Temporal Networks of Tobacco Withdrawal Symptoms During Smoking Cessation Treatment

David M. Lydon-Staley1, 2, Adam M. Leventhal3, 4, Megan E. Piper5, Robert A. Schnoll6, 7, and Danielle S. Bassett2, 8
1 Annenberg School for Communication, University of Pennsylvania
2 Department of Bioengineering, School of Engineering and Applied Science, University of Pennsylvania
3 Department of Preventive Medicine, Institute for Addiction Science, University of Southern California Keck School of Medicine
4 Department of Psychology, University of Southern California
5 Department of Medicine, University of Wisconsin Center for Tobacco Research and Intervention, University of Wisconsin School of Medicine and Public Health
6 Department of Psychiatry, Perelman School of Medicine, University of Pennsylvania
7 Abramson Cancer Center, University of Pennsylvania
8 The Santa Fe Institute, Santa Fe, New Mexico, United States

A recently developed network perspective on tobacco withdrawal posits that withdrawal symptoms causally influence one another across time, rather than simply being indicators of a latent syndrome. Evidence supporting a network perspective would shift the focus of tobacco withdrawal research and intervention toward studying and treating individual withdrawal symptoms and intersymptom associations. Here we construct and examine temporal tobacco withdrawal networks that describe the interplay among withdrawal symptoms across time using experience-sampling data from 1,210 participants (58.35% female, 86.24% White) undergoing smoking cessation treatment. We also construct person-specific withdrawal networks and capture individual differences in the extent to which withdrawal symptom networks promote the spread of symptom activity through the network across time using impulse response analysis. Results indicate substantial moment-to-moment associations among withdrawal symptoms, substantial between-person differences in withdrawal network structure, and reductions in the interplay among withdrawal symptoms during combination smoking cessation treatment. Overall, findings suggest the utility of a network perspective and also highlight challenges associated with the network approach stemming from vast between-person differences in symptom networks.

Keywords: networks, withdrawal, nicotine, tobacco, symptoms

General Scientific Summary
This study uses a network approach to examine how symptoms of tobacco withdrawal may impact one another from one moment to the next during a smoking cessation attempt. We find substantial moment-to-moment associations among withdrawal symptoms. Participants undergoing combination treatment have networks in which symptom activity is less likely to linger across time due to moment-to-moment intersymptom associations relative to participants in a placebo condition.

Supplemental materials: https://doi.org/10.1037/abn0000650.supp

This article was published Online First November 30, 2020.
David M. Lydon-Staley https://orcid.org/0000-0001-8702-3923
Robert A. Schnoll https://orcid.org/0000-0002-4089-856X

David M. Lydon-Staley acknowledges support from the National Institute on Drug Abuse (K01-A047417-01A1), Danielle S. Bassett and David M. Lydon-Staley acknowledge support from the John D. and Catherine T. MacArthur Foundation, the Alfred P. Sloan Foundation, the ISI Foundation, the Paul Allen Foundation, the Army Research Laboratory (W911NF-10-2-0002), the Army Research Office (W911NF-14-1-0679, W911NF-16-1-0474, W911NF-17-2-0181), the Office of Naval Research, the National Institute of Mental Health (2-R01-DC-009209-11, R01-MH12847, R01-MH107235, R21-MH106799), the National Institute of Child Health and Human Development (1R01HD086888-01), National Institute of Neurological Disorders and Stroke (R01 NS099348), and the National Science Foundation (BCS-1441502, BCS-1430087, NSF PHY-1554488, and BCS-1631550). Megan E. Piper acknowledges support from the National Institute on Drug Abuse (P50 DA019706), the General Clinical Research Centers Program of the National Center for Research Resources NIH (M01 RR03186), and an Institutional Clinical and Translational Science Award (UW-Madison; KL2 RR025012). Adam M. Leventhal acknowledges support from Grant K24DA048160 and RSG-13-163-01 and the American Cancer Society. Robert A. Schnoll acknowledges support from Grant K24 DA045244. The content is solely the responsibility of the authors and does not necessarily represent the official views of any of the funding agencies. A pre-peer reviewed version of this article is available on PsyArXiv 10.31234/osf.io/2a9q. Collection of the data used in this study was approved by the University of Wisconsin Health Sciences Institutional Review Board, study title A Randomized Placebo-Controlled Clinical Trial of 5 Smoking Cessation Pharmacotherapies. Correspondence concerning this article should be addressed to David M. Lydon-Staley, Annenberg School for Communication, University of Pennsylvania, 3620 Walnut Street, Philadelphia, PA 19104, United States. Email: david.lydonstaley@asc.upenn.edu

Journal of Abnormal Psychology
2021, Vol. 130, No. 1, 89–101
https://doi.org/10.1037/abn0000650
During efforts to quit smoking, aversive withdrawal symptoms appear that are prominent as primary motives for the resumption of smoking in many theories of addiction (Baker et al., 2004; Piper, 2015; Solomon & Corbit, 1974). Withdrawal symptoms are often treated as interchangeable indicators of an underlying syndrome (Toll et al., 2007). During diagnosis (American Psychiatric Association, 2013), for example, criteria for tobacco withdrawal is met if a patient presents with four or more withdrawal symptoms (anger, anxiety, depressed mood, difficulty concentrating, increased appetite, insomnia, and restlessness). Yet, there is increasing evidence that individual withdrawal symptoms are not interchangeable; rather, they exhibit distinct time profiles during the course of cessation attempts (Hendricks et al., 2006; Leventhal et al., 2010; Piasecki et al., 2000; West et al., 1987) and unique responses to treatment (Foulds et al., 2013).

A network approach to tobacco withdrawal has recently been proposed to complement a syndrome perspective on withdrawal (Lydon-Staley et al., 2020). A network theory of psychopathology proposed to complement a syndrome perspective on withdrawal symptoms (Borsboom, 2017; Bringmann et al., 2013; Schmitt-highlights symptoms of disorders as units that causally impact one another across time, forming networks of causally connected symptoms (Borsboom, 2017; Bringmann et al., 2013; Schmittmann et al., 2013). From this perspective, the experience of tobacco withdrawal can be described as a network in which the nodes of the network represent symptoms and the edges (or lines between nodes) represent temporal associations among symptoms across time. Studies provide preliminary support for the network perspective’s proposal that individual tobacco withdrawal symptoms influence one another in a potentially causal fashion over time. For example, positive associations between anhedonia, negative affect, and craving have been observed over the course of a smoking cessation attempt (Cook et al., 2017) and strong positive associations have been observed between sleep problems and restlessness, as well as among affective symptoms (anger, anxiety, and depressed mood), in smokers undergoing smoking cessation treatment (Lydon-Staley et al., 2020). Recent work making use of experience sampling protocols has considered the dynamic associations among cessation fatigue, negative affect, nicotine craving, and self-efficacy in smokers undergoing cessation treatment (Bekiroglu et al., 2017). This work, which makes use of the temporal precedence available in time series data to model how the reporting of symptoms at a previous time point (t-1) predicts symptom intensity at the current time point (t), found that these four constructs changed over time in response to each other during the course of smoking cessation.

Although there is empirical support for the notion that individual symptoms of withdrawal influence one another over time, key questions about a network approach to withdrawal remain to be addressed. We use experience-sampling data from participants providing momentary reports of withdrawal symptoms over 14 days to estimate a temporal network that describes the moment-to-moment interplay among withdrawal symptoms. The use of experience-sampling data to construct temporal networks (Holme & Saramäki, 2012) is an important complement to existing studies modeling cross-sectional withdrawal networks (Lydon-Staley et al., 2020). The edges in cross-sectional networks indicate between-person associations among symptoms, with positive edges indicating that if, for example, an individual experiences high levels of anxiety, then they are also likely to experience high levels of anger. This symptom co-occurrence, from a network perspective, is theorized to result from causal associations among symptoms. Yet, between-person associations (i.e., people high in anxiety experience high levels of anger) do not always translate to within-person associations (i.e., at times when a person is more anxious than usual, they are also angrier than usual; Bos et al., 2017; Hamaker et al., 2005). One of the strengths of experience-sampling data is that it can probe time-lagged, within-person associations between symptoms.

Further, the use of temporal networks allows a test for the presence of a potential source of distress experienced during smoking cessation, unique to a network approach to withdrawal, that may undermine efforts to remain abstinent. In particular, individuals undergoing cessation may experience continued symptom activation long after the cause of symptom activity (i.e., nicotine deprivation) has disappeared due to symptom interplay across time (Borsboom, 2017; Cramer et al., 2016). Network theory posits that this failure for symptoms to deactivate arises in strongly connected symptom networks that facilitate the reverberation of symptom activity through the network across time as activity in one symptom is transferred to another symptom via causal associations. This spreading of symptom activity through networks is analogous to the behavior of two sets of domino tiles: One in which the tiles are far apart and a second in which the tiles are close together. In a densely connected symptom network, the set of domino tiles are close together, and when one tile falls it causes other tiles to topple and activity ripples through the system (Borsboom & Cramer, 2013). In a less densely connected symptom network, the set of domino tiles are far apart, and when one tile falls the other tiles are unaffected and remain standing. Such an explanation of spreading activity following symptom network perturbation would fit with empirical evidence of extended withdrawal symptom experiences among some smokers (Piasecki et al., 1998, 2003) when other research suggests that withdrawal should peak after the first 14 days (Hughes, 2007).

As yet, the phenomenon of self-perpetuating symptom networks in the context of tobacco withdrawal awaits empirical examination. Complicating this examination is the need to identify an appropriate statistical framework that can capture the complex theoretical notion of self-perpetuating symptom networks. Existing efforts to capture the reverberation of symptom activity through symptom networks have focused on node centrality and network density. Centrality indices have been hypothesized to identify the most important nodes in a network. Degree centrality, for example, is a common measure of node centrality and is defined as the number of edges emanating from a node (Newman, 2010). Network density is defined as a fraction of possible edges that exist in a network or, in networks with weighted edges, network strength. Both node centrality and network density have been identified as potential indices for identifying networks that would facilitate the spread of activity through the network, activating self-perpetuating sequences of symptom activity across time (Bello et al., 2017; Fried et al., 2016; Hasmi et al., 2017; Lydon-Staley, Xia, et al., 2019; Pe et al., 2015). Yet, the use of both centrality and density are limited. Centrality and density focus on direct connections and contain little information about how symptoms might affect each other indirectly as activity flows through the entire network (Bringmann et al., 2019).

Impulse response analysis is a tool with the potential to capture the complex notion of the spread of symptom activity through a
Once a temporal network indicating the predictive associations between symptoms across time has been constructed, impulse response analysis simulates an instantaneous, exogenous impulse (sometimes termed a shock) to certain network symptoms. Through simulation, the propagation of this impulse through the network, along the time-lagged symptom network edges, is charted. The impulse response function shows the hypothetical change in a symptom in response to a simulated increase in one of the other symptoms over a horizon of several time points. Importantly, activity in a symptom observed after the symptom network receives a shock contains system-level information. The impulse experienced by nonshocked symptoms is due to the flow of activity along the network’s edges emanating from the shocked node.

Impulse response analysis has recently been applied to networks of symptoms and behaviors to capture individual differences in the spread of activity through networks of time-lagged edges. In a sample of subclinically depressed individuals, impulse response analysis was used to quantify the extent to which stress impacted activity in addicted symptoms relative to participants without addicted symptoms (Bos et al., 2018). A shock to stress was simulated and the spread of activity associated with this simulated increase in stress along time-lagged edges in a network of six variables (high-arousal positive effect, low-arousal positive affect, high-arousal positive affect, low-arousal negative affect, stress, and physical activity) was charted over a horizon of 10 time points. When an increase in stress was simulated, affect in non-anhedonic individuals showed a stronger increase relative to affect in anhedonic individuals. By using impulse response analysis, the observed association between stress and affect consisted of the effect of a simulated increase in stress filtering through direct pathways to impact affect, but also indirect pathways emerging from the complex interplay between stress and affect with other variables in the network, (e.g., physical activity) across time.

In addition to capturing the spread of symptom activity through networks, impulse response analysis has been used to capture individual differences in the extent to which networks facilitate the persistence of activity through complex systems of temporally associated components, long after the initiating shock has ended. In 13 variable, person-specific, temporal networks indicating the moment-to-moment associations between a range of socioemotional processes (e.g., anger, sadness, ratings of interpersonal behaviors and perceptions), impulse response analysis was used to simulate a shock to sadness (Yang et al., 2018). Changes in sadness intensity was charted over 150 time steps and the time step at which sadness intensity returned a level consistent with its pre-shock level was identified. This time step was taken as a measure of the recovery time of sadness following a perturbation and, importantly, consisted of activity related to the exogenous shock, the autoregressive effect of sadness on itself from one timepoint to the next, but also indirect effects on sadness as activity spread to other nodes and returned via time-lagged edges to impact sadness. Participants with networks associated with longer recovery times of sadness exhibited higher levels of depressive symptoms, especially during periods of higher than usual stressful life events.

Impulse response analysis, then, is a framework capable of capturing the potential for symptom activity to spread through a network via time-lagged edges and for quantifying the time it takes for symptoms to return to baseline levels of intensity following hypothetical perturbations. Here, we extend this work to tobacco withdrawal. We estimate temporal networks of tobacco withdrawal, articulating the moment-to-moment associations between withdrawal symptoms across time. After constructing temporal networks, we quantify individual differences in networks by using impulse response analysis to simulate a shock to network symptoms and chart the spread of symptom activity along time-lagged edges through each network. To quantify the extent to which symptom networks facilitate persistent symptom activity, we identify the number of timesteps after the initial shock at which symptom activity returns to a baseline level of intensity comparable to the intensity of activity preceding the shock to the symptom network. We test whether participants undergoing smoking cessation treatment show a lower potential for symptom activity spread and reverberation through symptom networks across time relative to participants in a placebo condition. We hypothesize that treatment is associated with a quicker return to symptom intensity baseline (i.e., shorter recovery time) following a simulated shock.

**Method**

**Participants**

Participants were 1,210 smokers (58.35% female, 86.24% White), previously reported on in an existing study (Bolt et al., 2012; Piper et al., 2009). Participants were recruited via TV, radio, and newspaper advertisements, community flyers, and earned media (e.g., radio and TV interviews, press releases) in the greater Madison and Milwaukee (Wisconsin) areas. The primary inclusion criteria were smoking at least 10 cigarettes per day for the past 6 months and being motivated to quit smoking. Exclusion criteria included: use of certain medications (including MAO inhibitors, bupropion); any history of psychosis, bipolar disorder, or an eating disorder; consuming six or more alcohol beverages daily 6 or 7 days a week; pregnancy or breast-feeding; and a serious health condition that might prevent study completion. The study was approved by the University of Wisconsin Health Sciences Institutional Review Board, study title A Randomized Placebo-Controlled Clinical Trial of 5 Smoking Cessation Pharmacotherapies.

**Procedure**

Participants passing a phone screen were invited to an information session at which a study description was provided and written informed consent was obtained. Participants then completed multiple baseline screenings, including a medical history screening, vital signs measurements, and a carbon monoxide (CO) breath test. Participants also completed demographic, smoking history, and tobacco dependence questionnaires.

Eligible participants were randomized to one of six treatment conditions: bupropion sustained-release (SR; n = 215), nicotine lozenge (n = 202), nicotine patch (n = 210), nicotine patch + nicotine lozenge (n = 228), bupropion SR + nicotine lozenge (n = 214), or placebo (n = 141). Five placebo conditions matched the five active conditions, with three monotherapy placebo conditions (n = 80) and two combination therapy placebo conditions (n = 61). All medications were provided for 8 weeks postquit except the nicotine lozenge, which was provided for 12 weeks.
postquit, consistent with prescribing instructions at the time (Fiore et al., 2008).

**Measures**

Participants completed ecological momentary assessments (EMA) four times a day (once just after waking, once prior to going to bed, and twice at randomly chosen times between waking and sleeping) for 1 week prequit and 2 weeks postquit. We analyzed data from the 2-week postquit period. The EMA prompted participants to rate their withdrawal symptoms within the last 15 min using 11 items from the Wisconsin Smoking Withdrawal Scale (Welsch et al., 1999) and one item adapted from the Questionnaire of Smoking Urges (Sweeney et al., 1996) to assess six symptoms: anxiety (tense or anxious, impatient), craving (bored by desire to smoke, urge to smoke), sadness (sad or depressed, hopeless or discouraged), irritability (irritable or easily angered, bothered by negative moods such as anger, frustration, and irritability), hunger (hungry, thinking about food a lot), and difficulty concentrating (hard to pay attention, difficult to think clearly). The mean of the two items making up each symptom scale was computed. Participants also reported the number of cigarettes smoked since the last report at each prompt. Prior to the EMA protocol, participants completed the Fagerström Test for Nicotine Dependence (Heatherton et al., 1991). CO confirmed 7-day point-prevalence abstinence data was collected at 8 weeks and 6 months postquit. Alveolar CO was assessed using a Bedfont Smokerlyzer and smokers with a CO <10ppm were considered abstinent.

**Data Preparation**

Temporal network models assume stationarity of symptoms such that symptoms do not exhibit a trend in the mean or variance of their intensity across time (Lütkepohl, 2005). We removed linear trends from each symptom by regressing each participants’ ratings of symptom intensity on time in the study to remove the linear trend, and then by taking the residuals forward, which is a common practice to handle violations of stationarity in the means. A second assumption of the models used is that there are equal time lags between consecutive measurements (de Haan-Rietdijk et al., 2017). To accommodate this assumption, we did not regress the first measurement of each day on the last measurement of the previous day.

**Data Analysis**

We used the R package mlVAR (Epskamp et al., 2018) to estimate the dynamic associations among withdrawal symptoms at the group level (Epskamp et al., 2016). Each symptom was regressed on the values of all other symptoms at the previous time point. To accommodate the nested nature of the data, with participants providing multiple reports, a multilevel extension of VAR was used (Bringmann et al., 2013; Lydon-Staley, Xia, et al., 2019). The resulting coefficients represent the degree to which changes in one symptom at time \( t \) predict changes in another symptom at time \( t + 1 \) for the average individual in the sample. Importantly, data are person-mean centered by default, allowing a consideration of within-person associations between variables that are disentangled from between-person differences in levels of symptom intensity (Curran & Bauer, 2011). From this within-person perspective, edges may be interpreted as how deviations in symptom intensity from one’s average level of symptom intensity is associated with change in another symptom’s intensity. In addition to producing a temporal network, the mlVAR approach also estimates a contemporaneous network. Edges in the temporal network encode information regarding how symptoms at one measurement occasion predict symptoms at the next measurement occasion. This network has the potential to provide information about the directionality of effects (Epskamp et al., 2018), and thus estimates of influence are depicted as directed edges with arrows indicating the direction of effects across time. Edges in the contemporaneous network show associations between symptoms occurring within consecutive measurements and indicate partial correlations between nodes at the same time, controlling for both temporal effects and all other variables within the same measurement occasion. Edges in the contemporaneous network are computed by correlating the residuals of the temporal effects (Dahluh & Eichler, 2003). The contemporaneous network provides insight into the co-occurrence of symptoms, and may also indicate the presence of causal associations occurring on a shorter timescale than the timescale of measurement (Epskamp et al., 2018). One of the advantages of multilevel models is their tolerance to heterogeneity of time points across participants. As such, data from all participants were included in the model.

After estimating a group level dynamic withdrawal network, we used the same participants and measures to estimate person-specific dynamic networks of withdrawal. The raw time series data for the six withdrawal symptoms were regressed on time in order to remove trends and meet the assumption of stationarity. These data were then within-person standardized so that each symptom for each person had a mean of 0 and a standard deviation of 1. This standardization was performed to render the coefficients representing different edges in the network comparable with one another (Bulter et al., 2016). Within-person standardization also allows us to focus on within-person associations among the various symptoms when comparing networks between participants. The edges in the network represent how fluctuations in one symptom beyond a participant’s typical level (captured by 0 due to the within-person standardization) are associated with fluctuations in a different symptom (Curran & Bauer, 2011). We then modeled each participant’s six-symptom time series as a 6-node network using a unified structural equation model (uSEM; Gates et al., 2010).

In this approach, the observed multivariate time series \( \gamma(t) \) is modeled as the output of a latent variable time series \( \eta(t) \),

\[
\gamma(t) = \Lambda \eta(t) + \varepsilon(t),
\]

where \( \Lambda \) is a factor loading matrix and \( \varepsilon(t) \) is a time series of residuals given by a matrix \( \Theta \), that is assumed diagonal. We then model the temporal associations among the set of latent withdrawal symptoms represented in \( \eta(t) \) as,

\[
\eta(t) = A \eta(t) + \Phi \eta(t - 1) + \zeta(t),
\]

where \( \eta(t - 1) \) is a vector of the lag-1 version of the multivariate latent withdrawal symptom time series, \( A \) is a matrix of regression parameters describing the contemporaneous associations among the latent withdrawal symptom variables, \( \Phi \) is a matrix of regression parameters describing the lag-1 associations (auto- and cross-
regressions) among the latent withdrawal symptom variables, and \( \zeta(t) \) contains the errors in prediction. Both the contemporaneous associations in \( A \) and the autoregressive and cross-regressive associations in \( \Phi_1 \) indicate directed temporal associations among the withdrawal symptoms through which exogenous input can diffuse. Note that although the contemporaneous networks estimated in mlVAR are undirected, in the uSEM approach, directed edges are estimated in the contemporaneous networks (e.g., Wright et al., 2019; Yang et al., 2018; Yang et al., 2019). Directed contemporaneous edges in structural equation models provide insight into the co-occurrence of symptoms and may also indicate the presence of causal associations occurring on a faster timescale than the timescale of symptoms and may also indicate the presence of causal associations occurring on a faster timescale than the timescale of symptoms (Granger, 1969), especially when autoregressive influences are simultaneously estimated (Gates et al., 2010), as in the current case.

The uSEM model is estimated using an iterative search process. A series of models is constructed and improvements to model fit are determined. With each iteration, Lagrange Multiplier tests (Sörbom, 1989) are used to select the edge which, if free to be estimated, would have the maximum improvement on model fit. We constrained the iteration process such that only \( A \) and \( \Phi_1 \) blocks of the parameter matrix may be freed. This choice serves to maintain the time-series structure of the model. We avoided bidirectional edges in the contemporaneous associations by including all potential autoregressive associations in the initial model and deeming the reverse association between two nodes unavailable when any edge in \( A \) was freed (see also Yang et al., 2018). By specifying unidirectional edges in the contemporaneous network, only one causal edge between two variables is allowed. This restriction still allows reciprocal edges in withdrawal networks given that the reverse association can be observed in the lagged network. We configured the factor loading matrix \( A \) as an identity matrix \( I \) and we fixed all elements of \( \Theta \), the variance-covariance matrix of \( \epsilon_1 \) and \( \epsilon_n \), to 0. Model fit is determined according to a predetermined \( \alpha \) level on a \( \chi^2 \)-square distribution with one degree of freedom. Here, as elsewhere (Beltz et al., 2013), we set \( \alpha \) to 0.05. This edge is freed, the model is reestimated, and a new set of modification indices are calculated. Edges are freed or added until further addition does not significantly improve model fit.

To quantify individual differences in symptom activity spread across symptom network edges, we used impulse response analysis (Lütkepohl, 2005) to provide insight into how the experience of one symptom propagates through the withdrawal symptom network by predicting other, connected symptoms. In this approach, an impulse is given to an individual node and the behavior of the symptom network is observed, through simulation, over many time steps. This dynamical process simulates how a symptom’s activity moves through the symptom network along edges when it becomes transiently increased due to an external perturbation. Formally, the impulse response simulation is derived from Equation 2 as a one step ahead forecasting process through conversion into a vector autoregression model (Amisano & Giannini, 2012). The system is set in motion with an initial impulse vector, \( \zeta(0) \), and latent states are calculated for \( t = 0 \) as:

\[
\eta(0) = (I - A)^{-1}\zeta(0),
\]

(3)

to accommodate the contemporaneous associations among latent states and for each subsequent \( t \) as:

\[
\eta(t) = [(I - A)^{-1}\Phi_1]\eta(t - 1) = [(I - A)^{-1}\Phi_1][I - A)^{-1}\zeta(0),
\]

(4)

and the system evolution over 100 time steps is simulated. The time profile, showing how symptom intensity changes over the time steps following the initial impulse, is examined. Specifically, the recovery time of symptom intensity, defined as the time taken to return to within \(+1/0.01\) of equilibrium, is derived via a backward search. We searched backward from the end of the time profile to identify the time step \( k \) where the intensity of a symptom is first outside the \(+1/0.01\) boundary. Recovery time is then quantified as the number of time steps from perturbation to equilibrium. Implementation of person-specific uSEM and impulse response analysis was done using the R package pompom (Yang et al., 2018).

We next quantified individual differences in the extent to which withdrawal symptom networks facilitate self-perpetuating symptom activity across time by estimating system recovery time. Impulses were given to all six symptoms, each time tracking changes in all six symptoms. The result is a \( 6 \times 6 \) matrix of symptom activity trajectories. We took the mean value of the column means in the \( 6 \times 6 \) matrix as an indication of system recovery time, indicating the average time taken for a symptom to return to equilibrium after a hypothetical perturbation.

To explore the extent to which system recovery time may be impacted by treatment, we compared the system recovery time of participants receiving combination therapy during the quit attempt (nicotine patch + nicotine lozenge and buproprion SR + nicotine lozenge conditions) to participants in combination therapy placebo conditions using independent samples \( t \) tests. A second independent samples \( t \) test was used to determine whether system recovery time differed between participants in the monotherapy conditions (buproprion SR, nicotine lozenge, and nicotine patch) and the associated placebo conditions.

Results

We provide descriptive statistics of the study sample before presenting the results for the group and person-specific withdrawal networks.

Descriptive Statistics

The average intensity of withdrawal symptoms across the EMA period are shown in Table 1. Smoking lapses were common with only 400 participants (33.06%) reporting not smoking since the previous EMA report. On average, smoking since the previous prompt was reported at 18.00% of prompts (\( SD = 27.01 \)). The mean number of EMA reports available per participant was 27.91 (\( SD = 21.51 \)). Participants completing a greater number of EMA reports exhibited lower intensity of withdrawal symptoms (all \( rs < 0.20 \) indicating weak associations) during the EMA protocol (see Table 1).

Group-Level Withdrawal Networks

Results from the multilevel vector autoregressive model are in Table S1. The edges of the temporal network (Figure 1A) represent the prediction of symptoms at one measurement occasion by
symptoms at the previous measurement occasion for the prototypical individual in the sample. We find that all estimated edges are positive, such that withdrawal symptoms at one moment predict increases (rather than decreases) in the experience of other withdrawal symptoms (including themselves) at the next moment. Craving has many outgoing edges, indicating that increased craving at one moment has a relatively strong prediction of other withdrawal symptoms from moment to moment. Both sadness and irritability have the largest number of ingoing edges, indicating that increases in sadness and irritability are predicted by increases in the experience of other withdrawal symptoms. The majority of symptoms show positive self-loops (edges pointing toward themselves), which indicate that current symptom experiences are predictive of subsequent experiences. Notable is the presence of reciprocal edges between sadness and irritability, and between anxiety and irritability. These reciprocal edges represent positive feedback loops whereby the experience of one symptom (e.g., irritability) predicts increases in the experience of another symptom (e.g., anxiety), which in turn predicts the experience of the original symptom (i.e., irritability). Also notable is hunger, which is disconnected from the other withdrawal symptoms; the lack of incoming or outgoing edges to and from hunger indicates that fluctuations in the experience of hunger over time do not play a large role in predicting other withdrawal symptoms.

The edges of the contemporaneous network (Figure 1B) represent the co-occurrence of symptoms, controlling for both temporal effects and the effects of all other variables at the same measurement occasion. Edges are unidirectional (without arrows) but they may indicate causal associations playing out at timescales shorter than the timescale of symptom measurement (Epskamp et al., 2018). Similar to the temporal network, all edges are positive, indicating that when an increase in one symptom is experienced

### Table 1

**Descriptive Statistics**

<table>
<thead>
<tr>
<th>Variables</th>
<th>1</th>
<th>2</th>
<th>3</th>
<th>4</th>
<th>5</th>
<th>6</th>
<th>7</th>
<th>8</th>
</tr>
</thead>
<tbody>
<tr>
<td>1. Anxietya</td>
<td>—</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>2. Irritabilitya</td>
<td>0.85</td>
<td>—</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>3. Difficulty concentratinga</td>
<td>0.75</td>
<td>0.72</td>
<td>—</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>4. Cravinga</td>
<td>0.52</td>
<td>0.47</td>
<td>0.41</td>
<td>—</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>5. Sadnessa</td>
<td>0.69</td>
<td>0.73</td>
<td>0.65</td>
<td>0.35</td>
<td>—</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>6. Hungera</td>
<td>0.44</td>
<td>0.41</td>
<td>0.40</td>
<td>0.34</td>
<td>0.32</td>
<td>—</td>
<td></td>
<td></td>
</tr>
<tr>
<td>7. Smokingb</td>
<td>0.14</td>
<td>0.17</td>
<td>0.14</td>
<td>0.30</td>
<td>0.24</td>
<td>0.05</td>
<td>—</td>
<td>—</td>
</tr>
<tr>
<td>8. EMA completion rate</td>
<td>−0.17</td>
<td>−0.16</td>
<td>−0.14</td>
<td>−0.15</td>
<td>−0.14</td>
<td>−0.08</td>
<td>−0.25</td>
<td>—</td>
</tr>
</tbody>
</table>

**Variables**

| M     | 1.90  | 1.57  | 1.37  | 4.38  | 1.06  | 2.57  | 0.18  | 27.91 |
| SD    | 1.69  | 1.60  | 1.62  | 2.55  | 1.60  | 1.40  | 0.27  | 12.51 |

**Note.** EMA = ecological momentary assessments.

a Variables represent the intraindividual mean of participant’s repeated measures; n = 1,210. All correlations greater than 0.05 are significant at p < .05. b Indicates the proportion of prompts at which smoking since the previous prompt was reported.

Figure 1

**Dynamic Networks of Tobacco Withdrawal**

**A**

- Hunger
- Sadness
- Craving
- Difficulty Concentrating
- Irritable
- Anxious

**B**

- Hunger
- Sadness
- Craving
- Difficulty Concentrating
- Irritable
- Anxious

**Note.** (A) A temporal network in which the nodes (circles) represent symptoms and the edges represent the association between symptoms from one measurement occasion to the next. (B) A contemporaneous network in which the edges represent the co-occurrence of symptoms at the same measurement occasion while controlling for temporal effects and the effects of other symptoms. Only green edges are observed in the networks indicating that the experience of increases in withdrawal symptoms tend to be associated with increases in the experience of other symptoms (e.g., when sadness is higher than usual, anxiety is higher than usual). Both temporal and contemporaneous networks are estimated by pooling data across all participants in the sample and, thus, edges represent the prototypical associations among symptoms.
(e.g., sadness) an increase in a second symptom is experienced (e.g., irritability).

**Person-Specific Withdrawal Networks**

We next moved from a group model indicating the temporal associations between symptoms across time for the prototypical individual in the sample to the estimation of person-specific temporal networks. The estimation of person-specific networks requires substantially more data than the estimation of a group model that pools data across individuals. A person-specific model of the associations among the six withdrawal symptoms fit the data of 290 individual participants well; the goodness of fit is indicated by at least three of four following criteria: RMSEA $\leq 0.08$, SRMRs $\leq 0.08$, CFIs $\geq 0.95$, NNFI $\geq 0.95$ (Beltz et al., 2013; Yang et al., 2018). Comparing the number of days available for participants for whom a person-specific model fit well relative to those for whom a person-specific model could not be fit well indicate that, in line with previous work (Yang et al., 2018), the length of the time series impacted model fit, with a significantly greater amount of time series data available for participants with good ($M = 37.21, n = 290$) versus poor ($M = 23.16, n = 998$) model fit, $t(745.72) = 21.692, p < .001, d = 1.14$.

Substantial heterogeneity in symptom network structure is evident (Figure 2A; see Figures S1–S7 for distributions of edge estimates). We quantify individual differences in participants’ withdrawal symptom networks and their implications for the extent to which symptom activity persists across time due to the predictive associations among network symptoms by using impulse response analysis (Lütkepohl, 2005; Yang et al., 2018). The average system recovery time (Figure 2B) for the sample is 3.18 ($SD = 2.55$) time steps and exhibits substantial positive skew ($min = 0.89, max = 26.86, skew = 4.23$). We used the common

**Figure 2**

**Person-Specific Networks of Tobacco Withdrawal**

Note. (A) Two graphs estimated using a unified structural equation model on experience-sampling data show the temporal associations (edges) among individual withdrawal symptoms (nodes) across time for two different participants. Red edges show negative associations; green edges show positive associations; dashed edges are lagged associations; and continuous edges are contemporaneous associations. Notable is the heterogeneity in network structure across the two participants. (B) A sample matrix of recovery times associated with the network on the left in Panel A, estimated using impulse response analysis, indicates the time it takes for a symptom to return to equilibrium after being perturbed. For example, when an increase in anxiety is simulated, it takes 4.92 time steps for anxiety to return to baseline (as indicated by the Anx. $\rightarrow$ Anx label). A subsample of three symptoms (anxiety, irritability, and sadness) of the full six symptom matrix is shown to enhance readability. (C) Independent samples $t$ tests show higher log system recovery times in the symptom networks of participants in the combination therapy placebo condition (nonlog mean value = 3.55) relative to participants in the active combination therapy condition (nonlog mean value = 3.05). No differences in system recovery time emerges between participants in the active (nonlog mean value = 2.88) and placebo monotherapy (nonlog mean value = 2.39) groups. Anx. = anxiety; Irr. = irritability; Sad. = sadness; NS = nonsignificant. $^p = .03$. 

This document is copyrighted by the American Psychological Association or one of its allied publishers. This article is intended solely for the personal use of the individual user and is not to be disseminated broadly.
This document is copyrighted by the American Psychological Association or one of its allied publishers. This article is intended solely for the personal use of individual users and is not to be disseminated broadly.

logarithm to reduce the positive skew and remove three outliers (all greater than 3 SDs above the mean). In the sample of 287 participants, there were no differences across the treatment conditions in the average number of EMA data points available, in self-reported cigarettes smoked per day reported at baseline, in FTND total score reported at baseline, or in the average intensity of the different withdrawal symptoms across the EMA period (all \( p \) values > 0.05). There was a difference in the proportion of prompts during the EMA period for which participants across the different treatment groups reported smoking since the previous report such that participants in the monotherapy placebo group reported the most smoking occasions (\( M = 0.33, SD = 0.34 \)), followed by the combination placebo group (\( M = 0.14, SD = 0.19 \)), then the monotherapy group (\( M = 0.13, SD = 0.22 \)), and finally the combination therapy group (\( M = 0.10, SD = 0.17 \)). Only the difference between the combination therapy and the monotherapy placebo group, however, was statistically significant, \( t(12.45) = -2.43, p = .03, d = 1.25 \).

Using independent samples \( t \) tests we found that participants undergoing combination treatment have faster recovery times relative to participants in a matched placebo condition, \( t(26.76) = -2.23, p = .03, d = 0.43 \), a small effect size (Cohen, 1988; Figure 2C). There was no significant difference between participants in monotherapy treatment and a matched placebo condition, \( r(21.04) = 1.00, p = .33, d = 0.24 \). Beyond these planned contrasts, we also observed that participants in the monotherapy condition had shorter recovery time relative to participants in the combination placebo group, \( r(34.60) = -2.29, p = .03, d = 0.52 \), and that participants in the monotherapy placebo group had shorter recovery time relative to participants in the combination placebo group, \( r(28.67) = -2.69, p = .01, d = 0.92 \). There was no significant difference in recovery time between the combination therapy and monotherapy groups, \( r(155.39) = 0.34, p = .74, d = 0.04 \) or between the combination therapy and monotherapy placebo group, \( r(16.40) = 1.29, p = .21, d = 0.26 \).

Average system recovery time was not associated with CO-verified smoking abstinence at end of treatment (8 weeks postquit; \( r(277.31) = 1.05, p = .30, d = 0.12 \) or at 6 months postquit, \( r(228.19) = 0.87, p = .38, d = 0.11 \). Correlations revealed that participants with longer system recovery time experience more intense anxiety, \( r(285) = 0.12, p = .04 \), sadness, \( r(285) = 0.12, p = .04 \), irritability, \( r(285) = 0.14, p = .03 \), and difficulty concentrating, \( r(285) = 0.15, p = .01 \), but did not report more intense craving, \( r(285) = 0.05, p = .37 \), or hunger, \( r(285) = -0.01, p = .83 \), during the EMA protocol.

**Discussion**

Smoking cessation is notoriously difficult, with the vast majority of quit attempts ending in relapse (Alterman et al., 2001; Fiore et al., 2000). Withdrawal symptoms emerge following reductions in smoking that are associated with decreased quit intentions as well as increased smoking relapse (Allen et al., 2008; Orleans et al., 1991; West et al., 1987). The present study examined the potential utility of a network perspective on tobacco withdrawal, particularly in providing insight into smokers’ withdrawal experiences.

A key contribution of the present article was to extend cross-sectional network findings of associations among symptoms of tobacco withdrawal (Lydon-Staley et al., 2020) to model the interplay among withdrawal symptoms using experience-sampling data containing temporal information on the moment-to-moment activity of withdrawal symptoms. The edges in cross-sectional networks indicate between-person associations, with strong edges indicating that if an individual experiences high levels of anxiety, for example, they are likely to also experience high levels of anger. This symptom co-occurrence, from a network perspective, is theorized to result from causal associations among symptoms. Yet, between-person associations (i.e., people high in anxiety experience high anger) do not always translate to the within-person associations (i.e., at times when a person is more anxious than usual, they are also angrier than usual; Bos et al., 2017; Hamaker et al., 2005).

To address this open challenge in the field, we estimated networks in which the edges indicated the within-person association between symptom activity at one time point and symptom activity at the next time point. The group-level networks of withdrawal exhibited substantial interplay among symptoms from moment to moment. Comparison of temporal networks to cross-sectional withdrawal networks estimated in previous work (Lydon-Staley et al., 2020) is difficult due to the minimal overlap between symptoms examined across studies. However, the positive associations among anxiety, sadness, and difficulty concentrating observed in the group-level contemporaneous network in the current study were also observed in cross-sectional withdrawal networks that also included these three constructs (Lydon-Staley et al., 2020), suggesting that some findings from cross-sectional networks may translate to within-person associations. However, no associations were observed between sadness and anxiety and difficulty concentrating and anxiety in the present study in the group-level temporal network wherein edges represented the extent to which symptom intensity at the current timepoint predicted symptom intensity at the next timepoint. Thus, although some edges present in cross-sectional networks may be observed in temporal networks, the limited overlap suggests that, like previous work in the context of depression (Bos et al., 2017), cross-sectional networks do not reflect how symptoms are associated with one another across time. This limited overlap between cross-sectional and temporal networks limits the extent to which cross-sectional networks, which require data that are more easily obtained than repeated measures data, provide information relevant for clinical applications aimed at identifying potentially causal associations between symptoms that exacerbate illness (Fisher & Boswell, 2016; Kroeze et al., 2017).

Additional considerations raised by examining the group-level networks are the insights gained from the group-level temporal network. The temporal network contained many positive edges, such that if symptom activity was high at one time point, it was more likely to increase, rather than decrease, in activity at the next time point. Indeed, the group temporal network contained numerous positive potential positive feedback loops that might render symptom activity vulnerable to exhibiting self-perpetuation of symptom activity due to the reverberation of symptom activity through the network. For example, anxiety at one time point was associated with increased irritability and anger at the next time point and, in turn, irritability and anger at one time point was associated with increased anxiety at the next time point. The capacity for one’s current emotions to give rise to other emotions has long been
emphasized in emotion theory (e.g., Gross & Muñoz, 1995) and there is empirical support for the moment-to-moment transfer of emotions across time and states (Anand et al., 2017; Lydon-Staley, Xia, et al., 2019; Pe & Kuppens, 2012).

Additional findings of interest include the many out-going edges from craving to other withdrawal symptoms. Although the association between negative affect and craving is prominent in models of withdrawal, the greatest emphasis is placed on the effect of affect to craving rather than the reverse direction (Baker et al., 2004; Conklin & Perkins, 2005; Delfino et al., 2001; Otto et al., 2007). The current findings suggest that the experience of craving at one moment predicts the emergence of aversive states (e.g., anxiety, irritability) at the next moment. Thus, craving may be a useful treatment target, not simply to avoid drug use in response to craving, but also to ameliorate aversive states resulting from its experience (Tiffany & Wray, 2012). We also note that the current findings do not preclude the possibility that negative affect predicts craving given the edges between affect and craving in the contemporaneous group network.

Estimation of person-specific networks revealed that the group model masks substantial individual differences. The most striking difference includes observing negative edges in the networks of individual participants (red edges in Figure 2A) when the networks estimated at the group level contained only positive edges. The extensive heterogeneity in withdrawal symptom structure echoes heterogeneity in temporal symptom networks observed in disorders beyond tobacco withdrawal (e.g., Fisher et al., 2017; Reeves & Fisher, 2020) as well as long-standing calls for the need for an idiographic science to inform psychological processes (Molenaar, 2004). The heterogeneous experience of tobacco withdrawal is increasingly appreciated (Piper, 2015). There are individual differences in the extent to which smokers endorse the experience of various withdrawal symptoms during periods of smoking abstinence (Pang et al., 2019; Weinberger et al., 2017) and tracking withdrawal symptoms across prequit and postquit periods reveals substantial individual differences in the trajectories of craving and negative affect intensity (McCarthy et al., 2006; see also Robinson et al., 2007). The heterogeneity in withdrawal experiences, observed in this study and in others, may impact the extent to which treatments can successfully target withdrawal in the service of facilitating smoking cessation. Developing the type of data and methodologies that capture person-specific withdrawal experiences provides a pathway to precision medicine, in which treatments are tailored to individuals (Allenby et al., 2016).

In addition to allowing an appreciation for the vast heterogeneity of tobacco withdrawal, constructing person-specific, temporal networks of withdrawal symptoms enabled us to probe the notion of self-perpetuating symptom networks and its implications for the experience and treatment of withdrawal. At the core of network theory in psychology are notions of individual differences in the extent to which symptom networks are vulnerable to the reverberation of symptom activity across time as symptoms influence one another. We quantified the extent to which withdrawal symptom networks were vulnerable to self-perpetuation by capturing the time it took for symptom activity to recover after a simulated increase in activity that spread along the estimated directed paths of the symptom network. Findings showed that combination nicotine replacement smoking cessation treatment was associated with reduced symptom network recovery time relative to a placebo condition. While combination nicotine replacement has been shown to work by reducing craving (Bolt et al., 2012), this research suggests that this treatment may also reduce the time it takes for a smoker to recover and return to baseline levels of withdrawal (i.e., negative affect, cognitive disturbance, craving) after an acute increase in withdrawal symptom activity. Further evidence for the validity of symptom recovery time are observations that longer system recovery time are associated with more intense experiences of anxiety, sadness, irritability, and difficulty concentrating during the EMA protocol.

Perhaps surprisingly, the monotherapy placebo group showed faster recovery time relative to the combination placebo group. One potential explanation for this observation is that the monotherapy placebo group exhibited the most smoking lapses during the quit period. As such, the short network recovery time observed in the monotherapy group may stem less from treatment effects (or lack thereof in this case) and more from continued smoking which alleviated the monotherapy placebo group’s withdrawal experiences to a greater extent than the other groups showing fewer smoking lapses.

In considering differences between the group-level and person-specific networks, it is important to highlight a number of methodological differences. The group temporal networks incorporate data from all individuals in estimating edges and, thus, estimates represent the prototypical individual in the sample. In contrast, the person-specific networks are estimated in an approach that models only data from the individual. Additionally, estimates from six different models are combined to create the group temporal networks. The person-specific approach, in contrast, leverages structural equation modeling to estimate all potential edges in the network simultaneously. Advantages of the person-specific approach are not, however, without limitations. Estimating a well-fitting person-specific model requires substantially more data than approaches that aggregate data across groups of people. Indeed, in the present case, person-specific networks could not be constructed for each individual due to the length of the time series required for estimation.

Going forward, it will be important to identify the mechanisms underlying the network edges. In the context of the cross-talk observed between emotions across time, for example, emerging work suggests the relevance of individual differences in the neural correlates of cognitive control as well individual differences in the use of emotion regulation strategies involving the adoption of an accepting and open attitude to one’s current experience for understanding the spread of emotions to other emotions across time (Drake et al., 2019; Lydon-Staley, Kuehner, et al., 2019; Lydon-Staley, Xia, et al., 2019).

Limitations and Future Outlook

Despite observing associations between symptom recovery time and treatment, we observe no association between symptom recovery time and biochemically confirmed smoking abstinence at end of treatment (8 weeks postquit) or at 6 months
postquit. The lack of association between symptom network recovery time and smoking cessation abstinence may be interpreted in a number of ways. We emphasize that recovery time provides information on how individual differences in tobacco withdrawal symptom network structure might promote symptom spread across time following a hypothetical, simulated perturbation. As such, recovery time represents a potential risk factor through which events external to the withdrawal network may act to promote persistent symptom activity. Indeed, in a study using impulse response analysis to examine the recovery time of sadness to a simulated perturbation, longer recovery time was associated with overall higher levels of depression symptoms in participants experiencing high exposure to stressful life events but not in participants experiencing low exposure to stressful life events. In examining the association between withdrawal network recovery time and smoking cessation outcomes going forward, it will be important to concurrently measure the occurrence of events outside of the withdrawal network that could lead to increases in symptom activity in ways that mirror the simulated shock of the current study.

The person-specific approach allowed an appreciation for the heterogeneous experience of withdrawal as well as the computation of system recovery time, allowing us to capture theoretical notions of self-perpetuating symptom networks. However, estimating good-fitting person-specific models required substantially more data than approaches that aggregate data across groups of people. In the present case, person-specific networks could not be constructed for each individual in the sample due to the length of the time series required for estimation, a difficulty observed in applications beyond tobacco withdrawal (Yang et al., 2018). We hope that highlighting the difficulties of estimating person-specific networks using the type of time series data that has provided substantial insights into withdrawal dynamics to date, as well as the important insights associated with the heterogeneous experience of withdrawal, encourages future collection efforts of denser time series data of tobacco withdrawal. An additional potential limitation is the use of random assessments twice per day between waking and sleeping. The use of random assessments minimizes potential problems with participant anticipation of measurement occasions but also complicates the assumption of equal intervals between pairs of EMA observations.

Smoking lapses are very common during smoking quit attempts (Ashare et al., 2013; Shiffman et al., 2006) and the prototypical participant in the present study reported smoking since the last EMA report at approximately 18% of their EMA prompts. As such, the symptom networks should be interpreted as withdrawal networks estimated during a quit attempt marked by occasional lapses. With denser withdrawal symptom time series data, extensions of the unified structural equation modeling approach used here to examine person-specific networks could incorporate information on lapses into the modeling approach, treating smoking lapses as events that may moderate network edges (Gates et al., 2011).

Conclusions

The present study examined the temporal network structure of tobacco withdrawal. In doing so, we find that there are substantial associations between individual withdrawal symptoms across time, and that smoking cessation treatments may impact the extent to which withdrawal symptom activity lingers across time due to moment-to-moment intersymptom associations. These findings suggest that a network perspective of tobacco withdrawal is a promising approach to understanding withdrawal and its treatment to help all smokers succeed in quitting smoking. The study highlights the challenges associated with using cross-sectional networks to inform within-person associations between symptoms across time (Bos et al., 2017), reinforcing the need to collect intensive repeated measures data to articulate theories of within-person variation in withdrawal symptoms. The vast heterogeneity observed in withdrawal symptom experiences also provides challenges for generalizability, necessitating consideration of methodological approaches capable of capturing both idiographic experiences as well as experiences that may be common to most smokers (Beltz et al., 2016).

References


Received November 5, 2019
Revision received September 8, 2020
Accepted September 24, 2020