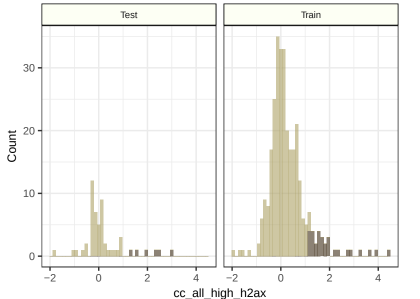
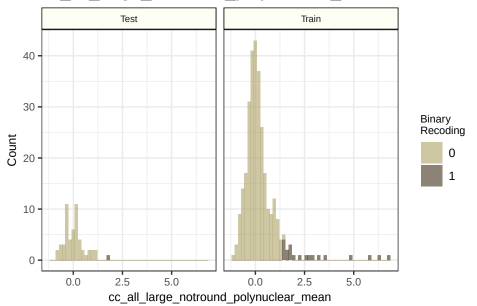
# cc\_all\_high\_h2ax

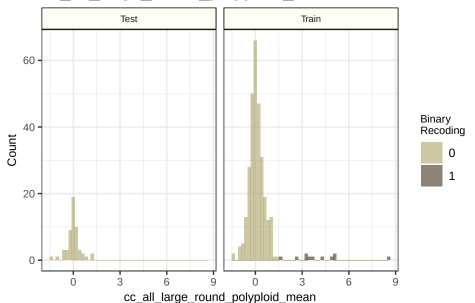




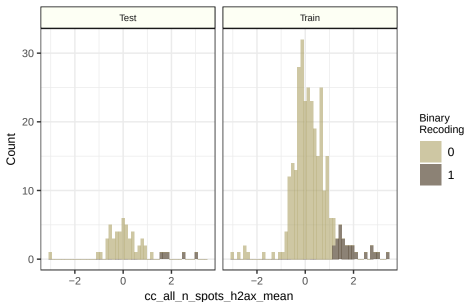
#### cc\_all\_large\_notround\_polynuclear\_mean



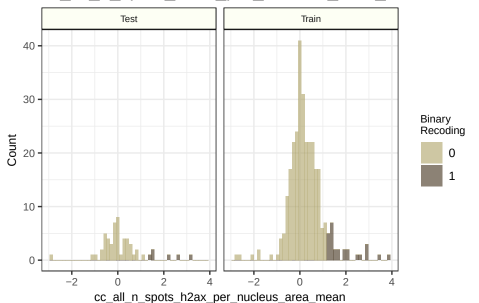
#### cc\_all\_large\_round\_polyploid\_mean



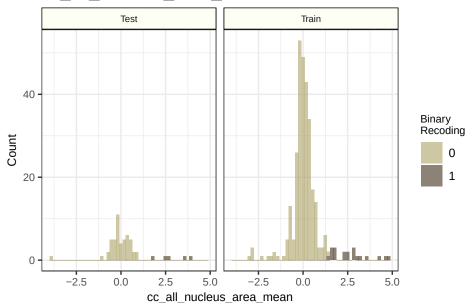
# cc\_all\_n\_spots\_h2ax\_mean



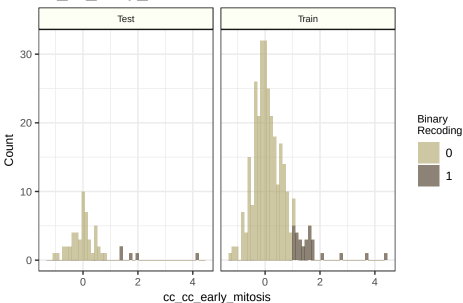
#### cc\_all\_n\_spots\_h2ax\_per\_nucleus\_area\_mean



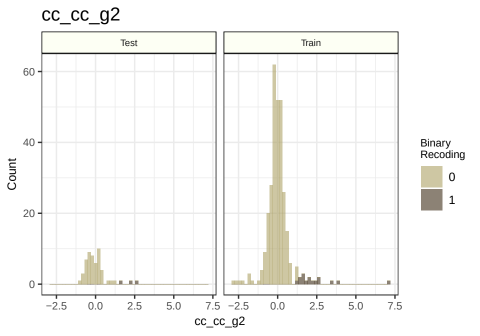
#### cc\_all\_nucleus\_area\_mean



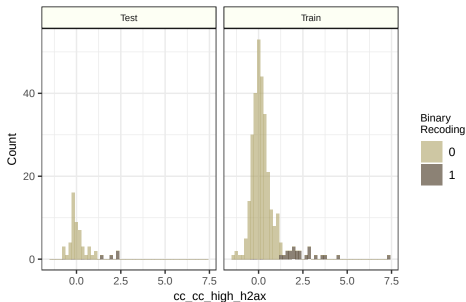
# cc\_cc\_early\_mitosis



cc\_cc\_g1 Test Train 40 -30 -Binary Recoding Count 0 10 -0 2.5 -2.5 0.0 -2.5 0.0 2.5 cc\_cc\_g1

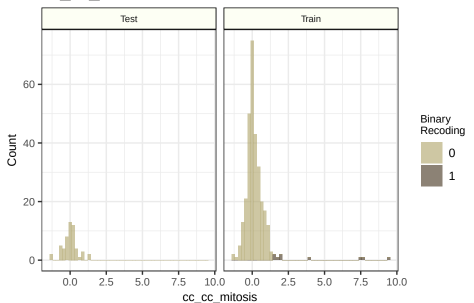


# cc\_cc\_high\_h2ax

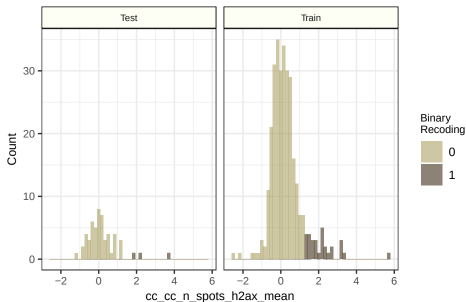


0

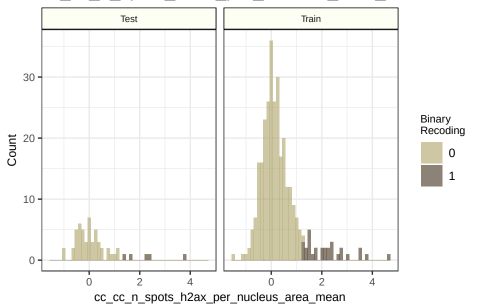
### cc\_cc\_mitosis

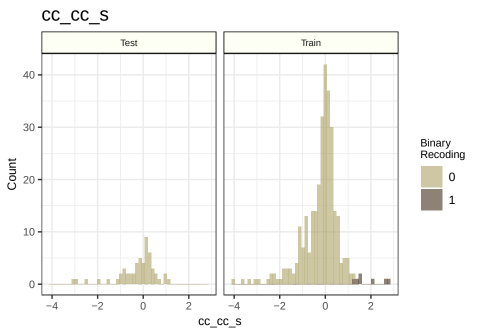


#### cc\_cc\_n\_spots\_h2ax\_mean

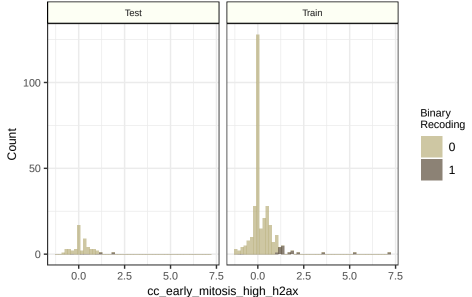


#### cc\_cc\_n\_spots\_h2ax\_per\_nucleus\_area\_mean

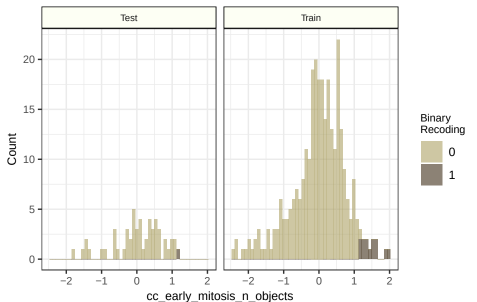




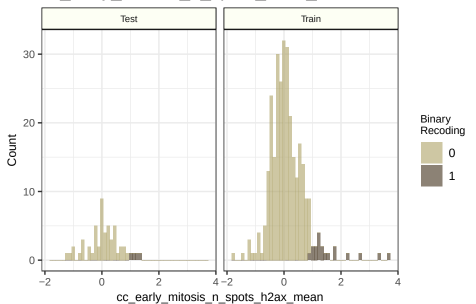
# cc\_early\_mitosis\_high\_h2ax



# cc\_early\_mitosis\_n\_objects



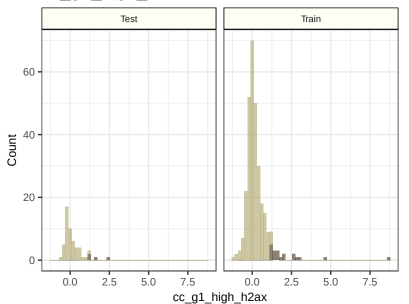
#### cc\_early\_mitosis\_n\_spots\_h2ax\_mean



cc\_early\_mitosis\_n\_spots\_h2ax\_per\_nucleus\_area\_i Test Train 30 . 20 -Binary Recoding Count 0 10 -0

cc\_early\_mitosis\_n\_spots\_h2ax\_per\_nucleus\_area\_mean

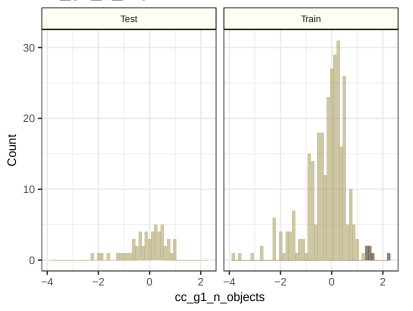
# cc\_g1\_high\_h2ax



Binary

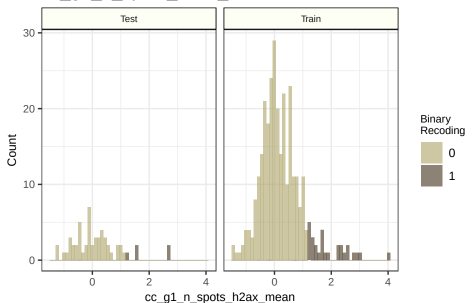
Recoding 0

# cc\_g1\_n\_objects

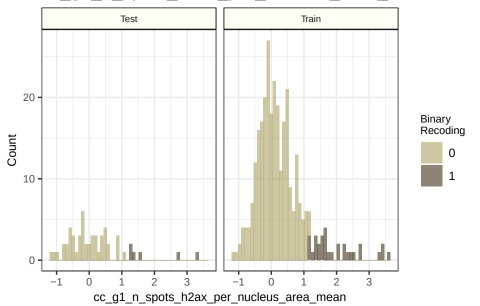




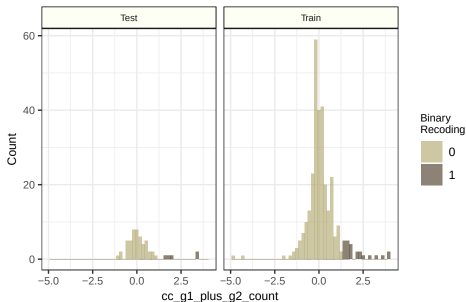
# cc\_g1\_n\_spots\_h2ax\_mean



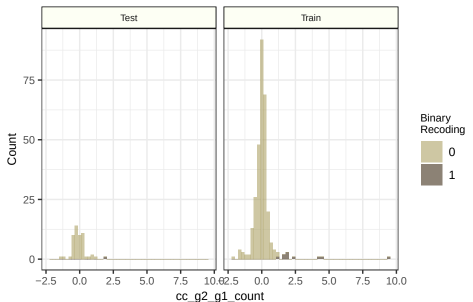
#### cc\_g1\_n\_spots\_h2ax\_per\_nucleus\_area\_mean



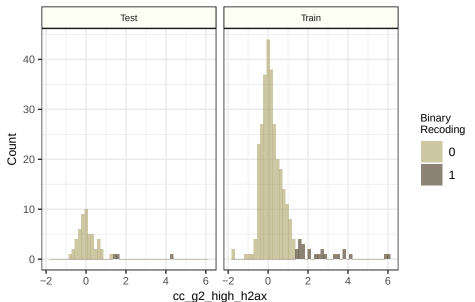
# cc\_g1\_plus\_g2\_count



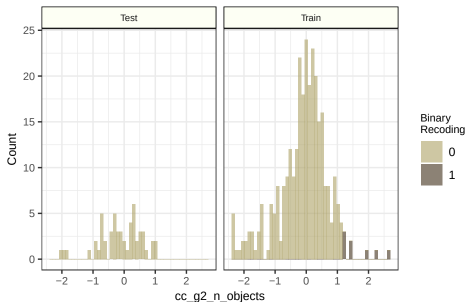
# cc\_g2\_g1\_count



# cc\_g2\_high\_h2ax

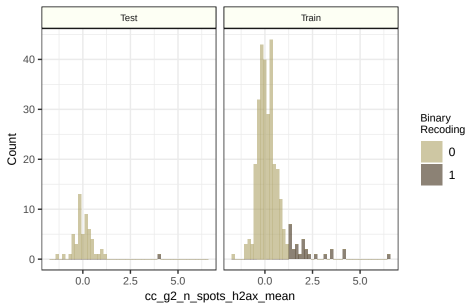


# cc\_g2\_n\_objects

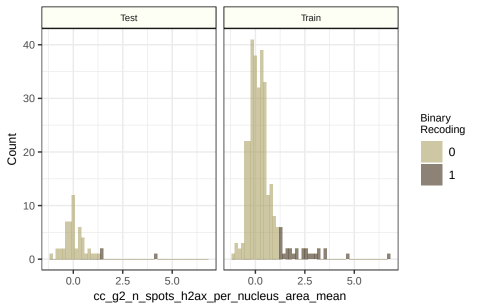


0

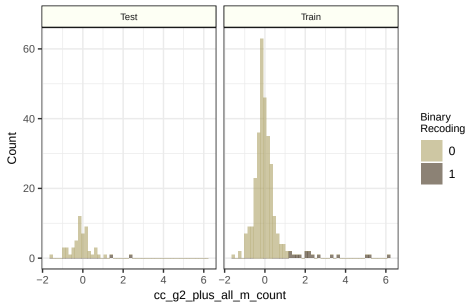
# cc\_g2\_n\_spots\_h2ax\_mean



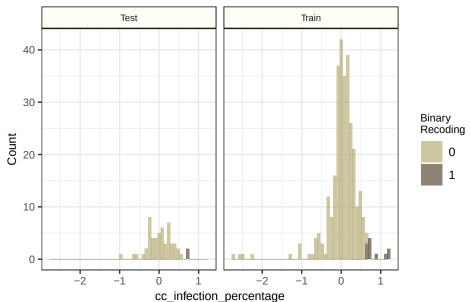
#### cc\_g2\_n\_spots\_h2ax\_per\_nucleus\_area\_mean



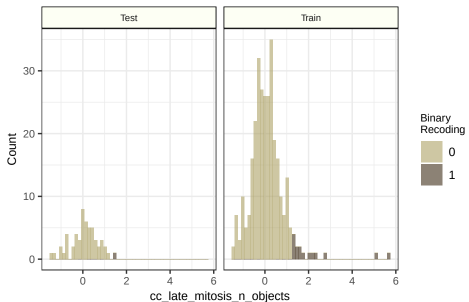
# cc\_g2\_plus\_all\_m\_count



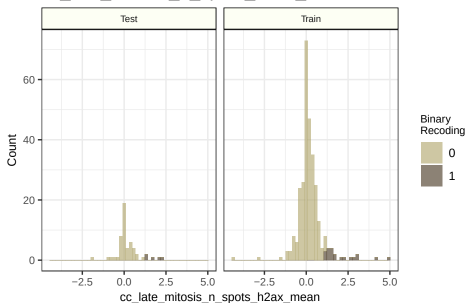
# cc\_infection\_percentage



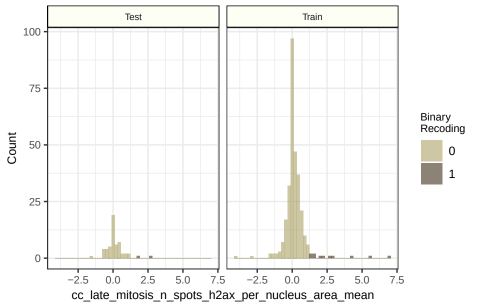
# cc\_late\_mitosis\_n\_objects



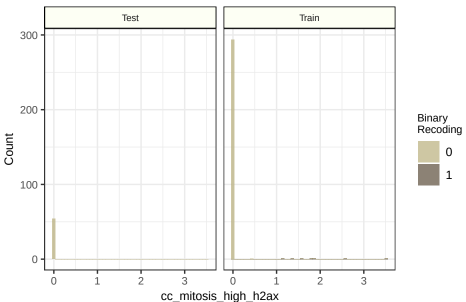
#### cc\_late\_mitosis\_n\_spots\_h2ax\_mean



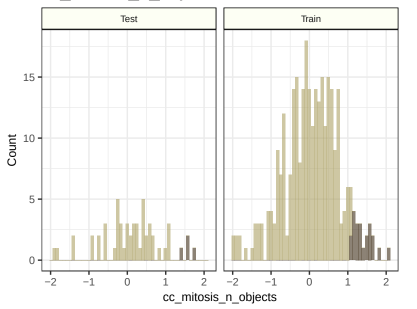
cc\_late\_mitosis\_n\_spots\_h2ax\_per\_nucleus\_area\_n



# cc\_mitosis\_high\_h2ax

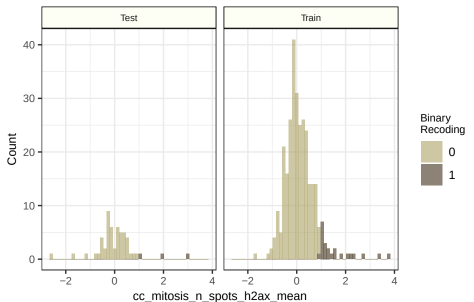


# cc\_mitosis\_n\_objects

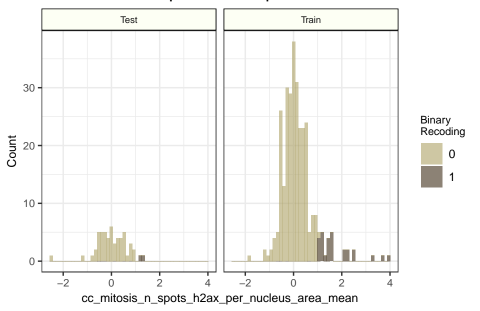




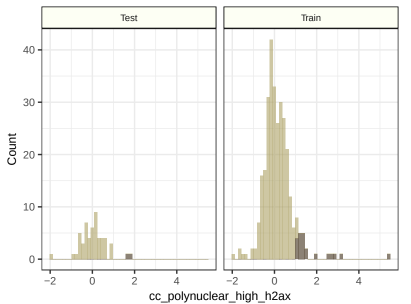
#### cc\_mitosis\_n\_spots\_h2ax\_mean



#### cc\_mitosis\_n\_spots\_h2ax\_per\_nucleus\_area\_mean

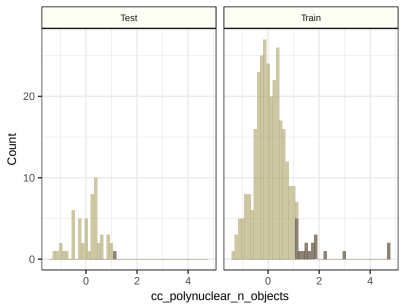


## cc\_polynuclear\_high\_h2ax



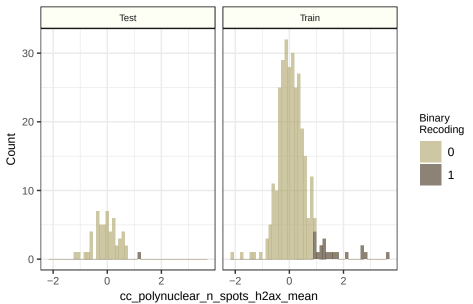


## cc\_polynuclear\_n\_objects





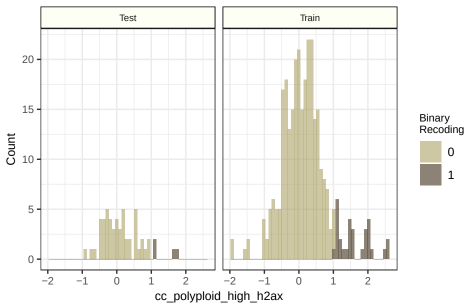
### cc\_polynuclear\_n\_spots\_h2ax\_mean



cc\_polynuclear\_n\_spots\_h2ax\_per\_nucleus\_area\_m Test Train 30 -Binary Count 50 Recoding 0 10 -0

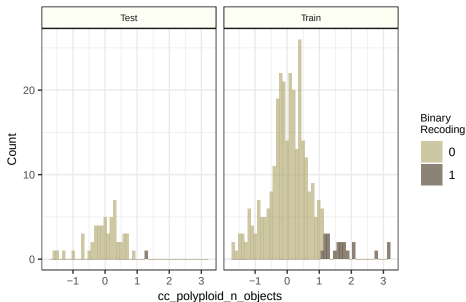
cc\_polynuclear\_n\_spots\_h2ax\_per\_nucleus\_area\_mean

# cc\_polyploid\_high\_h2ax



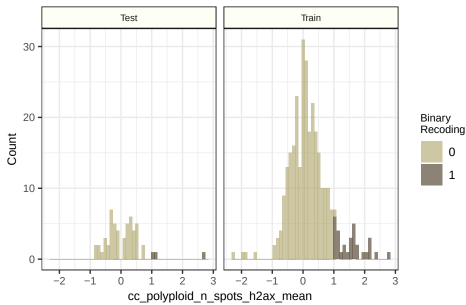
0

## cc\_polyploid\_n\_objects

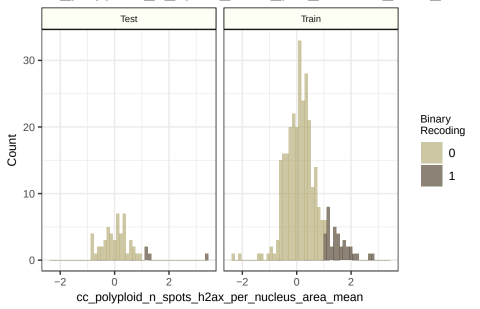


0

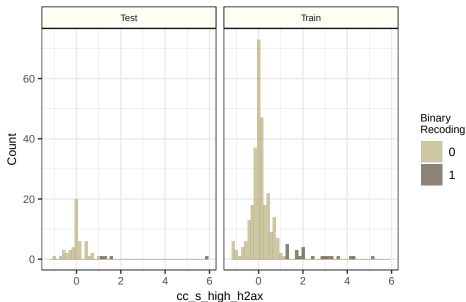
#### cc\_polyploid\_n\_spots\_h2ax\_mean



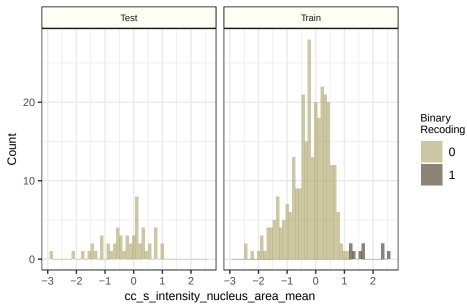
cc\_polyploid\_n\_spots\_h2ax\_per\_nucleus\_area\_mea



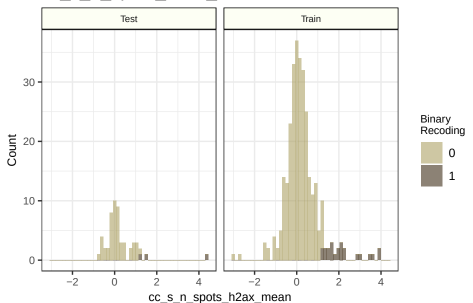
## cc\_s\_high\_h2ax



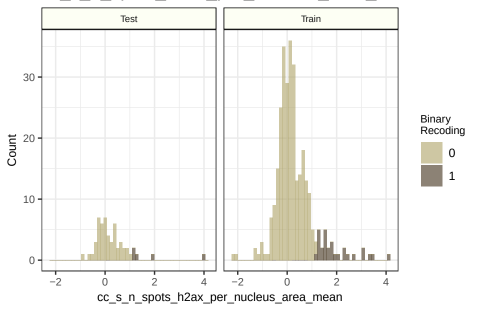
## cc\_s\_intensity\_nucleus\_area\_mean



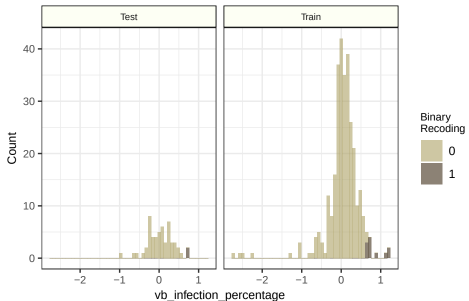
#### cc\_s\_n\_spots\_h2ax\_mean



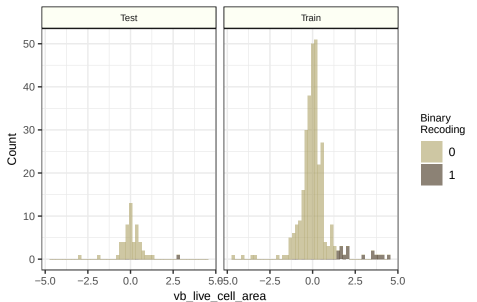
#### cc\_s\_n\_spots\_h2ax\_per\_nucleus\_area\_mean



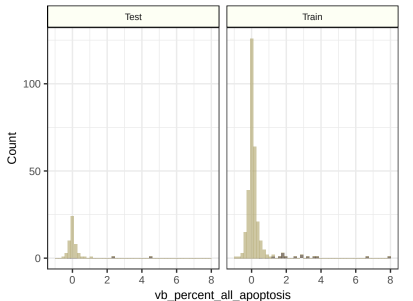
# vb\_infection\_percentage



vb\_live\_cell\_area



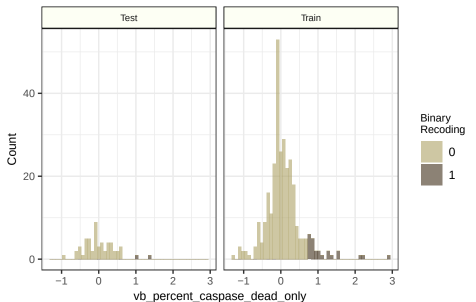
# vb\_percent\_all\_apoptosis



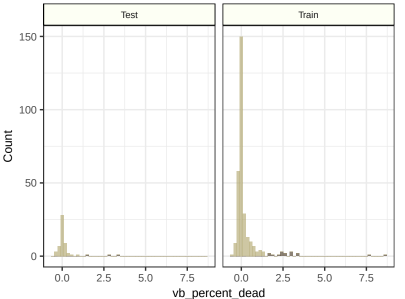
Binary Recoding

0

## vb\_percent\_caspase\_dead\_only



# vb\_percent\_dead



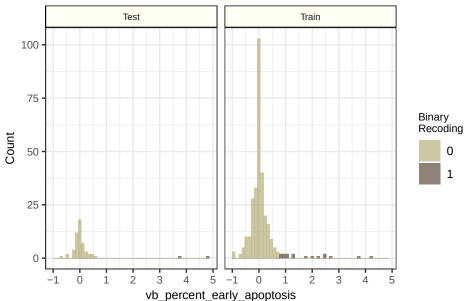
Binary Recoding

0

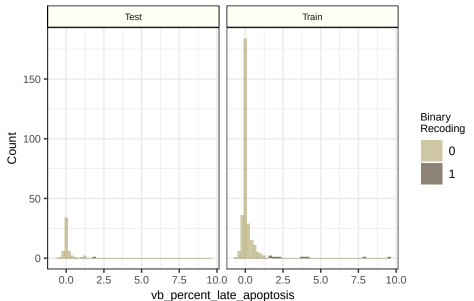
# vb\_percent\_dead\_only Test Train 80 -60 -Binary Recoding count 0 20 -

vb\_percent\_dead\_only

# vb\_percent\_early\_apoptosis



# vb\_percent\_late\_apoptosis



#### vb\_ros\_mean

