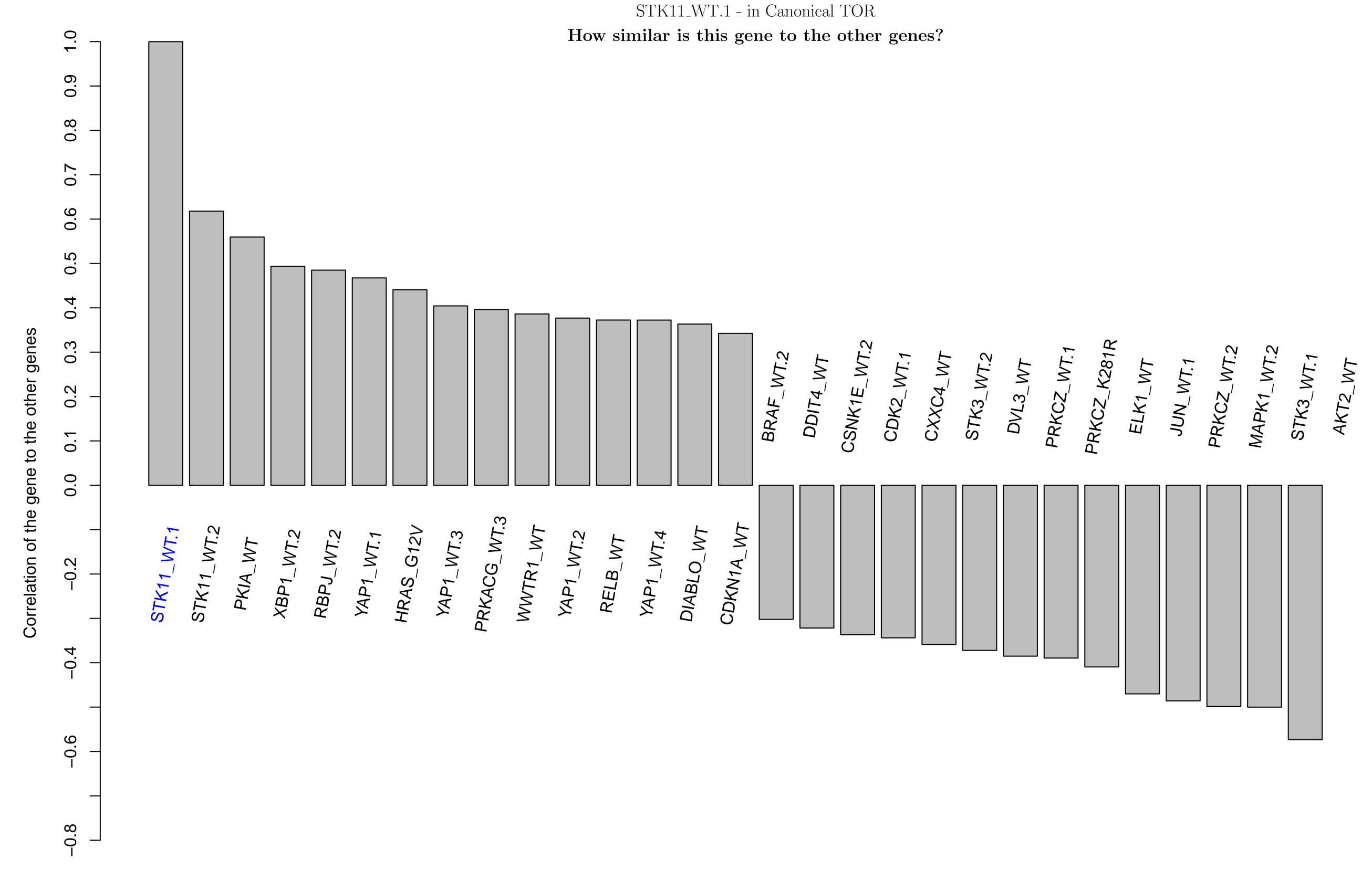
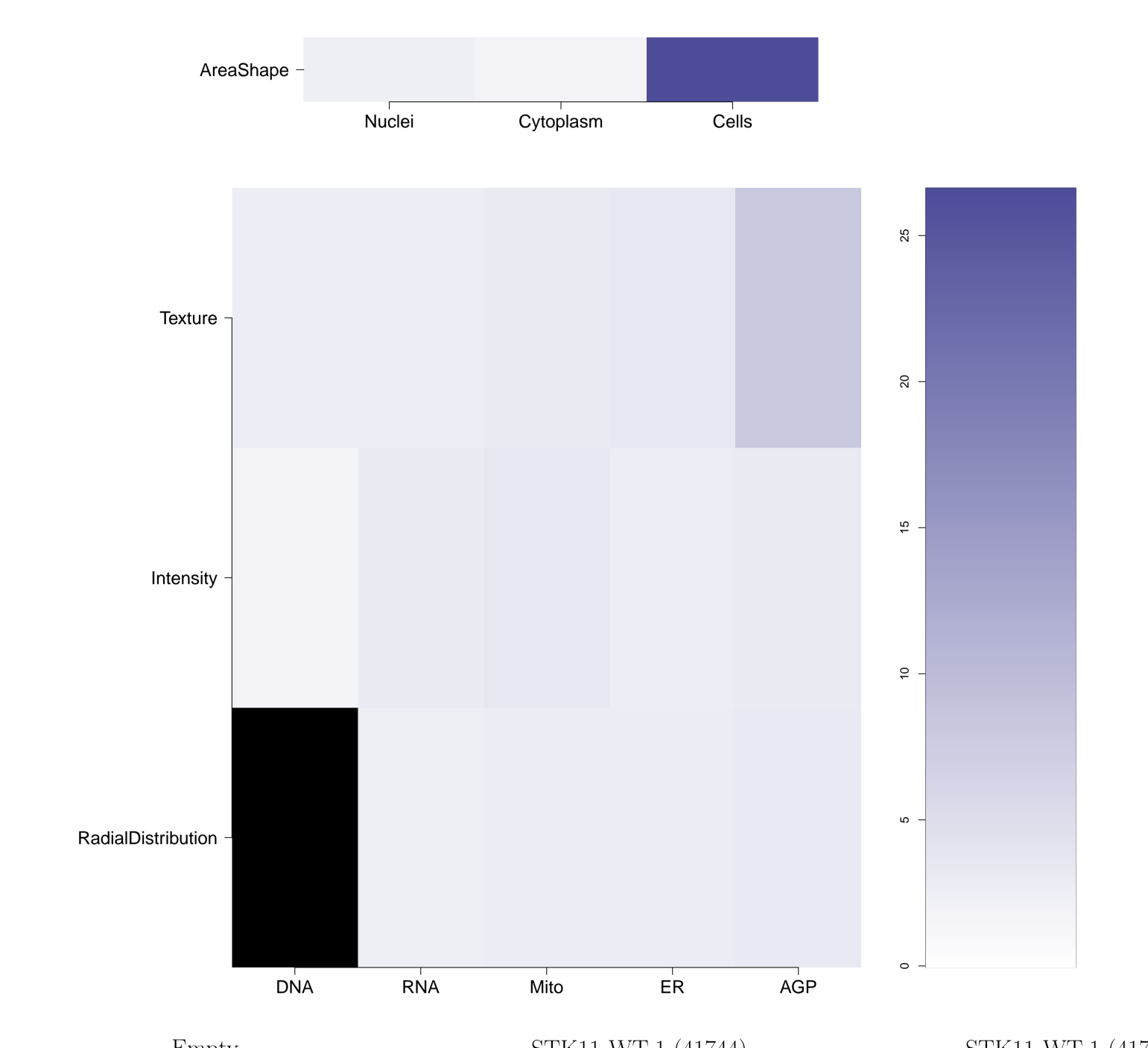
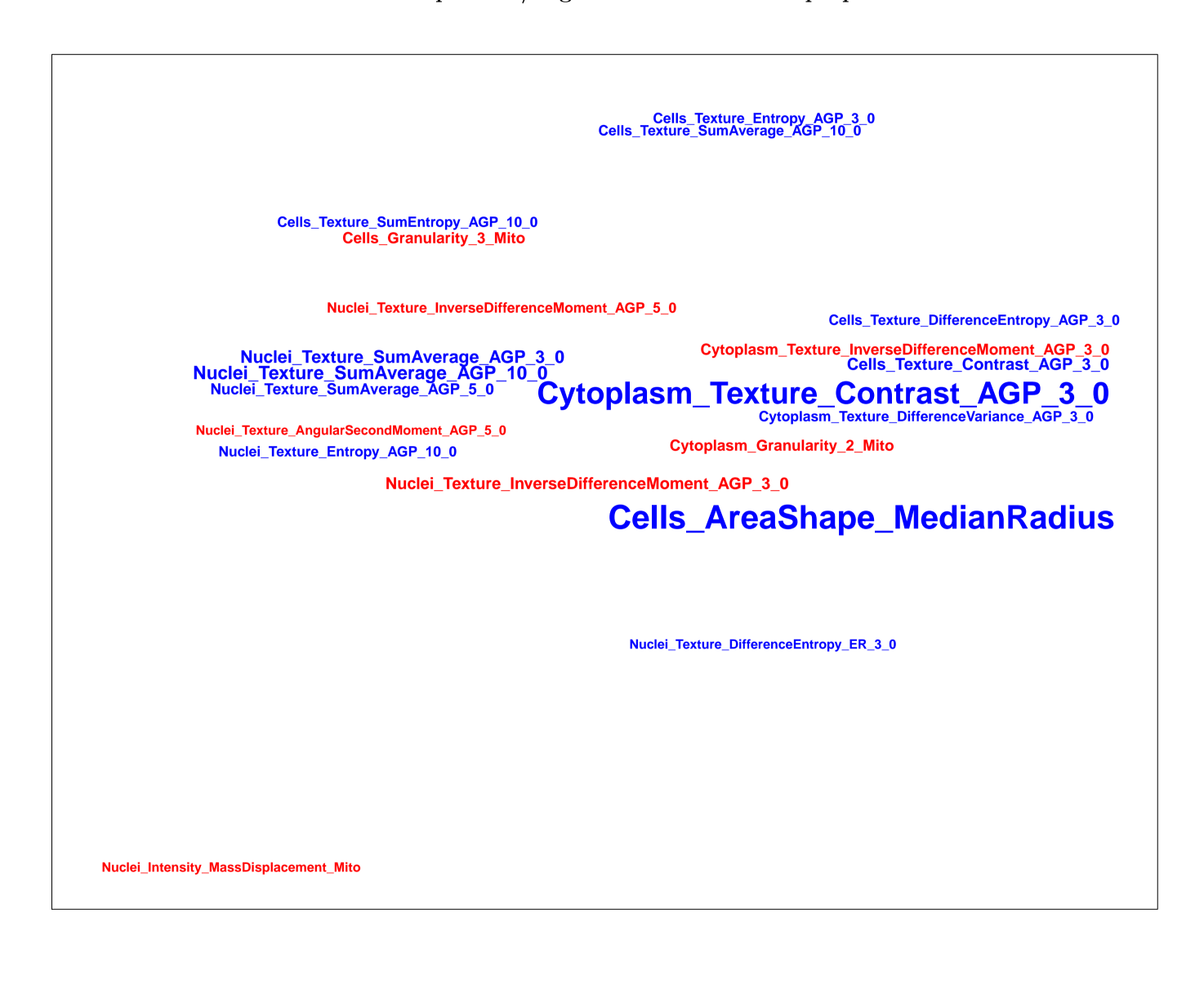
CONFIDENTIAL, contact the Imaging Platform to collaborate on the findings herein

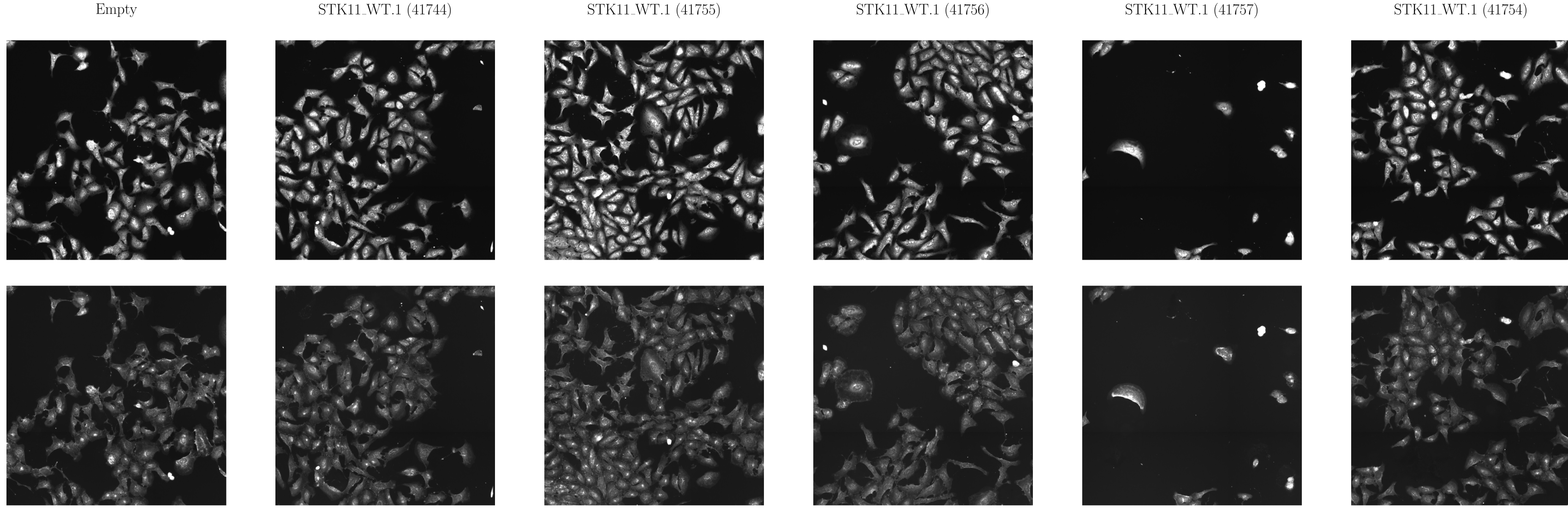


What groups of morphological features are distinguishing in the cluster relative to the untreated samples? (maximum of absolute m-score for the features belonging to the same category; m-score defined as median of a feature z-score across genes in the cluster) Black means no feature is available in the category



Which individual morphological features are distinguishing in the gene relative to the untreated samples? Blue/Red means the feature has a positive/negative z-score. Size is proportional to the z-score value.





Compound IDs and common names (where available); blue/red colored box means the matching compound is positively/negatively correlated with the cluster	Chemical structure co	Mean pairwise replicates correlation of the compound signature (95th DMSO replicate correlation is 0.52)	Correlation between compound the gene		How similar is the compound signature to the genes in this experiment? (Yellow and red lines correspond to top/bottom 1st and 5th percentile DMSO correlation to all the genes)	Transfer of the composition of the composition of the composition and the composition of	Distinguishing individual features for the compound relative to untreated samples. Black means a mismatch; i.e. active (= high z-score in magnitude) in the compound, and either inactive (= small z-score in magnitude) or oppositely active in the gene	the compound was tested; assays in
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AreaShape -Cytoplasm Nuclei\_Texture\_SumVariance\_AGP\_3\_0 Cytoplasm\_Intensity\_MaxIntensityEdge\_AGP BRD-K57745505-001-05-6 Cytoplasm\_Texture\_InfoMeas1\_DNA\_5\_0 ZINC00318849 Cytoplasm\_Texture\_DifferenceVariance\_AGP\_5\_0 STK11\_WT.1
RELB\_WT
XBP1\_WT.2
HSPA5\_WT
NFKBIA\_WT
YAP1\_WT.1
SMO\_WT.1
SMO\_WT.1
SMO\_WT.1
REL\_WT.1
REL\_WT.1
XBP1\_WT.2
BMP2\_WT
XAP1\_WT.2 Cytoplasm\_Texture\_AngularSecondMoment\_AGP\_3\_0
Nuclei\_Texture\_DifferenceVariance\_AGP\_5\_0 AC1LFSIA Nuclei\_Texture\_DifferenceEntropy\_Mito\_3\_0 Total number of assays tested in: 667. Active in the following assays: MLS000575147 0.63 (in 4 replicates) -0.42 HMS2325K20 • CYP2C9 Assay (AID 777) ZINC318849 Intensity -STL199970 SMR000196381 PubChem CID: 808132 Nuclei\_Texture\_AngularSecondMoment\_AGP\_5\_0 RadialDistribution -Mito ER