


Highlights of This Issue 209**REVIEW**

- 211**  **Telomere-Regulating Genes and the Telomere Interactome in Familial Cancers**
Carla Daniela Robles-Espinoza,
Martin del Castillo Velasco-Herrera, Nicholas K. Hayward,
and David J. Adams

MCR RapidIMPACT

- 223** **A Mechanism for Asymmetric Cell Division Resulting in Proliferative Asynchronicity**
Ipsita Dey-Guha, Cleidson P. Alves, Albert C. Yeh,
Salony, Xavier Sole, Revati Darp, and
Sridhar Ramaswamy

CELL CYCLE AND SENESCENCE

- 231** **A p53/ARF-Dependent Anticancer Barrier Activates Senescence and Blocks Tumorigenesis without Impacting Apoptosis**
Vidya C. Sinha, Lan Qin, and Yi Li
- 239** **Global Increase of p16^{INK4a} in APC-Deficient Mouse Liver Drives Clonal Growth of p16^{INK4a}-Negative Tumors**
Elke Ueberham, Pia Glöckner, Claudia Göhler,
Beate K. Straub, Daniel Teupser, Kai Schönig,
Albert Braeuning, Anne Kathrin Höhn, Boris Jerchow,
Walter Birchmeier, Frank Gaunitz, Thomas Arendt,
Owen Sansom, Rolf Gebhardt, and Uwe Ueberham


CELL DEATH AND SURVIVAL

- 250** **Identification of TRIML2, a Novel p53 Target, that Enhances p53 SUMOylation and Regulates the Transactivation of Proapoptotic Genes**
Che-Pei Kung, Sakina Khaku, Matthew Jennis, Yan Zhou,
and Maureen E. Murphy
- 263** **Hypoxia Promotes Dissemination and Colonization in New Bone Marrow Niches in Waldenström Macroglobulinemia**
Barbara Muz, Pilar de la Puente, Fedá Azab,
Irene M. Ghobrial, and Abdel Kareem Azab

- 273** **Relocation of CLIC1 Promotes Tumor Cell Invasion and Colonization of Fibrin**
Lisa A. Gurski, Lynn M. Knowles, Per H. Basse,
Jodi K. Maranchie, Simon C. Watkins, and Jan Pilch

- 281** **Bile Acids Regulate Nuclear Receptor (Nur77) Expression and Intracellular Location to Control Proliferation and Apoptosis**
Ying Hu, Thinh Chau, Hui-Xin Liu, Degui Liao,
Ryan Keane, Yuqiang Nie, Hui Yang, and
Yu-Jui Yvonne Wan

CHROMATIN, GENE, AND RNA REGULATION

- 293**  **WHSC1 Promotes Oncogenesis through Regulation of NIMA-Related Kinase-7 in Squamous Cell Carcinoma of the Head and Neck**
Vassiliki Saloura, Hyun-Soo Cho, Kazuma Kiyotani,
Houda Alachkar, Zhixiang Zuo, Makoto Nakakido,
Tatsuhiko Tsunoda, Tanguy Seiwert, Mark Linggen,
Jonathan Licht, Yusuke Nakamura, and
Ryuji Hamamoto
- 305** **Transcriptome-wide Landscape of Pre-mRNA Alternative Splicing Associated with Metastatic Colonization**
Zhi-xiang Lu, Qin Huang, Juw Won Park, Shihao Shen,
Lan Lin, Collin J. Tokheim, Michael D. Henry, and
Yi Xing

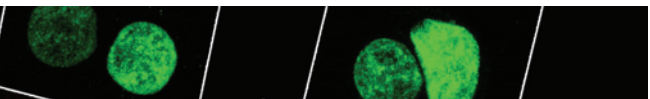
DNA DAMAGE AND REPAIR

- 319** **TGFβ1 Protects Cells from γ-IR by Enhancing the Activity of the NHEJ Repair Pathway**
Mi-Ra Kim, Jeeyong Lee, You Sun An, Yeung Bae Jin,
In-Chul Park, Eunkyung Chung, Incheol Shin,
Mary Helen Barcellos-Hoff, and Jae Youn Yi

GENOMICS

- 330** **lincRNA-RoR and miR-145 Regulate Invasion in Triple-Negative Breast Cancer via Targeting ARF6**
Gabriel Eades, Benjamin Wolfson, Yongshu Zhang,
Qinglin Li, Yuan Yao, and Qun Zhou
- 339** **The Landscape of Somatic Chromosomal Copy Number Aberrations in GEM Models of Prostate Carcinoma**
Daniella Bianchi-Frias, Susana A. Hernandez,
Roger Coleman, Hong Wu, and Peter S. Nelson

Table of Contents



ONCOGENES AND TUMOR SUPPRESSORS

348 RSK Promotes Prostate Cancer Progression in Bone through ING3, CKAP2, and PTK6-Mediated Cell Survival



Guoyu Yu, Yu-Chen Lee, Chien-Jui Cheng, Chuan-Fen Wu, Jian H. Song, Gary E. Gallick, Li-Yuan Yu-Lee, Jian Kuang, and Sue-Hwa Lin

358 DAPK3 Suppresses Acini Morphogenesis and Is Required for Mouse Development

Brandon A. Kocher, Lynn S. White, and David Piwnica-Worms

SIGNAL TRANSDUCTION

368 ERG Oncoprotein Inhibits ANXA2 Expression and Function in Prostate Cancer

Nicholas B. Griner, Denise Young, Pankaj Chaudhary, Ahmed A. Mohamed, Wei Huang, Yongmei Chen, Taduru Sreenath, Albert Dobi, Gyorgy Petrovics, Jamboor K. Vishwanatha, Isabell A. Sesterhenn, Shiv Srivastava, and Shyh-Han Tan

380 The Chemokine (CCL2–CCR2) Signaling Axis Mediates Perineural Invasion

Shizhi He, Shuangba He, Chun-Hao Chen, Sylvie Deborde, Richard L. Bakst, Natalya Chernichenko, William F. McNamara, Sei Young Lee, Fernando Barajas, Zhenkun Yu, Hikmat A. Al-Ahmadie, and Richard J. Wong

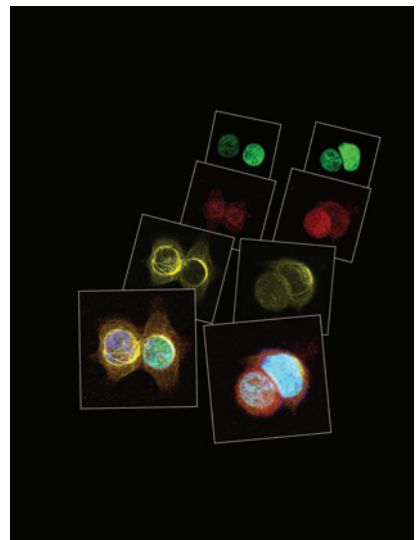


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ABOUT THE COVER

Slowly proliferating cancer cells can be hard to eradicate with currently available drugs, difficult to detect, and are thought to be a cause of late disease relapse. The Rapid Impact study within the current issue used confocal immunofluorescence microscopy to identify an MCF7 breast cancer cell dividing asymmetrically to produce a slowly proliferating daughter cell. The cover images show stains for H3K9me2 (green), phospho-AKT1-S473 (red), tubulin (yellow), and all three merged. See the Rapid Impact article by Dey-Guha, Alves and colleagues (beginning on page 223) for more information.



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