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ABSTRACT

In their recent paper, Forbes et al. (2019; FWMK) evaluate the replicability of network models in two studies. They identify considerable replicability issues, concluding that “current ‘state-of-the-art’ methods in the psychopathology network literature […] are not well-suited to analyzing the structure of the relationships between individual symptoms”. Such strong claims require strong evidence, which the authors do not provide. FWMK identify low replicability by analyzing point estimates of networks; contrast low replicability with results of two statistical tests that indicate higher replicability, and conclude that these tests are problematic. We make four points. First, statistical tests are superior to the visual inspection of point estimates, because tests take into account sampling variability. Second, FWMK misinterpret the statistical tests in several important ways. Third, FWMK did not follow established recommendations when estimating networks in their first study, underestimating replicability. Fourth, FWMK draw conclusions about methodology, which does not follow from investigations of data, and requires investigations of methodology. We conclude by discussing important recent simulation work that guides researchers to use models appropriate for their data, such as nonregularized estimation routines.

KEYWORDS

gaussian graphical model; network model; regularization; replicability

In their paper entitled “Quantifying the reliability and replicability of psychopathology network characteristics”, Forbes et al. (2019)—from here on FWMK—conducted two studies. First, they estimated Gaussian Graphical Models (GGMs) of 16 depression and anxiety symptoms in two waves of data from an observational study (n = 403) one week apart. Second, they re-analyzed GGMs of 16 posttraumatic stress disorder symptoms across 4 datasets reported by Fried et al. (2018).

In both studies, FWMK highlight replicability issues of network models, and conclude that the “low reliability of partial correlations and high rates of false negatives related to LASSO regularization in such data suggest that the current ‘state-of-the-art’ methods in the psychopathology network literature […] are not well-suited to analyzing the structure of the relationships between individual symptoms of mental illness” (p. 14).

To assess replicability, FWMK compare point estimates of network parameters, and contrast their results of low replicability with the results of two statistical tests that indicate higher replicability. The two tests are part of two R packages, which were developed with the express purpose so that researchers stop overinterpreting point estimates when conducting comparisons. The first package is bootnet (Epskamp et al., 2018), which performs nonparametric and subsampling bootstraps to assess the stability and accuracy of estimated networks. The second package is the Network Comparison Test (NCT; van Borkulo et al., 2017), which uses permutations to test if two samples feature different underlying network structures.

We agree with FWMK that there are numerous challenges in this emerging field, both of substantive and statistical nature. However, we disagree with some of the conclusions the authors draw, and focus on four issues below.
Sampling variability

Statistical tests aim to avoid overinterpretation of point estimates. Suppose we want to compare standardized neuroticism levels across two small samples of 30 participants each, with means .24 (SD = 1.1) and .28 (SD = 1.2). Point estimates clearly differ from each other, but a t-test indicates that this difference is not significant; \( t(57.9) = 1.6, p = 0.12 \) (Figure 1, left).

Or suppose that point estimates of the factor loadings of an item x1 differ slightly across two factor models estimated in two samples, but measurement invariance tests indicate that the null hypothesis that the models are the same cannot be rejected (Meredith, 1993). Researchers would rely on the results of the statistical tests—not the point estimates—to draw inferences. This also applies to network models: point estimates of two specific edges might be different (Figure 1, right), but whether this difference is meaningful can only be determined in the light of sampling variability, which requires statistical tests to find out if a difference in two parameter estimates is more extreme than one would expect under the null hypothesis. Bootnet and NCT were developed with the purpose to conduct such tests—bootnet for comparing edge weights within one network, NCT for comparing edge weights across different networks.

In their paper, FWMK analyze and visualize differences in point estimates in detail, and show that statistical tests provided by bootnet and NCT arrive at different conclusions than the authors’ inferences of point estimates. For example, they state: “all NCT results indicated that the depression and anxiety symptom networks had no significant differences when in fact they had a multitude of differences” (p. 15, our highlight). This inference is no different than concluding that the t-test for neuroticism in Figure 1 reaches a non-significant result when in fact point estimates of neuroticism differ across samples (Figure 1)—it ignores sampling variability. Of note, FWMK only apply this logic to parameters derived from network models, not to other statistical parameters. For instance, they state that polychoric correlation matrices of the longitudinal data “could be constrained to equality without affecting model fit ([…] p = 0.666)”.

Misinterpretation of bootnet and NCT

Three conclusions of FWMK regarding bootnet and NCT do not follow from their investigation. First, a central claim of FWMK is that “existing suite of methods tended to suggest that the networks were accurately estimated” (p. 12), and that “the interpretation guidelines for bootnet results err toward indicating stability and interpretability in networks” (p. 15). This is not true: bootnet results clearly indicate lack of stable estimates for study 1. The CS coefficient—a metric bootnet provides researchers an idea about the stability of the order of centrality estimates—was 0.13 for the depression and anxiety networks, implying that the centrality order is unstable and should thus not be interpreted (Epskamp et al., 2018). FWMK write: “the CS-coefficient was below the minimum recommended cutoff [for accurate estimation] at both
waves (CS(0.7) = .13), which notably represents the only clear guideline available for interpreting bootnet output” (p. 9). In other words, FWMK 1) identified only one clear benchmark for accurate estimation in bootnet, the CS-coefficient; 2) demonstrate that this one benchmark is below the recommended threshold for accurate estimation; and 3) conclude nonetheless that bootnet guidelines suggest networks were estimated accurately. Thus, their conclusion that bootnet errs toward indicating stability and interpretability does not follow from the evidence FWMK present.

Second, FWMK correctly state in the manuscript that “the authors of the [bootnet] package emphasize that the CIs do not represent significance tests” (p. 3). However, the authors then continue to interpret bootnet results in precisely this way. They consider an edge to be “bootnet-accurate”—a novel concept invented by FWMK that has not been used before in the literature—if its bootstrapped CI does not contain zero, irrespective of how large the bootstrapped CI is. This definition is inconsistent with common definitions of parameter accuracy: a parameter estimate is accurate when it has small confidence regions, regardless of whether these confidence regions contain zero. For example, a standardized parameter that is reliably estimated to be non-zero but fluctuates between 0.1 and 0.8 in bootstrap samples indicates the estimate is not accurate, but falls under the definition of accurate according to FWWK. A related concern is that regularization already pulls parameters to zero. Bootstrapping to test the null-hypothesis that an edge weight is different from zero after using regularization leads to a considerable drop in statistical power to detect if an edge weight is nonzero, and the probability increases that an estimated edge weight is exactly zero, which in turn leads to a loss of statistical power to detect an edge to be nonzero using such a confidence region (i.e. increased type 2 error rate). It is not surprising that when comparing many such low powered tests, many edges will not be considered different from zero in both datasets—this is expected by chance alone. FWMK seem aware of both problems, as is evident from the quote above and footnote 5, but used “bootnet-accurate” to inform their most important conclusions nonetheless.

Third, with regard to the NCT, FWMK state that “The simulations in van Borkulo et al. (under review) indicated that the NCT omnibus test should be adequately powered to compare these networks” (p. 15). This is not correct: the simulation study investigated continuous, independent data, not ordinal, dependent (i.e., longitudinal) data as used by the authors. It is therefore unclear how much power FWMK have to detect differences. In this case, the authors obtain a non-significant p-value, and argue that there is something wrong with the method because it arrives at different conclusions than the comparison of parameter point estimates. We refer back to our discussion above on sampling variation for the logic of this argument, or to measurement invariance in structural equation models (Meredith, 1993). Comparing factor models across two samples is not done by visualizing differences in point estimates of factor loadings, but by using statistical tests to see if parameters differ more than would be expected under sampling variability.

### Problems of polychoric correlations in skewed ordinal data

The results of study 1, including the switching of signs in edges, are partly due to the specific estimation algorithm used: regularized GGM estimation based on polychoric correlations. As described in the tutorial paper cited by FWMK (Epskamp & Fried, 2018), polychoric correlations can lead to biased estimates in case of skewed ordinal data, especially in smaller samples: “When the sample size is relatively low, some cells in the item by item frequency table can be low or even zero […] The estimation of polychoric correlations is […] biased whenever an expected frequency is too small (i.e., below 10; Olsson, 1979). Low frequencies can thus lead to biased polychoric correlations, which can compound to large biases in the estimated partial correlations.

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**Table 1. Replication of depression and anxiety networks based on polychoric and Spearman correlations.**

<table>
<thead>
<tr>
<th>Replication metric proposed by FWMK</th>
<th>GGMs based on polychoric correlations</th>
<th>GGMs based on Spearman correlations</th>
</tr>
</thead>
<tbody>
<tr>
<td>Edges of N1 replicated with same sign in N2</td>
<td>55 (70.5%)</td>
<td>65 (81.3%)</td>
</tr>
<tr>
<td>Edges of N1 that switch sign in N2</td>
<td>4 (5.1%)</td>
<td>0 (0%)</td>
</tr>
<tr>
<td>Absent edges of N1 replicated in N2</td>
<td>23 (54.8%)</td>
<td>28 (70%)</td>
</tr>
<tr>
<td>Bridging edges of N1 replicated with same sign in N2</td>
<td>23 (56.1%)</td>
<td>23 (67.6%)</td>
</tr>
<tr>
<td>Bridging edges of N1 switched sign in N2</td>
<td>3 (7.3%)</td>
<td>0 (0%)</td>
</tr>
<tr>
<td>Absent bridging edges of N1 replicated in N2</td>
<td>14 (32.6%)</td>
<td>22 (75.9%)</td>
</tr>
</tbody>
</table>

Note: Higher replicability marked bold. GGM: Gaussian Graphical Model; N1: network estimated on first time point in depression and anxiety data; N2: network estimated on second time point. Total number of edges per network: N1 (Spearman) = 80; N1 (polychoric) = 78; N2 (Spearman) = 78; N2 (polychoric) = 42.
Another situation in which one might obtain low frequencies is when the scores are highly skewed (Rigdon & Ferguson, 1991), which unfortunately often is the case in psychopathology data. Again, the network based on polychoric correlations should be compared with a network based on Spearman correlations” (p. 14).

Such estimation problems can be expected in the data of the FMWK: for timepoint 1, all 16 items are skewed (mean skewness = 1, range 0.44 to 2.59; mean kurtosis = 0.54, range −0.82 to 6.33), with an average item mean of 0.86 (range 0.29 to 1.31) on a scale from 0 to 4. As expected, such data shows floor effects, with a correlation of 0.86 between item means and standard deviations. Timepoint 2 shows even more pronounced skew and floor effects. Following the recommendation of using Spearman correlations to obtain more stable estimates (Epskamp & Fried, 2018), replicability is consistently higher, on 6 out of 6 metrics calculated by FWMK (Table 1). This highlights that the results from FMWK appear to be specific to the estimation method used rather than generalize to network estimation methods in general (Figure 2).

Figure 2. Networks estimated on timepoints 1 and 2 from the depression and anxiety data used by FMWK. Top row: GGMs estimated by FWMK, based on polychoric correlations. Bottom row: GGMs based on Spearman correlations.
Replicability is a substantive, not a statistical phenomenon

Finally, we want to highlight that conclusions about methodology cannot follow from investigations of data—they require investigations of methodology, which can be done via simulation studies or mathematical proofs (Borsboom et al., 2017, 2018). This means that replications or non-replications of substantive findings cannot be informative about the quality of statistical models. One can fairly criticize researchers for using t-tests incorrectly, but one would not criticize t-tests as a statistical tool because large-scale collaborative projects in psychology that primarily relied on t-tests have shown limited replicability (Camerer et al., 2018; Open Science Collaboration, 2015). This is the conclusion FWMK draw about network models: “Poor replicability underpins our concern surrounding the use of these methods” (p. 1).

Conclusion

We show that the “limited reliability of the detailed characteristics of networks” (p. 1) observed by FWMK occurs in part due to sampling variability, and in part because polychoric correlations can be unstable in skewed ordinal data, especially in small samples.

Network models are high-dimensional, multivariate models with many parameters. Exploratory search in this parameter space will come with sampling variability and differences in the performance across specific algorithms. Prior work has discussed these aspects in some detail, and put forward ways to address sampling variability using statistical tests (Epskamp et al., 2018; Epskamp et al., 2016; Fried & Cramer, 2017; Williams & Rast, 2018; Williams et al., 2019). Recent methodological studies have led to an increased understanding of the performance of specific network estimation methods under different conditions. For instance, the regularization algorithm commonly used was specifically designed for sparse underlying network structures. Under dense structures, regularization leads to a higher false positive rate than was previously known (Williams & Rast, 2018; Williams et al., 2019). We welcome such simulation studies that are important methodological contributions to the literature, as they strengthen the knowledge about methodology. Network psychometrics is no different in this regard than psychometrics in general.

Overall, such methodological insights encourage researchers to use models appropriate for their data. For network models, researchers may, under specific circumstances, consider alternative estimation routines that do not rely on regularization, such as non-regularized estimation procedures (e.g., Isvoranu et al., 2019). But rather than to conclude this is a problem for network psychometrics, we conclude that researchers should be aware of the assumptions inherent in the methods such as regularization, and choose estimation methods that are most appropriate for their data and research question.

Finally, it is unclear if e.g. 75% replicated edges imply comparably good or bad model performance, as we are not aware of simulation studies on the expected number of replicated present and absent edges given conditions such as sample size, skewness of data, and number of variables. That is, even if data for two samples come from the same true model, it is unclear how many edges can be expected to replicate—the same way one would not expect for all factor loadings to replicate across two samples in all possible conditions, even if the data generating mechanism is the same factor model across both. We have implemented the function replicationSimulator in the bootnet package to facilitate research on the topic, which allows researchers to conduct such simulation studies across a range of network estimation methods and compare their performance. Results may differ substantially per estimation method, and the optimal method may depend on the particular needs of the researcher.

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