**Journal of Rare Cardiovascular Diseases in EuroPub database**

**Dear Readers,**

It is my great pleasure to inform you that the *Journal of Rare Cardiovascular Diseases (JRCD)* has been accepted for inclusion in EuroPub (https://europub.co.uk). This comprehensive and multipurpose database covers scholarly literature from all over the world and indexes articles from highly active and authoritative journals. As such, it may greatly facilitate research by considerably reducing the time of data search and review.

In addition to EuroPub, *JRCD* is indexed in multiple other databases including Scopus, Directory of Open Access Journals (DOAJ), Arianta, POL-index, Index Copernicus, JournalTOCs, Google Scholar, PKP Index, Polish Medical Bibliography, Free Medical Journals – Geneva Foundation for Medical Education and Research, as well as J-Gate. Please also note that, as of this year, manuscripts published in *JRCD* are granted 20 points by the Polish Ministry of Science and Higher Education (formerly 8 points).

In view of these successes and to ensure the high quality of published papers, the editorial board has decided to publish the journal twice a year, starting in 2020.

The current issue of the journal contains a summary of papers published in *JRCD* in 2018 and 2019. It also features an original article entitled “Survival analysis of time to develop cardiovascular complications and its predictors among hypertensive patients treated in the Ayder Comprehensive Specialized Hospital, Ethiopia: a retrospective cohort study (RCD code: VIII)”. The study was performed in 578 patients with hypertension, 25.4% of whom developed cardiovascular complications within the 5-year follow-up, which translates to an incidence rate of over 8 per 1000 persons per month. The predictors of these complications included baseline blood pressure, baseline cardiovascular complications, age, and proteinuria. This study is particularly valuable because it adds to the current knowledge on the African population of hypertensive patients, which has been much less well described than European or North American populations. Finally, this issue contains an editorial on amyloid cardiomyopathy, an interesting case report on autoimmune hepatitis induced by bosentan in a patient with pulmonary arterial hypertension, and another case report on a rare association of congenital unicuspid aortic valve with left ventricular noncompaction.

On behalf of the editors and the publisher, I would like to thank you for the ongoing support of our journal. I hope that you will find this issue of JRCD interesting.

We hope that our readers will find this issue interesting and that it will update the knowledge of health care providers and scientists related to the field of rare cardiovascular diseases and disorders.

Piotr Podolec  
Editor-in-Chief  
Journal of Rare Cardiovascular Diseases