Neurobiological Divergence of the Positive and Negative Schizophrenia Subtypes Identified on a New Factor Structure of Psychopathology Using Non-negative Factorization

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Neurobiological Divergence of the Positive and Negative Schizophrenia Subtypes Identified on a New Factor Structure of Psychopathology Using Non-negative Factorization: An International Machine Learning Study


ABSTRACT

BACKGROUND: Disentangling psychopathological heterogeneity in schizophrenia is challenging, and previous results remain inconclusive. We employed advanced machine learning to identify a stable and generalizable factorization of the Positive and Negative Syndrome Scale and used it to identify psychopathological subtypes as well as their neurobiological differentiations.

METHODS: Positive and Negative Syndrome Scale data from the Pharmacotherapy Monitoring and Outcome Survey cohort (1545 patients; 586 followed up after 1.35 ± 0.70 years) were used for learning the factor structure by an orthonormal projective non-negative factorization. An international sample, pooled from 9 medical centers across Europe, the United States, and Asia (490 patients), was used for validation. Patients were clustered into psychopathological subtypes based on the identified factor structure, and the neurobiological divergence between the subtypes was assessed by classification analysis on functional magnetic resonance imaging connectivity patterns.

RESULTS: A 4-factor structure representing negative, positive, affective, and cognitive symptoms was identified as the most stable and generalizable representation of psychopathology. It showed higher internal consistency than the original Positive and Negative Syndrome Scale subscales and previously proposed factor models. Based on this representation, the positive–negative dichotomy was confirmed as the (only) robust psychopathological subtypes, and these subtypes were longitudinally stable in about 80% of the repeatedly assessed patients. Finally, the individual subtype could be predicted with good accuracy from functional connectivity profiles of the ventromedial frontal cortex, temporoparietal junction, and precuneus.

CONCLUSIONS: Machine learning applied to multisite data with cross-validation yielded a factorization generalizable across populations and medical systems. Together with subtyping and the demonstrated ability to predict subtype membership from neuroimaging data, this work further disentangles the heterogeneity in schizophrenia.

Keywords: Brain imaging, Machine learning, Multivariate classification, Non-negative factorization, Schizophrenia, Subtyping

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Schizophrenia is a heterogeneous disorder with marked interindividual variability of psychopathology, which is related to treatment response and long-term outcomes (1,2). Earlier clinical subtypes (e.g., hebephrenic, paranoid) were eliminated in recent nosological classifications owing to poor diagnostic stability, validity, and utility (3). Considerable efforts have been devoted to better understand and categorize schizophrenia phenomenology by factorizing symptoms into cardinal dimensions or clustering patients into psychopathological subtypes based on scales such as the Positive and Negative Syndrome Scale (PANSS), a well-established assessment of schizophrenia psychopathology (4).
The 3 PANSS subscales (negative, positive, and general psychopathology) are generally suggested to not optimally and adequately capture the latent organization of schizophrenia symptomatology; items within a subscale show modest internal consistency (5), while those across subscales are strongly correlated (6–8). Previous factorizations of the PANSS have been inconclusive, advocating solutions between 4 and 7 factors (6–11). Although a 5-factor structure was most frequently proposed (9–12), it has continuously failed to be confirmed in independent samples (12–15). Interpretations of previous factor models are, furthermore, complicated by lack of sparsity (all items contribute to any factor) (16) and by coexistence of positive and negative weights (17). Finally, most previous studies investigated rather small and geographically restricted samples, raising doubts over generalization to different populations and medical systems because systematic cross-validation to assess stability and generalizability have rarely been performed. Previous work on psychopathological subtyping is likewise inconclusive (18–20), with added concerns related to longitudinal stability and neurobiological differentiability. These aspects are particularly relevant in the emerging context of precision psychiatry and raise the following questions: Do psychopathological subtypes represent stable patient characteristics, and do they relate to divergent neurobiological substrates that are identifiable from brain imaging data? Functional magnetic resonance imaging (fMRI) parameters may serve as an endophenotype, underpinning the symptomatic heterogeneity (21), which has added ample valuable insights into the neural pathophysiology of schizophrenia and its relation to clinical presentations (22,23). However, whether and to what extent the brain functional connectivity (FC) could discriminate psychopathological subtypes remains unknown. A successful classification using endophenotypical characteristics would support distinctiveness of symptomatically derived schizophrenia subtypes expressed along the cardinal axes of psychopathology.

In the current study, we addressed the aforementioned questions as follows: 1) a robust, cross-validated, and interpretable factor structure of schizophrenia psychopathology was identified based on PANSS scores of more than 2000 patients using an unsupervised machine learning approach (orthonormal projective non-negative matrix factorization [OPNMF]) (24–27); 2) core schizophrenia subtypes were derived by applying soft clustering to the identified factor structure, whose longitudinal stability was evaluated in repeatedly assessed patients; and 3) neurobiological differentiation of those subtypes based on resting-state FC (rsFC) patterns was investigated by cross-validated classification analysis, serving as a biological validation of a clinical (multivariate) construct.

**METHODS AND MATERIALS**

**Sample**

We used 2 large datasets collectively providing individual-item PANSS scores for 2035 patients with schizophrenia: 1) a subset of 1545 patients (586 followed up after 1.35 ± 0.70 years) with complete individual-item PANSS scores and a diagnosis of schizophrenia (DSM-IV criteria) retrieved from the Pharmacotherapy Monitoring and Outcome Survey (PHAMOUS) database (28,29) (this dataset was recruited from 4 institutions located in The Netherlands and assessed with a uniform protocol); 2) a deliberately heterogeneous sample pooled from 9 centers located in Europe, the United States, and Asia (490 patients) (Table 1 and Supplemental Table S1). This international dataset covers a broad range of clinical states, settings, and medical systems, making it ideal to evaluate the generalization of our factor model to new and diverse populations. Diagnoses in the international sample were established based on the DSM-IV, DSM-IV-TR, or DSM-5 criteria (Supplement). At all sites, data were acquired in accordance with the Declaration of Helsinki and after obtaining informed consent from the patients. Approval for the pooled reanalysis was obtained from the ethics committee of the Heinrich Heine University Düsseldorf.

**Factorization of PANSS Using OPNMF**

OPNMF (25,27) decomposes given data (PANSS) into 2 non-negative matrices: 1) a basis matrix (dictionary) with factors as columns that can be readily generalized to new data owing to the projective constraint and 2) a factor-loading matrix representing symptomatology of individual patients along these factors. The orthonormality constraint promotes a sparse, and hence interpretable, representation. For choosing the number of factors, a set of sophisticated evaluation strategies was implemented (see Supplement and Supplemental Figures S1 and S2).

We first applied OPNMF to the PANSS scores from the 1545 PHAMOUS patients with the number of factors ranging from 2 to 11. The optimal number of factors was identified by using cross-validation in 10,000 split-half analyses. The PHAMOUS sample was split into two halves, and on each split sample OPNMF was performed to derive the dictionary. The congruency between item-to-factor assignments, based on its largest coefficient, was assessed using the adjusted Rand index (30) and variation of information (31) along with the concordance index (32) between the dictionaries. We also quantified out-of-sample reconstruction error by projecting the data of one split sample onto the dictionary from the other split sample. A lower increase in out-of-sample error compared with within-sample reconstruction error indicates better generalizability. This split-sample analysis was repeated on the international dataset. Additional bootstrap and 10-fold cross-validation analyses were conducted on each of the 2 samples independently.

Most critically, we assessed stability and generalizability between the factorizations of the PHAMOUS sample (good for learning a structure owing to size) and the international sample (good for validation owing to heterogeneity). We performed OPNMF independently on the bootstrapped samples from each dataset. The resulting factorizations were then compared using the approaches described above. That is, for each number of factors, we assessed stability by comparing the dictionaries obtained from factorization of bootstrapped samples (PHAMOUS vs. international). Most important, we also evaluated generalization to new data by measuring the increase in reconstruction error for the international data following projection onto the PHAMOUS dictionary. This cross-sample evaluation was repeated after accounting for between-dataset differences in sample size, age, and illness duration.
Internal Consistency and Relationship Among Variables

Internal consistency of the optimal OPNMF model, as well as the PANSS subscales (as reference), was assessed using Cronbach’s alpha, where higher values indicate more closely related items within a set. Relationships between the OPNMF factor loadings were assessed using linear and partial correlations (controlling for symptom severity, i.e., total PANSS score), including bootstrap stability analyses (Supplement). The OPNMF factor loadings were correlated with the 3 PANSS subscales. Correlations between individual items were also computed. For comparison, we performed an exploratory factor analysis (EFA) on the PHAMOUS sample and a confirmatory factor analysis on the international sample as well as a principal component analysis on both samples (Supplement). Effects of gender, age, illness duration, and symptom severity on the OPNMF factor loadings were analyzed in the international sample (N = 393 with complete information). Following a multivariate analysis of variance to assess effects on all the loadings, individual 4-way analyses of variance (ANOVA) were performed on each loading to identify its association with the demographic and clinical features (corroborated by bootstrap and leave-one-site-out analyses) (Supplement). Current drug dosages of antipsychotic medication, available for 149 patients, were olanzapine-equivalent transformed (33) and included in the 4-way ANOVA models for a supplementary analysis.

Table 1. Demographic and Clinical Characteristics of the Patients With Schizophrenia

<table>
<thead>
<tr>
<th>Characteristic</th>
<th>PHAMOUS Sample (N = 1545)</th>
<th>International Dataset From Nine Centers (N = 490)</th>
<th>International Dataset With Imaging (N = 147)</th>
<th>Statistic</th>
<th>p Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Demographic</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Age, years</td>
<td>44.15 (11.42)</td>
<td>33.82 (10.28)</td>
<td>34.89 (11.67)</td>
<td>183.51</td>
<td>&lt;.001</td>
</tr>
<tr>
<td>Gender, male/female, n</td>
<td>1108/437</td>
<td>333/157</td>
<td>102/45</td>
<td>2.45</td>
<td>.292</td>
</tr>
<tr>
<td>Illness duration, years</td>
<td>18.22 (10.54)</td>
<td>9.13 (8.98)</td>
<td>11.37 (10.36)</td>
<td>134.71</td>
<td>&lt;.001</td>
</tr>
<tr>
<td>PANSS</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Positive</td>
<td>12.48 (4.91)</td>
<td>14.24 (5.76)</td>
<td>15.36 (5.50)</td>
<td>37</td>
<td>&lt;.001</td>
</tr>
<tr>
<td>Negative</td>
<td>14.60 (6.20)</td>
<td>14.67 (7.21)</td>
<td>15.07 (6.06)</td>
<td>0.375</td>
<td>.687</td>
</tr>
<tr>
<td>General</td>
<td>26.70 (8.16)</td>
<td>29.10 (11.34)</td>
<td>30.93 (10.97)</td>
<td>23.67</td>
<td>&lt;.001</td>
</tr>
<tr>
<td>Symptom severity, total PANSS score</td>
<td>53.78 (16.35)</td>
<td>58.01 (21.87)</td>
<td>61.36 (19.57)</td>
<td>19.48</td>
<td>&lt;.001</td>
</tr>
<tr>
<td>P3 item score</td>
<td>2.30 (1.47)</td>
<td>2.66 (1.83)</td>
<td>3.22 (1.91)</td>
<td>28.18</td>
<td>&lt;.001</td>
</tr>
<tr>
<td>Medication</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Atypical antipsychotics</td>
<td>NA</td>
<td>167 (34.1%)</td>
<td>110 (74.8%)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Typical antipsychotics</td>
<td>NA</td>
<td>26 (5.3%)</td>
<td>8 (5.4%)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Both atypical and typical antipsychotics</td>
<td>NA</td>
<td>16 (3.3%)</td>
<td>9 (6.1%)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>None or unknown</td>
<td>NA</td>
<td>281 (57.3%)</td>
<td>20 (25.9%)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Current antipsychotic medication</td>
<td>NA</td>
<td>19.64 (14.15)</td>
<td>19.30 (12.57)</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Data are mean (SD) or n (%). p Values of <.001 indicate a significance of p < .05. Except for gender, which was based on χ² test, all other statistics were based on 1-way analyses of variance. Of note, because the detailed medication information was missing for several patients in different proportions for those with or without imaging data in the international dataset, statistical comparisons were not conducted.

Post hoc analysis after 1-way analyses of variance showing significant pairwise differences among the 3 datasets are indicated in the footnotes. NA, not available; PANSS, Positive and Negative Syndrome Scale; PHAMOUS, Pharmacotherapy Monitoring and Outcome Survey; P3 item measures hallucinatory behavior.

(Supplement). Leave-one-site-out validation was performed on both the PHAMOUS and international samples to check for site bias. We also repeated all analyses after removing outliers or including the repeated PANSS measurements (Supplement). Factorizations of the pooled (PHAMOUS + international) sample, as well as the stability and accuracy of PHAMOUS-generated dictionaries in estimating out-of-sample loadings or item scores, were also assessed (Supplement).

After identifying the optimal OPNMF factorization, the international sample was projected onto this PHAMOUS-derived dictionary to obtain factor loadings for subsequent analyses (except for longitudinal analysis because reassessments were available only in the PHAMOUS sample) to avoid double dipping or leakage that would occur if scores were analyzed in the same dataset used to derive the dictionary.

Machine Learning Models of Schizophrenia Psychopathology
Psychopathological Subtypes
After adjusting for age, gender, illness duration, and symptom severity, factor loadings were used as features for clustering patients into psychopathological subtypes. After confirming the data clusterability (34), we applied fuzzy c-means clustering (35), which provided cluster membership likelihoods for all patients. The optimal cluster number was determined based on the fuzzy silhouette index (36), the Xie and Beni index (37), and partition entropy (35). Stability was tested by leave-one-site-out replication, subsampling, and bootstrap resampling (Supplement). Given the heterogeneous nature of schizophrenia and the observation of multiple patients with ambiguous memberships, a cutoff over the membership likelihoods was adopted to remove cluster-ambiguous patients. For this, additional evidence from Gaussian mixture modeling (GMM) was considered. Specifically, patients were clustered again using GMM, and the optimal cluster number was determined by Bayesian information criterion (38). After assigning patients to the clusters, we took the intersection of the c-means and GMM results. A cutoff was chosen, based on the c-means membership likelihoods that well discriminated the patients inside the intersection from those outside, while also retaining a decent sample size for classification. This filtering step is critical because ambiguous patients might obscure otherwise classifiable rsFC patterns for the identified subtypes. A forward-backward scheme between subtypes regarding factor loadings and demographic and clinical features were ascertained by permutation tests (39). To assess longitudinal stability, the same c-means clustering was applied to the repeated assessments of the PHAMOUS sample. The optimal dictionary, identified on the 1545 PHAMOUS patients without repeatedly assessed PANSS scores, was used for projection to yield the factor loadings. After excluding ambiguous cases, patients assigned to the same clusters in both initial and follow-up stages were regarded as longitudinally stable (Supplement). For comparison, the same clustering was also performed on the factor loadings without any covariates or symptom severity adjustment as well as on the PANSS subscales or items both with and without covariates adjustment (Supplement).

Classifying Psychopathological Subtypes From rsFC
Multivariate classification analysis was conducted on patients from the international sample for whom imaging data were available after excluding those with ambiguous subtype assignment, low image quality, or excessive head motion (n = 84) (Supplemental Figure S24). After standard preprocessing (Supplement), regional time series were extracted based on a parcellation scheme with 600 cortical parcels (40) and 36 subcortical parcels (41), adjusted for confounders (42), and were used to compute the functional connectome. We tested each parcel for whether its pattern of rsFC to all other parcels allowed classifying subtype membership in novel subjects. Resulting parcelwise accuracies yielded a whole-brain map indicating the classifiable power of each parcel's connectivity profile. The radial basis function kernel support vector machine classifier, which can deal with the potentially nonlinear relationship between the psychopathological and neural spaces, was employed. A stratified 10-fold cross-validation was implemented to assess the out-of-sample classification performance (Supplemental Figure S25). Effects of age, gender, site, illness duration, symptom severity, and head-motion parameters were adjusted using a linear regression model fitted only in the training sample (43). Significance of the parcelwise accuracy was estimated by permutation tests, followed by false discovery rate correction for multiple comparisons (Supplement). Parcels surviving false discovery rate were functionally characterized (http://brainmap.org/) (44) (Supplement).

RESULTS
Dimensions of Psychopathology
The most robust and generalizable model consisted of 3 factors for the PHAMOUS sample (Figure 1A), which effectively combined the positive and affective symptoms compared with the optimal 4-factor model for the international dataset (Figure 1B). The additional factor in the international dataset may relate to the higher prevalence of psychotic symptoms, particularly auditory hallucinations, compared with the PHAMOUS sample, which contains more long-term patients (Table 1). Consequently, a 4-factor model was identified as the most stable and, importantly, generalizable model of psychopathology in the cross-sample evaluation (Figure 1C). The first factor mainly represents negative symptoms such as blunted affect and affective (Figure 1D). The second factor represents positive symptoms such as delusions and hallucinations. The third factor comprises symptoms such as depression, anxiety, and tension, reflecting an affective dimension. The fourth factor represents cognitive impairments. Notably, only a few items contributed to multiple dimensions (e.g., active social avoidance contributed to both negative and affective factors).

All findings were fully confirmed by 1) bootstrap and 10-fold cross-validation, 2) removing outliers (18 patients), 3) adding PANSS data from follow-up examination in the PHAMOUS sample, 4) leave-one-site-out validation, 5) accounting for between-dataset differences in sample size, age, and illness duration, 6) pooling the 2 datasets with cross-validation and out-of-sample generalization assessments, and 7) loading or item score predictions across factor solutions, bootstrapped samples, and sites (Supplemental Figures S3–S11).

Internal Consistency and Relationship Among Variables
Items within a factor showed higher and more homogeneous positive correlations (and fewer anticorrelations) for the OPNMF factors than for the PANSS subscales (Figure 2A, B and Supplemental Figure S12). Internal consistency of our OPNMF 4-factor structure (positive: Cronbach’s alpha = .75; negative: .92; affective: .85; cognitive: .83) was on average higher than that of the PANSS subscales (positive: .72; negative: .87; general psychopathology: .87), previously reported factor models (ranging from .60 [excited] to .90 [negative]) (7–9,45), and the EFA models derived from the current sample (.49–.91) (Supplemental Table S2). All PHAMOUS-derived 4- to 7-factor EFA models could not be confirmed in the international sample owing to inadequate fit (Supplemental Table S2). Compared with principal component analysis, OPNMF showed
better generalizability (Supplemental Figure S13). The positive and negative factors were highly correlated with the positive and negative PANSS subscales, respectively, both before (r = .92 and r = .97) and after (r = .85 and r = .89) controlling for symptom severity (Supplemental Figure S14). Interestingly, after adjusting for symptom severity, the cognitive factor did not correlate with general psychopathology (r = .02).

Over individual patients, the loadings on our 4 factors were significantly intercorrelated, with negative and positive factors showing the lowest correlation (r = .32 averaged over 10,000 bootstraps) and negative and affective factors showing the highest correlation (r = .70) (Figure 2C). After controlling for symptom severity, positive and negative factors became anticorrelated (r = −.59) (Supplemental Figure S15).

Multivariate analysis of variance revealed a significant influence of symptom severity on the joint factor loadings (p < .001). Follow-up 4-way ANOVAs showed that symptom severity had a significant effect on each factor (all ps < .001). A significant negative relationship with age (β = −.15) and a significant negative relationship with age (p = .033, β = −.015) (Figure 2D), although both covariates were collinear (r = .65). In contrast, loadings on the negative factor were higher for older individuals (p = .11, β = .016) and lower for those with longer illness duration (p = .18, β = −.015).

Gender differences were not observed in any factor. Bootstrap (Figure 2E) and leave-one-site-out analyses corroborated the aforementioned ANOVA findings (Supplemental). Adding

**Figure 1.** Split-half cross-validation (10,000 repetitions) of stability and generalizability of the factor solutions derived by orthonormal projective non-negative matrix factorization. The 3 indices—adjusted Rand index (aRI), variation of information (VI), and concordance index (CI)—demonstrate the factor stability, while out-of-sample increased reconstruction error (RE) reflects the performance of generalizability. Box plots show stability and generalizability results of the factor solutions. Higher values for aRI and CI (upper row) indicate higher stability. Lower values for VI and out-of-sample increase in RE (bottom row) indicate better stability and generalizability, respectively. For the box plots, the red line depicts the median, the green diamond depicts the mean, and the whiskers represent the 5th and 95th percentiles. For the factor models, the weight of an item in assigning to a specific psychopathological factor (columns of the matrix) is color-coded according to the coefficients by a heat color map from gray (minimum) to dark red (maximum). (A) The best factor solution derived from the Pharmacotherapy Monitoring and Outcome Survey (PHAMOUS) data (1545 patients) is shown. According to the 4 aforementioned evaluation indices, a 3-factor model was indicated as the best because both the mean and median values for VI and out-of-sample increase in RE achieve the lowest, while the aRI and CI reach the highest, at that point. (B) The best factor solution derived from the international sample (490 patients) is shown. Four factors is the optimal solution because the mean and median values of VI and out-of-sample increase in RE achieve the local minimum, while the aRI reaches maximum and the CI reaches a local maximum. (C) The best factor solution identified by the bootstrap comparison of the two datasets (PHAMOUS vs. international) is shown. A 4-factor solution is optimal because the mean and median values of the aRI and CI reach the maximum, while the mean and median values of VI and the median value of out-of-sample increase in RE achieve the minimum. (D) The most stable and generalizable 4-factor structure derived from the PHAMOUS sample, serving as the best basis for future studies, is shown. This 4-factor model consists of a negative (factor 1), a positive (factor 2), an affective (factor 3), and a cognitive (factor 4) factor that were named based on the items they contained. PANSS, Positive and Negative Syndrome Scale.
olanzapine-equivalent dosage to the 4-way ANOVA did not reveal any significant association with medication.

**Psychopathological Subtypes**

Fuzzy c-means clustering on the adjusted loadings revealed an optimal 2-cluster solution (Figure 3A, B and Supplemental Figures S16 and S17). Although GMM demonstrated an optimal 3-cluster solution, one of the clusters was diffusely distributed in space containing patients from both c-means clusters. This GMM cluster was excluded because it would not represent any specific subtype (Figure 3C). Patients inside the c-means GMM intersection had higher c-means membership likelihoods (roughly .70) to belong to their own cluster than those outside the intersection (p < .01, Wilcoxon rank-sum test) (Figure 3D). We chose the cluster cores using the likelihoods of c-means with a cutoff of .70. As a result, 2 core subtypes were defined after filtering out 50 ambiguous patients.
from each of the 2 c-means clusters (Figure 3E). The first subtype showed a psychopathological profile dominated by negative and affective symptomatology (subtype A). The other subtype featured prominent positive symptoms (subtype B; all ps < .001 in permutation tests) (Figure 3F and Supplemental Figure S18). Importantly, subtypes did not differ in gender distribution, age, or illness duration (all ps > .05), but subtype B showed higher symptom severity (p = .008). The same 2-cluster solution was replicated on the PHAMOUS patients with complete demographic and clinical information (N = 1326; 56% of the 603 ambiguous patients in subtype B when hard-clustered) and on those with repeatedly assessed PANSS scores (n = 527; 45% ambiguous). Nearly 80% of the reassessed patients retained their subtype, with subtype A being more stable (85%) (Supplement). The additional clustering analyses supported our 4-factor model with covariates...
adjustment for a clinically meaningful subtyping (Supplemental Figures S19–S23).

Classifying Psychopathological Subtypes From rsFC

The rsFC profile of the parcel located in the right ventromedial prefrontal cortex yielded the highest out-of-sample classification accuracy (70% of patients not used for training were assigned to the correct psychopathological subtypes), followed by parcels in the right temporo-parietal junction, bilateral precuneus, and posterior cingulate cortex (Figure 4C). Permutation tests showed that the top 104 classifiable parcels were significant (p < .05) against chance (i.e., randomized subtype labels), and 53 parcels survived false discovery rate correction (q < .05) (Figure 4B). Of note, parcels are labeled by their microanatomical location with their functional implications (Supplemental Table S4). Classification with additional global mean signal removal or with an rsFC-based subcortical parcellation replacement (7 parcels [46] as a control analysis instead of the finer Brainnetome subcortical parcellation) replicated these results (see Supplement and Supplemental Figure S26).

DISCUSSION

By factoring the PANSS scores from a large sample using OPNMF and cross-validating the results in a heterogeneous multisite dataset, we revealed a robust, replicable, and generalizable 4-factor structure comprising negative, positive, affective, and cognitive dimensions across populations, settings, and medical systems. Based on this 4-factor structure, 2 core psychopathological subtypes were obtained that showed good longitudinal stability and could be discriminated by regional rsFC patterns, with the right ventromedial prefrontal cortex showing the highest (70%) classification accuracy.

Relationship to Previous Factor Models of PANSS

The 3 PANSS subscales do not reflect the latent structure of this inventory well (5–7). In turn, our model represents a stable, generalizable, and well-interpretable description of schizophrenia psychopathology suited for representing the full range of acute and chronic symptoms. Resonating with this view, a pyramidal model proposed by the PANSS developers comprised 4 components (6). Three of these (negative, positive, and affective dimensions) showed good agreement with our model. The fourth component, however, isolated only excitement, while cognitive disturbances were distributed across all dimensions or discarded. Such a representation is obviously at odds with the importance of cognitive dysfunction and has prompted the proposal of more complex models (7–11), for example, a recent 5-factor model reflecting negative, positive, depressed, excited, and cognitive dimensions (45). However, replicability, external validity, and generalization remain a concern for these models (47). As a striking example, White et al. (13) found that none of 20 tested models fit their data adequately, and they put forward a new pentagonal model that later (together with 24 other models) also could not be confirmed (14). In the same study (12), the authors developed an improved 5-factor model using 10-fold cross-validation. However, it still failed to be confirmed, along with 31 other 5-factor models, in a later study involving 2 large Chinese samples (15). In our sample, inadequate fit for EFA models with 4 to 7 factors was also manifested, and the fifth OPNMF factor, compared with the 4 factors, showed the poorest out-of-sample loading predictions (Supplemental Figure S10B). These facts, as a whole, point to a fundamental instability of 5-factor models (9–15,44). Addressing these concerns, the current work not only was based on a large sample for model identification but also, importantly, focused strongly on cross-validated stability and out-of-sample generalization. Critically, the external validation was based on a heterogeneous international sample, and the optimal model suggested a single factor to combine both the cognitive and excited symptoms. This view is corroborated by observations that cognitive and excited symptomatologies are highly correlated (45) and share similar neurobiological substrates (48,49).

Internal Consistency and Relationship Among Variables

Although we identified the optimal representation by its robustness and ability to generalize to new populations, the positive and negative dimensions of our model also showed better internal consistency than the PANSS subscales while differentiating the broad general psychopathology. Moreover, our affective and cognitive factors showed higher internal consistency than those reported in previous factor models (7,8,10,45). Finally, correlations between individual items within OPNMF factors were higher and more homogeneous compared with the PANSS subscales.

Matching previous reports (4,8,45), negative and positive factors from our model were least related before, and showed a strong anticorrelation after, controlling for symptom severity. The inverse age versus illness duration effect on negative symptoms implies that this effect may be more related to age than to illness duration. This is intriguing from the perspective of early aging or degeneration, but it needs to be viewed with caution because age and illness duration are highly correlated. Collinear variables in a single linear regression model make it difficult to disentangle their respective effects on the negative factor as well as on the cognitive factor (50).

Psychopathological Subtypes, Longitudinal Stability, and Neurobiological Differentiability

Our results revealed 2 distinct schizophrenia subtypes featuring predominantly positive and negative symptoms, respectively. The subtypes were longitudinally stable and could be classified from neuroimaging data. Such a positive-negative dichotomy has been widely supported (51,52). Finer distinctions have been proposed but show poor replicability (18–20). The inconsistency of finer subtyping may relate to idiosyncrasies in small samples from a single geographical region and to the lack of explicit analyses of stability and replicability. Moreover, longitudinal stability of our new subtypes was higher than that reported for traditional clinical subtypes or for a positive/negative/mixed topology (53–55). Interestingly, we found subtype A to be particularly stable. Previous studies indicated that both mixed and negative symptom states increase over time, whereas psychotic
expressions usually diminish outside acute episodes and over time (54,55). Future studies with a longer follow-up duration are desired; the mean of 1.35 years’ follow-up assessed in the current study is not a long period in schizophrenia. In addition, the employed soft-clustering method better accommodates ambiguous patients compared with previous hard-clustering methods, and furthermore, patients with ambiguous memberships can be filtered out with appropriate cutoffs to improve the ability of detecting neurobiological distinctions between subtypes. Nonetheless, the cutoff value chosen in the current

Figure 4. Classifying psychopathological subtypes from resting-state functional connectivity. (A) The brain parcellation scheme (600 cortical parcels plus 36 subcortical parcels) and the resting-state functional connectivity matrix that was constructed based on this parcellation system are illustrated. In parcelwise classification analysis, one column of the connectivity matrix was taken to represent the functional connectivity pattern for a single parcel. (B) Cortical surface rendering and subcortical axial slices show parcelwise classification results for those parcels that survived false discovery rate (FDR) correction ($q < .05$), demonstrating a neurobiological divergence between the two identified psychopathological subtypes of schizophrenia. (C) Cortical surface rendering and subcortical axial slices show parcelwise classification results for the whole brain. The balanced classification accuracy is color-coded from light gray to dark red.
study should be noted as heuristic. The cluster-ambiguous patients might represent a transient group lying between the two more differentiated subtypes.

The current top classifiable brain regions all are implicated in schizophrenia pathophysiology and processes relevant to the psychopathological distinction (22,56–60). Previous findings in the literature relating fMRI parameters to differential symptoms exclusively relied on group-level analyses, while our approach bridged an important gap between neurobiological divergence and distinct symptomatic patterns at the individual level. To our knowledge, it is the first study to successfully classify psychopathological subtypes in schizophrenia. Of note, the current classification accuracy was similar to accuracy levels previously reported for classifications of patients with schizophrenia versus healthy participants (61–63). The demonstrated neurobiological differentiability corroborates the currently identified schizophrenia subtypes expressed along the 4 OPNMF dimensions.

**Limitations and Considerations**

We assessed factor structure, subtypes, and their neurobiological differentiations with a particular emphasis on robustness and generalization. This conservative approach seems necessary given current concerns of nonreplicability in biomedical research, but it might have contributed to the fact that we corroborated the clinically well-established positive–negative distinction rather than identifying more differentiated subtypes. We note, however, that a recent imaging-based clustering also provided evidence for 2 subtypes (64), and we stress that the current analysis of a large heterogeneous sample did not reveal any evidence for a more fine-grained differentiation among the patients. Thus, it remains to be seen whether an additional differentiation between these 2 core subtypes may be robustly revealed by analyzing substantially larger samples or whether previously proposed additional subtypes represent distinctions that could be found in a particular dataset but are not universally present. We also acknowledge that patients were on their regular medication as prescribed by the attending psychiatrists, and thus the current results might be confounded by direct and indirect medication effects thereof. However, it stands to reason that a multisite study, pooling patients from different psychiatrists with differential medication strategies, will render medication largely as a source of random variation in our data. Such noise would effectively make it harder to identify generalizable psycho-pathological factors and robust subtypes and, in particular, to train models that work well for out-of-sample classification of subtype membership. Thus, we would argue that the current results should not be driven by medication effects but rather represent general structures of psychopathology and schizophrenia subtypes. In addition, rsfMRI has its own limitations such as variability across scanning sessions and the issue of confounding factors (42,65,66). We focused on rsfMRI because it could temporally better map the likewise state-dependent psychopathology compared with structural MRI. Using advanced machine learning with cross-sample validation, the current study suggested a stable and generalizable 4-factor model of PANSS. This representation allowed for the definition of a reliable positive–negative subtype differentiation that showed good longitudinal stability and a neurobiological divergence in rsFC. Overall, the current work further disentangled the heterogeneity of schizophrenia, possibly allowing for the design of more specifically targeted treatments.

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A Dimensions and Clustering Tool for assessing schizophrenia Symptomatology (DCTS) is available at http://webtools.inm7.de/sczDCTS/.

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