

Guest Editorial for Selected Papers From BIOKDD 2021

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THE 20th International Workshop on Data Mining in Bioinformatics (BIOKDD 2021) was held virtually on August 15, 2021 due to the COVID-19 pandemic. BIKDD 2021 featured the special theme of "Artificial Intelligence in Medicine" which particularly welcomed paper submissions and invited talks related to the use of machine learning and data mining techniques for the analysis of large amounts of heterogeneous complex biological and medical data, with a particular focus on deep learning methods that see fast advancement and wider adoption in Bioinformatics. As a whole-day workshop, BIKDD 2021 accepted 9 submissions for oral presentation, and has 7 additional invited talks by domain experts. These presentations were divided into 4 sessions: (1) Structural Bioinformatics, (2) Clinical Informatics, (3) Bioinformatics, and (4) Network Biology and Machine Learning.

This special section of TCBB features the extended versions of 5 quality papers presented in BIKDD 2021. Each of the 5 invited papers was reviewed by 3 reviewers invited by the TCBB guest editors, and the BIKDD workshop reviews were shared with the TCBB reviewers. The papers also went through 1 to 2 rounds of revisions.

The first invited paper, "A Knowledge Graph-Enhanced Tensor Factorisation Model for Discovering Drug Target," by Cheng Ye, Rowan Swiers, Stephen Bonner, and Ian Barrett explores the use of machine learning classification algorithms and both matrix and tensor factorisation techniques to predict the clinical outcomes of unseen gene target-disease pairs. A 3D data tensor was created consisting of 1048 gene targets, 860 diseases and 230,011 evidence attributes and clinical outcomes connecting them, using data extracted from the Open Targets and PharmaProjects databases. The data is enriched with gene target representations learned from a drug discovery-oriented knowledge graph. Their results show that incorporating knowledge graph embeddings significantly improves the prediction accuracy and that training tensor

factorisation alongside a dense neural network outperforms all other baselines.

The second invited paper, "MuCoMiD: A Multitask Graph Convolutional Learning Framework for miRNA-Disease Association Prediction," by Ngan Dong, Stefanie Mücke, and Megha Khosla studies the use of a multitask graph convolutional learning framework called MUCOMID for the problem of predicting miRNA-disease associations. Their approach allows automatic feature extraction while incorporating knowledge from five heterogeneous biological information sources in a multitask setting: associations between miRNAs/diseases and protein-coding genes (PCGs), interactions between protein-coding genes, miRNA family information, and disease ontology. Incorporating multiple sources of information helps compensate for the lack of information in any single source and, at the same time, enables the model to generate predictions for any new miRNA or disease. Their model can be employed in both transductive and inductive settings.

The third invited paper, "Heterogeneous Multi-Task Learning with Expert Diversity," by Raquel Aoki, Frederick Tung, and Gabriel L. Oliveira predicts multiple heterogeneous biological and medical targets simultaneously using multi-task learning (MTL). Their model, Multi-gate Mixture-of-Experts with Exclusivity (MMoEEx), optimizes multiple tasks with different characteristics by inducing more diversity among experts, thus creating representations more suitable for highly imbalanced and heterogeneous MTL learning. This is realized with two mechanisms, exclusion and exclusivity, under which some experts only contribute to some tasks, while other experts are shared among all tasks. A two-step optimization approach inspired by MAML is also used to balance the tasks at the gradient level. The approach is validated on three MTL benchmark datasets, including UCI-Census-income dataset, Medical Information Mart for Intensive Care (MIMIC-III) and PubChem BioAssay (PCBA).

The fourth invited paper, "Biocode: A Data-Driven Procedure to Learn the Growth of Biological Networks," by Emre Sefer proposes Biocode, a framework to automatically discover novel biological growth models matching user-specified graph attributes in directed and undirected biological graphs. Such probabilistic biological network growth models have been utilized for tasks such as capturing mechanism and dynamics of biological growth activities, and capturing anomalies. Biocode designs a basic set of instructions which are common enough to model a number of well-known biological graph growth models such as Kronecker model, preferential attachment model, and duplication-based model. These instruction-wise representation are combined with a genetic

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algorithm based optimization procedure to encode models for various biological networks. The performance of Biocode has been evaluated in discovering models for biological collaboration networks, gene regulatory networks, and protein interaction networks with features such as assortativity, clustering coefficient, degree distribution closely match with the true ones in the corresponding real biological networks.

The fifth invited paper, "Finding Overlapping Rmaps via Clustering," by Kingshuk Mukherjee, Daniel Dole-Muinos, Massimiliano Rossi, Ayomide Ajayi, Mattia Prosperi, and Christina Boucher considers the context of optical mapping which is a method for creating high resolution restriction maps of an entire genome. Optical mapping first produces single molecule restriction maps, called Rmaps, which are assembled to generate genome wide optical maps. This work develops a method, called OMCLUST, for finding overlapping Rmaps that uses Gaussian mixture model clustering, and does not require any quantization. Their work demonstrates that OMCLUST not only achieves the highest precision and was more efficient than competing methods, but can also be integrated into the error correction methods to improve their quality of error correction. OMCLUST may serve as a filtering step for finding related Rmaps for error correction, assembly or other applications of optical mapping data.

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