

Special Issue

Personalized Medicine in Clinical Practice

Message from the Guest Editors

This Special Issue of the *Journal of Personalized Medicine* aims to highlight the current state of the utilization (or implementation) of genomic technologies (including family history) in various clinical practices and showcase innovative and novel approaches to integrating genomic technologies into practice. In particular, we are interested in examples of innovative clinical programs and evidence of provider utilization, patient acceptance, cost-effectiveness, and/or feasibility. Papers may address use of telemedicine, as well as other aspects of digital health to solve challenges in resource-limited settings (e.g., limited access to geneticists, genetic counselors), molecular review boards, team-based care, and other clinical delivery approaches. Case reports, reviews, and original articles are welcome.

Guest Editors

Dr. Susanne B. Haga

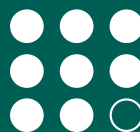
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closed (31 August 2020)



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About the Journal

Message from the Editor-in-Chief

Journal of Personalized Medicine (JPM), ISSN 2075-4426) is an international, open access journal aimed at bringing all aspects of personalized medicine to one platform. *JPM* publishes cutting edge, innovative preclinical and translational scientific research and technologies related to personalized medicine (e.g., precision medicine, pharmacogenomics/proteomics, systems biology, 'omics association analysis). *JPM* is covered in Scopus, the Science Citation Index Expanded (SCIE), PubMed, PMC, Embase, and other databases.

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Author Benefits

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Rapid Publication:

manuscripts are peer-reviewed and a first decision is provided to authors approximately 21.5 days after submission; acceptance to publication is undertaken in 3.5 days (median values for papers published in this journal in the first half of 2025).