Assignment 3 - Network Analysis 2022

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Conceptual Questions

Question 1 (1 point)

Are the following statements true or false? Explain why (0.5 point per statement).

1. If ergodicity holds, results from between-person analysis are expected to equal results from within-person analysis.

Yes, if ergodicity holds, there should be no difference between individuals (i.e., no difference in between-person), and correspondingly it aligns with the within-person analysis of every individual (Molenaar, 2004).

2. It is generally recommended to always remove trends (such as linear trends) prior to analyzing your N=1 time series.

No, even though some simulation stuides show detrending helps increasing the performance of estimated networks (Epskamp et al., 2018), it is often the case that these changing trends are of main interest. So it is not correct to say it is *always* recommended to detrend (Isvoranu et al., 2022).

Question 2 (1.5 points)

Suppose a therapist measures a patient about 75 times on a set of depression symptoms, including a question on "suicidal thought". Suppose that you estimate a graphical VAR model from this data and find that the node "suicidal thought" is not connected to any of the other nodes in your network, neither in the temporal nor in the contemporaneous network.

List three potential reasons why the node "suicidal thought" may be disconnected in the resulting network.

- 1. Probably the patient answered "no" every time on "suicidal thought, since it is quite an extreme symptom to develop. It is likely that the patient never developed"suicidal thought" and that leads to no variance, which will result in no connections in both temporal and contemporaneous network (Hoekstra, 2022).
- 2. It could be the case that the estimated network model was quite large (> 10 nodes) and it just did not have enough power to retrieve the edges connecting to "suicidal thought". It is in fact not established whether GVAR model can be reliably estimated from the N = 1 data sets (Isvoranu et al., 2022), especially when the model is large and sample size is relatively small, like in this case (n = 75).
- 3. Lastly, another potential scenario is that the measurement period was too short to capture the occurrence of "suicidal thought". For example, the 75 observations were measured by 5 times a day over 2 weeks, which is relatively a short period of time. It could be that "suicidal thought" only started occurring after the measurement period had ended, and hence left the item score on "suicidal thought" constant. Or there was a mismatch between the chosen times scale and "suicidal thought" processes, such that suicidal thought process is much slower than the chosen measurement frequency. It could be that the therapist measured the symptoms every 1-2 hrs, when the suicidal thought process operates over several days or so, which require more distance between assessments (Isvoranu et al., 2022). This could result in barely any variations on "suicidal thought" item and correspondingly leads to empty temporal and contemporaneous network.

Question 3 (1 point)

Give an example of a relationship that can only be studied at the between-person level.

Things that can only be studied at the *between-person* level have to be something that stay stable and do not fluctuate over time. A couple of example relationships that can be studied only in *between-person* level are:

- genetic structure and cancer: do people with a certain genetic structure have higher risk for cancer?
- personality trait and zodiac sign: do people with different zodiac signs have different personality traits?
- average education level and race/nationality: does the average education level differ across different races/nationalities?

Question 4 (2 points)

During the lecture we discussed multiple challenges regarding time-series modeling in the network approach. Pick your favorite challenge and explain this challenge in your own words. Make sure to not only explain what the challenge is, but also why this is a challenge: in what way may this challenge impact your results (i.e., the network model you estimate) and how may this jeopardize your conclusions?

One of the challenges in time-series network modelling is the difficulty of incorporating variables that operate on different time-scales. For example, the symptom network (e.g., network consist of symptoms) evolves relatively fast, as symptoms generally fluctuate over days. Whereas, psychological resilience, which is one of the important protective factors against developing mental disorder, evolves rather slowly and gradually (Lunansky et al., 2020). Obviously, resilience is part of the dynamic of psychopathology, but there exists no such model that can incorporate variables operating on the different time-scales. Absence of such a model prevents us from analyzing the whole network that integrates the slow-changing variables (e.g., resilience) with the fast-changing symptoms. Note that Lunansky et al. (2020) looked into the interaction between slow and fast network processes but it was studied in such a way that the slow changing variables influence the nodes in the symptom network instead of incorporating them all in one network model.

According to Borsboom & Van Borkulo (2021), omitting this relevant factor that operates on slower time-scale from the symptom network could lead to a much denser network. The reasoning is as follows. If the slow-changing variable (SV) is omitted in the symptom network model, then the edges will be estimated without controlling for the SV. Assuming that the SV has consistent influences on the symptoms, not controlling for the SV is likely to exaggerate the edges (i.e., partial correlations) between the symptoms. Therefore, it is expected that the symptom network model excluding the SV would appear denser (i.e., many thicker edges) than the integrated network model including the SV.

Hence, basing the network model only on the symptoms without incorporating the relevant SV can result in misleading model that overestimates the strength of relationships between symptoms. This could be especially detrimental to studies that look into a particular connection between two symptoms. If a SV that has a considerable influence on the symptom dynamics is not included in the model, the researcher may draw a conclusion based on one of the spurious connections in the network, which are resulted from overlooking the effects of the SV.¹

Question 5 (2 points)

In a recent study by Haslbeck et al., (preprint) it was shown that use of a VAS or Likert scale has an affect on the observed distributions. Take a look at their article. Give (a) a short summary of the problem described in their paper, (b) reflect on this problem (e.g., why is this an issue and how

 $^{^{1}}Note$. Reference list can be found at the end of the document.

does it affect the interpretation of our results?) and (c) think of a study to specifically test whether this phenomenon (different scales lead to different distributions) is a methodological artifact or a "true" phenomenon (max 250 words).

Haselbeck et al. assessed the distributional modality and skewness in emotional measurements using seven different emotion ESM datasets and investigated whether there are any patterns of the distributional forms on the level of items, individuals, and measurement designs. They found that skewed unimodal distributions as well as multimodality are highly prevalent across the datasets and the presence of multimodality was found to be strongly associated with the measurement designs such that multimodality was more prevalent in VAS (visual analog scale) than in the Likert scales (e.g., 1-5, or 1-7). In addition, they found some other associations on the item level such as negative emotions exhibit higher skewness than position emotions.

This has a great implication on multiple aspects. First, with regard to theorizing emotion dynamics, if the found multimodality and high skewness is indeed true dynamics of emotion, then it implies that a person experiences emotions by going through multiple states (intensity) instead of experiencing it one typical intensity all the time (i.e., unimodal). Additionally, if we presume the multimodality, then the measurement design should preferably include a broader scale (e.g., VAS with 0 – 100 scale) rather than a limited scale such as Likert scale of 1 – 5, so that it can capture the variabilities across multiple peaks. Lastly, the VAR model might not be deemed appropriate if the true underlying dynamic follows a skewed/multimodal distribution as the VAR model assumes a single equilibrium state that is drawn from a normal distribution. Accordingly, a VAR model will fit poorly on the skewed or multimodally-distributed time-series emotion data, and correspondingly interpreting its parameters would be misleading.

However, it is difficult to reason whether the found result by Haselbeck et al. actually reflects the true emotion dynamics or it might be just some sorts of methodological artifacts (e.g., specific measurement design induce multimodality). In order to test this, we could design a study such that participants are asked to report on each item with different measurement scales. For example, a participant reports each item twice, once with VAS (1 - 100 scale) and once with Likert scale (e.g., 1 - 7) at every occasion and in the end we can compare the distribution of items to see if they approximate to each other or not. If the distributions of item based on VAS and Likert-scale are turned out to be similar (i.e., multimodal), then we can be more confident in presuming that the found multimodality and skewness do reflect the true phenomenon.

Question 6 (1 point)

What do the *day* and *beep* arguments in the packages graphicalVAR, psychonetrics, and mlVAR do? Suppose you have a dataset containing only one observation per day for every weekday (but not weekends) in a year long study. Which argument would you use (and why) to make sure that measures from Mondays are not regressed on measures from Fridays?

- dayvar argument indicates assessment day and when you add this, it ensures that the first measurement of a day is not regressed on the last measurement of the previous day (it removes pairs of observations that cross a night).
- beepvar argument indicates the assessment beep per day, and when you add this, it will treat the non-consecutive beeps as missing (removes pairs of observations that are not consecutive).
- In this case, we want to regress measures of each weekdays on measures of the previous days but there are breaks between Mondays and Fridays. We can then use dayvar argument and specify the same number to each week (e.g., Monday Friday of week 1: dayvar = 1 and Monday Friday of week 2: dayvar = 2, ...) then the measures from Mondays are not going to be regressed on the measures from Fridays of the previous week.

Practical Questions

```
# load data
load("clean_network.RData")

# Variables to investigate:
vars <- paste0("Q",1:18)

# Labels:
varLabs <- c("Relax","Irritable","Worry","Nervous","Future","Anhedonia",
"Tired","Hungry","Alone","Angry","Social_offline","Social_online",
"Music","Procrastinate","Outdoors","C19_occupied","C19_worry",
"Home")

# Rename columns in data:
names(Data2) [names(Data2) %in% vars] <- varLabs

# Remove items:
Data2 <- Data2 %>% select(-Hungry,-Angry,-Music,-Procrastinate)
varLabs <- varLabs[!varLabs %in% c("Hungry","Angry","Music","Procrastinate")]</pre>
```

N = 1 time series

```
## select a random participant
student_number <- 12183881
set.seed(student_number)
subject <- sample(Data2$id, 1)
my_data_copy <- my_data <- Data2[which(Data2$id == subject),]

I chose 4 variables: "Irritable", "Worry", "Nervous", "Alone".
chosen_var <- c("Irritable", "Worry", "Nervous", "Alone")</pre>
```

Question 7 (1.5 points)

Estimate a *saturated* (no model selection) GVAR model on your sampled subject using the 3 or 4 variables you chose above using psychonetrics. Plot the estimated temporal (partial directed correlations) and contemporaneous networks with the same layout (1 point). Which edges are significant (0.5 point)?

```
## estimate GVAR model
res1 <- gvar(my_data, vars = chosen_var, dayvar = "day", beepvar = "beep", estimator="FIML") %>% runmodel
## temporal network
temporal1 <- getmatrix(res1, "PDC") # PDC = Partial Directed Correlations
## contemporaneous network
contemporaneous1 <- getmatrix(res1, "omega_zeta")</pre>
```

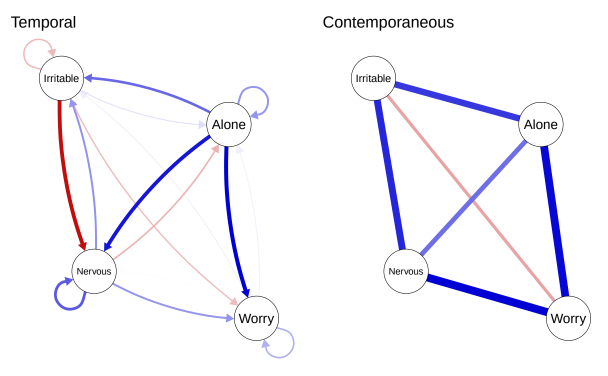


Figure 1: Estimated graphical vector auto-regression (GVAR) model

Assuming that $\alpha = 0.05$,

- there is no significant edge found in the temporal network as all p > 0.05 in the beta matrix, as shown below.
- the significant edges in the contemporaneous network are: Nervous -- Irritable and Alone--Worry as p < 0.05 in the omega-zeta matrix.

```
Parameters for group fullsample
  - mu
         var1 op var2 est
                                      p row col par
Irritable_lag1 ~1
                     1.62 0.13 < 0.0001
                                          1
                                                  1
 Worry_lag1 ~1
Nervous_lag1 ~1
                     2.02 0.10 < 0.0001
                                                  2
                   1.95 0.13 < 0.0001 3 1
                                                 3
   Alone_lag1 ~1
                   1.39 0.089 < 0.0001 4 1
                                                  4
    Irritable ~1
                     1.49 0.10 < 0.0001
                                          5 1
                                                 5
                     1.86 0.096 < 0.0001
        Worry ~1
                                                  6
```

```
Nervous ~1
                     1.82 0.11 < 0.0001
                                         7
        Alone ~1
                     1.29 0.070 < 0.0001
  exo_cholesky
         var1
                             var2
                 op
                                    est
                                          se
                                                    p row col par
Irritable_lag1 ~chol~ Irritable_lag1  0.80 0.096 < 0.0001</pre>
   Worry_lag1 ~chol~ Irritable_lag1 0.17 0.11
                                                 0.11
                                                           1 10
 Nervous_lag1 ~chol~ Irritable_lag1 0.41 0.12 0.00097
                                                          1 11
   Alone_lag1 ~chol~ Irritable_lag1 0.22 0.090
                                                0.015
                                                          1 12
                                                          2 13
   Worry_lag1 ~chol~
                        Worry_lag1
                                   0.62 \ 0.075 < 0.0001
                        Worry_lag1 0.34 0.11
                                                        3 2 14
 Nervous_lag1 ~chol~
                                               0.0014
   Alone_lag1 ~chol~
                        Worry_lag1 0.24 0.080
                                               0.0030
                                                        4 2 15
 Nervous_lag1 ~chol~
                    Nervous lag1 0.58 0.070 < 0.0001
                                                        3 3 16
                      Nervous lag1 0.078 0.075
   Alone lag1 ~chol~
                                                 0.30
                                                        4 3 17
                        Alone_lag1 0.44 0.053 < 0.0001
   Alone_lag1 ~chol~
                                                        4 4 18
  - beta
    var1 op
                var2
                        est
                             se
                                    p row col par
Irritable <- Irritable -0.13 0.25 0.61
                                              19
   Worry <- Irritable -0.099 0.24 0.68
                                            1
                                              20
 Nervous <- Irritable -0.46 0.28 0.10
                                           1 21
   Alone <- Irritable 0.043 0.16 0.79
                                           1 22
Irritable <-</pre>
               Worry 0.044 0.25 0.86
                                           2 23
                                           2 24
   Worry <-
               Worry
                     0.16 0.25 0.53
                                       2
 Nervous <-
               Worry 0.0054 0.27 0.98
                                           2 25
   Alone <-
               Worry 0.029 0.17 0.86
                                          2 26
Irritable <-
             Nervous 0.24 0.26 0.36
                                           3 27
                                       1
   Worry <-
             Nervous 0.19 0.25 0.45
                                           3 28
 Nervous <-
             Nervous 0.35 0.28 0.21
                                       3 3 29
   Alone <-
             Nervous -0.12 0.17 0.46
                                      4 3 30
                                          4 31
Irritable <-</pre>
              Alone 0.43 0.34 0.20 1
                                       2 4 32
   Worry <-
              Alone 0.62 0.35 0.075
 Nervous <-
               Alone 0.66 0.38 0.079 3 4 33
   Alone <-
               Alone 0.22 0.22 0.32
                                      4 4 34
  - omega_zeta (symmetric)
              var2 est
  var1 op
                         se
                                 p row col par
  Worry -- Irritable -0.13 0.18 0.46
                                         1 35
Nervous -- Irritable 0.32 0.15 0.033 3
                                        1 36
  Alone -- Irritable 0.29 0.16 0.061 4
                                       1 37
Nervous --
             Worry 0.37 0.20 0.063 3 2 38
  Alone --
             Worry 0.36 0.17 0.032 4 2 39
  Alone -- Nervous 0.21 0.16 0.19 4
                                        3 40
  - delta_zeta (diagonal)
    var1 op
                 var2 est
                                      p row col par
```

se

Worry 0.42 0.067 < 0.0001

Alone $0.37 \ 0.051 < 0.0001$

Irritable ~/~ Irritable 0.56 0.062 < 0.0001

Nervous ~/~ Nervous 0.48 0.063 < 0.0001

Worry ~/~

Alone ~/~

1

2

1 41

2 42

4 44

3 3 43

Question 8 (1 point)

Estimate a GVAR model on your sampled subject using the 3 or 4 variables you chose above using the graphicalVAR package (use $\gamma = 0$), and compare your results to the results of the previous question.

The resulting GVAR model estimated by graphicalVAR (Figure 2) is more sparse compared to the GVAR model estimated by psychonetrics::gvar (Figure 1). That is because the temporal network model is estimated with LASSO regularization and the contemporaneous network model is estimated with GLASSO using BIC (as $\gamma = 0$) in graphicalVAR, whereas the networks in Figure 1 are estimated without any model selection/regularization.

Partial Directed Correlations

Partial Contemporaneous Correlations

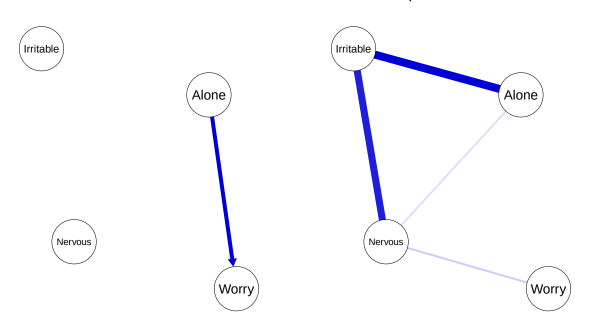


Figure 2: Estimated graphical vector auto-regression (GVAR) model using graphical VAR function

```
\#plot(res2, include = c("PCC", "PDC"), titles = TRUE, sameLayout = TRUE, theme = "colorblind")
```

Question 9 (1.5 points)

Test for significant trends ($\alpha=0.05$) for your selected variables and detrend these variables if the trends are significant. Then, re-estimate the networks using either graphicalVAR or psychonetrics. Did your estimated networks change?

```
## Detrending significant linear trends
for (v in seq_along(chosen_var)){
    ff <- as.formula(pasteO(chosen_var[[v]]," ~ conc"))
    fit <- lm(ff, data = my_data)
    if (anova(fit)$P[1] < 0.05){
        message(paste("Detrending variable:", chosen_var[v], "as the p-value is", round(anova(fit)$P[1],3)))
        # detrend the found variables with significant linear trends
        my_data[[chosen_var[v]]][!is.na(my_data[[chosen_var[[v]]]])] <- residuals(fit)
    }
}</pre>
```

Detrending variable: Worry as the p-value is 0.049

Detrending variable: Nervous as the p-value is 0.008

Re-estimated networks after detrending the variable Worry and Nervous are shown below in Figure 3. Overall, the networks estimated after detrending barely show any changes. The ones estimated with psychonetrics seems more or less the same with the networks estimated on non-detrended data (Figure 1) except for a few changes in the weak edges (faded ones) in the temporal network. The networks estimated with grahicalVAR() are also shown to be almost the same as the ones estimated on non-detrended data (Figure 2). The only visible change is

that the edge between Nervous -- Worry is gone in the contemporaneous network after detrending.²

```
## after detrending, estimate GVAR model using psychonetrics:
res3 <- gvar(my_data, vars = chosen_var, dayvar = "day", beepvar = "beep", estimator="FIML") %>% runmodel
# temporal network
temporal3 <- getmatrix(res3, "PDC")</pre>
# contemporaneous network
contemporaneous3 <- getmatrix(res3, "omega_zeta")</pre>
## after detrending, estimate GVAR model using graphicalVAR:
res4 <- graphicalVAR(my_data, vars = chosen_var, gamma = 0, dayvar = "day", beepvar = "beep")
## plot networks after detrending
layout(t(matrix(1:4, 2, 2)))
# psychonetrics GVAR networks
qgraph(temporal3, layout = L1, theme = "colorblind", directed=TRUE, diag=TRUE,
       vsize = 12, mar = rep(6,4), asize = 5, labels = labs)
title(main = "Detrended Temporal (psychonetrics)", font.main = 1, cex.main=1)
qgraph(contemporaneous3, layout = L1, theme = "colorblind", vsize = 12,
       mar = rep(6,4), asize = 5, labels = labs)
title(main = "Detrended Contemporaneous (psychonetrics)", font.main = 1, cex.main=1)
# graphicalVAR networks
plot(res4, "PDC", layout = L1, main = "Temporal", vsize = 12, mar = rep(6,4), asize = 5,
       labels = labs, theme="colorblind", titles = FALSE)
title(main = "Detrended Temporal (graphicalVAR)", font.main = 1, cex.main=1)
plot(res4, "PCC", layout = L1, main = "Contemporaneous", vsize = 12, mar = rep(6,4), asize = 5,
       labels = labs, theme = "colorblind", titles=FALSE)
title(main = "Detrended Contemporaneous (graphicalVAR)", font.main = 1, cex.main=1)
```

 $^{^{2}}Note$. See the *Appendix* for the full summary results of linear models.

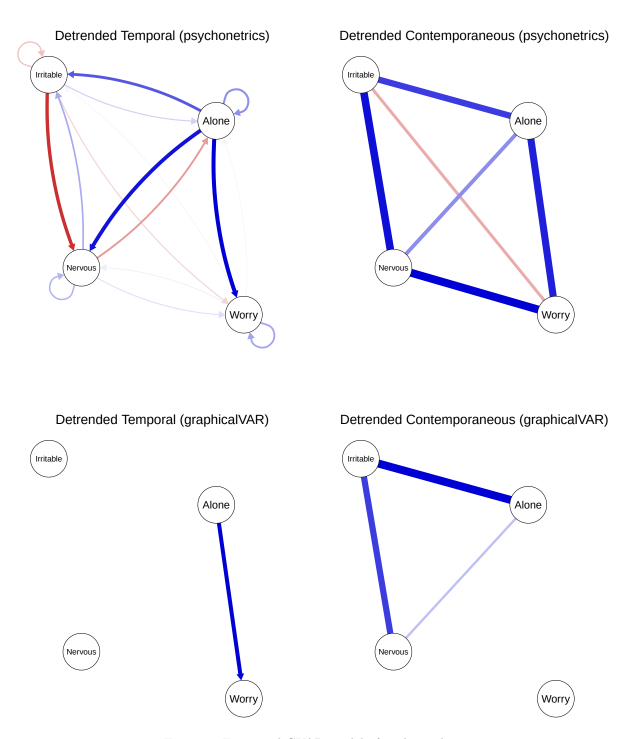


Figure 3: Estimated GVAR model after detrending

Question 10 (1 point)

For question 8 you estimated a graphical VAR network containing 3 or 4 variables for a sampled subject using the graphical VAR package. Repeat this process for another subject. Now that you have estimated two graphical VAR models on non-detrended data, plot the resulting networks and inspect the similarities and differences between these estimated graphical VAR networks. Can you conclude there is heterogeneity between these two people? Why (not)?

Just at a glance, it seems two people are different as the estimated network structures look different (see Figure 4). However, we cannot really conclude that there is heterogeneity based on visually comparing the network structures, because we cannot ensure the these estimated GVAR models are reliable. It could be that they are just random noises, then we would compare a set of random noises to another set of random noises. According to Hoekstra et al. (2022), even when the generating structure is the same, it is likely that we would find different network structures. Hence, it is not advisable to interpret the differences between individual network models as an evidence for heterogeneity, especially when the networks are sparse as in this case.

```
## Another seed: select another random subject
set.seed(13294889)
subject2 <- sample(Data2$id, 1)</pre>
my data2 <- Data2[which(Data2$id == subject2),]</pre>
## Estimate GVAR model for another subject
res5 <- graphicalVAR(my_data2, vars = chosen_var, gamma = 0, dayvar = "day", beepvar = "beep")
## plot networks and compare
layout(t(matrix(1:4, 2, 2)))
\# subject from Q8 (id = 68)
plot(res2, "PDC", layout = L1, vsize = 12, mar = rep(6,4), asize = 5,
       labels = labs, theme="colorblind", details=T, titles=FALSE)
title(main = "Temporal (subject id=68)", cex.main = 1, font.main = 1)
plot(res2, "PCC", layout = L1, vsize = 12, mar = rep(6,4), asize = 5,
       labels = labs, theme = "colorblind",details=T, titles=FALSE)
title(main = "Contemporaneous (subject id=68)", cex.main = 1, font.main = 1)
# another subject (id = 33)
plot(res5, "PDC", layout = L1, vsize = 12, mar = rep(6,4), asize = 5,
       labels = labs, theme="colorblind",details=T, titles=FALSE)
title(main = "Temporal (subject id=33)", cex.main = 1, font.main = 1)
plot(res5, "PCC", layout = L1, vsize = 12, mar = rep(6,4), asize = 5,
       labels = labs, theme = "colorblind", details=T, titles=FALSE)
title("Contemporaneous (subject id=33)", cex.main = 1, font.main = 1)
```

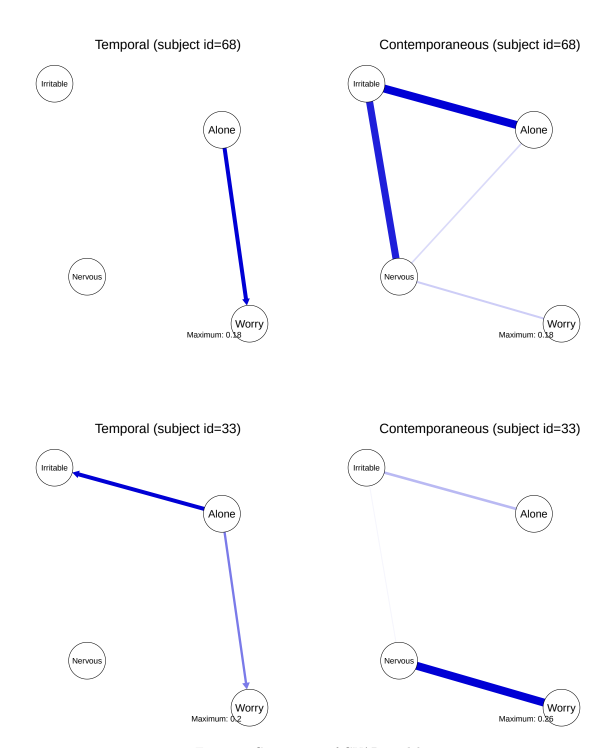


Figure 4: Comparison of GVAR model

N > 1 time-series

Question 11 (1 point)

Look at the help file for the mlVAR function and estimate a multilevel GVAR model on the entire dataset on 4 to 6 variables of your choice.

Question 12 (1 point)

Look at the mIVAR plot method help file (?plot.mlVAR), and plot the estimated fixed-effect temporal, contemporaneous and between-subjects networks. Plot all networks with the same layout (circle layout or average layout), and hide (threshold) edges that are not significant at $\alpha = 0.05$. For the contemporaneous and between-subjects networks, use an "and"-rule to minimize type-1 error rate in showing edges.

```
# Get networks:
temp <- getNet(mlVAR res, "temporal", nonsig = "hide")</pre>
cont <- getNet(mlVAR_res, "contemporaneous", nonsig = "hide", rule = "and")</pre>
bet <- getNet(mlVAR res, "between", nonsig = "hide", rule = "and")
# Get average layout
L <- averageLayout(cont,bet,temp)</pre>
## Plot networks
par(mfrow=c(1,3))
plot(mlVAR_res, "temporal", title="(a) Temporal (Lag-1)", layout = L, nonsig = "hide",
     alpha = 0.05, rule = "and", theme='colorblind', vsize = 16, asize = 5,
     mar = rep(7,4), title.cex = 1.3)
plot(mlVAR_res, "contemporaneous", title="(b) Contemporaneous", layout = L, nonsig = "hide",
     alpha = 0.05, rule = "and", theme='colorblind', vsize = 16, asize = 5,
     mar = rep(7,4), title.cex = 1.3)
plot(mlVAR_res, "between", title="(c) Between-subjects", layout = L, nonsig = "hide",
     alpha = 0.05, theme='colorblind', vsize = 16, asize = 5, mar = rep(7,4), title.cex = 1.3,)
```

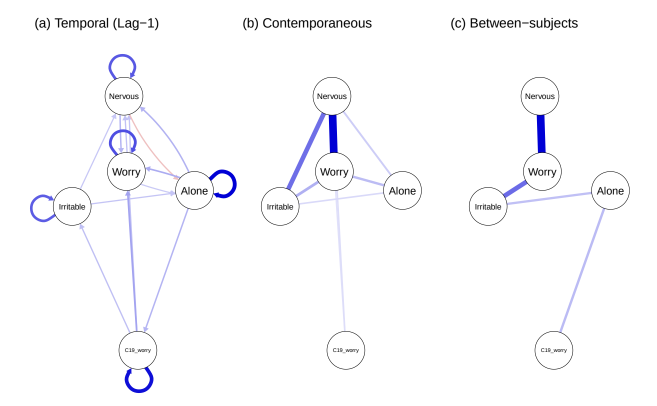


Figure 5: Estimated fixed effects of network structures

Question 13 (2 points)

Plot the estimated individual differences for both the temporal and the contemporaneous network. Explain what the edges in these networks represent. Inspect the plotted networks, what conclusion can you draw about heterogeneity within this sample based on these network structures?

Figure 6 shows the estimated networks of the individual differences, where the edges represent the standard deviation of random effects in the temporal and in the contemporaneous network, respectively.

In the temporal network, the minimum argument is set to 0.1, meaning that only the standard deviation of temporal random effects above 0.1 are shown with a non-transparent arrow, following the advice of Bringmann et al. (2013). From Figure 6 (a) we can see that the largest individual differences are found in the auto-correlations, given that the self-loops of Alone and C19-worry are the most pronounced. It implies that there is a high individual variability over those items such that for some individuals, once they feel alone they tend to feel lonely for long time, but for the other individuals, it is rather momentary. Among the cross-lagged relations, Nervous -> Irritable turns out to have relatively larger individual differences.

When looking at the contemporaneous network in in Figure 6 (b), the largest individual differences are found again in the relationship between Nervous and Irritable, followed by the relationship between Nervous and Worry.

All in all, we can conclude that there is some amount of individual heterogeneity observed in our data. In general, auto-regressions have relatively larger individual differences compared to cross-lagged regressions and the relation between Nervous and Irritable seems to vary a lot across individuals, considering that it is pronounced in both temporal and contemporaneous networks.

```
minimum = 0.1)
## contemporaneous: random effects SD
plot(mlVAR_res , type = "contemporaneous", SD = TRUE, title ="Individual differences - Contemporaneous",
    layout = L, labels = chosen_var2, theme='colorblind', vsize = 12, asize = 4, mar = rep(7,4))
```

Individual differences - Temporal

Individual differences - Contemporaneous

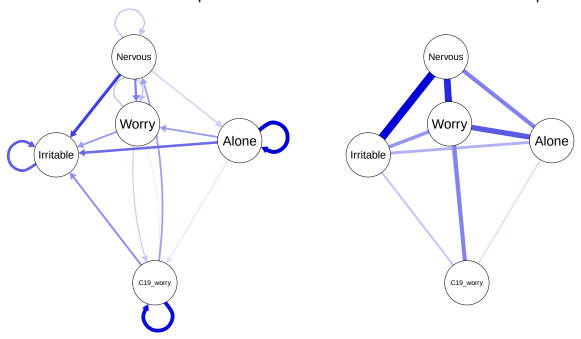


Figure 6: Individual differences networks

Essay Question

Question 14 (2 points)

Inspect these network measures and choose one that would be interesting to compute on one of the estimated networks from any of the previous questions. Report your chosen measure and what it represents for the network you have chosen, explain your choice, and what research question it could answer in the form of a short essay (max 250 words).

After inspecting various measures from Brain Connectivity Toolbox, I decided to look at *in-strength & out-strength* for the fixed effect lag-1 *temporal network* from Figure 5 (a). The overall strength is the sum of absolute edge weights for a given node (Costantini et al., 2015). For the temporal network, which includes directionality of edges, strength can be sub-divided into *in-strength* and *out-strength*. In-strength is the sum of inward edge weights and out-strength is the sum of outward edge weights. In the temporal network, in-strength represents the *relative susceptibility* (i.e., the extent to which a node is influenced by the previous state of the other nodes) and out-strength represents the *relative predictability* (i.e., the extent to which a node has an influence on the following state of the other nodes) (Black et al., 2022).

The motivation to look at in-/out-strength for the temporal network is that in temporal networks, it is interesting to see whether a node is a strong affector (activate other nodes) or an effector (activated by other nodes). Given

that a temporal network contains information not only about the direction of influence but also the strength of influence, in-/out-strength is appropriate measure as it can capture both aspects.

Using in-/out-strength we can answer research questions such as: what are the nodes that have highest predictive power or what are the nodes with highest susceptibility? And based on the result, we may be able to draw an inference on the dynamics of the system by figuring out, for example, which node takes a main role when it comes to activating the system.

Figure 7 shows the in-strength and out-strength of the temporal network from Figure 5 (a). We can see that Alone and C19_worry have relatively high predictive power as they have stronger influence on the following states of other nodes, while Nervous and Worry have high susceptibility as they are affected strongly by the previous state of the other nodes. Based on that, we can assume that the central dynamic in this network consists of Alone and C19Pworry mainly activating Nervous and Worry variables. Lastly, Irritable is shown to be low on both outstrength and in-strength, which implies that it probably does not play a crucial role in the corresponding temporal network dynamic. Note that the stability of centrality measures is not tested. Hence, any inferences based on in-/out-strength values should be drawn with caution.

```
## compute in-/out-strength
inoutStrength <- temp %>% centrality_auto() %>%
    .$node.centrality %>%
    select(InStrength, OutStrength)

## plot in-/out-strength
p1 <- inoutStrength %>%
    ggplot(aes(y = OutStrength, x=reorder(rownames(.), OutStrength))) +
    geom_line(aes(group=1)) + geom_point() + coord_flip() +
    labs(x="", y = "", title = "Out-Strength") + ylim(0.1, 0.26) + theme_bw()
p2 <- inoutStrength %>%
    ggplot(aes(y = InStrength, x=reorder(rownames(.), OutStrength))) +
    geom_line(aes(group=1)) + geom_point() + coord_flip()+
    labs(x="", y = "", title = "In-Strength") + ylim(0.05, 0.26) + theme_bw()
ggarrange(p1, p2)
```

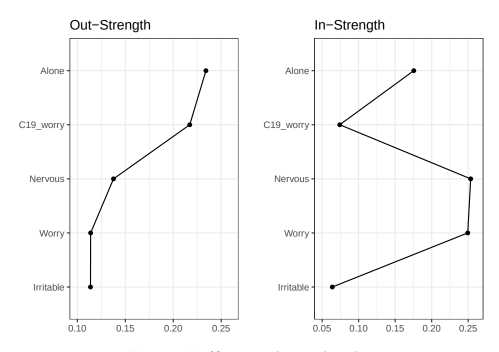


Figure 7: In-/Out-strengh centrality plot

Appendix

Summary results of linear models (Question 9)

```
## run linear regression on chosen variables
lms <- chosen_var %>%
 paste(., '~ conc') %>%
 map(as.formula) %>%
  map(lm, data = my_data_copy)
names(lms) <- chosen_var</pre>
## get the summary results
lapply(lms, summary)
$Irritable
Call:
.f(formula = .x[[i]], data = ..1)
Residuals:
                            ЗQ
   Min
            1Q Median
                                   Max
-0.6114 -0.4959 -0.4076 0.4905 2.4611
Coefficients:
            Estimate Std. Error t value Pr(>|t|)
(Intercept) 1.624942 0.223492 7.271 2.53e-09 ***
          -0.004527 0.006664 -0.679
conc
                                             0.5
Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
Residual standard error: 0.7353 on 49 degrees of freedom
  (5 observations deleted due to missingness)
Multiple R-squared: 0.00933, Adjusted R-squared: -0.01089
F-statistic: 0.4615 on 1 and 49 DF, p-value: 0.5001
$Worry
Call:
.f(formula = .x[[i]], data = ..1)
Residuals:
              1Q Median
                                3Q
-1.19171 -0.57716 0.04182 0.28764 1.28764
Coefficients:
            Estimate Std. Error t value Pr(>|t|)
(Intercept) 2.22859 0.20458 10.894 1.09e-14 ***
           -0.01229 0.00610 -2.015 0.0494 *
conc
```

```
$Nervous
Call:
.f(formula = .x[[i]], data = ..1)
Residuals:
              1Q Median
                               3Q
-1.27329 -0.48327 0.01729 0.33512 2.25339
Coefficients:
            Estimate Std. Error t value Pr(>|t|)
(Intercept) 2.364097 0.218806 10.805 1.45e-14 ***
           conc
Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
Residual standard error: 0.7199 on 49 degrees of freedom
  (5 observations deleted due to missingness)
Multiple R-squared: 0.1365,
                              Adjusted R-squared: 0.1189
F-statistic: 7.748 on 1 and 49 DF, p-value: 0.007617
$Alone
Call:
.f(formula = .x[[i]], data = ..1)
Residuals:
   Min
            1Q Median
                           3Q
                                  Max
-0.3968 -0.3236 -0.2433 0.6550 1.6675
Coefficients:
            Estimate Std. Error t value Pr(>|t|)
(Intercept) 1.400390 0.153096 9.147 3.56e-12 ***
           -0.003570 0.004565 -0.782
                                         0.438
conc
Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
Residual standard error: 0.5037 on 49 degrees of freedom
  (5 observations deleted due to missingness)
Multiple R-squared: 0.01233, Adjusted R-squared: -0.007827
F-statistic: 0.6117 on 1 and 49 DF, p-value: 0.4379
Individual differences manually figured out
                                            18
```

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

Adjusted R-squared: 0.05767

Residual standard error: 0.6731 on 49 degrees of freedom

F-statistic: 4.06 on 1 and 49 DF, p-value: 0.04942

(5 observations deleted due to missingness)

Multiple R-squared: 0.07651,

```
# ## checked with summary result
# summary(mlVAR_res)

# ## temporal: random effects SD
# temp_ran_sd <- mlVAR_res[["results"]][["Beta"]][["SD"]] # mlVAR_res$results$Beta$SD
# tem_inddif <- cbind(rep(1:5,each=5), rep(1:5,5), weight=as.vector(temp_ran_sd))
# agraph(tem_inddif, layout = L, labels = chosen_var2, theme='colorblind', vsize = 12, asize = 4, mar = ref
# ## contemporaneous: random effects SD
# contemp_ran_sd <- mlVAR_res[["results"]][["Theta"]][["pcor"]][["SD"]]
# agraph(contemp_ran_sd, layout = L, labels = chosen_var2, theme='colorblind', vsize = 12, asize = 4, mar</pre>
```

References

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