ACE Models with the NLSY

William Howard Beasley (Howard Live Oak LLC, Norman)
Joseph Lee Rodgers (Vanderbilt University, Nashville)
David Bard (University of Oklahoma Health Sciences Center, OKC)
Kelly Meredith (University of Oklahoma, Norman)
Michael D. Hunter (University of Oklahoma, Norman)

November 15, 2013

Abstract

We describe how to use the NlsyLinks package to examine various biometric models, using the NLSY79.

Contents

1	Terminology	1
2	Example: DF analysis with a Simple Outcome for Gen2 Subjects, Using a Package Variable	3
3	Example: DF analysis with a univariate outcome from a Gen2 Extract	5
4	Example: Multiple Group SEM of a Simple Outcome for Gen2 Subjects	7
5	Example: Multiple Group SEM of a Simple Outcome for Gen1 Subjects	9
6	Example: Midstream data manipulation with SAS	11
A	Appendix: Receiving Help for the NlsyLinks Package	11
В	Appendix: Creating and Saving R Scripts	11
\mathbf{C}	Appendix: Installing and Loading the NlsyLinks Package	11
D	Appendix: References	12
\mathbf{E}	Notes	12
\mathbf{F}	Version Information	12

1 Terminology

This package considers both Gen1 and Gen2 subjects. **Gen1** refers to subjects in the original NLSY79 sample (http://www.bls.gov/nls/nlsy79.htm). **Gen2** subjects are the biological offspring of the Gen1 females -i.e., those in the NLSY79 Children and Young Adults sample (http://www.bls.gov/nls/nlsy79ch.htm). The NLSY97 is a third dataset that can be used for behavior genetic research (http://www.bls.gov/nls/nlsy97.htm), although this vignette focuses on the two generations in the NLSY79.

Standard terminology is to refer Gen2 subjects as 'children' when they are younger than age 15 (NSLY79-C), and as 'young adults' when they are 15 and older (NLSY79-YA); though they are the same respondents, different funding mechanisms and different survey items necessitate the distinction. This cohort is sometimes abbreviated as 'NLSY79-C', 'NLSY79C', 'NLSY-C' or 'NLSYC'.

The SubjectTag variable uniquely identify NLSY79 subjects when a dataset contains both generations. For Gen2 subjects, the SubjectTag is identical to their CID (i.e., C00001.00 -the ID assigned in the NLSY79-Children files). However for Gen1 subjects, the SubjectTag is their CaseID (i.e., R00001.00), with "00" appended. This manipulation is necessary to identify subjects uniquely in inter-generational datasets. A Gen1 subject with an ID of 43 becomes 4300. The SubjectTags of her four children remain 4301, 4302, 4303, and 4304.

The **expected coefficient of relatedness** of a pair of subjects is typically represented by the variable R. Examples are: Monozygotic twins have R=1; dizygotic twins have R=0.5; full siblings (*i.e.*, those who share both biological parents) have R=0.5; half-siblings (*i.e.*, those who share exactly one biological parent) have R=0.25; adopted siblings have R=0.0. Other possibilities exist too. The font (and hopefully their context) should distinguish the variable R from the software R.

A subject's ExtendedID indicates their extended family. Two subjects will be in the same extended family if either: [1] they are Gen1 housemates, [2] they are Gen2 siblings, [3] they are Gen2 cousins (*i.e.*, they have mothers who are Gen1 sisters in the NLSY79), [4] they are mother and child (in Gen1 and Gen2, respectively), or [5] they are (aunt—uncle) and (niece—nephew) (in Gen1 and Gen2, respectively).

An **outcome variable** is directly relevant to the applied researcher; these might represent constructs like height, IQ, and income. A **plumbing variable** is necessary to manage BG datasets; examples are R, a subject's ID, and the date of a subject's last survey.

An ACE model is the basic biometrical model used by Behavior Genetic researchers, where the genetic and environmental effects are assumed to be additive. The three primary variance components are (1) the proportion of variability due to a shared genetic influence (typically represented as a^2 , or sometimes h^2), (2) the proportion of variability due to shared common environmental influence (typically e^2), and (3) the proportion of variability due to unexplained/residual/error influence (typically e^2).

The variables are scaled so that they account for all observed variability in the outcome variable; specifically: $a^2 + c^2 + e^2 = 1$. Using appropriate designs that can logically distinguish these different components (under carefully specified assumptions), the basic biometrical modeling strategy is to estimate the magnitude of a^2 , c^2 , and e^2 within the context of a particular model. For gentle introductions to Behavior Genetic research, we recommend Plomin (1990) and Carey (2003). For more in-depth ACE model-fitting strategies, we recommend Neale & Cardon (1992). //This paragraph may get moved to the yet-to-be-written introduction that precedes the Terminology section.

The NLS Investigator (http://www.nlsinfo.org/investigator/) is the best way to obtain the NLSY79 and NLSY97 datasets. See our vignette dedicated to the NLS Investigator by typing vignette("NlsInvestigator") or by visiting http://cran.r-project.org/web/packages/NlsyLinks/.

Before starting the real examples, first verify that the NlsyLinks package is installed correctly. If not, refer to Appendix C.

```
any(.packages(all.available = TRUE) == "NlsyLinks") #Should evaluate to TRUE.
## [1] TRUE
require(NlsyLinks) #Load the package into the current session.
## Loading required package: NlsyLinks
## Loading required package: Matrix
```

The package's documentation manual can be opened by typing ?NlsyLinks in R or RStudio.

2 Example: DF analysis with a Simple Outcome for Gen2 Subjects, Using a Package Variable

The vignette's first example uses a simple statistical model and all available Gen2 subjects. The CreatePairLinksDoubleEntered function will create a data frame where each represents one pair of siblings, respective of order (i.e., there is a row for Subjects 201 and 202, and a second row for Subjects 202 and 201). This function examines the subjects' IDs and determines who is related to whom (and by how much). By default, each row it produces has at least six values/columns: (i) ID for the older member of the kinship pair: Subject1Tag, (ii) ID for the younger member: Subject2Tag, (iii) ID for their extended family: ExtendedID, (iv) their estimated coefficient of genetic relatedness: R, (v and beyond) outcome values for the older member; (vi and beyond) outcome values for the younger member.

A DeFries-Fulker (**DF**) Analysis uses linear regression to estimate the a^2 , c^2 , and e^2 of a univariate biometric system. The interpretations of the DF analysis can be found in Rodgers & Kohler (2005) and Rodgers, Rowe, & Li (1999). This vignette example uses the newest variation, which estimates two parameters; the corresponding function is called **DeFriesFulkerMethod3**. The steps are:

- 1. Use the NLS Investigator to select and download a Gen2 dataset.
- 2. Open R and create a new script (see Appendix B) and load the NlsyLinks package. If you haven't done so, first install the NlsyLinks package (see Appendix C).
- 3. Within the R script, load the linking dataset. Then select only Gen2 subjects. The 'Pair' version of the linking dataset is essentially an upper triangle of a symmetric sparse matrix.
- 4. Load and assign the ExtraOutcomes79 dataset.
- 5. Specify the outcome variable name and filter out all subjects who have a negative value in this variable. The NLSY typically uses negative values to indicate different types of missingness (see 'Further Information' below).
- 6. Create a double-entered file by calling the CreatePairLinksDoubleEntered function. At minimum, pass the (i) outcome dataset, the (ii) linking dataset, and the (iii) name(s) of the outcome variable(s). (There are occasions when a single-entered file is more appropriate for a DF analysis. See Rodgers & Kohler, 2005, for additional information.)
- 7. Use DeFriesFulkerMethod3 function (i.e., general linear model) to estimate the coefficients of the DF model.

```
### R Code for Example DF analysis with a simple outcome and Gen2 subjects
#Step 2: Load the package containing the linking routines.
require(NlsyLinks)
#Step 3: Load the LINKING dataset and filter for the Gen2 subjects
dsLinking <- subset(Links79Pair, RelationshipPath=="Gen2Siblings")</pre>
summary(dsLinking) #Notice there are 11,088 records (one for each unique pair).
##
      ExtendedID
                      Subject1Tag
                                        Subject2Tag
                                                                 R
          :
                2
                            :
                                               :
                                                    202
                                                                  :0.250
##
                    Min.
                                 201
                                       Min.
                                                          Min.
##
    1st Qu.: 3155
                    1st Qu.: 315501
                                       1st Qu.: 315503
                                                          1st Qu.:0.250
   Median: 6114
                                       Median : 611404
                                                          Median :0.500
                    Median: 611402
##
##
           : 5933
                    Mean
                            : 593658
                                       Mean
                                               : 593660
                                                          Mean
                                                                  :0.417
##
    3rd Qu.: 8511
                    3rd Qu.: 851101
                                       3rd Qu.: 851103
                                                          3rd Qu.:0.500
##
           :12673
                            :1267301
                                       Max.
                                               :1267302
                                                                  :1.000
                    Max.
                                                          Max.
##
          RelationshipPath
    Gen1Housemates:
    Gen2Siblings
##
                  :11088
    Gen2Cousins
```

```
## ParentChild : 0
## AuntNiece
##
#Step 4: Load the OUTCOMES dataset, and then examine the summary.
dsOutcomes <- ExtraOutcomes79 #'ds' stands for 'Data Set'
summary(dsOutcomes)
##
                                                  AfqtRescaled2006Gaussified
     SubjectTag
                     SubjectID
                                      Generation
  Min. : 100
                   Min. :
                               1
                                   Min. :1.00
                                                  Min. :-3
   1st Qu.: 314025
                   1st Qu.:
                            5998
                                   1st Qu.:1.00
                                                 1st Qu.:-1
                   Median: 12000
## Median : 620050
                                    Median :1.00
                                                 Median: 0
## Mean : 618600 Mean : 289254
                                   Mean :1.48
                                                 Mean : 0
## 3rd Qu.: 914501 3rd Qu.: 577403
                                    3rd Qu.:2.00
                                                  3rd Qu.: 0
## Max. :1268600 Max. :1267501 Max. :2.00
                                                 Max. : 3
                                                  NA's
                                                        :12276
## HeightZGender HeightZGenderAge WeightZGender
                                               WeightZGenderAge
                                                                   Afi
              Min. :-3
## Min. :-3
                              Min. :-3
                                               Min. :-4
                                                             Min. : 2
## 1st Qu.:-1
                 1st Qu.:−1
                                1st Qu.:-1
                                               1st Qu.:-1
                                                              1st Qu.:15
## Median : 0
                Median: 0
                                Median: 0
                                               Median : 0
                                                              Median:17
## Mean : 0
               Mean : 0
                                Mean : 0
                                               Mean : 0
                                                             Mean :17
## 3rd Qu.: 1
                 3rd Qu.: 1
                                3rd Qu.: 0
                                               3rd Qu.: 0
                                                              3rd Qu.:18
## Max. : 3
                 Max. : 3
                                Max. : 8
                                               Max. : 7
                                                              Max.
                                                                     :27
## NA's :5029 NA's :5030
                                NA's :12104
                                              NA's :12105
                                                              NA's
                                                                     :12740
##
      Afm
                MathStandardized
## Min. : 0
                Min. : 65
## 1st Qu.:12
                 1st Qu.: 91
## Median :13
                 Median:100
## Mean :13
                 Mean :100
## 3rd Qu.:14
                  3rd Qu.:110
## Max. :19
                 Max. :135
## NA's :18165 NA's :15048
#Step 5: This step isn't necessary for this example, because Kelly Meredith already
# groomed the values. If the negative values (which represent NLSY missing or
# skip patterns) still exist, then:
dsOutcomes$MathStandardized[dsOutcomes$MathStandardized < 0] <- NA
#Step 6: Create the double entered dataset.
dsDouble <- CreatePairLinksDoubleEntered(</pre>
 outcomeDataset=dsOutcomes,
 linksPairDataset=dsLinking,
 outcomeNames=c('MathStandardized')
summary(dsDouble) #Notice there are 22176=(2*11088) records now (two for each unique pair).
                    Subject2Tag
                                      ExtendedID
   Subject1Tag
## Min. :
             201
                   Min. :
                              201
                                    Min. : 2
                                                 Min. :0.250
## 1st Qu.: 315502
                   1st Qu.: 315502
                                    1st Qu.: 3155
                                                  1st Qu.:0.250
## Median : 611404
                   Median : 611404
                                    Median: 6114
                                                 Median :0.500
## Mean : 593659
                   Mean : 593659
                                    Mean : 5933
                                                  Mean :0.417
## 3rd Qu.: 851102
                   3rd Qu.: 851102
                                    3rd Qu.: 8511
                                                  3rd Qu.:0.500
                   Max. :1267302
## Max. :1267302
                                   Max. :12673
                                                 Max. :1.000
##
        RelationshipPath MathStandardized_1 MathStandardized_2
```

```
Gen1Housemates:
                       0
                           Min.
                                  : 65
                                               Min.
                                                      : 65
##
    Gen2Siblings :22176
                           1st Qu.: 89
                                               1st Qu.: 89
##
    Gen2Cousins
                       0
                           Median: 98
                                               Median: 98
                       0
##
   ParentChild
                                  : 98
                           Mean
                                               Mean
                                                     : 98
##
    AuntNiece
                           3rd Qu.:108
                                               3rd Qu.:108
##
                           Max.
                                   :135
                                               Max.
                                                       :135
##
                           NA's
                                   :3815
                                               NA's
                                                       :3815
#Step 7: Estimate the ACE components with a DF Analysis
ace <- DeFriesFulkerMethod3(</pre>
    dataSet=dsDouble.
    oName_1="MathStandardized_1",
    oName_2="MathStandardized_2")
## [1] "Results of ACE estimation: [show]"
  ASquared CSquared ESquared CaseCount
## 8.480e-01 4.477e-02 1.073e-01 1.678e+04
```

Further Information: If the different reasons of missingness are important, further work is necessary. For instance, some analyses that use item Y19940000 might need to distinguish a response of "Don't Know" (which is coded as -2) from "Missing" (which is coded as -7). For this vignette example, we'll assume it's safe to clump the responses together.

3 Example: DF analysis with a univariate outcome from a Gen2 Extract

The vignette's second example differs from the previous example in two ways. First, the outcome variables are read from a CSV (comma separated values file) that was downloaded from the NLS Investigator. Second, the DF analysis is called through the function AceUnivariate; this function is a wrapper around some simple ACE methods, and will help us smoothly transition to more techniques later in the vignette.

The steps are:

- 1. Use the NLS Investigator to select and download a Gen2 dataset. Select the variables 'length of gestation of child in weeks' (C03280.00), 'weight of child at birth in ounces' (C03286.00), and 'length of child at birth' (C03288.00), and then download the *.zip file to your local computer.
- 2. Open R and create a new script (see Appendix B) and load the NlsyLinks package.
- 3. Within the R script, load the linking dataset. Then select only Gen2 subjects.
- 4. Read the CSV into R as a data.frame using ReadCsvNlsy79Gen2.
- 5. Verify the desired outcome column exists, and rename it something meaningful to your project. It is important that the data.frame is reassigned (i.e., ds <- RenameNlsyColumn(...)). In this example, we rename column C0328800 to BirthWeightInOunces.
- 6. Filter out all subjects who have a negative BirthWeightInOunces value. See the 'Further Information' note in the previous example.
- 7. Create a double-entered file by calling the CreatePairLinksDoubleEntered function. At minimum, pass the (i) outcome dataset, the (ii) linking dataset, and the (iii) name(s) of the outcome variable(s).
- 8. Call the AceUnivariate function to estimate the coefficients.

```
### R Code for Example of a DF analysis with a simple outcome and Gen2 subjects #Step 2: Load the package containing the linking routines.
```

```
require(NlsyLinks)
#Step 3: Load the linking dataset and filter for the Gen2 subjects
dsLinking <- subset(Links79Pair, RelationshipPath=="Gen2Siblings")
#Step 4: Load the outcomes dataset from the hard drive and then examine the summary.
# Your path might be: filePathOutcomes <- 'C:/BGResearch/NlsExtracts/Gen2Birth.csv'
filePathOutcomes <- file.path(path.package("NlsyLinks"), "extdata", "Gen2Birth.csv")</pre>
dsOutcomes <- ReadCsvNlsy79Gen2(filePathOutcomes)</pre>
summary(dsOutcomes)
##
     SubjectTag
                      SubjectID
                                      ExtendedID
                                                      Generation SubjectTagOfMother
                               201 Min. : 2 Min. :2
## Min. : 201
                   Min. :
                                                               Min. :
## 1st Qu.: 310302
                   1st Qu.: 310302
                                    1st Qu.: 3101
                                                    1st Qu.:2
                                                                1st Qu.: 310300
## Median : 604607 Median : 604607 Median : 6045
                                                   Median :2
                                                              Median: 604600
## Mean : 601313 Mean : 601313
                                    Mean : 6007
                                                    Mean :2
                                                              Mean : 601311
## 3rd Qu.: 876202 3rd Qu.: 876202
                                     3rd Qu.: 8757
                                                    3rd Qu.:2
                                                                3rd Qu.: 876200
## Max. :1267501 Max. :1267501 Max. :12675
                                                    Max. :2
                                                              Max. :1267500
##
                                     NA's :2
      C0005300
                   C0005400
                                   C0005700
                                                 C0328000
                                                               C0328600
##
                                                           Min. : -7
## Min. :1.00 Min. :-3.00 Min. :-3 Min. :-7.0
## 1st Qu.:2.00 1st Qu.: 1.00 1st Qu.:1981 1st Qu.:37.0
                                                            1st Qu.: 99
## Median: 3.00 Median: 1.00 Median: 1985 Median: 39.0
                                                           Median:115
                                                            Mean :104
## Mean :2.34
                Mean : 1.49 Mean :1986 Mean :33.5
## 3rd Qu.:3.00
                 3rd Qu.: 2.00 3rd Qu.:1990 3rd Qu.:39.0
                                                            3rd Qu.:128
## Max. :3.00
                Max. : 2.00 Max. :2008 Max. :51.0 Max. :768
##
      C0328800
##
## Min. :-7.0
## 1st Qu.:18.0
## Median :20.0
## Mean :16.5
## 3rd Qu.:21.0
## Max. :48.0
##
#Step 5: Verify and rename an existing column.
VerifyColumnExists(dsOutcomes, "CO328600") #Should return '10' in this example.
## [1] 10
dsOutcomes <- RenameNlsyColumn(dsOutcomes, "CO328600", "BirthWeightInOunces")
#Step 6: For this item, a negative value indicates the parent refused, didn't know,
# invalidly skipped, or was missing for some other reason.
# For our present purposes, we'll treat these responses equivalently.
  Then clip/Winsorized/truncate the weight to something reasonable.
dsOutcomes$BirthWeightInOunces[dsOutcomes$BirthWeightInOunces < 0] <- NA
dsOutcomes$BirthWeightInOunces <- pmin(dsOutcomes$BirthWeightInOunces, 200)
#Step 7: Create the double entered dataset.
dsDouble <- CreatePairLinksDoubleEntered(</pre>
 outcomeDataset=dsOutcomes,
 linksPairDataset=dsLinking,
 outcomeNames=c('BirthWeightInOunces')
```

```
#Step 8: Estimate the ACE components with a DF Analysis
ace <- AceUnivariate(
   method="DeFriesFulkerMethod3",
   dataSet=dsDouble,
   oName_1="BirthWeightInOunces_1",
   oName_2="BirthWeightInOunces_2"
)
ace
## [1] "Results of ACE estimation: [show]"
## ASquared CSquared ESquared CaseCount
## 5.042e-01 1.777e-01 3.182e-01 1.744e+04</pre>
```

For another example of incorporating CSVs downloaded from the NLS Investigator, please see the "Race and Gender Variables" entry in the FAQ.

4 Example: Multiple Group SEM of a Simple Outcome for Gen2 Subjects

The example differs from the first one by the statistical mechanism used to estimate the components. The first example uses multiple regression to estimate the influence of the shared genetic and environmental factors, while this example uses structural equation modeling (SEM).

The CreatePairLinksSingleEntered function will create a data.frame where each row represents one unique pair of siblings, *irrespective of order*. Other than producing half the number of rows, this function is identical to CreatePairLinksDoubleEntered.

The steps are:

(Steps 1-5 proceed identically to the first example.)

- 6. Create a *single*-entered file by calling the CreatePairLinksSingleEntered function. At minimum, pass the (i) outcome dataset, the (ii) linking dataset, and the (iii) name(s) of the outcome variable(s).
- 7. Declare the names of the outcome variables corresponding to the two members in each pair. Assuming the variable is called 'ZZZ' and the preceding steps have been followed, the variable 'ZZZ_1' corresponds to the first members and ZZZ_2' corresponds to the second members.
- 8. Create a GroupSummary data.frame, which identifies the R groups that should be considered by the model. Inspect the output to see if the groups show unexpected or fishy differences.
- 9. Create a data.frame with cleaned variables to pass to the SEM function. This data.frame contains only the three necessary rows and columns.
- 10. Estimate the SEM with the lavaan package. The function returns an S4 object, which shows the basic ACE information.
- 11. Inspect details of the SEM, beyond the ACE components. In this example, we look at the fit stats and the parameter estimates. The lavaan package has additional methods that may be useful for your purposes.

```
### R Code for Example lavaan estimation analysis with a simple outcome and Gen2 subjects
#Steps 1-5 are explained in the vignette's first example:
require(NlsyLinks)
dsLinking <- subset(Links79Pair, RelationshipPath=="Gen2Siblings")</pre>
```

```
dsOutcomes <- ExtraOutcomes79
dsOutcomes$MathStandardized[dsOutcomes$MathStandardized < 0] <- NA
#Step 6: Create the single entered dataset.
dsSingle <- CreatePairLinksSingleEntered(outcomeDataset=dsOutcomes,</pre>
  linksPairDataset=dsLinking, outcomeNames=c('MathStandardized'))
#Step 7: Declare the names for the two outcome variables.
oName_1 <- "MathStandardized_1" #Stands for Outcome1
oName_2 <- "MathStandardized_2" #Stands for Outcome2
#Step 8: Summarize the R groups and determine which groups can be estimated.
dsGroupSummary <- RGroupSummary(dsSingle, oName_1, oName_2)</pre>
dsGroupSummary
         R Included PairCount O1Mean O2Mean O1Variance O2Variance O1O2Covariance
                         2718 94.64 95.60
## 1 0.250
              TRUE
                                                 169.7
                                                            207.8
                                                                           41.08
                         139 92.60 93.17
## 2 0.375
              TRUE
                                                 172.5
                                                            187.1
                                                                           40.48
## 3 0.500
              TRUE
                         5511 99.89 100.18
                                                 230.5
                                                            233.0
                                                                          107.37
                           2 108.50 106.00
## 4 0.750
           FALSE
                                                220.5
                                                            18.0
                                                                          63.00
                           22 98.64 95.55
## 5 1.000
              TRUE
                                                 319.2
                                                            343.1
                                                                          277.59
   Correlation Determinant PosDefinite
## 1
        0.2188
                      33573
                                    TRUE
## 2
         0.2253
                       30639
                                    TRUE
## 3
         0.4633
                       42172
                                    TRUE
## 4
         1.0000
                           0
                                   FALSE
## 5
         0.8388
                       32466
                                    TRUE
#Step 9: Create a cleaned dataset
dsClean <- CleanSemAceDataset(dsDirty=dsSingle, dsGroupSummary, oName_1, oName_2)
#Step 10: Run the model
ace <- AceLavaanGroup(dsClean)</pre>
ace
## [1] "Results of ACE estimation: [show]"
## ASquared CSquared ESquared CaseCount
##
     0.6673
               0.1188
                         0.2139 8390.0000
#Notice the `CaseCount' is 8,390 instead of 17,440.
# This is because (a) one pair with R=.75 was excluded, and
# (b) the SEM uses a single-entered dataset instead of double-entered.
#Step 11: Inspect the output further
require(lavaan) #Load the package to access methods of the lavaan class.
GetDetails(ace)
## lavaan (0.5-14) converged normally after 62 iterations
##
##
    Number of observations per group
                                                     2718
##
##
    2
                                                      139
##
    3
                                                     5511
##
                                                       22
##
## Estimator
                                                       MT.
```

```
Minimum Function Test Statistic
                                                    447.272
##
     Degrees of freedom
                                                         16
                                                      0.000
##
     P-value (Chi-square)
##
  Chi-square for each group:
##
##
##
     1
                                                    282.194
     2
##
                                                     32.858
##
     3
                                                     127.880
##
                                                      4.339
#Examine fit stats like Chi-Squared, RMSEA, CFI, etc.
fitMeasures(GetDetails(ace)) #'fitMeasures' is defined in the lavaan package.
##
                fmin
                                   chisq
                                                         df
                                                                       pvalue
##
               0.027
                                447.272
                                                    16.000
                                                                        0.000
                                           baseline.pvalue
##
      baseline.chisq
                            baseline.df
                                                                           cfi
##
            1499.091
                                  4.000
                                                     0.000
                                                                        0.712
##
                 tli
                                   nnfi
                                                       rfi
                                                                          nfi
##
               0.928
                                  0.928
                                                      0.925
                                                                        0.702
##
                                    ifi
                                                                          logl
                pnfi
                                                       rni
               2.807
                                                      0.712
##
                                  0.709
                                                                   -68369.132
## unrestricted.logl
                                   npar
                                                       aic
                                                                           bic
##
          -68145.496
                                                136746.264
                                                                   136774.404
                                   4.000
##
              ntotal
                                   bic2
                                                               rmsea.ci.lower
                                                     rmsea
            8390.000
                                                                        0.104
##
                             136761.692
                                                      0.113
      rmsea.ci.upper
##
                           rmsea.pvalue
                                                       rmr
                                                                   rmr_nomean
##
               0.123
                                  0.000
                                                    13.895
                                                                       17.820
##
                srmr
                            srmr_nomean
                                                      cn_05
                                                                        cn_01
##
               0.128
                                   0.093
                                                   494.269
                                                                      601.260
##
                 gfi
                                                                          mfi
                                   agfi
                                                      pgfi
               0.999
                                  0.999
                                                      0.799
                                                                        0.975
#Examine low-level details like each group's individual parameter estimates and standard
# errors. Uncomment the next line to view the entire output (which is roughly 4 pages).
```

5 Example: Multiple Group SEM of a Simple Outcome for Gen1 Subjects

#summary(GetDetails(ace))

The example differs from the previous one in three ways. First, Gen1 subjects are used. Second, standardized height is used instead of math. Third, pairs are dropped if their R is zero; we return to this last issue after the code is run.

```
### R Code for Example lavaan estimation analysis with a simple outcome and Gen1 subjects
#Steps 1-5 are explained in the vignette's first example:
require(NlsyLinks)
dsLinking <- subset(Links79Pair, RelationshipPath=="Gen1Housemates")
dsOutcomes <- ExtraOutcomes79
#The HeightZGenderAge variable is already groomed

#Step 6: Create the single entered dataset.
dsSingle <- CreatePairLinksSingleEntered(outcomeDataset=dsOutcomes,</pre>
```

```
linksPairDataset=dsLinking, outcomeNames=c('HeightZGenderAge'))
#Step 7: Declare the names for the two outcome variables.
oName_1 <- "HeightZGenderAge_1"
oName_2 <- "HeightZGenderAge_2"
#Step 8: Summarize the R groups and determine which groups can be estimated.
dsGroupSummary <- RGroupSummary(dsSingle, oName_1, oName_2)
rGroupsToDrop <- c( 0 )
dsGroupSummary[dsGroupSummary$R %in% rGroupsToDrop, "Included"] <- FALSE
dsGroupSummary
##
          R Included PairCount
                                  01Mean
                                            O2Mean O1Variance O2Variance O1O2Covariance
## 1 0.0625
                            37 -0.39797 0.019119
                                                                   0.9773
                                                                                 0.37990
                TRUE
                                                       0.7872
## 2 0.1250
                TRUE
                            63 -0.15982 -0.455057
                                                       0.9499
                                                                   0.9529
                                                                                 0.09187
## 3 0.2500
                TRUE
                           267 0.05476 0.006209
                                                                                 0.26929
                                                       0.9853
                                                                   1.1350
                                                                                 0.44371
## 4 0.5000
                TRUE
                           3778 -0.02323 -0.023449
                                                       0.9769
                                                                   1.0194
                            11 0.02395 0.146080
                                                       0.2909
                                                                   0.6106
                                                                                 0.37473
## 5 1.0000
                TRUE
##
     Correlation Determinant PosDefinite
## 1
         0.43312
                     0.62499
                                     TRUE
## 2
         0.09656
                     0.89677
                                     TRUE
## 3
         0.25465
                     1.04577
                                     TRUE
## 4
         0.44465
                     0.79890
                                     TRUE
## 5
         0.88917
                     0.03719
                                     TRUE
#Step 9: Create a cleaned dataset
dsClean <- CleanSemAceDataset(dsDirty=dsSingle, dsGroupSummary, oName_1, oName_2)
#Step 10: Run the model
ace <- AceLavaanGroup(dsClean)</pre>
ace
## [1] "Results of ACE estimation: [show]"
    ASquared CSquared ESquared CaseCount
      0.8834
##
                0.0087
                           0.1079 4156.0000
#Step 11: Inspect the output further (see the final step in the previous example).
```

Most of them responded they were Non-relatives to the explict items asked in 1979 (i.e., NLSY79 variables R00001.50 through R00001.59). Yet their height's observed correlations is far larger than would be expected for a sample of unrelated subjects. Since our team began BG research with the NLSY in the mid-1990s, the R=0 group has consistently presented higher than expected correlations, across many domains of outcome variables. For a long time, we have substantial doubts that subject pairs in this group share a low proportion of their selective genes. Consequently, we suggest applied researchers consider excluding this group from their biometric analyses. **Joe, is there anything else you'd like to say about this here?**

If you wish to exclude additional groups from the analyses, one line of code in Step 8 should change. For instance, two exclude ambiguous sibs (in addition to R=0 pairs), change

```
rGroupsToDrop <- c( 0 )
to
rGroupsToDrop <- c( 0, .375 ).
```

6 Example: Midstream data manipulation with SAS

The example differs from the previous one substantial way: After R is used to link the related pairs, and connect them to their outcome values, the dataset is exported so that the user can further manipulate the data in SAS.

After a presentation, several audience members at the 2012 BGA meeting informed us that this vignette example would help them be more efficient. This approach is also consistent with our feeling that analysts should use the workflow tools that are best suited to their needs and capabilities.

Should we export/import as CSVs (which would make the R code much simpler, because the foreign package wouldn't have to be loaded), but increase the code necessary on the SAS side? I really don't want to mess with sasbdats and xports.

A Appendix: Receiving Help for the NlsyLinks Package

A portion of our current grant covers a small, part-time support staff. If you have questions about BG research with our kinship links, or questions about our package, we'd like to hear from you.

We provide personal support for researchers in several ways. Perhaps the best place to start are the forums on R-Forge (http://r-forge.r-project.org/forum/?group_id=1330); there are forums for people using R, as well as other software such as SAS. This post is a good overview of the current project is, which originally was an email Joe sent to previous users of our kinship links (many of them are/were SAS users).

B Appendix: Creating and Saving R Scripts

There are several options and environments for executing R code. Our current recommendation is RStudio, because it is easy to install, and has features targeting beginnner and experienced R users. We've had good experiences with it on Windows, OS X, and Ubuntu Linux.

RStudio allows you to create and save R files; these are simply text files that have an file extension of '.R'. RStudio will execute the commands written in the file. Help documentation for RStudio can be found at http://www.rstudio.com/ide/docs/.

C Appendix: Installing and Loading the NlsyLinks Package

There are three operations you'll typically do with a package: (a) install, (b) load, and (c) update.

The simplest way to **install** NlsyLinks is to type **install.packages("NlsyLinks")**. You may be asked to select a CRAN mirror to download the package from; if so, choose a close location.

R then will download NlsyLinks on your local computer. It may try to save and install the package to a location that you don't have permission to write files in. If so, R will ask if you would like to install it to a better location (*i.e.*, somewhere you do have permission to write files). Approve this decision (which is acceptable for everyone except for some network administrators).

For a given computer, you'll need to *install* a package only once for each version of R (new versions of R are released every few months). However, you'll need to *load* a package in every session that you call its functions. To **load** NlsyLinks, type either library(NlsyLinks) or require(NlsyLinks); (the difference between the two commands is likely irrelevant for your uses.) Loading reads NlsyLinks information from the hard drive and places it in temporary memory. Once it's loaded, you won't need to load it again until R is closed and reopened later.

Developers are continually improving their packages by adding functions and documentation. These newer versions are then uploaded to the CRAN servers. You may **update** all your installed packages at once by typing **update.packages**(). The command checks a CRAN server for newer versions of the packages installed on your local machine. Then they are automatically downloaded and installed.

The grant supporting NlsyLinks extends until Summer 2014. Until then, we'll be including new features and documentation, as we address additional user needs (if you have suggestions, we'd like to hear from you). When the NLSY periodically updates its data, we'll update our kinship links (embedded in NlsyLinks) with the newest information.

D Appendix: References

Carey, Gregory (2002). Human Genetics for the Social Sciences. Sage.

Plomin, Robert (1990). Nature and nurture: an introduction to human behavioral genetics. Brooks/Cole Publishing Company.

Rodgers, Joseph Lee, & Kohler, Hans-Peter (2005). Reformulating and simplifying the DF analysis model. Behavior Genetics, 35 (2), 211-217.

Rodgers, Joseph Lee, Rowe, David C., & Li, Chengchang (1994). Beyond nature versus nurture: DF analysis of nonshared influences on problem behaviors. *Developmental Psychology*, 30 (3), 374-384.

Neale, Michael C., & Cardon, Lou R. (1992). Methodology for genetic studies of twins and families. Norwell, MA: Kluwer Academic Publishers. (Also see Neale & Maes: http://www.vipbg.vcu.edu/OpenMxFall09/NMbook05.pdf).

E Notes

This package's development was largely supported by the NIH Grant 1R01HD65865, "NLSY Kinship Links: Reliable and Valid Sibling Identification" (PI: Joe Rodgers; Vignette Construction by Will Beasley)

F Version Information

- R version 3.0.2 (2013-09-25), x86_64-w64-mingw32
- Locale: LC_COLLATE=English_United States.1252, LC_CTYPE=English_United States.1252, LC_MONETARY=English_United States.1252, LC_NUMERIC=C, LC_TIME=English_United States.1252
- Base packages: base, datasets, graphics, grDevices, methods, stats, utils
- Other packages: boot 1.3-9, knitr 1.5, lavaan 0.5-14, MASS 7.3-29, Matrix 1.1-0, mnormt 1.4-5, NlsyLinks 1.220, pbivnorm 0.5-1, quadprog 1.5-5
- Loaded via a namespace (and not attached): evaluate 0.5.1, formatR 0.10, grid 3.0.2, highr 0.3, lattice 0.20-24, stats4 3.0.2, stringr 0.6.2, tools 3.0.2