Special Issue

Advance in the Mechanism and Treatment of Lysosomal Storage Disorders

Message from the Guest Editor

Lysosomal storage disorders (LSDs) are inherited metabolic diseases usually caused by the deficiency of an enzyme, resulting in substrate accumulation. The advent of enzyme replacement therapy (ERT) in the 1990s changed the natural history of Gaucher disease, the first treated LSD. Nowadays, many other ERTs are commercially available, but the morbidity and mortality of LSDs are still high. We invite researchers to contribute original studies, as well as review articles, addressing recent advances in the treatment of LSDs. Your ground-breaking research will contribute to the improvement of patients' lives in the near future

Guest Editor

Dr. Annalisa Sechi

Regional Coordinating Centre for Rare Diseases, University Hospital of Udine, 33100 Udine, Italy

Deadline for manuscript submissions

closed (15 August 2022)



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

mdpi.com/journal/ life





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Message from the Editor-in-Chief

Life (ISSN 2075-1729) is an international, peer-reviewed open access journal that publishes scientific studies related to fundamental themes in life sciences. Some papers are published individually, while others are submitted for inclusion in special issues with guest editors. You are invited to contribute a research article, essay, or a review to be considered for publication.

Editor-in-Chief

Prof. Dr. Lluís Ribas de Pouplana

Institute for Research in Biomedicine (IRB Barcelona), The Barcelona Institute of Science and Technology, 08028 Barcelona, Spain

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