Deriving symptom networks from digital phenotyping data in serious mental illness

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Background
Symptoms of serious mental illness are multidimensional and often interact in complex ways. Generative models offer value in elucidating the underlying relationships that characterise these networks of symptoms.

Aims
In this paper we use generative models to find unique interactions of schizophrenia symptoms as experienced on a moment-by-moment basis.

Method
Self-reported mood, anxiety and psychosis symptoms, self-reported measurements of sleep quality and social function, cognitive assessment, and smartphone touch screen data from two assessments modelled after the Trail Making A and B tests were collected with a digital phenotyping app for 47 patients in active treatment for schizophrenia over a 90-day period. Patients were retrospectively divided up into various non-exclusive subgroups based on measurements of depression, anxiety, sleep duration, cognition and psychosis symptoms taken in the clinic. Associated transition probabilities for the patient cohort and for the clinical subgroups were calculated using state transitions between adjacent 3-day timesteps of pairwise survey domains.

Results
The three highest probabilities for associated transitions across all patients were anxiety-inducing mood (0.357, P < 0.001), psychosis-inducing mood (0.276, P < 0.001), and anxiety-inducing poor sleep (0.268, P < 0.001). These transition probabilities were compared against a validation set of 17 patients from a pilot study, and no significant differences were found. Unique symptom networks were found for clinical subgroups.

Conclusions
Using a generative model using digital phenotyping data, we show that certain symptoms of schizophrenia may play a role in elevating other schizophrenia symptoms in future timesteps. Symptom networks show that it is feasible to create clinically interpretable models that reflect the unique symptom interactions of psychosis-spectrum illness. These results offer a framework for researchers capturing temporal dynamics, for clinicians seeking to move towards preventative care, and for patients to better understand their lived experience.

Keywords
mhealth; EMA; networks; psychosis; smartphones.

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Serious mental illnesses, such as schizophrenia, which are still often diagnosed using largely static symptom reports, are increasingly viewed as network illnesses existing along a spectrum of severity and time. The dynamic and multidimensional nature of psychosis is clear, but the static labels and current diagnoses are unable to capture this complexity. New tools and approaches, such as smartphone digital phenotyping, offer promise in more effectively characterising these dynamic illnesses.

When approaching psychosis spectrum illnesses through a network model, comorbid conditions such as mood and anxiety disorders are not static labels but rather impermanent states that a patient may experience with varying frequency along the course of illness. These comorbidities affect one another across time, as high rates of anxiety and depression in those at risk for developing psychosis suggest these conditions may be related to the underlying psychopathology, and a wealth of cross-sectional data confirms anxiety and depression are related to psychosis severity. A network model allows psychosis symptoms to be viewed in the context of other mental health symptoms that are concurrently assessed, providing a conceptual framework for how a patient’s prior symptoms may be related to future ones. For example, ecological momentary assessment (EMA) research suggests that psychosis’ impact on quality of life is mediated through depression and social functioning as well as anxiety. Research also suggests that depressive symptoms may act as a moderator between psychosis and suicidality and that mood itself may predict psychosis symptoms. Teasing apart the causal effects within networks of mental health symptoms is complex, but has strong potential for personalised and preventative psychiatric care.

Digital phenotyping
One useful tool to embrace this complexity is digital phenotyping. This method uses smartphones to capture EMA data, such as self-reported symptoms, as well as functional data from sensors embedded in mobile devices, such as step count and survey response time for each EMA question. For example, information from smartphone screen interactions (for example latency) can be used as a proxy for cognitive state and measurements of sleep/physical activity/sedentary activity can be derived from a smartphone’s accelerometer. Additionally, this novel smartphone data are captured longitudinally, enabling the observation of temporal dynamics across different physical and social environments.

Current applications of digital phenotyping in mental health research are expanding. In schizophrenia, they have already been used to explore the relationships between anticipation and experience of pleasure as well as geolocation and mood. However, few studies have explored how the relationships between the variables measured with digital phenotyping tools can be coalesced into a single context – i.e. a symptom network. In prior work, our team has combined different digital phenotyping signals using anomaly detection to predict a specific event (relapse), and
another team’s digital phenotyping work on suicidal ideations has suggested unique clusters of phenotypes.\textsuperscript{17} Still, there is a need to model and understand symptom interactions in their own right, without having to consider a prediction or clustering task that seeks to inform a single clinical outcome.

**Generative models**

Discriminative models, such as logistic regression and support vector machines, use data to discern between discrete outcomes – for example whether an exercise regimen will prevent a future stroke. However, they do not provide information regarding the underlying distribution of the data. Generative models, on the other hand, are able to provide information regarding the distribution of two or more variables, and from this distribution new samples can be generated – for example a cancer patient’s oncogenic profile can be used to predict what their profile will look like 1 month into the future. With their ability to learn the distributions of longitudinal, multivariate data, generative models may be able to elucidate the underlying relationships that characterise networks of mental health symptoms, as experienced on a moment-by-moment basis. The neuroscience research community has already realised the potential of generative models\textsuperscript{18} and has used such as a foundation for computational-based approaches to neuropsychiatry.\textsuperscript{19,20} However, the potential of this digital phenotyping approach is unknown and challenging to explore because few studies have offered the tools or code to enable others to reproduce experiments or data pipelines.

**Study aims**

In this paper we explore how generative models utilising digital phenotyping data can be used to capture unique symptom interactions in severe mental illness. Utilising open source digital phenotyping tools and fully sharable methods, we present an example of replicable research in the hope others will expand, challenge and adapt our efforts.

**Method**

**Recruitment and participation**

For this study, 47 patients in active treatment for schizophrenia were recruited from a community mental health centre in the metro Boston area, USA and from several satellite programmes that patients in care at this community health centre attend during the day. A total of 43 healthy controls were recruited from Craigslist and local colleges. Four patients were removed from the study because of study drop-out, and one patient was removed because of complete non-engagement – i.e. they completed no digital assessments. Five controls were removed from the data-set because of study drop-out. Demographic information for the remaining participants is listed in Table 1.

Clinical diagnoses of schizophrenia were confirmed with the treating psychiatrists, and healthy controls were screened for current or previous mental illness. Inclusion criteria for those with schizophrenia included: age 18 or older, in active treatment at a community mental health centre, owning a smartphone able to run the study app and the ability to sign informed consent. Comorbid illness was not an exclusion factor. Inclusion criteria for study controls were: age 18 or older, no reported mental illness (both current and prior) and owning a smartphone able to run the study app.

All procedures involving human patients were approved by both the Beth Israel Deaconess Medical Center and State of Massachusetts Department of Mental Health Institutional Review Boards. Written informed consent was obtained from all participants.

| Table 1 Demographic information for patient and controls* |
|----------------|----------------|----------------|
|               | Control group | Patient group |
| n              | 38            | 42            |
| Age, years: mean (s.d.) | 32.30 (16.36) | 38.09 (14.64) | 0.1 |
| Male gender, n (%)   | 20 (52.6)     | 23 (54.8)     | 1   |
| Ethnicity, n (%)     |               |               | <0.001 |
| American Indian or Alaskan | 0 (0.0)      | 2 (4.8)       |
| Native              |               |               |
| Asian               | 26 (68.4)     | 0 (0)         |
| Black or African American | 4 (10.5)     | 15 (35.7)     |
| Multiracial or other | 3 (7.9)       | 2 (4.8)       |
| White               | 5 (13.1)      | 23 (54.8)     |

*Patients in active treatment for schizophrenia were recruited from several mental health community centres in the metro Boston area, and healthy controls were recruited from Craigslist and local colleges. Participants were removed from the study because of study drop-out or complete non-engagement – i.e. they completed no digital assessments. The participants who were removed are not included in the demographic information.

In the initial visit, participants were asked to take a series of surveys assessing their lifestyle and physical and mental health, including the Patient Health Questionnaire (PHQ-9),\textsuperscript{21} Generalized Anxiety Disorder (GAD-7),\textsuperscript{22} Social Functioning Scale,\textsuperscript{23} Short Form Health Survey,\textsuperscript{24} Behavior and Symptom Identification Scale 24,\textsuperscript{25} Warning Signals Scale\textsuperscript{26} and Pittsburgh Sleep Quality Index (PSQI).\textsuperscript{27} For the patient group, the Brief Assessment of Cognition in Schizophrenia (BACS)\textsuperscript{28} battery was performed, and the Positive and Negative Syndrome Scale (PANSS)\textsuperscript{29} was used to record symptoms.

The mindLAMP smartphone app,\textsuperscript{3} a digital health platform developed by our group, was downloaded onto participants’ phones. The code and instructions are available online at digitalpsych.org/lamp. Each weekday, participants were notified via the app to undertake a batch of surveys. Notifications for mood, sleep and social functioning surveys were sent to users on Mondays, Wednesdays and Fridays and notifications for anxiety, psychosis and social functioning surveys were sent on Tuesdays and Thursdays. Participants were also prompted on weekdays to complete smartphone-based cognitive assessments, modelled after the Trail Making A and B tests used in prior neuropsychiatric research.\textsuperscript{30} Although symptom expression may change faster than on a day-by-day basis, previous smartphone studies also assessing longitudinal trajectories have used a similar resolution.\textsuperscript{31}

Complete assessment information, including question content and the notification schedule, are listed in Supplementary Table 1 available at https://doi.org/10.1192/bjo.2020.94.

After 90 days, study participants were asked to return to the clinic for a follow-up visit, including the same surveys and batteries as at the first visit. Their participation in the study concluded at this time.

EMA data from a previous pilot study of 17 patients with schizophrenia were also used as validation data.\textsuperscript{41} This study assessed mood, anxiety, psychosis and sleep via self-reported surveys similar to those in main study. Sociability and cognition were not assessed in the pilot study. Details for how this data was used to validate results from the main study are listed in the Statistical analysis section below.

**Data normalisation and discretisation**

Using the LAMP application programming interface,\textsuperscript{32} participant data was preprocessed so that daily values were found for each participant in every data domain. Because data was collected naturally, meaning users could engage with the app as they saw fit, there were instances where a participant completed a particular survey more than once on a given day. If this was the case, the
daily value for the given survey was set to the average of all completed surveys of that type on that day.

Data in each domain was normalised to zero mean and unit variance across the patient cohort. Using the domain means and variances derived from the patient group, the control and validation cohorts were also normalised. For each patient, normalised daily survey results were grouped into 3-day bins, as patients were expected to report symptoms in each domain once every 2–4 days (Supplementary Table 1). If there existed several results in a given domain and a given bin, the average of the results was used; if there were no events in a given domain and a given bin, imputation was performed by taking the mean of the bin directly preceding and following the bin in question; if both adjacent bins were empty, the bin was ignored.

Scores for each bin were discretised into one of two states: elevated or stable:

(a) elevated results are those that are equal to or greater than one s.d. above the domain mean;
(b) stable results are those that are less than one s.d. above the domain mean.

Although we acknowledge that this threshold has its limitations, as it does not necessarily differentiate between acute psychiatric episodes and innocuous symptom fluctuation, this 1 s.d. threshold has been used in previous papers to dichotomise EMA results into commonly stable and uncommonly elevated states.33

Transition probabilities

Adapting from Betzel & Bassett,18 we calculated transition probabilities for each domain (Fig. 1). We define a ‘transition’ as the elevated/stable state of an assessed domain at a specified time unit compared with the state of the domain in the subsequent timestep.

All combinations of pairwise adjacent bins were generated for all participants in the cohort, and the transition events were counted across all of these pairs, for each domain. Conditioning on the initial state in each pair, the transition events were converted into probabilities:

\[
P[x_{t1}^d = I | x_{t0}^d] = \frac{\# \text{events } (x_{t0}^d = x_{t0}^d \text{ AND } x_{t1}^d = x_{t1}^d)}{\# \text{events } (x_{t0}^d = x_{t0}^d \text{ AND } x_{t1}^d \neq x_{t1}^d) + \# \text{events } (x_{t0}^d = x_{t0}^d \text{ AND } x_{t1}^d = x_{t1}^d)}
\]

where \(x_t^d\) is a binary elevated or stable score in domain \(d\) at timestep \(t \in \{t_0, t_1\}\); \(x_t^I\) is a cohort-wide variable, being one of the two dichotomous elevated/stable states in domain \(d\) at timestep \(t \in \{t_0, t_1\}\); and \(I\) is a dummy variable indicating a certain state \(I \in \{\text{elevated}, \text{stable}\}\).

Fig. 1 Generating transition events from semi-continuous ecological momentary assessment (EMA) data.

Self-reported symptom scores were categorised as being elevated (dark green) or stable (light green) based on the predefined threshold of 1 s.d. above the study mean (a); scores were grouped into 3-day windows, and window state categorisations were determined from the mean of all scores in the respective window (b). Pairs of adjacent windows were generated, and similar pairs were grouped together based on the category of both the initial window and the next 3-day window (c). Probabilities were then calculated based on the initial (time \(t_0\)) state (d).

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The mean elevated bout duration, or the number of days in which a participant is expected to remain in an elevated state, was calculated for each domain across the cohort.

After calculating the transition probabilities within each domain (Fig. 2(a)), we calculated the transition events of domain pairs (Fig. 2(b)). By looking at transition events in a pairwise fashion, we double the number of initial states, as well as double the number of states in the \( t+1 \) timestep. While the number of states grows exponentially with the number of domains in the joint set – \( 2^n \), where \( n \) is the number of domains that are being observed concurrently – we reduced this set size by only looking at a subset of transition events – specifically, those in which a stable domain at time \( t \) transitions to an elevated state in the \( t+1 \) time period, with the complement domain already existing in an elevated state at time \( t \). We call these events associated transitions, as the presence of an elevated state in one domain and the occurrence of a future elevated state in a complement domain suggests that the former domain may be associated with a pathological change in the latter.

Probabilities were calculated for these associated transitions, for patients and controls. Associated transition probabilities for patients were compared with those of the validation cohort, which consisted of 17 patients from a pilot study.

**Clinical subgroups**

In order to move towards personalisation while still retaining enough data to find associated transitions, models were produced for subgroups of participants experiencing similar symptoms. Participants were divided into these subgroups based on the following in-clinic measures: PHQ-9 (mood/depression), GAD-7 (anxiety), PSQI (sleep duration), BACS (cognition), composite PANSS (psychosis) and component PANSS (positive and negative symptoms). Participants with more severe symptoms – those scoring at or above the cohort median for the given measure – were grouped together, and those below the cohort median were also grouped together for further analysis. Subgroups are non-exclusive, so participants can belong to any number of groups (Fig. 3). For each subgroup, mean elevated bout duration and associated transition probabilities were calculated.

**Statistical analysis**

When comparing mean survey scores and bout lengths, 2-sided \( t \)-tests were performed, with a 5% level of significance. Chi-squared was performed to determine the significance of associated transition probabilities, with the null hypothesis stating that the likelihood of a pathological transition in one domain is agnostic to the state of the complement domain. In order to further validate the associated transition probabilities of the patient group, we compared them with the associated transition probabilities of the validation group – i.e. the 17 patients from the pilot study. This comparison was performed with a \( \chi^2 \) test, in which a two-way table was used to classify all of the associated and ‘non-associated’ transition events of both the ‘patient’ and ‘validation’ groups. The null hypothesis stated that both the patient set and the validation set would have the same frequencies of transition events, and failure to reject the null hypothesis suggests that the frequencies of transition events does not significantly differ between the patient and validation groups.

All statistical analysis was performed with the SciPy library in Python.

**Results**

Self-reported survey scores were higher in patients than in controls, in every domain (Fig. 4(a)). The greatest difference occurred in self-
reported psychosis and sleep. Self-reported survey scores were also higher in validation participants than in controls. The mean elevated bout duration of patients tended to be higher than that of controls (Fig. 4(c)); however, this difference could not be determined statistically, as there were very few (if any) elevated bouts in controls. There was no significant difference in mean elevated bout length between patients and validation participants. See Supplementary Table 1 for details on participant engagement.

There was no significant difference in attention scores between patients and controls for the easier attention task (the one modelled after Trails A smartphone cognitive assessment). However, patients performed significantly worse than controls at the more complex task, modelled after Trails B smartphone cognitive assessment (Supplementary Table 2). There were no significant associated transitions found between either of the cognitive scores and associated transitions between these cognitive measures and other self-reported symptom domains. Thus, aside from Fig. 5, we do not utilise this data further in figures and analysis.

Elevated PHQ-9, GAD-7, PANSS composite, and PANSS positive subgroups reported higher self-reported survey scores in all five assessed domains compared with the general patient cohort (Fig. 4(b)). The poor sleep subgroup reported the lowest number of significantly elevated self-reported survey scores relative to the patients, with only anxiety and mood being elevated relative to the patient cohort. There was no significance difference in mean elevated bout length between patients and any of the subgroups (Fig. 4(d)).

For single domain transitions, psychosis had the highest probability of remaining in an elevated state in the following timestep, whereas sleep had the lowest (Supplementary Table 3). However, the psychosis domain had the lowest probability of transitioning into an elevated state.

The three highest probabilities for associated transitions were anxiety-inducing mood (0.357, \( P < 0.001 \)), psychosis-inducing mood (0.276, \( P < 0.001 \)) and anxiety-inducing poor sleep (0.268, \( P < 0.001 \)), respectively (Fig. 5). The three lowest probabilities for associated transitions were psychosis-inducing social (0.09, \( P < 0.02 \)), mood-inducing psychosis (0.162, \( P < 0.001 \)) and sleep-inducing psychosis (0.189, \( P < 0.189 \)). There were even lower transition probabilities for various other
domain pairs – notably those including social functioning – but they were non-significant.

None of the probabilities for associated transitions in the patient set were significantly different than those in the validation set, although the validation probabilities did trend to higher values (Supplementary Table 4).

Discussion

Main findings

In this article we have shown that using digital phenotyping data, it is feasible to create generative models based on transition probabilities for associated transitions in the patient set were significantly different than those in the validation set, although the validation probabilities did trend to higher values (Supplementary Table 4).
probabilities for patients with psychosis. Our results suggest that patients with elevated symptoms in specific domains may experience downstream effects in other symptom domains, with anxiety-inducing mood as the highest probability of associated transitions (Fig. 5). Our finding suggests that psychosis may be a moderator of mood symptoms, while itself being induced by anxiety, supporting a network approach towards understanding real-time and dynamic psychopathophysiology. Although the probabilities of associated transitions between either of the smartphone cognitive measures and other self-reported

Fig. 6 Node graphs of patient (a-i) and validation cohorts (a-ii), along with clinical subgroups: all patients (b-i), Patient Health Questionnaire-9 (b-ii), Generalized Anxiety Disorder-7 (b-iii), Brief Assessment of Cognition in Schizophrenia (b-iv), Sleep duration (b-v), The Positive and Negative Syndrome Scale (PANSS) (b-vi), PANSS: positive symptoms (b-vii), and PANSS: negative symptoms (b-viii).

The node diameter represents the average bout duration (number of days) once an elevated clinical state is reached in that domain. Edge (arrow) diameter represents the probability of the target node transitioning into an elevated state in the next timestep. Only significant transition probabilities were included as edges. Edges with probabilities less than 0.2 were pruned in patient and validation graphs. There were no significant transitions in the control cohort.
Clinical relevance
Our results offer eventual clinical relevance in their ability to offer both patients and clinicians generative models to guide preventative psychiatric care. Clinical experience supports heterogeneity in interactions between and severity of mood, anxiety and psychosis symptoms, which are reflected in our transition probability models (Fig. 6(b)). Although the population-level results presented in this paper reflect broad interactions applicable to all patients (Fig. 6(a)), the numerous symptom-specific models (Fig. 6(b)) offer a tool that could help guide an individual’s care.

For example, patients in the GAD-7 subgroup, who reported high anxiety symptoms in clinic, had a 50% higher chance of elevated mood symptoms inducing elevated anxiety compared with the cohort (Fig. 6). This conceptual framework may (a) offer a target for prevention and (b) guide treatment goals towards minimising risk of future transitions.

Although our generative models are not causal, as associated transitions do not account for other variables which may confound – for example the state of other complement symptoms; physical and social activity, measured passively; and demographic information – they offer insights beyond the correlational, cross-sectional relationships. By considering the initial state of the induced symptom domain, they account for autocorrelative effects that may confound results from other correlational models. Prior results on symptoms causality in psychosis supports our finding that both mood and anxiety have an impact on the severity of psychotic symptoms.4,5,7,8

While we report a range in the probabilities of associated symptoms (Fig. 5), all symptoms interactions could potentially induce elevated states in one another. This highly connected symptom network highlights the potential for creating more personalised models (Fig. 6(b)).

Although our results are novel, they are also reproducible. This is especially important as network approaches to psychopathology symptom networks are complex, and the ability to reanalyse such data often leads to novel insights and competitive interpretations.34,35 Thus, our results can be explored in future studies from a neuroscience perspective, as the generative models presented here may offer targets for generative models of the brain proposed in the Bayesian brain hypothesis36 and structural as well as functional brain networks. These digital phenotyping-derived generative models could thus serve as a bridge between units of analysis in the National Institute of Mental Health’s Research Domain Criteria model,37 especially self-reported symptoms/behaviour and circuits.

Limitations
In our model, associated state changes were derived from symptoms pair at 3-day intervals, offering a window large enough to offer a valid transition probability but still small enough to enable early clinical intervention. Our choice of pairwise symptoms enables us to move towards causality while allowing the models to remain clinically interpretable. These pairwise symptom interactions are not independent and may occur concurrently, which could mask more complex symptom interactions. However, a model that utilises more specific combinations of symptoms may not be feasible without a substantially large amount of data. Defining relevant states that are both data-driven and clinically actionable will be necessary for feasible generative models.

Results from validation on a distinct data-set of 17 different patients with schizophrenia show similarity in the duration of elevated symptoms and the interactions connecting them (Figs 5(c) and 6). Findings that transitional probabilities for the training set tend to be lower than those of the validation set may be related to the small size of the latter. Definitions for elevated self-reported clinical states were derived from retrospective data collection, given the lack of prior research using these methods.

Future directions
In this study we focused on active data from surveys as well as passive data from cognitive measures to generate clinically interpretable and actionable models. Future efforts with prospective methods will expand this work to include more multivariate data with sensors that have been used in previous digital phenotyping studies, such as passively derived physical activity,43 GPS44 and sleep measurements.45

Although the generative models presented here are feasible and clinically relevant, other models may further elucidate symptom interactions in severe mental illness. In generating associated transitions, we consider only a 3-day time lag. However, mental health symptoms may affect future symptoms at varying timescales, and these effects, as a function of the time lag, may be nonlinear and non-stationary. A hidden Markov model, another type of generative model that works on discrete states, may better learn the longitudinal nature of the data in its entirety, as it considers more than
just adjacent pairwise events. An autoregressive model, such as the autoregressive integrated moving average, may characterise the time-lagged effect that a symptom may have on itself – i.e. the ‘memory’ of symptoms. Future efforts will implement these models to provide deeper insight into severe mental illness as quantified through digital phenotyping.

In our clinical subgroups, we clustered patients based on similar symptomatology, retaining clinical interpretability. However, there are other methods of clustering that may better characterise patients’ disease states and thus move closer toward personalisation. For example, latent class analysis clusters patients by transforming observed, co-dependent variables – such as mental health symptoms – into underlying classes, which are independent of one another.46 These classes may be used as a new method for diagnostic labelling,47,48 offering a data-driven approach to psychiatric nosology.

Digital phenotyping and probabilistic transitional models based on associated state change offers a feasible means to approach network interactions of symptoms in a clinically actionable manner. Further research with larger sample sizes using the foundation and sharable tools outlined in this article offers a method towards generating models personalised to each patient, supporting preventative care and improving clinically relevant knowledge.

Supplementary material
Supplementary material is available online at http://doi.org/10.1192/bjo.2020.94

Data availability
The data that support the findings of this study are available from J.T. upon reasonable request.

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