in Advanced Triple-Negative Breast Cancer and/or BRCA Mutation-Associated Breast Cancer
Eve T. Rodler, Brenda F. Kurland, Melissa Griffin, Julie R. Gralow, Peggy Porter, Rosa F. Yeh, Vijayakrishna K. Gadi, Jamie Guenthoer, Jan H. Beumer, Larissa Korde, Sandra Strychor, Brian F. Kiesel, Hannah M. Linden, John A. Thompson, Elizabeth Swisher, Xiaoyu Chai, Stacie Shepherd, Vincent Giranda, and Jennifer M. Specht
- 2865** Endometrial Carcinomas with POLE Exonuclease Domain Mutations Have a Favorable Prognosis
Melissa K. McConechy, Aline Talhouk, Samuel Leung, Derek Chiu, Winnie Yang, Janine Senz, Linda J. Reha-Krantz, Cheng-Han Lee, David G. Huntsman, C. Blake Gilks, and Jessica N. McAlpine
- 2874** Phase I Study of Apatolisib (GDC-0980), Dual Phosphatidylinositol-3-Kinase and Mammalian Target of Rapamycin Kinase Inhibitor, in Patients with Advanced Solid Tumors
Saoirse O. Dolly, Andrew J. Wagner, Johanna C. Bendell, Hedy L. Kindler, Lee M. Krug, Tanguy Y. Seiwert, Marjorie G. Zauderer, Martijn P. Lolkema, Doris Apt, Ru-Fang Yeh, Jill O. Fredrickson, Jill M. Spoerke, Hartmut Koeppe, Joseph A. Ware, Jennifer O. Lauchle, Howard A. Burris 3rd, and Johann S. de Bono
- 2885** Vaccination with Irradiated Autologous Tumor Cells Mixed with Irradiated GM-K562 Cells Stimulates Antitumor Immunity and T Lymphocyte Activation in Patients with Recurrent Malignant Glioma
William T. Curry, Jr, Ramana Gorrepati, Matthias Piesche, Tetsuro Sasada, Pankaj Agarwalla, Pamela S. Jones, Elizabeth R. Gerstner, Alexandra J. Golby, Tracy T. Batchelor, Patrick Y. Wen, Martin C. Mihm, and Glenn Dranoff
- 2897** Phase I Dose-Escalation Study of Linsitinib (OSI-906) and Erlotinib in Patients with Advanced Solid Tumors
Valentine M. Macaulay, Mark R. Middleton, S. Gail Eckhardt, Charles M. Rudin, Rosalyn A. Juergens, Richard Gedrich, Sven Gogov, Sean McCarthy, Srinivasu Poondru, Andrew W. Stephens, and Shirish M. Gadgeel

Table of Contents

PERSONALIZED MEDICINE AND IMAGING

2908 Baseline Peripheral Blood Biomarkers Associated with Clinical Outcome of Advanced Melanoma Patients Treated with Ipilimumab

Alexander Martens, Kilian Wistuba-Hamprecht, Marnix Geukes Foppen, Jianda Yuan, Michael A. Postow, Phillip Wong, Emanuela Romano, Amir Khammari, Brigitte Dreno, Mariaelena Capone, Paolo A. Ascierto, Anna Maria Di Giacomo, Michele Maio, Bastian Schilling, Antje Sucker, Dirk Schadendorf, Jessica C. Hassel, Thomas K. Eigentler, Peter Martus, Jedd D. Wolchok, Christian Blank, Graham Pawelec, Claus Garbe, and Benjamin Weide

2919 Next-Generation Sequencing in Diffuse Large B-Cell Lymphoma Highlights Molecular Divergence and Therapeutic Opportunities: a LYSA Study

Sydney Dubois, Pierre-Julien Vially, Sylvain Mareschal, Elodie Bohers, Philippe Bertrand, Philippe Ruminy, Catherine Maingonnat, Jean-Philippe Jais, Pauline Peyrouze, Martin Figeac, Thierry J. Molina, Fabienne Desmots, Thierry Fest, Corinne Haioun, Thierry Lamy, Christiane Copie-Bergman, Josette Brière, Tony Petrella, Danielle Canioni, Bettina Fabiani, Bertrand Coiffier, Richard Delarue, Frédéric Peyrade, André Bosly, Marc André, Nicolas Ketterer, Gilles Salles, Hervé Tilly, Karen Leroy, and Fabrice Jardin

See related commentary, p. 2829

2929 A Novel Tumor-Specific Agent for Intraoperative Near-Infrared Fluorescence Imaging: A Translational Study in Healthy Volunteers and Patients with Ovarian Cancer



Charlotte E.S. Hoogstins, Quirijn R.J.G. Tummers, Katja N. Gaarenstroom, Cor D. de Kroon, J. Baptist M.Z. Trimbos, Tjalling Bosse, Vincent T.H.B.M. Smit, Jaap Vuyk, Cornelis J.H. van de Velde, Adam F. Cohen, Philip S. Low, Jacobus Burggraaf, and Alexander L. Vahrmeijer

2939 Incorporation of Next-Generation Sequencing into Routine Clinical Care to Direct Treatment of Head and Neck Squamous Cell Carcinoma

Nicole G. Chau, Yvonne Y. Li, Vickie Y. Jo, Guilherme Rabinowits, Jochen H. Lorch, Roy B. Tishler, Danielle N. Margalit, Jonathan D. Schoenfeld, Don J. Annino, Laura A. Goguen, Tom Thomas, Hailey Becker, Tyler Haddad, Jeffrey F. Krane, Neal I. Lindeman, Geoffrey I. Shapiro, Robert I. Haddad, and Peter S. Hammerman

2950 Alternate Metabolic Programs Define Regional Variation of Relevant Biological Features in Renal Cell Carcinoma Progression

Samira A. Brooks, Amir H. Khandani, Julia R. Fielding, Weili Lin, Tiffany Sills, Yueh Lee, Alexandra Arreola, Mathew I. Milowsky, Eric M. Wallen, Michael E. Woods, Angie B. Smith, Mathew E. Nielsen, Joel S. Parker, David S. Lalush, and W. Kimryn Rathmell

2960 Circulating Cell-Free Tumor DNA Analysis of 50 Genes by Next-Generation Sequencing in the Prospective MOSCATO Trial

Cécile Jovelet, Ecaterina Ileana, Marie-Cécile Le Deley, Nelly Motté, Silvia Rosellini, Alfredo Romero, Celine Lefebvre, Marion Pedrero, Noémie Pata-Merci, Nathalie Droin, Marc Deloger, Christophe Massard, Antoine Hollebecque, Charles Féré, Amélie Boichard, Sophie Postel-Vinay, Maud Ngo-Camus, Thierry De Baere, Philippe Vielh, Jean-Yves Scoazec, Gilles Vassal, Alexander Eggermont, Fabrice André, Jean-Charles Soria, and Ludovic Lacroix

CANCER THERAPY: PRECLINICAL

2969 Augmentation of Immune Checkpoint Cancer Immunotherapy with IL18



Zhifeng Ma, Wen Li, Shinichi Yoshiya, Yunfeng Xu, Masaki Hata, Yosif El-Darawish, Tzvetanka Markova, Kyosuke Yamanishi, Hiromichi Yamanishi, Hideaki Tahara, Yoshimasa Tanaka, and Haruki Okamura

2981 Not all NOTCH Is Created Equal: The Oncogenic Role of NOTCH2 in Bladder Cancer and Its Implications for Targeted Therapy

Tetsutaro Hayashi, Kilian M. Gust, Alexander W. Wyatt, Akihiro Goriki, Wolfgang Jäger, Shannon Awrey, Na Li, Htoo Zarni Oo, Manuel Altamirano-Dimas, Ralph Buttyan, Ladan Fazli, Akio Matsubara, and Peter C. Black

2993 PIGF/VEGFR-1 Signaling Promotes Macrophage Polarization and Accelerated Tumor Progression in Obesity






Joao Incio, Josh Tam, Nuh N. Rahbari, Priya Suboj, Dan T. McManus, Shan M. Chin, Trupti D. Vardam, Ana Batista, Suboj Babykutty, Keehoon Jung, Anna Khachatryan, Tai Hato, Jennifer A. Ligibel, Ian E. Krop, Stefan B. Puchner, Christopher L. Schlett, Udo Hoffmann, Marek Ancukiewicz, Masabumi Shibuya, Peter Carmeliet, Raquel Soares, Dan G. Duda, Rakesh K. Jain, and Dai Fukumura

BIOLOGY OF HUMAN TUMORS

3005 Tumor-Infiltrating Plasma Cells Are Associated with Tertiary Lymphoid Structures, Cytolytic T-Cell Responses, and Superior Prognosis in Ovarian Cancer

David R. Kroeger, Katy Milne, and Brad H. Nelson

Table of Contents

- 3016** **Inflammatory Marker Testing Identifies CD74 Expression in Melanoma Tumor Cells, and Its Expression Associates with Favorable Survival for Stage III Melanoma**
 Suhendan Ekmekcioglu, Michael A. Davies, Keiji Tanese, Jason Roszik, Myung Shin-Sim, Roland L. Bassett Jr, Denái R. Milton, Scott E. Woodman, Victor G. Prieto, Jeffrey E. Gershenwald, Donald L. Morton, Dave S. Hoon, and Elizabeth A. Grimm
- 3025** **Neoadjuvant Chemotherapy Modulates the Immune Microenvironment in Metastases of Tubo-Ovarian High-Grade Serous Carcinoma**
 Steffen Böhm, Anne Montfort, Oliver M.T. Pearce, Joanne Topping, Probir Chakravarty, Gemma L.A. Everitt, Andrew Clear, Jackie R. McDermott, Darren Ennis, Thomas Dowe, Amanda Fitzpatrick, Elly C. Brockbank, Alexandra C. Lawrence, Arjun Jeyarajah, Asma Z. Faruqi, Iain A. McNeish, Naveena Singh, Michelle Lockley, and Frances R. Balkwill
- 3037** **Reduced Expression of SMAD4 Is Associated with Poor Survival in Colon Cancer**
Pu Yan, Dirk Klingbiel, Zenia Saridaki, Paola Ceppa, Monica Curto, Thomas Alexander McKee, Arnaud Roth, Sabine Tejpar, Mauro Delorenzi, Fredrik T. Bosman, and Roberto Fiocca
- 3048** **MET Amplification and Exon 14 Splice Site Mutation Define Unique Molecular Subgroups of Non-Small Cell Lung Carcinoma with Poor Prognosis**
Joanna H. Tong, Sai F. Yeung, Anthony W.H. Chan, Lau Y. Chung, Shuk L. Chau, Raymond Wai Ming Lung, Carol Y. Tong, Chit Chow, Edith K.Y. Tin, Yau H. Yu, Hui Li, Yi Pan, Wing P. Chak, Calvin S.H. Ng, Tony S.K. Mok, and Ka F. To
See related commentary, p. 2832
- 3057** **T-Cell Immunoglobulin and ITIM Domain (TIGIT) Associates with CD8⁺ T-Cell Exhaustion and Poor Clinical Outcome in AML Patients**
Yaxian Kong, Liuluan Zhu, Todd D. Schell, Jianhong Zhang, David F. Claxton, W. Christopher Ehmann, Witold B. Rybka, Melissa R. George, Hui Zeng, and Hong Zheng
- 3067** **Circulating DNA as a Strong Multimarker Prognostic Tool for Metastatic Colorectal Cancer Patient Management Care**
Safia El Messaoudi, Florent Mouliere, Stanislas Du Manoir, Caroline Bascoul-Mollevi, Brigitte Gillet, Michelle Nouaille, Catherine Fiess, Evelyne Crapez, Frederic Bibeau, Charles Theillet, Thibault Mazard, Denis Pezet, Muriel Mathonnet, Marc Ychou, and Alain R. Thierry
- 3078** **IgG Glycome in Colorectal Cancer**
 Frano Vučković, Evropi Theodoratou, Kujtim Thaçi, Maria Timofeeva, Aleksandar Vojta, Jerko Štambuk, Maja Pučić-Baković, Pauline M. Rudd, Lovorka Đerek, Dražen Servis, Annika Wennerström, Susan M. Farrington, Markus Perola, Yurii Aulchenko, Malcolm G. Dunlop, Harry Campbell, and Gordan Lauc
- 3087** **HMGB1 and Its Hyperacetylated Isoform are Sensitive and Specific Serum Biomarkers to Detect Asbestos Exposure and to Identify Mesothelioma Patients**
Andrea Napolitano, Daniel J. Antoine, Laura Pellegrini, Francine Baumann, Ian Pagano, Sandra Pastorino, Chandra M. Goparaju, Kirill Prokrym, Claudia Canino, Harvey I. Pass, Michele Carbone, and Haining Yang
- 3097** **Epigenetic Regulation of the Homeobox Gene MSX1 Associates with Platinum-Resistant Disease in High-Grade Serous Epithelial Ovarian Cancer**
Nair A. Bonito, Jane Borley, Charlotte S. Wilhelm-Benartzi, Sadaf Ghaem-Maghami, and Robert Brown
- 3105** **NDUFA4L2 Fine-tunes Oxidative Stress in Hepatocellular Carcinoma**
Robin Kit-Ho Lai, Iris Ming-Jing Xu, David Kung-Chun Chiu, Aki Pui-Wah Tse, Larry Lai Wei, Cheuk-Ting Law, Derek Lee, Chun-Ming Wong, Maria Pik Wong, Irene Oi-Lin Ng, and Carmen Chak-Lui Wong
- CORRECTION**
- 3118** **Correction: Pretreatment microRNA Expression Impacting on Epithelial-to-Mesenchymal Transition Predicts Intrinsic Radiosensitivity in Head and Neck Cancer Cell Lines and Patients**

 AC icon indicates Author Choice

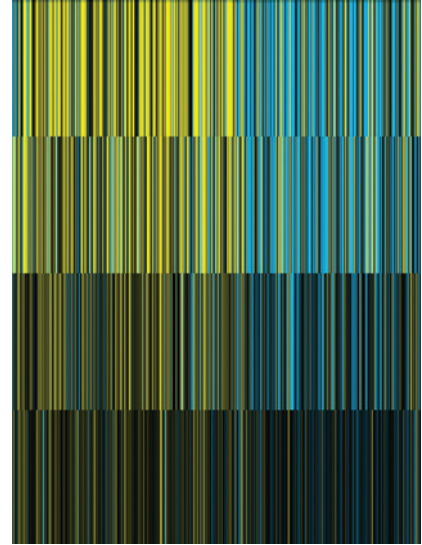
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Table of Contents

ABOUT THE COVER

The cover depicts gene expression differences between clear cell Renal Cell Carcinoma (ccRCC) patients from the Cancer Genome Atlas (TCGA) classified as either the clear cell A (ccA) or B (ccB) subtype. ccA and ccB patients separated by the activation of distinct metabolic genes associated with increased glycogen synthesis. For details, see the article by Brooks and colleagues on page 2950 of this issue.



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