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Exploring comorbid depression and physical health trajectories: A case-based computational modelling approach

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Abstract

While comorbid depression/physical health is a major clinical concern, the conventional methods of medicine make it difficult to model the complexities of this relationship. Such challenges include cataloguing multiple trends, developing multiple complex aetiological explanations, and modelling the collective large-scale dynamics of these trends. Using a case-based complexity approach, this study engaged in a richly described case study to demonstrate the utility of computational modelling for primary care research. N = 259 people were subsampled from the Diamond database, one of the largest primary care depression cohort studies worldwide. A global measure of depressive symptoms (PHQ-9) and physical health (PCS-12) were assessed at 3, 6, 9, and 12 months and then annually for a total of 7 years. Eleven trajectories and 2 large-scale collective dynamics were identified, revealing that while depression is comorbid with poor physical health, chronic illness is often low dynamic and not always linked to depression. Also, some of the cases in the unhealthy and oscillator trends remain ill without much chance of improvement. Finally, childhood abuse, partner violence, and negative life events are greater amongst unhealthy trends. Computational modelling offers a major advance for health researchers to account for the diversity of primary care patients and for developing better prognostic models for team-based interdisciplinary care.

KEYWORDS

artificial intelligence, case-based modelling, cluster analysis, comorbid depression and physical health, complexity theory, differential equations, longitudinal analysis, nonlinear dynamics, primary care

1 | INTRODUCTION

Making clinical sense of the comorbid evolution of depressive symptoms and physical health in primary care is a major concern given their consistent association.¹⁻⁶

For example, a cross-sectional review of 1.7 million primary care patients (18 or older) found that over 68.3% of depressed patients (10% of total sample, mean = 52.7 years) had at least 1 comorbid health condition⁷—compared with the rest of the sample (nondepressed, mean = 47.5 years), which had a 41.1% comorbid

physical condition. Also, a 10-year longitudinal study suggested that for men, perceived poor health and chronic illness at baseline was associated with a higher risk of developing depression symptoms.⁸ Further, in some instances, physical health interacts with the coexistence of depression, often resulting in worse health outcomes and an increased burden on provision and cost of health care.⁹⁻¹⁵

Despite these recent insights, research into the comorbid evolution of depression/physical health continues to deal with 2 key challenges.¹⁶ The first is clinical, and the second is methodological.

The clinical challenge is that while the temporal evolution of depression and physical health is often comorbid, this relationship is not singular in its aetiological pattern, but is multiple and aetiologically complex.^{17,18} For example, a review of these interactions suggests that chronic physical illness can sometimes cause depression through physiological mechanisms,¹⁹ particularly changes in allostatic load or the amount of pain experienced or other psychological mechanisms.²⁰ Changes in social circumstances due to disability are also a causal mechanism, although social support modifies this effect.²¹ Furthermore, for many chronic health conditions, where people have concurrent depression, there is no evidence of improvement in the physical condition if the depression is treated.¹⁹ although pain may be better controlled and quality of life may improve if depressive symptoms are reduced. For example, for people living with arthritis who are also depressed, treatment of depression with medication and/or psychological therapy improved pain, function and guality of life.²² There is also evidence that depression can predate and increase risk of developing a range of chronic illnesses, and underlying physiological mechanisms have been identified for this.¹⁹ In turn, however, there are cases where patients with chronic illness do not have a comorbid mental health condition, as in the case of depression.

In short, it appears that not only is the comorbid relationship between depressive symptoms and physical health based on different complex combinations of sociodemographic and clinical factors, this relationship also evolves along multiple and different trajectories, and in some instances, as in the case of some chronic illnesses, there may be no significant negative relationship at all.²³⁻²⁵ The challenge for the current literature, therefore, is to engage in a series of exploratory analyses to create a catalogue of these multiple comorbid trajectories, particularly for primary care and, in turn, to explore what differences in their complex aetiologies account for them, all of which takes us to the issue of method.

The methodological challenge is that while significant advance has been made in primary care and mental health and clinical evaluation research, the study of the complexities of comorbid depression and physical health continues to be beset by a number of methodological challenges.^{2,16,24} On the one hand, there is the issue of data. As discussed in Gunn et al,^{1,2,24} most studies in the field focus on clinical depression rather than depressive symptoms in general and emphasize specific populations instead of the diverse range of patients in primary care. Research also tends to be cross-sectional, and when longitudinal, the number of time-stamps examined is often too low or too spread out. Also, greater emphasis needs to be placed on the continuous (as opposed to discrete) changes in these comorbid trajectories.²³

On the other hand, there is the current methodological mindset of the clinical and mental health fields and the conventions of their statistical techniques. For example, as of 2018, the vast majority of primary care, clinical evaluation, and mental health journals have yet to advocate for (and very seldom publish) studies that employ the latest advances in computational methods and complex systems thinking.²⁶⁻²⁸ In contrast, these same computational methods and systems approaches are being used widely in other scientific fields, including biomedical and health systems research—mainly because they address the aforementioned clinical and methodological limitations.²⁹⁻³⁴ Because of their power, these methods are also used daily to run the cyber-infrastructure world(s) in which we all now live, including the ever-growing smart machinery upon which contemporary health care depends, from MRIs and diagnostic software to surgical robotics and medical informatics to the nudgewear used to change health behaviours.^{23,26,29,35}

Examples of these new computational methods include artificial intelligence, machine learning, systems mapping, visual complexity, genetic algorithms, complex network analysis, agent-based modelling, and dynamical systems modelling.^{26,29,35} For those interested in an intuitive introduction to these methods, see the following map of the complexity sciences (http://www.art-sciencefactory.com/complexity-map_feb09.html).

Among these computational methods, of significant note for clinical evaluation research is the approach known as *case-based complexity*,^{26,27} and more specifically the Sociology and Complexity Science Toolkit^{23,30-34}—which is specifically useful for modelling multiple comorbid trends across time, particularly those that are high dynamic, as in the case of severe depression. Given the SACS Toolkit's utility for such inquiries, we used it for the current study.

1.1 Case-based computational modelling

The SACS Toolkit is an established case-based, computationally grounded, mixed methods framework for modelling complex topics.^{33,34} It is part of the wider methodological field of study known as case-based complexity and case-comparative methods, specifically qualitative comparative analysis.^{26,27}

Given its computational approach, however, the SACS Toolkit provides some useful advantages over qualitative comparative analysis, growth mixture modelling (GMM), and other statistical methods.^{23,25} To begin, as already suggested, the SACS Toolkit allows comorbid depression-physical health trajectories to be studied as continuous (as opposed to discrete), so that the modelling process focused not only on how trends differ but also on how trends change across time, particularly when high dynamic, as in the case of severe depression. To do so, it employs a novel combination of casecomparative method in conjunction with vector quantization, genetic algorithms, ordinary differential equations, and nonequilibrium statistical mechanics, specifically transport theory and the continuity (advection) partial differential equation. Second, rather than fitting comorbid depressive symptoms and physical health to a function-as is done in GMM-the SACS Toolkit fits its complex functions to these trajectories, which allows for the type of highly refined curve fitting shown in Figures 1 and 2 later in the paper, as well as the identification of minor (small size) and major trends. The result is a multistep, multilevel procedure for transforming the nonlinear dynamics of complex trajectories into cases, clusters, and densities. In the current paper, we provide a quick step-by-step overview of how we used the SACS Toolkit in the current study, sufficient for readers to gain an appreciation of this approach. (For those interested in a complete overview, see http://www.art-sciencefactory.com/cases.html.)

1.2 | Case study: the Diamond cohort

In addition to employing the SACS Toolkit, the current study examined a subsample of the *Diamond* prospective longitudinal study²⁴—which



FIGURE 1 Self-organizing topographical map of 11 major and minor trends. Figure 1 reads as follows: u-matrix and component maps for final 11 exploratory trends (comorbid depression/physical health trajectories). This solution was a reduction of the k-means 18-cluster solution—which is why all 3 maps above show 18 different cluster numbers and their respective cluster name (which is one of the final 11 clusters). Map A and map B are graphic representations of the cluster solution arrived at by the self-organizing map (SOM) neural net, referred to as the u-matrix. In terms of the information they provide, map A is a 3-dimensional (topographical) u-matrix: For it, the SOM adds hexagons to the original map to allow for visual inspection of the degree of similarity among neighbouring map units; the dark blue areas indicate neighbourhoods of cases that are highly similar; in turn, bright yellow and red areas, as in the lower right corner of the map, indicate highly defined cluster boundaries. Map A side view gives a more visually intuitive sense of the topography of the map. Map B is a 2-dimensional version of map A that allows for visual inspection of how the SOM clustered the individual cases. Cases on this version of the u-matrix (as well as map A) were labelled according to their k-means cluster membership

explores the natural history of depressive symptoms over time. Diamond was useful because it is informed by a social model of depressive symptoms and physical health insomuch as it examines the aetiological role that clinical, sociodemographic (marital status, age, drug usage, abuse history, etc.), and health service factors (mental health treatment received, current medications, etc.) play in depressive symptoms and chronic illness—which we explored in the current study.³⁶⁻³⁸ We also chose this database because it is 1 of the largest primary care depression cohort studies worldwide and because it addresses many of the aforementioned methodological challenges, including (a) studying persistent depressive symptoms in general (from subsyndromal to clinical depression) and (b) conducting a longitudinal assessment at 3, 6, 9, and 12 months and then an annual follow-up for a total of 7 years—see Section 2 for details.

1.3 | Research questions

Given our methodological and clinical concerns, for our study, we sought to explore the following 4 research questions:

- First, what are the major and minor trends along which comorbid depressive symptoms and physical health evolve?
- Second, are there trends where depression and physical health are not comorbid? For example, do any such trends exist where



FIGURE 2 Eleven major and minor comorbid depression/physical trends. Figure 2 reads as follows: Each graph provides the longitudinal cluster centroids for the 11 major and minor trends in the data. On the left side are the centroids for depression (our first trace variable), and on the right side are the centroids for physical health (our other trace variable). The *x*-axis represents time, moving from time = 0 to time = 84 months. In terms of the *y*-axis for depression, a high score on PHQ-9 indicates poor health; in contrast, a high score on PCS-12 (physical health) indicates excellent health. The graphs also include labels for each region, going from "severe" to "mild" for depression, and "poor health" to "good health" for physical health. In terms of sample statistics, for PHQ-9, the mean was 7.93 (standard deviation = 6.04), and for PCS-12, the mean was 45.16 (standard deviation = 11.58)

chronic illness is not positively correlated with high rates of depression or clinical depression? Or, alternatively, are there instances where clinical depression is not associated with chronic illness or significant negative physical health?

- Third, when examined together, do these trends exhibit any largescale collective dynamics? For example, are there large-scale similarities among certain trends that cannot be identified when looking at the individual trajectories alone?
- Finally, what combination of clinical and sociodemographic factors account for the different trends identified or large-scale collective dynamics?

1.4 | Methodological caveat

Before proceeding to our methods, we need to clarify a few things. First, the current study, while inferential in nature, was primarily exploratory (as opposed to confirmatory). That is, while we made descriptive claims for what appeared to be key comorbid trends, along with their collective large-scale dynamics and their corresponding complex aetiology, our tentative conclusions require further replication. We also acknowledge, as discussed in Section 2.3 below, that in order to model the dynamics of multiple comorbid depression and physical health trends across time, we required a complete subsample (N = 259) for all time points of the Diamond study (N = 789). As such, further inquiry may be useful to explore the entire dataset for other trends-however, as discussed in Section 2.4 below, our baseline results were similar to those found in the missing data. Finally, to demonstrate the importance that complexity science gives to minor (longtail) trends, we also explored the small-n trajectories typically treated by conventional health research as outliers, mainly in order to identify high-dynamic minor trends that would be otherwise missed. As such, these trends, while providing important insights, may not be replicable in other studies, given that, while vetted by expert analysis, they are the result of cluster analysis and unsupervised machine intelligence, and could be therefore a statistical artefact.

2 | METHODS

2.1 | Design and clinical setting

As stated in Section 1.2, this study used the Diamond database,³⁶⁻³⁸ which was created by recruiting subjects in 2005 in Victoria, Australia from a group of 30 randomly selected family practices, ranging from small private practices to multidisciplinary community health centres. Time-stamps included initial assessment (t = 0), and then follow-up at t = 3, 6, 9, and 12 months and then annually for 7 years. The University of Melbourne's Human Research Ethics Committee approved the Diamond study and consent to publish (reference number: 030613X). For the current study, we examined all 11 time-stamps.

2.2 | Enrolment, retention, and subsample

Subjects were eligible for the study if they scored a 16 or higher on the well-known *Center for Epidemiological Studies-Depression* (CES-D) and read English—see³⁷ for details. Of the N = 789 subjects that were provided informed consent, N = 129 (16.3%) were lost to follow-up between 3 and 12 months—a common problem for community-based cohort studies—dropping the total to N = 449 subjects. Diamond's dropout for the first year nonetheless compared favourably to similar studies.³⁹

2.3 | Justification for and validity of current sample

As mentioned in Section 1.4, the justification in the current study for reducing the Diamond sample to N = 259 cases was based on its exploratory (albeit inferential) purpose: We sought to examine complete longitudinal data to gain new insights into how global measures of comorbid depression and physical health coevolve differently across time. Furthermore, we sought to examine continuous (as opposed to discrete) change, which also required that our data be complete at all sample points. Finally, given that we also explored

the larger vector field formed by these multiple trajectories, again complete interpolated data were necessary. For more on these data requirements, see Castellani et al.²³

Still, as shown in Table 1, the N = 259 subsample was, overall, satisfactorily similar to the original N = 789 cohort in Gunn et al,²⁴ with the following differences highlighted. The current subsample was slightly less educated, had a higher rate of marriage, was less likely supported by a pension, slightly more employed, significantly lower in current smoking, and self-assessed health status was slightly better on the lower end of things (fair/poor health). The current subsample was also slightly lower on number of participants having been told they are depressed by a doctor. And yet the subsample had the same number of participants taking antidepressants and antianxiety medications as in the cohort and had almost the exact same CES-D baseline scores for depression. Also, the subsample and cohort were similar in terms of physical health. Still, caution needs to be given to any argument that the current study constitutes anything more than a working catalogue which requires further corroboration, editing.

2.4 | Missing data

Given that missing data in a longitudinal cohort may be related to clinical state, we explored further the N = 530 missing data cases. Cluster analysis grouped them into 4 cluster trends—each based on when data became missing in the 7 years of the study. As shown in Table 2, the baseline depression and physical health scores for the study sample (N = 259) are roughly similar to the 4 missing data clusters, suggesting that those who left the study did not vary significantly in their depression or physical health. Still, differences across time could have existed. Nonetheless, it seems that the exploratory results of the current study, at least at baseline, are reasonably valid with respect to the Diamond cohort.

2.5 | Measures

In terms of measures, we examined a rather exhaustive list of 40 variables used in Castellani et al's²³ study of the Diamond database. For more on the study design, they used see also Boardman et al,³⁶ Gunn et al,³⁷ and Potiriadis et al.³⁸

2.5.1 | Global outcome variables

Because our study was designed to explore and catalogue the different trajectories of comorbid depressive symptoms and physical health, we chose the following 2 well-known global outcome variables. For depressive symptoms, we used the Primary Care Evaluation of Mental Disorders Patient Health Questionnaire (PHQ-9).⁴⁰ The PHQ-9 is a global, multipurpose instrument for screening, diagnosing, and clinically measuring depression severity. The more severe the depression is, the higher the score. For physical health, we used the physical health component (PCS-12) of the SF-12 Heath Survey.⁴¹ The PCS-12 is a 6-item, multipurpose global assessment of physical health. The higher the score, the better the physical health. With these 2 global outcome variables identified, we could then model and explore their intersection across time and then group them into a working catalogue of their respective major and minor trends. A final note: Once

Characteristic	Cohort (n = 789)	Sample (N = 259)
	Mean (SD)	Mean (SD)
Age in years	48.0 (13.1)	49.81 (12.14)
SEIFA advantage deciles (IRSAD)	6.8 (2.4)	6.9 (2.39)
CES-D score (baseline)	27.2 (9.4)	26.24 (9.21)
	Number (%)	Number (%)
Gender (female)	563 (71.4)	185 (71.4)
Marital status		
Never married	184 (23.5)	50 (19.3)
Widowed/divorced/separated	228 (29.1)	78 (30.1)
Married	371 (47.4)	131 (50.6)
Lives alone	167 (21.3)	61 (23.6)
Highest level of education		
Left school before year 10	134 (17.0)	32 (12.4)
Completed year 10, 11, or 12	300 (38.0)	93 (35.9)
Certificate of diploma	190 (24.1)	72 (27.8)
Bachelors degree or higher	163 (30.7)	61 (23.6)
Pension/benefit main source of income	281 (36.0)	79 (30.5)
Employment		
Employed/student	475 (60.2)	169 (65.3)
Unemployed	200 (25.3)	64 (24.7)
Unable to work due to sickness/ disability	111 (14.1)	24 (9.3)
Hazardous drinking in past 12 months	180 (23.0)	52 (20.1)
Current smoker	249 (31.7)	60 (23.2)
Long-term illness/health problem/ disability	405 (52.5)	128 (49.4)
At least 1 chronic physical condition in past 12 months	542 (68.8)	180 (70.0)
Self-assessed health status		
Excellent/very good	171 (21.7)	62 (23.9)
Good	296 (37.5)	115 (44.4)
Fair/poor	322 (40.8)	79 (30.5)
Ever told by doctor had depression	530 (70.5)	167 (64.5)
Currently taking depression medication	317 (40.2)	104 (40.2)
Currently taking antianxiety medication	77 (9.8)	24 (9.3)

the final database was set, all scores on PHQ9 and PCS-12 were converted to z-scores to remove scale bias.

2.5.2 | Clinical profile variables

Given that our second goal was to explore the set of clinical and sociodemographic factors accounted for the differences in the major and minor trends identified in the first part of our study, we examined 38 baselines variables from the Diamond database, which were broken

TABLE 2 Cluster an	alysis of	N = 530	missing	data cases
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down into 16 sociodemographic factors, 18 psychological factors, and 4 physical factors.²³ See Tables 3 and 4 for details.

2.6 | Case-based computational modelling

For this study, we employed a combination of statistical and computational techniques under the general methodology of *case-based complexity*.^{26,27,42} Case-based complexity seeks to advance current statistical and computational methods by studying cases in complex systems terms.^{23,26,27} Presently, a variety of techniques exist. The particular platform we used was the case-based computational modelling framework known as the *SACS Toolkit*.^{23,30} For more on this approach, see Section 1.1 above.

2.6.1 | Analytic procedure

Our order of analysis is as follows. First, to identify and catalogue our major and minor comorbid trends, we followed current convention, employing a combination of (a) k-means cluster analysis, (b) a self-organizing topographical neural net (SOM), and (c) expert knowledge. Next, to determine how the 38 clinical and sociodemographical variables combined to uniquely account for different comorbid trends, we used a combination of ANOVA (for our continuous measures) and chi-square (for our discrete nominal measures)—see Tables 3 and 4. This multistep approach, which the SACS Toolkit employs, has proven highly useful, as it follows a rather rigorous process for corroboration.^{23,43} Steps are as follows:

- Step 1: Creating longitudinal clusters. Modelling multiple comorbid trends involves clustering case trajectories. To do so, we treat each time instance as a measure, and the total of time instances/measures as the longitudinal k-dimensional vector profile for each case. The result is a database where the rows on the right-hand side of the database are cases (ie, c1, c2, c3, ... n), and the columns across the top, in turn, are the discrete scores on depression and then physical health, at each time (t) instance (ie, t1, t2, t3 ... n) for each case—for details on this approach, see Castellani et al.²³ In turn, these trajectories are combined (appended to one another) so that the cluster solution is based on similarities in evolution across our 2 global outcome measures. For the current study, we appended the 7-year trajectories for PHQ9 and PCS-12 to each other for each of our N = 259 cases.
- Step 2: K-means. Analysis begins with k-means, which requires researchers to postulate the expected number of cluster trends, based on current theory. For the current study, given our literature review, we assumed that there would be several large and mostly healthy cluster trends, followed by a handful of smaller and more pathological trends, ending with a group of highdynamic, minor cluster trends with high rates of comorbid

Cluster Name	Missing Data from Start of Study Forward	Missing Data from Middle of Study Forward	Missing Data from End of Study Forward	Data Missing but No Major Trend	N = 259 Valid Cases in the Study
Count	N = 114	N = 84	N = 159	N = 173	N = 259
PHQ9 mean	10.39	10.11	10.80	11.68	10.03
PCS-12 mean	80.67	81.03	78.94	78.21	82.20

Longitudinal Trend												ELLA
	Healthy (N = 58)	Okay Vacillate (N = 20)	Okay Same (N = 27)	Okay Improving (N = 26)	Moderate Depression Improving (N = 18)	Episodic Depression 1 (N = 16)	Episodic Depression 2 (N = 22)	Moderate Depression Poor Health (N = 14)	Unhealthy (N = 9)	Chronic III (N = 23)	Oscillators (N = 17)	NI ET AL.
Psychological												
Days out of role due	6.34***	8.7***	3.84***	4.38***	16.11 ^{***}	7.81***	8.24***	19.64***	55.43***	10.22***	36.33***	
to emotional problems***	12.96 sd	15.39 sd	8.86 sd	12.14 sd	24.08 sd	13.83 sd	9.53 sd	29.10 sd	37.80 sd	22.15 sd	32.59 sd	
Hazardous drinking in last 12 months**	20.7%** (0.1) se	20.0%** (0.0) se	11.5%** (-1.0) se	38.5%** (2.1) se	5.6%** (-1.4) se	37.5%** (1.5) se	19.0%** (-0.1) se	35.7%** (1.3) se	22.2%** (0.1) se	8.7%** (-1.2) se	11.5%** (-1.0) se	
Any substance abuse ^a	12.3% (-0.7) se	15.0% (-0.1) se	7.7% (-1.0) se	24.0% (1.0) se	11.1% (-0.5) se	6.7% (-0.9) se	19.0% (0.4) se	21.4% (0.5) se	22.2% (0.5) se	17.4% (0.2) se	24.0% (1.0) se	
Never smoked	46.6% (0.1) se	40.0% (-0.4) se	51.9% (0.5) se	42.3% (-0.2) se	47.1% (0.1) se	43.8% (-0.1) se	54.5% (0.6) se	42.9% (-0.1) se	44.4% (0.0) se	43.5% (-0.1) se	38.5% (-0.5) se	
Currently smoke	22.4% (-0.1) se	15.0% (-0.8) se	11.1% (-1.3) se	15.4% (-0.8) se	29.4% (0.5) se	25.0% (0.1) se	22.7% (-0.1) se	35.7% (1.0) se	44.4% (1.3) se	13.0% (-1.0) se	42.3% (2.0) se	
Depression a current	39.3% ^a	25.0% ^a	34.6% ^a	42.3% ^a	72.2% ^a	66.7% ^a	71.4% ^a	78.6% ^a	88.9% ^a	47.8% ^a	88.0% ^a	\
problem ^a	(-1.5) se	(-1.8) se	(-1.4) se	(-0.8) se	(1.0) se	(0.7) se	(1.1) se	(1.2) se	(1.4) se	(-0.4) se	(2.3) se	
Ever told by doctor	50.9% ^a	57.9% ^a	50.0% ^a	57.7% ^a	70.6% ^a	86.7% ^a	85.7% ^a	92.9% ^a	100% ^a	61.9% ^a	88.0% ^a	NIL
had depression ^a	(-1.5) se	(-0.5) se	(-1.1) se	(-0.6) se	(0.2) se	(0.9) se	(1.0) se	(1.2) se	(1.2) se	(-0.3) se	(1.3) se	
Ever told by doctor	41.8% ^a	58.8% ^a	30.4% ^a	42.9% ^a	62.5% ^a	91.7% ^a	70.6% ^a	77.8% ^a	100% ^a	50.0% ^a	73.9% ^a	EY-
had anxiety ^a	(-1.4) se	(0.2) se	(-1.6) se	(-0.8) se	(0.3) se	(1.7) se	(0.8) se	(0.9) se	(1.6) se	(-0.3) se	(1.2) se	
Dysthymia ^{ba}	1.8% ^a (-2.1) se	5.0% ^a (-0.8) se	0.0% ^a (-1.7) se	8.0% ^a (-0.4) se	11.1% ^a (0.1) se	6.7% ^a (-0.5) se	9.5% ^a (-0.2) se	14.3% ^a (0.4) se	44.4% ^a (3.1) se	21.7% ^a (1.6) se	28.0% ^a (2.7) se	Jour
Currently taking	24.6% ^a	30.0% ^a	26.9% ^a	23.1% ^a	33.3% ^a	37.5% ^a	50.0% ^a	57.1% ^a	55.6% ^a	26.1% ^a	73.1% ^a	nal of
depression meds ^a	(-1.5) se	(-0.5) se	(-0.8) se	(-1.1) se	(-0.2) se	(0.1) se	(1.0) se	(1.3) se	(0.9) se	(-0.8) se	(3.1) se	
Currently taking	8.6%	5.0%	3.7%	0.0%	22.2%	6.3%	22.7%	0.0%	22.2%	8.7%	11.5%	Evalu
antianxiety meds	(-0.2) se	(-0.6) se	(-0.9) se	(-1.6) se	(1.8) se	(-0.4) se	(2.1) se	(-1.1) se	(1.3) se	(-0.1) se	(0.4) se	
Currently taking	1.7% ^a	10.0% ^a	0.0% ^a	0.0% ^a	0.0% ^a	0.0% ^a	0.0% ^a	7.1% ^a	0.0% ^a	0.0% ^a	15.4% ^a	ation
antipsychotic meds ^a	(-0.6) se	(1.8) se	(-0.9) se	(-0.9) se	(-0.7) se	(-0.7) se	(-0.8) se	(0.9) se	(-0.5) se	(-0.8) se	(3.6) se	c Health Po
Currently taking	3.4% ^a	5.0% ^a	0.0% ^a	3.8% ^a	0.0% ^a	6.3% ^a	27.3% ^a	0.0% ^a	0.0% ^a	0.0% ^a	7.7% ^a	in Clir
sedative meds ^a	(-0.5) se	(0.0) se	(-1.2) se	(-0.3) se	(-1.0) se	(0.2) se	(4.7) se	(-0.8) se	(-0.7) se	(-1.1) se	(0.6) se	olicy and H
Partner abuse, severe ^{ca}	1.8% ^a (-1.7) se	5.0% ^a (-0.5) se	4.2% ^a (-0.7) se	4.0% ^a (-0.8) se	11.8% ^a (0.5) se	20.0% ^a (1.5) se	9.1% ^a (0.1) se	14.3% ^a (0.7) se	44.4% ^a (3.7) se	0.0% ^a (-1.3) se	16.7% ^a (1.4) se	ical P
Childhood sexual	12.1% ^a	20.0% ^a	26.9% ^a	23.1% ^a	35.3% ^a	31.3% ^a	38.1% ^a	42.9% ^a	44.4% ^a	9.1% ^a	46.2% ^a	ractic
abuse ^{da}	(-2.1) se	(-0.5) se	(0.1) se	(-0.3) se	(0.7) se	(0.4) se	(1.1) se	(1.2) se	(1.1) se	(-1.6) se	(2.0) se	
Childhood sexual	12.1% ^a	20.0% ^a	23.1% ^a	19.2% ^a	35.3% ^a	25.0% ^a	28.6% ^a	35.7% ^a	44.4% ^a	9.1% ^a	46.2% ^a	e
abuse, severe ^{da}	(-1.8) se	(-0.4) se	(-0.1) se	(-0.5) se	(1.0) se	(0.1) se	(0.4) se	(0.9) se	(1.3) se	(-1.4) se	(2.3) se	ch
Childhood physical	27.6% ^a	35.0% ^a	46.2% ^a	50.0% ^a	33.3% ^a	56.3% ^a	50.0% ^a	64.3% ^a	55.6% ^a	36.4% ^a	69.2% ^a	
abuse ^{da}	(-1.9) se	(-0.6) se	(0.1) se	(0.4) se	(-0.7) se	(0.7) se	(0.4) se	(1.1) se	(0.5) se	(-0.6) se	(1.9) se	
Childhood physical	12.1% ^a	30.0% ^a	19.2% ^a	30.8% ^a	16.7% ^a	37.5% ^a	27.3% ^a	46.2% ^a	44.4% ^a	31.8% ^a	42.3% ^a	
abuse, severe ^{d**}	(-2.2) se	(0.3) se	(-0.8) se	(0.4) se	(-0.8) se	(0.8) se	(0.0) se	(1.3) se	(1.0) se	(-0.4) se	(1.5) se	
											(Continues)	129

 TABLE 3
 Eleven clinical profiles for major and minor trends on key psychological and physical factors

1299

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Longitudinal Trend											
	Healthy (N = 58)	Okay Vacillate (N = 20)	Okay Same (N = 27)	Okay Improving (N = 26)	Moderate Depression Improving (N = 18)	Episodic Depression 1 (N = 16)	Episodic Depression 2 (N = 22)	Moderate Depression Poor Health (N = 14)	Unhealthy (N = 9)	Chronic III (N = 23)	Oscillators (N = 17)
Physical											
Chronic illness or	12.1% ^a	35.0% ^a	63.0% ^a	34.6% ^a	27.8% ^a	56.3% ^a	63.6% ^a	92.9% ^a	88.9% ^a	95.7% ^a	70.8% ^a
disability ^{ea}	(-4.1) se	(-0.9) se	(1.0) se	(-1.1) se	(-1.3) se	(0.4) se	(0.9) se	(2.3) se	(1.7) se	(3.1) se	(1.5) se
Chronic condition	58.6%	85.0%	74.1%	73.1%	55.6%	75.0%	81.8%	78.6%	66.7%	73.9%	76.9%
last 12 months ^f	(-1.1) se	(0.7) se	(0.2) se	(0.1) se	(-0.8) se	(0.2) se	(0.6) se	(0.3) se	(-0.2) se	(0.2) se	(0.4) se
Self-health rating ^{5***}	2.54***	2.90***	3.35***	2.81***	2.83***	3.00***	3.32***	3.79***	3.89***	3.68***	3.54***
	0.76 sd	0.64 sd	0.75 sd	0.80 sd	0.99 sd	0.63 sd	0.65 sd	0.70 sd	1.05 sd	0.84 sd	1.24 sd
Days out of role for	2.95***	6.95***	18.96***	13.81***	3.06***	7.07***	3.10***	19.21***	45.71***	22.37***	22.65***
physical health***	8.52 sd	16.54 sd	25.57sd	26.63 sd	9.35 sd	12.84 sd	3.71 sd	28.27 sd	41.86 sd	28.55 sd	31.07 sd
old-factore bad similification	t arobability will	ni orchi-ranora	chidee percenta	pacts pac (%) en	ordized error (ce)	or standardized	Posidual (er). ANC	acem epiderio VV	and standard de	viation (cd)	

(nc) D N 0 280 $^{\rm a}{\rm Any}$ substance abuse (CIDI 12-month disorders) including alcohol. $^{\rm 55}$ Bold-faced factors had significatit prov-

^bDysthymia (CIDI 12-month disorder).⁵⁵

^cPartner abuse (CAS)⁵² (scale used by Gunn et al¹⁴ was 0: no abuse, 1: other abuse, 3: severe abuse.

 $^{\rm d}{\rm Childhood}$ sexual and physical abuse, MacMillan et al.^{53}

^eLong-term illness, health problem or disability.

fAt least 1 chronic physical condition in the past 12 months, including asthma, emphysema, arthritis, hypertension, back problems, chronic sinusitis, lipid disorder, cancer, stroke, and dermatitis.

⁸Self-health rating: 1 = excellent health, 2 = very good, 3 = good, 4 = fair, 5 = poor health.

*Chi-square significant at 0.05 or higher.

**Chi-square significant at $P \leq 0.08$.

***ANOVA significant at 0.001 or higher.

Longitudinal Trend											
	Healthy (N = 58)	Okay Vacillate (N = 20)	Okay Same (N = 27)	Okay Improving (N = 26)	Moderate Depression Improving (N = 18)	Episodic Depression 1 (N = 16)	Episodic Depression 2 (N = 22)	Moderate Depression Poor Health (N = 14)	Unhealthy (N = 9)	Chronic III (N = 23)	Oscillators (N = 17)
Income ^{a***}	4.89***	3.55***	3.77***	4.13***	4.67***	4.00***	4.62***	2.54***	1.71***	3.43***	3.24***
	2.09 sd	2.14 sd	2.46 sd	2.38 sd	2.25 sd	2.34 sd	2.01 sd	1.76 sd	1.50 sd	2.34 sd	2.45 sd
Socioeconomic	7.50 [†]	7.35 [†]	6.63 [†]	7.12 [†]	7.83 [†]	7.06 [†]	7.24 [†]	6.50 [†]	6.00 [†]	6.04 [†]	5.96 [†]
advantage ^{b†}	2.05 sd	2.30 sd	2.53 sd	2.18 sd	1.98 sd	1.98 sd	2.55 sd	2.96 sd	3.16 sd	2.82 sd	2.25 sd
Highest level of education ^{c***}	3.84***	3.70***	2.93***	3.12***	3.17***	3.31***	3.64***	2.93***	2.33***	2.57***	3.27***
	1.19 sd	1.30 sd	1.14 sd	1.28 sd	1.34 sd	1.45 sd	1.40 sd	1.39 sd	1.58 sd	1.31 sd	1.49 sd
Visits to health	7.09***	8.28***	9.06***	8.06***	9.06***	12.5***	13.46***	14.71***	17.33 ***	8.33***	14.67***
professional ^{d***}	5.20 sd	6.90 sd	5.41 sd	6.71 sd	6.46 sd	8.58 sd	10.07 sd	10.19 sd	16.05 sd	5.20 sd	7.94 sd
Age***	43.58***	51.73***	57.17***	48.35***	47.33***	54.32***	44.05***	58.59***	49.78***	57.07***	48.70***
	11.49 sd	11.65 sd	9.84 sd	10.14 sd	11.01 sd	14.07 sd	11.91 sd	7.26 sd	7.33 sd	12.34 sd	11.00 sd
Negative life events score (0 to 13)***	1.74*** 1.26 sd	1.95*** 1.47 sd	1.70*** 1.38 sd	2.12*** 1.51 sd	1.56*** 1.29 sd	1.69*** 1.62 sd	2.41*** 1.50 sd	2.43*** 1.83 sd	4.11*** 2.37 sd	1.96*** 1.30 sd	2.46*** 1.70 sd
SSQ number of	2.24***	2.05***	2.08***	2.12***	2.00***	2.19***	1.71***	1.64***	1.67***	1.87***	1.73***
supporters ^{e***}	0.63 sd	0.76 sd	0.70 sd	0.65 sd	0.77 sd	0.54 sd	0.96 sd	0.63 sd	0.71 sd	0.63 sd	0.67 sd
Social participation	32.22***	29.50***	25.19***	29.27***	20.67***	24.19***	31.82***	24.14***	19.22***	22.00***	20.54***
score ^{f***}	10.32 sd	14.00 sd	9.48 sd	9.15 sd	8.44 sd	9.14 sd	11.43 sd	11.17 sd	8.20 sd	12.81 sd	10.23 sd
Unable to work ^{8*}	0.0%*	0.0%*	3.7%*	4.0%*	0.0%*	12.5%*	0.0%*	21.4%*	75%*	21.7%*	23.1%*
	(-2.3) se	(-1.4) se	(-1.0) se	(-0.9) se	(-1.3) se	(0.4) se	(-1.4) se	(1.5) se	(6.1) se	(1.9) se	(2.3) se
Employed ^{8*}	81.0%*	60.0%*	51.9%*	76.0%*	83.3%*	43.8%*	86.4%*	50.0%*	41.2%*	47.8%*	61.5%*
	(1.4) se	(-0.3) se	(-0.9) se	(0.6) se	(0.9) se	(-1.1) se	(1.2) se	(-0.7) se	(-1.2) se	(-1.1) se	(-0.1) se
General practice	29.3%	45.0%	29.6%	23.1%	22.2%	31.3%	31.8%	28.6%	22.2%	43.5%	34.6%
location (rural)	(-0.2) se	(1.1) se	(-0.1) se	(-0.7) se	(-0.7) se	(0.0) se	(0.1) se	(-0.2) se	(-0.7) se	(1.1) se	(0.3) se
Private health	60.3%	60.0%	66.7%	65.4%	55.6%	62.5%	59.1%	50.0%	27.8%	52.2%	35.3%
insurance	(0.4) se	(0.2) se	(0.7) se	(0.6) se	(0.0) se	(0.3) se	(0.2) se	(-0.3) se	(-1.6) se	(-0.2) se	(-1.1) se
Per cent female	23.8%	9.7%	10.8%	9.2%	6.5%	4.3%	9.2%	4.9%	3.2%	7.6%	10.8%
(N = 185) ^h	(-0.6) sr	(1.0) sr	(0.2) sr	(-0.4) sr	(-0.2) sr	(-1.0) sr	(0.3) sr	(-0.3) sr	(-0.2)	(-0.6) sr	(0.3) sr
Per cent male	18.9%	2.7%	9.5%	12.2%	8.1%	10.8%	6.8%	6.8%	4.1%	12.2%	8.1%
(N = 74) ^h	(-0.6) sr	(-1.6) sr	(-0.3) sr	(0.6) sr	(0.4) sr	(1.6) sr	(-0.5) sr	(0.5) sr	(0.3) sr	(0.9) sr	(-0.5) sr

 TABLE 4
 Eleven clinical profiles for major and minor trends on key sociological factors

1301

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TABLE 4 (Continued)	(
Longitudinal Trend											
	Healthy (N = 58)	Okay Vacillate (N = 20)	Okay Same (N = 27)	Okay Improving (N = 26)	Moderate Depression Improving (N = 18)	Episodic Depression 1 (N = 16)	Episodic Depression 2 (N = 22)	Moderate Depression Poor Health (N = 14)	Unhealthy (N = 9)	Chronic III (N = 23)	Oscillators (N = 17)
Lives alone	15.5% (-1.3) se	30.0% (0.6) se	22.2% (-0.1) se	23.1% (0.0) se	33.3% (0.9) se	25.0% (0.1) se	9.1% (-1.4) se	42.9% (1.5) se	33.3% (0.9) se	13.0% (-1.0) se	41.6% (1.2) se
Married	48.3% (-0.2) se	45.0% (-0.4) se	66.7% (1.2) se	50.0% (0.0) se	33.3% (-1.0) se	56.3% (0.3) se	59.1% (0.6) se	42.9% (-0.4) se	38.9% (-0.7) se	60.9% (0.7) se	47.1% (-0.2) se
Satisfied with support ^{i**}	77.6% (0.9) se	73.7% (0.3) se	68.0% (0.0) se	80.0% (0.8) se	64.7% (-0.1) se	68.8% (0.1) se	50.0% (-1.0) se	50.0% (-0.8) se	38.9% (-1.5) se	63.6% (-0.2) se	80.0% (0.6) se
Bold-faced factors ha	d significant probak	oility values. Chi-s	quare includes p	ercentage (%) and	d standardized err	or (se) or standardi	ized residual (sr); AN	JOVA includes me	an and standard (deviation (sd).	
^a Income: $1 = 20\ 000$ (or less; 7 = 70 000	or higher (Austra	lian currency).								
^b Socioeconomic adva	ntage: index of rela	tive socioeconom	ic advantage and	l disadvantage (se	e Gunn et al ¹⁴ fo	r more details.)					
^c Highest level of educ	cation: 1 = left schc	ool before year 1C); 2 = completed	year 10; 3 = con	pleted year 12; 4	= certificate/diplo	ima; 5 = bachelors o	ır higher.			
^d Self reported visits to	o health profession	al includes numbe	er of visits to any	r and all health pr	oviders, from gen	eral practitioner to	social worker.				
^e SSQ number of supp	orters (Sarason et ;	al ⁵⁴) mean out of	9.								

^eFor employment, we reported 2 different statistics 1 = employed/student or 2 = unable to work due to sickness or disability. ^fSocial participation score (Baum et al⁵¹) score range 0 to 90.

^hThis statistics provides the percentage of each gender for each cluster, relative to the total sample for each gender. Also, reported here is the chi-square standard residual, which is the difference between the observed count and the expected count and the expected count.

Satisfied with support, 1 = yes.

*Chi-square significant at 0.05 or higher.

**Chi-square significant at P = 0.06.

***ANOVA significant at 0.05 or higher.

[†]ANOVA significant at 0.08.

depression and physical unhealthiness. As discussed in the results, k-means arrived at an 18-cluster solution.

- Step 3: SOM. The next step is to corroborate the k-means 18-cluster solution using the SOM. The current study used the SOM Toolkit.⁴⁴ Because the SOM engages in unsupervised cluster analysis, it decides which cluster solution is optimal—based on 2 validity measures: quantization error and topographical error.^{45,46} While these error measures are unstandardized, the closer to zero the better, with topographical error scores less than 10 considered a good fit. Similar to Google Analytics, if the unsupervised SOM is a good fit and arrives at a solution similar to the k-means, it provides effective corroboration.
- Step 4: Visual inspection of SOM. As shown in Figure 1, the SOM graphs its cluster solution onto a multidimensional surface called the u-matrix. On the u-matrix, comorbid depressive symptoms and physical health cases most like one another are graphically positioned as nearby neighbours, with the most unlike cases placed furthest apart. As shown later in Figure 1 (map A top and side view), the u-matrix is also topographical: Valley (darker coloured) areas represent comorbid cluster trends that are more similar, while hilly, brighter coloured areas show comorbid cluster trends that are more distinct.
- Step 5: Comparing k-means to SOM. Map B is a 2-dimensional version of map A, which allows for visual inspection of how the SOM clustered the N = 259 cases for the current study. Cases on this version of the u-matrix (as well as map A) were labelled according to their k-means cluster membership (the 18-cluster solution) to see if the SOM arrived at a similar solution, which it did.
- Step 6: Expert corroboration: With the k-means and SOM corroborated, an expert panel is assembled to review the results. To facilitate this process, a visualization of the comorbid depressive symptoms and physical health trends was also created, as shown in Figure 2. This allowed the panel to visually inspect the trends and name and catalogue their differences, as well as get rid of or collapse trends into one another. For the current study, our panel of primary care physicians and mental health professionals (which included 3 of the current authors for this study) collapsed several of the small-n, high-dynamic, minor trends together, resulting in the final exploratory 11-cluster solution shown in Figures 1 and 2.
- The names of these clusters, in order, were Healthy (n = 58), Okay Vacillate (n = 20), Okay Same (n = 27), Okay Improving (n = 26), Moderate Depression Improving (n = 18), Episodic Depression 1 (n = 16), Episodic Depression 2 (n = 22), Moderate Depression Poor Health (n = 14), Unhealthy (n = 9), Chronic III (n = 23), and a collection of small-n trends grouped together to form the Oscillators cluster (n = 17). Still, our expert panel did agree that despite the important insights they provide on high-dynamic depression, the Unhealthy and Oscillating clusters, given their small-n size, could be a statistical artefact of using k-means cluster analysis and unsupervised artificial intelligence to arrive at them.
- Step 7: Aetiology of comorbid trajectories. To determine how the various combinations of our 38 clinical and sociodemographic variables accounted for our 11 trends, a combination of ANOVA and

chi-square was used. However, given the database is only N = 259, we did not engage in any posthoc *t* tests, as the chance for error with an 11-cluster solution across 38 variables, even with the most conservative of statistics, would be too high.

- Step 8: Modelling collective large-scale dynamics. To model the global temporal dynamics of the vector field. CBDM creates what it calls the microscopic model. For the purposes of normalization, all data for the microscopic model were converted to z-scores. We used differential equations and smooth curve fitting techniques to compute the velocities for PHQ9 and PCS12, followed by using a genetic algorithm to fit a polynomial differential equation to the velocities-see Figure 3. The microscopic model (vector field, V) uses the 18 trajectories upon which the 11 trends are based to construct a state-space of all possible trajectories, for all 7 years of the study, starting with the baseline (t = 0). The form of V, which is a part of the ordinary differential equations, is not known to us ahead of time, as our goal is to fit curves to data, rather than the GMM approach, which fits data to curves. This is key to the novelty of our approach. To run our genetic algorithm, we used Eurega's software.⁴⁷ The component functions of the vector field are constrained to have a polynomial form. We choose a polynomial fit without any constraint on the degree, and use the mean squared error with the Akaike information criterion as an error measure. The software provides a measure of stability and maturity: "Stability" refers proportionally to how long ago the top solutions were modified among the multiple solutions provided; "maturity" refers to how long ago any of the solutions have improved. Stability and maturity values close to 100% suggest that the solutions cannot be improved. The software shows multiple solutions ordered according to their level of complexity of polynomials and fit. The midrange solutions are, generally speaking, the best.
- Step 9: Constructing comorbid trend narratives. The last step was to use our expert panel to construct a clinical narrative for each of the 11 trends.

3 | RESULTS

3.1 | Cluster trends

As outlined in step 6 of Section 2.6.1, including the names for each trend, the current study identified 11 major and minor comorbid trends across the 11 time-stamps explored (baseline; 3, 6, and 9 months; and then years 1 through 7). As a reminder from Section 2.6.1, this solution was an expert-based reduction of the original 18 clusters identified by k-means; also, some of the smaller trends, such as Unhealthy and Oscillators, given their small-n size, could be, in part, a statistical artefact of the analyses. Still, the validity estimate for our exploratory SOM solution was satisfactory: quantization error = 2.56; topographical error = $0.02^{45,46}$ —see Section 2 for interpreting these statistics.

The 11 trends and their respective groupings are found in Figure 1;. (See Section 2 for review on how to read Figure 1.) Looking



FIGURE 3 Microscopic model of global-temporal dynamics across 11 trends. Figure 3 reads as follows: The microscopic model uses the nonclustered data to construct a state-space of all possible trajectories, for all 7 years of the study, starting with the baseline (time = 0) and proceeding, across the next 8 snapshots, to the end (time = 84 months). All possible trajectories are visualized in the form of arrows (shown in grey between the blue trend lines), which show direction and velocity; the larger the arrow, the faster the trajectory. For the purposes of normalization, all data were converted to *z*-scores; as such, coordinate (0,0) is the centre, with the majority of the data falling within 2 standard deviations (the inner grey area on each graph). The graphs were cropped at 3 standard deviations, in order to visualize more fully the globally dynamic behaviour of the model, while none of the data actually fell outside the first 2 standard deviations. In other words, this model does not show the trajectories of specific cases and should not be read as such. Instead, it is read as a map of all possible trajectories, with the focus on identifying (as Section 3 does) major global dynamics. The *x*-axis is physical health (with poor physical health on the left and good physical health on the right); the *y*-axis is depression, going from low to high levels of depression as one moves upward along the axis. Two key global-dynamic behaviours were identified: a saddle point and a spiralling source. The graph for time = 0 was likewise labelled to give the reader a rough sense of the different quadrants, from high physical health and low depression to low physical health and high depression

at map B, the SOM placed the *Healthy* trend (N = 58) on the opposite side of the map from the *Unhealthy* (N = 9), *Moderate Depression Poor Health* (N = 14), and the *Oscillators* (N = 17). In map A, the *Unhealthy* (N = 9) and some of the *Oscillators* are on a ridge in the upper right,

separated from the rest of the cluster trends, suggesting that these cluster trends differ significantly from the rest.

Map B also suggests that Okay Vacillating (N = 20), Okay Improving (N = 26), and Okay Same (N = 27) are somewhat similar to the cases in

the Healthy trend, as they are all in the valley of map A, side view. However, while Moderate Depression Improving (N = 18) is also proximate to the Healthy trend, the topography of map A (top and side view) suggests that it is not as similar to Healthy as the other okay health trends as it is not as far down in the valley. It is also worth noting that the 2 episodic depression trends—*Episodic Depression* 1 (N = 16) and *Episodic Depression* 2 (N = 22)—were placed near each other and in the middle. Finally, consistent with the k-means solution, *Chronic III* (N = 23) is off to the left side, as a distinct cluster, separated by the green ridge shown in map A, from the rest of the poor health trends.

Figure 2 is a temporal visualization of Figure 1, showing how the 11 trends evolve across time. On the left side are the trends for depressive symptoms, and on the right side are their corresponding physical health trends. In terms of the y-axis for depressive symptoms, a high score indicates poor health; in contrast, a high score on physical well-being indicates excellent health. Also, as highlighted in grey on Figure 2, it is important to remember that there were a total of 4 time-stamps for the first year of the study (ie, 0, 3, 6, 9, 12 months); as such, one would expect to see a higher degree of dynamics in these first several months—which we do see.

Looking at Figure 2, the first 4 trends are the Healthy to Okay Healthy clusters, which were the largest and most stable, comprising 51% of the total cases in the study. From there, however, the dynamics begin to intensify and diversify, starting with the Moderate Health clusters. Finally, there were the Overall Poor Health clusters. In addition to making up the minor trends in the study, these clusters had a high degree of dynamic fluctuation. The only exception was Chronic III (N = 23; cluster 6), for which depressive symptoms were not dynamic or comorbid with poor physical health.

3.2 | Aetiological profiles

Tables 3 and 4 provide the sociodemographic and clinical profile for each of the 11 trends in our study. Here, we provide a narrative for these indicators. Still, given the richness and complexity of these different profiles, others may identify narratives different from the ones we found. Also, our exploratory goal here is to provide a quick sense of the clusters, knowing that exact aetiological algorithms (rule extraction and the proper weighting of each factor's influence) can be developed through further replication and study.

Our first clinical narrative is for the healthiest cluster trend. Healthy (N = 58), with high across-the-board rates of health and sociodemographic wellbeing, this trend is doing well. It is also the youngest.

Okay Vacillating (N = 20): As the specific indicators in Tables 3 and 4 show, this trend is struggling a bit, including declining physical health, but otherwise okay. Note: By "okay", we mean that the scores on PHQ9 and PCS12 for this trend (as well as others below that use the same term) are in the satisfactory range, but are not exceptionally or especially good. This trend is also middle aged, and at baseline, 85% reported a chronic condition in last 12 months. Ten per cent also report currently taking antipsychotics, the second highest rate among the trends, and 45% have a rural GP, the highest rate (along with the unhealthy group) among the trends.

Okay Same (N = 27): Looking at the indicators in Tables 3 and 4, this trend is doing okay without any significant decline in psychological or physical health. At baseline, 74.1% reported a chronic condition in the last 12 months. This is 1 of the older groups. There is a significant abuse history, with 46.2% reporting childhood physical abuse, and 23.1% reporting severe childhood sexual abuse.

Okay Improving (N = 26): This trend improves across time, and, overall, is younger and scoring well on social indicators. However, at baseline, it has the highest rate of hazardous drinking (38.5%) and 1 of the highest rates of substance abuse (24%).

Moderate Depression Improving (N = 18): struggles with moderate baseline depression that improves across time. However, this trend has the best physical health. Also, although over 80% are working, their social participation rates are among the lowest in the sample. The proportion reporting severe childhood sexual abuse is among the highest of the moderate heath trends, but severe childhood physical abuse is not.

Episodic Depression 1 (N = 16) and 2 (N = 22): These 2 trends mirror each other, with each going up in depression as the other goes down, and with physical health on a somewhat dynamic but steady decline. There are, however, differences. Episodic Depression 1 is older, has lower scores for social support and participation, and only 43.8% are employed; it also has a higher rate of social support satisfaction. However, it is more likely to have experienced partner abuse and severe childhood physical abuse. Still, both trends have considerable and similar sexual abuse exposure. Episodic Depression 2, the younger of the 2 trends, has a higher probability of a chronic health problem and is less likely to have been told they have anxiety by a provider (70.6%). But, this trend is more likely to use sedatives or antianxiety medication, and there is a higher rate of drug dependence.

Moderate Depression Poor Health (N = 14): older and not doing well socially or physically; however, this trend's depression trajectory improves across time, albeit dynamically. This trend also has the second highest annual visits to a health provider, and the second highest negative events score. Fifty per cent are currently employed, with 21.4% reporting that they cannot work. There are also drug dependence issues and some of the highest rates of childhood abuse. Finally, 92.9% reported a chronic illness or disability at baseline and the second worst self-health rating of all the groups.

Unhealthy (N = 9): This middle-aged trend has sustained poor physical and mental health. It also has, overall, the most disadvantaged sociodemographic profile. Psychological distress is also pronounced, with 100% being told by provider, at baseline; they have depression and anxiety. Childhood abuse exposure is second highest of all groups, and severe partner abuse is more than double the rate in any other group. In terms of physical health, they have the third highest rate of chronic illness, they have the worst self-health rating, and the highest rate of days missed for physical and emotional disorders.

Chronically III (N = 23): This older trend is struggling with chronic illness, but only mild depression. However, in terms of physical illness, 73.9% reported a past chronic illness or disability, 73.9% reported a current chronic condition, 21.7% cannot work, and only 47.8% are currently employed. This trend also had the third worst self-health rating and the third highest number of days missed for a physical condition.

Oscillators (N = 17): As a reminder, our study did not seek to remove or ignore small-n trends in order to explore trajectories where the comorbid relationship between depression and physical health was high dynamic. Such was the case with the oscillator trend. Each of these 8 minor trajectories—with the largest cluster added to this trend being N = 4 cases—fluctuates between moderate to severe levels of unhealthiness. As a group, the socioeconomic well-being of the Oscillators is average to below average. They also have (along with Okay Improving) the highest rates of substance abuse; very high rates of depression, anxiety, and dysthymia; and the highest rate of antipsychotic medication usage. Abuse history is also significant, with 69.2%reporting childhood physical abuse and 46.2% reporting severe childhood sexual abuse. They also have 1 of the worst baseline self-health ratings, missing a significant number of days due to emotional or physical issues.

3.3 | Collective large-scale (across trend) patterns

To examine the collective large-scale dynamics of our 11 trends, we created the vector field in Figure 3, which is read as follows. Data were converted to z-scores, with coordinate (0,0) as the average score for both depression (y-axis) and physical health (x-axis). The shaded box shows the standard deviations (sd) within which the 11 trends fell-which correspond, for comparison purposes, to the sd in Figure 2. In this shaded box are 5 pointers-poor health, moderate health, okay health, chronically ill, and moderate depression improving-to help readers locate the particular state-space occupied by the 11 trends. The lines in Figure 3 illustrate the contours of comorbid depression and physical health, as they coevolve across time/space. The arrows (some of which are highlighted in black) indicate the direction of the trajectories, as well as their velocity: The larger the arrow, the faster the trajectory is moving at that point in time. As a final point, Figure 3 is not useful for exploring individual trajectories. Instead, it is to be examined for global dynamic trends-that is, largescale collective patterns-that exist across all 11 trends in the study. In other words, it does not show the trajectory of a particular case, but the potential trajectories of all cases.

Looking at Figure 3, the vector field solution identified 2 unexpected global-temporal patterns, a spiralling source, and a saddle-point, which evolve across time. In terms of a basic definition, spiralling sources and saddle points identify critical attractor points in a system. If the trajectories around such a critical point repel and spiral away from it, it is called a spiralling source. If, however, trajectories follow conflicting courses of action, the critical point is called a saddle. As an example, consider the saddle for time = 84 months. On all 4 sides, the trajectories are converging; however, they never actually run into each other; instead, the saddle point repels them, forcing them back outward in different destinations. This is why it is called a saddle point: It roughly approximates the critical point at which the different trajectories stop increasing or decreasing in time-space. In other words, a saddle point functions like a local minima/maxima, constituting a barrier for how low or high a trajectory can go. In contrast, the spiralling source serves a different function. As its name implies, it is a continuously and gradually widening curve, which winds itself away from a critical point in time/space. Trajectories are repelled away from (or drawn into) a spiralling source, increasing or decreasing the velocity of the trajectories as they reach the outer or inner rings of the curve.

4 | DISCUSSION

While descriptive and inferential in nature, the purpose of this study was exploratory (as opposed to confirmatory): We sought to advance the study of the clinical complexities of comorbid depression and physical health by (a) creating a catalogue of their multiple comorbid trajectories, particularly for primary care; (b) looking for any noteworthy large-scale collective dynamics; and, in turn, (c) exploring the complex aetiologies that accounted for these results. In other words, we were trying to see if we could use complete data at all time points to model dynamics (and their complex aetiology) otherwise outside the conventional purview—as opposed to generalizing some set of confirmatory findings to the general primary care population. In turn, to create this tentative catalogue, we sought to overcome current methodological convention by employing case-based complexity, specifically the SACS Toolkit, which was created for modelling such complex health issues.²³

4.1 | Cluster trends

In terms of the utility of a complexity theory approach, our exploratory analysis concluded that the longitudinal evolution of comorbid depressive symptoms and physical health follows multiple major and minor trends, demonstrating that the more severe the depression, the more dynamic the trends. Most trends are somewhat similar to their neighbouring trends. The exceptions are the Oscillators and Unhealthy groups, which varied in dynamic between each other and are very different from other groups, but depression and physical health are comorbid, and the Chronically III, where physical health is often low dynamic and it is not necessarily comorbid with depression. What is also striking about the cluster solution is how stable most of the trends are, apart from the oscillators, which constitute a very small part of the total dataset. We do not know, however, whether depression symptoms would show a less stable pattern if the cohort participants had not received any treatment over the 7 years, given that we did not explore this issue. Also, we do not know if the missing cases not explored might have demonstrated a different set of dynamics.

4.2 | Clinical profiles

Looking across all 11 trends, we find that, similar to studies such as,²⁴ sexual abuse, childhood physical abuse, partner violence, and negative life events generally increase except for the chronic illness group. These well-known sociodemographic determinants of mental health seem to be keys in determining trend membership.

Another significant finding is the distinctness of the chronic illness trend. Cases in this trend seem to have physical illness that limits their function but they cope without necessarily getting depressed—beyond, perhaps, the transient low mood picked up on screening at baseline that resulted in them joining the cohort. This finding seems consistent with other research.^{20,48-50} Also striking, this trend had

_EY Journal of Evaluation in Clinical Practice

low levels of sexual abuse and partner violence compared to the Unhealthy and Oscillators trends.

4.3 | Collective large-scale dynamics

The clinical utility of the vector field is that it brings alive the reality of the evolution of depression and physical health, by depicting it dynamically at a large scale and across trends. Looking at the results, this stage of the analysis suggests that depression is more dynamic than typically portrayed by growth mixture modelling, and that some of the cases spiral in and out of depression, across time, regardless of their particular level of severity of depression, although the speed of change is slow towards the centre of the spiralling source. The identification of saddle points also suggests that there may be limits to what treatment can achieve for some people-particularly among the Unhealthy and Oscillator trends. However, it also suggests that saddle points are dynamic, so health care experts and public institutions (potentially through effective preventive policy) can potentially change them by, for example, reducing the physical and sexual abuse people (particularly women) experience. Again, these insights are exploratory and, therefore, further analysis and replication is necessary.

4.4 | Implications for interdisciplinary clinical practice

The concern in the current literature (as outlined in the introduction) that health providers are missing the multiple trends of comorbid depressive symptoms (including major depression) when treating people for their physical health seems to be well supported by the current study-and not just because depression is always comorbid with physical health, but because in some instances, as in the case of chronic illness, it is necessary to know when depression is neither significant or at a clinical level. There are also sociodemographic moderators predisposing many cases to the development of depression, particularly childhood abuse and partner abuse, to which clinicians need to give their attention. There are also, however, trajectories of depression/chronic illness that suggest that resilience can act as a barrier to the extremes of severe chronic illness/abuse. Conversely, the trajectories in the extremes of severe chronic illness/abuse tend not to lead back to improved physical and mental health; in other words, some cases, as found in the Unhealthy and Oscillator trends, remain ill without much chance of improvement-which, again, suggests different forms of treatment.

Given these results, it is likely that the different major and minor trends where depression and physical health do coevolve (and where we are too late to prevent causative factors such as childhood abuse) may benefit from different interdisciplinary, team-based approaches or combinations of approaches to treatment. Clinical examples include tackling hazardous drinking, providing medication to lift mood during a dip, strengthening strategies for resilience, and improving the management of the physical condition or providing social support.

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CONFLICT OF INTEREST

The authors declare no conflict of interest.

CONTRIBUTORS

All 4 authors were involved in conceptualizing the manuscript, conducting the analyses, interpreting the results, outlining the references, and writing the paper and consent to its publication.

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Journal of Evaluation in Clinical Practice

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