## Understanding the Sources of Performance in Deep Drug Response Models

Nikhil Branson Digital Environment Research Institute Queen Mary University of London Pedro R. Cutillas Barts Cancer Institute Queen Mary University of London

Conrad Bessant Digital Environment Research Institute Queen Mary University of London c.bessant@qmul.ac.uk

## Abstract

Anti-cancer drug response prediction (DRP) using cancer cell lines plays a vital role in stratified medicine and drug discovery. Recently there has been a surge of new deep learning (DL) models for DRP that improve on the performance of their predecessors. However, different models use different input data types and neural network architectures making it hard to find the source of these improvements. Here we consider multiple published DRP models that report state-of-the-art performance in predicting continuous drug response values. These models take the chemical structures of drugs and omics profiles of cell lines as input. By experimenting with these models and comparing with our own simple benchmarks we show that no performance comes from drug features, instead, performance is due to the transcriptomics cell line profiles. Furthermore, we show that, depending on the testing type, much of the current reported performance is a property of the training target values. See full paper at https://www.biorxiv.org/content/10.1101/2024.06.05.597337v1?versioned=true.