Special Issue

Viral Vectors Paired with Genome-Editing Tools: Evolving Technologies for the Treatment of Neurodegenerative Diseases and Disorders

Message from the Guest Editors

Diseases and disorders of the central nervous system (CNS) have historically been among the most difficult to combat using conventional pharmacological strategies. In this chapter, we describe the various tools available for genome editing and summarize in vitro and in vivo preclinical studies of CNS genome editing. We discuss the future progress of these technologies, as well as the current limitations and alternative approaches to overcoming some of their bottlenecks. We will aim here to devote significant attention to viral vectors derived from human immunodeficiency virus type 1 (lentiviral vectors; LVs) and adeno-associated virus (AAVs). The high interest in these viral delivery systems vectors is due to: (i) robust delivery and long-lasting expression; (ii) efficient transduction into postmitotic cells, including the brain; (iii) low immunogenicity and toxicity; and (iv) compatibility with advanced manufacturing techniques. Finally, this chapter will summarize the current therapeutic application and novel approaches of advance genome-editing technology for the prevention and treatment of neurodegenerative diseases (NDDs).

Guest Editors

Dr. Boris Kantor

Director of Viral Vector Core, Department of Neurobiology, Duke University School of Medicine, Durham, NC 27710, USA

Prof. Dr. Ornit Chiba-Falek

Division of Translational Brain Sciences, Department of Neurology & Pathology, and the Center for Genomic and Computational Biology, Duke University Medical Center, Durham, NC 27710, USA

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Cells
Editorial Office
MDPI, Grosspeteranlage 5
4052 Basel, Switzerland
Tel: +41 61 683 77 34
cells@mdpi.com

mdpi.com/journal/cells





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