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## Learning to swim the endless waves: examining self-management and risk factors for (chronic) depression

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## CHAPTER 6

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### **The 9-year clinical course of depressive and anxiety disorders: new NESDA findings.**

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### Abstract

**Background:** In longitudinal research, switching between diagnoses should be considered when examining patients with depression and anxiety. We investigated course trajectories of affective disorders over a nine-year period, comparing a categorical approach using diagnoses to a dimensional approach using symptom severity. **Method:** Patients with a current depressive and/or anxiety disorder at baseline ( $N = 1701$ ) were selected from the Netherlands Study of Depression and Anxiety (NESDA). Using psychiatric diagnoses, we described 'consistently recovered,' 'intermittently recovered,' 'intermittently recurrent,' and 'consistently chronic' at two-, four-, six-, and nine-year follow-up. Additionally, latent class growth analysis (LCGA) using depressive, anxiety, fear, and worry symptom severity scores was used to identify distinct classes. **Results:** Considering the categorical approach, 8.5% were chronic, 32.9% were intermittently recurrent, 37.6% were intermittently recovered, and 21.0% remained consistently recovered from any affective disorder at nine-year follow-up. In the dimensional approach, 66.6% were chronic, 25.9% showed partial recovery, and 7.6% had recovered. **Limitations:** 30.6% of patients were lost to follow-up. Diagnoses were rated by the interviewer and questionnaires were completed by the participant. **Conclusions:** Using diagnoses alone as discrete categories to describe clinical course fails to fully capture the persistence of affective symptoms that were observed when using a dimensional approach. The enduring, fluctuating presence of subthreshold affective symptoms likely predisposes patients to frequent relapse. The commonness of subthreshold symptoms and their adverse impact on long-term prognoses deserve continuous clinical attention in mental health care as well further research.

**Keywords** (3-6 words): Depression; Anxiety; Nine-year Course; Prognosis; Chronicity; Diagnostic Switching

## **Key findings**

- When using a categorical approach (i.e., diagnoses), the majority of patients recovered from affective disorders at nine-year follow-up.
- When using a dimensional approach (i.e., symptom severity), the majority of patients had a more chronic course.
- The majority of patients who did not meet criteria for an affective disorder still experienced subthreshold symptomatology.

## **1. Introduction**

Longitudinal research is essential to validate diagnostic classifications and tailor treatment plans for optimal effectiveness and efficiency for patients with affective disorders (Gillis et al., 1995; Kendell and Jablensky, 2003; Kraepelin, 1921; Lorenzo-Luaces et al., 2017; McGorry et al., 2016; Penninx et al., 2011). In longitudinal research, depressive and anxiety disorders should be investigated synchronously considering that major depressive disorder (MDD) and anxiety disorders often co-occur (Angst et al., 2009; Giandinoto and Edward, 2015; Hayden and Klein, 2001; Howland et al., 2009; Kessler et al., 2005; Kleiboer et al., 2016; Lamers et al., 2011; Moffitt et al., 2007; Rush et al., 2006), and as time passes, one disorder will often switch over to the other ("diagnostic switching") (Gregory et al., 2007; Hovenkamp-Hermelink et al., 2016; Scholten et al., 2016). For example, those recovered from MDD may then meet criteria for generalized anxiety disorder (GAD) and still experience functional impairment and a decreased quality of life (Angst et al., 2009; Maj et al., 2002; Wells et al., 1992). Patients may also switch over to dysthymic disorder, the more chronic but milder form of depression (Klein et al., 2008; Rhebergen et al., 2012).

The selected observation period may also reveal crucial insights into the course of depressive and anxiety disorders. A shorter window of less than two years has led to relatively positive findings with regard to prognosis (Gilchrist and Gunn, 2007; Hendriks et al., 2013; Richards, 2011; Steinert et al., 2014). For example, previous research has shown that more than half (50 – 70%) of patients recovered within one year from MDD (Richards, 2011; Steinert et al., 2014), and approximately half (43 – 73%) of patients recovered within two years from anxiety disorders (Hendriks et al., 2013). However, when extending the observation window, lower levels of recovery were found, which may be due to diagnostic switching (Caspi et al., 2020; Gregory et al., 2007; Hovenkamp-Hermelink et al., 2016; Plana-Ripoll et al., 2019; Scholten et al., 2016), relapse (Hardeveld et al., 2010; Mueller et al., 1999; Scholten et al., 2013; Verduijn et al., 2017), or continuous subthreshold or subclinical symptoms (Ormel et al., 1993; Wagner et al., 2000).

The course of depressive and anxiety disorders can be and have been described using two clinically-relevant approaches: with psychiatric diagnosis criteria according to the Diagnostic Statistical Manual of Mental Disorders (DSM-5) (American Psychiatric Association, 2013) (i.e. categorical approach) or scores on symptom severity scales (i.e. dimensional approach) (Wardenaar et al., 2014). Use of DSM diagnoses allows us to examine participants who (continue to) meet full DSM criteria over time, whether these are comorbid diagnoses or transitions to other diagnoses. Alternatively, symptom severity scores are regularly used throughout the treatment follow-up process to assess change of symptoms over time, such as in routine outcome monitoring (ROM) (e.g. de Beurs et al., 2011). Using symptom severity scores may also help reveal and describe the heterogeneity in (sub)clinical symptom profiles (Brunoni, 2017;

Fried and Nesse, 2015a). Considering that depression and anxiety DSM diagnoses can be viewed as discrete categorical syndromes imposed on a continuum of depressive or anxiety symptoms of varying severity and duration (Kendler and Gardner, 1998; Klein et al., 2006, 2008; Torpey and Klein, 2008), comparing both approaches is relevant. Latent class growth analysis (LCGA) with both depressive and anxiety symptoms would allow us to examine how symptoms change over time as a dimensional approach and to compare the data with DSM diagnoses as a categorical approach.

The present study aimed to describe and compare course trajectories of depressive and anxiety disorders over a nine-year period using a categorical approach for DSM-diagnosis trajectories and a dimensional approach for symptom pathways using LCGA. We considered depressive and anxiety comorbidity, switching to other diagnoses over time, and switching to other trajectories over time. We specified how participants intermittently recover and relapse over a nine-year period and how the DSM diagnoses concur with fluctuating symptom scores. For the categorical approach, we hypothesized that those with baseline comorbid depressive and anxiety disorders, compared to those with baseline depression or anxiety disorder, would have higher levels of chronicity at nine-year follow-up (Hendriks et al., 2013; Richards, 2011; Steinert et al., 2014). Likewise, for the dimensional approach, we hypothesized that those with higher baseline symptom levels would coincide with a more chronic course. Finally, we hypothesized that the trajectories according to the categorical approach would correspond highly with the dimensional symptom pathways.

## **2. Method**

### *2.1 Study Sample and Procedure*

Data were used from the Netherlands Study of Depression and Anxiety (NESDA) (Penninx et al., 2021), an ongoing longitudinal cohort study examining the course of depressive and anxiety disorders (for the extensive protocol, see Penninx et al., 2008). At baseline (2004-2007), a total of 2981 participants aged between 18 and 65 years were recruited from community (19.0%), primary care (54.0%), and specialized mental healthcare (27.0%). After baseline, face-to-face follow-up assessments were conducted at two years (87.1%,  $n = 2596$ ), four years (80.6%,  $n = 2402$ ), six years (75.5%,  $n = 2256$ ), and nine years (69.4%,  $n = 2069$ ). The nine-year follow-up was completed in 2016.

For the present study, we selected a total of 1701 participants with MDD, dysthymic disorder, and/or anxiety in the six months prior to baseline (i.e. a current six-month diagnosis) (see below).

### *2.2 Measures*

#### *2.2.1 Categorical course trajectories*

To determine the categorical trajectories, we examined the presence of depressive (MDD, dysthymic disorder) and anxiety disorders (panic disorder, social phobia, generalized anxiety disorder (GAD), and agoraphobia) according to the DSM-IV criteria at each assessment. The disorders were assessed in person using the Composite Interview Diagnostic Instrument (CIDI) version 2.1 by trained research personnel (Wittchen, 1994; Wittchen and Nelson, 1996). To extrapolate whether the CIDI diagnoses were consistently present during follow-up periods, we examined the continuous presence of diagnosis-related symptoms reported in the Life Chart Interview (LCI; Lyketsos et al., 1994) for

that same follow-up period. More information regarding the instruments can be found in the **Supplementary Material**.

Using the CIDI, participants were first divided into subgroups at baseline: 1) depression only (single or any combination of MDD and/or dysthymic disorder); 2) anxiety only (single or any combination of panic disorder, social phobia, GAD, and agoraphobia); and 3) comorbid depression-anxiety (any combination of both depressive and anxiety disorders). The following “primary” categorical diagnosis trajectories were then described at two, four-, six-, and nine-year follow-up for the total group and for each of the subgroups by examining whether participants had any CIDI diagnosis at each following assessment after baseline: ‘*consistently recovered*’ (fixed diagnosis-free period at each assessment), ‘*intermittently recovered*’ (variable diagnosis-free period preceded by a diagnosis-present period with or without discontinuous LCI symptoms), ‘*intermittently recurrent*’ (variable diagnosis-present period with discontinuous LCI symptoms) and ‘*consistently chronic*’ (fixed diagnosis-present period at each assessment with continuous LCI symptoms). The trajectories of ‘intermittently recovered’ and ‘intermittently recurrent’ allowed us to explore the ‘ebb and flow’ of diagnoses across the nine-year period. At nine-year follow-up, we defined three “comparison” trajectories, where each percentage was aggregated at each subsequent assessment point: ‘*recovered*’ (diagnosis-free period), ‘*recurrent*’ (diagnosis-present period with discontinuous LCI symptoms), and ‘*chronic*’ (diagnosis-present period with continuous LCI symptoms). This allowed us to compare our results to Verduijn et al. (2017), who examined the six-year follow-up NESDA data. An extended description of the trajectories can also be found in the **Supplementary Material** (see “Description of categorical trajectories”). The specific differences

with Verduijn et al. (2017) are also outlined in the **Supplementary Material** (see “Comparison with previous NESDA study”).

### *2.2.2 Dimensional course trajectories*

To determine dimensional trajectories, we used depressive and anxiety symptom severity measures. For depressive symptom severity, we used the Inventory of Depressive Symptomatology – Self-Report (IDS-SR; Rush et al., 1996; Trivedi et al., 2004). For anxiety symptom severity, we used the Beck Anxiety Inventory – Self-Report (BAI; Beck et al., 1988; De Ayala et al., 2005; Muntingh et al., 2011; Steer et al., 1993); the Fear Questionnaire (FQ; Marks and Mathews, 1979; Oei et al., 1991); and the Penn State Worry Scale (PSWQ; Meyer et al., 1990; Salarifar and Pouretamad, 2012; Verkuil and Brosschot, 2012). Finally, we examined functional (dis)ability, which was measured using the World Health Organization Disability Assessment Schedule (WHO-DAS-II; Chwastiak and Von Korff, 2003). More information regarding these instruments can be found in the **Supplementary Material**.

## *2.3 Statistical Analyses*

*Sociodemographic characteristics.* We reported baseline characteristics as percentages or as means with standard deviations (*SD*) (see **Table 1**). We inspected baseline symptom severity scores for clinically-relevant threshold levels (IDS > 13; BAI > 7; approximations FQ > 30; PSWQ > 48) (Gillis et al., 1995; Rush et al., 1996).

*Categorical approach using diagnoses.* For the categorical approach, we described the DSM-diagnoses trajectories using variable and aggregated percentages. We also examined the trajectories without any missing CIDI data

at all follow-up assessments. Moreover, we conducted a contingency table analysis with adjusted residuals (Beasley and Schumacker, 1995; García-Pérez and Núñez-Antón, 2003), to test whether there was an association between baseline diagnostic subgroup (i.e., depression only, anxiety only, and depression-anxiety) and the four course trajectories at nine-year follow-up.

*Dimensional approach using symptom severity.* For the dimensional approach, we used latent class growth analysis (LCGA), a type of mixture modeling with latent class analysis, to identify subgroups of participants with distinct patterns of change in their longitudinal symptom severity assessments relative to baseline (using the 'hlme' function of the 'lcmm' package in R version 3.6.1) (Proust-Lima et al., 2017). In LCGA, participants are clustered into classes representing trajectories with similar mean levels and mean-level changes of symptom severity by modeling a latent categorical variable. Unlike conventional growth mixture modeling (GMM), the analysis was conducted with the variance of the latent slope and intercept fixed to zero within class, thus limiting the heterogeneity within the latent classes, allowing us to identify course-trajectory classes that were optimally differentiated from each other (Armour et al., 2012; Asparouhov, 2006; Berlin et al., 2014; Cicchetti and Rogosch, 1999; Curran and Hussong, 2003; deRoos-Cassini et al., 2010; Jung and Wickrama, 2008; Muthén and Muthén, 2000; Nylund et al., 2007). For more information, see "Description and interpretation criteria of the latent class growth analysis" and the checklist for the Guidelines for Reporting on Latent Trajectory Studies (GRoLTS; van de Schoot, 2015) in the **Supplementary Material**.

*Comparison of categorical and dimensional approaches.* We compared the outcome of both the categorical and dimensional approaches at nine-year

follow-up. Hereafter, we examined the overlap across the identified dimensional symptom severity pathways and the primary and comparison DSM-diagnosis trajectories at nine-year follow-up. For the comparison, or aggregated, trajectories, where a 3 x 3 contingency table was possible, we tested the level of agreement between the categorical and dimensional approaches with the Kappa statistic as a complete-case analysis, where .01-.20 = none to slight/very poor, .21-.39 = minimal, .40-.59 = weak, .60-.79 = moderate and .80-.90 = strong (Cohen, 1960; de Raadt et al., 2019; McHugh, 2012). We then examined the average functional (dis)ability scores across the identified dimensional symptom severity pathways.

To further compare the symptom severity pathways with the DSM-diagnoses trajectories, we examined the trends of symptom severity per questionnaire across the assessment points based on baseline diagnosis subgroups and functional (dis)ability across the assessment points. We split the IDS-SR, BAI, FQ, PSWQ, and WHO-DAS-II scores into quartiles based on the baseline total score of that particular questionnaire and used mixed model regression analysis to yield marginal mean values at each follow-up assessment per questionnaire. Given that depression and anxiety frequently emerge in younger adulthood (Caspi et al., 2020; Ernst and Angst, 1992; Moffitt et al., 2007), we tested whether there was a potential effect of age and gender on depressive, anxiety, fear and worry severity symptoms. Age was split into quartiles, or four age groups. We then examined these trajectories over the nine-year follow-up.

*Statistics programs for the analyses.* The descriptive analyses and categorical approach analysis were conducted using IBM SPSS version 25.0. For the dimensional approach, we used R statistical software (R version 3.6.0; R

Foundation for Statistical Computing, Vienna, Austria, 2016. URL: <https://www.R-project.org/>). The mixed-model analyses were also conducted using R software using the 'lme4' package. Alpha was set at 0.05, except for the chi-square post hoc contrasts, where alpha was set to 0.0042 after a Bonferroni correction.

### 3. Results

#### 3.1 Baseline sociodemographic information

**Table 1** shows the baseline characteristics of the study sample ( $N = 1701$ ). Participants were on average 41.3 years old ( $SD = 12.4$ ); 67.0% were female ( $n = 1140$ ), and 65.3% had a spouse or partner ( $n = 1110$ ). The mean total IDS and BAI scores were 31.5 ( $SD = 10.9$ ) and 18.3 ( $SD = 10.6$ ), respectively, reflecting a moderate level of depressive and anxiety symptom severity at baseline. Nearly half of our participant sample (44.8%) had comorbid depression and anxiety at baseline.

#### 3.2 Categorical approach using depression and anxiety diagnoses

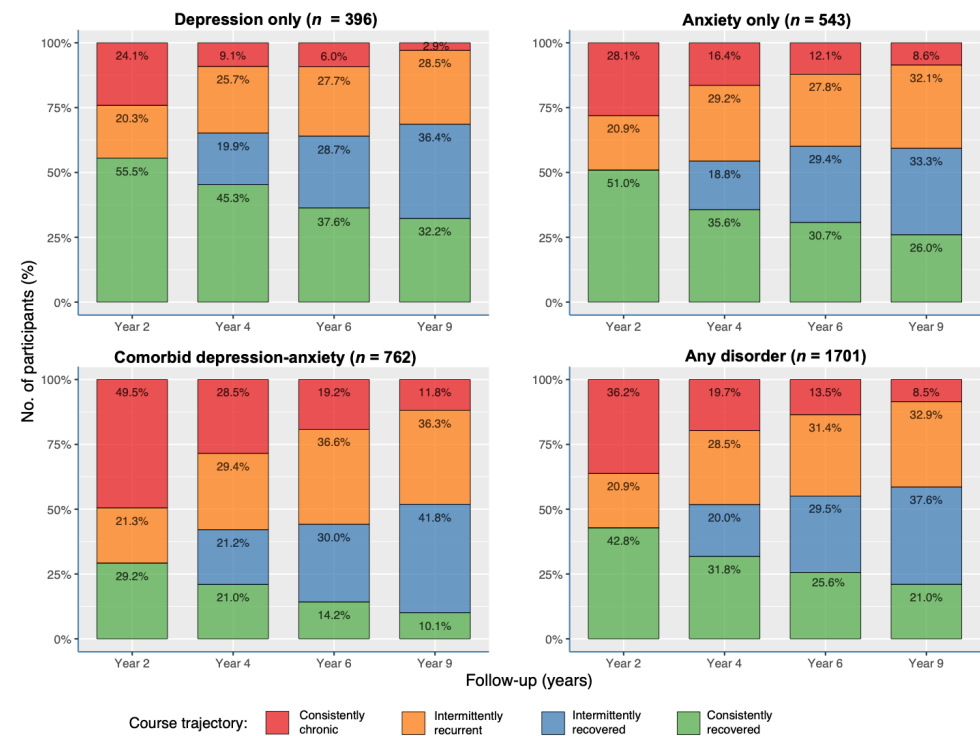
**Figure 1** presents four stacked bar charts showing the proportions of participants at each follow-up assessment who were 'consistently recovered,' 'intermittently recovered,' 'intermittently recurrent,' and 'consistently chronic,' divided by baseline diagnostic status. Over the course of nine years, the prognosis of each group was relatively positive: the percentage of participants who recovered increased, and the percentage of those with consistently chronic diagnoses decreased. Specifically, at nine-year follow-up, across all groups, between 51.9 and 68.6% had recovered from any disorder (when combining both consistent and intermittent recovery) and between 2.9 and 11.8%

maintained consistently chronic disorders. Approximately one-third of participants experienced intermittent recurrence across all groups at four-, six- and nine-year follow-up. Looking specifically at nine-year follow-up, baseline diagnostic status was significantly associated with course,  $\chi^2(6, N = 966) = 61.5, p < .0001$ , with a lower proportion of participants with baseline depression-anxiety comorbidity experiencing consistent recovery than participants with baseline depression ( $p < .0001$ ), and a higher proportion of participants with depression-anxiety comorbidity experiencing consistent chronicity than participants with baseline depression ( $p = .0012$ ). Additionally, when examining recovery more closely at nine-year follow-up, only 10.1% of participants with baseline comorbid depression-anxiety were consistently recovered, 26.0% of the anxiety-only group were consistently recovered, and 32.2% of the depression-only group were consistently recovered. Lastly, for the total group with any disorder, 21.0% were consistently recovered. When using a sample without missing CIDI diagnoses at any follow-up assessment ( $N = 956$ ), we found the same pattern in the categorical trajectories. For comparison with trajectories of Verduijn et al. (2017), see Figure A and “Comparison with previous NESDA study” in the **Supplementary Material**. The main finding of the comparison trajectories is that 78.5% of participants with any disorder experienced recurrence over a nine-year period.

**Table 1.**Baseline Socio-Demographic and Clinical Characteristics (*N* = 1701)

|                                                                    |             |
|--------------------------------------------------------------------|-------------|
| Age, years, mean ( <i>SD</i> )                                     | 41.3 (12.4) |
| Sex, female, % ( <i>n</i> )                                        | 67.0 (1140) |
| Education, years, mean ( <i>SD</i> )                               | 11.8 (3.3)  |
| Married or life partner, yes, % ( <i>n</i> )                       | 65.3 (1110) |
| Depression only, % ( <i>n</i> )                                    | 23.3 (396)  |
| Single depressive disorder, % ( <i>n</i> )                         | 19.9 (339)  |
| Comorbid depressive disorders, % ( <i>n</i> )                      | 3.4 (57)    |
| Severity of depressive symptoms (IDS-SR), mean ( <i>SD</i> )       | 27.6 (11.6) |
| Severity of anxiety symptoms (BAI), mean ( <i>SD</i> )             | 12.1 (8.9)  |
| Severity of fear symptoms (FQ), mean ( <i>SD</i> )                 | 21.7 (16.8) |
| Severity of worry symptoms (PSWQ), mean ( <i>SD</i> )              | 29.3 (15.7) |
| Severity of functional disability (WHO-DAS-II), mean ( <i>SD</i> ) | 31.6 (16.0) |
| Anxiety only, % ( <i>n</i> )                                       | 31.9 (543)  |
| Single anxiety disorder, % ( <i>n</i> )                            | 22.6 (384)  |
| Comorbid anxiety disorders (2 to 3), % ( <i>n</i> )                | 9.3 (159)   |
| Severity of depressive symptoms (IDS-SR), mean ( <i>SD</i> )       | 21.9 (9.8)  |
| Severity of anxiety symptoms (BAI), mean ( <i>SD</i> )             | 14.7 (9.5)  |
| Severity of fear symptoms (FQ), mean ( <i>SD</i> )                 | 31.1 (18.5) |
| Severity of worry symptoms (PSWQ), mean ( <i>SD</i> )              | 29.7 (14.4) |
| Severity of functional disability (WHO-DAS-II), mean ( <i>SD</i> ) | 25.1 (15.4) |
| Comorbid depression and anxiety, % ( <i>n</i> )                    | 44.8 (762)  |
| Severity of depressive symptoms (IDS-SR), mean ( <i>SD</i> )       | 34.2 (13.1) |
| Severity of anxiety symptoms (BAI), mean ( <i>SD</i> )             | 20.8 (11.5) |
| Severity of fear symptoms (FQ), mean ( <i>SD</i> )                 | 38.8 (21.9) |
| Severity of worry symptoms (PSWQ), mean ( <i>SD</i> )              | 32.4 (18.2) |
| Severity of functional disability (WHO-DAS-II), mean ( <i>SD</i> ) | 41.1 (17.2) |

*Note.* Depression only = depressive disorders such as major depressive disorder or dysthymia. Anxiety only = anxiety disorders such as social phobia and generalized anxiety disorder. Comorbid depression and anxiety = MDD, dysthymic disorder and anxiety disorders. *IDS-SR* = Inventory of Depressive Symptomatology – Self-Report; *BAI* = Beck Anxiety Inventory; *FQ* = Fear Questionnaire; *PSWQ* = Penn State Worry Scale; *WHO-DAS-II* = World Health Organization Disability Assessment Schedule (standardized).



**Figure 1.** Categorical trajectories of diagnoses at two-, four-, six-, and nine-year assessments, based on any CIDI affective disorder and LCI data. Depression only = depressive disorders such as major depressive disorder or dysthymia. Anxiety only = anxiety disorders such as social phobia and generalized anxiety disorder.

3.3 Dimensional approach using symptom severity

**Table 2** shows the fit indices resulting from the log-linear LCGA with one- to six-classes.

**Table 2.**  
Fit Indices of One- to Six-Class Latent Class Growth Analysis over a Nine-Year Follow-Up ( $N = 1693$ )

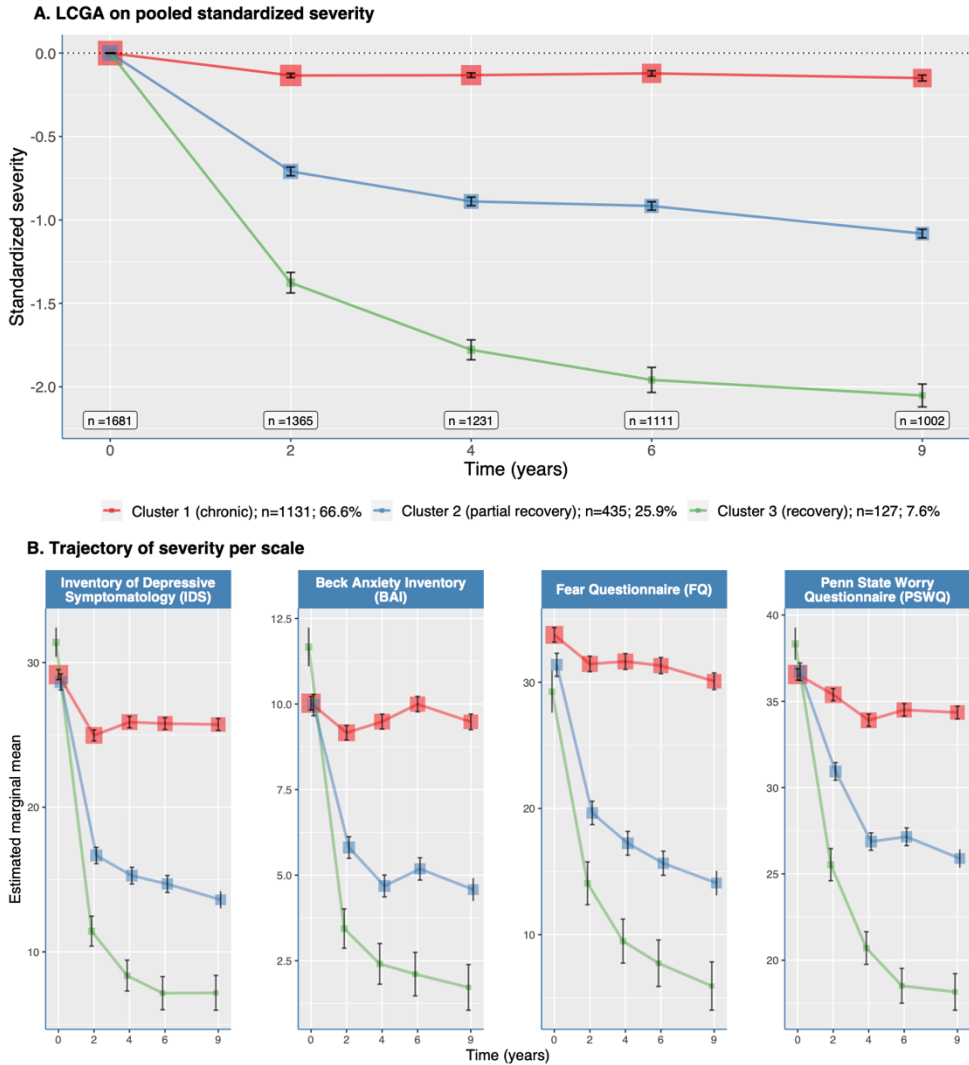
| Classes | Log-Likelihood | AIC     | BIC     | SA-BIC  | Entropy | Percentage of Individuals in Class |      |      |      |      |     |
|---------|----------------|---------|---------|---------|---------|------------------------------------|------|------|------|------|-----|
|         |                |         |         |         |         | 1                                  | 2    | 3    | 4    | 5    | 6   |
| 1       | -5735.8        | 11477.6 | 11493.9 | 11484.3 | 1.00    | 100                                | -    | -    | -    | -    | -   |
| 2       | -4827.9        | 9667.8  | 9700.5  | 9681.4  | 0.71    | 22.1                               | 77.8 | -    | -    | -    | -   |
| 3       | -4598.8        | 9215.6  | 9264.5  | 9235.9  | 0.69    | 7.5                                | 66.6 | 25.9 | -    | -    | -   |
| 4       | -4505.9        | 9035.9  | 9101.1  | 9063.0  | 0.62    | 4.7                                | 14.1 | 18.9 | 62.4 | -    | -   |
| 5       | -4452.1        | 8934.2  | 9015.7  | 8968.1  | 0.65    | 2.4                                | 7.6  | 62.3 | 19.5 | 8.2  | -   |
| 6       | -4427.9        | 8891.8  | 8989.7  | 8932.5  | 0.66    | 1.2                                | 2.4  | 61.4 | 7.5  | 19.8 | 7.8 |

Note. AIC = Akaike Information Criterion (Akaike, 1987); BIC = Bayesian Information Criterion (Schwartz, 1978); SA-BIC = sample-size-adjusted BIC (Scllove, 1987); Entropy (Ramasmamy et al., 1993).

Taking all fit indices into account, the three-class model was considered the optimal solution. The AIC and SA-BIC values showed a marked drop between one-, two- and three-class models (see elbow plots in **Figure B** in **Supplemental Material**), and hereafter, the differences in AIC and SA-BIC values were smaller, suggesting that the addition of more classes did not further improve the model. Also, solutions with four to six classes had very small numbers of participants (<5%) in some classes. Examining the entropy value of the three-class model (0.69) indicated that a moderate to high proportion of participants were correctly classified. The grid search also confirmed a three-class model.

**Figure 2A** delineates the three latent classes that were identified using pooled standardized mean severity scores of depressive, anxiety, fear, and worry symptom questionnaires: class 1: *chronic* (66.8%); class 2: *partially recovered* (25.7%); and class 3: *(sustained) recovered* (7.5%). Class 1 (red) remained stable and severe; whereas symptom levels decreased moderately in class 2 (blue) and markedly in class 3 (green). This was consistent with the latent classes of pooled symptom severity for depression, anxiety, fear, and worry symptom levels, viewing each questionnaire separately (see **Figure 2B**): the group with the highest symptom levels maintained high symptom levels throughout the nine-year period; the moderate level group showed slow decline and intermittent increases in anxiety and worry; and the lowest symptom level group showed a larger and steady decline.

NINE-YEAR CLINICAL COURSE OF DEPRESSIVE AND ANXIETY DISORDERS (NESDA)



**Figure 2.** Three latent class solution of pooled standardized mean severity scores of depressive, anxiety, fear, and worry symptoms (LCGA: latent class growth analysis). Panel A shows the average change in standardized severity over time as compared to baseline severity, and Panel B shows the standardized severity scores per scale over time. Error bars represent standard errors, and the box size is proportional to the number of participants.

Moreover, inspection of the classes, or symptom pathways, showed that there was a relatively greater change in symptom severity between baseline and the first follow-up, after which the course showed less change between follow-up assessments and began to stabilize, suggesting that the log-linear function best described these symptom pathways. For a comparison with alternative models, see “Alternative models using latent class growth analysis” in the **Supplementary Material**. Recomputing the model without any missing data and comparing these to the abovementioned classes resulted in highly corresponding models: 94.9% similarity for class 1 (chronicity), 89.9% similarity for class 2 (partial recovery), and 100% similarity for class 3 (recovery).

### *3.4 Comparison of categorical and dimensional approaches*

We compared the diagnosis trajectories of the total sample using the categorical approach at nine-year follow-up with the three identified symptom severity pathways of the dimensional approach. The ‘intermittently recurrent’ trajectory and the “partially recovered” pathway corresponded relatively well (32.9% versus 25.7%). However, the percentages of the ‘(consistently) chronic’ and ‘(consistently) recovered’ trajectories did not align. In the categorical approach compared to the dimensional approach, chronicity was lower (8.5% versus 66.8%), and (combined) recovery was higher (58.6% versus 7.5%). The ‘recovered’ dimensional pathway appeared to correspond more to the ‘consistently recovered’ diagnosis trajectory (10.1%) at nine-year follow-up.

We then compared the primary and comparison DSM-diagnosis trajectories and three identified symptom severity pathways using crosstabulation at nine-year follow-up (see **Table 3**).

**Table 3.**

Comparison of the Three Identified Dimensional Symptom Severity Pathways of the Latent Class Growth Analysis (LCGA) with the Three and Four Categorical Diagnosis Trajectories based on Overlap and Disability at Nine-Year Follow-Up

| Categorical diagnoses trajectories <sup>a</sup> | Dimensional symptom severity pathways or LCGA classes |                     |             |
|-------------------------------------------------|-------------------------------------------------------|---------------------|-------------|
|                                                 | Recovered                                             | Partially Recovered | Chronic     |
| Consistently Recovered                          | 33                                                    | 92                  | 78          |
| <i>% within categorical trajectories</i>        | 16.3%                                                 | 45.3%               | 38.4%       |
| <i>% within LCGA classes</i>                    | 52.4%                                                 | 29.7%               | 13.2%       |
| Intermittently Recovered                        | 25                                                    | 135                 | 203         |
| <i>% within categorical trajectories</i>        | 6.9%                                                  | 37.2%               | 55.9%       |
| <i>% within LCGA classes</i>                    | 39.7%                                                 | 43.5%               | 34.2%       |
| Intermittently Recurrent                        | 5                                                     | 75                  | 238         |
| <i>% within categorical trajectories</i>        | 1.6%                                                  | 23.6%               | 74.8%       |
| <i>% within LCGA classes</i>                    | 7.9%                                                  | 24.2%               | 40.1%       |
| Consistently Chronic                            | 0                                                     | 8                   | 74          |
| <i>% within categorical trajectories</i>        | 0.0%                                                  | 9.8%                | 90.2%       |
| <i>% within LCGA classes</i>                    | 0.0%                                                  | 2.6%                | 12.5%       |
| Categorical diagnoses trajectories <sup>b</sup> | Recovered                                             | Partially Recovered | Chronic     |
| Recovered                                       | <b>33</b>                                             | 92                  | 78          |
| <i>% within LCGA classes</i>                    | 40.7%                                                 | 24.3%               | 9.0%        |
| <i>% within categorical trajectories</i>        | 16.3%                                                 | 45.3%               | 38.4%       |
| Recurrent                                       | 48                                                    | <b>278</b>          | 715         |
| <i>% within LCGA classes</i>                    | 59.3%                                                 | 73.5%               | 82.5%       |
| <i>% within categorical trajectories</i>        | 4.6%                                                  | 26.7%               | 68.7%       |
| Chronic                                         | 0                                                     | 8                   | <b>74</b>   |
| <i>% within LCGA classes</i>                    | 0.0%                                                  | 2.1%                | 8.5%        |
| <i>% within categorical trajectories</i>        | 0.0%                                                  | 9.8%                | 90.2%       |
| Disability (WHO-DAS-II), mean (SD)              | 5.3 (7.9)                                             | 13.9 (12.6)         | 28.8 (17.6) |

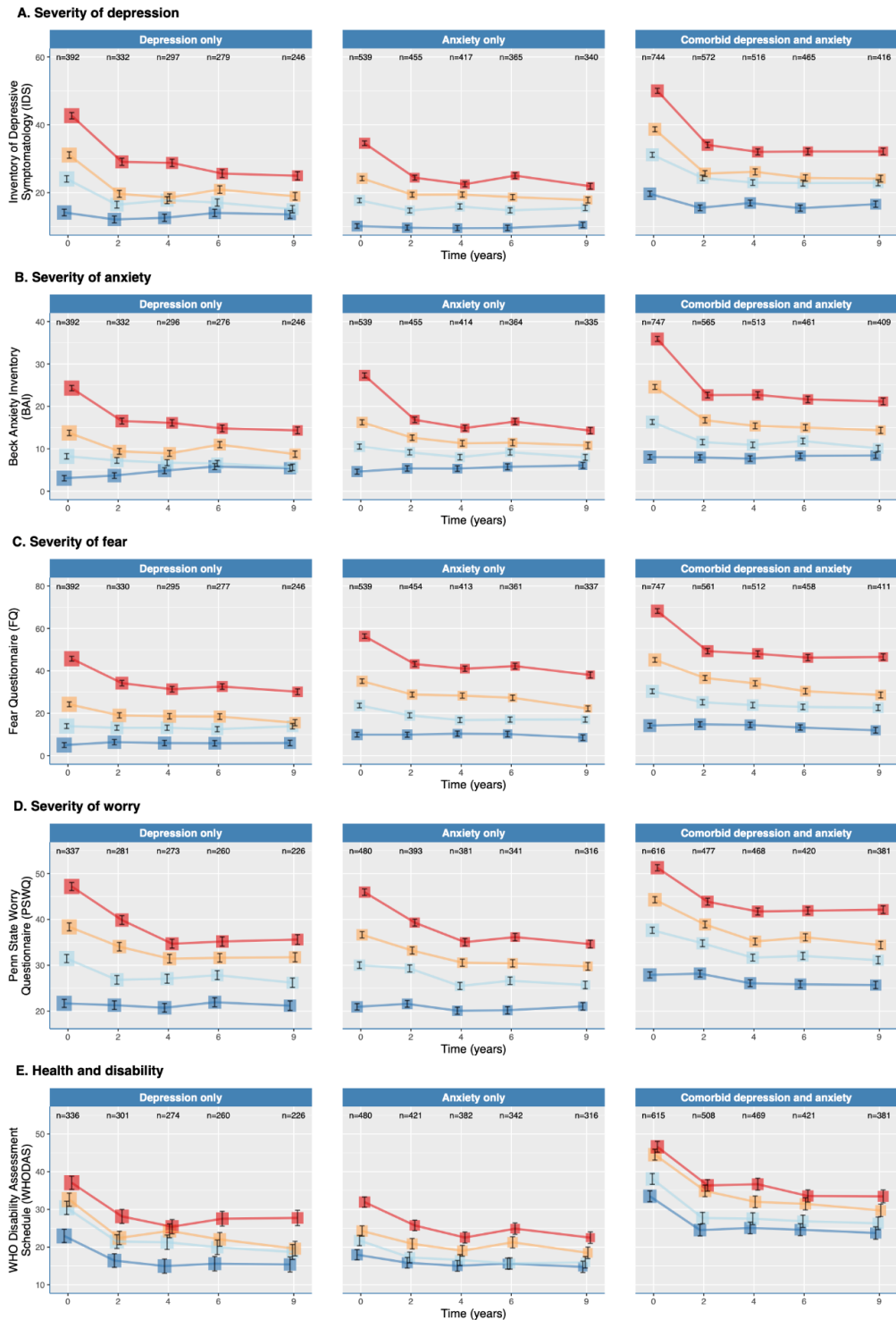
<sup>a</sup>4 primary trajectories of the categorical approach,  $N = 966$ .

<sup>b</sup>3 comparison/aggregated trajectories of the categorical approach,  $N = 1326$ .

In **bold** = overlap between the aggregated categorical trajectories and dimensional pathways.  
 WHO-DAS-II = World Health Organization disability assessment schedule.

First, we looked at the four primary/variable DSM-diagnosis trajectories. Participants who were 'consistently recovered,' 'intermittently recovered,' and 'intermittently recurrent' from a DSM diagnosis were spread across all three symptom severity pathways. No clear pattern of correspondence emerged. Second, we looked at the three comparison/aggregated DSM-diagnosis trajectories with the symptom severity pathways, and the overlap of was low, with very poor agreement,  $\kappa = .02$ . Participants from the categorical 'recovered' trajectory, for example, were also allocated to the 'partially recovered' and 'chronic' dimensional pathways. Regarding functional disability, participants in the 'chronic' symptom severity pathway experienced the highest functional disability, followed by the 'partial recovery', then 'recovered' pathway.

Finally, we tested whether there was a potential effect of age in quartiles and gender on depressive, anxiety, fear and worry severity symptoms (see **Figure C** in **Supplemental Material**). There were age group and gender differences, with age having a slightly higher effect on severity symptoms. **Figure 3** presents the trends of depressive (A), anxiety (B), fear (C), and worry (D) severity symptoms for subsamples of participants with depression only, anxiety only, or comorbid depression-anxiety, based on baseline severity scores split into quartiles. These were adjusted for the effects of age and gender. However, the adjusted mixed model regression analyses resulted in similar trajectories as the non-adjusted analyses.



**Figure 3.** Trajectories of depressive, anxiety, fear, and worry symptoms, controlled for the effect of age and gender. Error bars represent standard errors, and the box size is proportional to the number of participants.

Throughout the nine-year period, those with the highest baseline severity levels (red line) on average maintained the highest severity scores, remaining above clinically-relevant threshold levels. In general, all baseline quartiles had trends that ran rather parallel over time and remained stable after an initial drop in severity at two-year follow-up. Furthermore, comparing the three subsamples, those with comorbid depression-anxiety at baseline appeared to have the highest scores across all symptom measures compared to the depression only and anxiety only groups.

Finally, to assess the trends of functional disability, we reviewed the WHO-DAS-II scores at each follow-up assessment over the nine-year period for subgroups of depression only, anxiety only, and comorbid depression-anxiety based on baseline total scores split into quartiles (see **Figure 2E**). Those with comorbid depression-anxiety appeared to have higher disability than those with depression only or anxiety only.

#### **4. Discussion**

Our findings show that using DSM diagnoses alone fails to fully capture the persistence of depressive and anxiety symptoms over time. The categorical trajectories using diagnoses were not congruent with results of the identified dimensional pathways using symptom severity, further emphasizing the need to better capture clinical course (Batelaan et al., 2014). In other words, diagnostic recovery was not equivalent to symptomatic recovery. When using the categorical approach with any depressive or anxiety diagnosis, approximately 8% were consistently chronic, 33% had intermittently relapsed, 38% had intermittently recovered, and 21% remained consistently recovered at nine-year follow-up, showing a higher estimation of recovery compared to the dimensional

approach. However, when we examined participants with baseline comorbid depression and anxiety, only 10.1% remained consistently recovered at nine-year follow-up, which paralleled the 7.5% who were recovered using the dimensional approach. Additionally, compared to the categorical trajectories of comorbid depression and anxiety, the chronicity of the dimensional symptom severity pathway was much higher (11.8% versus 66.8%). When using aggregated trajectories, we found that 78.5% of participants with any disorder experienced recurrence over a nine-year period. This was much higher than the 25.7% who were partially recovered when using the dimensional approach.

The discrepancy between the trajectories could be explained by the following reasons.

First, the discrepancy may be due to the inherent differences of the methods used to assess the dimensional versus the categorical outcomes. Specifically, the dimensional approach used self-report questionnaires to determine the change in symptom severity over time while the categorical approach used a structured interview to establish DSM diagnoses and examine them over time. Although the self-report questionnaires encompassed the same symptoms as the depression and anxiety syndromes in the structured interview, the questionnaires assessed a shorter duration period of one to two weeks. For the diagnoses, we allowed a longer recency period of six months. The diagnosis, therefore, could no longer be present and might have been present up to five and a half months ago. Moreover, in the structured interview, the trained interviewer examined the number of symptoms, duration of the symptoms, and the patient's level of functioning. Thus, if a patient had the required number of symptoms yet a higher level of functioning, a DSM diagnosis was not recorded. At the outset, due to the specific diagnostic criteria, using a categorical

approach with DSM diagnoses may be less inclusive when compared to the dimensional approach, which may have resulted in higher recovery rates and lower chronicity. Because of the inclusion of the functional disability criterion, we consider DSM diagnoses to be a better indicator of psychological suffering than self-report questionnaires. Second, the discrepancy in results could be due to enduring, fluctuating subthreshold affective symptoms and the higher prevalence of individuals experiencing recurrence of diagnoses. Those with remitted affective diagnoses, for instance, may still suffer from subthreshold disease (e.g., minor depression), or they could have been assessed during a short period in which affective symptoms had slightly receded (Batelaan et al., 2014; Torpey and Klein, 2008). Previous research has shown that, compared to those without symptoms, those with subthreshold symptoms relapsed sooner (Cuijpers and Smit, 2004; Fawcett, 1994; Judd et al., 1998; Karsten et al., 2011) and were found to suffer more chronic episodes and fewer symptom-free weeks (Arnow and Constantino, 2003; Judd and Akiskal, 2000). Other studies have also found a recurrent course with (non)chronic episodes in at least 50% of participants (Bobes et al., 2018; Bruce et al., 2005; Scholten et al., 2016; Scholten et al., 2013; Verduijn et al., 2017; Yonkers et al., 2003) and up to 80% (Judd, 1997).

Furthermore, when using the categorical approach, as expected, those with comorbid depressive and anxiety disorders had a relatively poorer long-term course when compared to those baseline depressive disorders only. This finding is in line with previous research examining anxiety and (comorbid) depressive disorders (Batelaan et al., 2014; Bruce et al., 2005; Rhebergen et al., 2011; ter Meulen et al., 2021). According to Batelaan et al. (2014),

however, baseline severity, duration of symptoms, and disability appear to be better indicators of a poor prognosis than DSM-categories.

### *Clinical significance of the results*

Kraepelinian nosology (Kraepelin, 1921) emphasizes that course, functional outcome, and etiology support a dimensional approach of depression and anxiety. In light of our findings, the question arises whether we should view individuals with comorbidity as having two or more distinct DSM disorders or as having a single dimensional disorder, or symptoms on a continuum, in which myriad etiological factors may result in diverse syndromes that are modified with time and environmental exposures (Angst and Wicki, 1991; Cardno et al., 2002; Hyman, 2010; Kendler et al., 1992; Krueger and Markon, 2006). Previous studies (Fried and Nesse, 2015b; Fried et al., 2014; Lux and Kendler, 2010; van Eeden et al., 2019) have identified risk factors, such as neuroticism and baseline chronicity, that affected individual depressive symptoms to varying degrees, suggesting that depression is not one unified latent construct. On the other hand, abandoning the current DSM criteria would reduce reliability and leave psychiatry without a common language (Kendell and Jablensky, 2003; Patten, 2015). Therefore, until a better dimensional model is proposed, we suggest the integrative use of diagnostic classification, symptom severity, symptom duration, and levels of impairment (Batelaan et al., 2014; Hyman, 2010; Patten, 2015). An example is the Activity, Cognition, and Emotion (ACE) model that groups symptoms commonly present in mood disorders like depression and bipolar disorder according to functional domains (Malhi et al., 2018). Another option is the symptom-based framework which looks at individual symptoms and how they influence each other (Fried, 2015). However, more research is

needed to confirm whether a dimensional symptom-oriented tool will lead to more accurate symptom-specific tailored treatment plans and better outcomes. Moreover, clinicians may consider the adoption of longer-term treatment strategies (Vos et al., 2004) that may include relapse prevention strategies and psychiatric rehabilitation (i.e., functional recovery) for those with high levels of subthreshold symptoms and a high risk of recurrence of disorders over nine years. Well-designed interventions studies looking into the types and effectiveness of relapse prevention strategies are warranted.

### *Strengths and Limitations*

This study has some noteworthy strengths. First, NESDA was designed to gain more insight into the long-term clinical course of depression and anxiety in a large cohort study with access to the full range of depression and anxiety disorders from primary and secondary care. Patients were rigorously diagnosed using diagnostic interviews and followed up over a nine-year time period with multiple assessment points. Moreover, rather than investigating the course of single MDD or anxiety disorders, we examined variable course categories that allowed comorbidity and diagnostic switching, which may be more ecologically and clinically relevant. To strengthen our conclusions, we used the data-driven LCGA method, to distinguish groups with similar symptom severity trajectories, which handles missing data rather well. The findings were further confirmed by inspecting symptom severity scores, diagnoses, and functional (dis)ability. Additionally, we compared our models to those without missing data, and results were not significantly altered.

With regard to our study limitations and research recommendations, the following may be mentioned. First, the current study used data from NESDA, which concerns DSM-IV criteria rather than the newest DSM-5 criteria. In the DSM-5, there is a duration criterion for social phobia and agoraphobia, which could be useful in excluding patients with only transient fears (Batelaan et al., 2014). Second, NESDA was set up as an observational, naturalistic study, and therefore, the effects of treatment on outcomes could not be examined (e.g., due to confounding by indication effects). While general data were collected regarding medication use, treatment duration, and setting (e.g., primary or secondary care, or psychotropic use), data on the specifics of type, duration, and intensity of psychotherapeutic interventions were not collected. Third, to maximize our sample, we selected participants with a diagnosis in the six months prior to baseline, a period during which some participants may have remitted. Fourth, NESDA did not include participants with obsessive compulsive disorder (OCD), bipolar disorders, or other common mental disorders, which participants may also be experiencing, thereby potentially increasing prevalence rates of chronicity. Fifth, our sample included participants recruited from the community, primary care and specialized mental health care, resulting in a heterogeneous cohort (Patten, 2015; Regier et al., 1998). Thus, our results may not be generalizable to each population. Sixth, at each follow-up there was a selective loss of participants, which were those with the highest baseline severity levels. Thus, chronicity may have been underestimated. Seventh, the Life Chart Interview (LCI), which relies on the recall of past memories, was used to establish the continuity of symptoms between assessment points in the categorical approach. Recall error of autobiographical information may have had an impact on the accuracy of the sustained recovered or consistently chronic

categorical trajectories (Drasch and Matthes, 2013; Reimer and Matthes, 2007). Although the LCI uses an event history calendar with landmark events, which has been shown to be an effective tool for the reliable retrieval of older memories (Belli, 1998; Belli et al., 2001; Drasch and Matthes, 2013; Vaart and Glasner, 2011), memory is not always reliable, and any results using retrospective recall should be carefully considered (Glasner and Vaart, 2009; Patten et al., 2010). Eighth, NESDA used the CIDI to diagnose DSM-IV disorders. However, there is an alternative semi-structured clinical interview, namely the Structured Psychopathological Interview and Rating of the Social Consequences for Epidemiology (SPIKE) (Angst and Dobler-Mikola, 1985; Angst et al., 2009; Angst and Wicki, 1991; Ernst and Angst, 1992), which has a lower threshold than the CIDI to meet diagnostic criteria. The strict criteria of the CIDI may make it less inclusive, potentially leading to an underestimation of chronicity levels in the current study. Ninth, LCGA classes were calculated primarily with log-linear functions and with too few assessments. Therefore, our dimensional approach may not have been able to fully describe the “waxing and waning” of symptom severity. However, examining alternative models with other shape trajectories corresponded highly with the log-linear primary model. Tenth, while we did examine the functional disability in the dimensional approach using mixed model regression analyses, functional disability was not included in the main LCGA due to bias. Future research using a longitudinal design and a dimensional approach that includes both rater-based versus self-rated symptom scales would be of value and may help improve the prediction of the clinical course.

## **5. Conclusion**

This study showed that, when taking comorbidity, switching between diagnoses, and symptom duration into consideration to examine the long-term course of depression and anxiety, using categorical diagnoses alone to describe clinical course led to a high estimation of recovery, low estimation of chronicity, and appeared to inadequately capture the persistence of affective symptoms. Discrepancies in the clinical course using DSM-categories or symptom severity could be explained by the enduring, fluctuating presence of subthreshold affective symptoms that do not meet diagnosis criteria but still may lead to recurrence of disorders. The commonness of subthreshold symptoms and their adverse impact on long-term prognoses deserve continuous clinical attention in mental health care as well further research.

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**SUPPLEMENTARY MATERIAL****I- Instruments**

***Composite Interview Diagnostic Instrument (CIDI).*** The CIDI is a standardized diagnostic interview that is used worldwide to classify DSM-IV psychiatric diagnoses. It has been shown to have high interrater reliability, high test-retest reliability, and high validity for depressive and anxiety disorders (Andrews and Peters, 1998; Haro et al., 2006; Wittchen, 1994; Wittchen et al., 1991).

***Life Chart Interview (LCI).*** The LCI uses both age- and calendar-linked personal landmark events and an interactive life event chart to aid in memory recall. The methodology of the LCI has been shown to have high validity and reliability (Warshaw et al., 1994). Examining diagnosis-related symptoms was conditional upon meeting DSM-criteria for a psychiatric disorder as per the CIDI at an assessment point. Specifically, for each month between assessments, the presence of depressive and/or anxiety diagnostic symptoms was determined. If diagnostic symptoms were reported during  $\geq 85\%$  of the months between assessments (i.e., no more than two months without symptoms), we considered this continuous or chronic symptomatology and confirmation of the continuous presence of a psychiatric disorder.

***Inventory of Depressive Symptomatology - Self-Report (IDS-SR).*** The 30-item IDS-SR measures depressive symptoms regarding the previous week on a four-point Likert scale (0-3) (Rush et al., 1996; Trivedi et al., 2004). The IDS-SR has shown adequate internal consistency (Cronbach's alpha ranged from 0.92 to 0.94), test-retest stability, as well as adequate convergent and discriminant validity (Rush et al., 1996).

**Beck Anxiety Inventory self-report (BAI).** When completing the 21-item BAI, participants rated how much each anxiety symptom troubled them in the past week, on a four-point Likert scale (0-3). Total scores range from 0 to 63 (Muntingh et al., 2011). The BAI has been shown to have high internal validity (Cronbach's alpha ranged from 0.92 to 0.94) (Beck et al., 1988; Fydrich et al., 1992; Spinhoven et al., 2016) and acceptable test-retest reliability over one week, and good discriminant and convergent validity (Beck et al., 1988; Fydrich et al., 1992). **Fear Questionnaire (FQ).** When completing the 15-item FQ, participants reported how much they endorsed statements related to agoraphobia, blood/injury phobia, and social phobia. These subscales were previously shown to have moderate to high internal consistency (Cronbach's alpha ranged from 0.71 to 0.83) (Oei et al., 1991). Total scores range from 0 to 120 (Lee and Oei, 1994), with the total scale showing moderate high internal consistency (Cronbach's alpha = 0.83) and adequate reliability (Lee and Oei, 1994). The FQ has high retest reliability (Marks and Mathews, 1979).

**Penn State Worry Scale (PSWQ).** When completing the 16-item PSWQ, participants rated how much they endorsed statements related to their disposition to worry, including the frequency and burden of the worry complaints, on a five-point Likert scale (1-5) (Meyer et al., 1990). Total scores range from 16 to 80, with higher scores indicative of higher levels of worry. The PSWQ was previously found to have high moderate to high internal consistency (Cronbach's alpha ranged from 0.83 to 0.93) (Brown et al., 1992; Salarifar and Pouretmad, 2012) as well as good test-retest reliability and good convergent, discriminant, and criterion-related validity (Brown et al., 1992; Davey, 1993; Meyer et al., 1990).

**World Health Organization Disability Assessment Schedule (WHO-DAS-II).**

We used the 32-item total score of the WHO-DAS-II and excluded the four-item 'work disability' scale considering the high percentage of patients without work in this population (Bokma et al., 2017; Rizvi et al., 2015; Verduijn et al., 2017; Zimmerman and Katon, 2005). Total scores ranged from 0 to 100, with higher scores reflecting greater disability (Bokma et al., 2017). The total WHO-DAS-II has been shown to have excellent internal consistency (Cronbach's alpha ranged from 0.92 and 0.95) (Bokma et al., 2017; Chwastiak and Von Korff, 2003) and convergent validity (Chwastiak and Von Korff, 2003).

**II- Description of categorical trajectories**

We used the following definitions for the trajectories of the main analysis ("primary trajectories"):

- **Consistently recovered** (*fixed diagnosis-free period*): participants with no current six-month CIDI diagnosis at two-year follow-up nor at any subsequent follow-up assessment;
- **Intermittently recovered** (*variable diagnosis-free period*): participants with either (A) no current six-month CIDI diagnosis at four-, six-, or nine-year follow-up, preceded by any CIDI diagnosis with LCI symptoms (i.e., in the recurrent or chronic trajectory) at the two-, four-, six-year follow-up period, respectively, or (B) no current six-month CIDI diagnosis at six- or nine-year follow-up, preceded by a delayed recovery course at the four- or six-year follow-up period, respectively;
- **Intermittently recurrent** (*variable diagnosis-present period*): either (A) any current six-month CIDI diagnosis at two-year follow-up with discontinuous (LCI) symptoms in the preceding period, or (B) any six-

month CIDI diagnosis at four-, six-, or nine-year follow-up assessments, which was preceded by a recurrent episode (see 3A) or recovery period at two-, four-, or six-year follow-up, respectively;

- **Consistently chronic** (*fixed diagnosis-present period*): any current six-month CIDI diagnosis at each follow-up assessment with continuous preceding (LCI) symptoms between all assessment periods.

The “fixed” trajectories integrated each subsequent assessment point from two- to nine-year follow-up, resulting in aggregate percentages of participants with a consistent course trajectory over time, and the “variable” trajectories allowed switching to another course trajectories across the assessment points (course variability).

### III- Comparison with previous NESDA study

Our procedure was based on that of Verduijn et al. (2017) but deviated in the following ways: we did not incorporate the LCI (hypo)mania or avoidance symptoms; we did not define an additional trajectory of recurrence with chronic episodes; we integrated comorbidity and diagnostic switching across the assessment points, defining different diagnostic groups at baseline; and we allowed transitioning or course variability between trajectories across the assessment points, with two variable ‘intermittent recovery’ and ‘intermittent recurrence’ trajectories. To compare with previous results of Verduijn et al. (2017), we classified three aggregated trajectories across the follow-up assessment points, based on the CIDI diagnoses and LCI symptom duration: ‘recovered’, ‘recurrence’, and ‘chronicity’ (“comparison trajectories”). For the purposes of comparison, we used the following definitions for the aggregated

trajectories, where each percentage was aggregated at each subsequent assessment point from two- to nine-year follow-up:

- **Recovered:** participants with no current six-month CIDI diagnosis at two-year follow-up and at all subsequent follow-up assessments thereafter;
- **Recurrent:** participants experiencing a recurrent six-month CIDI diagnosis at any follow-up assessment;
- **Chronic:** participants with any current six-month CIDI diagnosis at each follow-up assessment with continuous or chronic preceding (LCI) symptoms between all assessment periods.

When calculating the aggregated trajectories based on Verduijn et al. (2017) and comparing the trajectories with those of the main analysis, we found at the nine-year follow-up in the any disorder group that there were slightly fewer participants with a chronic trajectory (6.2% vs 8.5%), and fewer with a sustained recovery trajectory (15.3% vs. 21.0%) (see *Figure A in Supplemental Material*). We also found that 78.5% of participants with any disorder experienced recurrence over a nine-year period. Therefore, at each subsequent follow-up, participants either did not meet the chronicity condition for the chronic trajectory nor the diagnosis-free condition for the sustained recovery trajectory.

#### **IV- Description and interpretation criteria of the latent class growth analysis**

For the latent class growth analysis (LCGA), all depression and anxiety severity scores were log-transformed to enable estimation of the log-linear relationships between severity and time. The scores were then standardized to

z-scores to allow comparison of the relative position of the scores within the distribution of each scale, after which these four z-scores were averaged per time point. Baseline values were centered to zero to remove the impact of baseline severity on the prognostic course, thus calculating the degree of change over the two-, four-, six- and nine-year follow-up period in relation to baseline. The aforementioned composite (pooled standardized mean) severity scores were analyzed using LCGA (Andruff et al., 2009; Jung and Wickrama, 2008; van de Schoot, 2015) to estimate one- to six-class models, using full information maximum likelihood, assuming that the missing data were missing at random (MAR). We examined a log-linear model as the primary model. This was compared to other alternative models with other trajectory shapes: quadratic, cubic, beta, and spline with three knots. Additionally, to test for the impact of missing data, we recalculated the LCGA classes without any missing data and compared the resulting trajectories. All models were estimated with 100 random starts to prevent solutions at a local maximum and 10 iterations in the optimization algorithm (Biernacki et al., 2003). Because LCGAs are susceptible to converging on local maximum, rather than global solutions, we conducted a grid search (Biernacki et al., 2003; Proust-Lima et al., 2017).

To determine which model best fitted the data, we inspected the log-likelihood, Akaike Information Criterion (AIC; Akaike, 1987), Bayesian Information Criterion (BIC; Schwarz, 1978), sample-size-adjusted BIC (SA-BIC; Sclove, 1987), entropy values (Ramaswamy et al., 1993), group membership probabilities, and posterior probabilities. Guidelines state that with reference to the AIC, BIC, and SA-BIC, the lower the values, the superior the fit (Armour et al., 2012; Jung and Wickrama, 2008; Muthén, 2003). As more classes are added to the model, fit tends to improve when the AIC, BIC, and SA-BIC values

continue to fall. If the difference between the values of one additional class is small, an additional class is said to add little to the model (DiStefano and Kamphaus, 2006). On the basis of parsimony, the solution with the fewest classes should be accepted (Armour et al., 2012). Moreover, Ramaswamy et al. (1993) reported that high entropy values indicate good classification, with a value of 1.0 indicating perfect model fit and classification. Interpretability and sufficient class size (all > 50) were also used as criteria. After identification of the best model using the aforementioned criteria, we examined how participants were assigned to classes by inspecting group membership probabilities and posterior probabilities using maximum likelihood techniques.

**V- The compliance in this manuscript with guidelines for reporting latent trajectory study, according to GRoLTS Checklist (Van de Schoot et al., 2017):**

1. Is the metric of time used in the statistical model reported? Yes.
2. Is information presented about the mean and variance of time within a wave?  
*No, but it is represented in the figure.*
- 3a. Is the missing data mechanism reported? Yes.
- 3b. Is a description provided of what variables are related to attrition/missing data? Yes.
- 3c. Is a description provided of how missing data in the analyses were dealt with? Yes.
4. Is information about the distribution of the observed variables included? Yes.
5. Is the software mentioned? Yes.
- 6a. Are alternative specifications of within-class heterogeneity considered (e.g., LGCA vs. LGMM) and clearly documented? *We did not explore within-class*

*heterogeneity as the primary model. We did explore other models with alternative shapes.*

6b. Are alternative specifications of the between-class differences in variance-covariance matrix structure considered and clearly documented? *In the LCGA the variance and covariance were assumed to be fixed to be zero.*

7. Are alternative shape/functional forms of the trajectories described? *Yes.*

8. If covariates have been used, can analyses still be replicated? *No covariates.*

9. Is information reported about the number of random start values and final iterations included? *Yes.*

10. Are the model comparison (and selection) tools described from a statistical perspective? *Yes.*

11. Are the total number of fitted models reported, including a one-class solution? *Yes.*

12. Are the number of cases per class reported for each model (absolute sample size, or proportion)? *Yes, proportion.*

13. If classification of cases in a trajectory is the goal, is entropy reported? *Yes.*

14a. Is a plot included with the estimated mean trajectories of the final solution? *Yes.*

14b. Are plots included with the estimated mean trajectories for each model? *Not for each model due to limited number of figures.*

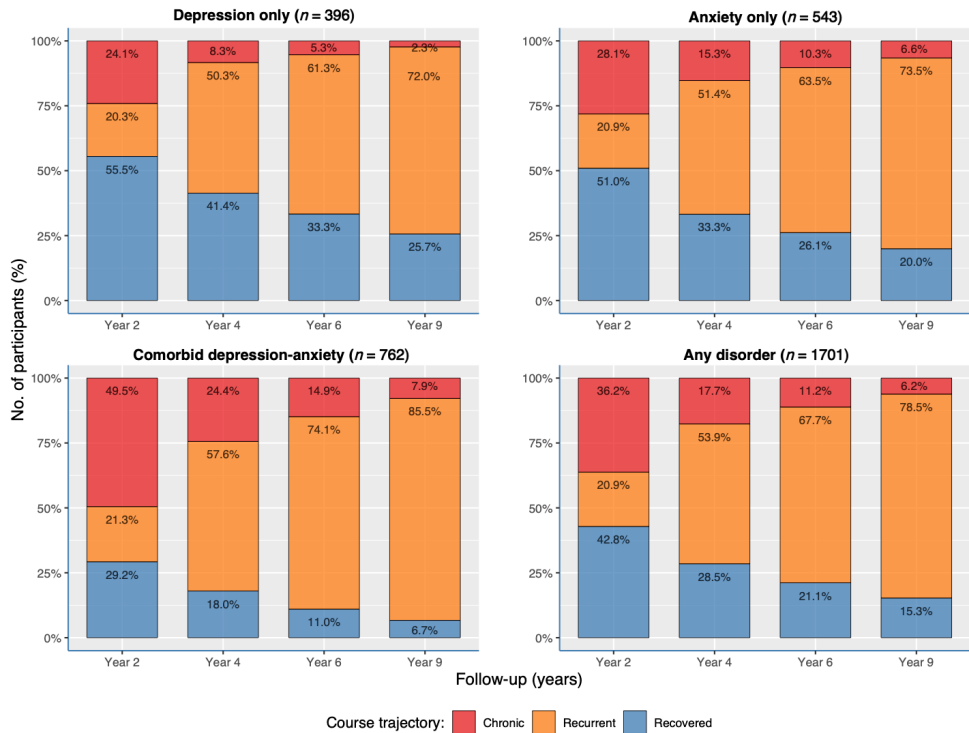
14c. Is a plot included of the combination of estimated means of the final model and the observed individual trajectories split out for each latent class? *We only show the individual trajectories as this is our goal in the paper.*

15. Are characteristics of the final class solution numerically described (i.e., means, SD/SE, n, CI, etc.)? *No, due to the word- and table limits of our manuscript. However, the figure and table show the final class solution.*

16. Are the syntax files available (either in the appendix, supplementary materials, or from the authors)? *They are available from the first author (Solis, corresponding author).*

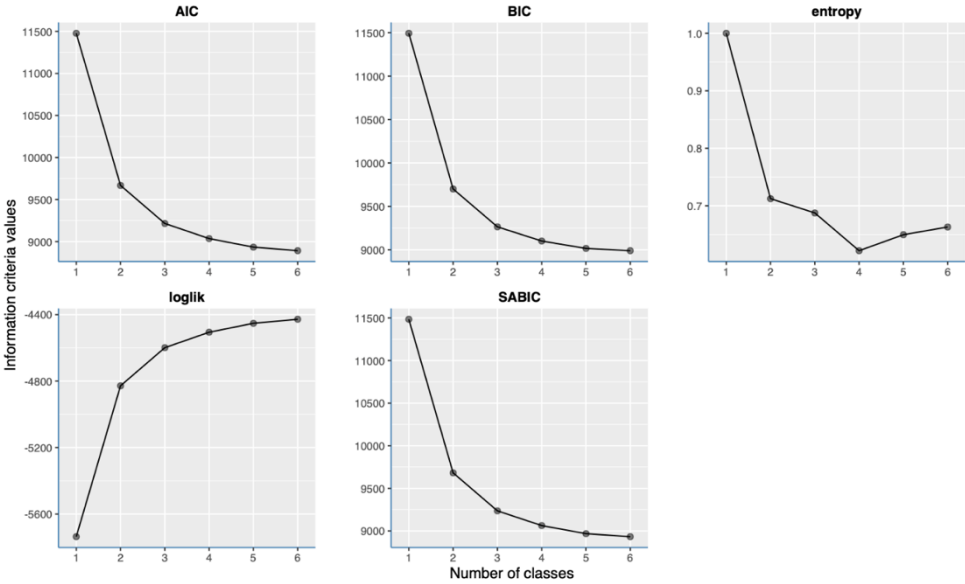
## **VI- Alternative models using latent class growth analysis**

Comparing the primary log-linear model to the alternative models resulted in high percentages of similarity across the classes, with approximately 96% similarity for class 1, 92% similarity for class 2, and 92% for class 3 in the quadratic and cubic models, and approximately 84% similarity for class 1, 81% similarity of class 2, and 100% for class 3 in the beta and 3-knot spline models. We thus retained the log-linear model.

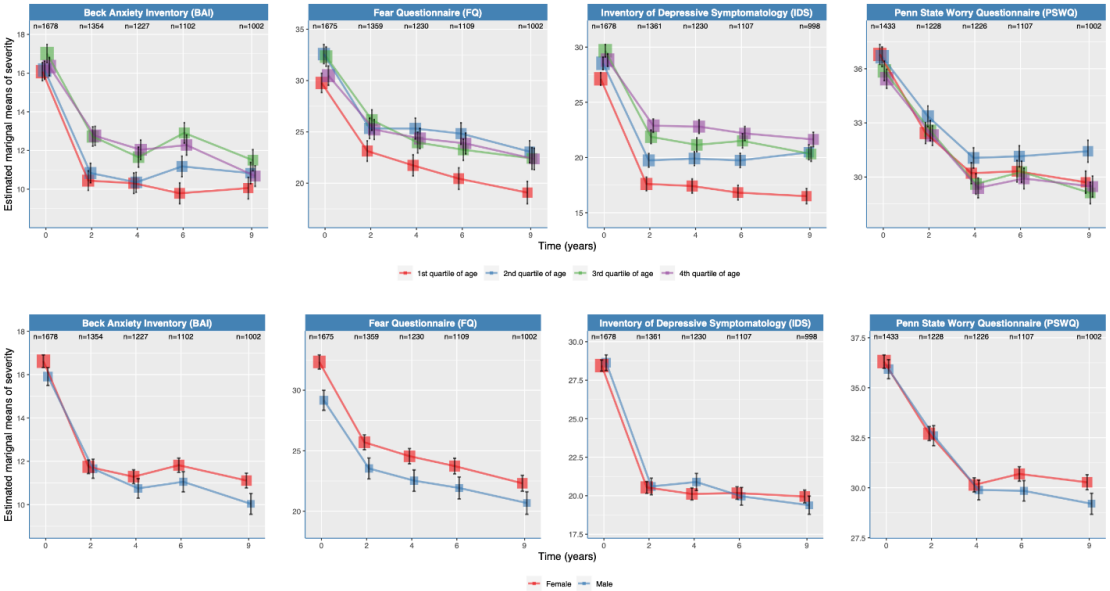


**Figure A.** Categorical trajectories of aggregate diagnoses at two-, four-, six-, and nine-year assessments, based on any CIDI affective disorder and LCI data, following a procedure similar to Verduijn et al (2017). Depression only = depressive disorders such as major depressive disorder or dysthymia. Anxiety only = anxiety disorders such as social phobia and generalized anxiety disorder.

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**Figure B.** Elbow plots of the information criteria values of the Latent Class Growth Analysis comparing one to six classes. AIC = Akaike Information Criterion; BIC = Bayesian Information Criterion; entropy = entropy values; Loglik = log-likelihood; SABIC = sample-size-adjusted BIC.



**Figure C.** Trajectories of depressive, anxiety, fear, and worry symptoms, split into age groups and gender, across the nine-year follow-up period. Error bars represent standard errors, and the box size is proportional to the number of participants.