INTRODUCTION

Bipolar disorder (BP) is characterized by profound pathological mood swings from mania to depression, interspersed with periods of subsyndromal symptoms (that is, symptoms that are insufficient to constitute a major mood episode). The Diagnostic and Statistical Manual of Mental Disorders (DSM-IV) and now DSM-5 enables a common clinical language between practitioners, but is yet to anchor an understanding of disease etiology of BP.

Longitudinal course of mood is the defining feature of BPI, determining how patients are diagnosed and subtyped. We propose that BPI patients can be objectively classified based solely on their longitudinal course of mood states. A data-driven approach is likely to better represent the random processes behind transitions between mood states and, in doing so, help identify classes of clinical significance.1-7

Here, we introduce a methodology that uses mood data to objectively identify new classes within DSM categories such as BP. It consists of descriptions of (i) a single patient’s mood as a random process and (ii) BPI patients as a Bayesian nonparametric hierarchical model with latent classes. We fit these models to self-reported longitudinal mood data from BPI patients (N=209) tracked over one or more years to determine whether the data substantiate BPI classes.

MATERIALS AND METHODS

Data

Data from BPI patients (N=209) were drawn from the Prechter Longitudinal Study of Bipolar Disorder at the University of Michigan.8 Patients were evaluated using the Longitudinal Interval Follow-up Evaluation (LIFE)9 administered by a clinician involving retrospective analysis of symptoms over the past 1–3 years. Patients were included provided they had a diagnosis of BPI and at least one LIFE. Patients were rated from 0 to 3 weekly for depression, mania and hypomania. A rating of zero was given if they had no history of symptoms in the respective category, one if they had a history of symptoms but no symptoms for the given week, two if they had one or more symptoms for the respective category (mania, hypomania and depression) and three if they met full DSM-IV criteria. Serial assessments were combined, resulting in 1–8 years of observation for study patients (see Supplementary Figure 1). Of the 209 patients, 141 were female, 179 patients were white, 10 black or African-American, 1 American Indian or Alaskan Native, 1 Asian, 11 more than one race and 7 patients of unknown ethnicity. Patients were an average 40.3 ± 12.2 (± s.d.) years of age and had an average 15.6±3.1 (± s.d.) years of education at initial interview for the Prechter study. Years of education was unknown for two patients. The University of Michigan’s Biomedical Institutional Review Board approved all recruitment, assessment and research procedures for the main study HUM00000606. Patients provided written informed consent after receiving a complete description of the study.

Patient model of mood

Mood was modeled as a discrete-time Markov Chain with finite states (DTMC). The finite states represented mood states (defined below). Central to DTMC are one-step transition probabilities, which define the probability that a patient observed in one-mood state is in a certain mood state at the next observation time. The underlying assumption of DTMC (the Markov assumption) is that a patient’s state depends on its history only through their current mood state. Time between observations represented 1 week with respect to LIFE.

We defined mood states in three ways using (i) full DSM-IV criteria, (ii) subsyndromal symptoms and (iii) subsyndromal symptoms with mixed states (Table 1). First, a patient was considered manic (M) if they met full DSM-IV criteria for mania or depressive (D) if they met the full DSM-IV criteria for depression but not mania. A non-episode (nE) state was defined as the absence of meeting criteria for either mania or depression. Second, a patient was considered in a subsyndromal state of mania (m) if manic/hypomanic symptoms (one or more) were present and, if not present, they were considered in a subsyndromal state of depression (d) if depressive symptoms were present. The euthymic state (Eu) was defined as the absence of m or d. Lastly, we included mixed states, where a patient
was considered in a mixed state (md) if they reported subsyndromal symptoms for both depression and mania/hypomania.

Bayesian nonparametric hierarchical model
Patient models of mood were embedded into a Bayesian nonparametric hierarchical model. A hierarchical model uses other patients’ data to improve inferences on an individual. A nonparametric hierarchical model allows for a variable number of classes (with high probability) and are used when it is important to determine the number of classes. The following hierarchy was evaluated:

Bipolar → Class → Patient → Mood.

which involves patient-specific mood dynamics. This hierarchy was found to be a better model, as measured using information criteria, than a hierarchy that has bipolar I, class, and mood levels, but no patient level (see Supplementary Methods).

Patient data were collected in a matrix Y counting week-to-week transitions between mood states (see Supplementary Methods). A patient’s mood is governed by a transition probability matrix X, which along with the DTMC formulation specifies the probability of observing data Y given X. The patient’s transition matrix X was modeled as a Bayesian nonparametric hierarchical model. The model assumes that a patient belongs to one of the classes. Transition probability matrices for patients in a class are identically distributed, leading to similar mood dynamics for a class. Probabilities of belonging to a class and class-specific parameters are drawn from an approximate Dirichlet process.

Model fitting
Markov Chain Monte Carlo estimated the posterior distribution of parameters. Markov Chain Monte Carlo is an iterative algorithm generating a sequence of samples that approximate a target probability distribution. Following a particular blocked Gibbs algorithm, each iteration updated sequentially (i) patient transition matrices, (ii) class parameters, (iii) patient assignments, (iv) probabilities of belonging to a class and (v) additional parameters. Gibbs’ sampling was used when possible (i, iii, iv); otherwise slice sampling was used (ii); Supplementary Methods. To determine whether the results generalize to a larger population, threefold cross-validation was performed in which the population was randomly divided into three equal-sized groups and the analysis repeated three times with one of the groups omitted in each case.

Post analysis
Classes were evaluated for over/under-representation of categorical variables and for differences in the means of certain covariate variables (complete list of variables is in Supplementary Tables 1–3). All variables were collected on year 5 of the Pechter study, resulting in the exclusion of 13 of the original 209 patients from post analysis for joining the study after year 5. In SPSS, significant associations were evaluated using cross-tabulation with a Person’s χ²-test for categorical variables and one-way analysis of variance for covariates. Homogeneity of variances was tested using a Levene test and, when violated, analysis of variance was replaced by a Welch test. Post hoc analysis was performed using standardized Pearson residuals for categorical variables; a Tukey’s test for covariates when homogeneity of variances was not violated; and a Games–Howell test for covariates when homogeneity of variances was violated. Significance was considered an alpha level of 0.05. Significance was not adjusted for multiple testing.

Code availability
Analysis was performed in Matlab (Mathworks, Natick, MA, USA). Code has been made available at https://sites.google.com/site/amylouisecochran/code.

RESULTS
No classes identified using DSM-IV categories
When mood states were defined using DSM-IV criteria, 208 of the 209 patients were assigned the same class, with assignments based on which class he/she was most likely to belong (Figure 1a). The lone patient was atypical for having over 15 episodes of each of mania (M) and depression (D) in 2 years. The large class provides expectations for any patient’s illness course. We define a ‘typical’ patient as one whose parameters take estimated averages for the class, where averages are taken with respect to the estimated posterior distribution. A typical patient spends ~88% of their time in nEp, followed by 9% of their time in the depression state D and 2% in the mania state M (Figure 1b; see Supplementary Table 1 for average transition probability matrix of the class). The propensity toward the depression state D over the mania state M is attributed to both a greater chance of transitioning from the non-episodic state nEp to the depression state D (74% versus 26%) and longer episodes of the depression state D (7 versus 4 weeks). For comparison, the non-episodic state nEp lasts ~52 weeks. Threefold cross-validation also revealed only one large class for each run made up of 139 of 140 patients in the run, 138 of 139 in the second run and 138 of 140 in the third run.

Distinct classes in the subsyndrome
Using partial criteria (that is, subsyndromal states m, d and Eu), three classes are identified, labeled as ‘stable’, ‘depressive’ and ‘rapid cycling’ in a decreasing order of size (Figure 2a). The stable class consists of an estimated 42 ± 7% (± s.d.) of the BPI population; 97 patients in our study are classified as the stable class. The depressive class has 28 ± 7% (± s.d.) of the BPI population and 62 study patients, whereas the rapid-cycling class has 25 ± 5% (± s.d.) of the BPI population and 44 study patients. Six patients were not assigned to one of the three classes. Threefold

Table 1. Criteria from LIFE data for defining mood states in three models

| Model                  | Mood states                          | Criteria based on LIFE                                      |
|------------------------|--------------------------------------|-----------------------------------------------------------|
| DSM-IV model           | Mania (M)                            | 3 On mania scale                                          |
|                        | Depression (D)                       | 0–2 On mania scale                                       |
|                        | Non-episodic (nEp)                   | 0–2 On mania scale                                       |
| Subsyndromal model     | Subsyndromal mania (m)               | 2–3 On mania scale or 2–3 on hypomania scale            |
|                        | Subsyndromal depression (d)          | 0–1 On mania scale or 0–1 on hypomania scale           |
|                        | Euthymia (Eu)                        | 0–1 On mania scale or 0–1 on hypomania scale           |
| Subsyndromal—mixed state model | Subsyndromal mania (m) | 2–3 On mania or 2–3 on hypomania scale; 0–1 on depression scale |
|                        | Subsyndromal depression (d)          | 0–1 On mania scale or 0–1 on hypomania scales          |
|                        | Euthymia (Eu)                        | 0–1 On mania scale or 0–1 on hypomania scales          |
|                        | Subsyndromal mixed state (md)        | 2–3 On mania or 2–3 on hypomania scale; 2–3 on depression scale |

Abbreviations: 0, no history of symptoms; 1, no symptoms over past week; 2, one or more symptoms over past week; 3, symptoms meet full DSM-IV criteria over past week; DSM, Diagnostic and Statistical Manual of Mental Disorder; LIFE, Longitudinal Interval Follow-Up Evaluations.
cross-validation revealed three classes in two of the three runs, and two classes in the remaining run. When allowing for a mixed state, we still identified three classes with 93% of patients separated into the same classes (Supplementary Figure 2). We focus on the classes from subsyndromal model without a mixed state.

Description of subsyndromal classes

The stable class is labeled ‘stable’ for spending the most time in the euthymic state Eu. A typical stable patient spends 86% of their time in the euthymic state Eu and only 8% of their time in the depressive state d and 6% in the manic state m (Figure 2b; see Supplementary Table 1 for average transition probability matrices of each class). They are more likely to transition to the depressive state d over the mania state m (54% versus 46%) after the Eu (Figure 2c) and spend an average of 83 weeks in the Eu, 10 weeks in the depressive state d and 7 weeks in the manic state m before transitioning to another state (Figure 2d).

Compared with the stable class, the depressive class is labeled ‘depressive’ because a typical patient spends 23% of their time in the depressive state d and only 70% in the Eu, 7% of their time in the manic state m. They last an average of 12 weeks in the depressive state d, 25 weeks in the Eu and 4 weeks in the manic state m before a transition to another state. Similar to the stable...
class, they also are more likely to transition to the depressive state d over the manic state m (54% versus 46%) after being in the euthymic state Eu.

The final class is ‘rapid-cycling’ because a typical patient has frequent transitions between states. A typical patient lasts an average of only 7 weeks in the euthymic state Eu compared with 3 weeks in the depressive state d and 2 weeks in the manic state m before a transition to another state. An estimated 13% of their time is spent in the manic state m, the most of the three classes. However, they spend a similar fraction of time in the euthymic state Eu (70%) to the depressive class, leaving 17% of their time in the depressive state d. They were least likely to transition to the depressive state d over the manic state m (49% versus 51%) after in the euthymic state Eu.

Each class is defined based on transition probabilities rather than the number of episodes in a year. Hence, there is always a chance that a particular patient in the rapid-cycling class has less subsyndromal mood episodes in a year than a patient in another class, even though on average the rapid-cycling class is characterized by more subsyndromal mood episodes in a year (see Figure 3). In other words, the number of episodes in a year cannot alone determine classes. For example, all three classes have a significant probability of having one subsyndromal mood episode.

Classes revealed from threefold cross-validation could also be distinguished based on whether patients were relatively more stable or not, and whether patients were relatively more depressive or not (Supplementary Figure 3).

Comorbid clinical phenotypes of subsyndromal classes
Depressive and rapid-cycling classes have a worse clinical outcome than the stable class (Figure 4). Suicide attempts were significantly different between classes \( P = 0.017 \), with suicide attempts more frequent in the depressive class and less in the stable class. Classes differed in overall chronicity (mood symptoms most of the time or otherwise; \( P = 0.009 \)) and general impairment (no loss of status or otherwise; \( P = 0.012 \)), with increased chronicity and general impairment in the rapid-cycling class and decreased chronicity in the depressive class. Classes did not differ with respect to rapid cycling (history or no history), functionality during depression or mania (incapacitated or otherwise) and mixed episodes (history or no history; Supplementary Table 2).

On the Hamilton Rating Scale for Depression, classes differed significantly in the mean scores \( P = 0.014 \) from Welch test; homogeneity of variance violated). The rapid-cycling class had a significantly higher mean score than the depressive class. On the Young Mania Rating Score, classes differed significantly in the mean scores \( P = 0.006 \) from Welch test; homogeneity of variance violated) and baseline score \( P = 0.002 \) from Welch test; homogeneity of variance violated). The rapid-cycling class had a higher mean score on the Young Mania Rating Score than the depressive class; the depressive class had a lower baseline score than both the depressive and rapid-cycling classes. Classes did not differ in age of onset and number of depressive, manic and hypomanic episodes (Supplementary Tables 3 and 4).

Classes were not significantly different with respect to sex \( P = 0.242 \) or age at baseline \( P = 0.052 \); the depressive class was slightly younger at baseline. Type of baseline medication (antidepressant, anti-epileptic, antipsychotic, benzodiazepine, hypnotic, lithium and stimulant) did not differ between the classes.

DISCUSSION
We have demonstrated the utility of objectively classifying bipolar I patients from course of mood. Our method consists of statistically fitting mood data to a Bayesian nonparametric hierarchy with latent states and patient-specific mood dynamics.
Our methodology was applied to 209 bipolar I patients, identifying three classes of similar size when mood states were defined using partial DSM-IV criteria. No classes were identified when relying on strict DSM-IV criteria. Each class shared clinical features, for example, the ‘depressive’ class had higher rates of suicide attempts.

Implication
Current classification often relies on categories whose distinctions are not supported scientifically. There was debate surrounding the development of DSM-5 about how to weigh scientific evidence against historical practice in establishing definitions. Arguments against historical definitions are that disease causes remain largely unknown and boundaries between categories are more fluid than is recognized. The NIMH even proposed a system to research mental illness, the Research Domain Criteria, which emphasizes measurements from genomics to self-reports over DSM categories.

The present study presents a method to objectively classify BPI patients using longitudinal course of mood. Our results show that data-driven classification from mood course is not only viable, but can divide patients into groups with shared clinical features and can agree with clinical intuition, for example, certain patients are often considered to be a more ‘depressive’ subtype. The presented method can be applied to other recurrent mood disorders and other data sets of longitudinal course of mood.

Because data-driven classes emerge based solely on data, rather than expert consensus, they may better reflect etiological differences in what causes mood transitions. For example, genetic studies have looked to better subtype bipolar patients, as patient groups that are defined using imprecise or broad criteria can weaken the statistical inferences that can be obtained. Moreover, when compared with other criteria for data-driven...
classification, longitudinal course of mood states has the unique advantage that it defines BP patients.

Data-driven classes may also better reflect etiological differences between patients because they emerge only when substantiated by data and cannot be determined by other means, for example, using age, sex or baseline medication. Note classes were not revealed using full DSM criteria and our ‘rapid-cycling’ class could not be determined from the number of subsyndromal mood episodes or a patient’s history of rapid cycling. Certain DSM boundaries may be unsupported by data. For example, rapid cycling based on DSM-5 criteria is defined as at least four episodes in 1 year. Kupka et al. find factors (for example, gender) that are increasingly associated with the number of episodes. They and others argue that there is no empirical basis for separating patients on number of episodes.20,21

Comparison with longitudinal studies
Identifying only subsyndromal classes supports monitoring symptoms weaker in severity, number and/or duration.22,23 Stronger criteria can focus clinical efforts on critical patients, but weaker criteria provide more information over time. For example, major episodes are rare and longer observation may be needed to classify patients from major episodes alone. In other words, subsyndromal symptoms offer extensive clinical information. Judd et al. and Paykel et al.24 also find that weaker symptoms predominate severe symptoms, depressive symptoms predominate hypomanic/manic symptoms and moods change frequently in severity and polarity. All these results are consistent with current findings. In addition, they find that BPI patients spend significant time with mood symptoms, for example, 47.3% of the time in Judd et al. and 53% of the time in Paykel et al.25 Depending on the class, a typical patient in our study spends ~12–32% of their time without these symptoms. Differences in chronicity may reflect discrepancies in how mood states are defined between the studies, sample populations or treatment.

Methodological comparison
Our classification method has advantages over structural equation modeling (for example, latent class or latent growth analysis), the current approach to subtype and classify BP and other recurrent mood disorders.17,18 First, we used a random model of course of symptoms to better represent the unpredictable (and nonlinear) nature of mood states. Second, we used a nonparametric hierarchical model to objectively determine the number of patient classes. Structural equation models are typically parametric in that the number of classes are specified for each model.

Random models of mood had previously been fitted to data for major depression,12 BP1–5 and youth BP.6,7 For example, van der Werf et al. fit a random model to describe time-to-recovery from depressive episodes for a cohort of depressive patient. They present a patient-specific model, but do not fit it to data. Bonsall et al. fit modifications of autoregressive models and extended in Moore et al. The theses of Lopez and Fan are more similar to the present study in their use of DTMC and in inferring patient clusters using Bayesian techniques. They find clusters in youth BP, but do not allow mood dynamics to be patient-specific. Patient-specific models outperformed class-specific models in this study.

Limitations
The data used in this study were weekly retrospective data based on the LIFE. Retrospective data can be less accurate than prospective data because memory can be unreliable and dependent on mood. Despite its limitations, retrospective data have one significant advantage: enough data could be collected in one visit to classify a patient. Future work will need to test data-driven classification using prospective data, with consideration for length of observation needed to classify. Weekly data have the disadvantage that they do not capture all the transitions within the week. Mood in BP will often shift dramatically in a matter of days. It may be that the inclusion of daily information could better define classes of clinical relevance.

Another limitation of our study is the use of relatively small (N=209) and homogeneous (mostly female Caucasian) population. Therefore, whereas the utility of data-driven classification can be established, the exact classes identified may not generalize to other populations. To explore this generalizability, our analysis was repeated on subsamples of our data set using a threefold cross-validation. This additional analysis also found no classes when using DSM-IV states and two or three classes when using subsyndromal states with classes separated based on whether individuals were more or less stable and more or less depressive. We also did not identify a mechanism to explain why individuals have different longitudinal patterns of mood, although results suggest that these differences are probably not explained by baseline medication, gender, age or rapid cycling. Identifying a mechanism may require exploring connections between classes and complete medical history that includes hospitalizations, medication response and comorbidities. This exploration, in turn, may also elucidate whether data-driven classes could be used to improve treatment.

Lastly, our model makes assumptions to simplify the analysis that may affect the utility of the model. For example, mood is divided into three or four states in the model, whereas mood may be better described with additional states or as continuous. In addition, the model does not capture factors that could cause changes in longitudinal patterns, such as life events and changes in medication, among others. This assumption and others (such as the Markov assumption) will need to be considered in the future.

CONFLICT OF INTEREST
The authors declare no conflict of interest.

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Supplementary Information accompanies the paper on the Translational Psychiatry website (http://www.nature.com/tp)