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DNA Damage and Repair in Cancer

Guest Editors:

Prof. Dr. Srinivasan Madhusudan

Professor of Medical Oncology, Translational DNA Repair Group, Nottingham Biodiscovery Institute, School of Medicine, Nottingham University Hospitals, University Park, Nottingham NG7 3RD, UK

Dr. Gianluca Tell

Laboratory of Molecular Biology and DNA repair, Department of Medicine, University of Udine, 33100 Udine, Italy

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Message from the Guest Editors

Genomic DNA is threaten by DNA-damaging agents. For genomic stability, mammalian cells have DNA damagesensor mechanisms that induce; apoptosis to eliminate damaged cells; transcriptional response; DNA damage tolerance; activation of DNA damage checkpoints and modulation of cell cycle progression; and/or initiation of DNA repair. To maintain genomic stability, cells detect and repair DNA damage by multiple pathways.

Failure of response to DNA damage can lead to the accumulation of mutagenic lesions associated with ageing, neurological disorders, and cancer. Furthermore, germline mutations in DNA repair genes lead to cancer predisposing syndromes.

Pharmacological targeting of DNA repair can increase the efficacy of treatments and overcome therapeutic resistance. Synthetic lethality, the loss of function of either one of two inter-related genes is not lethal, but loss of both genes results in cell death, provides an exciting new platform for pharmacological targeting of DNA repair.

In this Special Issue, we focus on recent advances in molecular biology of DNA repair, as well as its relevance as a key prognostic, predictive, and therapeutic targets in cancer.













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Editor-in-Chief

Prof. Dr. Selvarangan Ponnazhagan

Department of Pathology, The University of Alabama at Birmingham, 1825 University Blvd, SHEL 814, Birmingham, AL 35294-2182, USA

Message from the Editor-in-Chief

Genes are central to our understanding of biology, and modern advances such as genomics and genome editing have maintained genetics as a vibrant, diverse and fastmoving field. There is a need for good quality, open access journals in this area, and the *Genes* team aims to provide expert manuscript handling, serious peer review, and rapid publication across the whole discipline of genetics. Starting in 2010, the journal is now well established and recognised.

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