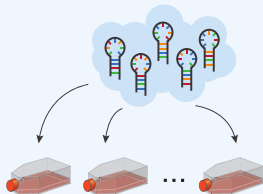
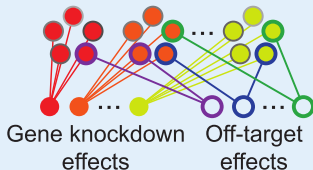


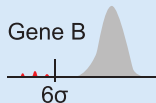
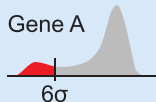
501 genome-scale RNAi loss-of-function screens



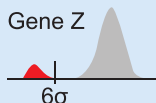
Removal of shRNA off-target effects



Identification of strong differential dependencies

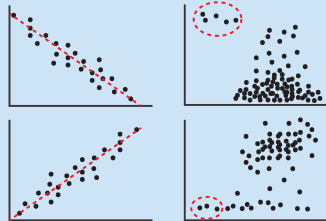


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Models predicting gene dependency from genomic features

Gene dependency



Genomic biomarker

