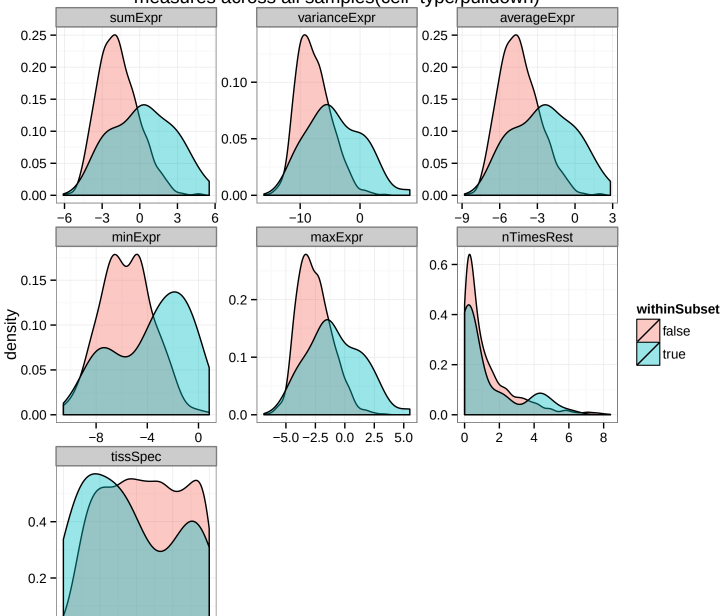
IncRNA group = IDRlessthan0_1 ::: data cols = bothPullDowns (45 / 1554) = (func/total IncRNA) ::: biotype = remove_antisense measures across all samples(cell-type/pulldown)



value of facet label

0.0

-2.0 -1.5 -1.0 -0.5