2018-10-19 11:23:28

Now that we have a bit better testing framework, let's repeat the within-modality assessment of datasets and see which ones fare best.

DTI

```
job_name=withinDTI_DL;
mydir=/data/NCR SBRB/baseline prediction/;
swarm_file=swarm.automl_${job_name};
rm -rf $swarm_file;
for f in dti ALL voxelwise n223 09212018.RData.gz
dti_ALL_voxelwise_n272_09212018.RData.gz \
    dti_fa_voxelwise_n223_09212018.RData.gz
dti fa voxelwise n272 09212018.RData.gz \
    dti ad voxelwise n223 09212018.RData.gz
dti_ad_voxelwise_n272_09212018.RData.gz \
    dti rd voxelwise n223 09212018.RData.gz
dti_rd_voxelwise_n272_09212018.RData.gz \
    dti_tracts_n223_10042018.RData.gz dti_tracts_n272_10042018.RData.gz;
do
    for target in nvVSper nvVSrem perVSrem nvVSadhd; do
        for i in {1..100}; do
            echo "Rscript --vanilla
~/research code/automl/uni test autoValidation DL.R ${mydir}/$f
${mydir}/long_clin_0918.csv ${target} ${mydir}/models_within_DL/${USER}
$RANDOM" >> $swarm_file;
        done;
    done;
done
sed -i -e "s/^/unset http_proxy; /g" $swarm_file;
split -l 1000 $swarm_file ${job_name}_split;
for f in `/bin/ls ${job_name}_split??`; do
    echo "ERROR" > swarm_wait_${USER}
    while grep -q ERROR swarm_wait_${USER}; do
        echo "Trying $f"
        swarm -f $f -g 60 -t 16 --time 3:00:00 --partition norm --logdir
trash_${job_name} --job-name ${job_name} -m R --gres=lscratch:10 2>
swarm_wait_${USER};
        if grep -q ERROR swarm_wait_${USER}; then
            echo -e "\tError, sleeping..."
            sleep 10m;
        fi;
    done;
done
```

Structural

```
job name=withinStruct DL;
mydir=/data/NCR SBRB/baseline prediction/;
swarm file=swarm.automl ${job name};
rm -rf $swarm file;
for f in struct thickness 09192018 260timeDiff12mo.RData.gz \
    struct_area_09192018_260timeDiff12mo.RData.gz \
    struct_volume_09192018_260timeDiff12mo.RData.gz \
    struct rois 09192018 260timeDiff12mo.RData.gz; do
    for target in nvVSper nvVSrem perVSrem nvVSadhd; do
        for i in {1..100}; do
            echo "Rscript --vanilla
~/research code/automl/uni test autoValidation DL.R ${mydir}/$f
${mydir}/long_clin_0918.csv ${target} ${mydir}/models_within_DL/${USER}
$RANDOM" >> $swarm_file;
        done;
    done:
done
sed -i -e "s/^/unset http_proxy; /g" $swarm_file;
split -l 1000 $swarm_file ${job_name}_split;
for f in `/bin/ls ${job_name}_split??`; do
    echo "ERROR" > swarm_wait_${USER}
    while grep -q ERROR swarm_wait_${USER}; do
        echo "Trying $f"
        swarm -f $f -g 60 -t 16 --time 3:00:00 --partition norm --logdir
trash_${job_name} --job-name ${job_name} -m R --gres=lscratch:10 2>
swarm wait ${USER};
        if grep -q ERROR swarm_wait_${USER}; then
            echo -e "\tError, sleeping..."
            sleep 10m;
        fi;
    done:
done
```

rsFMRI

```
done;
done
sed -i -e "s/^/unset http_proxy; /g" $swarm_file;
split -l 1000 $swarm_file ${job_name}_split;
for f in `/bin/ls ${job name} split??`; do
    echo "ERROR" > swarm wait ${USER}
    while grep -q ERROR swarm_wait_${USER}; do
        echo "Trying $f"
        swarm -f $f -q 60 -t 16 --time 3:00:00 --partition norm --logdir
trash_${job_name} --job-name ${job_name} -m R --gres=lscratch:10 2>
swarm_wait_${USER};
        if grep -g ERROR swarm wait ${USER}; then
            echo -e "\tError, sleeping..."
            sleep 10m;
        fi;
    done;
done
```

genetics

```
job name=withinSNPs DL;
mydir=/data/NCR SBRB/baseline prediction/;
swarm_file=swarm.automl_${job_name};
rm -rf $swarm file:
for f in geno3_snps1e04_09192018.RData.gz geno3_snps1e05_09192018.RData.gz
    geno3_snps1e05_09192018.RData.gz geno3_snps1e07_09192018.RData.gz \
    geno3_snps1e08_09192018.RData.gz geno3_snps1e09_09192018.RData.gz \
    geno3_prs_09192018.RData.gz; do
    for target in nvVSper nvVSrem perVSrem nvVSadhd; do
        for i in {1..100}; do
            echo "Rscript --vanilla
~/research_code/automl/uni_test_autoValidation_DL.R ${mydir}/$f
${mydir}/long clin 0918.csv ${target} ${mydir}/models within DL/${USER}
$RANDOM" >> $swarm file;
       done;
    done:
done
sed -i -e "s/^/unset http_proxy; /g" $swarm_file;
split -l 1000 $swarm_file ${job_name}_split;
for f in `/bin/ls ${job_name}_split??`; do
    echo "ERROR" > swarm_wait_${USER}
    while grep -q ERROR swarm_wait_${USER}; do
        echo "Trying $f"
        swarm -f $f -g 60 -t 16 --time 3:00:00 --partition norm --logdir
trash_${job_name} --job-name ${job_name} -m R --gres=lscratch:10 2>
swarm_wait_${USER};
        if grep -q ERROR swarm_wait_${USER}; then
            echo -e "\tError, sleeping..."
            sleep 10m;
        fi;
```

```
done;
done
```

2018-10-22 09:23:25

And now we need to grab the results:

```
echo "target,pheno,var,seed,nfeat,model,auc,f1,acc,spec,sens,prec,ratio" >
withinDomainAutoframeDL summary.csv;
for dir in DTI_DL Resting_DL SNPs_DL Struct_DL; do
    echo $dir
    for f in `ls trash_within${dir}/*o`; do
        phen=`head -n 2 $f | tail -1 | awk '{FS=" "; print $7}' | cut -
d"/" -f 5`:
        target=`head -n 2 $f | tail -1 | awk '{FS=" "; print $8}'`;
        seed=`head -n 2 $f | tail -1 | awk '{FS=" "; print $10}'`;
        var=`head -n 2 $f | tail -1 | awk '{FS=" "; print $5}' | cut -d"/"
-f 4 | sed -e "s/\.R//g"\;
       model=`grep -A 1 model id $f | tail -1 | awk '{FS=" "; print $2}'
| cut -d"_" -f 1`;
        auc=`grep -A 1 model_id $f | tail -1 | awk '{FS=" "; print $3}'`;
        nfeat=`grep "Running model on" $f | awk '{FS=" "; print $5}'`;
        ratio=`grep -A 1 "Class distribution" $f | tail -1 | awk '{FS=" ";
{for (i=2; i<=NF; i++) printf $i ";"}}'`;</pre>
        f1=`grep -A 2 "Maximum Metrics:" $f | tail -1 | awk '{FS=" ";
print $5}'`;
        acc=`grep -A 5 "Maximum Metrics:" $f | tail -1 | awk '{FS=" ";
print $5}'`;
        spec=`grep -A 8 "Maximum Metrics:" $f | tail -1 | awk '{FS=" ";
print $5}'`;
        sens=`grep -A 7 "Maximum Metrics:" $f | tail -1 | awk '{FS=" ";
print $5}'`;
        prec=`grep -A 6 "Maximum Metrics:" $f | tail -1 | awk '{FS=" ";
print $5}'`
        echo
$target,$phen,$var,$seed,$nfeat,$model,$auc,$f1,$acc,$spec,$sens,$prec,$ra
tio >> withinDomainAutoframeDL_summary.csv;
    done;
done
```

2018-10-25 11:50:35

Now that I know things are not doing so hot for PCA compared to raw, and while I play with ICA and AE on the side, let's check what's really the best phenotype within modality using raw, so that we can start to combine them and see if anything comes up that way.

```
job name=withinDTI rawDL;
mydir=/data/NCR SBRB/baseline prediction/;
swarm file=swarm.automl ${job name};
rm -rf $swarm file;
for f in dti ALL voxelwise n223 09212018.RData.gz
dti_ALL_voxelwise_n272_09212018.RData.gz \
    dti_fa_voxelwise_n223_09212018.RData.gz
dti_fa_voxelwise_n272_09212018.RData.gz \
    dti_ad_voxelwise_n223_09212018.RData.gz
dti_ad_voxelwise_n272_09212018.RData.gz \
    dti rd voxelwise n223 09212018.RData.gz
dti_rd_voxelwise_n272_09212018.RData.gz \
    dti_tracts_n223_10042018.RData.gz dti_tracts_n272_10042018.RData.gz;
do
    for target in nvVSper nvVSrem perVSrem nvVSadhd; do
        for i in {1..100}; do
            echo "Rscript --vanilla
~/research code/automl/raw autoValidation DL.R ${mydir}/$f
${mydir}/long_clin_0918.csv ${target}
${mydir}/models_raw_within_DL/${USER} $RANDOM" >> $swarm_file;
        done:
    done;
done
for f in dti_ALL_voxelwise_n223_09212018.RData.gz
dti_ALL_voxelwise_n272_09212018.RData.gz \
    dti_rd_voxelwise_n223_09212018.RData.gz
dti_rd_voxelwise_n272_09212018.RData.gz \
    dti_tracts_n223_10042018.RData.gz dti_tracts_n272_10042018.RData.gz;
do
    for target in nvVSper nvVSrem perVSrem nvVSadhd; do
        for i in {1..100}; do
            echo "Rscript --vanilla
~/research_code/automl/raw_autoValidation_DL.R ${mydir}/$f
${mydir}/long_clin_0918.csv ${target}
${mydir}/models_raw_within_DL/${USER} -$RANDOM" >> $swarm_file;
        done;
    done;
done
sed -i -e "s/^/unset http_proxy; /g" $swarm_file;
split -l 1000 $swarm_file ${job_name}_split;
for f in `/bin/ls ${job name} split??`; do
    echo "ERROR" > swarm_wait_${USER}
    while grep -q ERROR swarm_wait_${USER}; do
        echo "Trying $f"
        swarm -f $f -g 60 -t 16 --time 3:00:00 --partition norm --logdir
trash_${job_name} --job-name ${job_name} -m R --gres=lscratch:10 2>
swarm_wait_${USER};
        if grep -q ERROR swarm_wait_${USER}; then
            echo -e "\tError, sleeping..."
            sleep 10m;
        fi;
    done;
done
```

```
job_name=withinStruct_rawDL;
mydir=/data/NCR SBRB/baseline prediction/;
swarm_file=swarm.automl_${job_name};
rm -rf $swarm_file;
for f in struct_thickness_09192018_260timeDiff12mo.RData.gz \
    struct_area_09192018_260timeDiff12mo.RData.gz \
    struct_volume_09192018_260timeDiff12mo.RData.gz \
    struct_rois_09192018_260timeDiff12mo.RData.gz; do
    for target in nvVSper nvVSrem perVSrem nvVSadhd; do
        for i in {1..100}; do
            echo "Rscript --vanilla
~/research code/automl/raw autoValidation DL.R ${mydir}/$f
${mydir}/long_clin_0918.csv ${target}
${mydir}/models_raw_within_DL/${USER} $RANDOM" >> $swarm_file;
        done;
    done:
done
for f in struct_volume_09192018_260timeDiff12mo.RData.gz \
    struct rois 09192018 260timeDiff12mo.RData.qz018.RData.qz; do
    for target in nvVSper nvVSrem perVSrem nvVSadhd; do
        for i in {1..100}; do
            echo "Rscript --vanilla
~/research_code/automl/raw_autoValidation_DL.R ${mydir}/$f
${mydir}/long_clin_0918.csv ${target}
${mydir}/models_raw_within_DL/${USER} -$RANDOM" >> $swarm_file;
        done;
    done;
done
sed -i -e "s/^/unset http_proxy; /g" $swarm_file;
split -l 1000 $swarm_file ${job_name}_split;
for f in `/bin/ls ${job_name}_split??`; do
    echo "ERROR" > swarm_wait_${USER}
    while grep -q ERROR swarm_wait_${USER}; do
        echo "Trying $f"
        swarm -f $f -q 60 -t 16 --time 3:00:00 --partition norm --logdir
trash_${job_name} --job-name ${job_name} -m R --gres=lscratch:10 2>
swarm_wait_${USER};
        if grep -q ERROR swarm_wait_${USER}; then
            echo -e "\tError, sleeping..."
            sleep 10m;
        fi;
    done;
done
```

```
job_name=withinResting_rawDL;
mydir=/data/NCR_SBRB/baseline_prediction/;
swarm_file=swarm.automl_${job_name};
rm -rf $swarm_file;
```

```
for f in aparc.a2009s_n215_10252018.RData.gz \
    aparc.a2009s trimmed n215 10252018.RData.gz \
    aparc_n215_10252018.RData.gz \
    aparc_trimmed_n215_10252018.RData.gz; do
    for target in nvVSper nvVSrem perVSrem nvVSadhd; do
        for i in {1..100}; do
            echo "Rscript --vanilla
~/research code/automl/raw autoValidation DL.R ${mydir}/$f
${mydir}/long clin 0918.csv ${target}
${mydir}/models_raw_within_DL/${USER} $RANDOM" >> $swarm_file;
        done;
    done:
done
for f in aparc.a2009s_n215_10252018.RData.gz aparc_n215_10252018.RData.gz;
do
    for target in nvVSper nvVSrem perVSrem nvVSadhd; do
        for i in {1..100}; do
            echo "Rscript --vanilla
~/research code/automl/raw autoValidation DL.R ${mydir}/$f
${mydir}/long_clin_0918.csv ${target}
${mydir}/models_raw_within_DL/${USER} -$RANDOM" >> $swarm_file;
        done:
    done:
done
sed -i -e "s/^/unset http_proxy; /g" $swarm_file;
split -l 1000 $swarm_file ${job_name}_split;
for f in `/bin/ls ${job_name}_split??`; do
    echo "ERROR" > swarm wait ${USER}
    while grep -q ERROR swarm_wait_${USER}; do
        echo "Trying $f"
        swarm -f $f -g 60 -t 16 --time 3:00:00 --partition norm --logdir
trash_${job_name} --job-name ${job_name} -m R --gres=lscratch:10 2>
swarm_wait_${USER};
        if grep -q ERROR swarm_wait_${USER}; then
            echo -e "\tError, sleeping..."
            sleep 10m;
        fi;
    done:
done
```

```
job_name=withinSNP_rawDL;
mydir=/data/NCR_SBRB/baseline_prediction/;
swarm_file=swarm.automl_${job_name};
rm -rf $swarm_file;
for f in geno3_snps1e04_09192018.RData.gz geno3_snps1e05_09192018.RData.gz
\
    geno3_snps1e05_09192018.RData.gz geno3_snps1e07_09192018.RData.gz \
    geno3_snps1e08_09192018.RData.gz geno3_snps1e09_09192018.RData.gz \
    geno3_prs_09192018.RData.gz; do
    for target in nvVSper nvVSrem perVSrem nvVSadhd; do
        for i in {1..100}; do
```

```
echo "Rscript --vanilla
~/research code/automl/raw autoValidation DL.R ${mydir}/$f
${mydir}/long_clin_0918.csv ${target}
${mydir}/models_raw_within_DL/${USER} $RANDOM" >> $swarm_file;
        done:
    done;
done
for f in geno3 snps1e04 09192018.RData.gz geno3 snps1e05 09192018.RData.gz
    geno3_snps1e05_09192018.RData.gz geno3_snps1e07_09192018.RData.gz \
    geno3_snps1e08_09192018.RData.gz geno3_snps1e09_09192018.RData.gz \
    geno3 prs 09192018.RData.gz; do
    for target in nvVSper nvVSrem perVSrem nvVSadhd; do
        for i in {1..100}; do
            echo "Rscript --vanilla
~/research_code/automl/raw_autoValidation_DL.R ${mydir}/$f
${mydir}/long_clin_0918.csv ${target}
${mydir}/models_raw_within_DL/${USER} -$RANDOM" >> $swarm_file;
    done;
done
sed -i -e "s/^/unset http_proxy; /g" $swarm_file;
split -l 1000 $swarm_file ${job_name}_split;
for f in `/bin/ls ${job_name}_split??`; do
    echo "ERROR" > swarm_wait_${USER}
    while grep -q ERROR swarm_wait_${USER}; do
        echo "Trying $f"
        swarm -f $f -g 60 -t 16 --time 3:00:00 --partition norm --logdir
trash_${job_name} --job-name ${job_name} -m R --gres=lscratch:10 2>
swarm wait ${USER};
        if grep -g ERROR swarm wait ${USER}; then
            echo -e "\tError, sleeping..."
            sleep 10m;
        fi;
    done:
done
```

2018-10-26 14:50:56

DTI is the only one that is still running. So, we can go ahead and evaluate the other ones:

```
dir=withinSNP_rawDL;
echo "target,pheno,var,seed,nfeat,model,auc,f1,acc,ratio" >
${dir}_summary.csv;
for f in `ls -1 trash_${dir}/*o`; do
    phen=`head -n 2 $f | tail -1 | awk '{FS=" "; print $6}' | cut -d"/" -f
6`;
    target=`head -n 2 $f | tail -1 | awk '{FS=" "; print $8}'`;
    seed=`head -n 2 $f | tail -1 | awk '{FS=" "; print $10}'`;
    var=`head -n 2 $f | tail -1 | awk '{FS=" "; print $5}' | cut -d"/" -f
```

```
4 | sed -e "s/\.R//g"`;
    model=`grep -A 1 model_id $f | tail -1 | awk '{FS=" "; print $2}' |
cut -d"_" -f 1`;
    auc=`grep -A 1 model_id $f | tail -1 | awk '{FS=" "; print $3}'`;
    nfeat=`grep "Running model on" $f | awk '{FS=" "; print $5}'`;
    ratio=`grep -A 1 "Class distribution" $f | tail -1 | awk '{FS=" ";
    {for (i=2; i<=NF; i++) printf $i ";"}}'`;
    f1=`grep -A 2 "Maximum Metrics:" $f | tail -1 | awk '{FS=" "; print $5}'`;
    acc=`grep -A 5 "Maximum Metrics:" $f | tail -1 | awk '{FS=" "; print $5}'`;
    echo $target,$phen,$var,$seed,$nfeat,$model,$auc,$f1,$acc,$ratio >> ${dir}_summary.csv;
done;
```

Rerunning one of the fMRI trimmed results because of a typo. Also had to re-run all struct voxelwise because we ran out of memory.

The issue with structural raw is in converting to h2o... not sure what's going on:

```
dtrain = as.h2o(df[, x])
Error in h2o.parseSetup(data, pattern = "", destination_frame, header, :
  length of col.names must equal to the number of columns in dataset
```

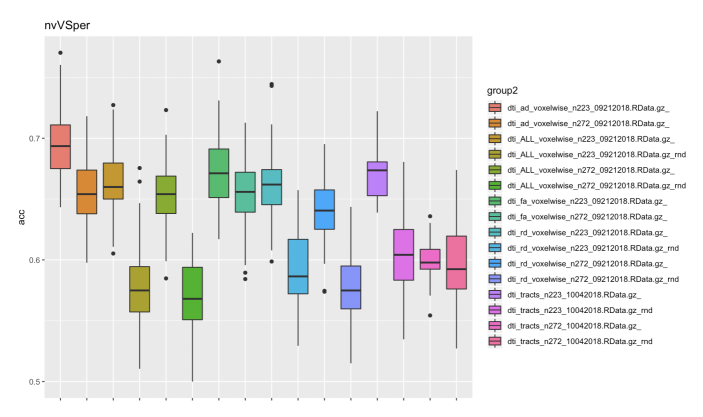
need to look more into it...

2018-10-29 10:17:03

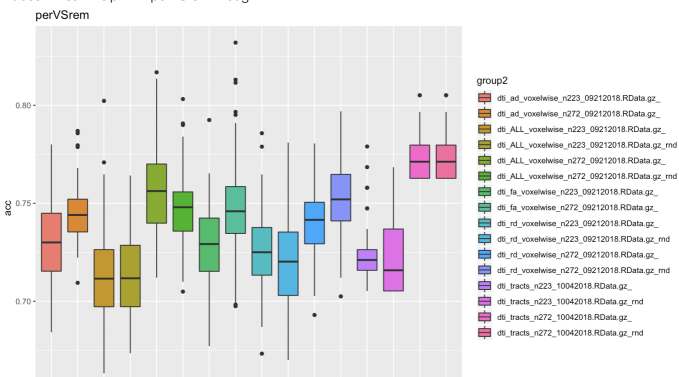
```
for dir in withinDTI rawDL withinResting rawDL withinSNP rawDL; do
    echo $dir;
    echo "target,pheno,var,seed,nfeat,model,auc,f1,acc,ratio" >
${dir}_summary.csv;
    for f in `ls -1 trash ${dir}/*o`; do
        phen=`head -n 2 $f | tail -1 | awk '{FS=" "; print $6}' | cut -
d"/" -f 6`;
        target=`head -n 2 $f | tail -1 | awk '{FS=" "; print $8}'`;
        seed=`head -n 2 $f | tail -1 | awk '{FS=" "; print $10}'`;
        var=`head -n 2 $f | tail -1 | awk '{FS=" "; print $5}' | cut -d"/"
-f 4 \mid sed -e "s/\.R//g";
        model=`grep -A 1 model_id $f | tail -1 | awk '{FS=" "; print $2}'
| cut -d"_" -f 1`;
        auc=`grep -A 1 model_id $f | tail -1 | awk '{FS=" "; print $3}'`;
        nfeat=`grep "Running model on" $f | awk '{FS=" "; print $5}'`;
        ratio=`grep -A 1 "Class distribution" $f | tail -1 | awk '{FS=" ";
{for (i=2; i<=NF; i++) printf $i ";"}}'`;
        f1=`grep -A 2 "Maximum Metrics:" $f | tail -1 | awk '{FS=" ";
print $5}'`;
        acc=`grep -A 5 "Maximum Metrics:" $f | tail -1 | awk '{FS=" ";
```

```
print $5}'`;
        echo $target,$phen,$var,$seed,$nfeat,$model,$auc,$f1,$acc,$ratio
>> ${dir}_summary.csv;
        done;
done;
```

So, apparently DTI_AD_n223 is still the best, and it does perform much better tha random... the actual results is just not as exciting.



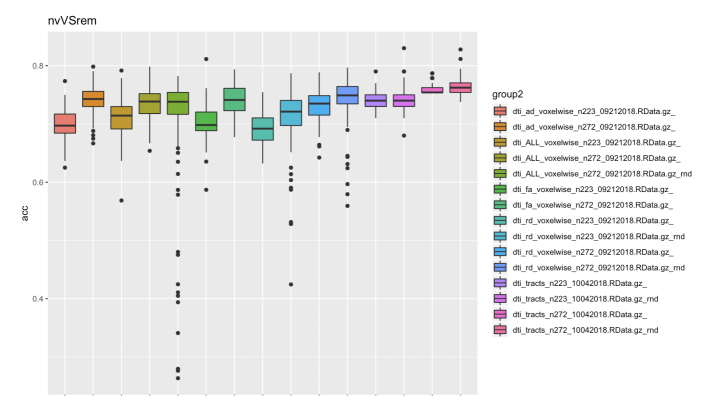
It doesn't reallt help with perVSrem though.

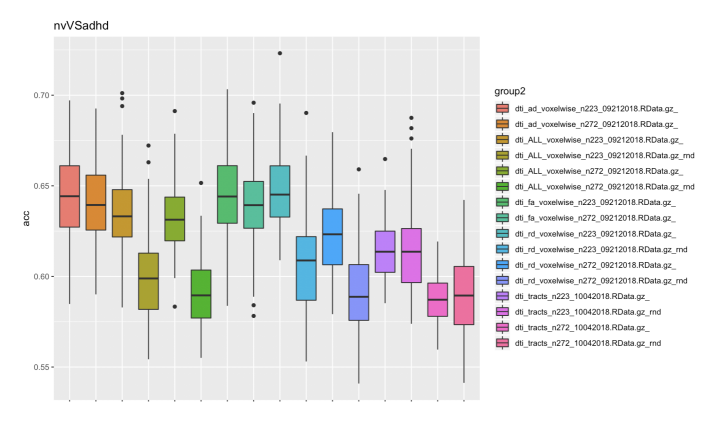


This is what I'm running to plot, btw:

```
data$group = ''
data[data$seed<0,]$group = 'rnd'
data$group2 = sapply(1:nrow(data), function(x) { sprintf('%s_%s',
data$pheno[x], data$group[x])} )
# then, for each target
target='nvVSper'
p1<-ggplot(data[data$target == target,], aes(x=group2, y=acc,
fill=group2))
print(p1+geom_boxplot() + ggtitle(target))</pre>
```

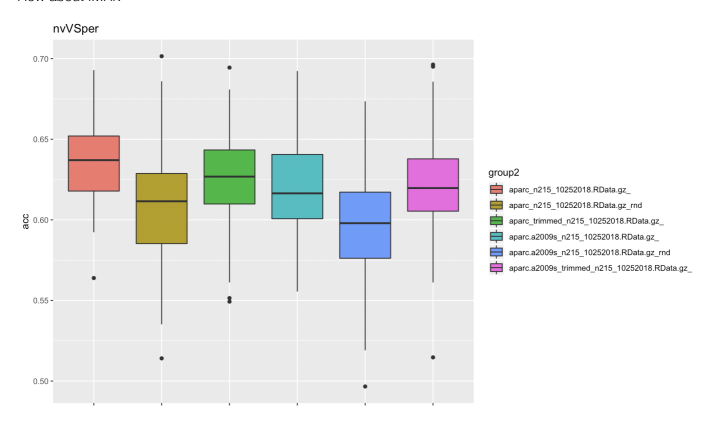
The other two plots, just for giggles:

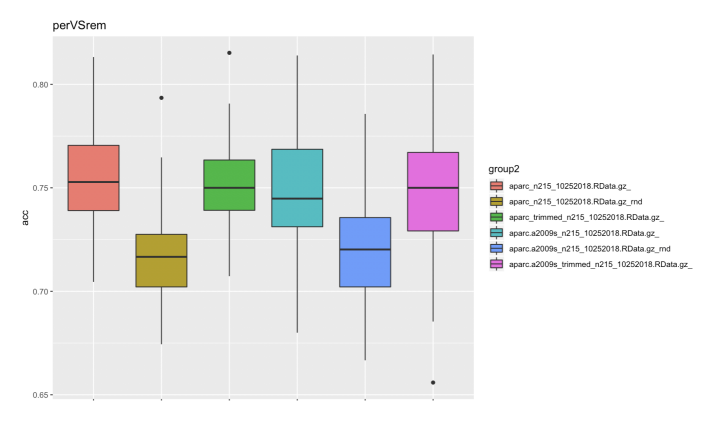




Plotting other metrics doesn't make that much difference either.

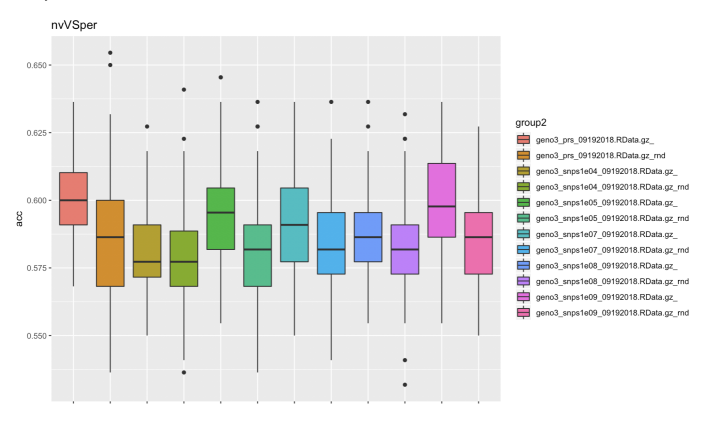
How about fMRI?

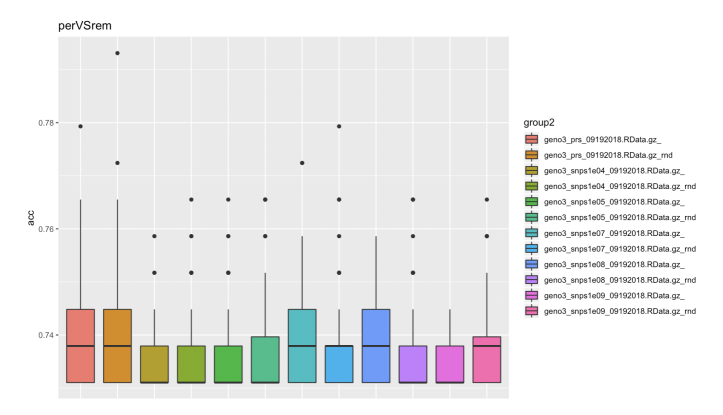




So, again, there is some signal there, a bit better than chance, but still nothing impressive.

Finally, SNPs.





As expected, barely better than chance.

BTW, it looks like it works to have the raw MR data firs tsaved to a tmp .csv and then read in with importFile. So, let's change the raw function to do that in those cases!

2018-10-30 07:54:01

I was looking at some old notes, and even if I manage to load the structural data in the raw script, it will fail during computations due to lack of memory, even in 120Gb machines. I could use a large memory node, but this might be overkill. Let's invest some time in the same space idea and see what we get.

2018-11-01 16:39:03

Going to fire up the within-domain tests using the spatial filter and test set approach, using 90-10 split. This only applies to DTI and spatial, but it's still worth trying out.

```
echo "Rscript --vanilla
~/research_code/automl/uni_spatial_test_autoValidation_DL.R ${mydir}/$f
${mydir}/long_clin_0918.csv ${target}
${mydir}/models_spatial_within_DL/${USER} $RANDOM 8" >> $swarm_file;
        done:
   done;
done
sed -i -e "s/^/unset http proxy; /g" $swarm file;
split -l 1000 $swarm file ${job name} split;
for f in `/bin/ls ${job_name}_split??`; do
   echo "ERROR" > swarm_wait_${USER}
   while grep -q ERROR swarm_wait_${USER}; do
        echo "Trying $f"
        swarm -f $f -g 40 -t 16 --time 3:00:00 --partition norm --logdir
trash ${job name} --job-name ${job name} -m R,afni --gres=lscratch:10 2>
swarm_wait_${USER};
        if grep -q ERROR swarm_wait_${USER}; then
            echo -e "\tError, sleeping..."
            sleep 10m;
        fi;
   done;
done
```

```
job_name=withinStruct_spatialTestDL;
mydir=/data/NCR SBRB/baseline prediction/;
swarm_file=swarm.automl_${job_name};
rm -rf $swarm file;
for f in struct thickness 09192018 260timeDiff12mo.RData.gz \
    struct area 09192018 260timeDiff12mo.RData.gz \
    struct_volume_09192018_260timeDiff12mo.RData.gz; do
    for target in nvVSper nvVSrem perVSrem nvVSadhd; do
        for i in {1..100}; do
            echo "Rscript --vanilla
~/research_code/automl/uni_spatial_test_autoValidation_DL.R ${mydir}/$f
${mydir}/long clin 0918.csv ${target}
${mydir}/models_spatial_within_DL/${USER} $RANDOM 35" >> $swarm_file;
        done;
    done:
done
sed -i -e "s/^/unset http_proxy; /g" $swarm_file;
split -l 1000 $swarm_file ${job_name}_split;
for f in `/bin/ls ${job_name}_split??`; do
    echo "ERROR" > swarm_wait_${USER}
   while grep -q ERROR swarm_wait_${USER}; do
        echo "Trying $f"
        swarm -f $f -g 60 -t 16 --time 3:00:00 --partition norm --logdir
trash_${job_name} --job-name ${job_name} -m R,afni --gres=lscratch:10 2>
swarm_wait_${USER};
        if grep -q ERROR swarm_wait_${USER}; then
            echo -e "\tError, sleeping..."
            sleep 10m;
```

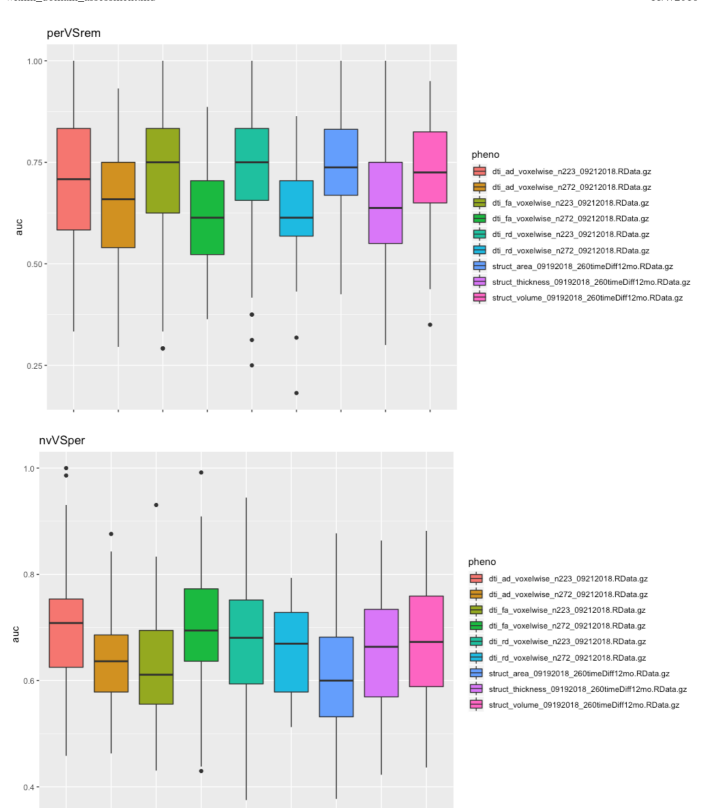
```
fi;
done;
done
```

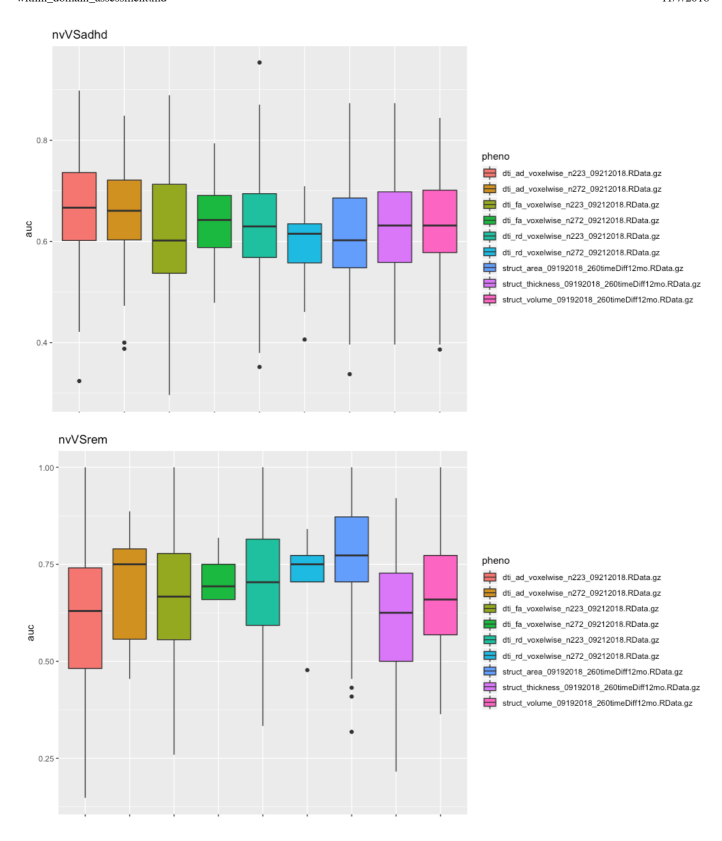
2018-11-02 09:36:57

Time to grab the results again:

```
echo "target,pheno,var,seed,nfeat,model,auc,f1,acc,ratio" >
spatialTest summary.csv;
for dir in withinDTI_spatialTestDL withinStruct_spatialTestDL; do
    echo $dir;
    for f in `ls -1 trash_${dir}/*o`; do
        phen=`head -n 2 $f | tail -1 | awk '{FS=" "; print $6}' | cut -
d"/" -f 6`;
        target=`head -n 2 $f | tail -1 | awk '{FS=" "; print $8}'`;
        seed=`head -n 2 $f | tail -1 | awk '{FS=" "; print $10}'`;
        var=`head -n 2 $f | tail -1 | awk '{FS=" "; print $5}' | cut -d"/"
-f 4 | sed -e "s/\.R//q"\;
        model=`grep -A 1 model_id $f | tail -1 | awk '{FS=" "; print $2}'
| cut -d"_" -f 1`;
        auc=`grep -A 1 model id $f | tail -1 | awk '{FS=" "; print $3}'`;
        nfeat=`grep "Running model on" $f | awk '{FS=" "; print $5}'`;
        ratio=`grep -A 1 "Class distribution" $f | tail -1 | awk '{FS=" ";
{for (i=2; i<=NF; i++) printf $i ";"}}'`;</pre>
        f1=`grep -A 2 "Maximum Metrics:" $f | tail -1 | awk '{FS=" ";
print $5}'`;
        acc=`grep -A 5 "Maximum Metrics:" $f | tail -1 | awk '{FS=" ";
print $5}'`;
        echo $target,$phen,$var,$seed,$nfeat,$model,$auc,$f1,$acc,$ratio
>> spatialTest_summary.csv;
    done;
done;
```

I'm plotting AUC because it's easier to see the differences there. But it looks like the n223 DTI is indeed better. For struct, area and volume are better!





But let's also plot accuracy, in case it helps with decisions:

