# Differences in Resting-State Functional MRI Quality Control Recommendations Alter the Representativeness of the Adolescent Brain Cognitive Development (ABCD) Study Supplemental Methods and Results

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## 1 Supplemental Methods

This section includes R code used to generate tables, figures, and statistics used in the main manuscript. Additional code (specifically, for pre-processing and coding the raw data) is available on osf: https://osf.io/57xer/.

```
recodesid <- function(x) {</pre>
  paste0("NDAR", substr(x, start = 6, stop = nchar(x)))
}
t1<-readRDS("ABCD-Data-Import/tabulateddata.Rds")
t2<-readRDS("ABCC_download/ABCCsubjectdata.RDS")
df<-left join(readRDS("ABCD-Data-Import/tabulateddata.Rds") %>%
                mutate(subjectkey=recodesid(as.character(subjectkey))),
              readRDS("ABCC download/ABCCsubjectdata.RDS"),
              by=c("subjectkey"="sname"))
df<-df %>%
  mutate(subjectkeychar=subjectkey,
         subjectkey=as.factor(subjectkey)) %>%
  # Pt's without ABCC data should have FALSE for the 'Scan Exists' booleans
  mutate(across(starts_with("Exists"),
                ~ case_when(. == "True" ~ TRUE,
                            . == "False" ~ FALSE,
                            is.na(.) ~ FALSE))) %>%
  # Pt's without ABCC data should have TRUE for > 5 minute data booleans.
  mutate(across(starts with("ltfive"),
                ~ replace_na(.,TRUE))) %>%
  #Standardize psychopathology scores
  mutate(across(pfactor:EXT,scale)) %>%
```

```
# Are there any rsfmri images in ABCC?
  mutate(any_rest = (Exists_rest_1 |
                     Exists_rest_2
                     Exists_rest_3
                     Exists_rest_4)) %>%
  #Are there >375 low motion frames per threshold
  mutate(across(starts_with("gframes"),\(x) \{x>375\},.names = "\{.col\}_375")) \%%
  mutate(across(ends with("375"),~replace na(.,FALSE)))
imgincl<-readRDS("ABCD-fastqc01/imginclusion.Rds")</pre>
df<-left_join(df,imgincl[,1:2],by=join_by(subjectkeychar==subjectkey)) %>%
  mutate(fastqcok=replace_na(fastqcok,FALSE))
# N.B. A few (8) Pt's have neuroimaging data but not tabular data.
# Possibly they have removed consent? They are not included in this
# analysis
options(tinytex.clean = FALSE)
# Seed for random number generation
set.seed(42)
knitr::opts chunk$set(cache=TRUE)
knitr::opts chunk$set(cache.extra = knitr::rand seed)
```

## 1.1 Additional Coding and Releveling

```
cvars <- c(
   "adi",
   "coi",
   "nihtbx_flanker_agecorrected",
   "nihtbx_cryst_agecorrected",
   "nihtbx_totalcomp_agecorrected",
   "pea_wiscv_tss",
   "pfactor",
   "INT",
   "EXT",
   "interview_age",
   "bmi"
)

# Z scores
df <- df %>% mutate(across(ends_with(cvars),scale,.names = "{.col}.z"))
```

```
# Relevel factors to set comparison to modal levels

df$p.edu<-relevel(factor(df$p.edu,ordered=FALSE),4)
df$household.income<-relevel(df$household.income,5)</pre>
```

## 1.2 Descriptives

1.2.1 Figure 1: Condition Descriptions, inclusion counts, and inconsistencies

Figure 1 panel b:

```
# This code also generates table 1
levelsdf<-data.frame(</pre>
  Level = c(
    "Full (F)",
    "ABCD 4 Tabulated (T)",
    "ABCC (C)",
    "ABCC < .5mm",
   "ABCC < .4mm",
   "ABCC < .3mm",
    "ABCD 4 Recommended (R)",
    "ABCD < .2mm",
    "ABCD < .1mm"
  ),
  Description = c(
    "Entire sample",
    "rsFMRI tabulated data available (at least 1 T1 and 1 rsFMRI were complete and passed visual inspection)",
    "rs-fMRI data in the ABCD Community Collection",
    "ABCC + censoring at threshold*",
    "\"",
    "\"",
```

```
"T1 and rs-fMRI recommended in 'ABCD Recommended Imaging Inclusion' table (includes 375 frames at <.2mm FD)",
    "ABCC + censoring at .2mm threshold",
    11 \ 11 11
  ),
  Count = c(
    nrow(df),
    sum(df$ABCD_rsfmri_QC1),
    sum(df$any_rest),
    sum(df$gframes_0.5_375),
    sum(df$gframes_0.4_375),
    sum(df$gframes 0.3 375),
    sum(df$ABCD_rsfmri_QC2),
    sum(df$gframes 0.2 375),
    sum(df$gframes_0.1_375)
)
levelsdf$Levelf<-factor(levelsdf$Level,levels=c(</pre>
    "Full (F)",
    "ABCD 4 Tabulated (T)",
    "ABCC (C)",
    "ABCC < .5mm",
    "ABCC < .4mm",
    "ABCC < .3mm",
    "ABCD 4 Recommended (R)",
    "ABCD < .2mm",
    "ABCD < .1mm"
  ),
  labels = c(
    "Full (**F**)",
    "ABCD 4 Tabulated (**T**)",
    "ABCC (**C**)",
    "ABCC < **.5**mm",
    "ABCC < **.4**mm",
    "ABCC < **.3**mm",
    "ABCD 4 Recommended (**R**)",
    "ABCC < **.2**mm",
    "ABCC < **.1**mm"
  ))
library(ggtext)
samplecounts<-ggplot(levelsdf,aes(x=Levelf,y=Count,label=Count)) +</pre>
  labs(title="B. n by Condition") +
  geom_point(color="#c5050c",size=4,alpha=0.6) +
```

Figure 1 panel C:

(panel A is added using a vector graphics program.)

# 1.2.2 Categorical Variable Figure (Figure 2)

```
"gframes_0.2_375",
          "gframes_0.1_375")
catvars<-c("sex",
              "household.income",
              "p.edu",
              "race_ethnicity.factor",
              "ksads factor",
              "pds_category")
#This replaces NA with 'missing' for tabulation and reverses the scrubbing vars.
dfcatna <- df %>% select(c(ends_with(catvars), ends_with(bools))) %>%
  mutate(across(
    ends_with(catvars),
    ~ factor(
      .x,
      levels = levels(addNA(.x)),
      labels = c(levels(.x), "Missing"),
      exclude = NULL
    )
  ))
catvar375figdf<-data.frame(variable=character(),</pre>
                           level=character())
for (var in catvars) {
  levels <- levels(dfcatna[[var]])</pre>
  var df <- data.frame(</pre>
    variable = var,
    level = levels
  # Append the data frame to the levels_df
  catvar375figdf <- rbind(catvar375figdf, var_df)</pre>
}
dfcatna$full=TRUE
for(b in bools) {
  n<-sum(dfcatna[,b])</pre>
  v<-sapply(1:nrow(catvar375figdf),\(x) {</pre>
    vname<-catvar375figdf[x,"variable"]</pre>
    tlevel<-catvar375figdf[x,"level"]</pre>
    vmatch<-dfcatna[[vname]]==tlevel</pre>
    sum(vmatch & dfcatna[[b]])/n
```

```
})
 catvar375figdf[,b] < -v*100
catvar375figdflong <- catvar375figdf %>%
 pivot_longer(full:gframes_0.1_375) %>%
 mutate(thresh=factor(name,levels=bools,labels=c("F",
                                                   "C",
                                                   ".5".
                                                  ".4",
                                                  ".3",
                                                  "R",
                                                  ".2",
                                                  ".1"))) %>%
 mutate(levelf=as.factor(level)) %>%
 mutate(variable=as.factor(variable))
cp1<-catvar375figdflong %>%
 filter(variable=="sex") %>%
 mutate(level=as.factor(level)) %>%
 ggplot(aes(x=thresh,y=value,fill=level)) +
     geom_bar(position="fill",stat="identity") +
     scale fill viridis(discrete = TRUE) +
     labs(title="Sex Assigned at Birth") +
     theme(axis.title.y=element_blank(),
            axis.text.y=element blank(),
            axis.title.x=element blank(),
            axis.ticks.y=element blank(),
            legend.title=element_blank())
cp2<-catvar375figdflong %>%
 filter(variable=="household.income") %>%
 mutate(level=factor(level,levels=levels(dfcatna$household.income))) %>%
 ggplot(aes(x=thresh,y=value,fill=level)) +
     geom_bar(position="fill",stat="identity") +
     scale fill viridis(discrete = TRUE) +
     labs(title="Household Income") +
     theme(axis.title.y=element blank(),
            axis.text.y=element_blank(),
            axis.title.x=element blank(),
           axis.ticks.y=element_blank(),
            legend.title=element blank())
```

```
cp3<-catvar375figdflong %>%
 filter(variable=="race_ethnicity.factor") %>%
 mutate(level=factor(level,levels=levels(dfcatna$race_ethnicity.factor))) %%
  ggplot(aes(x=thresh,y=value,fill=level)) +
      geom bar(position="fill",stat="identity") +
      scale fill viridis(discrete = TRUE) +
      labs(title="Census Race/Ethnicity") +
      theme(axis.title.y=element_blank(),
            axis.text.y=element blank(),
            axis.title.x=element blank(),
            axis.ticks.y=element blank(),
            legend.title=element_blank())
cp4<-catvar375figdflong %>%
 filter(variable=="ksads_factor") %>%
  mutate(level=as.factor(level)) %>%
  ggplot(aes(x=thresh,y=value,fill=level)) +
      geom_bar(position="fill",stat="identity") +
      scale fill viridis(discrete = TRUE) +
      labs(title="Trauma Count") +
      theme(axis.title.y=element blank(),
            axis.text.y=element_blank(),
            axis.title.x=element blank(),
            axis.ticks.y=element blank(),
            legend.title=element blank())
cp5<-catvar375figdflong %>%
 filter(variable=="pds category") %>%
  mutate(level=factor(level,levels=levels(dfcatna$pds_category))) %>%
  ggplot(aes(x=thresh,y=value,fill=level)) +
      geom_bar(position="fill",stat="identity") +
      scale_fill_viridis(discrete = TRUE) +
      labs(title="Pubertal Status") +
      theme(axis.title.y=element_blank(),
            axis.text.y=element_blank(),
            axis.title.x=element blank(),
            axis.ticks.y=element_blank(),
            legend.title=element blank())
cp6<-catvar375figdflong %>%
 filter(variable=="p.edu") %>%
  mutate(level=factor(level,levels=levels(dfcatna$p.edu))) %>%
  ggplot(aes(x=thresh,y=value,fill=level)) +
```

```
svg(filename="Manuscript/Figures/CatVarFig.svg",height=9,width=7.5,family="Times New Roman")

((cp1 / cp4 / cp3) |
    (cp2 / cp6 / cp5))
grid::grid.draw(grid::textGrob("Proportion at Threshold",x=.01,rot=90,gp = grid::gpar(fontfamily = "Times New Roman")))
grid::grid.draw(grid::textGrob("QC Threshold",y=.01,gp = grid::gpar(fontfamily = "Times New Roman")))

dev.off()

## cairo_pdf
```

## 1.2.3 Continuous Variables Figure (Figure 3)

```
# we need ymin, ymax, lower, upper, and middle for each variable in each condition.
#we need ymin, ymax, lower, upper, and middle for each variable in each condition.
bpstats <- function(var,bool) {</pre>
  #message(pasteO("var is ",var))
  #message(pasteO("bool is ",bool))
 x<-df[df[[bool]],var]
 minvalue=min(x,na.rm=TRUE)
 maxvalue=max(x,na.rm=TRUE)
 median=median(x,na.rm = TRUE)
 low=quantile(x,.25,na.rm=TRUE)
 high=quantile(x,.75,na.rm=TRUE)
 IQR=high-low
 mn=max(low-1.5*IQR,minvalue)
 mx=min(high+1.5*IQR,maxvalue)
 return(data.frame(lwhisk=mn,low=low,median=median,high=high,uwhisk=mx))
cvars <- c(
    "adi.z".
```

```
"coi.z",
    "nihtbx_flanker_agecorrected.z",
    "nihtbx_cryst_agecorrected.z",
    "nihtbx_totalcomp_agecorrected.z",
    "pea_wiscv_tss.z",
    "pfactor.z",
   "INT.z",
    "EXT.z",
    "interview_age.z",
    "bmiAgeZ"
  df$full=TRUE
ContFig375df<-expand.grid(as.factor(cvars),unique(bools)) %>%
  mutate(Var1 = as.character(Var1),
         Var2 = as.character(Var2)) %>%
  rowwise() %>%
  mutate(bpstats(Var1,Var2))
ContFig375df$measp<-factor(ContFig375df$Var1,</pre>
                            levels=cvars,
                           labels=c("Area Disadvantage",
                                     "Child Opportunity",
                                     "NIHTB Flanker",
                                     "NIHTB Crystalized",
                                     "NIHTB Total",
                                     "WISC V Matrix",
                                     "Psychopathology (p)",
                                     "Internalizing",
                                     "Externalizing",
                                     "Age",
                                     "BMI"))
ContFig375df$thresh <- factor(</pre>
  ContFig375df$Var2,
  levels = bools,
  labels = c("F",
             "T",
             "C",
             ".5",
             ".4",
             ".3",
             "R",
```

```
".2",
             ".1")
ggsave("Manuscript/Figures/ContVarFig.svg",
       ggplot(
         ContFig375df,
         aes(
           x = thresh,
           ymin = lwhisk,
           lower = low,
           middle = median,
           upper = high,
           ymax = uwhisk
         geom_boxplot(stat = "identity") +
         ylab("Z") +
         theme(axis.title.x = element_blank()) +
         facet_wrap( ~ measp, scales = "free"),
       height = 5, width = 7.5)
```

#### 1.2.4 Trauma Count

We describe trauma count statistics in our methods:

7723, 3004, 1149

## 1.3 Table: Associations of behavioral data with framewise displacement (Table 1 and 2)

```
ccor<-df %>% select("adi.z",
    "coi.z",
    "nihtbx_flanker_agecorrected.z",
    "nihtbx_cryst_agecorrected.z",
    "nihtbx_totalcomp_agecorrected.z",
    "pea_wiscv_tss.z",
    "pfactor.z",
    "INT.z",
    "EXT.z",
    "interview_age.z",
    "bmiAgeZ",
    filtered_fd_0.1:filtered_fd_0.5) %>%
```

```
sdlabFunctions::sdcorr()
#This is the version in the manuscript.
write.csv(ccor[1:11,19:30], "Manuscript/Tables/FDcorr_continuousDVs2.csv", row.names=TRUE)
# This will write tables of bivariate linear models of FD with both continuous and discrete variables::
# Updated colnames for vars
y_names = c("filtered_fd_0.1", "filtered_fd_0.2", "filtered_fd_0.3", "filtered_fd_0.4", "filtered_fd_0.5")
x_names = c("adi", "coi", "nihtbx_flanker_agecorrected", "nihtbx_cryst_agecorrected", "nihtbx_totalcomp_agecorrected",
             "pea_wiscv_tss", "pfactor", "INT", "EXT", "interview_age", "bmiAgeZ", "sex", "household.income", "p.edu",
             "race_ethnicity.factor", "ksads_factor", "pds_category")
contOut = data.frame("IV"=c(), "DV"=c(), "BETA"=c(), "STD BETA"=c(), "R SQU"=c(), "T"=c(), "P"=c())
discOut = data.frame("IV"=c(), "DV"=c(), "F"=c(), "DF_1"=c(), "DF_2"=c(), "MODEL_P"=c(), "CONTRAST"=c(), "BETA"=c(), "SE"=c(), "T-RATIO"=c(), "P"=c())
# Iterate dependent variables (motion thresholds)
for(y_name in y_names){
  # Iterate independent variables
  for(x_name in x_names){
    # Create dataframe with only variables of interest, remove NAs
    temp df = na.omit(data.frame(y name=df[y name], x name=df[x name]))
    # For continuous predictors:
    if( !(x name %in% c("sex", "household.income", "p.edu", "race ethnicity.factor", "ksads factor", "pds category"))){
      # Scale
      temp df[x name] = scale(temp df[x name],center=TRUE,scale=TRUE)
      # Build a model
      mod = lm(data=temp_df,paste0(y_name,"~",x_name))
      modTemp = summary(mod)
      # Append statistics
      contOut = rbind(contOut,data.frame(
        "IV"=y_name, "DV"= x_name, "BETA"=modTemp$coefficients[2], "STD_BETA"=as.numeric(QuantPsyc::lm.beta(mod)), "R_SQU"=modTemp$r.squared,
        "T"=modTemp$coefficients[6], "P"=modTemp$coefficients[8]
      ))
    }
    # For discrete predictors:
    else{
```

```
# Build a model
      mod = lm(data=temp_df,paste0(y_name,"~",x_name))
      modTemp = summary(mod)
      \#post\_hoc = data.frame(emmeans::emmeans(mod,reformulate(x\_name, 'pairwise'))$contrasts)[,c(1,2,3,5,6)]
      post hoc = data.frame(emmeans::emmeans(mod,reformulate(x name, 'trt.vs.ctrl1'))$contrasts)[,c(1,2,3,5,6)]
      # Format and append statistics
      colnames(post_hoc) = c("CONTRAST", "BETA", "SE", "T-RATIO", "P")
      mod stats = data.frame("IV"=rep(y name,nrow(post hoc)),
                              "DV"=rep(x name, nrow(post hoc)),
                              "F"=rep(as.numeric(modTemp$fstatistic[1]),nrow(post hoc)),
                             "DF_1"=rep(as.numeric(modTemp$fstatistic[2]),nrow(post_hoc)),
                             "DF_2"=rep(modTemp$fstatistic[3],nrow(post_hoc)),
                              "MODEL_P"=rep(pf(as.numeric(modTemp\fstatistic[1]),as.numeric(modTemp\fstatistic[2]),
                                                          as.numeric(modTemp$fstatistic[3]),lower.tail = FALSE),nrow(post_hoc)))
      discOut = rbind(discOut,cbind(mod_stats,post_hoc))
    }
  }
}
sdfd <- t(df %>%
  summarize(
    sd fd 0.1 = sd(filtered fd 0.1, na.rm = TRUE),
    sd fd 0.2 = sd(filtered fd 0.2, na.rm = TRUE),
    sd_fd_0.3 = sd(filtered_fd_0.3, na.rm = TRUE),
    sd fd 0.4 = sd(filtered fd 0.4, na.rm = TRUE),
    sd_fd_0.5 = sd(filtered_fd_0.5, na.rm = TRUE)
  ))
discOut$fdsd<-rep(sdfd,each=sum(discOut$IV=="filtered_fd_0.1"))</pre>
discOut$sBeta<-discOut$BETA/discOut$fdsd
# Write tables to csv This continuous table did not end up included in the manuscirpt.
write.csv(contOut, "Manuscript/Tables/FDcorr continuousDVs.csv", row.names=FALSE)
write.csv(discOut, "Manuscript/Tables/FDcorr_discreteDVs.csv", row.names=FALSE)
```

In the manuscript, F tests associated with the discrete variables tested are omitted for brevity. These are printed here:

```
discOut %>%
  filter(IV %in% c("filtered_fd_0.1","filtered_fd_0.2","filtered_fd_0.3")) %>%
  select(DV:MODEL_P) %>%
  distinct() %>%
```

```
as_hux() %>%
insert_column(c("Threshold",rep(c(".1mm",".2mm",".3mm"),each=6))) %>%
set_caption("F tests for relation of discrete DVs with FD after scrubbing, by threshold (pt. 1)")

discOut %>%
filter(IV %in% c("filtered_fd_0.4","filtered_fd_0.5")) %>%
select(DV:MODEL_P) %>%
distinct() %>%
as_hux() %>%
insert_column(c("Threshold",rep(c(".4mm",".5mm"),each=6))) %>%
set_caption("F tests for relation of discrete DVs with FD after scrubbing, by threshold (pt. 2)")
```

#### 1.4 Bivariate Models and Table

```
ORformat<-\(x) {DescTools::Format(exp(x),digits=2,ldigits=0)}</pre>
pformat<-\(x) {DescTools::Format(x,digits=3,ldigits=0,sci=30)}</pre>
varstring <-</pre>
  c(
    "sex",
    "household.income",
    "p.edu",
    "race ethnicity.factor",
    "ksads factor",
    "pds_category",
    "adi.z",
    "coi.z",
    "nihtbx_flanker_agecorrected.z",
    "nihtbx_cryst_agecorrected.z",
    "nihtbx_totalcomp_agecorrected.z",
    "pea_wiscv_tss.z",
    "pfactor.z",
    "INT.z",
    "EXT.z",
    "interview_age.z",
    "bmiAgeZ"
bvmodel \langle - \setminus (x, newline="\setminus n")  {
  cstrings <-
```

Table 1: F tests for relation of discrete DVs with FD after scrubbing, by threshold (pt. 1)

Threshold	DV	F	DF_1	$DF_2$	$MODEL_P$
0.1mm	sex	40.6	1	9.37e + 03	1.99e-10
0.1mm	household.income	20	5	8.6e + 03	6.39e-20
0.1mm	p.edu	17.9	4	9.36e + 03	1.3e-14
0.1mm	$race\_ethnicity.factor$	27.5	4	9.36e + 03	9.06e-23
0.1mm	ksads_factor	4.61	2	9.37e + 03	0.00996
$0.1 \mathrm{mm}$	$pds\_category$	2.36	4	9.29e + 03	0.0512
$0.2 \mathrm{mm}$	sex	88	1	9.58e + 03	7.92e-21
$0.2 \mathrm{mm}$	household.income	27.5	5	8.79e + 03	9.8e-28
$0.2 \mathrm{mm}$	p.edu	27.7	4	9.56e + 03	6.43e-23
$0.2 \mathrm{mm}$	$race\_ethnicity.factor$	32.9	4	9.57e + 03	2.95e-27
$0.2 \mathrm{mm}$	$ksads\_factor$	6.75	2	9.57e + 03	0.00118
$0.2 \mathrm{mm}$	$pds\_category$	1.42	4	9.49e + 03	0.224
0.3mm	sex	116	1	9.59e + 03	7.92e-27
0.3mm	household.income	29.5	5	8.8e + 03	8.34e-30
0.3mm	p.edu	31.7	4	9.57e + 03	2.97e-26
0.3mm	$race\_ethnicity.factor$	31.5	4	9.58e + 03	4.18e-26
0.3mm	ksads_factor	6.94	2	9.59e + 03	0.000977
$0.3 \mathrm{mm}$	pds_category	0.843	4	9.5e + 03	0.498

Table 2: F tests for relation of discrete DVs with FD after scrubbing, by threshold (pt. 2)

Threshold	DV	$\mathbf{F}$	DF_1	$DF_2$	$MODEL_P$
$0.4 \mathrm{mm}$	sex	128	1	9.59e + 03	2.21e-29
$0.4 \mathrm{mm}$	household.income	29.5	5	8.8e + 03	8.32e-30
$0.4 \mathrm{mm}$	p.edu	32.7	4	9.58e + 03	3.97e-27
$0.4 \mathrm{mm}$	race_ethnicity.factor	28.8	4	9.58e + 03	8.03e-24
$0.4 \mathrm{mm}$	ksads_factor	7.06	2	9.59e + 03	0.000859
$0.4 \mathrm{mm}$	pds_category	1.15	4	9.5e + 03	0.331
$0.5 \mathrm{mm}$	sex	136	1	9.59e + 03	3.02e-31
$0.5 \mathrm{mm}$	household.income	28.8	5	8.8e + 03	4.6e-29
$0.5 \mathrm{mm}$	p.edu	33.5	4	9.58e + 03	7.92e-28
$0.5 \mathrm{mm}$	race_ethnicity.factor	26.5	4	9.59e + 03	6.25 e-22
$0.5 \mathrm{mm}$	ksads_factor	7	2	9.59e + 03	0.000914
$0.5 \mathrm{mm}$	$pds\_category$	1.12	4	9.51e + 03	0.347

```
"!ABCD_rsfmri_QC1",
    "!any_rest",
    "!gframes_0.5_375",
    "!gframes_0.4_375",
    "!gframes_0.3_375",
    "!ABCD rsfmri QC2",
    "!gframes_0.2_375",
    "!gframes 0.1 375"
  )
model.list<-sapply(cstrings,\(y) glm(as.formula(paste0(y," ~ ",x)),data=df,family="binomial"),simplify=FALSE)</pre>
sapply(model.list,
       \(x) {
         lf<-length(coef(x))</pre>
         if(lf==2) {
           table<-t(c(coef=coef(x)[2:1f],
                       confint(x,2:lf,level=.95),
                       p=coef(summary(x))[2:lf,4]))
         } else {
           table<-cbind(coef=coef(x)[2:lf],</pre>
                       confint(x,2:1f,level=.95),
                       p=coef(summary(x))[2:1f,4])
         }
         #Bonferonni correction
         if(names(coef(x))[2] %in% c("adi.z",
                                        "coi.z",
                                        "nihtbx_totalcomp_agecorrected.z",
                                       "pea_wiscv_tss.z")) {
            table[,4] \leftarrow table[,4] *2
         m<-paste0(ORformat(table[,1]),</pre>
                                sdlabFunctions::starPs(table[,4]),
                                newline,
                                ORformat(table[,2]),
                                ORformat(table[,3]),
                                newline,
                                pformat(table[,4]))
         names(m) <-names(coef(x)[2:lf])</pre>
         sub(".000","<.001",m)</pre>
  },
  simplify=TRUE)
```

```
bvtable<-reduce(sapply(varstring,bvmodel),rbind)</pre>
alllevels<-c(
  "Sex (Male)",
     " $0-$25k",
     " $25-$50k",
     " $50-$75k",
     " $75-$100k",
      " >$200k",
     " <High School",
     " HS Grad.",
     " Some College",
      " Graduate",
     " Black",
      " Hispanic",
      " Asian",
     " Other",
     " 1 Trauma",
     " >=2 Trauma",
     " Early Puberty",
     " Mid Puberty",
     " Late Puberty",
     " Post Pubertal",
     "Area Disadvantage",
     "Child Opportunity",
     "NIHTB Flanker",
     "NIHTB Crystalized",
     "NIHTB Total",
     "WISC V Matrix",
     "Psychopathology",
      "Internalizing",
     "Externalizing",
      "Age",
      "BMI"
)
#1 column per level
bvhux <- as_hux(bvtable) %>%
 insert_column(alllevels) %>%
 insert_row(
   "Household Income (ref: $100-$200k)",
   after = 1,
   colspan = 9,
```

```
fill = ""
  ) %>%
  insert_row(
    "Highest Parental Education (ref: College Degree)",
    after = 7,
    colspan = 9,
    fill = ""
  ) %>%
  insert row(
    "Census Race/Ethnicity (ref: White)",
    after = 12,
    colspan = 9,
   fill = ""
  ) %>%
  insert_row(
    "KSADS Trauma Count (ref: 0 Exposures)",
    after = 17,
    colspan = 9,
    fill = ""
  ) %>%
  insert_row(
    "Pubertal Status (ref: pre-pubertal)",
    after = 20,
    colspan = 9,
   fill = ""
  ) %>%
  insert_row(c("Variable", rep(c("OR\n[90% CI]\np"), 8)), after = 0) %>%
  insert_row(c("", "T", "C", ".5", ".4", ".3", "R", ".2", ".1"))
tables.xlsx<-openxlsx::createWorkbook()
tables.xlsx<-as_Workbook(bvhux[1:20,],
                     Workbook=tables.xlsx,
                     sheet="bvmPage1")
tables.xlsx<-as_Workbook(bvhux[c(1:2,21:nrow(bvhux)),],
                     Workbook=tables.xlsx,
                     sheet="bvmPage2")
```

## 1.5 Condition (Adjusted) Models

```
#terms 22 and 23 (neighborhood factors), as well as term 26 and 27 (general cognition measures) need to be corrected.
pcorrect<-\(x) {
   x[22:23]<-x[22:23]*2</pre>
```

```
x[26:27] < -x[26:27] *2
  x
adjmodelrightterms<-"~ sex + household.income + p.edu + race ethnicity.factor +
ksads_factor + pds_category + adi.z + coi.z + nihtbx_flanker_agecorrected.z +
nihtbx cryst agecorrected.z + nihtbx totalcomp agecorrected.z +
pea_wiscv_tss.z + pfactor.z + INT.z + EXT.z + interview_age.z + bmiAgeZ"
model.QC1 <-
  glm(as.formula(paste("!ABCD_rsfmri_QC1 ", adjmodelrightterms)),
      data = df, family = "binomial")
model.QC1.results <-</pre>
  cbind(confint(model.QC1, level = .95), model.QC1$coefficients)[, c(1, 3, 2)]
model.QC1.results <-</pre>
  cbind(model.QC1.results, pcorrect(coef(summary(model.QC1))[, 4]))
model.ABCC <-
  glm(as.formula(paste("!any_rest ", adjmodelrightterms)), data = df, family =
        "binomial")
model.ABCC.results <-</pre>
  cbind(confint(model.ABCC, level = .95), model.ABCC$coefficients)[, c(1, 3, 2)]
model.ABCC.results <-
  cbind(model.ABCC.results, pcorrect(coef(summary(model.ABCC))[, 4]))
model.5mm <-
  glm(as.formula(paste("!gframes 0.5 375", adjmodelrightterms)), data = df, family =
        "binomial")
model.5mm.results <-</pre>
  cbind(confint(model.5mm, level = .95), model.5mm$coefficients)[, c(1, 3, 2)]
model.5mm.results <-</pre>
  cbind(model.5mm.results, pcorrect(coef(summary(model.5mm))[, 4]))
model.4mm <-
  glm(as.formula(paste("!gframes_0.4_375", adjmodelrightterms)), data = df, family =
        "binomial")
model.4mm.results <-</pre>
  cbind(confint(model.4mm, level = .95), model.4mm$coefficients)[, c(1, 3, 2)]
model.4mm.results <-
  cbind(model.4mm.results, pcorrect(coef(summary(model.4mm))[, 4]))
model.3mm <-
  glm(as.formula(paste("!gframes_0.3_375", adjmodelrightterms)), data = df, family =
        "binomial")
model.3mm.results <-</pre>
```

```
cbind(confint(model.3mm, level = .95), model.3mm$coefficients)[, c(1, 3, 2)]
model.3mm.results <-</pre>
  cbind(model.3mm.results, pcorrect(coef(summary(model.3mm))[, 4]))
model.QC2 <-
  glm(as.formula(paste("!ABCD rsfmri QC2 ", adjmodelrightterms)), data = df, family =
        "binomial")
model.QC2.results <-</pre>
  cbind(confint(model.QC2, level = .95), model.QC2$coefficients)[, c(1, 3, 2)]
model.QC2.results <-</pre>
  cbind(model.QC2.results, pcorrect(coef(summary(model.QC2))[, 4]))
model.2mm <-
  glm(as.formula(paste("!gframes_0.2_375", adjmodelrightterms)), data = df, family =
        "binomial")
model.2mm.results <-</pre>
  cbind(confint(model.2mm, level = .95), model.2mm$coefficients)[, c(1, 3, 2)]
model.2mm.results <-
  cbind(model.2mm.results, pcorrect(coef(summary(model.2mm))[, 4]))
model.1mm <-
  glm(as.formula(paste("!gframes 0.1 375 ", adjmodelrightterms)), data = df, family =
        "binomial")
model.1mm.results <-</pre>
  cbind(confint(model.1mm, level = .95), model.1mm$coefficients)[, c(1, 3, 2)]
model.1mm.results <-
  cbind(model.1mm.results, pcorrect(coef(summary(model.1mm))[, 4]))
```

#### 1.5.1 Adjusted Model Tables

```
Mformat(model.4mm.results),
                    Mformat(model.3mm.results),
                    Mformat(model.QC2.results),
                    Mformat(model.2mm.results),
                    Mformat(model.1mm.results))
h1modelstable[h1modelstable==".000"]<-"<.001"
alllevels<-c("Intercept",
     "Sex (Male)",
      " $0-$25k",
      " $25-$50k",
      " $50-$75k",
      " $75-$100k",
     " >$200k",
     " <High School",
     " HS Grad.",
      " Some College",
      " Graduate",
      " Black",
      " Hispanic",
     " Asian",
     " Other",
     " 1 Trauma",
     " >=2 Trauma",
     " Early Puberty",
     " Mid Puberty",
     " Late Puberty",
     " Post Pubertal",
     "Area Disadvantage",
     "Child Opportunity",
      "NIHTB Flanker",
      "NIHTB Crystalized",
      "NIHTB Total",
     "WISC V Matrix",
     "Psychopathology",
     "Internalizing",
      "Externalizing",
      "Age",
      "BMI"
     )
#1 column per level
h1hux <- as_hux(h1modelstable) %>%
```

```
insert_column(alllevels) %>%
  insert_row(
    "Household Income (ref: $100-$200k)",
    after = 2,
    colspan = 9,
    fill = ""
  ) %>%
  insert row(
    "Highest Parental Education (ref: College Degree)",
    after = 8,
   colspan = 9,
   fill = ""
  ) %>%
  insert_row(
    "Census Race/Ethnicity (ref: White)",
    after = 13.
    colspan = 9,
   fill = ""
  ) %>%
  insert_row(
    "KSADS Trauma Count (ref: 0 Exposures)",
    after = 18,
   colspan = 9,
   fill = ""
  ) %>%
  insert row(
   "Pubertal Status (ref: pre-pubertal)",
    after = 21,
    colspan = 9,
   fill = ""
  ) %>%
  insert_row(c("Variable", rep(c("OR\n[90% CI]\np"), 8)), after = 0) %>%
  insert_row(c("", "T", "C", ".5", ".4", ".3", "R", ".2", ".1"))
colnames(h1hux)[1]<-"h1"</pre>
tables.xlsx<-as_Workbook(h1hux[1:20,],
                     Workbook=tables.xlsx,
                     sheet="AdjustedModels1")
tables.xlsx<-as_Workbook(h1hux[c(1:2,21:nrow(h1hux)),],
                     Workbook=tables.xlsx,
                     sheet="AdjustedModels2")
```

## 1.5.2 Site and MRI Effects – sensitivity analysis.

These are run only in some conditions: ABCC, QC2, 2mm

```
model.ABCC.scanner<-glm(</pre>
  !any_rest ~ sex + household.income + p.edu + race_ethnicity.factor +
    ksads_factor + pds_category + adi.z + coi.z + nihtbx_flanker_agecorrected.z
    + nihtbx_cryst_agecorrected.z + nihtbx_totalcomp_agecorrected.z +
    pea_wiscv_tss.z + pfactor.z + INT.z + EXT.z + interview_age.z + bmiAgeZ +
    mri model,
  data=df,
  familv="binomial")
model.ABCC.scanner.results<-cbind(confint(model.ABCC.scanner,level=.95),model.ABCC.scanner$coefficients)[,c(1,3,2)]
model.ABCC.scanner.results<-cbind(model.ABCC.scanner.results,p.adjust(coef(summary(model.ABCC.scanner))[,4],method="fdr"))
model.ABCC.scanner.site<-glm(</pre>
  !any_rest ~ sex + household.income + p.edu + race_ethnicity.factor +
    ksads_factor + pds_category + adi.z + coi.z + nihtbx_flanker_agecorrected.z
    + nihtbx_cryst_agecorrected.z + nihtbx_totalcomp_agecorrected.z +
    pea_wiscv_tss.z + pfactor.z + INT.z + EXT.z + interview_age.z + bmiAgeZ +
    mri_model + site_id_l,
  data=df,
  family="binomial")
model.ABCC.scanner.site.results<-cbind(confint(model.ABCC.scanner.site,level=.95),model.ABCC.scanner.site$coefficients)[,c(1,3,2)]
model.ABCC.scanner.site.results<-cbind(model.ABCC.scanner.site.results,p.adjust(coef(summary(model.ABCC.scanner.site))[,4],method="fdr"))
model.2mm.scanner<-glm(
  !gframes 0.2 375 ~ sex + household.income + p.edu + race ethnicity.factor +
    ksads factor + pds category + adi.z + coi.z + nihtbx flanker agecorrected.z
    + nihtbx cryst agecorrected.z + nihtbx totalcomp agecorrected.z +
    pea_wiscv_tss.z + pfactor.z + INT.z + EXT.z + interview_age.z + bmiAgeZ +
    mri model,
  data=df,
  family="binomial")
model.2mm.scanner.results<-cbind(confint(model.2mm.scanner,level=.95),model.2mm.scanner$coefficients)[,c(1,3,2)]
model.2mm.scanner.results<-cbind(model.2mm.scanner.results,p.adjust(coef(summary(model.2mm.scanner))[,4],method="fdr"))
model.2mm.scanner.site<-glm(</pre>
  !gframes_0.2_375 ~ sex + household.income + p.edu + race_ethnicity.factor +
    ksads_factor + pds_category + adi.z + coi.z + nihtbx_flanker_agecorrected.z
    + nihtbx_cryst_agecorrected.z + nihtbx_totalcomp_agecorrected.z +
    pea_wiscv_tss.z + pfactor.z + INT.z + EXT.z + interview_age.z + bmiAgeZ +
    mri_model + site_id_l,
  data=df.
  family="binomial")
```

```
model.2mm.scanner.site.results<-cbind(confint(model.2mm.scanner.site,level=.95),model.2mm.scanner.site$coefficients)[,c(1,3,2)]
model.2mm.scanner.site.results<-cbind(model.2mm.scanner.site))[,4],method="fdr"))
model.QC2.scanner<-glm(</pre>
  !ABCD rsfmri QC2 ~ sex + household.income + p.edu + race ethnicity.factor +
    ksads factor + pds category + adi.z + coi.z + nihtbx flanker agecorrected.z
   + nihtbx cryst agecorrected.z + nihtbx totalcomp agecorrected.z +
    pea wiscv tss.z + pfactor.z + INT.z + EXT.z + interview age.z + bmiAgeZ +
    mri model,
 data=df.
 family="binomial")
model.QC2.scanner.results<-cbind(confint(model.QC2.scanner,level=.95),model.QC2.scanner$coefficients)[,c(1,3,2)]
model.QC2.scanner.results<-cbind(model.QC2.scanner.results,p.adjust(coef(summary(model.QC2.scanner))[,4],method="fdr"))
model.QC2.scanner.site<-glm(</pre>
  !ABCD_rsfmri_QC2 ~ sex + household.income + p.edu + race_ethnicity.factor +
    ksads_factor + pds_category + adi.z + coi.z + nihtbx_flanker_agecorrected.z
   + nihtbx_cryst_agecorrected.z + nihtbx_totalcomp_agecorrected.z +
    pea_wiscv_tss.z + pfactor.z + INT.z + EXT.z + interview_age.z + bmiAgeZ +
    mri_model + site_id_l,
 data=df,
 family="binomial")
model.QC2.scanner.site.results<-cbind(confint(model.QC2.scanner.site,level=.95), model.QC2.scanner.site$coefficients)[,c(1,3,2)]
model.QC2.scanner.site.results<-cbind(model.QC2.scanner.site))[,4],method="fdr"))
model.ABCC.site<-glm(</pre>
  !any rest ~ sex + household.income + p.edu + race ethnicity.factor +
    ksads_factor + pds_category + adi.z + coi.z + nihtbx_flanker_agecorrected.z
   + nihtbx cryst agecorrected.z + nihtbx totalcomp agecorrected.z +
    pea_wiscv_tss.z + pfactor.z + INT.z + EXT.z + interview_age.z + bmiAgeZ +
    site_id_l,
 data=df,
 family="binomial")
model.ABCC.site.results<-cbind(confint(model.ABCC.site,level=.95),model.ABCC.site$coefficients)[,c(1,3,2)]
model.ABCC.site.results<-cbind(model.ABCC.site.results,p.adjust(coef(summary(model.ABCC.site))[,4],method="fdr"))
model.2mm.site<-glm(</pre>
  !gframes_0.2_375 ~ sex + household.income + p.edu + race_ethnicity.factor +
   ksads factor + pds category + adi.z + coi.z + nihtbx flanker agecorrected.z
    + nihtbx_cryst_agecorrected.z + nihtbx_totalcomp_agecorrected.z +
    pea wiscv tss.z + pfactor.z + INT.z + EXT.z + interview age.z + bmiAgeZ +
    site id 1,
  data=df,
 family="binomial")
```

1.5.2.1 Supplementary Tables This is a table of the model output for just the scanner terms, with and without site control, which we reference in the text:

```
MformatCI <- \(m) {</pre>
  stars <- sdlabFunctions::starPs(m[, 4])</pre>
  cbind(
    paste0(
      ORformat(m[, 2]),
      stars,
      " [".
      ORformat(m[, 1]),
      ORformat(m[, 3]),
      "]"))
}
sitecoefs<-rbind(cbind(MformatCI(model.ABCC.scanner.results)[33:36,],</pre>
      MformatCI(model.QC2.scanner.results)[33:36.].
      MformatCI(model.2mm.scanner.results)[33:36,]),
      cbind(MformatCI(model.ABCC.scanner.site.results)[33:36,],
      MformatCI(model.QC2.scanner.site.results)[33:36,],
      MformatCI(model.2mm.scanner.site.results)[33:36,]))
as huxtable(sitecoefs) %>%
  insert_row(c("ABCC", "QC2", ".2mm")) %>%
  insert row(c("Without Site Control", "", "")) %>%
  merge_cells(1, 1:3) %>%
  merge cells(2, 1:3) %>%
  insert_row(c("With Site Control","",""),after = 6) %>%
  merge cells(7,1:3) %>%
  insert column(
```

```
c(
   "",
   "Scanner (ref: Philips Achieva)",
   "GE Discovery MR750",
   "Philips Ingenia",
   "Siemens Prisma",
   "Siemens Prisma fit",
   "",
   "GE Discovery MR750",
   "Philips Ingenia",
   "Siemens Prisma",
   "Siemens Prisma",
   "Siemens Prisma",
   "Siemens Prisma fit"
)
) %>%
set_caption("OR of scanner terms, with and without site control [95\\% CI]. ***: p<.001") %>%
set_wrap(TRUE) %>%
set_width(1)
```

Table 3: OR of scanner terms, with and without site control [95% CI]. \*\*\*: p<.001

	Without Site Control				
Scanner (ref: Philips Achieva)	ABCC				
GE Discovery MR750	.86 [.72–1.04]	.56*** [.4669]	.70*** [.59–.84]		
Philips Ingenia	.20*** [.1331]	.87 [.63–1.19]	.42*** [.3057]		
Siemens Prisma	.15*** [.1220]	.27*** [.2233]	.20*** [.1725]		
Siemens Prisma fit	.39*** [.3247]	.43*** [.3653]	.43*** [.36–.51]		
	With Site Control				
GE Discovery MR750	3894.69 [1.69e-14-NA]	44437.50 [.00-NA]	$13726.89 \; [.00\mathrm{-NA}]$		
Philips Ingenia	8698.06 [4.91e-09-NA]	7.24e+09 [5.44e-11-NA]	9.30e+09 [.01-NA]		
Siemens Prisma	3109.47 [1.16e-14-NA]	8613.49 [.00-NA]	$15297.36\ [.00\mathrm{-NA}]$		
Siemens Prisma fit	6030.51 [2.24e-14-NA]	8257.41 [.00-NA]	16617.52 [.00–NA]		

Without Site Control

1.5.2.2 Supplementary Forest Plots These forest plots show ORs with a significant (p<.05) model term in either the site-controlled or the site un-controlled adjusted model. They are here to visualize the effects of site control on the adjusted models.

```
ABCC.compdf <- rbind(
  model.ABCC.site.results %>%
    as.data.frame() %>%
    rownames_to_column("term") %>%
    rename_with( ~ c("term", "lowCI", "OR", "highCI", "p")) %>%
    mutate(
      lowCI = replace na(lowCI, min(lowCI, na.rm = TRUE)),
      highCI = replace na(highCI, max(highCI, na.rm = TRUE)),
      sig = p < .05,
      model = "Site Control"
    ) %>%
    mutate(across(lowCI:highCI, exp)),
  model.ABCC.results %>%
    as.data.frame() %>%
    rownames_to_column("term") %>%
    rename_with( ~ c("term", "lowCI", "OR", "highCI", "p")) %>%
    mutate(
      lowCI = replace_na(lowCI, min(lowCI, na.rm = TRUE)),
      highCI = replace_na(highCI, max(highCI, na.rm = TRUE)),
      sig = p < .05,
      model = "No Control"
    ) %>%
    mutate(across(lowCI:highCI, exp))
) %>%
  mutate(model = as.factor(model),
         alphaval=case when(sig ~ 1,
                            !sig ~ .5))
#terms where one model coefficient was sig
sigterms <-
  (ABCC.compdf %>% group_by(term) %>% summarise(anysig = sum(sig) != 0) %>% filter(anysig))$term
ABCC.compdf <-ABCC.compdf %>% mutate(alphaval=factor(alphaval,levels=c(.5,1),labels=c("p>.05","p<.05")))
test<-ggplot(ABCC.compdf %>% filter(term %in% sigterms),
       aes(x = term, color = model, alpha = alphaval)) +
  geom_point(aes(y = OR), position = position_dodge(.2)) +
  geom linerange(aes(ymin = lowCI, ymax = highCI, x = term),
                 position = position dodge(.2)) +
  ylim(-4, 4) +
  geom hline(lty = 2, aes(yintercept = 1), colour = 'black') +
  coord flip() + theme(legend.title = element blank())
```

```
QC2.compdf <- rbind(
  model.QC2.site.results %>%
    as.data.frame() %>%
    rownames_to_column("term") %>%
    rename with( ~ c("term", "lowCI", "OR", "highCI", "p")) %>%
    mutate(
      lowCI = replace na(lowCI, min(lowCI, na.rm = TRUE)),
      highCI = replace na(highCI, max(highCI, na.rm = TRUE)),
      sig = p < .05,
      model = "Site Control"
    ) %>%
    mutate(across(lowCI:highCI, exp)),
  model.QC2.results %>%
    as.data.frame() %>%
    rownames_to_column("term") %>%
    rename_with( ~ c("term", "lowCI", "OR", "highCI", "p")) %>%
    mutate(
      lowCI = replace_na(lowCI, min(lowCI, na.rm = TRUE)),
      highCI = replace_na(highCI, max(highCI, na.rm = TRUE)),
      sig = p < .05,
      model = "No Control"
    ) %>%
    mutate(across(lowCI:highCI, exp))
) %>%
  mutate(model = as.factor(model),
         alphaval=case when(sig ~ 1,
                            !sig ~ .5))
#terms where one model coefficient was sig
sigterms<-(QC2.compdf %>% group_by(term) %>% summarise(anysig=sum(sig)!=0) %>% filter(anysig))$term
QC2.compdf <-QC2.compdf %>% mutate(alphaval=factor(alphaval,levels=c(.5,1),labels=c("p>.05","p<.05")))
ggplot(QC2.compdf %>% filter(term %in% sigterms),
       aes(x = term, color = model, alpha = alphaval)) +
  geom_point(aes(y = OR), position = position_dodge(.2)) +
  geom_linerange(aes(ymin = lowCI, ymax = highCI, x = term),
                 position = position_dodge(.2)) +
  ylim(-4, 4) +
  geom hline(lty = 2, aes(yintercept = 1), colour = 'black') +
  coord flip() + theme(legend.title = element blank())
```

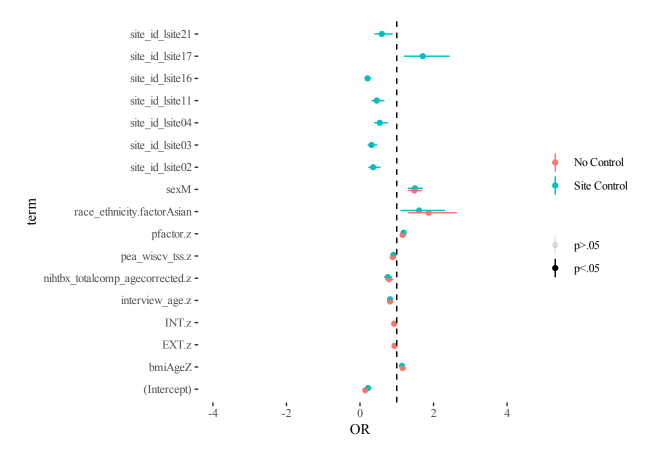


Figure 1: Significant ORs predicting DAIRC Recommended missingness with and without site control

```
twomm.compdf <- rbind(</pre>
  model.2mm.site.results %>%
    as.data.frame() %>%
    rownames_to_column("term") %>%
    rename_with( ~ c("term", "lowCI", "OR", "highCI", "p")) %>%
    mutate(
      lowCI = replace na(lowCI, min(lowCI, na.rm = TRUE)),
      highCI = replace na(highCI, max(highCI, na.rm = TRUE)),
      sig = p < .05,
      model = "Site Control"
    ) %>%
    mutate(across(lowCI:highCI, exp)),
  model.2mm.results %>%
    as.data.frame() %>%
    rownames_to_column("term") %>%
    rename_with( ~ c("term", "lowCI", "OR", "highCI", "p")) %>%
    mutate(
      lowCI = replace_na(lowCI, min(lowCI, na.rm = TRUE)),
      highCI = replace_na(highCI, max(highCI, na.rm = TRUE)),
      sig = p < .05,
      model = "No Control"
    ) %>%
    mutate(across(lowCI:highCI, exp))
) %>%
  mutate(model = as.factor(model),
         alphaval=case when(sig ~ 1,
                            !sig ~ .5))
#terms where one model coefficient was sig
sigterms <-
  (twomm.compdf %>% group_by(term) %>% summarise(anysig = sum(sig) != 0) %>% filter(anysig))$term
twomm.compdf<-twomm.compdf %>% mutate(alphaval=factor(alphaval,levels=c(.5,1),labels=c("p>.05","p<.05")))
ggplot(twomm.compdf %>% filter(term %in% sigterms),
       aes(x = term, color = model, alpha = alphaval)) +
  geom_point(aes(y = OR), position = position_dodge(.2)) +
  geom linerange(aes(ymin = lowCI, ymax = highCI, x = term),
                 position = position dodge(.2)) +
  ylim(-4, 4) +
  geom hline(lty = 2, aes(yintercept = 1), colour = 'black') +
  coord flip() + theme(legend.title = element blank())
```

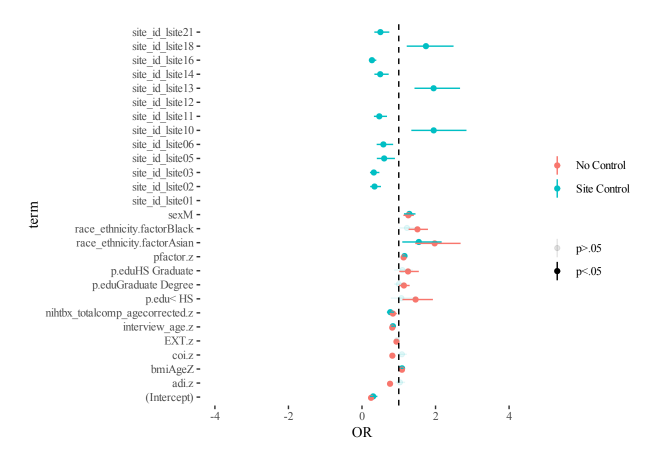


Figure 2: Significant ORs predicting .2mm missingness with and without site control

```
sitesummary <- df %>% group_by(site_id_1) %>%
  summarise(
    "ExcludedPerc.QC2" = mean(!ABCD_rsfmri_QC2),
    "ExcludedPerc.ABCC" = mean(!any rest),
   "ExcludedPerc.2mm" = mean(!gframes_0.2_375),
    "ExcludedPerc.fastqc" = mean(!fastqcok)
 ) %>%
 arrange(site_id_l) %>%
 mutate(
    sitelabel = recode factor(
      site id 1,
      `siteO1` = "Los Angeles, CA (CHLA)",
      `site02` = "Boulder, CO",
      `site03` = "Miami, FL",
      `site04` = "Tulsa, OK",
      `siteO5` = "Columbia, SC",
      `siteO6` = "Portland, OR",
      `site07` = "Rochester, NY",
      `site08` = "Menlo Park, CA",
      `site09` = "Los Angeles, CA (UCLA)",
      `site10` = "San Diego, CA",
      `site11` = "Gainsville, FL",
      `site12` = "Baltimore, MD",
      `site13` = "Ann Arbor, MI",
      `site14` = "Minneapolis, MN",
      `site15` = "Pittsburgh, Pa",
      `site16` = "Salt Lake City, UT",
      `site17` = "Burlington, VT",
      `site18` = "Milwaukee, WI",
      `site19` = "Richmond, VA",
      `site20` = "St. Louis, MO",
      `site21` = "New Haven, CT",
      `site22` = "New York, NY"
    )
 )
sitesummary2 <- sitesummary %>%
 mutate(sitelabel = fct_reorder(sitelabel, ExcludedPerc.QC2)) %>%
 pivot_longer(
   cols = starts with("ExcludedPerc"),
   names to = "sample",
   names prefix = "ExcludedPerc\\.",
   values to = "Percent Missing"
```

```
) %>%
 mutate(sample = factor(
    sample,
    levels = c("fastqc",
               "QC2",
               "ABCC",
               "2mm").
    labels = c("Current FastQC",
               "DAIC inclusion",
               "ABCC",
               "ABCC < .2mm FD")
 ))
sensplot1<-ggplot(sitesummary2,</pre>
       aes(x=sitelabel,y=Percent_Missing,fill=sample)) +
 geom_col(position="dodge") +
 theme(axis.title.x=element_blank(),
        legend.title= element_blank()) +
 vlab("% Excluded") +
 scale_x_discrete(guide = guide_axis(angle = 45)) +
 scale fill viridis(discrete=TRUE) +
 ggtitle("% Missing by site, inclusion criteria")
```

1.5.2.3 Figure 4: Missingness by site, inclusion criteria We reference the sites with the highest % exclusion, by site, in the text – they are given by this table.

Table 4: Sites with Highest % Excluded, by Condition

Site	Condition	% Excluded
Pittsburgh, Pa	DAIC inclusion	32.5
Pittsburgh, Pa	Current FastQC	30.3
Los Angeles, CA (CHLA)	$\mathrm{ABCC} < .2\mathrm{mm} \; \mathrm{FD}$	75.6
Baltimore, MD	ABCC	72.7

```
site id 1,
      `siteO1` = "Los Angeles, CA (CHLA)",
      `site02` = "Boulder, CO",
      `site03` = "Miami, FL",
      `site04` = "Tulsa, OK",
      `siteO5` = "Columbia, SC",
      `site06` = "Portland, OR",
      `site07` = "Rochester, NY",
      `site08` = "Menlo Park, CA",
      `site09` = "Los Angeles, CA (UCLA)",
      `site10` = "San Diego, CA",
      `site11` = "Gainsville, FL",
      `site12` = "Baltimore, MD",
      `site13` = "Ann Arbor, MI",
      `site14` = "Minneapolis, MN",
      `site15` = "Pittsburgh, Pa",
      `site16` = "Salt Lake City, UT",
      `site17` = "Burlington, VT",
      `site18` = "Milwaukee, WI",
      `site19` = "Richmond, VA",
      `site20` = "St. Louis, MO",
      `site21` = "New Haven, CT",
      `site22` = "New York, NY"
   )) %>%
 mutate(sitelabel = factor(sitelabel,levels=levels(sitesummary2$sitelabel)))
sensplot2<-ggplot(sitesummary3,</pre>
       aes(x=sitelabel,y=ExcludedN,fill=race_ethnicity.factor)) +
 geom_col(position="stack") +
 theme(axis.title.x=element_blank(),
       legend.title=element_blank()) +
 ylab("N Excluded") +
```

#### 1.6 H2 Model

This model specifically looks at changes associated with motion threshold choice in the ABCC data. In the pre-registration, we originally proposed a mixed effects logistic model where each participant had a case for each motion threshold, and the percentage of data missing from each threshold was used as a within-subject variable. In simulation, we discovered a number of problems with this approach:

- 1. The proposed mixed effects logistic regression created very poor predictions and model fits in simulation, and often did not converge. A linear model of probability proved more stable and accurate and we determined to use that approach (a logistic model is also presented below, but it is similarly flawed).
- 2. We proposed that an interaction between a variable and the 'pmiss' variable would indicate that bias was worsening as more data was excluded. However, in simulating this approach we discovered that in the case that there was a bias but it did not worsen as more data was excluded, as, for example, if 2 males were excluded for every female, this would surface as a linear interaction between pmiss and sex in the model. Instead, a worsening bias would be indicated by an interaction between a variable and a polynomial of pmiss, which would indicate a curvilinear trajectory.

(notes from the simulation are available on osf)

An important remaining question from the simulation study was whether to include an intercept and main effects in the model. As, at pmiss=0, the probability of exclusion ought to be 0 for everyone, one could argue the intercept should be omitted. We will try both methods and compare model fit.

```
names to="threshold",
                              values to="missing") %>%
  mutate(missing=!missing)
dflong$pmiss<-pmisssum[dflong$threshold]
# h2 model with intercepts.
m.h2 <- lmer(
  missing ~ poly(pmiss, 2) +
    sex + sex:poly(pmiss, 2) +
    household.income + household.income:poly(pmiss, 2) +
    p.edu + p.edu:poly(pmiss, 2) +
    race_ethnicity.factor + race_ethnicity.factor:poly(pmiss, 2) +
    ksads factor + ksads factor:poly(pmiss, 2) +
    pds_category + pds_category:poly(pmiss, 2) +
    adi.z + adi.z:poly(pmiss, 2) +
    coi.z + coi.z:poly(pmiss, 2) +
    nihtbx_flanker_agecorrected.z + nihtbx_flanker_agecorrected.z:poly(pmiss, 2) +
    nihtbx_cryst_agecorrected.z + nihtbx_cryst_agecorrected.z:poly(pmiss, 2) +
    nihtbx_totalcomp_agecorrected.z + nihtbx_totalcomp_agecorrected.z:poly(pmiss, 2) +
    pea_wiscv_tss.z + pea_wiscv_tss.z:poly(pmiss, 2) +
    pfactor.z + pfactor.z:poly(pmiss, 2) +
    INT.z + INT.z:poly(pmiss, 2) +
    EXT.z + EXT.z:poly(pmiss, 2) +
    interview_age.z + interview_age.z:poly(pmiss, 2) +
    bmiAgeZ + bmiAgeZ:poly(pmiss, 2) +
    (pmiss | subjectkey),
  data = dflong
# h2 model with intercepts removed
m.h2.stripped <- lmer(</pre>
  missing ~ pmiss + poly(pmiss, 2) +
    sex:poly(pmiss,2) +
    household.income:poly(pmiss,2) +
    p.edu:poly(pmiss,2) +
    race_ethnicity.factor:poly(pmiss,2) +
    ksads_factor:poly(pmiss,2) +
    pds_category:poly(pmiss,2) +
    adi.z:poly(pmiss,2) +
    coi.z:poly(pmiss,2) +
    nihtbx flanker agecorrected.z:poly(pmiss,2) +
    nihtbx cryst agecorrected.z:poly(pmiss,2) +
    nihtbx totalcomp agecorrected.z:poly(pmiss,2) +
```

```
pea_wiscv_tss.z:poly(pmiss,2) +
    pfactor.z:poly(pmiss,2) +
    INT.z:poly(pmiss,2) +
    EXT.z:poly(pmiss,2) +
    interview_age.z:poly(pmiss,2) +
    bmiAgeZ:poly(pmiss,2) +
    + 0 +
    (pmiss + 0 | subjectkey),
  data = dflong
# h2 model as a glm - this does not converge
# m.h2.qlm<- qlmer(
   missing ~ poly(pmiss, 2) +
      sex + sex:poly(pmiss, 2) +
      household.income + household.income:poly(pmiss, 2) +
      p.edu + p.edu:poly(pmiss, 2) +
      race_ethnicity.factor + race_ethnicity.factor:poly(pmiss, 2) +
      ksads_factor + ksads_factor:poly(pmiss, 2) +
      pds_category + pds_category:poly(pmiss, 2) +
      adi.z + adi.z:poly(pmiss, 2) +
      coi.z + coi.z:poly(pmiss, 2) +
#
#
      nihtbx flanker agecorrected.z + nihtbx flanker agecorrected.z:poly(pmiss, 2) +
#
      nihtbx_cryst_agecorrected.z + nihtbx_cryst_agecorrected.z:poly(pmiss, 2) +
#
      nihtbx totalcomp agecorrected.z + nihtbx totalcomp agecorrected.z:poly(pmiss, 2) +
#
      pea wiscv tss.z + pea wiscv tss.z:poly(pmiss, 2) +
#
      pfactor.z + pfactor.z:poly(pmiss, 2) +
#
      INT.z + INT.z:poly(pmiss, 2) +
      EXT.z + EXT.z:poly(pmiss, 2) +
#
      interview_age.z + interview_age.z:poly(pmiss, 2) +
      bmiAqeZ + bmiAqeZ:poly(pmiss, 2) +
      (pmiss | subjectkey),
   family="binomial",
   data = dflong,
   nAGQ=0.
    control=qlmerControl(optimizer="nloptwrap")
# )
performance::print md(performance::compare performance(m.h2,m.h2.stripped))
```

Table 5: Comparison of Model Performance Indices

Name	Model	AIC (weights)	AICc (weights)	BIC (weights)	R2 (cond.)	R2 (marg.)	ICC	RMSE	Sigma
m.h2 m.h2.stripped	${\bf lmerModLmerTest} \\ {\bf lmerModLmerTest}$	-1.8e+05 (>.999) 3.5e+05 (<.001)	-1.8e+05 (>.999) 3.5e+05 (<.001)	-1.8e+05 (>.999) 3.5e+05 (<.001)	0.91 0.51	0.19 0.28	0.89 0.31	$0.18 \\ 0.34$	$0.19 \\ 0.34$

The 'full' model has superior fit characteristics across indices. The GLM model, which is closer to what we originally proposed, performs poorly.

```
h2sum<-summary(m.h2)
temp = capture.output(h2sum)
End = grep("Fixed effects", temp)-1
print(unname(temp[0:End]))
    [1] "Linear mixed model fit by REML. t-tests use Satterthwaite's method ["
    [2] "lmerModLmerTest]"
    [3] "Formula: "
    [4] "missing ~ poly(pmiss, 2) + sex + sex:poly(pmiss, 2) + household.income + "
    [5] "
             household.income:poly(pmiss, 2) + p.edu + p.edu:poly(pmiss,
    [6] "
             2) + race_ethnicity.factor + race_ethnicity.factor:poly(pmiss, "
             2) + ksads_factor + ksads_factor:poly(pmiss, 2) + pds_category + "
    [7] "
    [8] "
             pds_category:poly(pmiss, 2) + adi.z + adi.z:poly(pmiss, 2) + "
             coi.z + coi.z:poly(pmiss, 2) + nihtbx_flanker_agecorrected.z + "
    [9] "
             nihtbx_flanker_agecorrected.z:poly(pmiss, 2) + nihtbx_cryst_agecorrected.z + "
## [10] "
## [11] "
             nihtbx cryst agecorrected.z:poly(pmiss, 2) + nihtbx totalcomp agecorrected.z + "
## [12] "
             nihtbx_totalcomp_agecorrected.z:poly(pmiss, 2) + pea_wiscv_tss.z + "
## [13] "
             pea wiscv tss.z:poly(pmiss, 2) + pfactor.z + pfactor.z:poly(pmiss, "
             2) + INT.z + INT.z:poly(pmiss, 2) + EXT.z + EXT.z:poly(pmiss, "
## [14] "
             2) + interview age.z + interview age.z:poly(pmiss, 2) + bmiAgeZ + "
## [15] "
             bmiAgeZ:poly(pmiss, 2) + (pmiss | subjectkey)"
## [16] "
## [17] "
            Data: dflong"
## [18] ""
## [19] "REML criterion at convergence: -176817.8"
## [20] ""
## [21] "Scaled residuals: "
   [22]
             Min
                      1Q Median
                                             Max "
                                      3Q
   [23] "-5.5260 -0.3032 0.0429 0.2260 5.0294 "
## [24] ""
## [25] "Random effects:"
## [26] " Groups
                                 Variance Std.Dev. Corr "
                     Name
## [27] " subjectkey (Intercept) 0.26699 0.5167
## [28] "
                     pmiss
                                 0.29719 0.5452
                                                   -0.94"
                                 0.03466 0.1862
## [29] " Residual
## [30] "Number of obs: 463488, groups: subjectkey, 9088"
## [31] ""
```

Table for the fixed effects:

```
pnames<-rownames(h2sum$coefficients)</pre>
h2table<-h2sum$coefficients %>% as.data.frame() %>%
  rownames_to_column(var = "Parameter") %>%
  left join(parameters::ci(m.h2)) %>%
  mutate(star = sdlabFunctions::starPs(`Pr(>|t|)`)) %>%
  select(c(Parameter, Estimate, CI_low, CI_high,star)) %>%
  mutate(level=c(1:3,rep(1,31),rep(2:3,31)),
         basep=c(rep("",3),pnames[4:34],rep(pnames[4:34],each=2))) \%
  pivot_wider(id_cols=basep,names_from=level,values_from = Estimate:star) %>%
  mutate(across(where(is.numeric).ORformat))
h2hux<-h2table[,c(1,2,5,8,11,3,6,9,12,4,7,10,13)] %>%
  as hux(add colnames=FALSE) %>%
  insert_row("","Intercept","","","","pmiss","","","",""pmiss^2","","","") %>%
  merge cells(1, 2:5) %>% merge cells(1, 6:9) %>% merge cells(1, 10:12) %>%
  insert_row("", "Main Eff.", "", "", "Interaction", "", "", "Q. Interaction", "", "", after=2) %>%
  merge_cells(3, 2:5) %>% merge_cells(3, 6:9) %>% merge_cells(3, 10:12) %>%
  insert_row("",rep(c("B","CI.1","CI.u","*"),3))
h2hux[,1] <- c(rep("",4),alllevels[-1])
h2hux <- h2hux %>% insert row(
    "Household Income (ref: $100-$200k)",
    after = 6,
    colspan = 9,
    fill = ""
  ) %>%
  insert row(
    "Highest Parental Education (ref: College Degree)",
    after = 11.
    colspan = 9,
   fill = ""
  ) %>%
  insert row(
    "Census Race/Ethnicity (ref: White)",
    after = 13+3,
    colspan = 9,
   fill = ""
  ) %>%
  insert row(
    "KSADS Trauma Count (ref: 0 Exposures)",
    after = 18+3.
```

```
colspan = 9,
  fill = ""
) %>%
insert_row(
   "Pubertal Status (ref: pre-pubertal)",
   after = 21+3,
   colspan = 9,
   fill = ""
)
h2hux[c(1:4,5:24),] %>% set_caption("H2 / Omnibus model parameters, pt. 1")
h2hux[c(1:4,25:nrow(h2hux)),] %>% set_caption("H2 / Omnibus model parameters, pt. 2")
```

## 1.7 H2 Marginal Means plotting (Figure 5)

```
# What ticks do we want for our plot?
xticks.mm<-c(.5,.45,.4,.35,.3,.25,.2,.15,.1,.05)
xticks.pmiss<-pmisssum[c("any_rest",paste0("gframes_",xticks.mm,"_375"))]</pre>
xticks.perc<-round(xticks.pmiss*100,0)</pre>
xticks.labela<-c("C",paste0(xticks.mm,"mm"))</pre>
xticks.label<-paste0(xticks.labela, "\n(", xticks.perc, "%)")</pre>
xticks.label[c(2:7)]<-""
# These functions all draw a plot with predicted probability of missingness
# (by group) - probability of missingness in the whole sample.
# 1 This one includes both model generated marginal means and simple mean
# inclusion.
detrend_catemplot_model<-function(x,label,model=NULL) {</pre>
  quosure <- quo(!!sym(x))
  if(!is.null(model)) {
  lmean <- emmeans(model,</pre>
                    as.formula(paste("~",x,"| pmiss")),
                      "pmiss",
                      at = list(pmiss=xticks.pmiss,
                                 z=levels(df %>%
                                            group_by(!!quosure) %>%
                                            filter(n() >= 50) %>%
                                            ungroup() %>%
                                            select(!!quosure) %>%
                                            unlist())),
```

Table 6: H2 / Omnibus model parameters, pt. 1

	В	CI.1	CI.u	*	В	CI.l	CI.u	*	В	CI.1	CI.u	*
	Interce	ept			pmiss				pmiss^2			
	1.36	1.33	1.39	***	1.40e + 87	9.68e + 83	2.04e + 90	***	4.66e + 08	1.18e + 08	1.84e + 09	***
	Main I	Eff.			Interaction				Q.Interaction	on		
Sex (Male)	1.03	1.01	1.05	***	1.73	.01	329.14		.00	6.56 e-08	.00	***
\$0-\$25k	1.02	.99	1.05		.00	.00	72.33		.69	.11	4.32	
Household Incor	ne (ref: S	\$100-\$20	00k)									
\$25-\$50k	.98	.96	1.01		105.65	.02	507780.08		44.64	9.04	220.55	***
\$50-\$75k	.99	.97	1.02		1165.69	.56	2426424.45		.01	.00	.06	***
\$75-\$100k	.99	.97	1.01		1.98	.00	2541.64		25.95	6.74	99.95	***
>\$200k	.99	.97	1.02		.64	.00	1587.92		12.81	2.94	55.89	***
Highest Parenta	l Educat	ion (ref:	College	Degree)								
<high school<="" td=""><td>1.07</td><td>1.02</td><td>1.12</td><td>**</td><td>8.22e-09</td><td>6.06 e-15</td><td>.01</td><td>**</td><td>.00</td><td>.00</td><td>.05</td><td>***</td></high>	1.07	1.02	1.12	**	8.22e-09	6.06 e-15	.01	**	.00	.00	.05	***
HS Grad.	1.04	1.00	1.07	*	.00	2.26e-08	18.33		.00	.00	.01	***
Some College	1.00	.98	1.03		1.64	.00	1525.01		.06	.02	.22	***
Graduate	1.02	1.00	1.04	*	.08	.00	29.44		.05	.02	.15	***
Census Race/Et	hnicity (	ref: Wh	ite)									
Black	1.07	1.05	1.10	***	3.79e-09	7.89e-13	.00	***	.00	.00	.01	***
Hispanic	1.02	1.00	1.04	*	.00	.00	1.51		.41	.11	1.46	
Asian	1.12	1.06	1.17	***	8.26e-12	.00	.00	**	.00	.00	.00	***
Other	1.03	1.01	1.05	*	.00	5.80e-09	.02	**	18.54	4.43	77.57	***
KSADS Trauma	Count (	ref: 0 E	xposures	)								
1 Trauma	1.00	.99	1.02		.23	.00	43.30		5.69	2.12	15.29	***
>=2 Trauma	1.02	.99	1.04		.07	.00	205.22		.00	.00	.00	***

Table 7: H2 / Omnibus model parameters, pt. 2

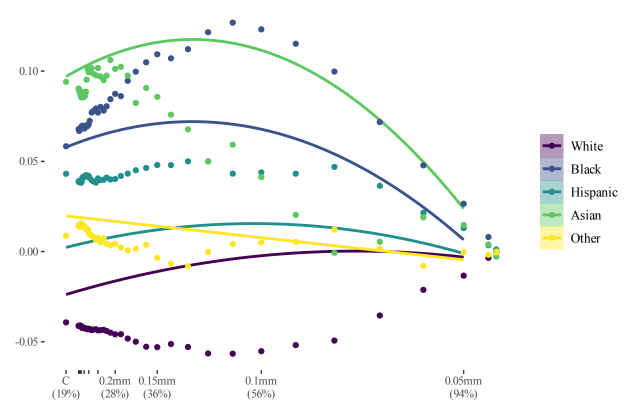
	В	CI.l	CI.u	*	В	CI.l	CI.u	*	В	CI.1	CI.u	*
	Interce	ept			pmiss				pmiss^2			
	1.36	1.33	1.39	***	1.40e + 87	9.68e + 83	2.04e + 90	***	4.66e + 08	1.18e + 08	1.84e + 09	***
	Main 1	Eff.			Interaction				Q.Interaction	on		
Pubertal Status (ref:	pre-pube	ertal)										
Early Puberty	1.00	.99	1.02		1.19	.00	359.61		.28	.10	.83	*
Mid Puberty	1.01	.99	1.03		2.05	.00	2067.75		.03	.01	.10	***
Late Puberty	1.06	1.00	1.13	*	1.91e-12	.00	.00	**	3926.89	109.53	140784.45	***
Post Pubertal	.94	.75	1.19		5.18e + 15	.00	2.07e + 48		1.14e-13	.00	.00	***
Area Disadvantage	.95	.94	.97	***	597703.26	13593.00	2.63e + 07	***	2.14	1.05	4.36	*
Child Opportunity	.97	.95	.98	***	4424.30	64.41	303924.31	***	11.79	5.31	26.16	***
NIHTB Flanker	1.00	.99	1.01		1.25	.07	23.95		.68	.39	1.19	
NIHTB Crystalized	1.01	1.00	1.03	*	7.21	.07	784.75		.00	.00	.00	***
NIHTB Total	.97	.96	.99	**	1.21	.00	301.16		32303.00	11419.24	91379.42	***
WISC V Matrix	1.00	.99	1.00		.68	.05	8.64		20.63	12.78	33.30	***
Psychopathology	1.02	1.01	1.03	***	.04	.00	.41	**	.01	.01	.02	***
Internalizing	.99	.98	1.00	*	.85	.08	9.31		104.30	66.47	163.66	***
Externalizing	.99	.98	1.00	**	5.66	.53	59.92		6.72	4.31	10.49	***
Age	.97	.96	.98	***	23.98	2.39	240.40	**	3247.86	2103.56	5014.63	***
BMI	1.01	1.01	1.02	***	18.73	3.73	94.21	***	.00	.00	.00	***

```
rg.limit = 400000)
  plot.dat<-emmip(lmean,</pre>
                  as.formula(paste(x,"~ pmiss")),
                  CIs=TRUE,
                  plotit = FALSE)
 }
  #Add detrended prediction
 plot.dat$ydt<-plot.dat$yvar-plot.dat$pmiss</pre>
  #Add the probability (average) without any cross control.
 mdf<-dflong %>%
    select(c(!!quosure,pmiss,missing)) %>%
    group_by(!!quosure,pmiss) %>%
    filter(n() >= 50) \%\% # Remove cases with n<50. Otherwise they are very noisy.
    summarise(avg=mean(missing)) %>%
    mutate(ydt=avg-pmiss,
           yvar=avg)
  names(mdf)[1]<-"tvar"</pre>
  ggplot(plot.dat,aes(x=pmiss,y=ydt,color=tvar,fill=tvar)) +
       geom_smooth(method="loess") +
       geom point(data=mdf) +
       scale color viridis(name=x,discrete = TRUE,na.value="gray") +
       scale fill viridis(name=x,discrete= TRUE,na.value="gray") +
       labs(title=paste("De-trended Mean Missingness by",label)) +
    scale x continuous(breaks=xticks.pmiss,labels = xticks.label) +
    theme(axis.title.x=element blank(),
          axis.title.y=element blank(),
          legend.title=element blank(),
          legend.text=element_text(size=10))# +
    \#qeom\_function(fun = (x) (-6.349e+00-3.078e-01)*x - (8.929e-01+1.996e+01)*x^2 + 3.078e-01)
#2 As above, but without specifying a model.
detrend_catemplot <- function(x, label) {</pre>
  quosure <- quo(!!sym(x))
  #The probability (average) without any cross control.
 mdf <- dflong %>%
    select(c(!!quosure, pmiss, missing)) %>%
    group_by(!!quosure, pmiss) %>%
    filter(n() >= 50) \%% # Remove cases with n<50. Otherwise they are very noisy.
    summarise(avg = mean(missing)) %>%
```

```
mutate(ydt = avg - pmiss,
           yvar = avg)
  names(mdf)[1] <- "tvar"</pre>
  ggplot(mdf, aes(x = pmiss,y = ydt,color = tvar,fill = tvar)) +
    geom point() +
    scale color viridis(name = x,
                        discrete = TRUE,
                        na.value = "gray") +
    scale fill viridis(name = x,
                       discrete = TRUE.
                       na.value = "gray") +
    labs(title = paste("De-trended Mean Missingness by", label)) +
    scale x continuous(breaks = xticks.pmiss, labels = xticks.label) +
    theme(
      axis.title.x = element_blank(),
      axis.title.y = element_blank(),
      legend.title = element_blank(),
      legend.text = element_text(size = 10)
# 3 As above, but for continuous variables.
detrend contemplot<-function(x,label) {</pre>
  quosure <- quo (!!sym(x))
  #Add the probability (average) without any cross control.
  mdf<-dflong %>%
    select(c(!!quosure,pmiss,missing)) %>%
    mutate(quantile=ntile(!!quosure,5)) %>%
    group by(quantile,pmiss) %>%
    summarise(avg=mean(missing)) %>%
    mutate(ydt=avg-pmiss,
           Quantile=as.factor(quantile))
  ggplot(mdf,aes(x=pmiss,y=ydt,color=Quantile)) +
       geom_point() +
       scale color viridis(discrete=TRUE) +
       labs(title=paste("De-Trended Mean Missingness by",label)) +
    scale_x_continuous(breaks=xticks.pmiss,labels = xticks.label) +
    theme(axis.title.x=element_blank(),
          axis.title.y=element blank(),
          legend.text=element_text(size=10))
}
```

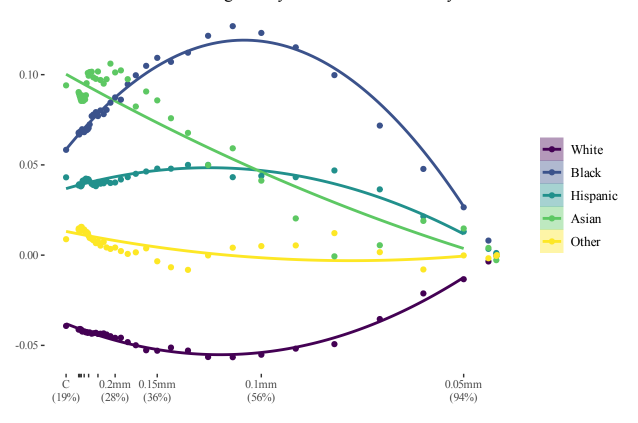
If we generate a plot with both model predictions and the average inclusion within each factor level and motion threshold (here, lines are the model marginal means and points are the averages from the data), we can see that they do not match, at least for this variable:

# De-trended Mean Missingness by Census Race/Ethnicity



If we use a model with only the race/ethnicity factor:

## De-trended Mean Missingness by Census Race/Ethnicity



That is much improved, so the distortions in the first plot are most likely due to interference from other variables. Regardless H2 is generally not supported by these results. The descriptive data is probably more helpful (and will be presented).

```
# Sub-Plots:
# Categorical:
h2p1<-detrend_catemplot("sex", "Sex")
h2p2<-detrend_catemplot("ksads_factor", "Trauma Count")
h2p3<-detrend_catemplot("race_ethnicity.factor", "Census Race/Ethnicity")
h2p4<-detrend_catemplot("household.income", "Household Income")
h2p5<-detrend_catemplot("p.edu", "Highest Parental Education")
h2p6<-detrend_catemplot("pds_category", "Pubertal Status")

# Continuous:
h2plot.adi<-detrend_contemplot("adi.z", "ADI")
h2plot.coi<-detrend_contemplot("coi.z", "COI")
h2plot.flank<-detrend_contemplot("nihtbx_flanker_agecorrected.z", "NIHTB Flanker Score")
h2plot.cryst<-detrend_contemplot("nihtbx_cryst_agecorrected.z", "NIHTB Crystalized")</pre>
```

```
h2plot.ntot<-detrend_contemplot("nihtbx_totalcomp_agecorrected.z","NIHTB Total")
h2plot.wisc<-detrend_contemplot("pea_wiscv_tss.z","Wisc V Matrix")
h2plot.pfac<-detrend_contemplot("pfactor.z","P-factor")
h2plot.int<-detrend_contemplot("INT.z","Internalizing")
h2plot.ext<-detrend_contemplot("EXT.z","Externalizing")
h2plot.age<-detrend_contemplot("interview_age.z","Age")
h2plot.bmi<-detrend_contemplot("bmiAgeZ","BMI")

# A few outstanding effects are saved to a plot for the manuscript:
svg(filename="Manuscript/Figures/h2catfig.svg",height=9,width=7.5,family="Times New Roman")
h2p4 / h2p5 / h2plot.ntot / h2plot.bmi
grid::grid.draw(grid::textGrob("Proportion of Participants Excluded - Total Proportion Excluded",x=.01,rot=90,gp = grid::gpar(fontfamily = "Times grid::grid.draw(grid::textGrob("Motion Threshold (Proportion of Sample Missing)",y=.01,gp = grid::gpar(fontfamily = "Times New Roman")))
dev.off()

## cairo_pdf
## cairo_pdf
## 2
```

For this supplement, we will also generate plots for the other variables in groups of four:

```
h2p1 / h2p2 / h2p3 / h2p6
grid::grid.draw(grid::textGrob("Proportion of Participants Excluded - Sample Mean",x=.01,rot=90,gp = grid::gpar(fontfamily = "Times New Roman")))
grid::grid.draw(grid::textGrob("Motion Threshold / Proportion of Sample Missing",y=.01,gp = grid::gpar(fontfamily = "Times New Roman")))

h2plot.adi / h2plot.coi / h2plot.flank / h2plot.cryst
grid::grid.draw(grid::textGrob("Proportion of Participants Excluded - Sample Mean",x=.01,rot=90,gp = grid::gpar(fontfamily = "Times New Roman")))
grid::grid.draw(grid::textGrob("Motion Threshold / Proportion of Sample Missing",y=.01,gp = grid::gpar(fontfamily = "Times New Roman")))

h2plot.wisc / h2plot.pfac / h2plot.int / h2plot.ext / h2plot.age
grid::grid.draw(grid::textGrob("Proportion of Participants Excluded - Sample Mean",x=.01,rot=90,gp = grid::gpar(fontfamily = "Times New Roman")))
grid::grid.draw(grid::textGrob("Motion Threshold / Proportion of Sample Missing",y=.01,gp = grid::gpar(fontfamily = "Times New Roman")))
```

# 1.8 QCFC Figures (Figure 6)

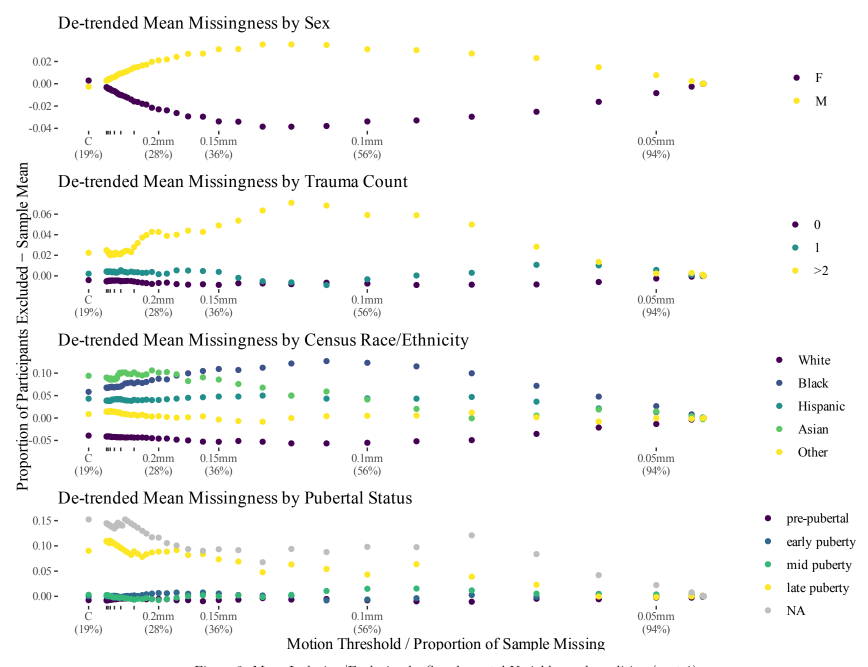


Figure 3: Mean Inclusion/Exclusion by Supplemental Variables and condition (part 1)



Figure 4: Mean Inclusion/Exclusion by Supplemental Variables and Condition (part 2)



Figure 5: Mean Inclusion/Exclusion by Supplemental Variables and Condition (part 3)

```
QCFCvals<-lapply(files,scan,sep=",")</pre>
# The distance matrix only has cortical (as opposed to subcortical distances, so
# we trim the QC-FC values to those dimensions. All of these matrices are
# symmetrical across the diagonal, so we can also omit the diagonal and lower
# triangle.
tdim=sqrt(length(QCFCvals[[1]]))
QCFCtrimvals <- lapply(QCFCvals,\(x) {
  m=matrix(x,ncol=sqrt(length(x)),byrow=TRUE)
  m2=m[1:tdim,1:tdim]
  m2[lower.tri(m2,diag = TRUE)] <- NA</pre>
  na.omit(c(m2))
  })
QCFCdf<-as.data.frame(reduce(QCFCtrimvals,cbind))</pre>
names(QCFCdf)<-c("distance","<.1mm","<.2mm","<.3mm","<.4mm","<.5mm","No Scrubbing")</pre>
QCFCdflong <- QCFCdf %>% pivot_longer(cols=-distance,
                                        names to="threshold",
                                        values_to="QCFC")
# Calculate Slopes
QCFCdfz <- QCFCdf %>%
  mutate(across(.fns=scale))
annotext<-function(x) {</pre>
  m<-lm(QCFCdfz$distance ~ QCFCdfz[,x])</pre>
  ci<-confint(m,level=.80)</pre>
  spec="%.2f"
  paste0(" = ",
         sprintf(spec,m$coefficients[2]),
         " 99% CI [".
         sprintf(spec,ci[2,1]),
         sprintf(spec,ci[2,2]),
         "]")
}
annot<-data.frame(threshold=unique(QCFCdflong$threshold),</pre>
                 text=sapply(unique(QCFCdflong$threshold),annotext))
annot$text<-gsub("0\\.", ".", annot$text)</pre>
annot$distance=mean(range(QCFCdflong$distance))
annot $QCFC=.1
```

#### 1.9 Multiply at Risk Cell Counts

```
subdf1 <- df %>%
 filter(race_ethnicity.factor!="White",
         pfactor.z \ge 1.5)
subtable1<-data.frame(</pre>
  "F"=nrow(subdf1).
 "T"=sum(subdf1$ABCD rsfmri QC1),
  "C"=sum(subdf1$any rest),
  ".5"=sum(subdf1$gframes 0.5 375),
 ".4"=sum(subdf1$gframes 0.4 375),
 ".3"=sum(subdf1$gframes 0.3 375),
  "R"=sum(subdf1$ABCD_rsfmri_QC2),
 ".2"=sum(subdf1$gframes_0.2_375),
 ".1"=sum(subdf1$gframes_0.1_375)
 ) %>%
 as hux() %>%
  set_header_rows(1,TRUE) %>%
  set_caption("Non-white (census) with psychopathology at z >= 1.5")
subtable1[1,]<-gsub("X\\.","\\.",subtable1[1,])
subdf2 <- df %>%
 filter(nihtbx_totalcomp_agecorrected.z<=-1.5,</pre>
         sex=="M")
subtable2<-data.frame(
```

```
"F"=nrow(subdf2),
"T"=sum(subdf2$ABCD_rsfmri_QC1),
"C"=sum(subdf2$any_rest),
".5"=sum(subdf2$gframes_0.5_375),
".4"=sum(subdf2$gframes_0.4_375),
".3"=sum(subdf2$gframes_0.3_375),
"R"=sum(subdf2$gframes_0.2_375),
".1"=sum(subdf2$gframes_0.2_375),
".1"=sum(subdf2$gframes_0.1_375)
) %>%
as_hux() %>%
set_header_rows(1,TRUE) %>%
set_caption("Male participants with NIH toolbox total scores of z <= -1.5")</pre>
subtable2[1,]<-gsub("X\\.","\\.",subtable2[1,])
```

subtable1

Table 8: Non-white (census) with psychopathology at z >= 1.5

F Τ С .5.3 .4 $\mathbf{R}$ .2 .1 373 399 180 552 510 407388385339

Vs. 0.1797951

subtable2

Table 9: Male participants with NIH toolbox total scores of z <= -1.5

F Τ С .5 .4 .3  $\mathbf{R}$ .1 248 382 343 25423523220794 275

Vs. 0.180442

# 2 Supplementary Results

#### 2.1 Categorical Variable Table by Condition

```
catvars<-c("sex",
             "household.income",
             "p.edu",
             "race_ethnicity.factor",
             "ksads factor",
             "pds category")
catTableAtThresh <- function(dataorig,</pre>
                              boolean,
                              base = NULL,
                              vars = catvars) {
  n <- sum(boolean)
  tl <- list(cbind(n, n * 100 / 11876))
  pdivisor <- n
  data <- dataorig[boolean, ]</pre>
  row.names(tl[[1]]) <- "Total"</pre>
  for (i in vars) {
    st <-
      cbind(table(data[, i], useNA = "always"), (table(data[, i], useNA = "always") *
                                                     100) / pdivisor)
    tl[[length(tl) + 1]] <- st
  t <- reduce(tl, rbind)
  if (is.null(base)) {
    t.
  } else {
    cbind(t, t[, 2] - base[, 2])
  }
}
CatVar375Table <- cbind(</pre>
  catTableAtThresh(df, rep(TRUE,nrow(df))),
  catTableAtThresh(df, df$ABCD_rsfmri_QC1),
  catTableAtThresh(df, df$any_rest),
  catTableAtThresh(df, df$gframes_0.5_375),
  catTableAtThresh(df, df$gframes_0.4_375),
  catTableAtThresh(df, df$gframes_0.3_375),
  catTableAtThresh(df, df$ABCD_rsfmri_QC2),
  catTableAtThresh(df, df$gframes 0.2 375),
```

```
catTableAtThresh(df, df$gframes_0.1_375)
)
#Try to draw as a huxtable
rownames(CatVar375Table) <-gsub("\\$","\\\\$",rownames(CatVar375Table))</pre>
CatVar375Hux<-as hux(CatVar375Table) %>%
  theme basic() %>%
  insert_row(value=c("n","\\%",rep(c("n","\\%"),8))) %>%
  insert row("Full","",
             "QC1","",
             "ABCC","",
             ".5","",
             ".4","",
             ".3mm","",
             "QC2","",
             ".2mm","",
             ".1mm","") %>%
  merge_cells(1,1:2) %>%
  merge_cells(1,3:4) %>%
  merge_cells(1,5:6) %>%
  merge_cells(1,7:8) %>%
  merge_cells(1,9:10) %>%
  merge cells(1,11:12) %>%
  merge cells(1,13:14) %>%
  merge cells(1,15:16) %>%
  merge cells(1,17:18) %>%
  insert column(value=c("","",rownames(CatVar375Table) %>% replace na("Missing"))) %>%
  insert_column(value=c("","","Total","Sex","","",
                        "Income","","","","","","",
                        "Parent Ed.","","","","","",
                        "Race/Ethnicity","","","","","",
                        "Trauma Count","","",",
                        "Puberty","","","","","")) %>%
  set_caption("Categorical Values at Each Level of Stringency (Subjects excluded with < 375 Frames), Pt 1") %>%
  set escape contents(FALSE) %>%
  set_font_size(8) %>%
  set number format(3:nrow(.),3+0:7*2,value=fmt_pretty()) %>%
  set_number_format(3:nrow(.),4+0:7*2,value="%.2g") %>%
  set all padding(1)
CatVar375Hux
```

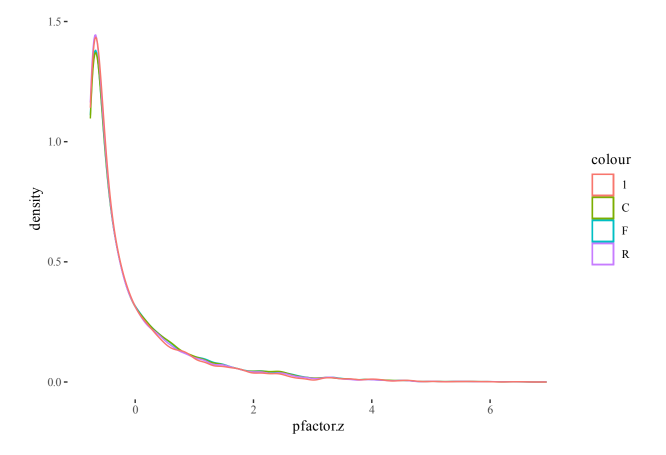
Table 10: Categorical Values at Each Level of Stringency (Subjects excluded with < 375 Frames), Pt 1

		Full		QC1		ABCC	;	0.5		0.4		0.3mm	ı	QC2		0.2mm	ı	0.1mm	
		n	%	n	%	n	%	n	%	n	%	n	%	n	%	n	%	n	%
Total	Total	11,876	1e+02	$11,\!355$	96	9,600	81	9,320	78	9,262	78	9,098	77	$9,\!627$	81	8,507	72	5.25e + 03	44.2
Sex	F	5,680	48	5,444	48	4,575	48	4,475	48	$4,\!458$	48	4,408	48	4,792	50	4,199	49	2.7e + 03	51.5
	M	6,196	52	5,911	52	5,025	52	4,845	52	4,804	52	4,690	52	4,835	50	4,308	51	2.55e + 03	48.5
	Missing	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0
Income	100k to $200k$	3,314	28	3,192	28	2,735	28	2,673	29	2,660	29	2,623	29	2,785	29	$2,\!470$	29	1.62e+03	30.9
	0 to 25k	1,635	14	1,526	13	1,206	13	1,139	12	1,126	12	1,106	12	1,188	12	988	12	538	10.2
	\$25k to $$50k$	1,588	13	1,524	13	1,291	13	1,250	13	1,239	13	1,212	13	$1,\!277$	13	1,135	13	676	12.9
	50k to $75k$	1,499	13	1,441	13	1,254	13	1,215	13	1,208	13	1,189	13	1,204	13	1,106	13	630	12
	75k to 100k	1,572	13	1,508	13	1,316	14	1,281	14	1,276	14	1,250	14	1,315	14	1,188	14	773	14.7
	Over $$200k$	1,250	11	1,206	11	1,014	11	995	11	992	11	978	11	1,056	11	930	11	626	11.9
	Missing	1,018	8.6	958	8.4	784	8.2	767	8.2	761	8.2	740	8.1	802	8.3	690	8.1	385	7.33
Parent Ed.	College Degree	3,015	25	2,877	25	2,504	26	2,441	26	2,429	26	2,393	26	2,486	26	2,268	27	1.44e + 03	27.4
	< HS	593	5	551	4.9	388	4	370	4	365	3.9	359	3.9	439	4.6	324	3.8	173	3.3
	HS Graduate	1,132	9.5	1,068	9.4	855	8.9	815	8.7	809	8.7	786	8.6	846	8.8	721	8.5	373	7.1
	Some College	3,079	26	2,957	26	$2,\!528$	26	2,432	26	2,410	26	2,361	26	$2,\!451$	25	$2,\!179$	26	1.29e + 03	24.6
	Graduate Degree	4,043	34	3,889	34	3,313	35	3,250	35	3,238	35	3,188	35	3,396	35	3,007	35	1.97e + 03	37.5
	Missing	14	0.12	13	0.11	12	0.12	12	0.13	11	0.12	11	0.12	9	0.093	8	0.094	3	0.0571
Race/Ethnicity	White	6,180	52	5,945	52	5,238	55	5,104	55	5,075	55	4,999	55	5,146	53	4,710	55	3.07e + 03	58.5
	Black	1,784	15	1,672	15	1,338	14	1,279	14	1,269	14	1,241	14	1,313	14	1,122	13	569	10.8
	Hispanic	2,411	20	2,312	20	1,845	19	1,799	19	1,788	19	1,746	19	1,966	20	1,630	19	960	18.3
	Asian	252	2.1	236	2.1	180	1.9	175	1.9	175	1.9	168	1.8	196	2	155	1.8	101	1.92
	Other	1,247	11	1,188	10	997	10	961	10	953	10	942	10	1,004	10	888	10	545	10.4
	Missing	2	0.017	2	0.018	2	0.021	2	0.021	2	0.022	2	0.022	2	0.021	2	0.024	2	0.0381
Trauma Count	0	7,723	65	7,383	65	$6,\!275$	65	6,102	65	6,059	65	5,957	65	$6,\!295$	65	$5,\!586$	66	3.47e + 03	66.1
	1	3,004	25	2,882	25	$2,\!422$	25	2,345	25	2,330	25	2,285	25	$2,\!437$	25	$2{,}147$	25	1.34e + 03	25.5
	>2	1,149	9.7	1,090	9.6	903	9.4	873	9.4	873	9.4	856	9.4	895	9.3	774	9.1	440	8.38
	Missing	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0
Puberty	pre-pubertal	5,938	50	5,690	50	4,845	50	4,707	51	4,671	50	$4,\!575$	50	4,818	50	4,284	50	2.67e + 03	50.9
	early puberty	2,815	24	2,680	24	$2,\!275$	24	2,207	24	$2{,}195$	24	2,160	24	$2,\!268$	24	1,998	23	1.26e + 03	24
	mid puberty	2,798	24	2,681	24	$2,\!253$	23	$2,\!190$	23	$2,\!181$	24	$2{,}150$	24	$2,\!289$	24	2,025	24	1.2e + 03	22.8
	late puberty	188	1.6	180	1.6	135	1.4	127	1.4	126	1.4	126	1.4	152	1.6	118	1.4	75	1.43
	post pubertal	12	0.1	10	0.088	10	0.1	9	0.097	9	0.097	9	0.099	7	0.073	7	0.082	3	0.0571
	Missing	125	1.1	114	1	82	0.85	80	0.86	80	0.86	78	0.86	93	0.97	75	0.88	43	0.819

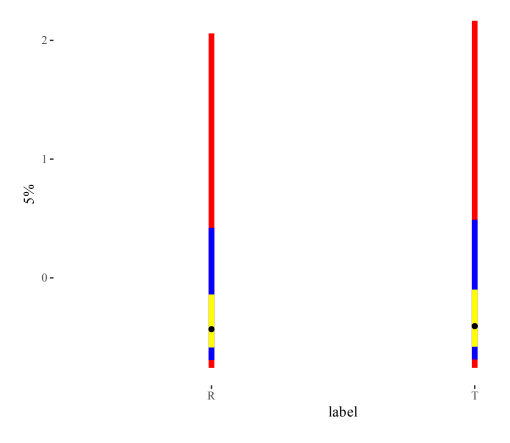
### 2.2 Alternative Continuous Variable Plots (psychopathology)

Here are some alternative ways of plotting continuous variables (using psychopathology as an example).

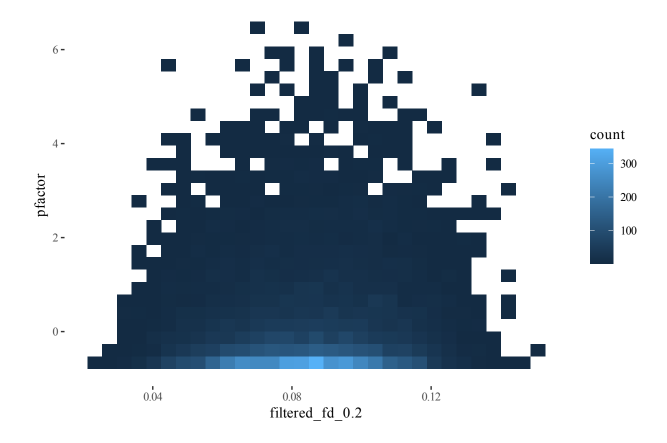
```
df %>% filter(ABCD_rsfmri_QC2==TRUE) %>%
ggplot(aes(x=pfactor.z)) +
  geom_density(aes(color = "F"), alpha = .2, data = df %>% filter(ABCD_rsfmri_QC1==TRUE)) +
  geom_density(aes(color = "C"), alpha = .2, data = df %>% filter(any_rest==TRUE)) +
  geom_density(aes(color = "R"), alpha = .2, data = df %>% filter(ABCD_rsfmri_QC2==TRUE)) +
  geom_density(aes(color = "1"), alpha = .2, data = df %>% filter(gframes_0.1_375==TRUE))
```



```
ggplot(test,aes(x=label)) +
   geom_segment(aes(xend=label,y=`5%`,yend=`95%`),color="red",size=2) +
   geom_segment(aes(xend=label,y=`20%`,yend=`80%`),color="blue",size=2) +
   geom_segment(aes(xend=label,y=`35%`,yend=`65%`),color="yellow",size=2) +
   geom_point(aes(y=`50%`))
```



```
ggplot(df,aes(x=filtered_fd_0.2,y=pfactor)) +
geom_bin2d()
```



### 2.3 Missingness in Behavioral Data

This table shows missingness counts in non-imaging variables with more than 50 missing cases.

```
df %>%
    select(c(interview_age:household.income,p.edu,race_ethnicity.factor:site_id_l)) %>%
    rename(c("Child Opportunity Index"="coi","Household Income"="household.income","Area Disadvantage Index"="adi","NIH Toolbox Total"="nihtbx_tota'
    summarise_all(~sum(is.na(.))) %>%
    pivot_longer(everything()) %>%
    filter(value>50) %>%
    mutate(`% missing`=(value/nrow(df))*100) %>%
    arrange(desc(value)) %>%
    rename(c("Variable"="name","n Missing"="value")) %>% as_hux() %>%
    set_header_rows(1,TRUE) %>%
    set_caption("Behavioral and Demographic variables with >50 missing values")
```

Table 11: Behavioral and Demographic variables with >50 missing values

Variable	n Missing	% missing
Child Opportunity Index	1093	9.2
Household Income	1018	8.57
Area Disadvantage Index	863	7.27
NIH Toolbox Total	397	3.34
NIH Toolbox Crystalized	338	2.85
WISC V	249	2.1
ethn.iden.hisp	153	1.29
NIH Toolbox Flanker	153	1.29
Pubertal Status	125	1.05

# 2.4 Adjusted Models – Forest Plot

A graphical depiction of effects from the adjusted/condition models.

```
MFformat<-\(m) {</pre>
  stars<-sdlabFunctions::starPs(m[,4])</pre>
  pasteO(ORformat(m[,2]),
         stars,
         "\n[",ORformat(m[,1]),"-",ORformat(m[,3]),"]")
}
allmodelterms<-c(
    "(Intercept)",
    "sexM",
    "household.incomeheader",
    "household.income$0 to $25k",
    "household.income$25k to $50k",
    "household.income$50k to $75k",
    "household.income$75k to $100k",
    "household.incomeOver $200k",
    "p.eduheader",
    "p.edu< HS",
```

```
"p.eduGraduate Degree",
    "p.eduHS Graduate",
    "p.eduSome College",
    "race_ethnicityheader",
    "race_ethnicity.factorAsian",
    "race_ethnicity.factorBlack",
    "race_ethnicity.factorHispanic",
    "race_ethnicity.factorOther",
    "ksadsheader",
    "ksads factor>2",
    "ksads_factor1",
    "pdsheader",
    "pds_categoryearly puberty",
    "pds_categorylate puberty",
    "pds_categorymid puberty",
    "pds_categorypost pubertal",
    "adi.z",
    "coi.z",
    "nihtbx_flanker_agecorrected.z",
    "nihtbx_cryst_agecorrected.z",
    "nihtbx_totalcomp_agecorrected.z",
    "pea_wiscv_tss.z",
   "pfactor.z",
    "INT.z",
    "EXT.z",
    "interview age.z",
    "bmiAgeZ")
alllevelsggplot<-c("Intercept",</pre>
      "Sex (Male)",
      "**Household Income**",
      " $0-$25k",
      " $25-$50k",
      " $50-$75k",
      " $75-$100k",
      " >$200k",
      "**Highest Parental Education**",
      " < High School",
      " HS Grad.",
      " Some College",
      " Graduate",
      "**Census Race/Ethnicity**",
      " Black",
      " Hispanic",
```

```
" Asian",
      " Other",
      "**Trauma Count**",
      " 1 Trauma",
      " >=2 Trauma",
      "**Pubertal Status**",
      " Early Puberty",
      " Mid Puberty",
      " Late Puberty",
      " Post Pubertal",
      "Area Disadvantage",
      "Child Opportunity",
      "NIHTB Flanker",
      "NIHTB Crystalized",
      "NIHTB Total",
      "WISC V Matrix",
      "Psychopathology",
      "Internalizing",
      "Externalizing",
      "Age",
      "BMI"
      )
modeltermsA<-allmodelterms[1:18]
modeltermLabelsA<-alllevelsggplot[1:18]</pre>
modeltermsB<-allmodelterms[19:length(allmodelterms)]</pre>
modeltermLabelsB<-alllevelsggplot[19:length(allmodelterms)]</pre>
forestdf <- rbind(</pre>
  model.QC1.results %>% data.frame() %>% rownames to column(),
  model.ABCC.results %>% data.frame() %>% rownames to column(),
  model.5mm.results %>% data.frame() %>% rownames_to_column(),
  model.4mm.results %>% data.frame() %>% rownames_to_column(),
  model.3mm.results %>% data.frame() %>% rownames to column(),
  model.QC2.results %>% data.frame() %>% rownames_to_column(),
  model.2mm.results %>% data.frame() %>% rownames to column(),
  model.1mm.results %>% data.frame() %>% rownames to column()
) %>% mutate(threshold=factor(rep(c("T", "C", "5", "4", "3", "R", "2", "1"), each=nrow(model.QC1.results)))) %>%
  rename with(~ c("term","lowCI","OR","highCI","p","threshold")) %>%
  mutate(across(lowCI:highCI,exp)) %>%
  add row(term="household.incomeheader") %>%
  add row(term="p.eduheader") %>%
  add row(term="ksadsheader") %>%
  add row(term="pdsheader") %>%
```

```
add_row(term="race_ethnicityheader") %>%
  mutate(threshold=factor(threshold))
maxsigCI<-max(forestdf[forestdf$lowCI>1,"highCI"],na.rm=TRUE)
forestdf[forestdf$highCI>maxsigCI & !is.na(forestdf$highCI),"highCI"]<-maxsigCI</pre>
forestdfA<- forestdf %>% filter(term %in% modeltermsA) %>%
  mutate(term=factor(term.
                     levels=modeltermsA.
                     labels=modeltermLabelsA))
forestdfB<- forestdf %>% filter(term %in% modeltermsB) %>%
  mutate(term=factor(term.
                     levels=modeltermsB,
                     labels=modeltermLabelsB))
dodge<-.8
ggplot(forestdfA,aes(y=term,color=threshold)) +
  geom_vline(lty=2,aes(xintercept=1),colour='black') +
  geom point(aes(x=0R),position=ggstance::position dodgev(height=dodge)) +
  geom linerange(aes(xmin=lowCI,xmax=highCI,y=term),position=ggstance::position dodgev(height=dodge)) +
  xlim(0,maxsigCI) +
  geom vline(lty=2,aes(xintercept=1),colour='black') +
  scale y discrete(drop=FALSE,limits=rev) +
  scale color discrete(limits=rev,drop=TRUE,na.translate = F) +
  theme(axis.text.y=element markdown(),
        axis.title.y=element blank()) +
  xlab("Odds Ratio of Missingness + 95% CI")
ggplot(forestdfB,aes(y=term,color=threshold)) +
  geom vline(lty=2,aes(xintercept=1),colour='black') +
  geom point(aes(x=OR), position=ggstance::position dodgev(height=dodge)) +
  geom linerange(aes(xmin=lowCI,xmax=highCI,y=term),position=ggstance::position dodgev(height=dodge)) +
  xlim(0,maxsigCI) +
  scale y discrete(drop=FALSE,limits=rev) +
  scale color discrete(limits=rev,drop=TRUE,na.translate = F) +
  theme(axis.text.y=element markdown(),
        axis.title.y=element blank()) +
  xlab("Odds Ratio of Missingness + 95% CI")
```

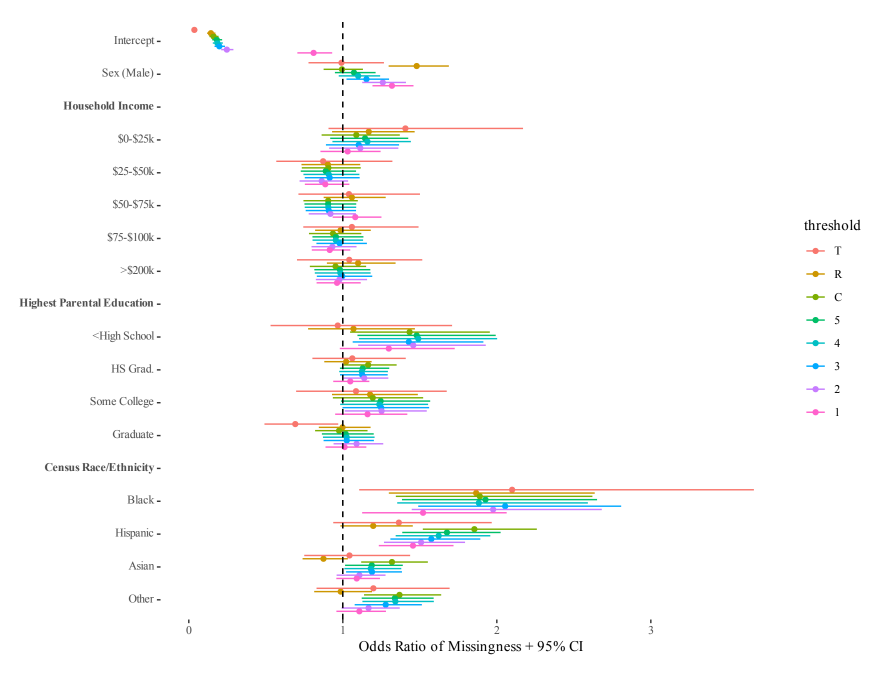


Figure 6: Odds ratios for missingness by threshold – Adjusted Models (part 1)

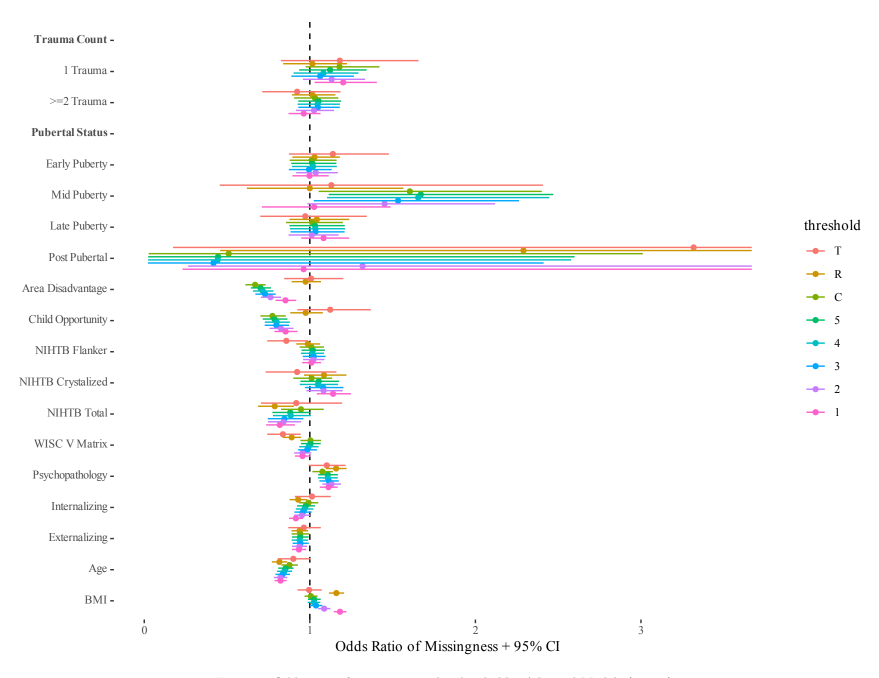


Figure 7: Odds ratios for missingness by threshold – Adjusted Models (part 2)

#### 2.5 Current Fast Track QC vs DAIRC recommendations

We state in the text that the currently published fastqc is non-overlapping with DAIRC inclusion recommendations. This table illustrates that non-overlap.

The variable 'fastqcok' is TRUE for participants who have at least one T1, rs-fMRI, and field map image marked useable in the current (Last modified 09/09/2019) abcd\_fastqc01.csv file. These recommendations are non overlapping with the tabular data list as well as DAIRC recommendations:

```
table(df$fastqcok,df$ABCD_rsfmri_QC1,dnn=c("FastQC","Tabulated"))
          Tabulated
##
## FastQC FALSE TRUE
     FALSE
             410 1378
     TRUE
             111 9977
table(df$fastqcok,df$ABCD_rsfmri_QC2,dnn=c("FastQC","DAIRC"))
##
          DAIRC
## FastQC FALSE TRUE
     FALSE
             822 966
     TRUE
           1427 8661
##
```

### 2.6 Association of QC missingness with Propensity weighting.

Gard (2020) notes that analyses using propensity weighting can be biased when missing data is correlated with population weights. They present data that association between missingness and population weights are small when considering structural and task-based brain data from ABCD. Here, we repeat these analyses considering rs-fMRI data in each condition.

Gard et al. appear to have used Welch's two-sample t-tests to examine differences in propensity weight between included and excluded groups, and we will do the same here.

```
gard_weighting_test<-function(boolvar) {
   formulastring=paste("acs_raked_propensity ~",boolvar)
   res<-broom::tidy(t.test(as.formula(formulastring), data=df, na.action="na.fail", conf.level=0.95))
   names(res)<-c("delta","Mexcluded","Mincluded","t","p","df","ci.low","ci.high","method","alternative")
   res %>% select(delta:ci.high)
}

weight_ttests<-reduce(lapply(bools[-1],gard_weighting_test),rbind)
weight_ttests$d<-weight_ttests$delta/sd(df$acs_raked_propensity)
as_hux(weight_ttests) %>%
   insert_column(c("",levelsdf$Level[-1]),after=0) %>%
   set_caption("Propensity weights of included vs. excluded participants by condition; t-test parameters")
```

Table 12: Propensity weights of included vs. excluded participants by condition; t-test parameters

	delta	Mexcluded	Mincluded	t	p	df	ci.low	ci.high	d
ABCD 4 Tabulated (T)	-0.487	691	691	-0.0309	0.975	568	-31.5	30.5	-0.00139
ABCC (C)	-3.68	688	692	-0.472	0.637	3.64e + 03	-19	11.6	-0.0105
ABCC < 0.5 mm	-4.63	688	692	-0.617	0.537	4.33e+03	-19.3	10.1	-0.0132
ABCC < 0.4mm	-3.49	689	692	-0.467	0.64	4.46e + 03	-18.1	11.2	-0.00994
ABCC < 0.3mm	-2.93	689	692	-0.4	0.689	4.89e + 03	-17.3	11.4	-0.00835
ABCD 4 Recommended (R)	22.4	709	687	2.72	0.00652	3.38e + 03	6.25	38.5	0.0637
ABCD < 0.2mm	8.1	697	689	1.16	0.248	6.44e + 03	-5.63	21.8	0.0231
ABCD < 0.1mm	41.6	710	668	6.42	1.45e-10	1.12e + 04	28.9	54.3	0.119

### 2.7 Association of Exclusion with Expanded Demographic Variables

Available information on household income and (especially) race/ethnicity was greatly simplified in the main analysis for the purpose of brevity and comparibility with existing literature. For the sake of completeness and inclusivity, we present descriptive tables and bivariate models for the more granular variables here:

#### 2.7.1 Household Income

```
#Reorder the factor in increasing order and relabel for descriptives.
df$household.income.full<-factor(df$household.income.full,
                                 levels=c("Less than $5,000",
                                           "$5,000 through $11,999",
                                          "$12,000 through $15,999",
                                           "$16,000 through $24,999",
                                          "$25,000 through $34,999",
                                           "$35,000 through $49,999",
                                          "$50,000 through $74,999",
                                           "$75,000 through $99,999",
                                           "$100,000 through $199,999",
                                          "$200,000 and greater"),
                                 labels=c("Less than \\$5k",
                                          "\\$5k through \\$11.9k",
                                          "\\$12k through \\$15.9k",
                                          "\\$16k through \\$24.9k",
                                           "\\$25k through \\$34.9k",
```

```
"\\$35k through \\$49.9k",
                                           "\\$50k through \\$74.9k",
                                           "\\$75k through \\$99.9k",
                                           "\\$100k through \\$199.9k",
                                           "\\$200k and greater"))
FullIncomeTableTable <- cbind(</pre>
  catTableAtThresh(df, rep(TRUE, nrow(df)), vars="household.income.full"),
  catTableAtThresh(df, df$ABCD rsfmri QC1,vars="household.income.full"),
  catTableAtThresh(df, df$any rest, vars="household.income.full"),
  catTableAtThresh(df, df$gframes 0.5 375,vars="household.income.full"),
  catTableAtThresh(df, df$gframes 0.4 375,vars="household.income.full"),
  catTableAtThresh(df, df$gframes 0.3 375, vars="household.income.full"),
  catTableAtThresh(df, df$ABCD_rsfmri_QC2,vars="household.income.full"),
  catTableAtThresh(df, df$gframes_0.2_375,vars="household.income.full"),
  catTableAtThresh(df, df$gframes_0.1_375,vars="household.income.full")
#Try to draw as a huxtable
\#rownames(FullIncomeTableTable) < -qsub("\\$","\\\$",rownames(FullIncomeTableTable))
FullIncomeTableHux<-as hux(FullIncomeTableTable) %>%
 theme basic() %>%
 insert_row(value=c("n","\\%",rep(c("n","\\%"),8))) %>%
  insert_row("Full","",
             "QC1","",
             "ABCC","",
             ".5","",
             ".4","",
             ".3mm","",
             "QC2","",
             ".2mm","",
             ".1mm","") %>%
 merge_cells(1,1:2) %>%
 merge_cells(1,3:4) %>%
 merge cells(1,5:6) %>%
 merge_cells(1,7:8) %>%
 merge cells(1,9:10) %>%
 merge_cells(1,11:12) %>%
 merge cells(1,13:14) %>%
 merge cells(1,15:16) %>%
 merge cells(1,17:18) %>%
 insert_column(value=c("","",rownames(FullIncomeTableTable) %>% replace_na("Missing"))) %>%
  set caption("Granular Household Income at Each Level of Stringency (Subjects excluded with < 375 Frames)") %>%
  set escape contents(FALSE) %>%
```

```
set_font_size(8) %>%
set_number_format(3:nrow(.),2+0:7*2,value=fmt_pretty()) %>%
set_number_format(3:nrow(.),3+0:7*2,value="%.2g") %>%
set_all_padding(1)
FullIncomeTableHux
```

Table 13: Granular Household Income at Each Level of Stringency (Subjects excluded with < 375 Frames)

	Full		QC1		ABCC	7	0.5		0.4		0.3mn	n	QC2		0.2mn	n	0.1mm	
	n	%	n	%	n	%	n	%	n	%	n	%	n	%	n	%	n	%
Total	11,876	1e+02	11,355	96	9,600	81	9,320	78	9,262	78	9,098	77	9,627	81	8,507	72	5.25e + 03	44.2
Less than \$5k	417	3.5	379	3.3	289	3	279	3	273	2.9	266	2.9	280	2.9	228	2.7	125	2.38
5k through \$11.9k	421	3.5	386	3.4	311	3.2	287	3.1	286	3.1	283	3.1	293	3	252	3	130	2.48
12k through $15.9k$	273	2.3	266	2.3	213	2.2	200	2.1	199	2.1	196	2.2	220	2.3	184	2.2	102	1.94
16k through 24.9k	524	4.4	495	4.4	393	4.1	373	4	368	4	361	4	395	4.1	324	3.8	181	3.45
25k through $34.9k$	654	5.5	628	5.5	521	5.4	503	5.4	499	5.4	491	5.4	537	5.6	459	5.4	269	5.12
35k through $49.9k$	934	7.9	896	7.9	770	8	747	8	740	8	721	7.9	740	7.7	676	7.9	407	7.75
50k through $74.9k$	1,499	13	1,441	13	1,254	13	1,215	13	1,208	13	1,189	13	1,204	13	1,106	13	630	12
75k through $99.9k$	1,572	13	1,508	13	1,316	14	1,281	14	1,276	14	1,250	14	1,315	14	1,188	14	773	14.7
100k through 200k	3,314	28	3,192	28	2,735	28	2,673	29	2,660	29	2,623	29	2,785	29	2,470	29	1.62e + 03	30.9
\$200k and greater	1,250	11	1,206	11	1,014	11	995	11	992	11	978	11	1,056	11	930	11	626	11.9
Missing	1,018	8.6	958	8.4	784	8.2	767	8.2	761	8.2	740	8.1	802	8.3	690	8.1	385	7.33

```
m.bvincome<-bvmodel("household.income.full")</pre>
#Reformat the table:
# Replace the first \n with " ["
m.bvincome.f <- sub("\n", " [", m.bvincome, fixed = TRUE)</pre>
# Replace the second \n with "]"
m.bvincome.f <- sub("\n", "]", m.bvincome.f, fixed = TRUE)</pre>
# Remove the p value
m.bvincome.f <- gsub("].*","]", m.bvincome.f)</pre>
as hux(m.bvincome.f) %>%
  insert_column(levels(df$household.income.full)[-1]) %>%
  insert row(
    "Household Income (ref: \\$100-\\$199.9k)", rep(c("OR [95\\% CI]"), 8)) %>%
  insert_row(c("", "T", "C", ".5", ".4", ".3", "R", ".2", ".1")) %>%
  set_caption("Bivariate Model of Granular Household Income and Missingness by Condition. Ref: \\$100k-\\$199k") %>%
  set escape contents(value=FALSE) %>%
  set wrap(TRUE) %>%
  set width(1)
```

#### 2.7.2 Detailed Race/Ethnicity

Census Race/Ethnicity Categories were used in the manuscript for the sake of brevity and because they are conventional and easy to compare to other work (which is of primary importance in a work commenting on methods). However, we acknowledge that the census categories are reductive both in the number of identifications present and in the lack of nuance they present (i.e., inability to pick multiple categories). ABCD collects information on participant race/ethnicity in considerably more detail, specifically as a series of yes/no questions about particular identities. We here present a comparison tables between the two measures, descriptives of the more granular race/ethnicity data, and bivariate models using those variables.

2.7.2.1 Descriptives This table presents a comparison of the census race/ethnicity measure included in ABCD and the extended race/ethnicity coding.

Table 14: Bivariate Model of Granular Household Income and Missingness by Condition. Ref: \$100k-\$199k

	Т	С	0.5	0.4	0.3	R	0.2	0.1
Household Income (ref: \$100-\$200k)	OR [95% CI]							
\$5k through \$11.9k	.90 [.56-1.46]	.80 [.59-1.08]	.94 [.71-1.26]	.89 [.67-1.19]	.86 [.65-1.14]	.89 [.67-1.19]	.81 [.61-1.06]	.96 [.71-1.29]
\$12k through \$15.9k	.26** [.1156]	.64* [.4490]	.74 [.53-1.03]	.70* [.5098]	.69* [.5096]	.49*** [.3470]	.58*** [.4280]	.72* [.5299]
\$16k through \$24.9k	.58* [.3596]	.75 [.56-1.00]	.82 [.62-1.08]	.80 [.61-1.06]	.80 [.61-1.04]	.67** [.5089]	.74* [.5797]	.81 [.61-1.07]
\$25k through \$34.9k	.41*** [.2469]	.58*** [.4376]	.61*** [.4680]	.59*** [.4577]	.58*** [.4576]	.45*** [.3359]	.51*** [.4066]	.61*** [.4779]
\$35k through \$49.9k	.42*** [.2767]	.48*** [.3763]	.51*** [.3966]	.50*** [.3864]	.52*** [.4067]	.54*** [.4169]	.46*** [.3659]	.55*** [.4371]
\$50k through \$74.9k	.40*** [.2662]	.44*** [.3457]	.47*** [.3760]	.46*** [.3658]	.46*** [.3658]	.50*** [.3964]	.43*** [.3454]	.59*** [.4774]
\$75k through \$99.9k	.42*** [.2865]	.44*** [.3456]	.46*** [.3658]	.44*** [.3556]	.45*** [.3657]	.40*** [.3151]	.39*** [.3149]	.44*** [.3556]
\$100k through \$200k	.38*** [.2656]	.48*** [.3860]	.48*** [.3961]	.47*** [.3758]	.46*** [.3758]	.39*** [.3149]	.41*** [.3351]	.45*** [.3656]
\$200k and greater	.36*** [.2357]	.53*** [.4168]	.52*** [.4166]	.49*** [.3963]	.49*** [.3962]	.38*** [.2949]	.42*** [.3352]	.43*** [.3454]

```
"American Indian",
                   "Alaska Native".
                   "Native Hawaiian",
                   "Guamanian",
                   "Samoan",
                   "Pacific Islander",
                   "Asian Indian",
                   "Filipino",
                   "Chinese",
                   "Japanese",
                   "Korean",
                   "Vietnamese",
                   "Other Asian",
                   "Other",
                   "Refuse",
                   "Don't Know",
                  "Hispanic")
retable1[,1:6]
```

Next, here are the count and percentages of each of the granular variables, by QC condition:

```
boolTable<-\(data=df,bool=TRUE) {</pre>
  df[bool,] %>% summarise(across(race.iden.white:ethn.iden.hisp,
            list(n= ~ sum(.,na.rm = TRUE),
                 Percent = ~ round(mean(. == 1, na.rm = TRUE)*100,2)
                 ))) %>%
    pivot_longer(cols=everything(),
                 names_to=c(".value", "measure"),
                 names sep=" ") %>% t()
}
retable2 <- cbind(
  boolTable(bool = TRUE),
  boolTable(bool = df$ABCD_rsfmri_QC1),
  boolTable(bool = df$any rest),
  boolTable(bool = df$gframes_0.5_375),
  boolTable(bool = df$gframes 0.4 375),
  boolTable(bool = df$gframes_0.3_375),
  boolTable(bool = df$ABCD_rsfmri_QC2),
  boolTable(bool = df$gframes_0.2_375),
  boolTable(bool = df$gframes_0.1_375))[-1,]
retable2 %>% as_hux() %>%
```

Table 15: Percentage Endorsing Granular Race/Ethnicity Variables within Census Race/Ethnicity Groups

Census Race/Ethnicity	White	Black	Hispanic	Asian	Other
White	99.9%	0.0%	68.0%	0.0%	79.3%
Black	0.0%	99.8%	8.9%	0.0%	41.9%
American Indian	0.0%	0.0%	4.7%	0.0%	23.4%
Alaska Native	0.0%	0.0%	0.1%	0.0%	0.2%
Native Hawaiian	0.0%	0.0%	0.2%	0.0%	1.4%
Guamanian	0.0%	0.0%	0.0%	0.0%	0.2%
Samoan	0.0%	0.0%	0.0%	0.0%	0.9%
Pacific Islander	0.0%	0.0%	0.3%	0.0%	2.6%
Asian Indian	0.0%	0.0%	0.2%	21.0%	4.4%
Filipino	0.0%	0.0%	0.7%	34.1%	8.1%
Chinese	0.0%	0.0%	1.5%	17.1%	7.1%
Japanese	0.0%	0.0%	0.2%	6.0%	5.1%
Korean	0.0%	0.0%	0.3%	10.3%	5.3%
Vietnamese	0.0%	0.0%	0.2%	10.3%	2.6%
Other Asian	0.0%	0.0%	0.4%	13.5%	3.7%
Other	0.0%	0.0%	27.2%	0.0%	11.5%
Refuse	0.0%	0.0%	1.2%	0.0%	2.5%
Don't Know	0.0%	0.0%	3.7%	0.0%	1.2%
Hispanic	0.0%	0.0%	100.0%	0.0%	0.0%

```
set_number_format(1:nrow(.),seq(1,ncol(.),2),value=fmt_pretty()) %>%
set_number_format(1:nrow(.),seq(2,ncol(.),2),value=fmt_pretty(digits=2,format="f")) %>%
insert_column(c("White",
                "Black",
                "American Indian",
                "Alaska Native",
                "Native Hawaiian",
                "Guamanian",
                "Samoan",
                "Pacific Islander",
                "Asian Indian",
                "Filipino",
                "Chinese",
                "Japanese",
                "Korean",
                "Vietnamese",
                "Other Asian",
                "Other",
                "Refuse",
                "Don't Know",
                "Hispanic")) %>%
insert_row(c("",rep(c("n","%"),length(bools)))) %>%
insert_row(c("",
             "F","",
             "QC1","",
             "C","",
             ".5","",
             ".4","",
             ".3","",
             "QC2","",
             ".2","",
             ".1","")) %>%
set_caption("Granular Race/Ethnicity Data by Condition")
```

2.7.2.2 Bivariate Models Finally, we present bivariate models of the granular binary race/ethnicity coding variables:

Table 16: Granular Race/Ethnicity Data by Condition

	F		QC1		$\mathbf{C}$		0.5		0.4		0.3		QC2		0.2		0.1	
	n	%	n	%	n	%	n	%	n	%	n	%	n	%	n	%	n	%
White	8,804	74	8,461	75	7,320	76	7,127	76	7,083	76	6,977	77	7,300	76	6,557	77	4,211	80
Black	2,518	21	2,375	21	1,918	20	1,840	20	1,825	20	1,785	20	1,903	20	1,631	19	870	17
American Indian	406	3.4	387	3.4	326	3.4	314	3.4	310	3.4	303	3.3	320	3.3	289	3.4	182	3.5
Alaska Native	5	0.04	3	0.03	2	0.02	1	0.01	1	0.01	1	0.01	2	0.02	1	0.01	1	0.02
Native Hawaiian	23	0.19	22	0.19	20	0.21	20	0.21	20	0.22	20	0.22	19	0.2	18	0.21	6	0.11
Guamanian	2	0.02	2	0.02	2	0.02	2	0.02	2	0.02	2	0.02	0	0	2	0.02	1	0.02
Samoan	12	0.1	12	0.11	11	0.11	11	0.12	11	0.12	11	0.12	12	0.12	11	0.13	7	0.13
Pacific Islander	40	0.34	36	0.32	31	0.32	30	0.32	30	0.32	30	0.33	30	0.31	27	0.32	17	0.32
Asian Indian	114	0.96	108	0.95	92	0.96	88	0.94	88	0.95	86	0.95	90	0.93	79	0.93	50	0.95
Filipino	203	1.7	191	1.7	152	1.6	150	1.6	149	1.6	146	1.6	158	1.6	135	1.6	95	1.8
Chinese	167	1.4	157	1.4	124	1.3	121	1.3	121	1.3	119	1.3	135	1.4	116	1.4	63	1.2
Japanese	84	0.71	82	0.72	60	0.62	59	0.63	59	0.64	57	0.63	72	0.75	54	0.63	41	0.78
Korean	100	0.84	98	0.86	80	0.83	78	0.84	78	0.84	76	0.84	85	0.88	70	0.82	48	0.91
Vietnamese	63	0.53	60	0.53	47	0.49	45	0.48	44	0.48	43	0.47	50	0.52	42	0.49	28	0.53
Other Asian	90	0.76	86	0.76	74	0.77	69	0.74	69	0.74	67	0.74	69	0.72	62	0.73	39	0.74
Other	800	6.7	761	6.7	581	6	567	6.1	561	6.1	549	6	645	6.7	513	6	292	5.6
Refuse	59	0.5	56	0.49	41	0.43	39	0.42	39	0.42	37	0.41	44	0.46	33	0.39	16	0.3
Don't Know	104	0.88	97	0.85	77	0.8	75	0.8	75	0.81	74	0.81	84	0.87	72	0.85	38	0.72
Hispanic	2,411	21	2,312	21	1,845	19	1,799	20	1,788	20	1,746	19	1,966	21	1,630	19	960	18

```
"Native Hawaiian",
                  "Guamanian",
                  "Samoan",
                  "Pacific Islander",
                  "Asian Indian",
                  "Filipino",
                  "Chinese".
                  "Japanese",
                  "Korean",
                  "Vietnamese",
                  "Other Asian",
                  "Other".
                  "Refuse".
                  "Don't Know",
                  "Hispanic")) %>%
  insert row(
    "", rep(c("OR\\newline [90\\% CI]\\newline p"), 8)) %>%
  insert_row(c("", "T", "C", ".5", ".4", ".3", "R", ".2", ".1")) %>%
  set_escape_contents(value=FALSE) %>%
  set wrap(TRUE) %>%
  set_width(1)
remodeltable[c(1:2,3:8),] %% set caption("Granular Race/Ethnicity Variables: Bivariate Models, Pt. 1") %>% set label("gran.race.ethn.1")
remodeltable[c(1:2,9:14),] %>% set caption("Granular Race/Ethnicity Variables: Bivariate Models, Pt. 2") %>% set label("gran.race.ethn.2")
remodeltable[c(1:2,15:19),] %>% set_caption("Granular Race/Ethnicity Variables: Bivariate Models, Pt. 3") %>% set_label("gran.race.ethn.3")
```

# 3 Changes since pre-registration

The following changes were made to the analysis plan post-registration:

- BMI was added as a study variable, to enhance comparibility with Cosgrove et. al. (2020) and reflect the known relation between BMI and in-scanner motion.
- The Behavioral Inhibition scale was removed. It is not available in the ABCD baseline data and was included in error.
- More data was excluded prior to motion filtering than expected and there were more differences in inclusion criteria between ABCD versions than expected. Consequently, three 'QC' conditions were added in addition to the motion thresholds, to provide additional detail to inform study design.
- The originally planned approach to evaluate H2 was found to be infeasible (See 'H2 Model,' above). In practice, a visual inspection of the data did not support H2 (see H2 Marginal Means Plotting). Specifically, because so much data was missing, biases in the missing data were self-correcting as more data was excluded. I.e., in the event that males were more likely to be excluded than females (which appears to be true in this dataset), the bias is strongest as data is first excluded

Table 17: Granular Race/Ethnicity Variables: Bivariate Models, Pt. 1

	Т	$\mathbf{C}$	0.5	0.4	0.3	R	0.2	0.1
	OR							
	[90% CI]							
	p	p	p	p	p	p	p	p
White	.66***	.58***	.59***	.59***	.58***	.64***	.60***	.56***
	.5580	.5364	.5365	.5465	.5364	.5871	.5565	.5161
	<.001	<.001	<.001	<.001	<.001	<.001	<.001	<.001
Black	1.43***	1.43***	1.47***	1.47***	1.47***	1.53***	1.51***	1.67***
	1.17-1.74	1.29-1.59	1.33-1.62	1.33-1.63	1.33-1.62	1.37-1.70	1.37-1.65	1.52-1.83
	<.001	<.001	<.001	<.001	<.001	<.001	<.001	<.001
American Indian	1.07 .65-1.67 .769	1.04 .80-1.32 .779	1.07 .84-1.35 .570	1.10 .87-1.38 .419	1.12 .89-1.40 .338	1.16 .90-1.47 .240	1.02 .82-1.27 .838	.97 .80-1.19 .798
Alaska Native	14.58**	6.33*	14.61*	14.19*	13.12*	6.43*	10.11*	3.17
	1.92-88.19	1.05-48.11	2.16-285.80	2.10-277.71	1.94-256.66	1.06-48.83	1.50-197.83	.47-62.03
	.003	.043	.016	.018	.021	.042	.039	.302
Native Hawaiian	.99 .06-4.73 .993	.63 .15-1.85 .459	.55 .13-1.60 .329	.53 .13-1.55 .307	.49 .12-1.43 .250	.90 .26-2.40 .850	.70 .23-1.76 .483	2.25 .93-6.24 .088
Guamanian	.00	.00	.00	.00	.00	451815.50	.00	.79
	NA-3.46e+11	NA-125672.14	NA-108642.02	NA-105570.45	NA-97579.29	.00-NA	NA-75234.92	.03-20.04
	.967	.942	.941	.941	.941	.926	.939	.869

Table 18: Granular Race/Ethnicity Variables: Bivariate Models, Pt. 2

	T	С	0.5	0.4	0.3	R	0.2	0.1
	OR	OR	OR	OR	OR	OR	OR	OR
	[90% CI]	[90% CI]	[90% CI]	[90% CI]	[90% CI]	[90% CI]	[90% CI]	[90% CI]
	p	p	p	p	p	p	P	p
Samoan	.00	.38	.33	.32	.30	.00	.23	.57
	NA-835.53	.02-1.97	.02-1.70	.02-1.66	.02-1.53	NA34	.01-1.18	.17-1.77
	.964	.358	.290	.278	.246	.938	.158	.331
Pacific Islander	2.43	1.23	1.22	1.18	1.09	1.43	1.22	1.07
	.73-6.10	.55-2.47	.56-2.41	.55-2.34	.51-2.16	.66-2.83	.61-2.32	.57-2.04
	.093	.592	.592	.648	.810	.329	.562	.828
Asian Indian	1.21	1.01	1.08	1.05	1.07	1.14	1.12	1.01
	.47-2.54	.62-1.58	.68-1.65	.66-1.60	.68-1.62	.71-1.77	.74-1.66	.70-1.48
	.647	.971	.737	.837	.767	.563	.579	.940
Filipino	1.38	1.42*	1.29	1.29	1.28	1.22	1.28	.90
	.72-2.38	1.02-1.95	.94-1.76	.93-1.75	.94-1.74	.87-1.69	.95-1.71	.68-1.19
	.287	.030	.110	.112	.112	.237	.103	.454
Chinese	1.40	1.47*	1.39	1.35	1.33	1.01	1.11	1.31
	.69-2.53	1.03-2.07	.98-1.95	.95-1.89	.94-1.85	.68-1.48	.79-1.54	.96-1.81
	.311	.030	.058	.083	.101	.941	.531	.090
Japanese	.53	1.69*	1.55	1.51	1.56	.71	1.41	.83
	.09-1.68	1.03-2.69	.95-2.45	.93-2.38	.97-2.44	.37-1.26	.89-2.18	.54-1.28
	.376	.030	.067	.087	.059	.277	.136	.395

Table 19: Granular Race/Ethnicity Variables: Bivariate Models, Pt. 3

	T	C	0.5	0.4	0.3	R	0.2	0.1
	OR	OR	OR	OR	OR	OR	OR	OR
	[90% CI]	[90% CI]	[90% CI]	[90% CI]	[90% CI]	[90% CI]	[90% CI]	[90% CI]
	p	p	p	p	p	p	p	p
Korean	.44	1.05	1.03	1.00	1.03	.75	1.08	.86
	.07-1.40	.63-1.69	.62-1.62	.61-1.58	.64-1.61	.42-1.27	.70-1.65	.58-1.27
	.255	.831	.907	.998	.885	.314	.716	.444
Vietnamese	1.09	1.44	1.46	1.53	1.53	1.11	1.26	.99
	.27-2.95	.79-2.49	.82-2.48	.87-2.59	.88-2.56	.58-1.99	.73-2.11	.60-1.64
	.884	.210	.175	.120	.119	.730	.382	.970
Other Asian	1.01	.91	1.11	1.08	1.13	1.31	1.14	1.04
	.31-2.44	.51-1.52	.66-1.78	.65-1.73	.69-1.78	.78-2.09	.72-1.77	.68-1.58
	.979	.737	.675	.761	.627	.287	.563	.867
Other	1.13	1.65***	1.55***	1.56***	1.55***	1.03	1.45***	1.41***
	.79-1.55	1.40-1.94	1.32-1.81	1.33-1.83	1.32-1.81	.86-1.23	1.25-1.69	1.22-1.64
	.486	<.001	<.001	<.001	<.001	.744	<.001	<.001
Refuse	1.17	1.86*	1.88*	1.82*	1.95*	1.46	2.00**	2.14**
	.28-3.18	1.04-3.19	1.07-3.18	1.04-3.09	1.13-3.29	.79-2.57	1.18-3.34	1.23-3.91
	.793	.029	.023	.029	.013	.205	.009	.010

from the dataset. Once a significantly larger proportion of males have been excluded than females, the over-representation of females in the sample results in their being more likely to be excluded. At the extreme, when all of the data is excluded, there is no bias).

Here, we present the analyses originally proposed as written, excepting H2. Specifically, we present results from adjusted models without the BMI variable in the motion scrubbing conditions.

```
model.h1prereg.5mm<-glm(!gframes_0.5_375 ~ sex + household.income + p.edu + race_ethnicity.factor + ksads_factor + pds_category + adi.z + coi.z -
model.h1prereg.5mm.results<-cbind(confint(model.h1prereg.5mm,level=.95),model.h1prereg.5mm$coefficients)[,c(1,3,2)]
model.h1prereg.5mm.results<-cbind(model.h1prereg.5mm.results,pcorrect(coef(summary(model.h1prereg.5mm))[,4]))
model.h1prereg.4mm<-glm(</pre>
  !gframes_0.4_375 ~ sex + household.income + p.edu + race_ethnicity.factor + ksads_factor + pds_category + adi.z + coi.z + nihtbx_flanker_agecon
  data=df.
  family="binomial")
model.h1prereg.4mm.results<-cbind(confint(model.h1prereg.4mm,level=.95),model.h1prereg.4mm$coefficients)[,c(1,3,2)]
model.h1prereg.4mm.results<-cbind(model.h1prereg.4mm.results,pcorrect(coef(summary(model.h1prereg.4mm))[,4]))
model.h1prereg.3mm<-glm(</pre>
  !gframes 0.3 375 ~ sex + household.income + p.edu + race ethnicity.factor + ksads factor + pds category + adi.z + coi.z + nihtbx flanker agecon
  data=df,
  family="binomial")
model.h1prereg.3mm.results<-cbind(confint(model.h1prereg.3mm,level=.95),model.h1prereg.3mm$coefficients)[,c(1,3,2)]
model.h1prereg.3mm.results<-cbind(model.h1prereg.3mm.results,pcorrect(coef(summary(model.h1prereg.3mm))[,4]))
model.h1prereg.2mm<-glm(</pre>
  !gframes_0.2_375 ~ sex + household.income + p.edu + race_ethnicity.factor + ksads_factor + pds_category + adi.z + coi.z + nihtbx_flanker_agecon
  data=df,
  family="binomial")
model.h1prereg.2mm.results<-cbind(confint(model.h1prereg.2mm,level=.95),model.h1prereg.2mm$coefficients)[,c(1,3,2)]
model.h1prereg.2mm.results<-cbind(model.h1prereg.2mm.results,pcorrect(coef(summary(model.h1prereg.2mm))[,4]))
model.h1prereg.1mm<-glm(
  !gframes 0.1 375 ~ sex + household.income + p.edu + race ethnicity.factor + ksads factor + pds category + adi.z + coi.z + nihtbx flanker agecomes
  data=df,
  family="binomial")
model.h1prereg.1mm.results<-cbind(confint(model.h1prereg.1mm,level=.95),model.h1prereg.1mm$coefficients)[,c(1,3,2)]
model.h1prereg.1mm.results<-cbind(model.h1prereg.1mm.results,pcorrect(coef(summary(model.h1prereg.1mm))[,4]))
```

#### 3.0.1 Pre-registered Condition (Adjusted) Model Tables

```
Mformat2<-\(m) {
  stars<-sdlabFunctions::starPs(m[,4])
  paste0(ORformat(m[,2]),</pre>
```

```
stars,
         " [", ORformat(m[,1]), "-", ORformat(m[,3]), "]")
}
h1modelspreregtable<-cbind(</pre>
                     Mformat2(model.h1prereg.5mm.results),
                     Mformat2(model.h1prereg.4mm.results),
                     Mformat2(model.h1prereg.3mm.results),
                     Mformat2(model.h1prereg.2mm.results),
                     Mformat2(model.h1prereg.1mm.results))
#h1modelspreregtable <- gsub("\n", "\\newline", h1modelspreregtable)</pre>
h1modelspreregtable[h1modelspreregtable==".000"]<-"<.001"
alllevels<-c("Intercept",
      "Sex (Male)",
      " $0-$25k",
      " $25-$50k",
      " $50-$75k",
      " $75-$100k",
      " >$200k",
      " <High School",
     " HS Grad.",
      " Some College",
      " Graduate",
      " Black",
      " Hispanic",
      " Asian",
      " Other",
      " 1 Trauma",
      " >=2 Trauma",
      " Early Puberty",
      " Mid Puberty",
      " Late Puberty",
      " Post Pubertal",
      "Area Disadvantage",
      "Child Opportunity",
      "NIHTB Flanker",
      "NIHTB Crystalized",
      "NIHTB Total",
      "WISC V Matrix",
      "Psychopathology",
      "Internalizing",
```

```
"Externalizing",
      "Age"
h1prereghux <-as_hux(h1modelspreregtable) %>%
  insert column(alllevels) %>%
  insert row("Household Income (ref: $100-$200k)",after=2,colspan=6,fill="") %%
  insert row("Highest Parental Education (ref: College Degree)",after=8,colspan=6,fill="") %>%
  insert row("Census Race/Ethnicity (ref: White)", after=13, colspan=6, fill="") %>%
  insert row("KSADS Trauma Count (ref: 0 Exposures)", after=18, colspan=6, fill="") %>%
  insert row("Pubertal Status (ref: pre-pubertal)",after=21,colspan=6,fill="") %>%
  insert row(c("Variable",rep(c("OR [95% CI]"),5)),after=0) %>%
  insert_row(c("",".5",".4",".3",".2",".1")) %>%
  set wrap(TRUE) %>%
  set width(1) %>%
  set_font_size(value=8)
colnames(h1prereghux)[1] <- "h1"</pre>
h1prereghux[c(1:2,3:20),] %>%
  set_caption("Adjusted Models Output, as Pre-registered (Pt. 1). ***: p<.001; **: p<.01; *:p<.05") %>% set_label("PreRegModel1")
h1prereghux[c(1:2,21:38),] %>%
  set_caption("Adjusted Models Output, as Pre-registered (Pt. 2). ***: p<.001; **: p<.01; *:p<.05") %>% set_label("PreRegModel2")
```

## 4 Save Tables

```
openxlsx::saveWorkbook(tables.xlsx,"Manuscript/Tables/tables.xlsx",overwrite=TRUE)
```

### 5 SessionInfo

```
R version 4.3.3 (2024-02-29)

Platform: x86_64-pc-linux-gnu (64-bit)

locale: LC_CTYPE=en_US.UTF-8, LC_NUMERIC=C, LC_TIME=en_US.UTF-8, LC_COLLATE=en_US.UTF-8, LC_MONETARY=en_US.UTF-8, LC_MESSAGES=en_US.UTF-8, LC_PAPER=en_US.UTF-8, LC_NAME=C, LC_ADDRESS=C, LC_TELEPHONE=C, LC_MEASUREMENT=en_US.UTF-8 and LC_IDENTIFICATION=C

attached base packages: stats, graphics, grDevices, utils, datasets, methods and base
```

Table 20: Adjusted Models Output, as Pre-registered (Pt. 1). \*\*\*: p<.001; \*\*: p<.01; \*:p<.05

	0.5	0.4	0.3	0.2	0.1
Variable	OR [95% CI]				
Intercept	.18*** [.1522]	.18*** [.1622]	.20*** [.1723]	.25*** [.2129]	.80** [.7092]
Sex (Male)	1.09 [.96-1.23]	1.11 [.99-1.26]	1.17** [1.04-1.32]	1.31*** [1.17-1.46]	1.42*** [1.29-1.57]
Household Income (ref: \$100-\$	2200k)				
\$0-\$25k	1.15 [.93-1.44]	1.17 [.94-1.45]	1.11 [.90-1.38]	1.13 [.93-1.38]	1.06 [.88-1.28]
\$25-\$50k	.89 [.73-1.09]	.91 [.75-1.11]	.92 [.76-1.12]	.87 [.73-1.04]	.89 [.76-1.05]
\$50-\$75k	.92 [.76-1.10]	.92 [.76-1.10]	.92 [.77-1.10]	.93 [.79-1.10]	1.09 [.94-1.26]
\$75-\$100k	.96 [.80-1.14]	.96 [.81-1.13]	.98 [.83-1.16]	.94 [.80-1.09]	.92 [.80-1.05]
>\$200k	.98 [.81-1.18]	.98 [.82-1.18]	1.00 [.83-1.19]	.98 [.82-1.15]	.96 [.82-1.11]
Highest Parental Education (re	ef: College Degree)				
<high school<="" td=""><td>1.49** [1.10-2.00]</td><td>1.50** [1.11-2.01]</td><td>1.44* [1.07-1.93]</td><td>1.49** [1.12-1.97]</td><td>1.35* [1.02-1.79]</td></high>	1.49** [1.10-2.00]	1.50** [1.11-2.01]	1.44* [1.07-1.93]	1.49** [1.12-1.97]	1.35* [1.02-1.79]
HS Grad.	1.25 [.99-1.58]	1.24 [.99-1.56]	1.26* [1.01-1.58]	1.29* [1.04-1.59]	1.23* [1.01-1.50]
Some College	1.02 [.86-1.20]	1.03 [.87-1.21]	1.03 [.88-1.20]	1.10 [.95-1.27]	1.03 [.91-1.18]
Graduate	1.13 [.98-1.30]	1.12 [.98-1.29]	1.12 [.98-1.29]	1.14 [1.00-1.29]	1.04 [.93-1.17]
Census Race/Ethnicity (ref: W	White)				
Black	1.69*** [1.40-2.04]	1.63*** [1.35-1.97]	1.59*** [1.32-1.91]	1.53*** [1.29-1.82]	1.48*** [1.26-1.75]
Hispanic	1.20* [1.02-1.40]	1.19* [1.02-1.39]	1.21* [1.04-1.40]	1.14 [.99-1.32]	1.16* [1.02-1.32]
Asian	1.92*** [1.38-2.64]	1.88*** [1.35-2.58]	2.05*** [1.48-2.80]	1.96*** [1.44-2.66]	1.49** [1.10-2.01]
Other	1.34*** [1.13-1.59]	1.34*** [1.13-1.59]	1.28** [1.08-1.52]	1.18* [1.00-1.38]	1.12 [.97-1.29]

Table 21: Adjusted Models Output, as Pre-registered (Pt. 2). \*\*\*: p<.001; \*\*: p<.01; \*:p<.05

	0.5	0.4	0.3	0.2	0.1
Variable	OR [95% CI]				
KSADS Trauma Count (ref: 0	Exposures)				
1 Trauma	1.06 [.93-1.19]	1.05 [.93-1.19]	1.05 [.93-1.18]	1.03 [.92-1.15]	.96 [.87-1.06]
>=2 Trauma	1.12 [.93-1.34]	1.08 [.90-1.29]	1.06 [.89-1.26]	1.13 [.96-1.33]	1.20* [1.03-1.40]
Pubertal Status (ref: pre-puber	rtal)				
Early Puberty	1.02 [.90-1.17]	1.03 [.90-1.17]	1.01 [.89-1.15]	1.07 [.95-1.20]	1.07 [.96-1.19]
Mid Puberty	1.05 [.90-1.24]	1.06 [.90-1.24]	1.07 [.91-1.24]	1.08 [.94-1.25]	1.25*** [1.10-1.42]
Late Puberty	1.73** [1.16-2.55]	1.71** [1.15-2.52]	1.61* [1.08-2.38]	1.63* [1.11-2.37]	1.31 [.91-1.88]
Post Pubertal	.47 [.02-2.73]	.47 [.02-2.70]	.45 [.02-2.59]	1.55 [.31-6.50]	1.37 [.33-6.82]
Area Disadvantage	.70*** [.6477]	.72*** [.6678]	.73*** [.6779]	.76*** [.7183]	.86*** [.8093]
Child Opportunity	.78*** [.7186]	.80*** [.7388]	.80*** [.7387]	.82*** [.7590]	.84*** [.7892]
NIHTB Flanker	1.02 [.95-1.09]	1.01 [.95-1.09]	1.03 [.96-1.10]	1.02 [.96-1.09]	1.01 [.96-1.07]
NIHTB Crystalized	1.06 [.94-1.18]	1.05 [.94-1.17]	1.08 [.97-1.21]	1.09 [.99-1.21]	1.16** [1.06-1.27]
NIHTB Total	.88 [.77-1.00]	.88 [.78-1.00]	.84* [.7496]	.84** [.7494]	.81*** [.7390]
WISC V Matrix	1.00 [.95-1.06]	.99 [.93-1.05]	.98 [.93-1.04]	.95 [.90-1.00]	.95* [.9099]
Psychopathology	1.11*** [1.05-1.17]	1.11*** [1.05-1.17]	1.11*** [1.06-1.17]	1.13*** [1.07-1.18]	1.11*** [1.06-1.16]
Internalizing	.98 [.92-1.03]	.97 [.92-1.02]	.96 [.91-1.01]	.95 [.91-1.00]	.92*** [.8896]
Externalizing	.94* [.8999]	.94* [.8999]	.94* [.9099]	.94* [.8999]	.94** [.9098]
Age	.85*** [.8190]	.84*** [.8089]	.83*** [.7988]	.82*** [.7886]	.81*** [.7885]

 $\begin{array}{l} \textbf{other attached packages:} \ \ ggtext(v.0.1.2), \ \ emmeans(v.1.8.9), \ \ lmerTest(v.3.1-3), \ \ lme4(v.1.1-35.1), \ \ Matrix(v.1.6-5), \ \ ggupset(v.0.3.0), \ \ openxlsx(v.4.2.5.2), \ \ patchwork(v.1.1.3), \ \ viridis(v.0.6.4), \ \ viridisLite(v.0.4.2), \ \ huxtable(v.5.5.2), \ \ hexbin(v.1.28.3), \ \ pander(v.0.6.5), \ \ ggExtra(v.0.10.1), \ \ ggthemes(v.4.2.4), \ \ lubridate(v.1.9.3), \ \ forcats(v.1.0.0), \ \ stringr(v.1.5.0), \ \ dplyr(v.1.1.3), \ \ purrr(v.1.0.2), \ \ readr(v.2.1.4), \ \ tidyr(v.1.3.0), \ \ tibble(v.3.2.1), \ \ ggplot2(v.3.4.4), \ \ tidyverse(v.2.0.0) \ \ and \ \ rmarkdown(v.2.25) \end{array}$ 

loaded via a namespace (and not attached): gridExtra(v.2.3), gld(v.2.6.6), readxl(v.1.4.3), rlang(v.1.1.1), magrittr(v.2.0.3), e1071(v.1.7-13), compiler(v.4.3.3), mgcv(v.1.9-1), systemfonts(v.1.0.5), vctrs(v.0.6.4), pkgconfig(v.2.0.3), crayon(v.1.5.2), fastmap(v.1.1.1), backports(v.1.4.1), ellipsis(v.0.3.2), labeling(v.0.4.3), utf8(v.1.2.3), ggstance(v.0.3.6), promises(v.1.2.1), markdown(v.1.11), tzdb(v.0.4.0), nloptr(v.2.0.3), ragg(v.1.2.6), xfun(v.0.43), cachem(v.1.0.8), later(v.1.3.1), broom(v.1.0.5), DescTools(v.0.99.51), R6(v.2.5.1), stringi(v.1.7.12), boot(v.1.3-28), extrafontdb(v.1.0), cellranger(v.1.1.0), numDeriv(v.2016.8-1.1), estimability(v.1.4.1), Rcpp(v.1.0.11), assertthat(v.0.2.1), knitr(v.1.44), parameters(v.0.21.2), extrafont(v.0.19), httpuv(v.1.6.12), splines(v.4.3.3), timechange(v.0.2.0), tidyselect(v.1.2.0), rstudioapi(v.0.15.0), yaml(v.2.3.7), codetools(v.0.2-19), miniUI(v.0.1.1.1), plyr(v.1.8.9), lattice(v.0.22-4), shiny(v.1.7.5.1), withr(v.2.5.1), bayestestR(v.0.13.1), coda(v.0.19-4), evaluate(v.0.22), proxy(v.0.4-27), zip(v.2.3.0), xml2(v.1.3.5), pillar(v.1.9.0), insight(v.0.19.6), generics(v.0.1.3), sdlab-Functions(v.0.2.0), hms(v.1.1.3), munsell(v.0.5.0), commonmark(v.1.9.0), scales(v.1.2.1), rootSolve(v.1.8.2.4), minqa(v.1.2.6), xtable(v.1.8-4), class(v.7.3-22), glue(v.1.6.2), lmom(v.3.0), tools(v.4.3.3), data.table(v.1.14.8), Exact(v.3.2), mvtnorm(v.1.2-3), grid(v.4.3.3), Rttf2pt1(v.1.3.12), datawizard(v.0.9.0), colorspace(v.2.1-0), nlme(v.3.1-163), performance(v.0.10.5), cli(v.3.6.1), textshaping(v.0.3.7), fansi(v.1.0.5), expm(v.0.999-7), svglite(v.2.1.2), gtable(v.0.3.4), digest(v.0.6.33), farver(v.2.1.1), memoise(v.2.0.1), htmlools(v.0.5.6.1), lifecycle(v.1.0.3), httr(v.1.4.7), mine(v.0.12), gridtext(v.0.1.5) and MASS(v.7.3-60.0.1)