Transcriptional and imaging-genetic association of cortical interneurons, brain function, and schizophrenia risk

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Inhibitory interneurons orchestrate information flow across the cortex and are implicated in psychiatric illness. Although interneuron classes have unique functional properties and spatial distributions, the influence of interneuron subtypes on brain function, cortical specialization, and illness risk remains elusive. Here, we demonstrate stereotyped negative correlation of somatostatin and parvalbumin transcripts within human and non-human primates. Cortical distributions of somatostatin and parvalbumin cell gene markers are strongly coupled to regional differences in functional MRI variability. In the general population (n = 9,713), parvalbumin-linked genes account for an enriched proportion of heritable variance in in-vivo functional MRI signal amplitude. Single-marker and polygenic cell deconvolution establish that this relationship is spatially dependent, following the topography of parvalbumin expression in post-mortem brain tissue. Finally, schizophrenia genetic risk is enriched among interneuron-linked genes and predicts cortical signal amplitude in parvalbumin-biased regions. These data indicate that the molecular-genetic basis of brain function is shaped by interneuron-related transcripts and may capture individual differences in schizophrenia risk.
Ramon y Cajal theorized that the functional diversity of the human brain arises, in part, from the vast assortment of neurons that pattern cortex\(^2\). Inhibitory interneurons are the most varied neuronal class\(^3\), exhibiting divergent morphological and physiological properties while coordinating information flow across the brain’s collective set of connections (functional connectome)\(^4\). Converging animal and human work provides evidence for the role of interneurons in healthy brain function as well as their dysregulation in psychiatric illnesses, including schizophrenia\(^5\) and major depressive disorder\(^6\). The development of dense spatial transcriptional atlases now enables the study of cellular and molecular associates of functional brain networks. Early work in this area identifies genes encoding ion channels\(^7\) and those enriched in supragranular layers of cortex\(^8\) as correlates of large-scale network organization. Other research indicates that cortical resting-state signal fluctuations follow the expression of neuron-enriched genes\(^8\). However, the transcriptional correlates of brain function have rarely been validated in vivo\(^9\) and little is known about how the spatial distribution of specific interneuron subtypes shapes cortical function and associated risk for psychiatric illness in humans.

The spatial distribution of interneuron subtypes is theorized to contribute to regional specialization of cortex, partly by altering the relative balance of excitation and inhibition for a given cortical area\(^2\)–\(^11\). Interneurons comprise 20–30% of cortical neurons\(^12\) and form stereotyped connections with excitatory projection neurons\(^2,3\). The majority of interneurons express one of a limited set of genetic markers: somatostatin (SST), parvalbumin (PVALB), and vasoactive-intestinal peptide (VIP; a subset of HTR3A interneurons)\(^2\). Each subtype possesses unique synaptic and functional characteristics, leading to the hypothesis that the ratio of interneuron classes underpins local differences in neural activity\(^11,13\). For example, SST interneurons preferentially target dendrites of cortical projection neurons to regulate their input, whereas PVALB interneurons synapse on perisomatic regions to regulate output\(^2\). Consequently, increased relative density of SST interneurons may facilitate filtering of noisy or task-irrelevant cortical signals and promote recurrent excitation required for higher-order cognition\(^11,14\). Conversely, relative increases in PVALB may produce stronger feedback inhibition on excitatory neurons, leading to shorter activation timescales suited for processing constantly changing sensorimotor stimuli\(^11,15\).

Previous work documents cortical gradients of gene transcription that mirror the hierarchical organization of timescales from fast (unimodal cortex) to slow (multimodal cortex)\(^8,10,16\). Further, SST and PVALB interneuron markers are differentially expressed within distributed limbic and somato-motor cortico-striatal networks, respectively\(^9\). These observations suggest that spatial distributions of interneuron subtypes could underlie regional signaling differences across the cortical sheet, as indexed by blood oxygenation level-dependent (BOLD) functional magnetic resonance imaging (fMRI). However, the psychiatric and functional consequences of spatially variable SST- and PVALB-related transcription in cortex have yet to be fully characterized.

Establishing the principles by which cellular diversity influences brain function is a long-standing challenge in neuroscience and could reveal biological mechanisms of individual variability of the human brain. Consistent with this aim, cross-species evidence indicates the importance of PVALB interneurons for fMRI measures of brain function\(^16\). PVALB interneurons orchestrate gamma-band oscillations (30–80 Hz\(^19\)), a frequency range that is tightly coupled to spontaneous BOLD fluctuations\(^20\). Experimental optogenetic stimulation of PVALB interneurons in rodents drives gamma-band rhythms, impacting information processing through the synchronization of excitatory neurons\(^19\). In psychiatric illness, decreased PVALB-mediated inhibition may be a core locus of disruption in schizophrenia, leading to altered gamma-band signals and working memory deficits that are a hallmark of the disorder\(^21\). However, a direct link between PVALB-related genetic variation and human brain activity has yet to be established. Linking cortical interneurons to individual differences in human brain function would yield deep biological insight into the hemodynamic BOLD signal, providing an engine for the discovery of functional connectome-linked genes and associated risk for illness onset.

Here, we bridge genetic, transcriptional, and neuroimaging data to advance three lines of inquiry linking interneurons to human brain function. First, we describe the organization of SST and PVALB expression in human and non-human primates, demonstrating a robust pattern of anti-correlation across cortex and subcortex. We perform single-cell polygenic deconvolution\(^22\) of bulk cortical tissue data to infer spatial distributions of inhibitory, excitatory, and non-neuronal cells across cortex, providing converging evidence with single-marker analyses. Second, we establish that the relative density of SST and PVALB tracks regional differences in cortical brain activity. In a large sample (N = 9713)\(^23\), genetic variation among PVALB-correlated genes explained an enriched proportion of heritable variance in resting-state signal amplitude (RSFA), in a manner that mirrors the spatial expression of PVALB measured in independent post-mortem brain tissue. These discoveries suggest that the molecular-genetic basis of cortical function is spatially nonuniform and that genes linked to PVALB interneurons explain heritable aspects of the BOLD signal. Third, we link PVALB interneurons and psychiatric illness, demonstrating that genetic risk for schizophrenia is enriched among interneuron-linked genes and predicts reduced resting-state signal amplitude in a spatially heterogeneous manner that follows PVALB expression. These data help address a deep-rooted challenge in neuroscience to understand how cytoarchitecture shapes human brain function and related vulnerability for psychiatric illness.

### Results

**Anti-correlation of SST and PVALB interneuron markers across cortex.** The properties of interneuron subtypes emerge early in development and are partly determined by their spatial origin in the embryonic ganglionic eminence\(^24\). SST and PVALB interneurons originate in the medial ganglionic eminence (MGE) along negatively correlated spatial gradients\(^25\). That is, PVALB-and SST-directed neurons differentially cluster within the dorsal and ventral MGE, respectively\(^26\). Evidence in humans\(^9,10\) and rodents\(^11,17\) suggests that SST and PVALB transcripts maintain a negative spatial correlation in adulthood, indicating that embryonic organization may constitute a “proto-map” of mature cortex. Although prior research suggests that SST and PVALB markers are differentially expressed across cortex\(^9,10\), the current work directly establishes transcriptional anti-correlation between these two cell types across multiple techniques, human datasets, non-human primate data, and human neurodevelopment. This comprehensive profiling of SST and PVALB interneuron expression is necessary for deep profiling of their relationship to in-vivo brain function and subsequent schizophrenia risk through statistical genetic approaches. The functional consequences of a negative spatial SST to PVALB relationship are not well understood, but the presence of replicable and evolutionarily conserved expression patterns may indicate the importance of interneuron gradients.

To characterize interneuron marker topography across human and non-human primate cortex, we analyzed gene expression data from the Allen Human Brain Atlas (AHHBA)\(^27\) and NIH Blueprint Non-Human Primate (NHP) Atlas\(^28\). AHHBA cortical...
samples from the left ($n = 1265$) and right ($n = 418$) hemispheres were analyzed. Microarrays do not provide absolute estimates of gene transcription, but can measure within-probe differences across samples. SST and PVALB expression were thus z-transformed across cortical samples, and subtracted (i.e., SST–PVALB) to reveal relative expression differences (Fig. 1a).

Extending prior evidence9,10, SST and PVALB were anti-correlated across AHBA cortical samples (Fig. 1e) for both parametric ($r(1681) = −0.45, p < 2.2e−16$) and non-parametric tests (Spearman’s rho ($r_s = −0.40, p < 2.2e−16$). Relative to all possible gene-wise correlations, the SST to PVALB correlation was among the most negative (Fig. 1h, i; AUC$_{sst-pvalb} = 0.009$, AUC$_{pvalb-sst} = 0.033$). Further, the SST and PVALB relationship was among the most negative relative to all possible ($n = 152,207,628$) two-gene pairings (Fig. 1j; AUC = 0.001). SST and PVALB distributions were organized along an anterior to posterior gradient, with greater relative SST in orbitofrontal and medial prefrontal cortex, anterior insula, and the temporal lobe (Fig. 1a, c and Supplementary Fig. 1). By contrast, relative SST and PVALB expression was greatest within unimodal somato-motor, parietal and visual cortices (Fig. 1a, c and Supplementary Fig. 1).

Histologically defined anatomical categories were used to characterize regional differences of interneuron marker expression, reflected in Fig. 1c showing negatively correlated SST and PVALB median across cortical subregions ($r(39) = −0.88, p = 2.4e−14$). The negative spatial relationship between SST and PVALB was evolutionarily conserved in non-human macaque primates (Fig. 1f; $r(34) = −0.74, p = 2.2e−7$; $r_s = −0.60, p = 0.0001$), suggesting that interneuron marker gradients may reflect a core organizational feature of primate cortex. These data complement evidence for similarly negatively correlated SST and PVALB gradients in rodents$^{11,17}$. Given that SST and PVALB interneurons originate along stereotyped, anti-correlated gradients in the MGE$^{26}$, we analyzed RNAseq data from the Brainspan Atlas of the Developing Human Brain to test whether the emergence of SST to PVALB negative correlations coincides with major waves of interneuron colonization, approximately 10–25 post-conception weeks (pcw)$^{29,30}$. The negative correlation between SST and PVALB was absent in early-fetal (8–12 pcw; $\beta = 0.32, p = 0.039$) and early-midfetal (13–21 pcw; $\beta = 0.02, p = 0.85$) stages. Consistent with the hypothesis that mature interneuron distributions result from developmentally programmed migration patterns, we observed significant negative correlations between SST and PVALB emerge during late-fetal (24–37 pcw; $\beta = −0.47, p = 0.012$), early-infancy (4 months; $\beta = −0.60, p = 0.0033$), mid-late childhood (8–11 years; $\beta = −0.52, p = 0.0038$), and adult (18–40 years; $\beta = −0.35, p = 0.0014$) periods, as well as at a trend-level in adolescence (13–15 years; $\beta = −0.54, p = 0.057$). We did not observe a relationship in late infancy (10 months; $b = −0.37, p = 0.36$) or early childhood (1–4 years; $b = −0.11, p = 0.48$). These data provide developmental context as well as an external replication of the SST–PVALB cortical expression pattern observed in the AHBA (adult human) and NHP Atlas (adult macaque) samples.

Polysymmetric deconvolution of cell types across cortex. SST and PVALB are reliable genetic markers of their respective interneuron subtypes$^2$, however single-cell transcriptomics show that cell classes possess polysymmetric signatures of expression$^{31}$. To capture the molecular complexity of cell identity and quantify cellular-level associates of brain function, we conducted polysymmetric cellular deconvolution of bulk AHBA expression data using CIBERSORTx (https://cibersortx.stanford.edu)$^{32}$. This method leverages transcriptomic ‘signatures’ of cellular identity to estimate the relative abundance of cell types in bulk tissue data. Single-nucleus droplet-based sequencing (snDrop-seq) data from Lake and colleagues$^{33}$ was used, providing cell-level expression data in dorsal frontal cortex (BA6/BA10; 10,319 cells) and visual cortex (BA17; 19,368 cells). Collinearity among transcriptionally similar cell types was reduced by using 18 superordinate cell identities defined by Lake and colleagues$^{31}$. Cell signatures were created with CIBERSORTx (Supplementary Fig. 5) and cell abundance, expressed as a fraction, was estimated for each cortical tissue sample from AHBA donors (see Supplementary Data). AHBA tissue samples were projected to the cortical surface, allowing for vertex-level localization on a group-atlas (fsLR32k; see “Methods”). Fractional cell abundances were averaged for each of the 400 bi-hemispheric parcels from the atlas of Schaefer and colleagues$^{32}$ (Supplementary Figs. 6–8).

We focus on imputed spatial distributions of SST and PVALB interneurons across cortex, estimated separately from visual and frontal cortex single-cell data (Fig. 2). Similar to single-marker analyses (Fig. 1a and Supplementary Fig. 1), the inferred fractional abundance of SST interneurons was greatest in medial PFC, anterior insula, and temporal poles (Fig. 2a), which was highly consistent between frontal and visual cortex cell type derived maps ($r(337) = 0.73, p < 2.2e−16$). Conversely, PVALB fractional abundance was highest within visual, motor, and dorsal parietal cortex (Fig. 2b), which was also consistent across frontal and visual cortex derived estimates ($r(337) = 0.76, p < 2.2e−16$). Given the hypothesized importance of relative SST and PVALB abundance for cortical functional dynamics, Fig. 2c shows the spatial distribution of estimated SST- (red) and PVALB-dense (blue) areas of cortex. Figure 2d illustrates the spatial correlation of all deconvolved cell type fractions across cortex, recapitulating the negative spatial relationship between SST and PVALB single-gene markers (frontal cortex: $r(337) = −0.62, p < 2.2e−16$; visual cortex: $r(337) = −0.37, p = 1.15e−12$). Critically, cell fraction maps of PVALB and SST obtained through polygenic deconvolution were strongly correlated with the expression of their respective single-marker genes (Fig. 2f, g).

SST and PVALB anti-correlation in subcortical territories. We next examined whether the negative spatial relationship between SST and PVALB is also preserved across subcortex (see Supplementary Information for subcortical sample information). Although there is evidence for anti-correlated SST and PVALB interneuron gradients in striatum$^9$ and hippocampus$^{33}$, it remains unclear whether SST and PVALB inverse gradients are ubiquitous throughout the brain. Sample-wise expression was normalized and correlated separately for each of the seven areas: striatum, dorsal thalamus, hypothalamus, globus pallidus, amygdala, hippocampus, and combined substantia nigra/ventral tegmentum. The strength of the SST to PVALB correlation was benchmarked against all gene-wise correlations to SST and PVALB (Fig. 3a), and quantified as AUC of the distribution less than or equal to the SST–PVALB correlation. SST was significantly negatively correlated to PVALB in the hypothalamus ($r(100) = −0.72, q = 1.9e−16$, AUC$_{sst} = 0.002$, AUC$_{pvalb} = 0.01$), globus pallidus ($r(37) = −0.39, q = 0.011$, AUC$_{sst} = 0.03$, AUC$_{pvalb} = 0.03$), amygdala ($r(65) = −0.30, q = 0.019$, AUC$_{sst} = 0.06$, AUC$_{pvalb} = 0.03$), and thalamus ($r(173) = −0.19, q = 0.019$, AUC$_{sst} = 0.04$, AUC$_{pvalb} = 0.07$), but not the hippocampus ($r(156) = −0.12, q = 0.14$, AUC$_{sst} = 0.06$, AUC$_{pvalb} = 0.21$), ventral tegmentum/substantia nigra ($r(63) = 0.09, q = 0.50$, AUC$_{sst} = 0.39$, AUC$_{pvalb} = 0.32$), and striatum ($r(168) = 0.23, q = 0.011$, AUC$_{sst} = 0.65$, AUC$_{pvalb} = 0.72$).

These data identified subcortical territories with biased SST or PVALB expression in humans, which show striking
Fig. 1 Cortical expression of SST and PVALB are negatively correlated across species and developmental stages. a AHBA tissue samples mapped to the human cortical surface, and b an illustration of non-human primate tissue sample locations, colored by relative expression of SST (red) and PVALB (blue). Normalized expression difference reflects the sample-wise subtraction of z-transformed PVALB from SST. Relative SST–PVALB expression among anatomically defined groups from the c AHBA (n = 6 donors; n = 1683 samples; n = 41 regions) and d NIH Blueprint Non-Human Primate Atlas (n = 4 donors; n = 182 samples; n = 10 regions); circles = median, thick lines = interquartile range, thin lines = min and max values. e Sample-wise negative correlation of SST and PVALB in human cortex (r = −0.45, p < 2.2e−16; r₁ = −0.40, p < 2.2e−16) and f non-human primates (r(34) = −0.74, p = 2.2e−7; r₂ = −0.60, p = 0.0001). g Correlation of cortical SST and PVALB across nine developmental stages using data from the Brainspan Atlas of the Developing Human Brain (n = 42 donors; n = 362 samples; n = 9 developmental stages). h–j The SST to PVALB correlation is at the left tail of the distribution of all gene-wise correlations to SST (AUC = 0.009) and to PVALB (AUC = 0.033), as well as all possible two-gene spatial correlations (AUC = 0.001). *p ≤ 0.05, uncorrected; †p ≤ 0.10; error bars = standard error.
correspondence to non-human animal literature. Median \( z \)-transformed SST and PVALB was calculated for subdivisions of each subcortical region (Supplementary Fig. 9). Across 40 AHBA subcortical sub-regions, 31 were able to be mapped onto rodent homologs present in cell density (cells per \( \text{mm}^3 \)) data from Kim and colleagues. The relative presence of human SST and PVALB across sub-regions was significantly positively correlated to ground truth cell density estimates in rodents (Fig. 3b; \( r_s = 0.41, p = 0.025 \)). Results were consistent when Spearman-rank correlation was used to limit the influence of outliers, and when the ratio of SST/PVALB rodent cell densities was used rather than the difference (\( r_s = 0.49, p = 0.006 \)).
Fig. 2 Deconvolved cell type distributions are consistent with SST and PVALB single marker expression maps. Using CibersortX22, frontal and visual cortex snDrop-Seq data from Lake and colleagues31 were used to deconvolve cell type fractions from bulk AHBA microarray expression data. Deconvolved cell fractions of a somatostatin and b parvalbumin interneurons across cortex using single-cell data from frontal (left) and visual (right) cortex.

c Somatostatin and parvalbumin cell fraction maps were z-transformed and subtracted (SST-PVALB) to illustrate the relative density of each subtype across cortex. d Spatial correlations of each deconvolved cell type across cortex using frontal (top-left triangle) and visual (bottom-right triangle) cell signatures. e Example marker genes for each cell class. f Deconvolved parvalbumin fractions across cortex were positively spatially correlated (Pearson’s) to single-gene PVALB expression (frontal cortex: r(337) = 0.81, p < 2.2e−16; visual cortex: r(337) = 0.48, p < 2.2e−16). g Deconvolved somatostatin fractions across cortex are positively spatially correlated (Pearson’s) to single-gene SST expression (frontal cortex: r(337) = 0.72, p < 2.2e−16; visual cortex: r(337) = 0.60; p < 2.2e−16).

Fig. 3 Subcortical SST and PVALB correlation and consistency in rodents. a The correlation between SST and PVALB was estimated using AHBA data for each of seven subcortical territories. Red/blue circles denote Pearson correlations between SST and PVALB. Light and dark gray boxplots show the distribution of spatial correlations of all other genes (n = 17,447) to PVALB and SST, respectively (center = median, box = Q1–Q3, whiskers = 1.5*IQR, circles = outliers). The strength of the SST to PVALB relationship is quantified relative to gene-wide reference distributions. SST was more negatively correlated to PVALB than what is expected by chance in the hypothalamus (r(100) = −0.72, q = 1.9e−16, AUCst = 0.002, AUCpvalb = 0.01), globus pallidus (r(37) = −0.39, q = 0.011), amygdala (r(65) = −0.30, q = 0.19), and thalamus (r(173) = −0.19, q = 0.019), but not the hippocampus (r(156) = −0.12, q = 0.14), ventral tegmentum/substantia nigra (r(63) = 0.09, q = 0.50), and striatum (r(168) = 0.23, q = 0.011). b Relative expression of SST and PVALB in human subcortical areas when compared to ground truth cell densities in rodent homologs from Kim et al.31 (Spearman’s rd = 0.41, p = 0.025).

In particular, the central nucleus (CeA) of the amygdala was enriched for SST in both rodents (SSTdensity = 24,342.60, PVALBdensity = 119.42) and humans (SSTexpr(z) = 1.27, PVALBexpr(z) = −0.76), paralleling observations in primates34. We also found greater relative SST in the ventral tegmental area (VTA) for both rodents (SSTdensity = 1953.98, PVALBdensity = 426.10) and humans (SSTexpr(z) = 1.12, PVALBexpr(z) = −0.42), but greater relative presence of PVALB in the substantia nigra pars reticulata (SNr) in humans (PVALBexpr(z) = 0.22, PVALBexpr(z) = 0.72) and rodents (SSTdensity = 680.27, PVALBdensity = 4373.48). Supporting this VTA/SNr distinction, the VTA is connected to other SST-biased regions, including the nucleus accumbens (NAc), anterior cingulate cortex, and mediodorsal thalamus35, whereas functional neuroimaging indicates a preferential coupling of the SNr to motor areas and sensorimotor striatum36. Building on the previous work9, the extent to which SST may be relatively increased within a distributed limbic cortico-striato-thalamic network is explored in Supplementary Fig. 10.

Relative SST and PVALB covaries with resting-state signal amplitude. Computational rodent work suggests the relative presence of SST and PVALB interneurons is a determinant of functional differences and hierarchical organization across cortex11. Spiking activity in cortex progresses from shorter timescales in sensory and unimodal cortices to longer timescales in association and integrative cortices15,37. This functional organization may be indexed by variability in the resting-state BOLD signal. Accordingly, we examined whether the difference of cortical SST and PVALB expression covaries with an in-vivo measurement of cortical signal variability, resting-state functional amplitude (RSFA). Voxel-wise RSFA was calculated using the UK Biobank sample (n = 9713) and averaged across the 400 parcel functional atlas of Schaefer and colleagues (Fig. 4a)32. Between-subject hierarchical clustering was conducted to reduce data dimensionality and identify cortical territories with similar patterns of signal amplitude across individuals (Fig. 4b). A seven-cluster solution was selected, corresponding to limbic A (light beige), limbic B (dark beige), cingulo-opercular (teal), temporoparietal (orange), prefrontal (red), somato/motor (blue), and visual (purple) clusters. With the exception of Figs. 5a, b and 6b, clusters serve to aid in visualization and do not influence statistics. Consistent with recent work38, this data-driven dimensionality reduction broadly sorted association and unimodal aspects of cortex.
Computational models suggest that increased PVALB density may correspond to greater recurrent inhibition of excitatory neurons and faster timescales (i.e., higher RSFA). Likewise, increased relative presence of SST is predicted to correspond to longer timescales of activity (i.e., lower RSFA). Earlier work documents a correlation of interneuron marker expression with fractional Amplitude of Low-Frequency Fluctuations (fALFF), a metric closely tied to RSFA, within a circumscribed set of cortical areas. Consistent with these predictions, a negative spatial correlation was observed between RSFA and the relative difference of SST and PVALB expression (Fig. 4c; \( r(337) = -0.53, \ p < 2.2e^{-16}; \ r_s = -0.60, \ p < 2.2e^{-16} \)). Across 152,207,628 two-gene pairings, the SST–PVALB to RSFA correlation \( (r = -0.53) \) was the 54,303rd most negatively associated pair (top 0.04%). Analysis of deconvolved cell fractions across cortex revealed that SST had the strongest negative spatial association to RSFA (Fig. 4d; frontal cortex signature: \( r(337) = -0.48, \ p < 2.2e^{-16}; \) visual cortex signature: \( r(337) = -0.36, \ p = 1.2e{-11} \)), while PVALB cell fractions were the most positively correlated (frontal cortex signature: \( r(337) = 0.47, \ p < 2.2e^{-16}; \) visual cortex signature: \( r(337) = 0.34, \ p = 1.7e{-10} \)). Across frontal cortex \( (n = 272) \) and visual cortex \( (n = 240) \) cell type pairs, the relative difference of SST and PVALB cell fractions is most spatially associated with cortical RSFA.
Polygenic variation among PVALB-correlated genes underlies cortical brain function. Genome-wide association studies (GWAS) demonstrate that the genetic bases of many complex traits are due to the cumulative weight of genetic variants spread across the entire genome, each with a subtle effect. Brain phenotypes such as resting-state functional amplitude likely possess a similar polygenic architecture. However, functionally-relevant polymorphisms can cluster in genes expressed within associated tissue and cell types. We tested whether single-nucleotide polymorphisms (SNPs) that explain heritable variance in brain activity (i.e., RSFA) are enriched within genes linked to PVALB and SST, which would provide insight into the molecular basis of the resting-state BOLD fluctuations.

We established the SNP heritability of RSFA. A significant proportion of between-subject variation in cluster-wise RSFA was due to common genetic variants (Fig. 5; $h^2_{SNP}$: limbic A = 0.25 (SE 0.04), limbic B = 0.30 (SE 0.04), cingulo-opercular = 0.21 (SE 0.04), temporo-parietal = 0.29 (SE 0.04), prefrontal = 0.33 (SE 0.04), somato/motor = 0.21 (SE 0.04), visual = 0.07 (SE 0.04)). See Supplementary Fig. 11 for parcel-wise heritability estimates. Interneuron-correlated gene sets were nominated using a “guilt-by-association” logic. That is, genes that were spatially correlated...
to interneuron markers (i.e., SST, PVALB) were assumed to relate to each interneuron subtype. Using cortical AHBA data, genes were rank-ordered by their spatial correlation to each interneuron marker and the top 500 most-correlated genes were selected. Interneuron-related SNP lists were generated for each gene set by identifying variants within ±5000 base pairs from transcription start and stop site of each gene. PVALB and SST SNP sets were non-overlapping. The eQTL variants for each gene set were included, defined using cortical data from the CommonMind consortium and NIH GTEx. We denote the SNP lists for each interneuron gene set as PVALBSNP and SSTSNP (see Supplementary Data). Genetic relatedness matrices were calculated for the UKB sample using each SNP set, and heritability was estimated using GCTA-GREML simultaneously across three partitions: PVALBSNP, SSTSNP, and a partition containing all remaining genotyped variants.
Fig. 6 Schizophrenia polygenic risk predicts brain function and tracks PVALB expression. a Genes were rank-ordered by cortical spatial correlation to SST and PVALB, then divided into 500-gene bins. MAGMA competitive gene set analysis revealed enrichment of polygenic risk for schizophrenia in the top PVALB (p = 0.022), but not the top SST (p = 0.51). Enrichment decreased across ordered bins for PVALB (Spearman’s r = −0.48, p = 0.03) but not for SST (Spearman’s r = −0.01, p = 0.51). b Schizophrenia polygenic risk negatively predicts RSFA within the visual (β = 0.04) and prefrontal (β = 0.10) clusters at trend-levels (corrected for multiple-comparisons). c Parcel-wise prediction of RSFA by the schizophrenia PRS negatively correlated with cortical expression of PVALB (Pearson’s r = −0.33, p = 3.1e−10), which was also significant relative to all genes (PVALB = 145/17,448, AUC = 0.008). d Across all deconvolved cell type distributions, PVALB was the most negatively correlated to cortical SCZ-RSFA effects (frontal cortex: Pearson’s r(337) = −0.36, p = 9.5e−12; visual cortex: Pearson’s r(337) = −0.26, p = 9.0e−7). SCZ = schizophrenia; PRS = polygenic risk score; RSFA = resting state functional amplitude. *p ≤ 0.05. Error bars = standard error.

Indicating that genetic variance in RSFA, a measure of in-vivo functional variability, is explained by genes linked to PVALB interneurons, the PVALB

partitioned heritability (Fig. 5e; frontal cortex: r(329) = 0.35, p = 2.27e−11; r(36) = 0.39, p = 2.2e−13; visual cortex: r(329) = 0.31, p = 6.67e−9; r(36) = 0.34, p = 1.5e−10). These results were highly consistent across parallel techniques using linkage-disequilibrium score regression (LDSC) and estimates of heritability from GWAS analyses of RSFA (Supplementary Fig. 12)44. Cumulatively, these data indicate that the heritable basis of resting-state functional amplitude is spatially heterogeneous, demonstrating a particularly important role of genes co-expressed with PVALB.

Functional associates of schizophrenia genetic risk correlate with PVALB expression. Convergent evidence from animal models and post-mortem tissue analyses suggests that interneuron dysfunction is a core pathophysiological feature of schizophrenia21. To determine whether interneuron-related genetic variation is tied to disease liability, we tested whether polygenic risk for schizophrenia45 is greater among PVALB-SNP and SST-SNP variants, relative to the rest of the genome. Using a partitioned MAGMA analysis46, we divided rank-ordered PVALB and SST gene lists into bins of 500. We observed significant enrichment of schizophrenia polygenic risk for the top PVALB gene set (β = 0.094, p = 0.022), but not the top SST set (β = −0.001, p = 0.51). Suggesting that polygenic schizophrenia risk is greater among interneuron-related genes, we examined 20 gene bins and found that the enrichment of schizophrenia genetic risk decreased as gene sets became less spatially correlated with PVALB (r = −0.48, p = 0.03), but not SST (r = 0.027, p = 0.91; Fig. 6a).

To test whether polygenic risk for schizophrenia relates to cortical RSFA, we calculated schizophrenia polygenic risk scores (SCZ-PRS)47 using genotyped variants from individuals in the UK Biobank imaging sample. Across the seven RSFA clusters, SCZ-PRS negatively predicted RSFA in the visual cluster (Benjamini–Hochberg corrected q = 0.04; Fig. 6b; GWAS threshold p < 1.0), as well as trend-levels in somato/motor (q = 0.08) and prefrontal (q = 0.10) clusters. Consistent with the hypothesized link between PVALB interneurons and psychotic illness, the relationship between RSFA and polygenic schizophrenia risk was negatively correlated to PVALB expression across cortex (Fig. 6c; r(337) = −0.33, p = 3.1e−10; r(36) = −0.35, p = 5.5e−11). That is, regions with greater PVALB expression (e.g., motor and visual parcels) showed stronger negative relationships between SCZ-PRS and RSFA. This relationship remained significant after controlling for overall SNP heritability of each parcel (β = −0.28, t(336) = −5.32, p = 1.95e−7), indicating that the effect isn’t driven by parcel-wise explainable genetic variance. Comparing the RSFA-schizophrenia polygenic risk map to all genes, PVALB was the among the top 0.83% negatively correlated expression profiles (145 out of 17,448; Spearman = 105/17,448 = 0.60%), showing that this relationship is not statistically obligated. Across all deconvolved cell types, parcel-wise PVALB cell fractions were on average the most negatively spatially correlated to schizophrenia risk RSFA effects (Fig. 6d; frontal cortex: r(337) = −0.36, p = 9.5e−12; r(36) = −0.35, p = 2.4e−11; visual cortex: r(337) = −0.26, p = 9.0e−7; r(36) = −0.26, p = 1.3e−6). Ontological
enrichment analysis further revealed that the top 500 genes correlated with *PVALB* in the AHBA data contained genes associated to schizophrenia, neuronal signaling, and gated channel activity (Table 1). Together, these data suggest that schizophrenia-related genetic variants are associated with cell types, particularly parvalbumin interneurons, and could explain functional disruptions across cortex.

**Discussion**

Integrating genetic, transcriptional, and neuroimaging data, we demonstrate that spatial distributions of interneurons are stereotyped across species and development, and explain a substantial portion of the heritable variation in RSFA, a measure of in-vivo brain activity. Somatostatin and parvalbumin interneuron markers were negatively spatially correlated across cortex, a relationship that was robust in early developmental periods in humans and evolutionarily conserved in non-human primates (Fig. 1). Stereotyped patterns of SST and *PVALB* expression were observed in subcortex and were consistent with cell density estimates in rodents (Fig. 3). Computational models theorize that interneuron ratios underlie regional differences in cortical brain function\(^{11}\). Providing empirical support for this hypothesis, relative SST and *PVALB* expression in post-mortem brain tissue aligned with spatial RSFA in the general population (Fig. 4). Indicating the functional relevance of this spatial relationship, genetic polymorphisms linked to *PVALB* correlated genes accounted for an enriched proportion of heritable variance underlying cortical signal amplitude (Fig. 5). Critically, the amount of in-vivo variance explained by *PVALB*-linked SNPs positively tracked spatial expression of *PVALB* in independent post-mortem brain tissue, suggesting that common genetic polymorphisms influence brain function in a cell-preferential and regionally variable manner. Implicating genetic differences among interneurons in schizophrenia, schizophrenia-related polygenic risk was enriched among genes co-expressed with interneurons, and predicted reduced resting-state functional amplitude across cortex, following the spatial landscape of *PVALB* gene expression (Fig. 6).

Adaptive functioning depends on the integration of information across timescales. Higher-order cognition often requires information accumulation over time, whereas sensorimotor processing entails rapid adaption to changing external stimuli\(^{15,18}\). These computational demands are met, in part, through the hierarchical organization of anatomic and functional connections in cortex, as well as cytoarchitectural differences across brain regions\(^{40,48}\). Our data indicate that interneuron ratios, as indexed by SST and *PVALB* expression, are an important feature of regionally variable brain function. Due to unique functional and synaptic properties of somatostatin and parvalbumin interneurons, relative shifts in their density can alter the balance of inhibitory control\(^{11}\), SST interneurons generally synapse onto dendrites of pyramidal neurons to gate incoming cortical signals, whereas PVALB interneurons generally provide perisomatic inhibition that is well-suited for feedback inhibition and output regulation\(^2\). Increased dendritic (i.e., SST) over perisomatic (i.e., PVALB) inhibition could lead to more robust filtering of task-irrelevant information, allowing for greater recurrent excitation and integration of information over time\(^{19}\). Conversely, parvalbumin-dense sensorimotor regions may benefit from fast responses and lower recurrent excitation to adapt to rapidly changing inputs\(^{15}\).

Our analyses provide molecular genetic support for a relationship between parvalbumin interneurons and the hemodynamic signal. BOLD signals are preferentially coupled to gamma-band oscillations (30–80 Hz) relative to other frequency domains\(^{20}\). Importantly, fast-spiking parvalbumin interneurons have been experimentally linked to gamma oscillations\(^{49}\). Here, we provide initial evidence in humans for the influence of parvalbumin interneurons on fMRI signal variability. For instance, polygenic variation among parvalbumin correlated genes explained upwards of 23% of the heritable variance in RSFA in visual cortex.

Schizophrenia is one of the most heritable forms of psychiatric illnesses (\(h^2 \sim 81\%\))\(^{50}\), with converging lines of evidence pointing toward GABAergic and parvalbumin interneuron abnormalities as cardinal features of the disorder\(^{21,51}\). Patients with schizophrenia exhibit reduced levels of GAD67, an enzymatic precursor of GABA\(^{52}\), and are characterized by PVALB interneurons with atypical perineuronal nets\(^{53}\). These abnormalities are thought to underlie disorder-related disruption of gamma-band oscillations and working memory\(^{51}\). Linking these observations, we demonstrate that polygenic schizophrenia risk is increased among genes that are spatially correlated to *PVALB* (Fig. 6a), expanding upon cell transcriptomic work implicating cortical interneurons as an illness marker\(^{40}\). We further show a negative association between individual polygenic schizophrenia risk and RSFA in a large population-based sample (Fig. 6b). The topography of these effects follows the spatial profile of *PVALB* expression across cortex (Fig. 6c), highlighting the potential role of parvalbumin interneurons in mediating brain-based intermediate phenotypes associated with psychotic illness. This report also compliments evidence that schizophrenia risk gene expression is spatially correlated to disorder-related changes in brain morphology\(^{34,35}\).

Disruption of excitatory/inhibitory balance may reflect a cross-diagnostic marker of psychiatric illness\(^{36}\). For instance, decreased expression of parvalbumin cell markers is evident in both schizophrenia and bipolar disorder\(^{37}\), while major depressive disorder

### Table 1 Enrichment terms for interneuron-correlated genes.

| Gene list | Category | ID | Name | p   | q FDR-BH | Hits | Genes in GO |
|-----------|----------|----|------|------|----------|------|-------------|
| SST       | GO: BP   | GO:0099536 | Synaptic signaling | 1.37e–5 | 3.04e–2 | 37   | 687         |
| SST       | GO: BP   | GO:0099537 | Trans-synaptic signaling | 2.43e–5 | 3.04e–2 | 36   | 678         |
| SST       | GO: CC   | GO:0097458 | Neuron part | 1.26e–6 | 6.83e–4 | 66   | 1545        |
| SST       | GO: CC   | GO:0098793 | Presynapse | 4.03e–4 | 3.64e–2 | 20   | 341         |
| PVALB     | GO: MF   | GO:0005261 | Cation channel activity | 3.33e–12 | 1.74e–9 | 33   | 306         |
| PVALB     | GO: MF   | GO:0005249 | Voltage-gated potassium channel activity | 1.33e–10 | 1.69e–8 | 17   | 91          |
| PVALB     | GO: BP   | GO:0071805 | Potassium ion transmembrane transport | 1.53e–9 | 2.43e–6 | 21   | 181         |
| PVALB     | GO: BP   | GO:0098655 | Cation transmembrane transport | 3.23e–9 | 3.09e–6 | 48   | 738         |
| PVALB     | GO: CC   | GO:0034703 | Cation channel complex | 1.04e–10 | 5.07e–8 | 23   | 176         |
| PVALB     | GO: CC   | GO:0034702 | Ion channel complex | 3.03e–10 | 5.07e–8 | 29   | 291         |
| PVALB     | Disease  | C0036341 | Schizophrenia | 5.85e–7 | 1.6e–3 | 68   | 1561        |

Ontological enrichments analysis were conducted with TopGene on the 500 genes used to generate the PVALB50 and SST50 lists. See Supplementary Data for full enrichment tables.
is marked by preferential reductions in somatostatin interneurons. Delineating the region-specific roles of cortical interneuron subtypes will provide insight into cross-diagnostic patterns of both behavior and brain function. With regard to depressed mood, modulation of cortical somatostatin interneurons causally influences anxiety- and depression-like behavioral phenotypes in rodents. In line with this observation, we observed preferential expression of somatostatin within mPFC, NAcc, mediodorsal thalamus, and VTA (Supplementary Fig. 9), a distributed set of regions implicated in reward and affective information processing. Somatostatin-biased regions (ACC, mPFC, and insula) also correspond to areas of cortical thinning in patients with MDD and individuals reporting elevated negative affect.

The present findings should be interpreted in light of several limitations. First, we use single molecular markers to infer the relative presence of SST and PV/ALB interneurons, which are not sensitive to morphological and physiological differences among interneuron subgroups, a point we sought to address through the analyses of deconvolved cell-type distributions. Second, we employ the “global null association” approach to nominate interneuron-related gene sets. While we cannot conclude that genes within each identified interneuron group directly influence interneuron function, similar correlation-based nomination approaches have been shown to correspond well with a priori defined gene groups. However, the examination of enrichment terms (Table 1 and Supplementary Information) allows for a more precise understanding of the biological processes contributing to these results. Third, our measure of brain signal amplitude (RSFA) is likely explained by a mixture of neural and vascular signals. However, effects related to BOLD variability have been shown to be stable even after controlling for measures of vascular lumen and vascular confounds are a caveat of many BOLD measures of brain function. Last, our in vivo imaging and genetic analyses focus on an aging population of white British individuals. Future work should examine the stability of these results across diverse populations.

Inherited genetic variation shapes brain function within and across individuals. Analyses of spatially-dense, whole-genome expression atlas increasingly reveal transcriptional correlates of brain function, structure, functional connectivity, and psychiatric illness. With the parallel emergence of large-scale imaging genetic data, it is now possible to bridge structural genetic, transcriptional, and large-scale neuroimaging brain phenotypes. Here, we leverage these data to show that interneuron marker distributions are consistent across species, correlate with cortical signal amplitude, explain regional differences in heritable brain function, and associate with genetic risk for schizophrenia in the general population.

Methods

Allen Human Brain Atlas. Publicly available human gene expression data from six postmortem donors (1 female), aged 24–57 years (42.5 ± 13.4) were obtained from the Allen Institute. Data reflect the microarray normalization pipeline implemented in March 2013 (https://human.brain-map.org) and analyses were conducted according to the guidelines of the Yale University Human Subjects Committee. Microarray probes from eight overarching ontological categories were selected: cortex, dorsal thalamus, striatum, globus pallidus, hypothalamus, hippocampus, amygdala, and the combined substantia nigra and ventral tegmentum (see Supplementary Information). Probes without Entrez IDs were removed. Probe-wise noise for each donor was quantified as the number of above-threshold samples in cortex, divided by total cortical sample count. A probe-wise average was computed across all six donors, which was used to remove probes expressed in fewer than 20% of cortical samples. If more than one probe existed for a given gene, the one with the highest mean expression level was selected for further analysis, resulting in 17,448 brain-expressed genes.

Individual cortical tissue samples were mapped to each AHBA donor’s Freesurfer derived cortical surfaces, downloaded from Romero-Garcia and colleagues. Native space midthickness surfaces were transformed to a common fslR32k group space while maintaining the native cortical geometry of each individual donor. The native voxel coordinate of each tissue sample was mapped to the midsurface vertex coordinates using tools from the HCP workbench. The cortical tissue sample was not analyzed if it was greater than 4 mm from the nearest surface vertex, resulting in 1683 analyzable cortical samples. Microarray expression of each gene was mean- and variance-normalized separately for each of the eight analyzed regions, revealing relative expression differences within cortical and subcortical territories. For regional analyses (e.g., Fig. 1c), ontological categories from the AHBA were used to calculate the median, min–max, and interquartile range of relative expression in each region. Detailed information about the analyzed regions is provided in the Supplementary Data. Functional data visualization was conducted using wfl view from the HCP workbench. The MNI locations of striatal and thalamic regions were cross-referenced to functional atlases of Choi and colleagues and Hwang and colleagues. With AFNI, a single voxel (1 mm3) region of interest (ROI) was generated at the MNI location of each sample. A functional network label was assigned if the ROI fell within a volumetric parcel. If the sample did not overlap with the functional atlas, the associated ROI was expanded to 2 mm3 and the network with the most overlapping voxels in the ROI was assigned. If the expanded 2 mm3 ROI did not overlap, the process was repeated using a 3 mm3 ROI. A sample was omitted from analysis if the 3 mm3 ROI did not overlap with the associated functional atlas. Functional sub-regions with 3 or fewer samples were excluded from analyses. This process was repeated for coordinates aligned to MNI152 1 mm space using ANTs registration tools (https://github.com/chrisgoralloeninf/tree/master/alleninf/data).

Subcortical AHBA expression data was compared to rodent cell density counts published by Kim et al. Z-transformed AHBA expression values were summarized across major subdivisions of the seven subcortical regions analyzed (e.g., CA1, NAcc, etc.; Supplementary Data). Median AHBA expression values were expressed for analyses in Fig. 3b. Rodent homologs of each sub-region were manually identified and PV/ALB and SST cell densities were averaged across male and female samples.

UKB imaging processing. Minimally preprocessed resting-state fMRI data from the UK Biobank were analyzed, reflecting the following preprocessing steps: motion correction with MClflirt, grand-mean intensity normalization, highpass temporal filtering, fieldmap unwarping, and gradient distortion correction. Noise terms were identified and removed using FSL ICA+FIx. Full information on the UKB imaging processing is published. Additional processing was conducted in AFNI and consisted of 3dDespike, resampling to MNI152 space using the UKB generated linear and nonlinear transforms, FWHM blur of 4.0 mm, regression of CSF, WM, and global resting state signals, and first- and second-order trend removal. Voxelwise RSFA maps were generated with 3dsRSCF and then averaged within each of the approximately symmetrical 400 volumetric parcels from the 7-network parcellation of Schaefer and colleagues. Due to signal blurring between lateral striatum and insular cortex, supplemental resting-state functional connectivity analyses reflect an additional local white matter regression against gray matter using AFNI anatcor. Imaging analyses were conducted in volume, but visualized on the cortical surface. Resting-state functional connectivity was calculated in the thalamus, and cortex was estimated using AFNI’s 3dNetCorr, which calculated the Fisher-Z transformed correlation values of timeseries across the Choi 7-region striatal atlas, the Hwang 9-region thalamic atlas, and the Schaefer 400-region cortical atlas. TheAtlas of 13,236 UKB subjects were processed through the imaging pipeline. Subjects with mean run-wise frame-to-frame head motion greater than 0.20 mm, and inverted resting-state SNR greater than three standard deviations above the mean were removed. After filtering for white British subjects with usable genetic data, cryptic relatedness <0.025, and conducting row-wise deletion for the variables age, sex, height, weight, BMI, three head position coordinates (X, Y, Z), combined gray/white matter volume, combined ventricular/CSF volume, diastolic and systolic blood pressure, run-wise resting state motion, resting state inverse SNR, T1 inverse SNR, and UK Biobank assessment center, 9713 subjects remained for analyses (percent female = 54.33, mean age = 63.67, SD = 7.45, min/max age = 45–80). We also excluded the anthropometric measures of height, weight, and blood pressure. Analyses were conducted according to the guidelines of the Yale University IRB.

UKB genetics. UK Biobank genotype data was filtered to include only white British subjects with imaging data passing the quality control thresholds described above. Plink v1.9 was used to remove samples with missingness >0.10, SNPs with minor allele frequency <0.05, Hardy-Weinberg equilibrium p < 1 × 10−5, and call rate <0.02, resulting in 337,501 autosomal variants. GCTA software was used to calculate a genetic relatedness matrix to remove individuals with cryptic relatedness more than 0.025, leaving N = 9713 subjects for analysis. Ten genetic principal components were then calculated for use as covariates in polygenic risk score and heritability analyses. When calculating polygenic risk for schizophrenia, SNPs from the major histocompatibility complex were censored except for the most significantly associated variant from the region.
**RSFA between-subject clustering and heritability.** Voxel-wise RSFA data from the (N = 9173) UK Biobank sample was averaged within each of 400 roughly symmetric ROIs from the 7-Network cortical parcellation of Schaefer and colleagues.32. Parcell-wise RSFA values were residualized for the effect of age, sex, age \* sex, age\(^2\) \* sex, height, weight, BMI, combined gray/white matter volume (normalized for head size), combined ventricular/CSF volume (normalized for head size), diastolic and systolic blood pressure, run-wise rsMRI motion, rsMRI inverse SNR, T1 inverse SNR, three head position coordinates (X, Y, Z), and UK Biobank assessment center. Hierarchical clustering of residualized RSFA estimates was conducted using R in order to group regions with similar between-subject patterns of covariation. A 7-parcel RSFA clustering was selected. Raw RSFA values were then parceled for each of the partitioning for the 7 Network cortical sub-area for use in heritability analyses. SNP heritability of RSFA was estimated with genotyped data using GCTA-REML software. Age, sex, age\(^2\), height, weight, BMI, combined normed gray/white matter volume, combined normed ventricular/CSF volume, diastolic and systolic blood pressure, run-wise rsMRI motion, rsMRI inverse SNR, T1 inverse SNR, head coordinates (X, Y, Z), and UK Biobank assessment center, and 10 genetic principal components were included as covariates.

**Partitioned heritability analyses** were conducted for the seven RSFA clusters and for each of the 400 individual cortical parcels. Using AHBA expression data, genes were ranked ordered by the spatial correction to SST and PVALB. Genes without Entrez IDs were removed. The BioMart package36 was used to identify each gene’s transcription start and stop sites (±5000 base pairs) according to the GRCh37-hg19 genome Assembly. Otherwise, the gene was cross-referenced to the current eQTL databases from the NIH GTEx project43 and CommonMind consortium.44 Intragenic (±5000 base pairs) and eQTL SNPs associated with the top 500 SST (N = 9571) and PVALB (N = 9713) correlated genes were used for partitioned heritability analyses, respectively denoted SST\(_{SNP}\) and PVALB\(_{SNP}\). A small subset of genes in each bin did not have analyzable variants using these criteria and thus did not contribute to results. Genetic-relatedness matrices for the SST\(_{SNP}\) and PVALB\(_{SNP}\) partitions were generated, as well as one for all remaining genotyped SNPs. RSFA heritability accounted for each genetic relatedness matrix was estimated simultaneously for each of the three partitions using GCTA.45 Partitioned heritability was then defined as the phenotypic variance explained by each partition, divided by the total phenotypic variance. To calculate the significance of individual partitions, we consider the Wald test statistic against the null of \(h^2 = 0\), which follows a half-half mixture of \(\chi^2\) distributions with a probability mass at zero and \(\chi^2\) (a \(\chi^2\) distribution with 1 degree of freedom). Enrichment values were calculated to determine if the proportion of variation explained by a particular partition was greater than the proportion of variation partitioned, defined as:

\[
\text{enrich}_{\text{part}} = \frac{h^2_{\text{SNP}}/n_{\text{total}}}{h^2_{\text{bin}}/n_{\text{total}}}
\]

where \(h^2_{\text{SNP}}\) is the heritable variance explained by the SNP partition (e.g., PVALB\(_{SNP}\)), \(h^2_{\text{bin}}\) is the heritable variance explained by all partitions, \(n_{\text{total}}\) is the total number of genotyped SNPs. Standard error for SNP heritabilities were calculated by the Wald test, using precomputed LD scores from 1,158,800 HapMap3 variants that overlapped with AHBA expression data. Partitioned heritability analyses were conducted by examining SNPs occurring near the coding regions of variants in the UK Biobank GWAS, using precomputed allele frequencies (i.e., 1000G_frq). 500-gene bins were used to divide the genome into roughly equal parts, and the start and end positions of each bin were defined as the first and last positions of the most significant SNP (denoted by \(\chi^2\)). Partitioned heritability values were converted from log10 to log2. Data from two macaques (one female) were excluded due to sparse sampling across cortex. Samples from the following 10 cortical regions were included in our analyses: OFC, ACC, medial temporal lobe, temporal area, DLPFc, A1C, S1C, M1C, V1, and V2. The collapseRows function was used in R to select the probe with the highest mean expression. UK ComBat was used to remove residual batch effects. PVALB expression was mean and variance-normalized to reveal relative expression differences across cortex.

**BrainSpan processing.** Publicly available RNAseq reads per kilobase per million (RPKM) data from the Brainspan atlas were used to characterize patterns of interneuron-marker gene expression across development. Cortical tissue samples were collected from early/late fetal (21–27 post-conception weeks [pcw]; donors = 10, samples = 88), early/mid fetal (13–21 pcw, donors = 10, samples = 88), late fetal (24–37 pcw; donors = 5, samples = 27), early infancy (4 months; donors = 3, samples = 22), late infancy (10 months; donors = 10, samples = 8), early childhood (1–4 years; donors = 5, samples = 41), mid/late childhood (8–11 years; donors = 2, samples = 30), adolescence (13–15 years; donors = 2, samples = 14), and adult (18–40 years; donors = 8, samples = 85) developmental stages. RNAseq probes without Entrez IDs were excluded and duplicated probes were removed by selecting the probe with the highest mean expression. Data was log2 transformed and the effect of donor was removed separately for each age group using ComBat. Polygenic expression was then mean- and variance-normalized across cortical tissue samples separately for each developmental stage. When multiple ages were present in a development stage, age was included as a covariate in a linear regression predicting normalized SST expression from normalized PVALB expression.

**Single-cell analysis and deconvolution.** Single-nucleus droplet-based sequencing (snDrop-seq) data from Lake and colleagues51 was obtained from the Gene Expression Omnibus database (GSE87936). Expression values were converted from log10 to log2. To assess whether schizophrenia polygenic risk was enriched among corticopetal markers using single-cell data. Cell type fractions for each AHBA cortical sample were mapped to the cortical surface and summarized across cortex. The LDSC implementation with single-cell data and CibersortX (Fig.2). We also identify stromal markers in cortex, using microarray data from the AHBA. This effect was replicated with independent measures of cell density in rodents (Fig.3). The LDSC method was used to replicate patterns of SST and PVALB expression in subcortical territories, which were replicated with independent measures of cell density in rodents (Fig.3). Spatial associations between cortical function (i.e., RSFA) and markers of interneurons were also established with two techniques. The first approach examined highly cell-specific individual gene markers and the second approach utilized markers of interneurons in cortex, using microarray data from the AHBA. This effect was replicated with independent measures of cell density in rodents (Fig.3). Spatial associations between cortical function (i.e., RSFA) and markers of interneurons were also established with two techniques. The first approach examined highly cell-specific individual gene markers and the second approach utilized markers of interneurons in cortex, using microarray data from the AHBA. This effect was replicated with independent measures of cell density in rodents (Fig.3). Spatial associations between cortical function (i.e., RSFA) and markers of interneurons were also established with two techniques. The first approach examined highly cell-specific individual gene markers and the second approach utilized markers of interneurons in cortex, using microarray data from the AHBA. This effect was replicated with independent measures of cell density in rodents (Fig.3). Spatial associations between cortical function (i.e., RSFA) and markers of interneurons were also established with two techniques. The first approach examined highly cell-specific individual gene markers and the second approach utilized markers of interneurons in cortex, using microarray data from the AHBA. This effect was replicated with independent measures of cell density in rodents (Fig.3). Spatial associations between cortical function (i.e., RSFA) and markers of interneurons were also established with two techniques. The first approach examined highly cell-specific individual gene markers and the second approach utilized markers of interneurons in cortex, using microarray data from the AHBA. This effect was replicated with independent measures of cell density in rodents (Fig.3). Spatial associations between cortical function (i.e., RSFA) and markers of interneurons were also established with two techniques. The first approach examined highly cell-specific individual gene markers and the second approach utilized markers of interneurons in cortex, using microarray data from the AHBA. This effect was replicated with independent measures of cell density in rodents (Fig.3). Spatial associations between cortical function (i.e., RSFA) and markers of interneurons were also established with two techniques. The first approach examined highly cell-specific individual gene markers and the second approach utilized markers of interneurons in cortex, using microarray data from the AHBA. This effect was replicated with independent measures of cell density in rodents (Fig.3). Spatial associations between cortical function (i.e., RSFA) and markers of interneurons were also established with two techniques. The first approach examined highly cell-specific individual gene markers and the second approach utilized markers of interneurons in cortex, using microarray data from the AHBA. This effect was replicated with independent measures of cell density in rodents (Fig.3). Spatial associations between cortical function (i.e., RSFA) and markers of interneurons were also established with two techniques. The first approach examined highly cell-specific individual gene markers and the second approach utilized markers of interneurons in cortex, using microarray data from the AHBA. This effect was replicated with independent measures of cell density in rodents (Fig.3). Spatial associations between cortical function (i.e., RSFA) and markers of interneurons were also established with two techniques. The first approach examined highly cell-specific individual gene markers and the second approach utilized markers of interneurons in cortex, using microarray data from the AHBA. This effect was replicated with independent measures of cell density in rodents (Fig.3).

**Replication analyses.** In this paper, we sought to replicate findings with alternative methodologies and independent datasets whenever possible. In Fig. 1, we document a negative spatial relationship between SST and PVALB interneuron markers in cortex, using microarray data from the AHBA. This effect was replicated using RNAseq data from the AHBA (Supplementary Fig. 4), in an independent sample from the same location (Fig. 1b). Single-cell data were then analyzed using CIBERSORTx to impute cell type fractions in the AHBA data. Independent gene signature matrices were defined with visual cortex and frontal cortex cell data. Cell type fractions were then estimated separately for each AHBA donor, once using a visual cell signature and once using a frontal cortex cell signature. Cell type fractions for each AJBA cortical sample were mapped to the cortical surface and summarized among Schaefer et al., atlas ROIs in the same manner as single-gene expression data.
polygenic cellular deconvolution based on multivariate transcriptional signatures of cell types (Fig. 4). Further, heritability and partitioned heritability analyses of cortical RSFA were replicated with two techniques, GCTA and LD score regression (Fig. 5 and Supplementary Fig. 12). These methods yielded highly convergent results that were robust to genetic preprocessing choices (e.g., analysis of genotyped versus imputed variants) and model assumptions of underlying genetic architecture.

**Reporting summary**. Further information on research design is available in the Nature Research Reporting Summary linked to this article.

**Data availability**. The data that support these findings are either publicly available, provided with the paper, or are under third party restrictions. Publicly available data are accessible at the following locations: Allen Human Brain Atlas (https://human.brain-map.org/), UK Biobank (https://www.ukbiobank.ac.uk/), Brainscan Atlas of the Developing Human Brain (https://www.brainspan.org/), NIH Blueprint NHP Atlas (https://www.blueprintnhapsatlases.nih.gov), NIH GTEx (https://gtexportal.org/gtex), CommonMind (https://www.nlmgenetics.org/resources/commonmind), snDrop-seq data at Gene Expression Omnibus “GSE97942” (https://www.ncbi.nlm.nih.gov/geo/), and GWAS data from the Psychiatric Genomics Consortium (https://www.med.yale.edu/pgc/).

Data not under third-party restrictions are available at https://github.com/HolmesLab/2020_NatComm_interneurons_cortical_function_schizophrenia.

**Code availability**. Code not under third-party restrictions are available at https://github.com/HolmesLab/2020_NatComm_interneurons_cortical_function_schizophrenia.

Received: 24 October 2019; Accepted: 19 May 2020; Published online: 08 June 2020

**References**

1. Ramón y Cajal, S. Recollections of My Life (The MIT Press, 1989).
2. Kepes, A. & Fishell, G. Interneuron cell types are fit to function. Nature 505, 318–326 (2014).
3. Markram, H. et al. Interneurons of the neocortical inhibitory system. Nat. Rev. Neurosci. 5, 793–807 (2004).
4. Gonzalez-Burgos, S., Cho, R. Y. & Lewis, D. A. Alterations in cortical network oscillations and parvalbumin neurons in schizophrenia. Biol. Psychiatry 77, 1031–1040 (2015).
5. Fee, C., Banares, M. & Sible, E. Somatostatin-positive gamma-aminobutyric acid interneuron deficits in depression: cortical microcircuit and therapeutic perspectives. Biol. Psychiatry 82, 549–559 (2017).
6. Richichi, J. et al. Correlated gene expression supports synchronous activity in brain networks. Science 348, 1241–1244 (2015).
7. Krienen, F. M., Yeo, B. T. T., Ge, T., Buckner, R. L. & Sherwood, C. C. Transcriptional profiles of supragranular-enriched genes associate with corticocortical network architecture in the human brain. Proc. Natl. Acad. Sci. USA 113, E469–E478 (2016).
8. Wang, G. Z. et al. Correspondence between resting-state activity and brain gene expression, Neuron 88, 659–666 (2015).
9. Anderson, K. M. et al. Gene expression links functional networks across cortex and striatum. Nat. Commun. 9, 1428 (2018).
10. Burt, J. B. et al. Hierarchy of transcriptomic specialization across human cortical territory captured by structural neuroimaging topology. Nat. Neurosci. 21, 1251–1259 (2018).
11. Kim, Y. et al. Brain-wide maps reveal stereotyped cell-type-based cortical architecture and subcortical sex dimorphism. Cell 171, 456–459 (2017).
12. Hendry, S. H. C., Schwark, H. D., Jones, E. G. & Yan, J. Numbers and proportions of GABA-immunoreactive neurons in different areas of monkey cerebral cortex. J. Neurosci. 7, 1503–1519 (1987).
13. Kawaguchi, Y. & Kubota, Y. GABAergic cell subtypes and their synaptic connections in rat frontal cortex. Cereb. Cortex 7, 476–486 (1997).
14. Wang, X. J., Tegner, J., Constantinidis, C. & Goldman-Rakic, P. S. Division of labor among distinct subtypes of inhibitory neurons in a cortical microcircuit of working memory. Proc. Natl. Acad. Sci. USA 101, 1368–1373 (2004).
15. Chaudhuri, R., Knoblauch, K., Gariel, M.-A., Kennedy, H. & Wang, X.-J. A large-scale circuit mechanism for hierarchical dynamical processing in the primate cortex. Neuron 88, 419–431 (2015).
16. Cioli, C., Abdi, H., Beaton, D., Burnod, Y. & Mesoudi, S. Differences in human cortical gene expression match the temporal properties of large-scale functional networks. PLoS ONE 9, e115913 (2014).
Akbarian, S. et al. Gene expression for glutamic acid decarboxylase is reduced without loss of neurons in prefrontal cortex of schizophrenics. Arch. Gen. Psychiatry 52, 258–266 (1995).

Mauney, S. A. et al. Developmental pattern of perineuronal nets in the human prefrontal cortex and their deficit in schizophrenia. Biol. Psychiatry 74, 427–435 (2013).

Morgan, S. E. et al. Cortical patterning of abnormal morphometric similarity in schizophrenia is associated with brain expression of schizophrenia-related genes. Proc. Natl. Acad. Sci. USA 116, 9604–9609 (2019).

Romme, I. A. C., de Reus, M. A., Ophoff, R. A., Kahn, R. S. & van den Heuvel, M. P. Connectome disconnectivity and cortical gene expression in patients with schizophrenia. Biol. Psychiatry 81, 495–502 (2017).

Marin, O. Interneuron dysfunction in psychiatric disorders. Nat. Rev. Neurosci. 13, 107–120 (2012).

Toker, L., Mancarci, B. O., Tripathy, S. & Pavlidis, P. Transcriptomic evidence for alterations in astrocytes and parvalbumin interneurons in subjects with bipolar disorder and schizophrenia. Biol. Psychiatry 84, 787–796 (2018).

Fuchs, T. et al. Disinhibition of somatostatin-positive GABAergic interneurons results in an anxiolytic and antidepressant-like brain state. Mol. Psychiatry 22, 920–930 (2017).

Schmaal, L. et al. Cortical abnormalities in adults and adolescents with major depression based on brain scans from 20 cohorts worldwide in the ENIGMA Major Depressive Disorder Working Group. Mol. Psychiatry 22, 900–909 (2017).

Anderson, K. M. et al. Convergent molecular, cellular, and neural signatures of major depressive disorder. Preprint at https://doi.org/10.1101/2020.02.09.942277 (2020).

Holmes, A. J. et al. Individual differences in amygdala-medial prefrontal anatomy link negative affect, impaired social functioning, and polygenic depression risk. J. Neurosci. 32, 18087–18100 (2012).

Welch, G. S. & Butte, A. J. Systemic survey reveals general applicability of ‘guilt-by-association’ within gene coexpression networks. BMC Bioinformatics 6, 227 (2005).

Garrett, D. D., Lindenberger, U., Hoge, R. D. & Gauthier, C. J. Age differences in brain signal variability are robust to multiple vascular controls. Sci. Rep. 7, 10149 (2017).

Duncan, L. et al. Analysis of polygenic risk score usage and performance in diverse human populations. Nat. Commun. 10, 3328 (2019).

Elliot, M. L. et al. A polygenic score for higher educational attainment is associated with larger brains. Cereb. Cortex 491, 56–59 (2018).

Ge, T., Holmes, A. J., Buckner, R. L., Smoller, J. W. & Sabuncu, M. R. Heritability analysis with repeat measurements and its application to resting-state functional connectivity. Proc. Natl. Acad. Sci. USA 114, 5521–5526 (2017).

Reardon, P. K. et al. Normative brain size variation and brain shape diversity in humans. Science 360, 1222–1227 (2018).

Armatkevičiūtė, A., Fulcher, B. D. & Fornito, A. A practical guide to linking brain-wide gene expression and neuroimaging data. NeuroImage 189, 353–367 (2019).

Romero-Garcia, R. et al. Structural covariance networks are coupled to expression of genes enriched in supragranular layers of the human cortex. NeuroImage 171, 256–267 (2018).

Marcus, D. S. et al. Informatics and data mining tools and strategies for the Human Connectome Project. Front. Neuroinformatics 5, 1–12 (2011).

Choi, E. Y., Yeo, B. T. T. & Buckner, R. L. The organization of the human striatum estimated by intrinsic functional connectivity. J. Neurophysiol. 108, 2242–2263 (2012).

Hwang, K., Bertolero, M. A., Liu, W. B. & D’Esposito, M. The human thalamus is an integrative hub for functional brain networks. J. Neurosci. 37, 5594–5607 (2017).

Jenkinson, M., Bannister, P., Brady, M. & Smith, S. Improved optimization for the robust and accurate linear registration and motion correction of brain images. NeuroImage 17, 825–841 (2002).

Cox, R. W. AFNI: software for analysis and visualization of functional magnetic resonance images. Comput. Biomed. Res. 29, 162–173 (1996).

Purcell, S. et al. PLINK: a tool set for whole-genome association and population-based linkage analyses. Am. J. Hum. Genet. 81, 559–575 (2007).

Durinck, S. et al. BioMart and Bioconductor: a powerful link between biological databases and microarray data analysis. Bioinformatics 21, 3439–3440 (2005).

Jiang, L. et al. A resource-efficient tool for mixed model association analysis of large-scale data. Nat. Genet. 51, 1749–1755 (2019).

Bakken, T. E. et al. A comprehensive transcriptional map of primate brain development. Nature 535, 367–375 (2016).

Johnson, W. E., Li, C. & Rabinovic, A. Adjusting batch effects in microarray expression data using empirical Bayes methods. Biostatistics 8, 118–126 (2007).

Stuart, T. et al. Comprehensive integration of single-cell data. Cell 177, 1888–1902 (2019).