ABIDEI thickness Dx efffects

OK let's do the funner thing...let's try to calculate some effect sizes for diagnosis effects

```
library(dplyr)
library(tidyr)
library(broom)
library(ggplot2)
library(knitr)
```

read in the data

```
fs5.1 = read.csv('../data/cortical_fs5.1_measuresenigma_thickavg.csv', header = T)
#names(fs5.1) <- pasteO(names(fs5.1), '_fs5.1')
fs5.3 = read.csv('../data/ABIDE_fs5.3_thickness.csv', header = T)
#names(fs5.3) <- pasteO(names(fs5.3), '_fs5.3')
ants = read.csv('../data/ABIDE_ants_thickness_data_r.csv', header = T)
#names(ants) <- pasteO(names(ants), '_ants')
pheno = read.csv('../data/ABIDE_Phenotype.csv', header = T)
pheno$DX_GROUP <- factor(pheno$DX_GROUP, levels = c(1,2), labels = c("ASD", "TD"))</pre>
```

Starting with FS 5.3 let's see what diagnosis effect sizes we can get

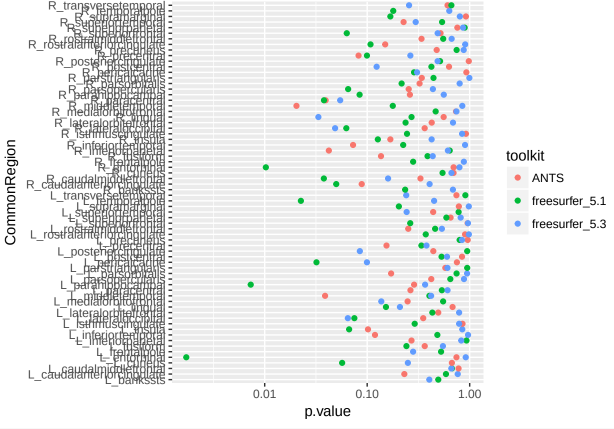
```
## Merge it all into one mega- data - frame
fs5.3_thick <- fs5.3 %>%
  gather(Region, thickness, ends_with("thickavg")) %>%
  mutate(CommonRegion = gsub('_thickavg','',Region)) %>%
  select(SubjID, CommonRegion, thickness)
fs5.3_thick$toolkit <- "freesurfer_5.3"
fs5.1_thick <- fs5.1 %>%
  gather(Region, thickness, ends_with("thickavg")) %>%
  mutate(CommonRegion = gsub('_thickavg','',Region)) %>%
  select(SubjID, CommonRegion, thickness)
fs5.1_thick$toolkit <- "freesurfer_5.1"
ants_thick <- ants %>%
  gather (Region, thickness, starts with ("right"), starts with ("left")) %%
  mutate(CommonRegion = gsub('\\.','', gsub('right','R_', gsub('left','L_',Region)))) %>%
 filter(CommonRegion %in% unique(fs5.3_thick$CommonRegion)) %>%
  select(Code, CommonRegion, thickness)
ants thick$toolkit <- "ANTS"
names(ants_thick) <- c("SubjID", "CommonRegion", "thickness", "toolkit")</pre>
allthickness <- rbind(fs5.3_thick, fs5.1_thick, ants_thick)
df <- merge(allthickness, pheno, by.x = "SubjID", by.y = "Subject_ID")</pre>
dx raw <- df %>%
 group_by(CommonRegion, toolkit) %>%
```

```
do(glance(lm(thickness ~ DX_GROUP, .)))

dx_wcovs<- df %>%
  group_by(CommonRegion, toolkit) %>%
  do(tidy(lm(thickness ~ DX_GROUP + AGE_AT_SCAN + SITE_ID, .)))

dx_effect_wcov <- filter(dx_wcovs, term == "DX_GROUPTD")
  age_effect_wcov <- filter(dx_wcovs, term == "AGE_AT_SCAN")

ggplot(dx_raw, aes(y=p.value , x=CommonRegion, color = toolkit)) +
  geom_point() +
  scale_y_log10() +
  coord_flip()</pre>
```



```
ggplot(dx_effect_wcov, aes(y=p.value , x=CommonRegion, color = toolkit)) +
  geom_point() +
  scale_y_log10() +
  coord_flip()
```

