Difficulties in advancing effective patient-specific therapies for psychiatric disorders highlights a need to develop a neurobiologically-grounded, quantitatively stable mapping between neural and symptom variation. This gap is particularly acute for psychosis-spectrum disorders (PSD). Here, in a sample of 436 cross-diagnostic PSD patients, we derived and replicated a data-driven dimensionality-reduced symptom space across hallmark psychopathology symptoms and cognitive deficits, which was predictive at the single patient level. In turn, these data-reduced symptom axes mapped onto distinct and replicable univariate brain maps. Critically, we found that multivariate brain-behavior mapping techniques (e.g. canonical correlation analysis) did not show stable results. Instead, we show that a univariate brain-behavioral space (BBS) mapping can resolve stable individualized prediction. Finally, we show a proof-of-principle framework for relating personalized BBS metrics with molecular targets via serotonin and glutamate receptor manipulations and gene expression maps. Collectively, these results highlight a stable and data-driven BBS mapping across PSD, which offers an actionable quantitative path that can be iteratively optimized for personalized clinical biomarker endpoints.

Mental health conditions cause profound disability with most treatments yielding limited efficacy across psychiatric symptoms (1–4). A key step towards developing more effective therapies for specific psychiatric symptoms is reliably mapping them onto underlying neural systems. This goal is challenging because neuropsychiatric diagnostics still operate under “legacy” categorical constraints, which were not informed by quantitative neural or symptom data. Critically, diagnostic systems in psychiatry, such as the Diagnostic and Statistical Manual of Mental Disorders (DSM) (5), were built to aid clinical consensus. However, they were not designed to guide quantitative mapping of symptoms onto neural alterations (6, 7). Consequently, the current diagnostic framework cannot, by definition, optimally map onto patient-specific brain-behavioral alterations. This challenge is particularly evident along the psychosis spectrum disorders (PSD) where there is notable symptom variation across what DSM considers distinct diagnostic categories (such as schizophrenia [SZP], schizo-affective [SADP], bipolar disorder with psychosis [BPP]). For instance, despite BPP being a distinct DSM diagnosis, BPP patients exhibit similar, but attenuated psychosis symptoms and neural alterations similar to SZP (e.g. thalamic functional connectivity (FC) (8)). It is essential to quantitatively map such shared clinical variation onto common neural alterations to circumvent constraints for biomarker development (6, 7, 9) – a key goal for development of neurobiologically-informed personalized therapies (8, 9).

Recognizing the limits of categorical frameworks, the NIMH’s Research Domain Criteria (RDoC) initiative introduced dimensional mapping of “functional domains” on to neural circuits (10). This motivated cross-diagnostic multisite studies to map PSD symptom and neural variation (11–14). In turn, multivariate neuro-behavioral analysis across PSD and mood spectra observed brain-behavioral relationships across diagnoses, with the goal of informing individualized treatment (15). A key challenge that these studies attempted to address is to move beyond a priori clinical scales, which provide composite scores (16) that may not optimally capture neural variation (7, 17). For instance, despite many data-driven dimensionality reduction symptom studies (18–24), a common approach in PSD neural research is still to sum “positive” or “negative” psychosis symptoms into a single score for mapping onto neural features. Importantly, altered symptom-to-neural variation may reflect a more complex weighted symptom combination beyond such composite scores.

As noted, multivariate neuro-behavioral studies have attempted to address this; however, such studies have failed to replicate due to overfitting (25), arising from high dimensionality of behavioral and neural features and a comparatively limited number of subjects (26). Current state-of-the-art large-scale clinical neuroimaging studies have target enrollment totals of ~200-600 subjects (https://www.humanconnectome.org/disease-studies). We
therefore wanted to test whether a stable, replicable neuro-behavioral geometry can be derived with a sample size that is on par with current large-scale studies.

It is possible that a linearly-weighted lower-dimensional symptom solution (capturing key disease-relevant information) produces a replicable and robust univariate brain-behavioral mapping. Indeed, recent work used dimensionality reduction methods successfully to compute a neural mapping across canonical SZP symptoms (23). However, it remains unknown if this approach generalizes across PSD. Moreover, it is unknown if incorporating cognitive assessment, a hallmark and untreated PSD deficit (17), explains neural feature variation that is distinct from canonical PSD symptoms. Finally, prior work has not tested if a low-dimensional symptom-to-neural mapping can be predictive at the single patient level – a prerequisite for individualized clinical endpoints.

To inform these gaps, we first tested two key questions: i) Can data-reduction methods reliably map symptom axes across PSD that include both canonical symptoms and cognitive deficits? ii) Do these lower-dimensional symptom axes map onto a replicable brain-behavioral solution across PSD? Specifically, we combined fMRI-derived resting-state measures with psychosis and cognitive symptoms (27, 28) obtained from a public multi-site cohort of 436 PSD patients and 202 healthy individuals collected by the Bipolar-Schizophrenia Network for Intermediate Phenotypes (BSNIP-1) consortium across 6 sites in North America (11). The dataset included included 150 patients formally diagnosed with BPP, 119 patients diagnosed SADP, and 167 patients diagnosed with SZP (Table S1). This cohort enabled symptom-to-neural cross-site comparisons across multiple psychiatric diagnostic categories, which we then mapped onto specific neural circuits. We tested if PSD symptoms map onto a low-dimensional solution that is stable for individual patient prediction. Next, we tested if this low-dimensional symptom solution yields novel and replicable neural mapping compared to canonical psychosis composite scores or DSM diagnoses. In turn, we tested if the computed symptom-to-neural mapping is replicable across symptom axes and actionable for individual patient prediction. Finally, we ‘benchmark’ the derived symptom-relevant neural maps by computing their similarity against independently collected pharmacological fMRI maps from healthy adults in response to putative PSD receptor treatment targets (glutamate via ketamine and serotonin via LSD) (29, 30). In turn, we used the Allen Human Brain Atlas (AHBA) to compute gene expression maps (31, 32) for targets implicated in PSD (i.e. interneurons, serotonin and GABA receptor genes). In turn, we tested if the gene targets map onto the derived symptom-relevant neural targets. Collectively, this study used data-driven dimensionality reduction methods to map orthogonal and stable symptom dimensions across 436 PSD patients. In turn, the goal was to map novel and replicable symptom-to-neural relationships across the PSD derived from a low-dimensional symptom solution. Finally, these effects were benchmarked against molecular imaging targets that may be actionable for individualized clinical endpoints. The key questions and results of this study are summarized in Fig. S1 and an overview of the workflow is presented in Fig. S2.

Results

Dimensionality-Reduced PSD Symptom Variation is Stable and Replicable. First, to evaluate PSD symptom variation we examined two instruments: the Brief Assessment of Cognition in Schizophrenia (BACS) and Positive and Negative Syndrome Scale (PANSS) instruments to capture “core” PSD psychopathology dimensions (Fig. 1A-B). We observed differences across DSM diagnoses (Fig. 1A, p<0.05, Bonferroni corrected), however, symptom distributions revealed notable overlap across the PSD sample that crossed diagnostic boundaries (11, 33). Furthermore, we observed marked collinearity between symptom items across the PSD sample (Fig. 1B), indicating that a lower-dimensional solution may better capture this symptom space. Specifically, we hypothesized that such a lower-dimensional symptom solution may improve PSD brain-behavior mapping as compared to a high-dimensional solution or pre-existing symptom scales. Here we report results from a principal component analysis (PCA) as it produces a deterministic solution with orthogonal axes (i.e. no a priori number of factors needs to be specified) and captures all symptom variance.

Results were highly consistent with prior symptom-reduction studies in PSD: we identified 5 PCs (Fig. 1C), which captured 50.93% of all variance (see Methods & Fig. S3) (23). Notably, complementary data-reduction procedures produced similar effects for this PSD sample (e.g. independent component analysis (ICA), see Supplementary Note 1 & Fig. S4). The key innovation here is the combined analysis of canonical psychosis symptoms (i.e. positive and negative) and cognitive deficits, which are a fundamental PSD feature (17). The 5 PCs revealed few distinct boundaries between DSM categories (Fig. 1D). Fig. 1E highlights symptom configurations forming each PC, which drove their naming via the pattern of the most strongly weighted individual symptoms (e.g. PC3 – “Psychosis Configuration”; see Supplementary Note 2). Critically, PC axes were not parallel with traditional aggregate symptom scales. For instance, PC3 is angled at ~45° to the dominant direction of PANSS Positive and Negative symptom variation (purple and blue arrows respectively in Fig. 1F).

Next, we show that the PCA solution was highly stable when tested across sites, k-fold cross-validations, and split-half replications. We show that the symptom-derived PCA solution remained highly robust across all sites (Fig. 2A) and 5-fold cross-validation iterations (see Methods). Notably, the total proportion of variance explained as well as the total number of significant PCs remained stable (Fig. 2A). Second, PCA loadings accurately and reliably predicted the scores of subjects in a hold-out sample (Fig. 2B). Specifically, for each left-out site, we tested if single-subject predicted scores were similar to the originally observed scores of these same subjects in the full PCA (i.e. performed with
Fig. 1. Quantifying data-driven low-dimensional variation of cross-diagnostic psychosis spectrum disorder symptoms and cognitive deficits. (A) Distributions of symptom severity for each of the DSM diagnostic groups across core psychosis symptoms (PANSS positive, negative, and general symptoms tracking overall illness) and cognitive deficits (BACS composite cognitive performance). BPP: bipolar disorder with psychosis (yellow, N=150); SADP: schizo-affective disorder (orange, N=119); SZP - schizophrenia (red, N=167); All PSD patients (black, N=436); Controls (white, N=202). Bar plots show group means; error bars show standard deviations. (B) Correlations between 36 psychosis and cognitive deficit symptoms for all PSD patients (N=436). (C) Screeplot showing the % variance explained by each of the principal components (PCs) from a principal component analysis performed using all 36 symptom measures across all 436 PSD patients. The size of each point is proportional to the variance explained. The first five PCs (green) survived permutation testing (p<0.05, 5,000 permutations). Together they capture 50.93% of all symptom variance (inset). (D) Distribution plots showing subject scores for the 5 significant PCs for each of the clinical groups, normalized relative to the control group. Note that control subjects (CON) were not used to derive the PCA solution. However, all subjects, including controls, can in principle be projected into the data-reduced symptom geometry. (E) Loading profiles shown in dark grey for the 36 PANSS/BACS symptom measures on the 5 significant PCs. Each PC (“Global Functioning”, “Cognitive Functioning”, “Psychosis Configuration”, “Affective Valence”, “Agitation/Excitement”) was labeled according to the pattern of symptom loadings (see Supplementary Note 2). See Fig. S3G for numerical values of the PC loadings. The PSD group mean score on each symptom measure is also shown, in light grey (scaled to fit on the same radarplots). Note that the group mean pattern resembles the PC1 loading profile most closely (as PC1 explains the most variance in the symptom data). (F) PCA solution projected into a 3D coordinate space defined by the first three significant PC axes. Colored arrows show a priori aggregate PANSS/BACS vectors projected into the data-reduced PC1-3 coordinate space. The a priori aggregate symptom vectors do not directly align with data-driven PC axes, highlighting that PSD symptom variation is not captured fully by any one aggregate a priori symptom score. Spheres denote centroids (i.e. center of mass) for each of the patient diagnostic groups and control subjects. Alternative views showing individual patients and controls projected into the PCA solution are shown in Fig. S3A-F.
all 436 subjects). This similarity was high for all sites (Fig. S6). Third, we verified that the predicted-versus-observed PCA score similarity also holds via a k-fold cross-validation (for k=2 to 10, Fig. 2C & Fig. S5). Finally, across all 5 PCs, predicted-to-observed similarity of actual loadings on to each PC was remarkably high. Moreover, PCA loadings were stable when testing via leave-site-out analysis, 5-fold cross-validation and across 1,000 independent split-half replications (Fig. 2D). For each run of split-half replication, the subjects were split into 2 sets and a PCA was performed independently in both sets. The loadings from both PCs were then compared using a Pearson’s correlation. Importantly, results were not driven by medication status or dosage (Fig. S7). Collectively, these data-reduction analyses strongly support a stable and replicable low-rank PSD symptom geometry.

**Dimensionality-Reduced PSD Symptom Geometry Reveals Novel and Robust Neuro-Behavioral Relationships.** Next, we tested if the dimensionality-reduced symptom geometry can identify robust and novel patterns of neural variation across the PSD. Across all neural analyses we used global brain connectivity (GBC), the column-wise mean of the full FC matrix because it yields a parsimonious metric reflecting how “globally” coupled a particular area is to the rest of the brain (35) (see Methods). Furthermore, we selected GBC because: i) the metric is agnostic regarding the location of “dysconnectivity” as it weights each area equally; ii) it yields an interpretable dimensionality-reduction of the full FC matrix; iii) unlike the full FC matrix or other abstracted measures, GBC produces a neural map, which can be related to other independent neural maps (e.g. gene expression or pharmacology maps, discussed below).

All 5 PCs captured unique GBC variation patterns across the PSD (Fig. S8). We highlight the PC3 solution (i.e. PC3 “Psychosis Configuration”) to illustrate the benefit of the low-rank PSD symptom geometry for neuro-behavioral mapping relative to traditional aggregate PANSS symptom scales. The relationship between total PANSS Positive scores and GBC across N=436 PSD patients (Fig. 3A) was statistically modest (Fig. 3B) and surprisingly no areas survived whole-brain type-I error protection (Fig. 3C, p<0.05). In contrast, regressing PC3 data-driven scores onto GBC across N=436 patients revealed a robust symptom-to-neural $\beta_{PC3\,GBC}$ map (Fig. 3E-F), which survived whole-brain type-I error protection. Of note, the PC3 “Psychosis Configuration” axis
is bi-directional whereby individuals who score either highly positively or negatively are symptomatic. Therefore, a high positive PC3 score was associated with both reduced GBC across insular and superior dorsal cingulate cortices, thalamus, and anterior cerebellum and elevated GBC across precuneus, medial prefrontal, inferior parietal, superior temporal cortices and posterior lateral cerebellum – consistent with the “default-mode network” (36). A high negative PC3 score would exhibit the opposite pattern. Critically, this robust symptom-to-neural mapping emerged despite no DSM diagnostic group differences in PC3 scores (Fig. 3D). The diverging nature of this axis may be captured in the ICA solution, when orthogonality is not enforced (Fig. S11).

Moreover, the PC3 neuro-behavioral map exhibited improved statistical properties relative to other GBC maps computed from traditional aggregate PANSS symptom scales (Fig. 3G-J).

**Univariate Neuro-Behavioral Map of Psychosis Configuration is Reproducible.** After observing improved PC3 neuro-behavioral statistics, we tested if these symptom-to-neural map patterns replicate. Recent attempts to derive stable symptom-to-neural mapping using multivariate techniques, while inferentially valid, have not replicated (25). This fundamentally relates to the tradeoff between the sample size needed to resolve multivariate neuro-behavioral solutions and the size of the feature space. To mitigate the feature size issue we re-computed the $\beta_{PC3 \text{GBC}}$ maps us-
Neural data were parcellated using a whole-brain functional parcellation (34) before computing GBC at the single patient level. This effects indicates neural feature data reduction via a functional neural parcellation (34) improves signal-to-noise ratio. Correlation between the value of each parcel in the regression model computed using patients in Fold 1 and the full PSD sample (N=436) model. Each point is a single parcel (718 parcels total). The resulting normalized coefficient map from the leave-one-fold-out model was highly similar to the map obtained from the full PSD sample model (r=0.924).

Summary of leave-one-site-out regression for symptom-to-GBC regression for the first half (H1) patients, shown here for one exemplar run out of 1000 split-half validations. (L) Correlation across 718 parcels between the H1 predicted coefficient map (i.e. panel K) and the observed coefficient map for H1 patient sample. (M-N) The same analysis as K-L is shown for patients in H2, indicating a striking consistency in Psychosis Configuration neural map correspondence.

Fig. 3  Parcellate 5-Fold Bootstrapping for All Maps

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Symptom-to-Neural Map

Observed betas w/o Site 3 (N=329)

Predicted betas

Full model betas

Fig. 4. Parcellated neuro-behavioral GBC maps reflecting psychosis configuration are statistically robust and replicable. (A) Z-scored PC3 Psychosis Configuration neural map at the ‘dense’ (full CIFTI resolution) level. (B) Neural data were parcellated using a whole-brain functional parcellation (34) before computing GBC at the patient level, which yields stronger statistical values in the Z-scored Psychosis Configuration GBC neural map as compared to when parcellation was performed after computing GBC for each patient. (C) A comparison of pre- and post-GBC parcellation Psychosis Configuration GBC maps shows a statistical improvement if parcellation is applied prior to computing GBC at the single patient level. This effects indicates neural feature data reduction via a functional neural parcellation (34) improves signal-to-noise for the group-level statistical map. (D) Summary of similarity for all symptom-to-GBC maps (PCs and traditional symptom scales) across 5-fold cross-validation. Boxplots show the range of r values between coefficient maps for each fold and the full model. (E) Normalized coefficient map from regression of individual patients’ Psychosis Configuration scores onto parcellated GBC data, shown here for a subset of patients from Fold 1 out of 5 (N=349). The greater the magnitude of the coefficient for a parcel, the stronger the statistical relationship between GBC of that parcel and Psychosis Configuration PC score. (F) Correlation between the value of each parcel in the regression model computed using patients in Fold 1 and the full PSD sample (N=436) model. Each point is a single parcel (718 parcels total). The resulting normalized coefficient map from the leave-one-fold-out model was highly similar to the map obtained from the full PSD sample model (r=0.924). (G) Summary of leave-one-site-out regression for all symptom-to-GBC maps. Regression of symptom scores onto parcellated GBC data, each time leaving out subjects from one site, resulted in highly similar maps. This highlights that the relationship between Psychosis Configuration and GBC variation is robust and not driven by a specific site. (H) Normalized coefficient map from regression of individual subject scores along Psychosis Configuration onto neural parcellated GBC data for all patients except one site. For example, Site 3 is excluded here given that it recruited the most patients (and therefore may have the greatest statistical impact on the full model). (I) Correlation between the value of each parcel in the regression model computed using all patients minus Site 3, and the full PSD sample model, again indicating a consistent neuro-behavioral relationship. (J) Split-half replication of Psychosis Configuration-to-GBC mapping. For each stratified split-half replication run, the full sample was randomly split into two halves (H1 and H2) with the proportion of each diagnostic group (BPP, SADP, SZP) preserved within each half. PCA scores for subjects in each half are computed using the PC loadings from the PCA conducted using subjects in the other half. Observed PCA scores are computed from a PCA on the same half-sample of subjects. Each set of scores were then regressed against parcellated GBC for subjects in a given half, resulting in a coefficient map for each PC (as in Fig. 3) reflecting the strength of the relationship between PC score and GBC across subjects. The coefficient maps for predicted and observed scores were then cross-correlated for PCs 1-5. This process was repeated 1,000 times. Bar plots show the mean correlation across the 1,000 runs; error bars show standard error of the mean. Note that the split-half effect for PC1 was exceptionally robust. The split-half consistency for PC3, while lower, was still highly robust and well above chance. (K) Normalized coefficient map from Psychosis Configuration-to-GBC regression for the first half (H1) patients, shown here for one exemplar run out of 1000 split-half validations. (L) Correlation across 718 parcels between the H1 predicted coefficient map (i.e. panel K) and the observed coefficient map for H1 patient sample. (M-N) The same analysis as K-L is shown for patients in H2, indicating a striking consistency in Psychosis Configuration neural map correspondence.
Fig. 5. Multivariate behavioral-to-neural feature mapping using canonical correlation analysis (CCA). (A) CCA computes a relationship between two sets of observed variables (in this case, a matrix $B$ of behavioral measures and a matrix $N$ of neural measures across all patients), by solving for two transformation matrices $\Psi$ and $\Theta$ such that the correlations between the latent variables are maximized (represented as columns along the transformed ‘latent’ matrices $U$ and $V$). Here each column in $U$ and $V$ is referred to as a canonical variate (CV); each corresponding pair of CVs (e.g. $U_i$ and $V_i$) is referred to as a canonical mode. (B) CCA maximized correlations between the CVs (i.e. matrices $U$ and $V$) (C) Screeplot showing canonical modes for the CVs obtained from 180 neural features (cortical GBC symmetric across hemispheres) and 36 single-item PANSS and BACS symptom measures. Inset illustrates the correlation ($r=0.85$) between the CV of the first mode, $U_1$ and $V_1$ (note that the correlation was not driven by a separation between diagnoses). Correlations between each subsequent pair of variates is computed from the residuals of the previous pair. Modes 9 and 12 remained significant after FDR correction across all 36 modes. (D) CCA was obtained from 180 neural features and 5 low-dimensional symptom scores derived via the PCA analysis. Here all modes remained significant after FDR correction. Dashed black line shows the null calculated via a permutation test with 5,000 shuffles; grey bars show 95% confidence interval. (E) Correlation between the behavioral data matrix $B$ and the neural data matrix weighted by the transformation matrix ($N\Theta$) reflects the amount of variance explained in $B$ that can be explained by the final latent neural matrix $V$. Put differently, this transformation calculates how much of the symptom variation can be explained by the latent neural features. (F) Proportion of symptom variance explained by each of the neural CVs in a CCA performed between 180 neural features and all 36 behavioral measures. Inset shows the total proportion of behavioral variance explained by the neural variates. (G) Proportion of total behavioral variance explained by each of the neural CVs in a CCA performed between 180 neural features and the 5 low-dimensional symptom scores derived via the PCA analysis. While CCA using symptom PCs has fewer dimensions and thus lower total variance explained (see inset), each neural variate explains a higher amount of symptom variance than seen in $F$, suggesting that CCA could be further optimized by first obtained a principled low-rank symptom solution. Dashed black line shows the null calculated via a permutation test with 5,000 shuffles; grey bars show 95% confidence interval. Behavioral-to-neural mapping is shown in Fig. S14. (H) Characterizing CV symptom configurations using CV3 as an example. Distributions of CV3 scores by each DSM diagnostic group. All scores are normalized to controls. (I) Behavioral canonical factor loadings for CV3. (J) Loadings of the original symptom items for CV3. (K) Neural canonical factor loadings for CV3. (L) Within-sample CCA cross-validation appeared robust (see Fig. S16). However, a split-half CCA replication using two independent non-overlapping patient samples was not reliable. Bar plots show the mean correlation for each CV between the first half (H1) and the second half (H2) CCA, each computed across 1,000 runs. Left, split-half replication of the behavioral PC loadings matrix $\Psi$; Middle: individual behavioral item loadings; and Right: the neural loadings matrix $\Theta$. Error bars show the standard error of the mean. The neural loadings matrix $\Theta$ in particular was not stable. Scatterplot shows the correlation between CV3 neural loadings for H1 vs. H2 for one example CCA run, illustrating lack of reliability. (M) Leave-one-subject-out cross-validation further highlights CCA instability. Here, CCA was computed for all except one patient (N=435). The loadings matrices were then used to compute the predicted neural ($N$) and behavioral ($B$) scores for the left-out patient. This was repeated across all patients. This yielded predicted neural and predicted behavioral scores for each patient for each of the five CVs. These correlations were far lower than the canonical correlation values obtained in the full CCA model (shown as red lines). This is highlighted in the scatterplot at far right for CV3 ($r=0.12$, whereas the full sample CCA was $r=0.68$). CCA solutions for subcortex parcels only (Fig. S12) and network-level neural feature reduction Fig. S13 are shown in the Supplement.
ing a functionally-derived whole-brain parcellation via the recently-validated CAB-NP atlas (34, 37) (Methods). Here, a functional parcellation is a principled way of neural feature reduction (to 718 parcels) that can also appreciably boost signal-to-noise (34, 37). Indeed, parcellating the full-resolution “dense” resting-state signal for each subject prior to computing GBC statistically improved the group-level neuro-behavioral maps compared to parcellating after computing GBC (Fig. 4A–C, all maps in Fig. S9). Results demonstrate that the univariate symptom-to-neural mapping was indeed stable across 5-fold bootstrapping, leave-site-out, and split-half cross-validations (Fig. 4D–N, see Supplementary Note 3), yielding consistent neuro-behavioral PC3 maps. Importantly, the neuro-behavioral maps computed using ICA showed comparable results (Fig. S11).

A Multivariate PSD Neuro-Behavioral Solution Can be Computed but is Not Reproducible with the Present Sample Size. Several studies have reported “latent” neuro-behavioral relationships using multivariate statistics (15, 38, 39), which would be preferable because they maximize covariation across neural and behavioral features simultaneously. However, concerns have emerged whether such multivariate results will replicate due to the size of the feature space (25). Nevertheless, we tested if results improve with canonical correlation analysis (CCA) (40) which maximizes relationships between linear combinations of symptom (B) and neural measures (N) across all subjects (Fig. 5A).

We examined two CCA solutions using symptom scores in relation to neural features: i) all item-level scores from the PANSS/BACS (Fig. 5C&F); ii) data-reduced PC scores (Fig. 5D&G, see Methods). In turn, to evaluate if the number of neural features affects the solution we computed CCA using: i) 718 bilateral whole-brain parcels derived from the brain-wide cortico-subcortical CAB-NP parcellation (34, 37); ii) 359 bilateral subcortex-only parcels; iii) 192 symmetrized subcortical parcels; iv) 180 symmetrized cortical parcels; v) 12 functional networks defined from the parcellation (34, 37). Notably, 4 out of the 5 CCA solutions were not robust: 718 whole-brain & 359 subcortex-only parcel solutions did not produce stable results according to any criterion (Fig. S14), whereas the symmetrized subcortical (192 features, Fig. S12) and network-level (12 features, Fig. S13) solutions captured statistically modest effects relative to the 180 symmetrized cortical CCA results (Fig. 5B-D). Therefore, we characterized the 180-parcel CCA solution further: Only 2 out of 36 CCA modes for the item-level symptom solution survived permutation testing and false discovery rate (FDR) correction (Fig. 5C). In contrast, all 5 modes computed using PC scores survived both permutation and FDR correction (Fig. 5D). Critically, we found that no single CCA mode computed on item-level symptoms captured more variance than the CCA modes derived from PC scores, suggesting that the PCA-derived dimensions capture more neurally-relevant variation than any one single clinical item (Fig. 5E-G). Additional detailed CCA results are presented in Supplementary Note 4 and Fig. S14.

We highlight an example canonical variate (CV) solution across both behavioral and neural effects for CV3. We show the CV3 scores across diagnostic groups normalized to controls Fig. 5H as well as how the CV3 loads onto each PC (Fig. 5I, full results are shown in Fig. S15). The negative loadings on to PCs 1, 4, and 5 and the high positive loadings on to PC3 in Fig. 5I indicate that CV3 captures some shared variation across PCs. This can also be visualized by computing how the CV3 projects onto the original 36 behavioral items (Fig. 5J). Finally, the neural loadings for CV3 are shown in Fig. 5K and data for all 5 CVs are shown in Fig. S14.

Next, we tested if the 180-parcel CCA solution is stable and replicable, as done with PC-to-GBC univariate results. The CCA solution was robust when tested with k-fold and leave-site-out cross-validation (Fig. S16) because these methods use CCA loadings derived from the full sample. However, the CCA loadings did not replicate in non-overlapping split-half samples (Fig. 5L, see Supplementary Note 5). Moreover, a leave-one-subject-out cross-validation revealed that removing a single subject from the sample affected the CCA solution such that it did not generalize to the left-out subject (Fig. 5M). As noted, the PCA-to-GBC univariate mapping was substantially more reproducible for all attempted cross-validations relative to the CCA approach, reflecting the fact that substantially more power is needed to resolve a stable multi-variate neuro-behavioral effect with these many features (26). Therefore, we leverage the univariate neuro-behavioral result for subsequent subject-specific model optimization and comparisons to molecular neuroimaging maps.

A Major Proportion of Overall Neural Variance May Not be Relevant for Psychosis Symptoms. Most studies look for differences between clinical and control groups, but to our knowledge no study has tested whether both PSD and healthy controls actually share a major portion of neural variance that may be present across all people. If the bulk of the neural variance is similar across both PSD and CON groups then including this clinically-irrelevant neural signal might obscure neuro-behavioral relationships that are clinically relevant. To test this, we examined the shared variance structure of the neural signal for all PSD patients (N=436) and all controls (N=202) independently by conducting a PCA on the GBC maps (see Methods). Patients’ and controls’ neural signals were highly similar for each of the first 3 neural PCs (>30% of all neural variance in each group) (Fig. S17A–J). These PCs may reflect a “core” shared symptom-irrelevant neural variance that generalizes across all people. These data suggest that the bulk of neural variance in PSD patients may actually not be symptom-relevant, which highlights the importance of optimizing symptom-to-neural mapping.

Optimizing Neuro-Behavioral Mapping Features for Personalized Prediction via Dimensionality-reduced Symptoms. Above we demonstrate that PC scores can be reliably predicted across sites and cross-validation approaches (Fig. 2). Here we show that leave-one-subject-out cross-validation yields reliable effects for the low-rank symptom PCA solution (Fig. 6A). This stable single-subject PC score...
prediction provides the basis for testing if the derived neural maps can yield an individually-reliable subset of features. To this end, we developed a univariate quantitative neural feature selection framework (Fig. S18) based on data-reduced PC scores (i.e. PC3 score). Specifically, we computed a dot product GBC metric ($d_{pGBC}$) that provides an index of similarity between an individual GBC topography relative to a “reference” group-level PC GBC map (see Methods and Fig. S18). Using this $d_{pGBC}$ index we found, via a feature selection step-down regression, a subset of parcels for which the symptom-to-neural feature statistical association was maximal (Fig. 6A). For PC3 we found $P = 39$ maximally predictive parcels out of the group map. Specifically, the relationship between PC3 symptom scores and $d_{pGBC}$ values across subjects was maximal (Fig. 6B, top panel, $r = 0.36$) as was the relationship between predicted $d_{pGBC}^{pred}$ vs. observed $d_{pGBC}^{obs}$ (Fig. 6B, bottom panel, $r = 0.31$) (see Fig. S19 for all PCs). Importantly, the “subset” feature map (i.e. $d_{pGBC}^{obs}$) exhibited improved statistical properties relative to the full map (i.e. $d_{pGBC}^{obs}$) (see Fig. 6C). Furthermore, the relationship between observed vs. predicted subset feature maps (i.e. $r[d_{pGBC}^{obs}, d_{pGBC}^{pred}]$) was highly consistent across DSM diagnoses (Fig. 6E). Finally, a single patient is shown for whom the correlation between their predicted and observed subset feature maps was high (i.e. $r[d_{pGBC}^{pred}, d_{pGBC}^{obs}]$), Fig. 6F), demonstrating that the dimensionality-reduced symptom scores can be used to quantitatively optimize individual neuro-behavioral map features.

Single Patient Evaluation via Neuro-Behavioral Target Map Similarity. We demonstrated a quantitative framework for computing a neuro-behavioral model at the single-patient level. This brain-behavior space (BBS) model was optimized along a single dimensionality-reduced symptom axis (i.e. PC score). Next, we tested a hybrid patient selection strategy by first imposing a PC-based symptom threshold, followed by a target neural similarity threshold driven by the most highly predictive neuro-behavioral map features. This is described in Methods, Supplementary Note 6, and Fig. S20-S21. We found that for patients with either a high positive or negative PC symptom score, the neuro-behavioral relationship was robust (Fig. S20A-D), and these patients could be reliably selected for that particular axis (Fig. S20E,G). Conversely, patients with a low absolute PC score showed a weak relationship with symptom-relevant neural features. This is intuitive because these individuals do not vary along the selected PC symptom axis. We also tested this patient selection strategy on an independent cross-diagnostic “replication” sample, which yielded consistent results (Fig. S20F,H). Collectively, these results show that data-driven symptom scores can pinpoint individual patients for whom their neural varia-
were selected from “discovery” PSD dataset for illustrative purposes. Patient PC5 loads highly negatively on the PC5 axis (blue) and Patient YPC3 loads highly positively (yellow). Both of these patients scored above the neural and behavioral thresholds for PC3 defined in the “discovery” PSD dataset. Patient XPC3 loads highly negatively on the PC5 axis and Patient YPC3 loads highly positively. Density plots show the projected PC scores for Patients XPC3 and YPC3 overlaid on distributions of PC scores from the discovery PSD sample. (B) Neural maps show cortical and subcortical $\Delta GBC_{obs}$ for the two patients XPC3 and YPC3 specifically reflecting a difference from the mean PC3. The similarity of $\Delta GBC_{obs}$ and the $\beta_{PC3}$GBC map within the most predictive neural parcels for PC3 (outlined in green). Note that the sign of neural similarity to the reference PC3 map and the sign of the PC3 score is consistent for these two patients. (C) The selected PC3 map (parcels outlined in green) is spatially correlated to the neural map reflecting the change in GBC after ketamine administration ($p = 0.76$, Methods). Note that Patient XPC3, who exhibits $\Delta GBC_{obs}$ that is anti-correlated to the ketamine map also expresses depressive moods symptoms (panel A). This is consistent with the possibility that this person may clinically benefit from ketamine administration, which may elevate connectivity in areas where they show reductions ($p = 41$). In contrast, Patient YPC3 may exhibit an exacerbation of their psychosis symptoms given that their $\Delta GBC_{obs}$ is positively correlated with the ketamine map. (D) Data for two individual patients from the replication dataset are highlighted for PC3: $Q_{PC5}$ (blue) and $Z_{PC5}$ (yellow). Note that no patients in the replication dataset were selected for PC5 so both of these patients were selected from “discovery” PSD dataset for illustrative purposes. Patient $Q_{PC5}$ loads highly negatively on the PC5 axis and Patient $Z_{PC5}$ loads highly positively. Density plots show the projected PC scores for Patients $Q_{PC5}$ and $Z_{PC5}$ overlaid on distributions of PC scores from the discovery PSD sample. (E) Neural maps show cortical and subcortical $\Delta GBC_{obs}$ for Patients $Q_{PC5}$ and $Z_{PC5}$, which are highly negatively and positively correlated with the selected PC5 map respectively. (F) The selected PC5 map (parcels outlined in green) is spatially anti-correlated with the LSD response map ($p = 0.44$, see Methods), suggesting that circuits modulated by LSD (i.e. serotonin, in particular 5-HT2A) may be relevant for the PC5 symptom expression. Here a serotonin receptor agonist may modulate the neuro-behavioral profile of Patient $Q_{PC5}$, whereas an antagonist may be effective for Patient $Z_{PC5}$.
tion strongly maps onto a target neural reference map. These data also highlight that both symptom and neural information for an independent patient can be quantified in the reference ‘discovery’ BBS using their symptom data alone.

**Subject-Specific PSD Neuro-Behavioral Features Track Neuropharmacological Map Patterns.** Next, we use the personalized BBS selection in a proof-of-concept framework for informing molecular mechanism of possible treatment response by relating subject-specific BBS to independently-acquired pharmacological neuroimaging maps. Here we examine two mechanisms implicated in PSD neuropathology via ketamine, a N-methyl-D-aspartate (NMDA) receptor antagonist (42), and lysergic acid diethylamide (LSD), primarily a serotonin receptor agonist (29, 43, 44). We first quantified individual subjects’ BBS “locations” in the established reference neuro-behavioral geometry. The radarplot in Fig. 7A shows original symptoms whereas Fig. 7B shows $\Delta GBC_{obs}$ maps for two patients from the replication dataset (denoted here with $X_{PCA}$ and $Y_{PCA}$, see Fig. S10 for other example patients). Both of these patients exceeded the neural and behavioral BBS selection indices for PC3 (defined independently in the “discovery” dataset, Fig. S20C). Furthermore, both patients exhibited neuro-behavioral variation in line with their expected locations in the BBS geometry. Specifically, patient $X_{PCA}$ from the replication dataset scored highly negatively on the PC3 axis defined in the “discovery” PSD sample (Fig. 7A). In contrast, patient $Y_{PCA}$ scored positively on the PC3 axis. Importantly, the correlation between the $\Delta GBC_{obs}$ map for each patient and the group-reference $\beta_{PC3}GBC$ was directionally consistent with their symptom PC score (Fig. 7B-C).

We then tested if the single-subject BBS selection could be quantified with respect to a neural map reflecting glutamate receptor manipulation, a hypothesized mechanism underlying PSD symptoms (45). Specifically, we used an independently collected ketamine infusion dataset, collected in healthy adult volunteers during resting-state fMRI (46). As with the clinical data, here we computed a $\Delta GBC$ map reflecting the effect of ketamine on GBC relative to placebo (Methods). The maximally-predictive PC3 parcels exhibited high spatial similarity with the ketamine map ($p=0.76$, see Methods), indicating that the $\Delta GBC$ pattern induced by ketamine tracks with the GBC pattern reflecting PC3 symptom variation.

Critically, because $X_{PCA}$ is negatively loaded on the PC3 symptom axis, an NMDA receptor antagonist like ketamine may modulate symptom-relevant circuits in a way that reduces similarity with the PC3 map. This may in turn have an impact on the PC3-like symptoms. Consistent with this hypothesis, $X_{PCA}$ expresses predominantly depressive symptoms (Fig. 7A), and ketamine has been shown to act as an anti-depressant (41). This approach can be applied for patients that load along another axis, such as PC5. Fig. 7D-E shows the symptom and neural data for two patients whom met thresholds for PC5 selection (Fig. S20C). Notably, the selected PC5 map is anti-correlated with a $\Delta GBC$ map reflecting LSD vs. placebo effects (29) ($\rho=-0.44$, Fig. 7F). Hence areas modulated by LSD may map onto behavioral variation along PC5. Consequently, serotonergic modulation may be effective for treating $Q_{PC5}$ and $Z_{PC5}$, via an antagonist or an agonist respectively. These differential similarities between pharmacological response maps and BBS maps (Fig. S23) can be refined for quantitative patient segmentation.

**Group-Level PSD Neuro-Behavioral Features Track Neural Gene Expression Patterns.** To further inform molecular mechanism for the derived BBS results, we compared results with patterns neural gene expression profiles derived from the Allen Human Brain Atlas (AHBA) (31, 32)(Fig. 8A, Methods). Specifically, we tested if BBS cortical topographies, which reflect stable symptom-to-neural mapping along PSD, covary with the expression of genes implicated in PSD neuropathology. We focus here on serotonin and GABA receptors as well as interneuron markers (SST, somatostatin; PV ALB, parvalbumin). Fig. 8B shows the distribution of correlations between the PC3 map and the cortical expression patterns of 20,200 available AHBA genes (results for other PCs are shown in Fig. S24). Seven genes of interest are highlighted, along with their cortical expression topographies and their similarity with the PC3 BBS map (Fig. 8C-E). This BBS-to-gene mapping can potentially reveal novel therapeutic molecular targets for neuro-behavioral variation. For example, the HTR1E gene, which encodes the serotonin 5-HT$_{1E}$ receptor, is highly correlated with the PC3 BBS map. This could drive further development of novel selective ligands for this receptor, which are not currently available (47).

**Discussion**

We found a robust and replicable symptom-to-neural mapping across the psychosis spectrum that emerged from a low-dimensional symptom solution. Critically, this low-rank symptom solution was predictive of a neural circuit pattern, which reproduced at the single-subject level. In turn, we show that the derived PSD symptom-to-neural feature maps exhibit spatial correspondence with independent pharmacological and gene expression neural maps that are directly relevant for PSD neurobiology.

**Deriving an Individually Predictive Low-Dimensional Symptom Representation Across the Psychosis Spectrum.** Psychosis spectrum is associated with notable clinical heterogeneity such deficits in cognition as well as altered beliefs (i.e. delusions), perception (i.e. hallucinations), and affect (i.e. negative symptoms) (24). This heterogeneity is captured by clinical instruments that quantify PSD symptoms across dozens of specific questions and ratings. This yields a high-dimensional symptom space that is intractable for reliable mapping of neural circuits (26). Here we show that a low-rank solution captures principal axes of PSD symptom variation, a finding in line with prior work in schizophrenia (18–24, 48).
These results highlights two key observations: i) Existing symptom reduction studies (even those in schizophrenia specifically) have not evaluated solutions that include cognitive impairment – a hallmark deficit across the psychosis spectrum (17). Here we show that cognitive function captures a notable portion of the symptom variance independent of other axes. We observed that cognitive variation captured 10% of PSD sample variance even after accounting for ‘General’ psychopathology. ii) No study has quantified cognitive deficit variation via dimensionality reduction across multiple PSD diagnoses along with core psychosis symptoms. We found that cognitive deficits load across several PCs, but the pattern of loading was particularly evident for executive function sub-scores on certain axes (e.g. PC5 symptom axis solution was not stable if a single composite cognitive score was used). While existing studies evaluated stability of data-reduced solutions within a single DSM category (18, 24, 49, 50), current results show that lower-dimensional PSD symptoms solutions can be reproducibly obtained across DSM diagnoses. For each data-reduced axis some PSD patients received a score near zero. This does not imply that these patients were unimpaired; rather, the symptom configurations for these patients were orthogonal to variation along this specific symptom axis. The observation that PSD is associated with multiple complex symptom dimensions highlights an important intuition that may extend to other mental health spectra. Also, the PSD symptom axes reported here are neither definitive nor exhaustive. In fact, close to 50% of all clinical variance was not captured by the symptom PCA – an observation often overlooked in symptom data-reduction studies, which focus on attaining ‘predictive accuracy’. Such studies rarely consider how much variance remains unexplained in the final data-reduced model and, relatedly, if the proportion of explained variance is reproducible across samples. This is a key property for reliable symptom-to-mapping. Thus, we tested if this replicable

Fig. 8. Psychosis spectrum neuro-behavioral maps track neural gene expression patterns computed from the Allen Human Brain Atlas (AHBA). (A) The symptom loadings and the associated neural map jointly reflect the PC3 brain-behavioral space (BBS) profile, which can be quantitatively related to human cortical gene expression patterns obtained from the AHBA (32). (B) Distribution of correlation values between the PC3 BBS map and ˜20,000 gene expression maps derived from the AHBA dataset. Specifically, AHBA gene expression maps were obtained using DNA microarrays from six postmortem brains, capturing gene expression topography across cortical areas. These expression patterns were then mapped onto the cortical surface models derived from the AHBA subjects’ anatomical scans and aligned with the Human Connectome Project (HCP) atlas, described in prior work and methods (32). Note that because no significant inter-hemispheric differences were found in cortical gene expression all results were symmetrized to the left hemisphere, resulting in 180 parcels. We focused on a select number of psychosis-relevant genes – namely genes coding for the serotonin and GABA receptor subunits and interneuron markers. Seven genes of interest are highlighted with dashed lines. Note that the expression pattern of HTR2C (green dashed line) is at the low negative tail of the entire distribution, i.e. highly anti-correlated with PC3 BBS map. Conversely, GABRA1 and HTR1E are on the far positive end, reflecting a highly similar gene-to-BBS spatial pattern. (C) Upper panels show gene expression patterns for two interneuron marker genes, somatostatin (SST) and parvalbumin (PVALB). Positive (yellow) regions show areas where the gene of interest is highly expressed, whereas negative (blue) regions indicate low expression values. Lower panels highlight all gene-to-BBS map spatial correlations where each value is a symmetrized cortical parcel (180 in total) from the HCP atlas parcellation. (D) Gene expression maps and spatial correlations with the PC3 BBS map for two GABA\(_\alpha\) receptor subunit genes: GABRA1 and GABRA5. (E) Gene expression maps and spatial correlations with the PC3 BBS map for three serotonin receptor subunit genes: HTR1E, HTR2C, and HTR2A.
low-dimensional PSD symptom space robustly mapped onto neural circuit patterns.

Leveraging a Robust Low-Dimensional Symptom Representation for Mapping Brain-Behavior Relationships. We show that the dimensionality-reduced symptom space improved the mapping onto neural circuit features (i.e. GBC), as compared to a priori item-level clinical symptoms (Fig. 3). This symptom-to-neural mapping was highly reproducible across various cross-validation procedures, including split-half replication (Fig. 4). The observed statistical symptom-to-neural improvement after dimensionality reduction suggests that data-driven clinical variation more robustly covaried with neural features. As noted, the low-rank symptom axes generalized across DSM diagnoses. Consequently, the mapping onto neural features (i.e. GBC) may have been restricted if only a single DSM category or clinical item was used. Importantly, as noted, traditional clinical scales are uni-directional (i.e. zero is asymptomatic, hence there is an explicit floor). Here, we show that data-driven symptom axes (e.g. PC3) were associated with bi-directional variation (i.e., no explicit floor effect). Put differently, patients who score highly on either end of these data-driven axes are severely symptomatic but in very different ways. If these axes reflect clinically meaningful phenomena at both tails, then they should more robustly map to neural feature variation, which is in line with reported effects. Therefore, by definition, the derived map for each of the PCs will reflect the neural circuitry that may be modulated by the behaviors that vary along that PC (but not others). For example, we named the PC3 axis “Psychosis Configuration” because of its strong loadings onto conventional “positive” and “negative” PSD symptoms. This PC3 “Psychosis Configuration” showed strong positive variation along neural regions that map onto the canonical default mode network (DMN), which has frequently been implicated in PSD (51–56). In turn, this bi-directional “Psychosis Configuration” axis showed strong negative variation along neural regions that map onto the sensory-motor and associative control regions, also strongly implicated in PSD (? ). This ‘bi-directional’ neuro-behavioral map property may be desirable for identifying neural features that support individual patient selection.

Deriving Individually Actionable Brain-Behavior Mapping Across the Psychosis Spectrum. Deriving a neuro-behavioral mapping that is resolvable and stable at the individual patient level is a necessary benchmark for deploying symptom-to-neural ‘biomarkers’ in a clinically useful way. Therefore, there is increasing attention placed on the importance of achieving reproducibility in the psychiatric neuroimaging literature (57–60), which becomes especially important for individualized symptom-to-neural mapping. Recently, several efforts have deployed multivariate methods to quantify symptom-to-neural relationships (15, 38, 39, 61–63), highlighting how multivariate techniques may provide clinically innovative insights. However, such methods face overfitting risk for high-dimensional but underpowered datasets (25), as recently shown via formal generative modeling (26).

Here we attempted to use multivariate solutions (i.e. CCA) to quantify symptom and neural feature co-variation. In principle, CCA is theoretically well-suited to address the brain-behavioral mapping problem. However, symptom-to-neural mapping using CCA across either parcel-level or network-level solutions was not reproducible even when using a low-dimensional symptom solutions as a starting point. Therefore, while CCA (and related multivariate methods such as partial least squares) are theoretically appropriate (and regularization methods such as sparse CCA may help), in practice many available psychiatric neuroimaging datasets may not provide sufficient power to resolve stable multivariate symptom-to-neural solutions (26). A key pressing need for forthcoming studies will be to use multivariate power calculators to inform sample sizes needed for resolving stable neuro-behavioral geometries at the single subject level. Consequently, we tested if a low-dimensional symptom solution can be used in a univariate symptom-to-neural model to optimize individually predictive features. Indeed, we found that a univariate brain-behavioral space (BBS) relationship can result in neural features that are stable for individualized prediction. Critically, we found that a patient exhibited a high PC symptom score, they were more likely to exhibit a topography of neural ΔGBC (i.e. difference relative to the group mean reference) that was topographically similar to PC symptom neural map. This suggests that optimizing such symptom-to-neural mapping solutions (and ultimately extending them to multivariate frameworks) can inform cross-diagnostic patient segmentation with respect to symptom-relevant neural features. Importantly, this could directly inform patient identification based on neural targets that are of direct symptom relevance for clinical trial design.

Utilizing Independent Molecular Neuroimaging Maps to ‘Benchmark’ Symptom-Relevant Neural Features. Selecting single patients via stable symptom-to-neural mapping BBS solutions is necessary for individual patient segmentation, which may ultimately inform treatment indication. However, are the derived symptom-to-neural maps related to a given mechanism? Here we highlight two ways to ‘benchmark’ the derived symptom-to-neural maps related to a given mechanism. We show a proof-of-principle framework for quantifying derived symptom-to-neural reference maps with two PSD-relevant neuropsychological manipulations derived in healthy adults via LSD and ketamine. These analyses revealed that selecting single patients, via the derived symptom-to-neural mapping solution, can yield above-chance quantitative correspondence to a given molecular target map. These data highlight an important effect: it is possible to construct a ‘strong inference’ (64) evaluation of single patients’ differential similarity to molecular target map. For instance, this approach could be applied to maps associated with already approved PSD treatments (such as clozapine, olanzapine, or chlorpromazine (65, 66)) to identify patients with symptom-to-neural configurations that best cap-
ture available treatment-covarying neural targets.

Relatedly, AHBA gene expression maps (32) may provide an a priori benchmark for treatment targets that may be associated with a given receptor profile. Here we show that identified BBS maps exhibit spatial correspondence with neural gene expression maps implicated in PSD – namely serotonin, GABA and interneuron gene expression. This gene-to-BBS mapping could be then used to select those patients that exhibit high correspondence to a given gene expression target. Collectively, this framework could inform empirically testable treatment selection methods (e.g. a patient may benefit from ketamine, but not serotonin agonists such as LSD/psilocybin). In turn, this independent molecular benchmarking framework could be extended to other approaches (e.g. positron emission tomography (PET) maps reflecting specific neural receptor density patterns (67, 68)) and iteratively optimized for quantitative patient-specific selection against actionable molecular targets.

**Considerations for Generalizing Solutions Across Time, Severity and Mental Health Spectra.** There are several constraints of the current result that require future optimization – namely the ability to generalize across time (early course vs. chronic patients), across a range of symptom severity (e.g. severe psychotic episode or persistent low-severity psychosis) and across distinct symptom spectra (e.g. mood). This applies to both the low-rank symptom solution and the resulting symptom-to-neural mapping. It is possible that the derived lower-dimensional symptom solution, and consequently the symptom-to-neural mapping solution, exhibits either time-dependent (i.e. state) or severity-dependent (i.e. trait) re-configuration. Relatedly, medication dose, type, and timing may also impact the solution.

Critically, these factors should constitute key extensions of an iteratively more robust model for individualized symptom-to-neural mapping across the PSD and other psychiatric spectra. Relatedly, it will be important to identify the ‘limits’ of a given BBS solution – namely a PSD-derived effect may not generalize into the mood spectrum (i.e. both the symptom space and the resulting symptom-to-neural mapping is orthogonal). It will be important to evaluate if this framework can be used to initialize symptom-to-neural mapping across other mental health symptom spectra, such as mood/anxiety disorders.

These types questions will require longitudinal and clinically diverse studies that start prior to the onset of full-blown symptoms (e.g. the North American Prodrome Longitudinal Study (NAPLS) (69, 70)). A corollary of this point is that "50% of unexplained symptom variance in the current PCA solution necessitates larger samples with adequate power to map this subtle but perhaps clinically essential PSD variation. Notably, the current cohort was adequately powered for symptom data reduction that drove univariate neural mapping. However, this sample was insufficiently powered for resolving stable multivariate symptom-to-neural relationships even with low-dimensional symptom features. Consequently, the limited sample size necessitated choices for dimensionality-reduction of the neural feature space in this study even for univariate analyses. While both parcellation and GBC constitute principled choices, symptom-relevant neural information may have been lost (which may be imbedded in a higher-dimensional space). One obvious solution is to increase sample sizes (e.g. via datasets such as the UK Biobank (71)). However, in parallel, it will be critical to develop neurobiologically-informed feature space reduction and/or to optimize the stability of multivariate solutions via regularization methods. Another improvement would be to optimize neural data reduction sensitivity for specific symptom variation (72).

Here we focused on the neural blood oxygen level dependent (BOLD) signal from fMRI. However, other modalities such as diffusion-weighted imaging (DWI), PET imaging, or electroencephalography (EEG) could be leveraged. Additional clinically-relevant information could be derived from genetics (such as polygenic risk scores (73–75)) or ecological momentary assessment (EMA) (76, 77), especially to parse state vs. trait biomarker variation.

Lastly, building on the proof-of-concept molecular neuroimaging comparisons, it will be imperative to eventually test such predictions in formal clinical trials. An actionable next step would be to optimize patient selection against existing treatments, which could result in higher success rates for drug development trials and potentially massive impact for developing new interventions. Critically, the opportunity to develop, validate, and refine an individual-level quantitative framework could deliver a more rapid and cost-effective way of pairing patients with the right treatments.

**Conclusions.** We show that complex and highly heterogeneous PSD symptom variation can be robustly reduced into a low-rank symptom solution that is cross-diagnostic, individually predictive, generalizable and incorporates cognitive deficits. In turn, the derived PSD symptom axes robustly mapped onto distinct yet replicable neural patterns, which were predictive at the single-patient level. Leveraging these stable results, we show a proof-of-principle framework for relating the derived symptom-relevant neural maps, at the individual patient level, with molecular targets implicated in PSD via LSD and ketamine neuro-pharmacological manipulations. Lastly, we used AHBA gene expression maps to show that identified PSD symptom-relevant neural maps covary with serotonin, GABA and interneuron gene expression patterns. This set of symptom-to-neural mapping results can be iteratively and quantitatively optimized for personalized treatment segmentation endpoints.

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Bibliography


Methods

Overall Data Collection and Study Design. We used publicly available behavioral and neural data from the Bipolar-Schizophrenia Network on Intermediate Phenotypes (BSNIP) consortium (11). All data were obtained from the National Data Archive (NDA) repository (https://nda.nih.gov/edit_collection.html?id=2274). Participants were collected at 6 sites across the United States. Full recruitment details are provided in prior publications (11, 78, 79). In brief, participants were recruited through advertising in Baltimore MD, Boston MA, Chicago IL, Dallas TX, and Hartford CT. All assessments were standardized across sites. Participants were excluded if they had i) a history of seizures or head injury resulting in >10 minutes loss of consciousness, ii) positive drug screen on the day of testing, iii) a diagnosis of substance abuse in the past 30 days or substance dependence in the past 6 months, iv) history of serious medical or neurological disorder that would likely affect cognitive functioning, v) history of serious medical or neurological disorder that would likely affect cognitive functioning, vi) insufficient English proficiency, or vii) an age-corrected Wide-Range Achievement Test (4th edition) reading test standard score <65. Additionally, participants were required to have had no change in medication and been clinically stable over the past month. Participants completed a SCID interview and were given a diagnosis via consensus from study clinicians; participants with an Axis I clinical psychosis diagnosis were additionally given assessments including the Positive and Negative Symptom Scale (PANSS) (28). Note that apart from the measures used in this study, the full BSNIP dataset includes a rich characterization of measures from multiple modalities, including electrophysiological, eye tracking, structural and diffusion neuroimaging, as well as a number of clinical batteries, which are not quantified in this study. We used data from a total of 638 participants with complete behavioral and neural data after preprocessing and quality control, including 202 healthy controls (CON), 167 patients diagnosed with schizophrenia (SZP), 119 patients with schizoaffective disorder (SADP), and 150 patients with bipolar disorder with psychosis (BPP). For full demographic and clinical characteristics of the sample see Table S1.

Neural Data Acquisition and Preprocessing. Participants completed a neural magnetic resonance imaging (MRI) scan at 3T, including resting-state functional blood-oxygen-level-dependent imaging (BOLD) and a magnetization-prepared rapid gradient-echo (MP-RAGE) sequence for T1 weighted data. Full details on scanners and acquisition parameters used at each of the sites have previously been described (80) and are summarized here in Table S2. Neuroimaging data were preprocessed using the Human Connectome Project (HCP) minimal preprocessing pipeline (81), adapted for compatibility with “legacy” data, which are now featured as a standard option in the HCP pipelines provided by our team (https://github.com/Washington-University/HCPpipelines/pull/156). These modifications to the HCP pipelines were necessary as the BSNIP data did not include a standard field map and did not incorporate a T2w high-resolution image. The adaptations for single-band BOLD acquisition have previously been described in detail (82).

A summary of the HCP pipelines is as follows: the T1-weighted structural images were first aligned by warping them to the standard Montreal Neurological Institute-152 (MNI-152) brain template in a single step, through a combination of linear and non-linear transformations via the FM-RIB Software Library (FSL) linear image registration tool (FLIRT) and non-linear image registration tool (FNIRT) (83). Next, FreeSurfer’s recon-all pipeline was used to segment brain-wide gray and white matter to produce individual cortical and subcortical anatomical segmentations (84). Cortical surface models were generated for pial and white matter boundaries as well as segmentation masks for each subcortical grey matter voxel. Using the pial and white matter surface boundaries, a ‘cortical ribbon’ was defined along with corresponding subcortical voxels, which were combined to generate the neural file in the Connectivity Informatics Technology Initiative (CIFTI) volume/surface ‘greyordinate’ space for each individual subject (81). BOLD data were motion-corrected by aligning to the middle frame of every run via
FLIRT in the initial NIFTI volume space. In turn, a brain-mask was applied to exclude signal from non-brain tissue. Next, cortical BOLD data were converted to the CIFTI gray matter matrix by sampling from the anatomically-defined gray matter cortical ribbon and subsequently aligned to the HCP atlas using surface-based nonlinear deformation (81). Subcortical voxels were aligned to the MNI-152 atlas using whole-brain non-linear registration and then the Freesurfer-defined subcortical segmentation applied to isolate the subcortical grayordinate portion of the CIFTI space.

After the HCP minimal preprocessing pipelines, movement scrubbing was performed (85). As done in our prior work (86), all BOLD image frames with possible movement-induced artifactual fluctuations in intensity were flagged using two criteria: frame displacement (the sum of the displacement across all six rigid body movement correction parameters) exceeding 0.5 mm (assuming 50 mm cortical sphere radius) and/or the normalized root mean square (RMS) (calculated as the RMS of differences in intensity between the current and preceding frame, computed across all voxels and divided by the mean intensity) exceeding 1.6 times the median across scans. Any frame that met one or both of these criteria, as well as the frame immediately preceding and immediately following, were discarded from further preprocessing and analyses. Subjects with more than 50% frames flagged using these criteria were excluded. Next, a high-pass filter (threshold 0.008 Hz) was applied to the BOLD data to remove low frequency signals due to scanner drift. In-house Matlab code was used to calculate the average variation in BOLD signal frequency signals due to scanner drift. In-house Matlab code was used to calculate the average variation in BOLD signal in the ventricles, deep white matter, and across the whole grey matter, as well as movement parameters. These signals, as well as their first derivatives to account for delayed effects, were then regressed out of the grey matter BOLD time series as nuisance variables (as any change in the BOLD signal due to these variables are persistent and likely not reflecting neural activity) (87). It should be noted that using global signal regression to remove spatially persistent artifact is highly controversial in neuroimaging (88, 89), but it remains the field-wide gold standard (though see other recent and emerging approaches at (90, 91)).

**Behavioral Symptom and Cognitive Data.** The behavioral measures analyzed included the PANSS, an assessment of psychosis symptom severity (28), and the Brief Assessment of Cognition in Schizophrenia (BACS) battery, which provided an assessment of cognitive functioning (92). Only control subjects with complete BACS measures were used for analyses; PANSS symptom scores were imputed for control subjects for whom the PANSS had not been administered under the assumption that these subjects were asymptomatic. Only patient subjects with complete PANSS and BACS measures were used in analyses. The BACS scores used here are presented as standardized Z-scores normalized to the mean and standard deviation of the control group for each BSNIP site, as done in prior work (11). The full PANSS battery is conventionally divided into three sub-scales: Positive symptom scale (7 items), Negative symptom scale (7 items), and General Psychopathology symptom scale (16 items). The BACS battery consists of 6 individual sub-scores (92). In total, this yielded 36 symptom variables per participant. Effects of symptom variation across assessment sites have been rigorously characterized in prior publications (93). Nevertheless, we explicitly tested for site effects in our analyses, described in detail below.

**Principal Component Analysis of Behavioral Symptoms and Cross-validation.** The principal component analysis (PCA) of behavioral data was computed by including all 36 symptom variables across all N=436 patients. Variables were first scaled to have unit variance across patients before running the PCA. Significance of the derived principal components (PCs) was computed via permutation testing. For each permutation, patient order was randomly shuffled for each symptom variable before re-computing PCA. This permutation was repeated 5,000 times to establish the null model. PCs which accounted for a proportion of variance that exceeded chance (p<0.05 across all 5000 permutations) were retained for further analysis.

To evaluate if there were site differences that uniquely drove the PCA solution, we performed a leave-site-out cross-validation analysis. Specifically, we re-ran the PCA using all patients except those from one site, which was held out. This process was repeated for each of the 6 sites. To further evaluate the stability of the derived PCA solutions, we performed 1,000 runs of k-fold cross-validation for values of k between 2 and 10. For each k-fold run, the full sample of patients was randomly divided into equally-sized k sets and a PCA was re-computed using subjects in k-1 sets (as the left-out set was used as the held-out sample). For each run of leave-site-out and k -fold cross-validations significance was assessed via permutation testing as described above. The number of significant PCs and the total proportion of variance explained by these significant PCs remained highly stable across all runs (see Fig. 2A and Fig. S6 and Fig. S5). Additionally, we compared observed and predicted PCA scores for the held-out sample of patients. Predicted scores for the held-out sample of patients were computed by weighing their raw symptom scores with the loadings from the PCA computed in all other patients. Observed scores for held-out patients were obtained from the original PCA computed on the full sample presented in the main text. The similarity between predicted and observed scores was high for all five significant PCs across all runs of leave-site-out and k-fold cross-validation, exceeding r=0.9 in most analyses (see Fig. 2B-C and Fig. S6 and S5). Notably, the PCA solution did not show medication status or dosage effects (Fig. S7).

We also assessed the similarity of the PCA loadings using leave-site-out and 1,000 runs of 5-fold cross-validation frameworks (Fig. 2D). Importantly, this cross-validation was designed to test if the observed loadings remained stable (as opposed to predicted patient-level scores). The loadings for significant PCs from each leave-site-out PCA solution as well as each run of the 5-fold cross-validation were highly correlated (Fig. 2D). The leave-site-out and k-fold cross-validation PCA analyses...
by definition use overlapping samples of patients for each iteration. Therefore, we additionally conducted a full split-half replication using entirely non-overlapping sets of patients in each iteration. For each split-half iteration, the full patient sample was randomly divided into two sets with equal proportions of each of the three diagnostic groups (BPP, SADP, SZP). Then, a PCA was computed using each of the split-half patient samples. The loadings from the two PCA solutions were then evaluated for reproducibility. This process was repeated 1,000 times. The loadings for significant PCs were again highly similar even when comparing PCA solutions derived from completely non-overlapping patient samples (Fig. 2D).

To predict individual patient PC scores for the leave-one-out analysis (Fig. S20A), a PCA was computed using all patients except one held-out patient (N=435). In turn, the derived loadings were then used to compute the predicted PC scores for the left-out patient. This process was repeated until predicted PC scores were calculated for every patient. Finally, the predicted score for each patient was evaluated for reproducibility relative to the observed score obtained from the PCA solution computed using the full N=436 sample of patients. In addition, we computed an independent component analysis (ICA) to evaluate the consistency of the behavioral data reduction solution across methods (see Methods).

**Global Brain Connectivity Calculation.** Following preprocessing, the functional connectivity (FC) matrix was calculated for each participant by computing the Pearson’s correlation between every grayordinate in the brain with all other grayordinates. A Fisher’s r-to-Z transform was then applied. Global brain connectivity (GBC) was calculate by computing the mean FC strength with all other grayordinates. A Fisher’s r-to-Z transform was then applied.

Global Brain Connectivity Calculation. Following preprocessing, the functional connectivity (FC) matrix was calculated for each participant by computing the Pearson’s correlation between every grayordinate in the brain with all other grayordinates. A Fisher’s r-to-Z transform was then applied. Global brain connectivity (GBC) was calculated by computing the mean FC strength with all other grayordinates (i.e. the mean, per row, across all columns of the FC matrix). GBC is a data-driven summary measure of connectedness that is unbiased with regards to the location of a possible alteration in connectivity (94) and is therefore a principled way for reducing the number of neural features while assessing neural variation across the entire brain.

\[
GBC(x) = \frac{1}{N} \sum_{y=1}^{N} r_{xy}
\]

- where \(GBC(x)\) denotes the GBC value at grayordinate \(x\);
- where \(N\) denotes the total number of grayordinates;
- where \(\sum_{y=1}^{N} \) denotes the sum from \(y = 1\) to \(y = N\);
- where \(r_{xy}\) denotes the correlation between the time-series of grayordinates \(x\) and \(y\);

For parcel-wise GBC maps (described below) we first computed the mean BOLD signal within each parcel (see section below for parcellation details) for each participant and then computed the pairwise FC between all parcels. Finally, to obtain the parcellated GBC metric we computed the mean FC for each parcel and all other parcels. This order of operations (first parcellating the dense data series and then computing GBC) was chosen because it resulted in stronger statistical values due to increased within-parcel signal-to-noise of the BOLD data (Fig. 4A).

\[
GBC(p) = \frac{1}{N} \sum_{q=1}^{N} r_{pq}
\]

- where \(GBC(p)\) denotes the GBC value at parcel \(p\);
- where \(N\) denotes the total number of parcels;
- where \(\sum_{y=1}^{N} \) denotes the sum from \(q = 1\) to \(q = N\);
- where \(r_{pq}\) denotes the correlation between the time-series of parcels \(p\) and \(q\);

**Neural Data Reduction via Functional Brain-wide Parcellation.** Here we applied a recently developed Cole-Anticevic Brain Network Parcellation (CAB-NP) parcellation (34), which defines functional networks and regions across cortex and subcortex that leveraged the Human Connectome Project’s Multi-Modal Parcellation (MMP1.0) (34, 37). The final published CAB-NP 1.0 parcellation solution can be visualized via the Brain Analysis Library of Spatial maps and Atlases (BALSa) resource (https://balsa.wustl.edu/rrg5v) and downloaded from the public repository (https://github.com/ColeLab/ColeAnticevicNetPartition). The cortex component of the parcellation solution is comprised of 180 bilateral parcels (a total of 360 across both left and right hemispheres), consistent with the Human Connectome Project’s Multi-Modal Parcellation (MMP1.0) (37). The subcortex component is comprised of 358 parcels defined using resting-state functional BOLD covariation with the cortical network solution (34).

**Mass Univariate Behavioral-to-Neural Mapping.** Behavioral scores (a priori and PCA) were quantified in relation to individual GBC variation (either dense or parcellated) via a mass univariate regression procedure. The resulting map of regression coefficients reflected the strength of the relationship between patients’ behavioral PC score and GBC at every neural location, across all 436 patients. The greater the magnitude of the coefficient for a given location, the stronger the statistical relationship between GBC and the behavioral variation across patients. The coefficients were then Z-scored for each map. Significance of the maps was assessed via non-parametric permutation testing, 2000 random shuffles with TFCE (95) type-I error-protection computed via the Permutation Analysis of Linear Models program (96).

**Cross-validation of Mass Univariate Behavioral-to-Neural Mapping.** Behavioral-to-Neural mapping followed the same cross-validation logic as described for the symptom-driven PCA solutions. Specifically, five-fold cross-validation was performed by first randomly partitioning all patients...
shown to be highly similar ((32, 81)). First, we computed the left and right hemispheres of this parcellation have been averaged for each pair of analogous parcels in the left and right hemispheres to 180 symmetrized cortical parcels (i.e. GBC was averaged across patients across the full sample replication). Here, the number of neural features in principled manner via the de-significance of the neural PCA solution was assessed via permutation testing (1,000 random shuffles of parcels within subject). Notably, both the 358 subcortical parcels and the 12 functional networks (which span both cortex and subcortex) were obtained from the parcellation in (34).

CCA Cross-validation. Five-fold cross-validation of the CCA was performed by first randomly partitioning all patients (N=436) into 5 subsets. The CCA was then performed between neural and behavioral features using all but one of the subsets. The results of this CCA was then compared to the full sample CCA. Fig. S16A-D shows the comparisons across all 5 five-fold cross-validation runs of the CCA compared to the full model, including neural factor loadings, behavioral factor loadings, and projected behavioral item loadings. For leave-one-out cross-validation, all subjects except those from one site were used in the CCA. The resulting outputs were then compared to those from the full sample CCA. This process was repeated for all 6 sites. These data are shown in Fig. S16E-H.

For leave-one-out cross-validation of the CCA, one subject was held out as a CCA was performed using neural and behavioral features from the other 435 subjects. The loadings matrices \( \hat{N} \) and \( \hat{B} \) from the CCA were then used to calculate the “predicted” neural and behavioral latent scores for all 5 canonical modes for the holdout subject. This was repeated for every subject, such that predicted neural and behavioral latent score matrices \( \hat{N} \) and \( \hat{B} \) were computed, of the same dimensions as (N and B) respectively. The corresponding CVs in \( \hat{N} \) and \( \hat{B} \) were then compared across subjects, as shown in Fig. 5M. If the CCA solution is stable, these correlations should be comparable to the canonical correlations of the full CCA (Fig. 5D).

Independent Replication Dataset. To illustrate the generalizability of the single-subject prediction results across inde-
dependent datasets and DSM disorders, we use an independent collected dataset consisting of 30 patients with a formal diagnosis of schizophrenia (SZP) and 39 patients diagnosed with obsessive compulsive disorder (OCD). These patients were recruited via clinician referral and regional advertising and assessed at the Connecticut Mental Health Center in New Haven, CT. Demographic and symptom data are shown in Table S3.

The behavioral assessment for the replication dataset included the PANSS and the Penn Computerized Neurocognitive Battery (PennCNB)(97). Items from the PennCNB were matched to items from the BACS as follows: BACS digital sequencing (working memory) -> LNB (N-back Total score); BACS symbol coding (attention) -> Penn Continuous Performance Test; BACS token motor task (motor) -> CTAP; BACS Tower of London (abstraction and mental flexibility) -> PC or RAVEN (PENN Conditional Exclusion Test); BACS verbal fluency (verbal ability) -> SPVRT (Penn Verbal Reasoning Test); BACS verbal memory test -> PLLT (Penn List Learning Test) (98).

Neural data were collected using a Siemens 3T scanner with a 64 channel head coil at the Yale Center for Biomedical Imaging. Imaging acquisition parameters were aligned with those of the Human Connectome Project (HCP) (99).

High-resolution T1w and T2w structural images were acquired in 224 AC-PC aligned slices, 0.8mm isotropic voxels. T1w images were collected with a magnetization-prepared rapid gradient-echo (MP-RAGE) pulse sequence (TR=3200 ms, TE=564 ms, flip angle mode=T2 var, field of view=256 x 256 mm). T2w images were collected with a SCP pulse sequence (TR=800 ms, time echo (TE)=31.0 ms, flip angle=55°, field of view=210 x 210 mm, matrix=84 x 84, bandwidth=2290 Hz); 54 interleaved axial slices aligned to the anterior-posterior commissure (AC-PC) with 2.5mm isotropic voxels. Resting-state BOLD images were collected with a multi-band accelerated fast gradient-echo, echo-planar sequence (acceleration factor=6, time repetition (TR)=800 ms, time echo (TE)=31.0 ms, flip angle=55°, field of view=210 x 210 mm, matrix=84 x 84, bandwidth=2290 Hz); 54 interleaved axial slices aligned to the anterior-posterior commissure (AC-PC) with 2.5mm isotropic voxels. Additionally, a pair of reverse phase-encoded spin-echo field maps (anterior-to-posterior and posterior-to-anterior) were acquired (voxel size=2.5 mm isotropic, TR=7220 ms, TE=73 ms, flip angle=90°, field of view=210 x 210 mm, bandwidth=2290 Hz). These neural data were preprocessed using the HCP minimal preprocessing pipeline as described in the section “Neural data acquisition and preprocessing” above, with the same motion scrubbing and nuisance signal regression parameters as were used for the BSNIP dataset.

**Neural Parcel ΔGBC Feature Selection and Prediction.**

For the purposes of patient selection, we were focused on individual differences in neural features as they co-vary in relation to individual symptom scores. However, this individual variation in neural features may be small compared to the overall group mean. Hence for each patient, we compute the difference in each brain location (parcel) between the patient’s actual GBC and the group mean GBC for that location. This is denoted by ΔGBC. Importantly, using a de-meaned GBC metric “standardizes” the data and helps to correct for possible differences in scanners/protocols across different datasets. Next, we developed an optimized univariate regression framework leveraging a “dot product” metric to relate a vector of neural features with a single symptom scalar value. This process for neural feature selection (results in Fig. S20) is shown as a systems flow diagram in Fig. S18. The observed dot product GBC metric (dpGBC<sub>obs</sub>) is computed as follows:

\[
\text{dpGBC}_{i}^{\text{obs}} = \left( \begin{array}{c} \Delta GBC_{i}^{\text{obs}} \\ \vdots \\ \Delta GBC_{i}^{\text{obs}} \end{array} \right) \cdot \left( \begin{array}{c} \beta GBC_{i}^{\text{obs}} \\ \vdots \\ \beta GBC_{i}^{\text{obs}} \end{array} \right)
\]

where for a given subject \( i \) dpGBC<sub>i</sub><sup>obs</sup> denotes the dot product GBC value of the two vectors \( \Delta GBC_{i}^{obs} \) and \( \beta GBC_{i}^{obs} \) across all \( P_{select} \) parcels,

\[
\Delta GBC_{i}^{obs} = \text{vector of length } P_{select} \text{ denoting a difference map of subject } i \text{'s } GBC^{obs} \text{ map relative to the group mean } GBC^{obs} \text{ map, within a given number of parcels } P_{select}.
\]

the \( \beta GBC_{i}^{obs} \) vector denotes the PCA-to-GBC statistical group-level \( \beta \) map for a low-dimensional PC symptom score across selected parcels \( P_{select} \) for \( N \) subjects excluding subject \( i \).

This calculation is then repeated for each subject \( i \), resulting in a final vector \( \text{dpGBC}_{i}^{obs} = [\text{dpGBC}_{1}^{obs}, \ldots, \text{dpGBC}_{N}^{obs}] \) for \( N \) subjects. There are several key properties of the dpGBC<sub>obs</sub> statistic: i) it is not inflated by individual GBC map similarity to the group map because each subject’s \( \Delta GBC^{obs} \) map is demeaned relative to the low-dimensional symptom score across all subjects (see Eq. 2). Put differently, the final evaluation considers the relationship between \( \text{dpGBC}_{i}^{obs} \) and the low-dimensional PC symptom scores across individuals; ii) the dot product statistic can yield both positive and negative values – a property which some map similarity measures lack (e.g. \( R^2 \)); iv) It is unbounded (unlike a correlation), which is key to maximize co-variation with low-dimensional symptom scores across subjects (see Eq. 2); v) The \( \Delta GBC_{i}^{obs} \) map for a given individual is projected onto the basis set of the \( \beta GBC_{N-i}^{obs} \) map, which is independent of the left-out individual but directly related to the low-dimensional PC symptom score variance, thus maximizing the dot product optimization.

Next, we select \( N - i \) individuals and compute a univariate regression where participants’ low dimensional symptom scores are regressed onto the \( \text{dpGBC}_{N-i}^{obs} \) values:

\[
\left( \begin{array}{c} \text{dpGBC}_{i}^{obs} \\ \vdots \\ \text{dpGBC}_{N-i}^{obs} \end{array} \right) = \alpha \left( \begin{array}{c} z_{i}^{obs} \\ \vdots \\ z_{N-i}^{obs} \end{array} \right) + \epsilon
\]
• where for $dpGBC_{obs}^{\text{pred}}$ each element denotes the dot product value of the $\Delta GBC_{i,obs}^\alpha \cdot \beta GBC_{N-1}^{obs}$ vectors per subject across $P$ parcels,
• $\alpha$ denotes the regression coefficient in the univariate linear model,
• where for $S_{obs}$ each element denotes the observed low-dimensional symptom score (e.g. PC3 score) per subject,
• $\epsilon$ denotes the error term in the univariate linear model.

After the $dpGBC_{obs}^{\text{pred}} = \alpha S_{obs}^{\text{pred}} + \epsilon$ regression is computed on $N$ subjects, it is applied to the left-out-subject $i$. This is repeated for all $N$ subjects and the model is evaluated for the number of parcels that maximize two key dot product evaluation metrics (Fig. 6B):

**Metric A**

$$\text{For } p = \{1, \ldots, P\} \text{ find } p \text{ where }$$

$$A = \max_r \left[ \begin{array}{c} (dpGBC_1)_r^{\text{pred}} \\ \vdots \\ (dpGBC_N)_r^{\text{pred}} \end{array} \right], \left[ \begin{array}{c} S_1^{\text{pred}} \\ \vdots \\ S_N^{\text{pred}} \end{array} \right]$$

**Metric B**

$$\text{For } p = \{1, \ldots, P\} \text{ find } p \text{ where }$$

$$B = \max_r \left[ \begin{array}{c} (dpGBC_1)_r^{obs} \\ \vdots \\ (dpGBC_N)_r^{obs} \end{array} \right], \left[ \begin{array}{c} (dpGBC_1)_r^{\text{pred}} \\ \vdots \\ (dpGBC_N)_r^{\text{pred}} \end{array} \right]$$

• where $A$ denotes the maximum $r$ correlation value for the two vectors of $dpGBC^{\text{pred}}$ values and the predicted $S^{\text{pred}}$ low-dimensional symptom scores (e.g. PC3 axis) for $N$ subjects from the leave-one-out cross-validation,
• where $B$ denotes the maximum $r$ correlation value for the two vectors of $obs$ and $\text{pred} dpGBC$ values for $N$ subjects.

In the initial step in the step-down model all $P = 718$ parcels are retained in the initial dot product calculation. For each iteration of $P$ selected parcels, the least predictable parcel $P$ (i.e. the parcel with the weakest value in the PC3 map) is eliminated from the map. Then, the step-down regression is repeated until $P = 1$.

**Pharmaceutical Neuroimaging Acquisition in Healthy Volunteers - LSD.** Methods for the lysergic acid diethylamide (LSD) neuroimaging study are described in detail in prior publications (29). The use of LSD in humans was authorized by the Swiss Federal Office of Public Health, Bern, Switzerland. The study protocol was approved by the Cantonal Ethics Committee of Zurich ($KEK - ZH\_NO : 2014\_0496$). Participants received both written and oral descriptions of the study procedures and the effects and possible risks of LSD. All participants provided written informed consent in accordance with the Declaration of Helsinki. The study employed a fully double-blind, randomized, within-subject cross-over design with 3 conditions: (1) placebo + placebo (Pla) condition: placebo (179 mg Mannitol and Aerosil 1 mg po) after pretreatment with placebo (179 mg Mannitol and Aerosil 1 mg po); (2) Pla+LSD (LSD) condition: LSD (100 µg po) after pretreatment with placebo (179 mg Mannitol and Aerosil 1 mg po), or (3) Ketanserin+LSD (Ket+LSD) condition: LSD (100 µg po) after pretreatment with the 5-HT2A antagonist Ket (40 mg po). Data were collected for all subjects in a randomized counterbalanced order at three different sessions each two weeks apart. For all conditions, the first substance was administered 60 minutes before the second substance, and the first neural scan was conducted 75 minutes after the second administration, with a second scan conducted at 300 minutes post-administration. In the present study, only data from the two neural scans for the LSD and Pla conditions were evaluated.

Briefly, neuroimaging data acquisition details for the LSD study are as follows. MRI data were acquired on a Philips Achieva 3.0T whole-body scanner (Best, The Netherlands). A 32-channel receiver head coil and MultiTransmit parallel radio frequency transmission was used. Images were acquired using a whole-brain gradient-echo planar imaging (EPI) sequence (repetition time=2,500 ms; echo time=27 ms; slice thickness=3 mm; 45 axial slices; no slice gap; field of view=240 mm; in-plane resolution=3 mm × 3 mm; sensitivity-encoding reduction factor=2.0). 240 volumes were acquired per resting state scan resulting in a scan duration of 10 mins. Additionally, two high-resolution anatomical images were acquired using T1-weighted and T2-weighted sequences. T1-weighted images were collected via a 3D magnetization-prepared rapid gradient-echo sequence (MP-RAGE) with the following parameters: voxel size=0.7 mm$^3$, time between two inversion pulses=3123 ms, inversion time=1055 ms, inter-echo delay=12 ms, flip angle=8°, matrix=320×335, field of view=224×235 mm, 236 sagittal slices. Furthermore T2-weighted images were collected using via a turbo spin-echo sequence with the following parameters: voxel size=0.7 mm$^3$, repetition time=2500 ms, echo time=415 ms, flip angle=90°, matrix=320×335, field of view=224 mm × 235 mm, 236 sagittal slices.

**Pharmaceutical Neuroimaging Acquisition in Healthy Volunteers - Ketamine.** Similar to the LSD study, the ketamine pharmacological neuroimaging protocol employed a within-subject design where all healthy volunteer participants underwent a single scanning session consisting of two infusions: i) placebo (saline solution) followed by ii) ketamine infusion. The study was approved by the Yale University institutional review board. Participants received both written and oral descriptions of the study procedures and the effects and possible risks of ketamine, and all participants provided written informed consent before beginning the study. Healthy volunteers were informed prior to scanning that they would undergo one placebo run and one ketamine run but
were blinded to the order of administration. Because of the sustained effects of ketamine, this infusion was always the second of the two runs, consistent with prior work (86). A doctor and two nurses as well as a study coordinator remained present for the duration of the scan. A bolus of ketamine (0.3 mg/kg of bodyweight) or saline were delivered via infusion 5 sec after the start of the run and then continuously at a rate of 0.65 mg/kg per hour through the duration of the session. The sequence of scans in each for either the placebo or ketamine infusion was as follows: i) resting state (4.67 min); ii) blood draw (sham if saline condition); iii) a cognitive working memory task (total 14 min); iv) blood draw (sham if saline condition); v) a cognitive working memory task (total 14 min); vi) blood draw (sham if saline condition); vii) a cognitive working memory task (total 8.63 min). Data from the the cognitive working memory task were not used in the present study and are actively undergoing a distinct analysis. Participants were scanned at Yale University on a Siemens Trio 3T whole-body scanner and 32-channel receiver head coil. High-resolution structural T1-weighted images were acquired using an MP-RAGE sequence and the following parameters: voxel size=0.8 mm$^3$, time between two inversion pulses=3123 ms, inversion time=1055 ms, inter-echo delay=12 ms, flip angle=$8^\circ$, matrix=320 $\times$ 335, field of view=227 $\times$ 272 mm, 227 sagittal slices. T2-weighted images were acquired with the following parameters: voxel size=0.8 mm$^3$, repetition time=2500 ms, echo time=415 ms, flip angle=$90^\circ$, matrix=320 $\times$ 335, field of view=227 $\times$ 272 mm, 227 sagittal slices. BOLD images were acquired using a whole-brain gradient-echo planar imaging (EPI) sequence (400 frames at TR=0.7 ms; TE=30 ms; slice thickness=2.5 mm; 54 axial slices; no slice gap; field of view=250 mm$^2$; in-plane resolution=2.5 mm $\times$ 2.5 mm). In addition, a field map and pair of spin-echo gradients were collected at the end of every scanning session.

Pharmacological Neuroimaging Processing and Analysis for LSD and Ketamine Samples. All data were preprocessed to be consistent with the BSNIP and replication dataset processing steps. Specifically, we used the HCP minimal preprocessing pipeline and with the same motion scrubbing and nuisance signal regression parameters as were used for the BSNIP dataset as described in the section “Neural data acquisition and preprocessing” above. Parcel-wise GBC maps were computed as described in the section “Resting-state functional connectivity and global brain connectivity” above, by first parcellating the dense time-series for each subject and then computing the parcel-level GBC. As with the BSNIP and replication analyses, parcel GBC was first calculated by parcellating the dense time-series for each subject and then computing GBC across all 718 parcels from the whole-brain functional parcellation (34, 37). Group-level GBC maps were computed for every participant for all conditions. We computed the contrast map for “LSD-Placebo” as well as “Ketamine-Placebo” conditions as a t-test between both the pharmacological scans versus placebo scans. The Z-scored t-contrast maps between pharmacological and placebo conditions were used as a pharmacological target map in relation to the PC-derived neural GBC maps Fig. 7 and Fig. S23.

Neural Gene Expression Mapping. The gene mapping analyses in this study utilize the procedure described in (32). Briefly, we used cortical gene expression data from the publicly available Allen Human Brain Atlas (AHBA, RRID:SCR_007416), mapped to cortex (32). Specifically, the AHBA quantified expression levels across 20,737 genes obtained from six postmortem human brains using DNA microarray probes sampled from hundreds of neuroanatomical loci. Recent studies demonstrated the ability to map expression of each gene onto neuroimaging-compatible templates (31, 32). Building on these innovations we mapped gene expression on to 180 symmetrized cortical parcels from the HCP atlas (37) in line with recently published methods (32). This yielded a group-level map for each gene where the value in each parcel reflected the average expression level of that gene in the AHBA dataset. These group-level maps were in turn as a gene expression target map in relation to the PC-derived neural GBC maps.

Data Availability. All primary results derive from data that is publicly available from sources described above. All symptom and neuroimaging data for the BSNIP-1 consortium was obtained via the NDA (https://nda.nih.gov/edit_collection.html?id=2274).

Code Availability. All code used in this paper are available from the corresponding author upon reasonable request.