

Causal Inference for Cognitive Decline Using Mixed Effects Models

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Abstract

In [1], the authors analyze a sample of longitudinal data from the OASIS (Open Access Series of Imaging Studies, see [2]) dataset, consisting of 150 subjects between the ages of 60 to 96. Subjects are categorized by their CDR (Clinical Dementia Ratio), being either nondemented ($CDR = 0$), having very mild Alzheimer's disease, ($CDR = 0.5$) or having mild Alzheimer's disease ($CDR = 1$). The authors analyze the progression of brain volume atrophy in Alzheimer's disease. In this report, we use the same dataset (retrieved from [3]) to study the progression of cognitive decline in Alzheimer's disease using MMSE (Mini-Mental State Examination) score as a proxy for cognitive decline. We model MMSE using a linear mixed effects model on various predictors in the dataset for the purpose of examining causes of cognitive decline, and in doing so we explore the limitations of mixed effect models on non-normal, heavily skewed data.

Introduction

Clinical Dementia Rating (CDR) is a system used to stratify the severity of Alzheimer's disease into stages. Stages include 0: nondemented, 0.5: questionable (or "very mild" as described in [1]), 1: mild, 2: moderate, 3: severe, and scoring is based on an interview with the subject, assessing impairment in the categories of memory, orientation, judgement and problem solving, community affairs, home and hobbies, and personal care [4].

The Mini-Mental State Examination (MMSE) is a test used to assess cognitive ability of patients with dementia or various psychiatric conditions. It has been found to accurately distinguish patients with cognitive deficiencies from those without, and MMSE scores correlate with scores in other standard cognition tests [5], making it a suitable proxy for modelling cognitive decline.

We will model MMSE using features in the dataset to assess predictors for cognitive decline. Determining strongest predictors for cognitive decline allows for earlier diagnosis of Alzheimer's disease. Earlier diagnosis allows for earlier treatment, slowing disease progression and reducing financial strain on healthcare systems [6]. We will build a linear mixed effects (LME) model. LMEs are effective at modelling data that is "clustered", such as longitudinal data; in this case, the "clusters" are the individual subjects, each having multiple visits. LMEs combine "fixed effects" and "random effects", hence the "mixed" part of the name. Fixed effects are effects that are assumed to be consistent across individuals while random effects are effects that differ for individuals, allowing us to account for the fact that different subjects have different baselines. LMEs are typically most effective when the response variable is normally distributed. With our data, we will see that MMSE is not normally distributed, and we will explore the limitations that arise and attempt to improve our model with various diagnostic methods. We will discuss possible alternative models that offer some hope of addressing these limitations.

Dataset

Features of interest in the dataset for our modelling include Subject ID (Subject.ID), visit number (Visit), sex (M.F), age at visit (Age), years of education (EDUC), socioeconomic status (SES), MMSE score (MMSE), CDR score (CDR), and normalized whole brain volume (nWBV). All subjects are right handed. Subjects were screened to ensure exclusion of subjects with physiological causes of dementia other than Alzheimer's disease [1].

```
df <- read.csv("D:/Daniel/Documents/MATH6642/final_project/oasis_longitudinal.csv")
dc <- df[!is.na(df$MMSE) & !is.na(df$SES), ]

#Note: SES is ordered lowest = 5 and highest = 1; we will reverse the ordering
dc$SES <- 5 - dc$SES

dc <- dc[order(dc$Subject.ID, dc$Age), ]

#The head of the dataframe:
print(head(dc))
```

##	Subject.ID	MRI.ID	Group	Visit	MR.Delay	M.F	Hand	Age	EDUC	SES
## 1	OAS2_0001	OAS2_0001_MR1	Nondemented	1	0	M	R	87	14	3
## 2	OAS2_0001	OAS2_0001_MR2	Nondemented	2	457	M	R	88	14	3
## 6	OAS2_0004	OAS2_0004_MR1	Nondemented	1	0	F	R	88	18	2
## 7	OAS2_0004	OAS2_0004_MR2	Nondemented	2	538	F	R	90	18	2
## 8	OAS2_0005	OAS2_0005_MR1	Nondemented	1	0	M	R	80	12	1
## 9	OAS2_0005	OAS2_0005_MR2	Nondemented	2	1010	M	R	83	12	1

##	MMSE	CDR	eTIV	nWBV	ASF
## 1	27	0.0	1987	0.696	0.883
## 2	30	0.0	2004	0.681	0.876
## 6	28	0.0	1215	0.710	1.444
## 7	27	0.0	1200	0.718	1.462
## 8	28	0.0	1689	0.712	1.039
## 9	29	0.5	1701	0.711	1.032

Education level is slightly right-skewed. Socioeconomic status is concentrated towards higher levels. Sex is slightly skewed towards females.

```

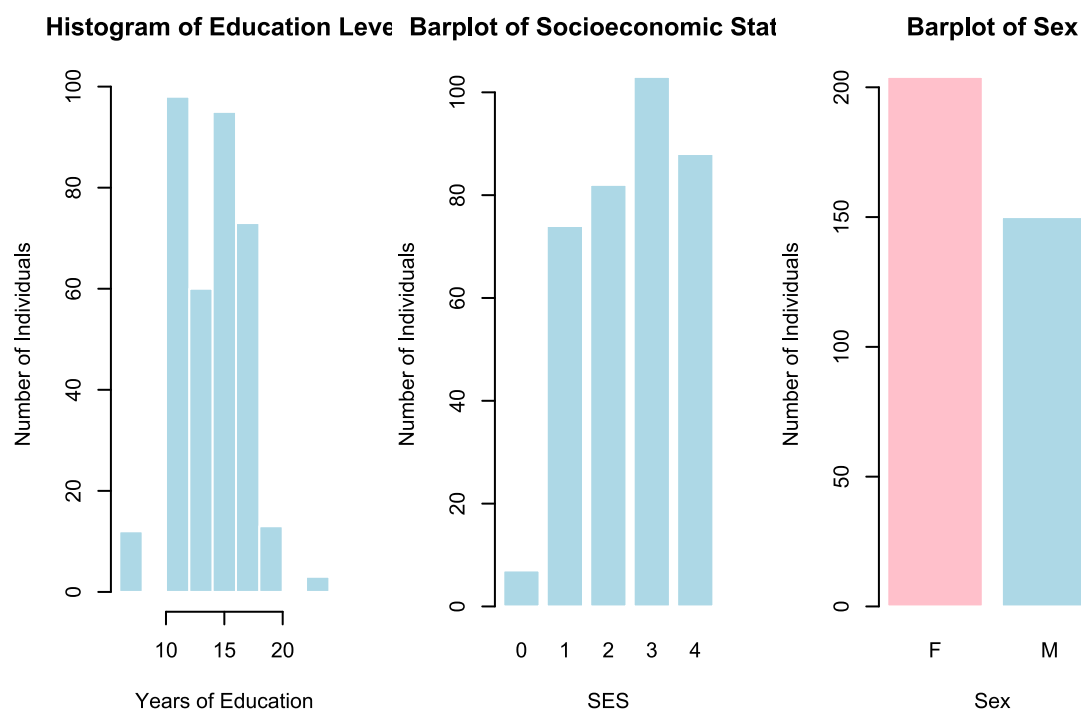
par(mfrow = c(1, 3))

hist(dc$EDUC,
     breaks = 10,
     main = "Histogram of Education Level",
     xlab = "Years of Education",
     ylab = "Number of Individuals",
     col = "lightblue",
     border = "white")

barplot(table(dc$SES),
       main = "Barplot of Socioeconomic Status",
       xlab = "SES",
       ylab = "Number of Individuals",
       col = "lightblue",
       border = "white")

barplot(table(dc$M.F),
       main = "Barplot of Sex",
       xlab = "Sex",
       ylab = "Number of Individuals",
       col = c("pink", "lightblue"),
       border = "white")

```



MMSE is heavily left-skewed. Comparing initial visit to final visit MMSE distribution, we see a decline in average MMSE.

```

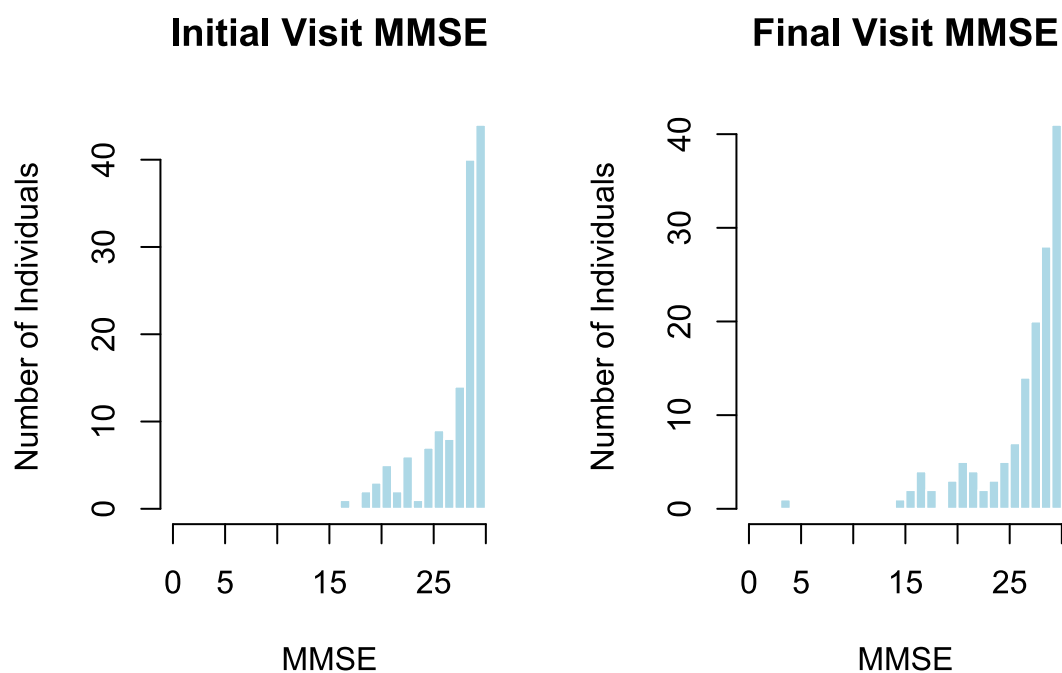
initial_visits <- dc[ave(dc$Visit, dc$Subject.ID, FUN = min) == dc$Visit, ]
final_visits <- dc[ave(dc$Visit, dc$Subject.ID, FUN = max) == dc$Visit, ]

#Set layout: 1 row, 2 columns
par(mfrow = c(1, 2))

#Histogram for initial visits
hist(initial_visits$MMSE,
      main = "Initial Visit MMSE",
      breaks = 0:30,
      xlab = "MMSE",
      ylab = "Number of Individuals",
      col = "lightblue",
      border = "white",
      xlim = c(0, 30))

#Histogram for final visits
hist(final_visits$MMSE,
      main = "Final Visit MMSE",
      breaks = 0:30,
      xlab = "MMSE",
      ylab = "Number of Individuals",
      col = "lightblue",
      border = "white",
      xlim = c(0, 30))

```



We plot the longitudinal trajectories of nWBV, recreating the same plot as shown in [1]:

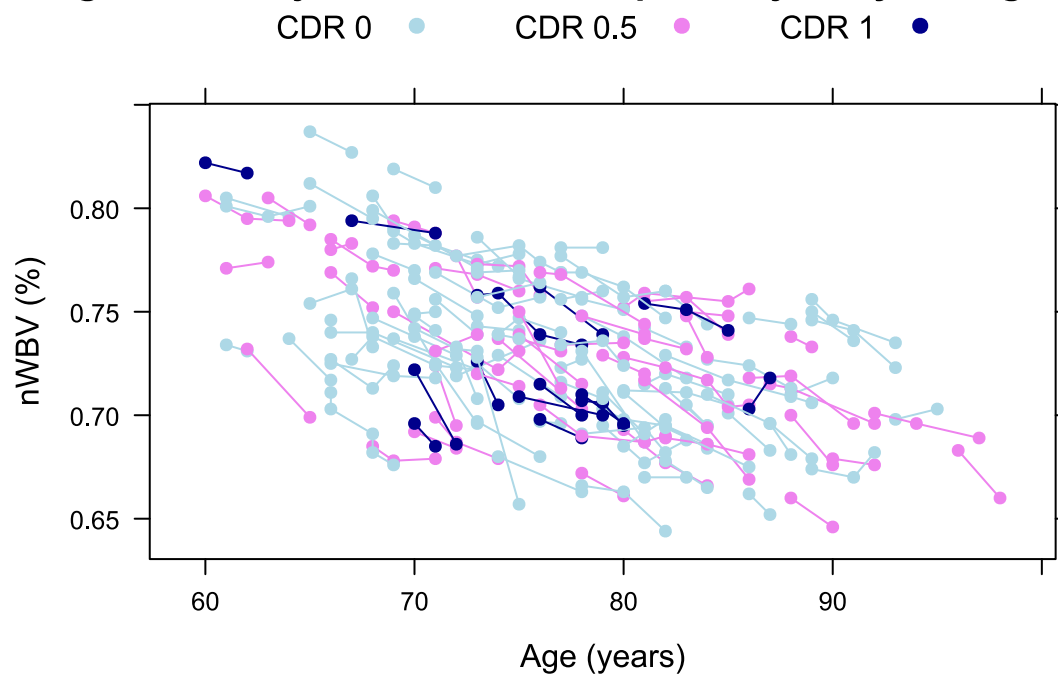
```

cdr_colors <- c("0" = "lightblue", "0.5" = "violet", "1" = "darkblue")

#nWBV trajectories
xyplot(nWBV ~ Age, data = dc,
      groups = Subject.ID,
      type = "b",
      lwd = 1,
      pch = 16,
      col = cdr_colors[as.character(dc$CDR)],
      xlab = "Age (years)",
      ylab = "nWBV (%)",
      main = "Longitudinal trajectories of nWBV per subject by CDR group",
      key = list(text = list(c("CDR 0", "CDR 0.5", "CDR 1")),
                points = list(pch = 16, col = c("lightblue", "violet", "darkblue")),
                columns = 3))

```

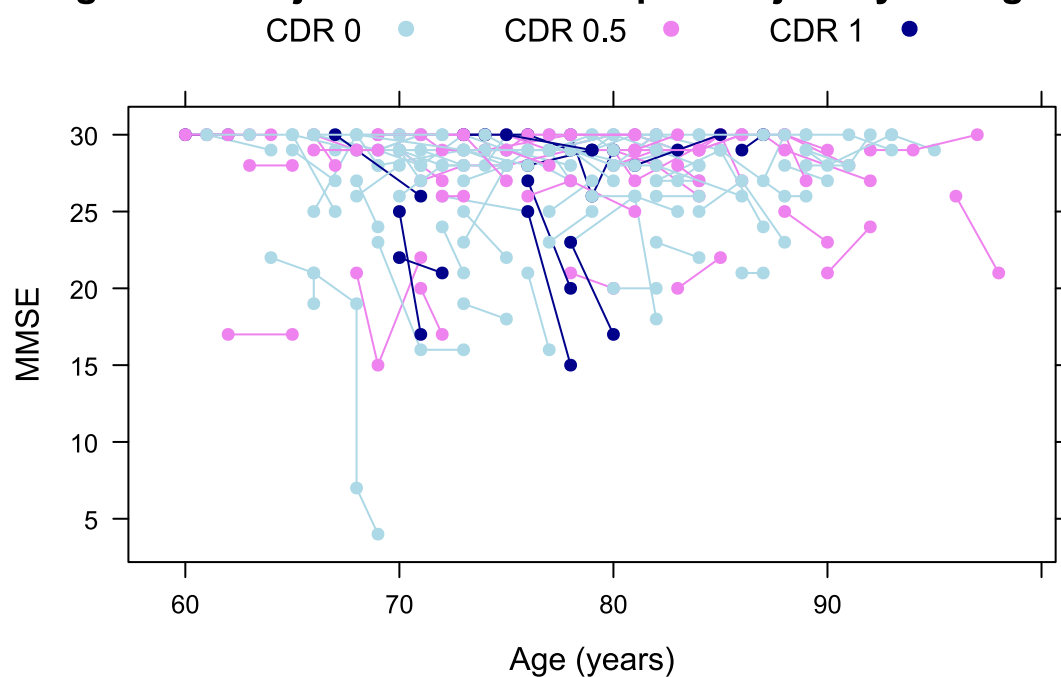
Longitudinal trajectories of nWBV per subject by CDR group



And we analogously plot the longitudinal trajectories of MMSE:

```
#MMSE trajectories
xyplot(MMSE ~ Age, data = dc,
      groups = Subject.ID,
      type = "b",
      lwd = 1,
      pch = 16,
      col = cdr_colors[as.character(dc$CDR)],
      xlab = "Age (years)",
      ylab = "MMSE",
      main = "Longitudinal trajectories of MMSE per subject by CDR group",
      key = list(text = list(c("CDR 0", "CDR 0.5", "CDR 1")),
                points = list(pch = 16, col = c("lightblue", "violet", "darkblue")),
                columns = 3))
```

Longitudinal trajectories of MMSE per subject by CDR group



Results

Modelling Random Effects

For our fixed effects, we tentatively include terms nWBV, Age, EDUC, SES, M.F. We begin by testing random effects. For random effects, we consider time-variant variables; in this case, nWBV and Age. We make the reasonable assumption that EDUC and SES are time-invariant for an elderly population.

Between random effects 1, 1 + Age and 1 + nWBV, ANOVA testing produces the lowest AIC for 1 + nWBV. So we proceed with the following model:

```
fit2 <- lme(MMSE ~ nWBV + Age + M.F + SES + EDUC,  
           data = dc,  
           random = ~ 1 + nWBV | Subject.ID,  
           correlation = corAR1(form = ~ 1 | Subject.ID),  
           control = lmeControl(opt = "optim", maxIter = 200, msMaxIter = 200))
```

Examining this model produces p-values > 0.05 for all predictors other than nWBV.

```
summary(fit2)
```

```

## Linear mixed-effects model fit by REML
## Data: dc
##      AIC      BIC    logLik
## 1632.026 1674.4 -805.0128
##
## Random effects:
## Formula: ~1 + nWBV | Subject.ID
## Structure: General positive-definite, Log-Cholesky parametrization
##           StdDev   Corr
## (Intercept) 30.125603 (Intr)
## nWBV         38.256604 -1
## Residual     1.882025
##
## Correlation Structure: AR(1)
## Formula: ~1 | Subject.ID
## Parameter estimate(s):
##      Phi
## 0.3503827
## Fixed effects: MMSE ~ nWBV + Age + M.F + SES + EDUC
##           Value Std.Error DF   t-value p-value
## (Intercept) 0.294103 7.157217 210 0.041092 0.9673
## nWBV         29.823705 7.116504 210 4.190781 0.0000
## Age          0.043803 0.031988 210 1.369333 0.1724
## M.FM         -0.666225 0.442148 138 -1.506791 0.1342
## SES          0.093875 0.262842 138 0.357152 0.7215
## EDUC         0.138319 0.104192 138 1.327536 0.1865
## Correlation:
##      (Intr) nWBV  Age    M.FM  SES
## nWBV -0.947
## Age  -0.786 0.597
## M.FM -0.223 0.232 0.148
## SES  -0.027 0.103 0.017 0.093
## EDUC -0.097 -0.072 0.010 -0.138 -0.696
##
## Standardized Within-Group Residuals:
##      Min      Q1      Med      Q3      Max
## -3.9190676 -0.2946575 0.1071537 0.4679001 2.9629229
##
## Number of Observations: 354
## Number of Groups: 142

```


Modelling Fixed Effects

For our fixed effects, we now consider interaction terms. Plausible three-way interaction terms and corresponding hypotheses for consideration include the following:

1. nWBV * Age * SES: Higher SES allows for access to higher quality healthcare, slowing decay in nWBV during the aging process
2. nWBV * Age * M.F: Decay in nWBV during the aging process differs between male and female for biological reasons
3. nWBV * Age * EDUC: Higher education slows the decay in nWBV during the aging process

Building separate models considering each interaction term and running ANOVA tests revealed the following model to have the lowest AIC:

```
fit3 <- lme(MMSE ~ nWBV + Age + M.F + SES + EDUC + nWBV * Age * SES
           + nWBV * Age + nWBV * SES + Age * SES,
           data = dc,
           random = ~ 1 + nWBV | Subject.ID,
           correlation = corAR1(form = ~ 1 | Subject.ID),
           control = lmeControl(opt = "optim", maxIter = 200, msMaxIter = 200))
```

However, all predictors in this model have p-value > 0.05, making it unsuitable for causal inference:

```
summary(fit3)
```

```

## Linear mixed-effects model fit by REML
##   Data: dc
##       AIC      BIC    logLik
## 1679.897 1737.507 -824.9487
##
## Random effects:
## Formula: ~1 + nWBV | Subject.ID
## Structure: General positive-definite, Log-Cholesky parametrization
##           StdDev   Corr
## (Intercept) 0.1954965 (Intr)
## nWBV         0.2027344 -0.371
## Residual     3.2745790
##
## Correlation Structure: AR(1)
## Formula: ~1 | Subject.ID
## Parameter estimate(s):
##      Phi
## 0.7649814
## Fixed effects: MMSE ~ nWBV + Age + M.F + SES + EDUC + nWBV * Age * SES + nWBV * Age + n
WBV * SES + Age * SES
##           Value Std.Error  DF    t-value p-value
## (Intercept) -109.64519  93.31457 206 -1.1750061  0.2413
## nWBV         159.75950 127.16399 206  1.2563266  0.2104
## Age          1.02400   1.22193 206  0.8380188  0.4030
## M.FM         -0.50329   0.53906 138 -0.9336385  0.3521
## SES          29.71438  32.38970 138  0.9174020  0.3605
## EDUC          0.20386   0.12943 138  1.5750715  0.1175
## nWBV:Age     -1.10089   1.68083 206 -0.6549678  0.5132
## nWBV:SES     -36.11864  44.43754 206 -0.8127956  0.4173
## Age:SES      -0.26514   0.42249 206 -0.6275692  0.5310
## nWBV:Age:SES  0.30722   0.58460 206  0.5255143  0.5998
## Correlation:
##           (Intr) nWBV   Age    M.FM   SES    EDUC   nWBV:Ag nWBV:S Ag:SES
## nWBV        -0.997
## Age         -0.991  0.994
## M.FM         0.115 -0.120 -0.126
## SES         -0.919  0.920  0.910 -0.153
## EDUC        -0.090  0.065  0.077 -0.091  0.060
## nWBV:Age     0.982 -0.991 -0.997  0.129 -0.905 -0.066
## nWBV:SES     0.913 -0.920 -0.910  0.151 -0.997 -0.057  0.911
## Age:SES     0.912 -0.918 -0.920  0.148 -0.990 -0.066  0.921  0.993
## nWBV:Age:SES -0.900  0.912  0.914 -0.145  0.981  0.055 -0.921 -0.990 -0.997
##
## Standardized Within-Group Residuals:
##           Min           Q1           Med           Q3           Max
## -6.6203240 -0.2772330  0.2388486  0.6347886  1.7999549
##
## Number of Observations: 354
## Number of Groups: 142

```

We discard three-way interaction terms and instead consider two-way interaction terms. Plausible two-way interaction terms and corresponding hypotheses include the following:

1. nWBV * Age: Brain volume changes with age
2. nWBV * EDUC: Education acts as a buffer against brain volume loss
3. nWBV * SES : Higher SES allows for better healthcare access, reducing brain volume loss
4. nWBV * M.F : Sex affects change in brain volume for biological reasons
5. Age * EDUC : Education acts as a buffer against age effects
6. Age * M.F : Aging affects males and females differently
7. Age * SES : Higher SES allows for better healthcare access, reducing aging effects

Modelling these hypotheses and assessing AIC values from ANOVA tests found this model to be the best performing:

```
fit5 <- lme(MMSE ~ nWBV + Age + SES + nWBV * SES + Age * SES,  
  data = dc,  
  random = ~ 1 + nWBV | Subject.ID,  
  correlation = corAR1(form = ~ 1 | Subject.ID),  
  control = lmeControl(opt = "optim", maxIter = 200, msMaxIter = 200))
```

And we see that all terms here are significant (p-value < 0.05). Note that sex and years of education were dropped as ANOVA testing found them to be insignificant:

```
summary(fit5)
```

```

## Linear mixed-effects model fit by REML
## Data: dc
##      AIC      BIC    logLik
## 1621.422 1663.797 -799.7112
##
## Random effects:
## Formula: ~1 + nWBV | Subject.ID
## Structure: General positive-definite, Log-Cholesky parametrization
##           StdDev   Corr
## (Intercept) 32.020859 (Intr)
## nWBV         40.961855 -1
## Residual     1.844624
##
## Correlation Structure: AR(1)
## Formula: ~1 | Subject.ID
## Parameter estimate(s):
##      Phi
## 0.312536
## Fixed effects: MMSE ~ nWBV + Age + SES + nWBV * SES + Age * SES
##           Value Std.Error DF   t-value p-value
## (Intercept) -54.94656 16.877249 208 -3.255658 0.0013
## nWBV         85.06516 16.963782 208  5.014516 0.0000
## Age          0.24594  0.079577 208  3.090611 0.0023
## SES          21.81791  6.125116 140  3.562041 0.0005
## nWBV:SES     -21.04513  6.232539 208 -3.376654 0.0009
## Age:SES      -0.07695  0.028539 208 -2.696233 0.0076
## Correlation:
##           (Intr) nWBV   Age    SES   nWBV:S
## nWBV       -0.958
## Age        -0.785  0.575
## SES        -0.919  0.884  0.716
## nWBV:SES    0.873 -0.916 -0.518 -0.958
## Age:SES     0.725 -0.531 -0.926 -0.775  0.563
##
## Standardized Within-Group Residuals:
##           Min           Q1           Med           Q3           Max
## -3.9307210 -0.2692234  0.1472228  0.4643059  2.8280703
##
## Number of Observations: 354
## Number of Groups: 142

```

Inspecting the random effects coefficients reveals that the correlation between the random slope and intercept is -1, which is suggestive of numerical instability in the random effects. Within-subject centering made no improvements, but including non-linear terms was successful in correlation improvement.

Non-linear Terms

To introduce non-linearity to the model, we initially added a squared nWBV term to the fixed effects:

```
dc$nWBV_scaled <- scale(dc$nWBV)[,1]
dc$nWBV_sq_scaled <- dc$nWBV_scaled^2

fit6 <- lmer(MMSE ~ nWBV_scaled + nWBV_sq_scaled + Age + SES +
             nWBV_scaled * SES + Age * SES +
             (1 + nWBV_scaled | Subject.ID),
             data = dc)
```

All terms are significant (p-value < 0.05), and the correlation in the random effects becomes a slightly more stable -0.96:

```
summary(fit6)
```

```
## Linear mixed model fit by REML. t-tests use Satterthwaite's method [
## lmerModLmerTest]
## Formula: MMSE ~ nWBV_scaled + nWBV_sq_scaled + Age + SES + nWBV_scaled *
##      SES + Age * SES + (1 + nWBV_scaled | Subject.ID)
##      Data: dc
##
## REML criterion at convergence: 1617.9
##
## Scaled residuals:
##      Min       1Q   Median       3Q      Max
## -4.1474 -0.2784  0.1098  0.4405  3.2477
##
## Random effects:
##      Groups      Name      Variance Std.Dev. Corr
##      Subject.ID (Intercept) 5.347    2.312
##              nWBV_scaled 2.600    1.612   -0.96
##      Residual              2.660    1.631
## Number of obs: 354, groups: Subject.ID, 142
##
## Fixed effects:
##              Estimate Std. Error      df t value Pr(>|t|)
## (Intercept)      7.78698    5.88637 154.69655   1.323 0.187826
## nWBV_scaled       3.59539    0.63605 139.35550   5.653 8.6e-08 ***
## nWBV_sq_scaled    -0.34260    0.14499  89.00378  -2.363 0.020309 *
## Age               0.24043    0.07588 147.81877   3.169 0.001861 **
## SES               6.92411    2.09132 147.38808   3.311 0.001170 **
## nWBV_scaled:SES   -0.87827    0.22917 136.90383  -3.832 0.000193 ***
## Age:SES          -0.08314    0.02711 142.29317  -3.067 0.002587 **
## ---
## Signif. codes:  0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
##
## Correlation of Fixed Effects:
##              (Intr) nWBV_s nWBV__ Age      SES      nWBV_:
## nWBV_scaled  -0.598
## nWBV_sq_scl  -0.110 -0.176
## Age          -0.995  0.556  0.097
## SES          -0.924  0.561  0.050  0.923
## nWBV_sc:SES  0.559 -0.907  0.026 -0.521 -0.609
## Age:SES      0.919 -0.519 -0.048 -0.926 -0.995  0.565
```

```
anova(fit6)
```

```
## Type III Analysis of Variance Table with Satterthwaite's method
##              Sum Sq Mean Sq NumDF   DenDF F value    Pr(>F)
## nWBV_scaled    84.990   84.990     1 139.356 31.9532 8.595e-08 ***
## nWBV_sq_scaled  14.851   14.851     1  89.004  5.5834 0.0203093 *
## Age             26.706   26.706     1 147.819 10.0406 0.0018612 **
## SES             29.157   29.157     1 147.388 10.9619 0.0011696 **
## nWBV_scaled:SES 39.067   39.067     1 136.904 14.6876 0.0001926 ***
## Age:SES         25.023   25.023     1 142.293  9.4076 0.0025871 **
## ---
## Signif. codes:  0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
```

Using splines more effectively captured non-linearity:

```
dc$nWBV_scaled <- scale(dc$nWBV)[,1]

fit_spline <- lmer(MMSE ~ ns(nWBV_scaled, df=4) + Age + SES + nWBV_scaled*SES + Age:SES +
  (1 + nWBV_scaled | Subject.ID),
  data = dc,
  REML = TRUE)
```

```
## fixed-effect model matrix is rank deficient so dropping 1 column / coefficient
```

And again, we see that all terms are significant (p-values < 0.05) with correlation in the random effects reduced to -0.95:

```
summary(fit_spline)
```

```

## Linear mixed model fit by REML. t-tests use Satterthwaite's method [
## lmerModLmerTest]
## Formula: MMSE ~ ns(nWBV_scaled, df = 4) + Age + SES + nWBV_scaled * SES +
##      Age:SES + (1 + nWBV_scaled | Subject.ID)
##      Data: dc
##
## REML criterion at convergence: 1598.2
##
## Scaled residuals:
##      Min      1Q  Median      3Q      Max
## -3.8821 -0.3099  0.1315  0.4339  3.1968
##
## Random effects:
##      Groups      Name      Variance Std.Dev. Corr
##      Subject.ID (Intercept) 5.511      2.347
##                  nWBV_scaled 2.717      1.648   -0.95
##      Residual                2.542      1.594
## Number of obs: 354, groups: Subject.ID, 142
##
## Fixed effects:
##
##              Estimate Std. Error      df t value Pr(>|t|)
## (Intercept)      -6.06696      7.01980 169.91995   -0.864 0.388661
## ns(nWBV_scaled, df = 4)1 12.59680      1.98711 238.51073    6.339 1.14e-09 ***
## ns(nWBV_scaled, df = 4)2 12.74060      2.12443 149.17160    5.997 1.45e-08 ***
## ns(nWBV_scaled, df = 4)3 28.71832      4.56269 241.57876    6.294 1.44e-09 ***
## ns(nWBV_scaled, df = 4)4 16.68518      3.08572 124.96132    5.407 3.13e-07 ***
## Age                0.25381      0.07641 152.53105    3.322 0.001120 **
## SES                7.11217      2.10048 149.27436    3.386 0.000907 ***
## SES:nWBV_scaled     -0.90454      0.23153 134.96978   -3.907 0.000147 ***
## Age:SES            -0.08572      0.02722 144.25897   -3.149 0.001992 **
## ---
## Signif. codes:  0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
##
## Correlation of Fixed Effects:
##      (Intr) n(WBV_,d=4)1 n(WBV_,d=4)2 n(WBV_,d=4)3 n(WBV_,d=4)4 Age
## n(WBV_,d=4)1 -0.639
## n(WBV_,d=4)2 -0.700  0.871
## n(WBV_,d=4)3 -0.684  0.953      0.895
## n(WBV_,d=4)4 -0.718  0.773      0.769      0.797
## Age          -0.953  0.395      0.480      0.445      0.562
## SES          -0.894  0.397      0.499      0.440      0.538      0.921
## SES:nWBV_sc  0.667 -0.684     -0.831     -0.726     -0.850     -0.517
## Age:SES      0.881 -0.368     -0.461     -0.408     -0.500     -0.925
##      SES      SES:WB
## n(WBV_,d=4)1
## n(WBV_,d=4)2
## n(WBV_,d=4)3
## n(WBV_,d=4)4
## Age
## SES
## SES:nWBV_sc -0.605
## Age:SES     -0.995  0.561

```



```
## fit warnings:
## fixed-effect model matrix is rank deficient so dropping 1 column / coefficient
```

```
anova(fit_spline)
```

```
## Type III Analysis of Variance Table with Satterthwaite's method
##
##              Sum Sq Mean Sq NumDF  DenDF F value    Pr(>F)
## ns(nWBV_scaled, df = 4) 105.569  26.392     4 159.90 10.3835 1.679e-07 ***
## Age                    28.043  28.043     1 152.53 11.0330 0.0011202 **
## SES                    29.141  29.141     1 149.27 11.4649 0.0009066 ***
## SES:nWBV_scaled        38.796  38.796     1 134.97 15.2634 0.0001472 ***
## Age:SES                25.204  25.204     1 144.26  9.9162 0.0019916 **
## nWBV_scaled
## ---
## Signif. codes:  0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
```

Purely by AIC, our model which does not account for non-linearity performs the best.

```
anova(fit_spline, fit6)
```

```
## refitting model(s) with ML (instead of REML)
```

```
## Data: dc
## Models:
## fit6: MMSE ~ nWBV_scaled + nWBV_sq_scaled + Age + SES + nWBV_scaled * SES + Age * SES + (1 +
nWBV_scaled | Subject.ID)
## fit_spline: MMSE ~ ns(nWBV_scaled, df = 4) + Age + SES + nWBV_scaled * SES + Age:SES + (1 + n
WBV_scaled | Subject.ID)
##
##          npar    AIC    BIC logLik -2*log(L)  Chisq Df Pr(>Chisq)
## fit6         11 1620.7 1663.3 -799.35   1598.7
## fit_spline   13 1616.6 1666.9 -795.31   1590.6 8.0868  2    0.01754 *
## ---
## Signif. codes:  0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
```

```
AIC(fit_spline, fit6, fit5)
```

```
## Warning in AIC.default(fit_spline, fit6, fit5): models are not all fitted to
## the same number of observations
```

```
##          df      AIC
## fit_spline 13 1624.226
## fit6       11 1639.894
## fit5       11 1621.422
```

Discussion

Model Interpretation

We discuss the interpretation of the three models we constructed in the Results section. We will refer to the model with only linear terms (fit5) as Model 1. We will refer to the model with squared nWBV (fit6) as Model 2. We will refer to the model with splines (fit_spline) as Model 3.

Beginning with Model 1:

```
summary(fit5)
```

```

## Linear mixed-effects model fit by REML
##   Data: dc
##       AIC      BIC    logLik
## 1621.422 1663.797 -799.7112
##
## Random effects:
## Formula: ~1 + nWBV | Subject.ID
## Structure: General positive-definite, Log-Cholesky parametrization
##           StdDev   Corr
## (Intercept) 32.020859 (Intr)
## nWBV        40.961855 -1
## Residual    1.844624
##
## Correlation Structure: AR(1)
## Formula: ~1 | Subject.ID
## Parameter estimate(s):
##      Phi
## 0.312536
## Fixed effects: MMSE ~ nWBV + Age + SES + nWBV * SES + Age * SES
##              Value Std.Error DF   t-value p-value
## (Intercept) -54.94656 16.877249 208 -3.255658 0.0013
## nWBV         85.06516 16.963782 208  5.014516 0.0000
## Age          0.24594  0.079577 208  3.090611 0.0023
## SES          21.81791  6.125116 140  3.562041 0.0005
## nWBV:SES     -21.04513  6.232539 208 -3.376654 0.0009
## Age:SES      -0.07695  0.028539 208 -2.696233 0.0076
## Correlation:
##      (Intr) nWBV   Age    SES   nWBV:S
## nWBV      -0.958
## Age       -0.785  0.575
## SES       -0.919  0.884  0.716
## nWBV:SES   0.873 -0.916 -0.518 -0.958
## Age:SES    0.725 -0.531 -0.926 -0.775  0.563
##
## Standardized Within-Group Residuals:
##      Min      Q1      Med      Q3      Max
## -3.9307210 -0.2692234  0.1472228  0.4643059  2.8280703
##
## Number of Observations: 354
## Number of Groups: 142

```

In the random effects, the high standard deviation for nWBV is indicative of variance in baseline nWBV by subject. The correlation of -1 suggests that the slope and intercept are strongly negatively correlated, but as discussed earlier is indicative of numerical issues, so we will not make an attempt to interpret it here.

In the fixed effects, the coefficient for nWBV (~85.07) suggests that higher nWBV is associated with higher MMSE, which is to be expected. The coefficient for Age (~0.25) suggests that higher age is associated with higher MMSE. This is something to be cautious of, as one would expect cognitive ability to decrease with age; this could also allude to a limitation in the data. Though there is an interaction term involving Age; the coefficient for Age:SES (~-0.08) suggests that the effect of age on MMSE decreases as socioeconomic status increases, which is sensible, since higher socioeconomic status allows for better healthcare access, which one would expect would

slow down the affects of age. The coefficient for SES (~21.82) suggests that higher SES is associated with higher MMSE, which is again sensible for reasons of better healthcare access. Lastly, the coefficient for nWBV:SES (~-21.05) suggests that the positive effect of nWBV decreases as socioeconomic status increases. Note that all coefficients are significant with p-value < 0.01.

Now we consider Model 2:

```
summary(fit6)
```

```
## Linear mixed model fit by REML. t-tests use Satterthwaite's method [
## lmerModLmerTest]
## Formula: MMSE ~ nWBV_scaled + nWBV_sq_scaled + Age + SES + nWBV_scaled *
##      SES + Age * SES + (1 + nWBV_scaled | Subject.ID)
##      Data: dc
##
## REML criterion at convergence: 1617.9
##
## Scaled residuals:
##      Min       1Q   Median       3Q      Max
## -4.1474 -0.2784  0.1098  0.4405  3.2477
##
## Random effects:
##      Groups      Name      Variance Std.Dev. Corr
##      Subject.ID (Intercept) 5.347     2.312
##                nWBV_scaled 2.600     1.612   -0.96
##      Residual              2.660     1.631
## Number of obs: 354, groups: Subject.ID, 142
##
## Fixed effects:
##              Estimate Std. Error      df t value Pr(>|t|)
## (Intercept)    7.78698    5.88637 154.69655   1.323 0.187826
## nWBV_scaled     3.59539    0.63605 139.35550   5.653 8.6e-08 ***
## nWBV_sq_scaled  -0.34260    0.14499  89.00378  -2.363 0.020309 *
## Age             0.24043    0.07588 147.81877   3.169 0.001861 **
## SES             6.92411    2.09132 147.38808   3.311 0.001170 **
## nWBV_scaled:SES -0.87827    0.22917 136.90383  -3.832 0.000193 ***
## Age:SES        -0.08314    0.02711 142.29317  -3.067 0.002587 **
## ---
## Signif. codes:  0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
##
## Correlation of Fixed Effects:
##              (Intr) nWBV_s nWBV__ Age      SES      nWBV_:
## nWBV_scaled  -0.598
## nWBV_sq_scl  -0.110 -0.176
## Age          -0.995  0.556  0.097
## SES          -0.924  0.561  0.050  0.923
## nWBV_sc:SES   0.559 -0.907  0.026 -0.521 -0.609
## Age:SES       0.919 -0.519 -0.048 -0.926 -0.995  0.565
```

```
anova(fit6)
```

```
## Type III Analysis of Variance Table with Satterthwaite's method
##              Sum Sq Mean Sq NumDF   DenDF F value    Pr(>F)
## nWBV_scaled    84.990   84.990     1 139.356 31.9532 8.595e-08 ***
## nWBV_sq_scaled  14.851   14.851     1  89.004  5.5834 0.0203093 *
## Age            26.706   26.706     1 147.819 10.0406 0.0018612 **
## SES            29.157   29.157     1 147.388 10.9619 0.0011696 **
## nWBV_scaled:SES 39.067   39.067     1 136.904 14.6876 0.0001926 ***
## Age:SES        25.023   25.023     1 142.293  9.4076 0.0025871 **
## ---
## Signif. codes:  0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
```

In the random effects, the highly negative correlation between slope and intercept (-0.96) suggests that subjects with higher baseline MMSE experience greater decline in MMSE with loss of brain volume.

In the fixed effects, the coefficient for nWBV (~3.60) suggests that higher nWBV is associated with higher MMSE. However, the coefficient for squared nWBV (~-0.34) dampens this effect. In particular, since nWBV ranges from 0 to 1, squaring nWBV will reduce it, so this dampening effect will be lower for higher nWBV and higher for lower nWBV. Similar to Model 1, the coefficient for Age (~0.24) suggests a positive effect of age on MMSE, and the coefficient for SES (~6.92) suggests a positive effect of SES on MMSE. The interpretation of the interaction coefficients for nWBV:SES (~-0.88) and Age:SES (~-.008) are both the same as the interpretations in Model 1; as SES increases, the effect of nWBV and effect of age decreases. Again, all coefficients are significant with p-value ~ 0.02 for squared nWBV and p-value < 0.01 for all other terms.

Lastly we consider Model 3:

```
summary(fit_spline)
```

```
## Linear mixed model fit by REML. t-tests use Satterthwaite's method [
## lmerModLmerTest]
## Formula: MMSE ~ ns(nWBV_scaled, df = 4) + Age + SES + nWBV_scaled * SES +
##      Age:SES + (1 + nWBV_scaled | Subject.ID)
##      Data: dc
##
## REML criterion at convergence: 1598.2
##
## Scaled residuals:
##      Min      1Q  Median      3Q      Max
## -3.8821 -0.3099  0.1315  0.4339  3.1968
##
## Random effects:
##      Groups      Name      Variance Std.Dev. Corr
##      Subject.ID (Intercept) 5.511      2.347
##                  nWBV_scaled 2.717      1.648   -0.95
##      Residual                2.542      1.594
## Number of obs: 354, groups: Subject.ID, 142
##
## Fixed effects:
##
##              Estimate Std. Error      df t value Pr(>|t|)
## (Intercept)      -6.06696      7.01980 169.91995  -0.864 0.388661
## ns(nWBV_scaled, df = 4)1 12.59680      1.98711 238.51073   6.339 1.14e-09 ***
## ns(nWBV_scaled, df = 4)2 12.74060      2.12443 149.17160   5.997 1.45e-08 ***
## ns(nWBV_scaled, df = 4)3 28.71832      4.56269 241.57876   6.294 1.44e-09 ***
## ns(nWBV_scaled, df = 4)4 16.68518      3.08572 124.96132   5.407 3.13e-07 ***
## Age                0.25381      0.07641 152.53105   3.322 0.001120 **
## SES                7.11217      2.10048 149.27436   3.386 0.000907 ***
## SES:nWBV_scaled      -0.90454      0.23153 134.96978  -3.907 0.000147 ***
## Age:SES             -0.08572      0.02722 144.25897  -3.149 0.001992 **
## ---
## Signif. codes:  0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
##
## Correlation of Fixed Effects:
##      (Intr) n(WBV_,d=4)1 n(WBV_,d=4)2 n(WBV_,d=4)3 n(WBV_,d=4)4 Age
## n(WBV_,d=4)1 -0.639
## n(WBV_,d=4)2 -0.700  0.871
## n(WBV_,d=4)3 -0.684  0.953      0.895
## n(WBV_,d=4)4 -0.718  0.773      0.769      0.797
## Age          -0.953  0.395      0.480      0.445      0.562
## SES          -0.894  0.397      0.499      0.440      0.538      0.921
## SES:nWBV_sc  0.667 -0.684     -0.831     -0.726     -0.850     -0.517
## Age:SES      0.881 -0.368     -0.461     -0.408     -0.500     -0.925
##      SES      SES:WB
## n(WBV_,d=4)1
## n(WBV_,d=4)2
## n(WBV_,d=4)3
## n(WBV_,d=4)4
## Age
## SES
## SES:nWBV_sc -0.605
## Age:SES     -0.995  0.561
```

```
## fit warnings:
## fixed-effect model matrix is rank deficient so dropping 1 column / coefficient
```

```
anova(fit_spline)
```

```
## Type III Analysis of Variance Table with Satterthwaite's method
##
```

	Sum Sq	Mean Sq	NumDF	DenDF	F value	Pr(>F)
ns(nWBV_scaled, df = 4)	105.569	26.392	4	159.90	10.3835	1.679e-07 ***
Age	28.043	28.043	1	152.53	11.0330	0.0011202 **
SES	29.141	29.141	1	149.27	11.4649	0.0009066 ***
SES:nWBV_scaled	38.796	38.796	1	134.97	15.2634	0.0001472 ***
Age:SES	25.204	25.204	1	144.26	9.9162	0.0019916 **
nWBV_scaled						

```
## Signif. codes:  0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
```

The interpretation of random effects is similar to that of Model 2, with Model 3 having correlation (-0.95) between slope and intercept.

In the fixed effects, the coefficients for Age, SES, SES:nWBV and Age:SES are similar to those in Model 2, and can be interpreted in the same way. The high, positive coefficients for the nWBV spline terms suggest that the relationship between MMSE and nWBV is both positive and highly non-linear. All coefficients are statistically significant with p-values < 0.01.

In summary, our models all agree that higher brain volume, higher socioeconomic status, and higher age predict higher MMSE, with age having a smaller effect than brain volume and socioeconomic status, and the effects of brain volume and age being dampened through interactions with socioeconomic status.

Comparing Mean Trajectory Slopes

We compute the mean of the slopes of the trajectory lines stratified by CDR from our MMSE trajectory plot constructed in the Dataset section:

```
dc_copy <- dc

subject_slopes <- dc_copy %>%
  group_by(Subject.ID, CDR) %>%
  filter(n() > 1) %>% # need at least two points per subject
  summarise(
    slope = {
      fit <- lm(MMSE ~ Age, data = cur_data())
      coef(fit)["Age"]
    },
    .groups = "drop"
  )
```

```
## Warning: There was 1 warning in `summarise()`.
## i In argument: `slope = { ... }`.
## i In group 1: `Subject.ID = "OAS2_0001"` `CDR = 0`.
## Caused by warning:
## ! `cur_data()` was deprecated in dplyr 1.1.0.
## i Please use `pick()` instead.
```

```
avg_slopes_by_CDR_old <- subject_slopes %>%
  group_by(CDR) %>%
  summarise(
    avg_slope = mean(slope, na.rm = TRUE),
    n = n()
  )

print(avg_slopes_by_CDR_old)
```

```
## # A tibble: 3 × 3
##   CDR avg_slope    n
##   <dbl>   <dbl> <int>
## 1  0     -0.0306   76
## 2  0.5   -0.464    38
## 3  1     -0.826     9
```

We see that as CDR increases, expected decline rate of MMSE increases as well. This matches the trend for the mean nWBV trajectory slopes computed in [1], further verifying the association between MMSE and nWBV.

We can similarly compute the mean of the slopes of the trajectory lines generated by Model 1:


```
dc$fit_vals5 <- fitted(fit5)

dc_slope <- dc %>%
  select(Subject.ID, Age, fit_vals5, CDR) %>%
  group_by(Subject.ID, CDR) %>%
  arrange(Age, .by_group = TRUE) %>%
  summarise(
    slope = if (n() >= 2) coef(lm(fit_vals5 ~ Age))[2] else NA_real_,
    .groups = "drop"
  )

avg_slopes_by_CDR_5 <- dc_slope %>%
  group_by(CDR) %>%
  summarise(
    avg_slope = mean(slope, na.rm = TRUE),
    n = n()
  )

print(avg_slopes_by_CDR_5)
```

```
## # A tibble: 4 × 3
##   CDR avg_slope    n
##   <dbl>   <dbl> <int>
## 1  0     0.00224   86
## 2  0.5  -0.160    61
## 3  1    -0.946    22
## 4  2    NaN      3
```

And for those generated by Model 2:

```
dc$fit_vals6 <- fitted(fit6)

dc_slope <- dc %>%
  select(Subject.ID, Age, fit_vals6, CDR) %>%
  group_by(Subject.ID, CDR) %>%
  arrange(Age, .by_group = TRUE) %>%
  summarise(
    slope = if (n() >= 2) coef(lm(fit_vals6 ~ Age))[2] else NA_real_,
    .groups = "drop"
  )

avg_slopes_by_CDR_6 <- dc_slope %>%
  group_by(CDR) %>%
  summarise(
    avg_slope = mean(slope, na.rm = TRUE),
    n = n()
  )

print(avg_slopes_by_CDR_6)
```

```
## # A tibble: 4 × 3
##   CDR avg_slope    n
##   <dbl>   <dbl> <int>
## 1  0     -0.0135   86
## 2  0.5   -0.227    61
## 3  1     -1.20    22
## 4  2      NaN     3
```

And for those generated by Model 3:

```
dc$fit_spline <- fitted(fit_spline)

dc_slope <- dc %>%
  select(Subject.ID, Age, fit_spline, CDR) %>%
  group_by(Subject.ID, CDR) %>%
  arrange(Age, .by_group = TRUE) %>%
  summarise(
    slope = if (n() >= 2) coef(lm(fit_spline ~ Age))[2] else NA_real_,
    .groups = "drop"
  )

avg_slopes_by_CDR <- dc_slope %>%
  group_by(CDR) %>%
  summarise(
    avg_slope = mean(slope, na.rm = TRUE),
    n = n()
  )

print(avg_slopes_by_CDR)
```

```
## # A tibble: 4 × 3
##   CDR avg_slope    n
##   <dbl>   <dbl> <int>
## 1  0     -0.0198   86
## 2  0.5   -0.302    61
## 3  1     -1.29    22
## 4  2      NaN     3
```

Comparing the mean slopes per CDR group to those of the actual trajectories for each model, we see that Model 1 has the closest slope for CDR = 1 and the furthest for CDR = 0 and CDR = 0.5; in particular, for CDR = 0 it has a positive mean slope, making it unsuitable for predictions with this CDR group but quite suitable for CDR = 1. Model 3 appears to be the best model for CDR = 0 and CDR = 0.5, while Model 2 is slightly better than Model 3 for CDR = 1 but worse than Model 1. Note that these assessments are quite limited due to the small dataset size; in particular, many subject slopes were not able to be computed for the initial trajectories, so the CDR = 1 group is especially small (9 subjects).

Issues With Residuals

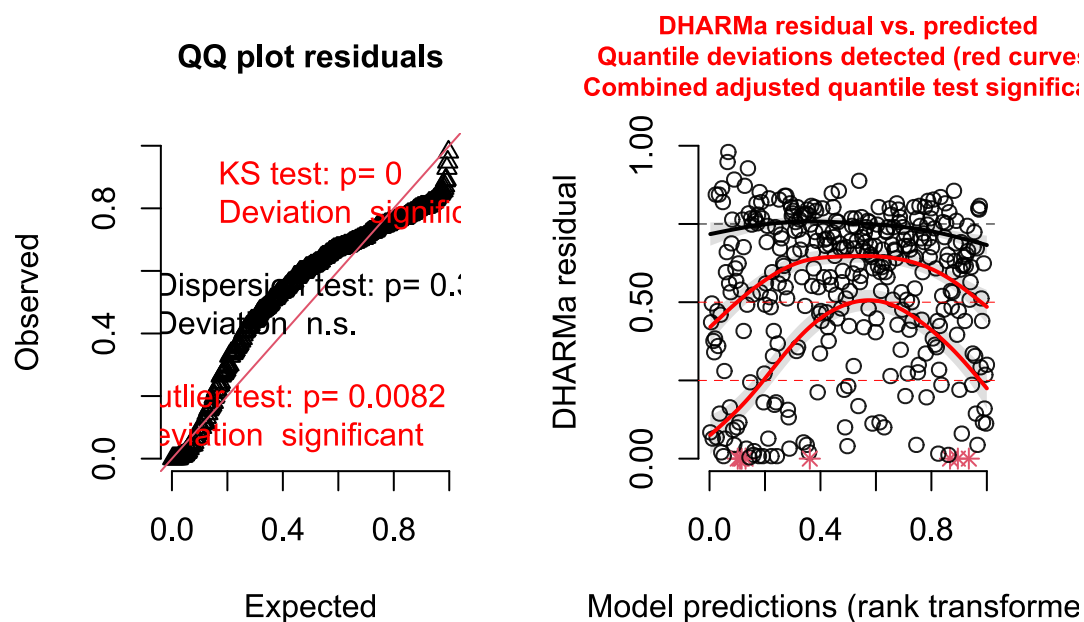
To test residuals, we reconstruct our models as generalized linear mixed models using the `glmmTMB` package and then use the `DHARMA` package to plot residuals.

For Model 1:

```
fit5_tmb <- glmmTMB(
  MMSE ~ nWBV + Age + SES + nWBV:SES + Age:SES + (1 + nWBV | Subject.ID),
  data = dc,
  REML = TRUE
)

res_5 <- simulateResiduals(fittedModel = fit5_tmb)
plot(res_5)
```

DHARMA residual

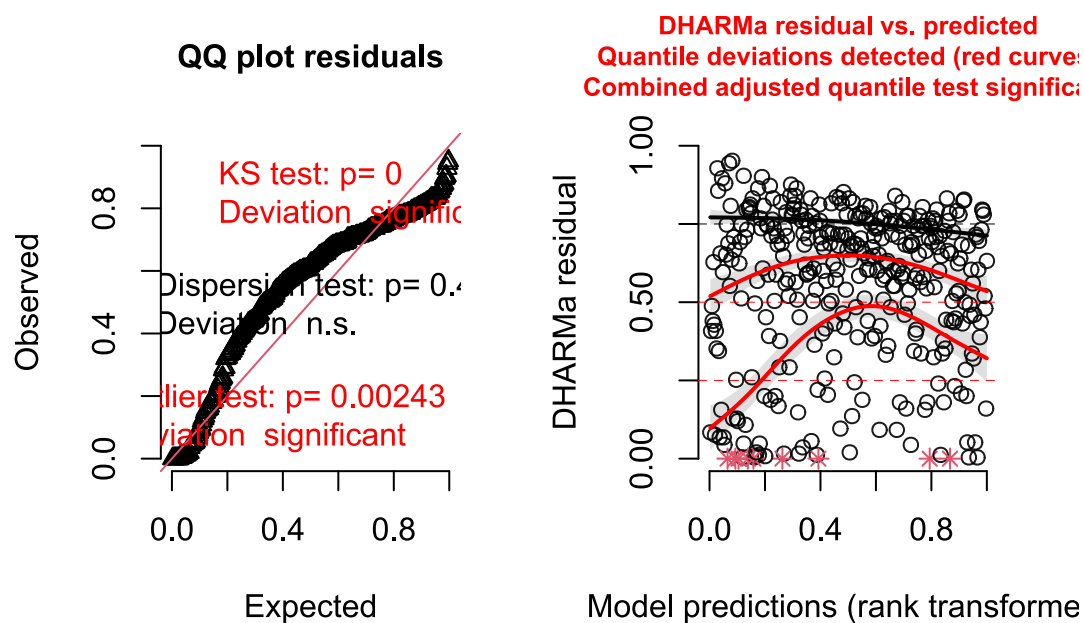


For Model 2:

```
fit6_tmb <- glmmTMB(
  MMSE ~ nWBV_scaled + nWBV_sq_scaled + Age + SES + nWBV_scaled*SES + Age*SES + (1 + nWBV | Subject.ID),
  data = dc,
  REML = TRUE
)

res_6 <- simulateResiduals(fittedModel = fit6_tmb)
plot(res_6)
```

DHARMA residual



For Model 3:

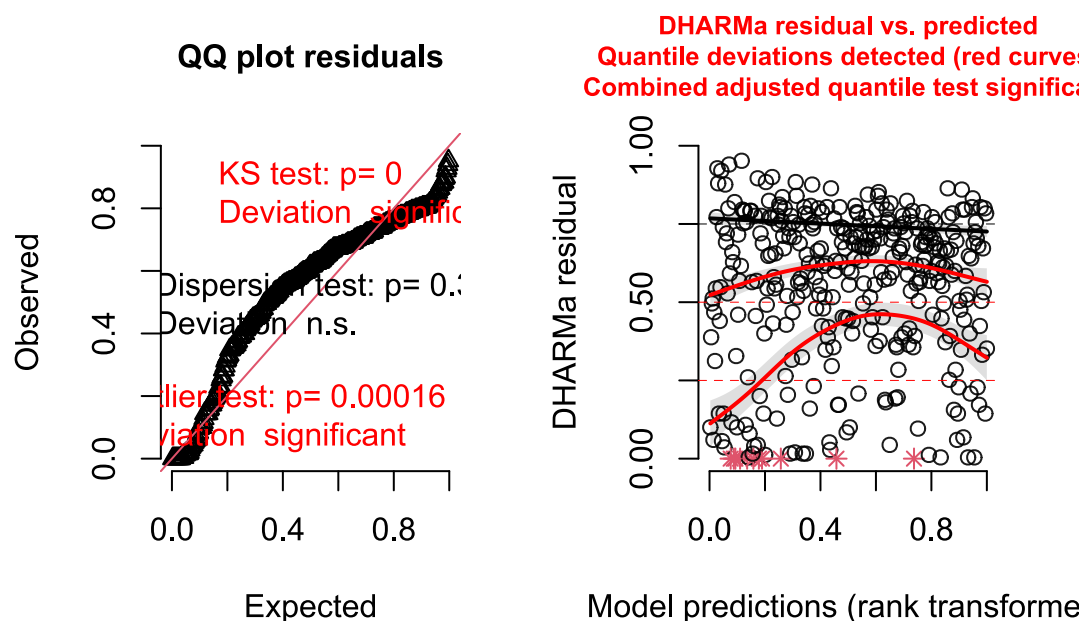
```
dc$ns_nWBV_scaled <- ns(dc$nWBV_scaled, df = 4)
ns_basis <- ns(dc$nWBV_scaled, df = 4)
ns_df <- as.data.frame(ns_basis)
colnames(ns_df) <- paste0("ns_nWBV_", 1:ncol(ns_df))

dc <- bind_cols(dc, ns_df)

fit_spline_tmb <- glmmTMB(
  MMSE ~ ns_nWBV_1 + ns_nWBV_2 + ns_nWBV_3 + Age + SES + nWBV * SES + Age * SES
  + (1 + nWBV | Subject.ID),
  data = dc,
  REML = TRUE
)

res_spline <- simulateResiduals(fit_spline_tmb)
plot(res_spline)
```

DHARMa residual



Note that for Model 3 we've excluded `ns_nWBV_4` as it is collinear with the other basis functions, which prevents DHARMa from simulating the residuals. This was unnecessary when we initially constructed the model in the results section, as LMER automatically dropped the collinear column.

In all three models, we see that the curves in the residual plots are not all flat, which is indicative of heteroskedasticity that the models fail to account for. Each model fails the KS test with $p\text{-value} < 0.01$, indicating that the residuals are not uniformly distributed, and each model fails the outlier test with $p\text{-value} < 0.01$, indicating that the model's performance is significantly affected by the presence of outliers. We see that even introducing non-linearity, as we have done in Model 2 and Model 3, does little to improve these issues. These are the limitations of LMEs that arise when working with non-normal data.

Possible Solutions for Residual Issues

In this particular case, the data is heavily left-skewed, with most MMSE scores being at or close to 30. In particular, this means that this data has a lot of individuals who do not experience cognitive decline. We can flip the MMSE scores so that most scores are at or close to 0, resulting in zero-inflation. Then we can use glmmTMB to construct a generalized linear mixed model that accounts for zero-inflation, i.e it accounts for those individuals who do not experience cognitive decline.

For example, we reconstruct Model 1 accounting for zero-inflation (note that we omit the random slope to resolve convergence issues):

```
dc_c <- dc
dc_c$MMSE <- 30 - dc_c$MMSE
dc_c$nWBV_scaled <- scale(dc_c$nWBV)[,1]
dc_c$Age_scaled <- scale(dc_c$Age)[,1]
dc_c$SES_scaled <- scale(dc_c$SES)[,1]

fit5_tmb <- glmmTMB(
  MMSE ~ nWBV_scaled + Age_scaled + SES_scaled +
    nWBV_scaled * SES_scaled + Age_scaled * SES_scaled +
    (1 | Subject.ID),
  data = dc_c,
  ziformula = ~ Age_scaled,
  family = poisson,
  REML = TRUE
)

AIC(fit5, fit5_tmb)
```

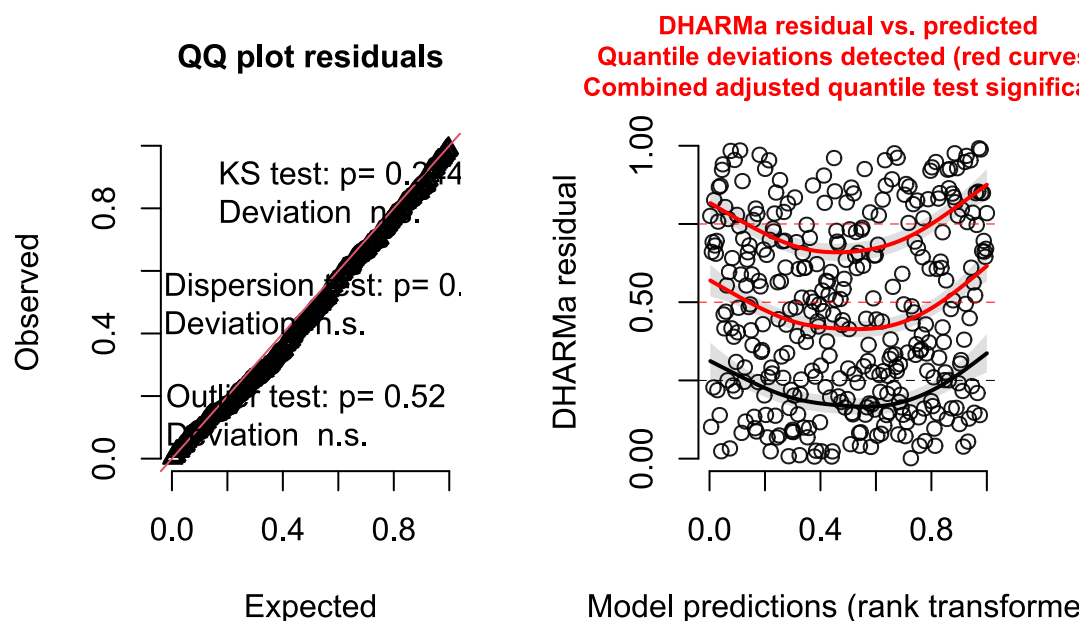
```
## Warning in AIC.default(fit5, fit5_tmb): models are not all fitted to the same
## number of observations
```

```
##          df      AIC
## fit5      11 1621.422
## fit5_tmb   9 1305.129
```

We see that this model performs significantly better than Model 1, as it has a significantly lower AIC. Moreover, we can plot the residuals:

```
res1 <- simulateResiduals(fit5_tmb)
plot(res1)
```

DHARMA residual



And we see that the model now passes the KS and outlier tests, suggesting that the residuals are uniformly distributed and that the model better handles outliers. The curves in the residual plots are still not perfectly flat indicating that the model still struggles with heteroskedasticity, though it is certainly an improvement to the previous plot.

A similar approach shows some promise in improving Models 2 and 3. It was intended that those improvements be showcased in this report, though the models that worked in the original R script are experiencing convergence issues in markdown. Due to time constraints, the convergence issues are unable to be resolved for the purpose of this report, and so those models are omitted.

On the use of AI

I would like to give credit to both ChatGPT and Claude for their assistance in completing this project. I used both LLMs for the following components of the project:

- Plotting and computing mean trajectory slopes
- Double-checking p-value/coefficient interpretation
- Model Diagnostics
- R Troubleshooting

I found both models to be quite helpful for plotting. For example, when recreating the nWBV trajectories that I originally found in [1], I gave ChatGPT a screenshot from the paper and asked it to recreate this plot, which it was able to do successfully using xyplot.

Both models were helpful in validating my impressions for p-values and interpreting model coefficients, though they were not perfect (for example, ChatGPT once interpreted a p-value to mean that the more complex model is a better fit when really it meant the simpler model is a better fit. Another example is Claude mixed up left and right skewed distributions when I showed it an image of my MMSE histogram.)

I found both models to be unhelpful with model diagnostics. For example, I asked Claude for suggestions on how to deal with the random effects being numerically unstable (correlation = -1), and it suggested response transformations for MMSE (log, square root, Box-Cox). When these transformations made no difference and I asked it for alternatives, it proceeded to give the same suggestions in a loop.

I found ChatGPT to be unhelpful for R troubleshooting, though Claude was quite helpful for R troubleshooting. For example, reproducing my R script involving non-linear terms in my LMEs was problematic in R markdown, as I started having singularity issues that did not come up when I ran the same lines of code in my original script. Claude suggested scaling nWBV and using LMER as an alternative to LME, and this resolved the issue.

References

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