

Instructions for Authors 2021

General Policy. ANTICANCER RESEARCH (AR) will accept original high quality works and reviews on all aspects of experimental and clinical cancer research. The Editorial Policy suggests that priority will be given to papers advancing the understanding of cancer causation, and to papers applying the results of basic research to cancer diagnosis, prognosis, and therapy. Each article should include a concrete conclusion constituting a “new piece of knowledge” backed up by scientific evidence. AR will also accept the following for publication: (a) Abstracts and Proceedings of scientific meetings on cancer, following consideration and approval by the Editorial Board; (b) Announcements of meetings related to cancer research; (c) Short reviews (of approximately 120 words) and announcements of newly received books and journals related to cancer, and (d) Announcements of awards and prizes.

AR provides for the prompt print and online publication of accepted articles, generally within 1-2 months from final acceptance. Manuscripts will be accepted on the understanding that they report original unpublished works in the field of cancer research that are not under consideration for publication by another journal, and that they will not be published again in the same form. All authors should sign a submission letter confirming the approval of their article contents. All material submitted to AR will be subject to peer-review, when appropriate, by two members of the Editorial Board and by one suitable outside referee. All manuscripts submitted to AR are urgently treated with absolute confidence, with access restricted to the Managing Editor, the journal’s secretary, the reviewers and the printers. The Editors reserve the right to improve manuscripts on grammar and style.

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Manuscripts. Submitted manuscripts exceeding 4 printed pages will be subject to excess page charges. The 4 printed pages correspond approximately to twelve (12) document pages (~250 words per double-spaced typed page in Arial 12), including abstract, text, tables, figures, and references. All manuscripts should be divided into the following sections: (a) *First page* including the title of the presented work [not exceeding fifteen (15) words], full names and full postal addresses of all Authors, name of the Author to whom proofs are to be sent, key words, an abbreviated running title, an indication “review”, “clinical”, “epidemiological”, or “experimental” study, and the date of submission. (Note: The order of the Authors is not necessarily indicative of their contribution to the work. Authors may note their individual contribution(s) in the appropriate section(s) of the presented work or before the Acknowledgements); (b) *Abstract* not exceeding 150 words, organized according to the following headings: Background/Aim – Materials and Methods/Patients and Methods – Results – Conclusion; (c) *Introduction*; (d) *Materials and Methods/Patients and Methods*; (e) *Results*; (f) *Discussion*; (g) *Conflicts of Interest*; (h) *Authors’ contributions*; (i) *Acknowledgements*; (j) *References*. All pages must be numbered consecutively. Footnotes should be avoided. Review articles may follow a different style according to the subject matter and the Author’s opinion. Review articles should not exceed 35 pages (approximately 250 words per double-spaced typed page) including all tables, figures, and references.

Figures (graphs and photographs). All figures should appear at the end of the submitted document file. Once a manuscript is accepted all figures should be submitted separately in either jpg, tiff or pdf format and at a minimum resolution of 300 dpi. Graphs must be submitted as pictures made from drawings and must not require any artwork, typesetting, or size modifications. Figures should be prepared at a width of 8 or 17cm with eligible symbols, lettering and numbers. The number of each figure must be indicated. Pages that include color figures are subject to color charges.

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References. Authors must assume responsibility for the accuracy of the references used. Citations for the reference sections of submitted works should follow the form below and must be numbered consecutively. In the text, references should be cited by number in parenthesis. Examples: 1 Kenyon J, Liu W and Dagleish A: Report of objective clinical responses of cancer patients to pharmaceutical-grade synthetic cannabidiol. *Anticancer Res* 38(10): 5831-5835, 2018. PMID: 30275207. DOI: 10.21873/anticancer.12924. (PMIDs and DOIs only if

applicable). 2 McGuire WL and Chamnes GC: Studies on the oestrogen receptor in breast cancer. In: Receptors for Reproductive Hormones. O' Malley BW, Chamnes GC (eds.). New York, Plenum Publ Corp., pp 113-136, 1973. 3 Global Health Estimates 2015: Disease Burden by Cause, Age, Sex, by Country and by Region, 2000-2015. Geneva, World Health Organisation, 2016. Available at http://www.who.int/healthinfo/global_burden_disease/estimates/en/index2.html. Last accessed on 3rd April 2018. (The web address should link directly to the cited information and not to a generic webpage).

Nomenclature and Abbreviations. Nomenclature should follow that given in “Chemical Abstracts”, “Index Medicus”, “Merck Index”, “IUPAC -IUB”, “Bergey’s Manual of Determinative Bacteriology”, The CBE Manual for Authors, Editors and Publishers (6th edition, 1994), and MIAME Standard for Microarray Data. Human gene symbols may be obtained from the HUGO Gene Nomenclature Committee (HGNC) (<http://www.gene.ucl.ac.uk/>). Approved mouse nomenclature may be obtained from <http://www.informatics.jax.org/>. Standard abbreviations are preferable. If a new abbreviation is used, it must be defined on first usage.

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For International Standard Randomised Controlled Trials (ISRCTN) Registry (a not-for-profit organization whose registry is administered by Current Controlled Trials Ltd.) the unique number must be provided in this format: ISRCTNXXXXXXXX (where XXXXXXXX represents the unique number, always prefixed by “ISRCTN”). Please note that there is no space between the prefix “ISRCTN” and the number. Example: ISRCTN47956475.

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1. Anticancer Research (AR) closely follows the new developments in all fields of experimental and clinical cancer research by (a) inviting reviews on topics of immediate importance and substantial progress in the last three years, and (b) providing the highest priority for rapid publication to manuscripts presenting original results judged to be of exceptional value. Theoretical papers will only be considered and accepted if they bear a significant impact or formulate existing knowledge for the benefit of research progress.
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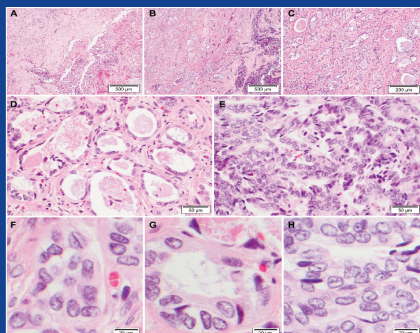
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- **CANCER GENOMICS & PROTEOMICS (CGP)** welcomes submissions of original high quality articles and reviews on all aspects of the application of genomic and proteomic technologies to experimental and clinical cancer research. The journal's scientific spectrum includes: (a) molecular causes of carcinogenesis, cancer progression and metastasis; (b) structural and functional aspects of genes in the cancer cell; (c) advances in genomic and proteomic technologies applicable to cancer research; (d) anticancer drug design and drug development. A main aim of CGP is to ensure the prompt and confidential review, and rapid publication of original works and reviews, generally within 1-3 months from submission.
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● Selection of Recent Articles

Micro RNAs Promoting Growth and Metastasis in Preclinical *In Vivo* Models of Subcutaneous Melanoma. U.H. WEIDLE, S. AUSLÄNDER, U. BRINKMANN (*Penzberg, Germany*)

Differential Proteomic Analysis of Hepatocellular Carcinomas from *Ppp2r5d* Knockout Mice and Normal (Knockout) Livers. C. LAMBRECHT, G.B. FERREIRA, J.D. OMELLA, L. LIBBRECHT, R. DE VOS, R. DERUA, C. MATHIEU, L. OVERBERGH, E. WAELEKENS, V. JANSSENS (*Leuven; Brussels, Belgium*)

Stem-like Cells from Invasive Breast Carcinoma Cell Line MDA-MB-231 Express a Distinct Set of Eph Receptors and Ephrin Ligands. M. LUCERO, J. THIND, J. SANDOVAL, S. SENAATI, B. JIMENEZ, R.P. KANDPAL (*Pomona, CA, USA*)

Circulating Tumor DNA in Biliary Tract Cancer: Current Evidence and Future Perspectives. A. RIZZO, A.D. RICCI, S. TAVOLARI, G. BRANDI (*Bologna, Italy*)

Whole-transcriptome Analysis of Fully Viable Energy Efficient Glycolytic-null Cancer Cells Established by Double Genetic Knockout of Lactate Dehydrogenase A/B or Glucose-6-Phosphate Isomerase. E. MAZZIO, R. BADISA, N. MACK, S. CASSIM, M. ZDRALEVIC, J. POUYSSEGUR, K.F.A. SOLIMAN (*Tallahassee, FL, USA; Monaco, Monaco; Nice, France*)

TIP60/P400/H4K12ac Plays a Role as a Heterochromatin Back-up Skeleton in Breast Cancer. M. IDRISOU, T. BOISNIER, A. SANCHEZ, F.Z.H. KHOUFAF, F. PENAULT-LLORCA, Y.-J. BIGNON, D. BERNARD-GALLON (*Clermont-Ferrand, France*)

STRA6 Expression Serves as a Prognostic Biomarker of Gastric Cancer. S. NAKAMURA, M. KANDA, D. SHIMIZU, K. SAWAKI, C. TANAKA, N. HATTORI, M. HAYASHI, S. YAMADA, G. NAKAYAMA, K. OMAE, M. KOIKE, Y. KODERA (*Nagoya; Fukushima, Japan*)

Expression Patterns of CD44 and AREG Under Treatment With Selective Tyrosine Kinase Inhibitors in HPV+ and HPV- Squamous Cell Carcinoma. B. KANSY, C. ADERHOLD, L. HUBER, S. LUDWIG, R. BIRK, A. LAMMERT, S. LANG, N. ROTTER, B. KRAMER (*Essen; Mannheim; Marburg, Germany*)

Chromobox 2 Expression Predicts Prognosis After Curative Resection of Oesophageal Squamous Cell Carcinoma. S. UEDA, M. KANDA, Y. SATO, H. BABA, S. NAKAMURA, K. SAWAKI, D. SHIMIZU, S. MOTOYAMA, T. FUJII, Y. KODERA, S. NOMOTO (*Nagoya; Akita; Toyama, Japan*)

Fusion of the Lumican (*LUM*) Gene With the Ubiquitin Specific Peptidase 6 (*USP6*) Gene in an Aneurysmal Bone Cyst Carrying a t(12;17)(q21;p13) Chromosome Translocation. I. PANAGOPOULOS, L. GORUNOVA, K. ANDERSEN, I. LOBMAIER, M. LUND-IVERSEN, F. MICCI, S. HEIM (*Oslo, Norway*)

Influence of Concurrent Mutations on Overall Survival in EGFR-mutated Non-small Cell Lung Cancer. M. CHEVALLIER, P. TSANTOULIS, A. ADDEO, A. FRIEDLAENDER (*Geneva, Switzerland*)

Long Noncoding RNA *ANROC* on the *INK4* Locus Functions to Suppress Cell Proliferation. Y. KOTAKE, T. TSURUDA (*Fukuoka, Japan*)

The KDR (VEGFR-2) Genetic Polymorphism Q472H and c-KIT Polymorphism M541L Are Associated With More Aggressive Behaviour in Astrocytic Gliomas. N. ZAMAN, S.S. DASS, P.D. PARCQ, S. MACMAHON, L. GALLAGHER, L. THOMPSON, J.S. KHORASHAD, C. LIMBÄCK-STANIC (*London, UK*)

KIF15 Expression in Tumor-associated Monocytes Is a Prognostic Biomarker in Hepatocellular Carcinoma. A. KITAGAWA, T. MASUDA, J. TAKAHASHI, T. TOBO, M. NODA, Y. KURODA, Q. HU, Y. KOUYAMA, Y. KOBAYASHI, S. KURAMITSU, K. SATO, A. FUJII, Y. YOSHIKAWA, H. WAKIYAMA, D. SHIMIZU, Y. TSURUDA, H. EGUCHI, Y. DOKI, M. MORI, K. MIMORI (*Oita; Osaka; Fukuoka, Japan*)

Contents

Reviews

Reasons to Reconsider Risk Associated With Power Morcellation of Uterine Fibroids. C. HOLZMANN, W. KIEPKE, B. ROMMEL, B. HELMKE, J. BULLERDIEK (Rostock; Baden Baden-Buehl; Bremen; Stade, Germany) 1

Three-dimensional Versus Two-dimensional Laparoscopic Surgery for Colorectal Cancer: Systematic Review and Meta-analysis. G. PANTALOS, D. PATSOURAS, E. SPARTALIS, D. DIMITROULIS, G. TSOUROUFLIS, N. NITTEAS (Athens, Greece) 11

Intraoperative Indocyanine Green (ICG) Angiography for the Identification of the Parathyroid Glands: Current Evidence and Future Perspectives. E. SPARTALIS, G. NTOKOS, K. GEORGIU, G. ZOGRAFOS, G. TSOUROUFLIS, D. DIMITROULIS, N.I. NITTEAS (Athens, Greece) 23

Experimental Studies

Fancsani Anemia Mouse Genotype-specific Mitigation of Total Body Irradiation by GS-Nitroside JPM-039. M.W. EPPERLY, R. FISHER, X. ZHANG, W. HOU, D. SHIELDS, P. WIPF, H. WANG, S. THERMOZIER, J.S. GREENBERGER (Pittsburgh, PA, USA) 33

Second-generation Probiotics Producing IL-22 Increase Survival of Mice After Total Body Irradiation. X. ZHANG, R. FISHER, W. HOU, D. SHIELDS, M.W. EPPERLY, H. WANG, L. WEI, B.J. LEBOWITZ, J. YU, L.M. ALEXANDER, J.-P. VAN PIJKEREN, S. WATKINS, P. WIPF, J.S. GREENBERGER (Pittsburgh, PA; Madison, WI, USA) 39

The Association of MMP7 Genotype With Pterygium. P.-S. HU, Y.-C. WANG, C.-H. LIAO, N.-Y. HSIA, M.-F. WU, J.-S. YANG, C.-C. YU, W.-S. CHANG, D.-T. BAI, C.-W. TSAI (Taichung; Changhua; Taipei, Taiwan, ROC) 51

Contents continued on the back cover

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General Policy

- **IN VIVO** is a multidisciplinary journal designed to bring together original high quality works and reviews on experimental and clinical biomedical research within the frames of human physiology, pathology and disease management. A special focus of the journal is the publication of works on: (a) Experimental development and application of new diagnostic procedures; (b) Pharmacological and toxicological evaluation of new drugs and drug combinations; (c) Clinical trials; (d) Development and characterization of models of biomedical research.
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● Selection of Recent Articles

Sutureless Surgical Orthotopic Implantation Technique of Primary and Metastatic Cancer in the Liver of Mouse Models. H. NISHINO, H.M. HOLLANDSWORTH, N. SUGISAWA, J. YAMAMOTO, Y. TASHIRO, S. INUBUSHI, K. HAMADA, Y. SUN, H. LIM, S. AMIRFAKHRI, F. FILEMONI, R.M. HOFFMAN, M. BOUVET (San Diego, CA, USA; Kyoto, Japan)

Hip Arthroplasty Following Subtotal Sacrectomy for Chordoma. M.R. CLAXTON, M.B. SHIRLEY, J.D. JOHNSON, K.I. PERRY, P.S. ROSE, M.T. HOUDEK (Rochester, MN, USA)

SMN Protein Contributes to Skeletal Muscle Cell Maturation Via Caspase-3 and Akt Activation. S. ANDO, M. TANAKA, N. CHINEN, S. NAKAMURA, M. SHIMAZAWA, H. HARA (Gifu, Japan)

Comparison of TMA Technique and Routine Whole Slide Analysis in Evaluation of Proliferative Markers Expression in Laryngeal Squamous Cell Cancer. U. CIESIELSKA, A. PIOTROWSKA, C. KOBIERZYCKI, W. PASTUSZEWSKI, M. PODHORSKA-OKOLOW, P. DZIEGIEL, K. NOWINSKA (Wroclaw, Poland; Namsos, Norway)

Leucocyte Count Does Not Improve the Diagnostic Performance of a Diagnostic Score (DS) in Distinguishing Acute Appendicitis (AA) from Nonspecific Abdominal Pain (NSAP). J. MEKLIN, M. ESKELINEN, K. SYRJANEN, M. ESKELINEN (Kuopio; Kaarina, Finland; Barretos, Brazil)

Evaluating the Decision-to-Delivery Interval in Emergency Cesarean Sections and its Impact on Neonatal Outcome. J.-A. BRANDT, B. MORGENSTERN, F. THANGARAJAH, B. GRÜTTNER, S. LUDWIG, C. EICHLER, J. RATIU, P. MALLMANN, D. RATIU (Cologne, Germany)

Cutaneous Stomal Recurrence of Colorectal Cancer After Curative Rectal Cancer Surgery – A Case Report and Systematic Review. S. DAVEY, K. MCCARTHY (Bristol, UK)

Knockout of TRPV1 Exacerbates Ischemia-reperfusion-induced Renal Inflammation and Injury in Obese Mice. B. ZHONG, S. MA, D.H. WANG (East Lansing, MI, USA)

In Vitro and *In Vivo* Biocompatibility Analysis of a New Transparent Collagen-based Wound Membrane for Tissue Regeneration in Different Clinical Indications. O. JUNG, M. RADENKOVIC, S. STOJANOVIĆ, C. LINDNER, M. BATINIC, O. GÖRKE, J. PISSAREK, A. PRÖHL, S. NAJMAN, M. BARBECK (Rostock; Berlin, Germany; Niš, Serbia)

Hepatocellular Carcinoma-associated microRNAs Induced by Hepatoma-derived Growth Factor Stimulation. H. ENOMOTO, H. NAKAMURA, H. NISHIKAWA, T. NISHIMURA, Y. IWATA, S. NISHIGUCHI, H. IJIMA (Hyogo; Osaka, Japan)

An Improved Encapsulation Method for Cryopreserving Hepatocytes for Functional Transplantation Using a Thermo-reversible Gelation Polymer. K. YAMADA, T. AOKI, Y. ENAMI, Y. TASHIRO, Z. ZEHAOU, T. KOIZUMI, T. KUSANO, K. MATSUDA, Y. WADA, H. SHIBATA, K. TOMIOKA, K. SIRIRATSIVAWONG, R.M. HOFFMAN, M. MURAKAMI (Tokyo, Japan; San Diego, CA, USA)

The Efficacy and Safety of Trifluridine/Tipiracil Treatment for Elderly Patients With Metastatic Colorectal Cancer in a Real-world Setting. M. SHIBUTANI, W. EN, Y. OKAZAKI, S. KASHIWAGI, T. FUKUOKA, Y. ISEKI, K. HIRAKAWA, M. OHIRA (<i>Osaka, Japan</i>).....	6211
Comparison of Treatment Completion Rate Between Conventional and Dose-dense Doxorubicin and Cyclophosphamide (AC) Followed by a Taxane in Patients With Breast Cancer: A Propensity Score-matched Analysis. K. MAMISHIN, Y. NAITO, S. NOMURA, G. OGAWA, K. NIGUMA, K. BABA, S. SAKAEDA, H. NAKAJIMA, S. KUSUHARA, C. FUNASAKA, T. NAKAO, Y. FUKASAWA, C. KONDOH, K. HARANO, T. KOGAWA, N. MATSUBARA, A. HOSONO, T. KAWASAKI, T. MUKOHARA (<i>Kashiwa; Tokyo, Japan</i>) ...	6217
Immune Checkpoint Inhibitor as a Therapeutic Choice for Double Cancer: A Case Series. H. AOKI, N. MATSUMOTO, H. TAKAHASHI, M. HONDA, T. KANEKO, S. ARIMA, T. ISHII, T. MIZUTANI, R. MASUZAKI, K. NIREI, H. YAMAGAMI, M. OGAWA, T. KANDA, M. MORIYAMA, K. MIURA (<i>Tokyo, Japan</i>).....	6225
Effect of Monoammonium Glycyrhizinate on the Development of Hepatotoxicity After Initial Intrathecal Chemotherapy for Leukemia. K. KISHIMOTO, D. HASEGAWA, S. UEMURA, S. NAKAMURA, A. KOZAKI, A. SAITO, T. ISHIDA, T. MORI, Y. KOSAKA (<i>Kobe, Japan</i>).....	6231
Prospective Randomized Trial of Early Postoperative Enteral and Total Parenteral Nutrition for Treating Esophageal Cancer. Y. HAMAI, J. HIHARA, M. EMI, Y. IBUKI, T. KUROKAWA, T. YOSHIKAWA, R. HIROHATA, M. OHSAWA, N. KITASAKI, M. OKADA (<i>Hiroshima, Japan</i>).....	6237
Phase II Study of Preoperative Chemoradiotherapy With S-1 Plus Oxaliplatin for Locally Advanced Rectal Cancer (PerSeUS-RC01). N. MATSUHASHI, T. TAKAHASHI, C. TANAKA, K. YAWATA, M. YAMADA, Y. IWATA, S. KIYAMA, C. MIZUTANI, J.Y. TAJIMA, T. ISHIHARA, K. YOSHIDA (<i>Gifu, Japan</i>).....	6247
Efficacy and Safety of Induction Chemotherapy and/or External Beam Radiotherapy Followed by Brachytherapy in Patients With Tongue Cancer. R.-I. YOSHIMURA, K. TODA, H. WATANABE, A. KAIDA, H. HARADA, T. ASAKAGE, M. MIURA (<i>Tokyo, Japan</i>).....	6259
CTLA-4 Expression in Tumor-infiltrating Lymphocytes Is Irrelevant to PD-L1 Expression in NSCLC. S. MUTO, S. INOMATA, H. YAMAGUCHI, H. MINE, H. TAKAGI, Y. OZAKI, M. WATANABE, T. INOUE, T. YAMAURA, M. FUKUHARA, N. OKABE, Y. MATSUMURA, T. HASEGAWA, J. OSUGI, M. HOSHINO, M. HIGUCHI, Y. SHIO, H. SUZUKI (<i>Fukushima, Japan</i>).....	6267
Doxorubicin Combined With Ifosfamide for Sarcoma Induces Muscle Atrophy and Sleep Disruption. H. KINOSHITA, Y. HAGIWARA, T. ISHII, H. KAMODA, T. TSUKANISHI, S. OHTORI, T. YONEMOTO (<i>Chiba, Japan</i>).....	6273
Effect of Concomitant Lafutidine on Adjuvant S-1 for Head and Neck Cancer: A Comparative Study. K. YOSHINO, I. OKAMOTO, H. SATO, T. OKADA, K. TOKASHIKI, T. KONDO, K. TSUKAHARA (<i>Tokyo, Japan</i>).....	6279
Applicability of the Histoculture Drug Response Assay to Predict Platinum Sensitivity and Prognosis in Ovarian Cancer. J. LEE, J.M. KIM, Y.H. LEE, G.O. CHONG, D.G. HONG (<i>Daegu, Republic of Korea</i>).....	6287
Index.....	6293

Combinational Anti-tumor Effects of Chemicals from <i>Paeonia lutea</i> Leaf Extract in Oral Squamous Cell Carcinoma Cells. S. NAKAMURA, Y. MUKUDAI, J. CHIKUDA, M. ZHANG, H. SHIGEMORI, K. YAZAWA, S. KONDO, T. SHIMANE, T. SHIROTA (<i>Tokyo; Fukuoka; Tsukuba, Japan</i>)	6077
A DNA Topoisomerase II Inhibitor Results in <i>Ex Vivo</i> Differentiation of THP-1 Cells and Activation of Dendritic Cells. Y.J. CHO, H. LEE, J. KIM, G. GONG, H.J. LEE, I.A. PARK (<i>Seoul, Republic of Korea</i>)	6087
Cetyltrimethylammonium Bromide Disrupts Mesenchymal Characteristics of Human Tongue Squamous Cell Carcinoma SCC4 Cells Through Modulating Canonical TGF- β /Smad/miR-181b/TIMP3 Signaling Pathway. C.-H. YUE, C.-H. CHEN, Y.-R. PAN, Y.-P. CHEN, F.-M. HUANG, C.-J. LEE (<i>Taichung; Miaoli; Changhua, Taiwan, ROC</i>)	6095
Differences in Transport Characteristics and Cytotoxicity of Epirubicin and Doxorubicin in HepG2 and A549 Cells. K. NAGAI, S. FUKUNO, M. SHIOTA, M. TAMURA, S. YABUMOTO, H. KONISHI (<i>Tondabayashi, Japan</i>)	6105
Suppressive Effects of Anisomycin on the Proliferation of B16 Mouse Melanoma Cells <i>In Vitro</i> . H. USHIJIMA, R. MONZAKI, A. ONODERA (<i>Iwate, Japan</i>)	6113
Flaccidoxide Induces Apoptosis Through Down-regulation of PI3K/AKT/mTOR/p70S6K Signaling in Human Bladder Cancer Cells. B.-S. WONG, W.-T. WU, J.-H. SU, Y.-G. GOAN, Y.-J. WU (<i>Pingtung; Kaohsiung, Taiwan, ROC</i>)	6123
RGS2 Suppresses Melanoma Growth <i>via</i> Inhibiting MAPK and AKT Signaling Pathways. S.-J. LIN, Y.-C. HUANG, H.-Y. CHEN, J.-Y. FANG, S.-Y. HSU, H.-Y. SHIH, Y.-C. LIU, Y.-C. CHENG (<i>Taoyuan, Taiwan, ROC</i>)	6135
Pongamol Inhibits Epithelial to Mesenchymal Transition Through Suppression of FAK/Akt-mTOR Signaling. H.E. PUTRI, B. SRITULARAK, P. CHANVORACHOTE (<i>Bangkok, Thailand</i>)	6147
Sequence of CX-4945 and Cisplatin Administration Determines the Effectiveness of Drug Combination and Cellular Response in Cholangiocarcinoma Cells <i>In Vitro</i> . J. LERTSUWAN, A. SAWASDICHAI, N. TASNAWIJITWONG, K. GASTON, P.-S. JAYARAMAN, J. SATAYAVIVAD (<i>Bangkok, Thailand; Nottingham, UK</i>)	6155
miR-3188 Enhances Sensitivity of Breast Cancer Cells to Ionizing Radiation by Down-regulating Rictor. S.-E. HONG, H.-O. JIN, S.-M. KIM, S.-K. JANG, C.S. PARK, M.-K. SEONG, H.-A. KIM, W.C. NOH, I.-C. PARK (<i>Seoul, Republic of Korea</i>)	6169
Sensitization Effects of Repurposed Blood Pressure-regulating Drugs on Drug-resistant Cancer Cells. C. JIANG, T. ZHENG, J.H. PARK, J.-S. LEE, Y. OH, A. KUNDU, H.S. KIM, S. YOON (<i>Suwon, Republic of Korea</i>)	6179
<i>Clinical Studies</i>	
A Case of Rare Matrix-producing Triple-negative Breast Carcinoma for Which Drug Response in a Patient-derived Orthotopic Xenograft Mouse Model Was Correlated With Patient Response. T. NOMURA, J. KUREBAYASHI, T. MORIYA, W. SAITO, T. MURATA, J. YAMAMOTO, C. HOZUMI, R.M. HOFFMAN (<i>Kurashiki, Japan; San Diego, CA, USA; Narita, Japan</i>)	6191
A Multicentre Retrospective Study of Nivolumab Plus Ipilimumab for Untreated Metastatic Renal Cell Carcinoma. R. KATO, T. KOJIMA, T. SAZUKA, H. YAMAMOTO, S. FUKUDA, K. YAMANA, Y. SUGINO, S. HAMAMOTO, N. NAKAIGAWA, K. KABU, H. MURAKAMI, W. OBARA (<i>Iwate; Ibaraki; Chiba; Aomori; Tokyo; Niigata; Mie; Nagoya; Kanagawa, Japan</i>)	6199

Activity in MCF-7 Estrogen-Sensitive Breast Cancer Cells of Capsicodendrin from <i>Cinnamosma fragrans</i> . U.M. ACUNA, N. EZZONE, L.H. RAKOTONDRAIBE, E.J. CARCACHE DE BLANCO (<i>Columbus, OH, USA</i>)	5935
Harnessing the Power of Kiwifruit for Radiosensitization of Melanoma. L. KOU, Z. ZHU, E. FAJARDO, Q. BAI, C. REDINGTON, H. XIAO, M. LEQUIO, N. SHAM, M.R. WAKEFIELD, Y. FANG (<i>St. Petersburg, FL; Des Moines, IA; Columbia, MO, USA</i>)	5945
Reduction in Copper Uptake and Inhibition of Prostate Cancer Cell Proliferation by Novel Steroid-based Compounds. F. XIE, F. PENG (<i>Dallas, TX, USA</i>)	5953
GLO 1 and PKC λ Regulate ALDH1-positive Breast Cancer Stem Cell Survival. H. MOTOMURA, S. TAMORI, M.-A. YATANI, A. NAMIKI, C. ONAGA, A. OZAKI, R. TAKASAWA, Y. MANO, T. SATO, Y. HARA, K. SATO, Y. XIONG, Y. HARADA, T. HANAWA, S.-I. TANUMA, K. SASAKI, S. OHNO, K. AKIMOTO (<i>Chiba; Tokyo, Japan</i>)	5959
The Novel, Orally Bioavailable CDK9 Inhibitor Atuveciclib Sensitises Pancreatic Cancer Cells to TRAIL-induced Cell Death. J.-P. RUFF, A.-L. KRETZ, M. KORNMANN, D. HENNE-BRUNS, J. LEMKE, B. TRAUB (<i>Ulm, Germany</i>)	5973
The Dual Histone Deacetylase-Proteasome Inhibitor RTS-V5 Acts Synergistically With Ritonavir to Induce Endoplasmic Reticulum Stress in Bladder Cancer Cells. K. OKUBO, N. REBING, W.A. SCHULZ, F.K. HANSEN, T. ASANO, A. SATO (<i>Tokorozawa, Japan; Bonn; Duesseldorf, Germany</i>)	5987
Effects of Rhenium(I)-diselenoether and of its Diselenide Ligand on the Production of Cathepsins B and S by MDA-MB231 Breast Malignant Cells. P. COLLERY, V. VEENA, D. DESMAËLE, A. HARIKRISHNAN, B. LAKSHMI (<i>Algajola; Châtenay-Malabry, France; Bangalore; Chennai, India</i>)	5997
Intravenous Administration of Dehydroxymethylepoxyquinomicin With Polymer Enhances the Inhibition of Pancreatic Carcinoma Growth in Mice. H. FUJISAKI, Y. NAKANO, S. MATSUDA, K. SUZUKI, O. ITANO, M. TANAKA, S. HORI, Y. HASEGAWA, Y. ABE, H. YAGI, M. KITAGO, T. KONNO, K. ISHIHARA, K. OHNO, S. KISHINO, K. UMEZAWA, Y. KITAGAWA (<i>Kanagawa; Tokyo; Tochigi; Chiba; Aichi, Japan</i>)	6003
The Thioredoxin-1 Inhibitor, PX-12, Suppresses Local Osteosarcoma Progression. H. KINOSHITA, O. SHIMOZATO, T. ISHII, H. KAMODA, Y. HAGIWARA, S. OHTORI, T. YONEMOTO (<i>Chiba, Japan</i>)	6013
Metabolic Alteration in Cancer Cells by Therapeutic Carbon Ions. N. OSU, H. MAKINOSHIMA, T. OIKE, T. OHNO (<i>Maebashi; Tsuruoka; Kashiwa, Japan</i>)	6023
Characteristics of Human Peripheral Blood $\gamma\delta$ T Cells Expanded With Zoledronate. M. KIM, H. KIM, M. HAN, H.J. HWANG, H. KIM, H.J. IM, N. KIM, K.-N. KOH (<i>Seoul, Republic of Korea</i>)	6031
Efficacy of a Novel Oral Chemotherapeutic Agent, TAS-102, Against Human Oral Squamous Cell Carcinoma Cells. K. HARADA, T. FERDOUS, K. MISHIMA (<i>Ube, Japan</i>)	6039
Evaluation of Erastin as a Therapeutic Agent Under Hypoxic Conditions in Pancreatic Cancer Cells. S. OWADA, H. ENDO, Y. SHIDA, T. KINOUE, H. FURUYA, M. TATEMACHI (<i>Isehara, Japan</i>)	6051
Activity of Free and Liposomal Antimony Trioxide in the Acute Promyelocytic Leukemia Cell Line NB4. A. ROSSATO VIANA, C. BORDIN DAVIDSON, B. SALLES, L. YAMAMOTO DE ALMEIDA, A. KRAUSE, C.A. BIZZI, E. MAGALHAES REGO, L.M. FONTANARI KRAUSE, S.R. MORTARI (<i>Santa Maria, RS; Ribeirão Preto; São Paulo, SP, Brasil</i>)	6061
Pharmacological Strategy for Selective Targeting of Glioblastoma by Redox-active Combination Drug – Comparison With the Chemotherapeutic Standard-of-care Temozolomide. A. SUMIYOSHI, S. SHIBATA, Z. ZHELEV, T. MILLER, D. LAZAROVA, G. ZLATEVA, I. AOKI, R. BAKALOVA (<i>Chiba, Japan; Stara Zagora; Sofia, Bulgaria; San Diego, CA, USA</i>)	6067