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# Clinical, Functional and Fitness Consequences of Genetic Variants

Guest Editor:

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Deadline for manuscript submissions:

closed (20 April 2024)

## **Message from the Guest Editor**

Dear Colleagues,

A major bottleneck in interpreting genetic data is estimating the impact of each single variant. Genome-wide association studies (GWASs) and family segregation studies (FSSs) may associate diseases with single variants or with genes. Experimental studies may associate variants with the loss or gain of protein function and indicate alterations in specific biological processes. Computational studies may use evolutionary and structural information to suggest variant effects on fitness and mechanistic insights, respectively. These studies improve our understanding of variant consequences and help in interpreting variants that are currently described as variants of unknown significance (VUS).

In this Special Issue, we welcome the submission of reviews, original articles, and short reports that cover any aspects of variant consequences. These may include works focused on specific genes, biological functions, phenotypes, or samples.

Dr. Panagiotis Katsonis *Guest Editor* 













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## **Message from the Editor-in-Chief**

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