

Stacked kinship CNN vs. GBLUP for genomic predictions of additive and complex continuous phenotypes

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Supplementary Figure S3: example of a training trajectory of the DNN (deep neural network) model. On the x-axis the training epochs. On the y-axis the metric of choice. Top panel: loss function (Mean Squared Error) used by the optimizer to tune the network. Bottom panel: Pearson correlation. Metrics are computed either on the training set (blue line) or the validation set (orange line).

