A recent publication reported an exciting polygenic effect of schizophrenia (SCZ) risk variants, identified by a large genome-wide association study (GWAS), on total brain and white matter volumes in schizophrenic patients and, even more prominently, in healthy subjects. The aim of the present work was to replicate and then potentially extend these findings. According to the original publication, polygenic risk scores—using single nucleotide polymorphism (SNP) information of SCZ GWAS—(polygenic SCZ risk scores; PSS) were calculated in 122 healthy subjects, enrolled in a structural magnetic resonance imaging (MRI) study. These scores were computed based on P-values and odds ratios available through the Psychiatric GWAS Consortium. In addition, polygenic white matter scores (PWM) were calculated, using the respective SNP subset in the original publication. None of the polygenic scores, either PSS or PWM, were found to be associated with total brain, white matter or gray matter volume in our replicate sample. Minor differences between the original and the present study that might have contributed to lack of reproducibility (but unlikely explain it fully), are number of subjects, ethnicity, age distribution, array technology, SNP imputation quality and MRI scanner type. In contrast to the original publication, our results do not reveal the slightest signal of association of the described sets of GWAS-identified SCZ risk variants with brain volumes in adults. Caution is indicated in interpreting studies building on polygenic risk scores without replication sample.

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Keywords: genome-wide association study; healthy subjects; MRI; risk genotypes; schizophrenia; SNP

INTRODUCTION

The Psychiatric Genome-Wide Association Study (GWAS) Consortium has recently published a large GWAS, including 9394 schizophrenia (SCZ) cases and 12 462 healthy controls, identifying common variants that contribute to SCZ susceptibility with relatively small odds ratios. Besides these genome-wide significant loci, evidence has been accruing that a significant proportion of the risk for SCZ may lie in markers not achieving genome-wide significance in GWAS. For instance, a quantitative polygenic SCZ risk score (PSS) was calculated based on the nominally associated alleles in a discovery sample. This polygenic score explained up to 3% of variance in SCZ in a number of independent samples. Several authors have subsequently explored whether such a polygenic effect might be associated not only with the disease but also with disease-relevant phenotypes. Although some studies described associations for example, with cognitive aging or with a functional imaging substrate of working memory processing, others reported a lack of association with psychosis dimensions or intelligence.

Along these lines, a recent study investigated the polygenic effect of SCZ-associated single nucleotide polymorphisms (SNPs) on brain volume (total brain, white matter and gray matter). The proportion of variance explained by the PSS was around 5% for both total brain and white matter volumes. The authors subsequently generated a polygenic white matter score (PWM) out of those 2020 SCZ-related SNPs showing the most significant associations with white matter volume in their sample. This PWM, that is, a final subset of 186 SNPs, influenced white matter volume most strongly. Importantly, effects were not only detected in patients but also in healthy control subjects, leading to the author’s assumption that ‘a relatively small subset of SCZ genetic risk variants is related to the normal development of white matter’.

Considering the potentially high general importance of these findings for the complex genetics and the disease-unrelated biological grounds of adult brain dimensions, the present study has been designed to replicate (i) the PSS and (ii) the PWM effects on total brain and white matter volume in healthy subjects.

MATERIALS AND METHODS

Participants and inclusion criteria

The study was approved by the Santa Lucia Foundation Ethical Committee and performed in accordance with the Helsinki Declaration. After signing an informed consent form, 122 healthy subjects of Italian origin were included. Participants were consecutively recruited by local advertisement from universities, community recreational centers and hospitals (personnel). Inclusion criteria were age 20–80 years and suitability for magnetic resonance imaging (MRI) scanning. Exclusion criteria included: (i) suspicion of cognitive impairment (score ≤26) or dementia based on Mini Mental State examination, the Mental Deterioration Battery and the NINCDS-ADRDIA criteria for dementia; (ii) subjective complaint of memory difficulties or of any other cognitive deficit, interfering with daily living;
(iii) presence of major non-stabilized medical illnesses (that is, non-stabilized diabetes, obstructive pulmonary disease or asthma, hematologic/oncologic disorders, vitamin B12 or folate deficiency, pernicious anemia, clinically significant and unstable active gastrointestinal, renal, hepatic, endocrine or cardiovascular disorders and recently treated hypothyroidism); (iv) known or suspected history of alcoholism, drug dependence and abuse, head trauma and mental disorders according to the Diagnostic and Statistical Manual of Mental Disorders, Fourth Edition–Text Revision (DSM-IV-TR) criteria (all subjects were interviewed by Structured Clinical Interview for DSM disorders–Non-Patient edition (SCID-NP), DSM-IV-TR); (v) presence of vascular brain lesions, brain tumor and/or marked cortical and subcortical atrophy on MRI scan. In particular, the prefrontal severity and location of vascular lesions were rated according to a protocol designed for the Rotterdam Scan Study.1,13 Generally, they were considered present in cases of hyperintense lesions on both proton-density and T2-weighted (see image acquisition) and rated semiquantitatively as 0 (none), 1 (pencil-thin lining), 2 (smooth halo) or 3 (large confluent) for three separate regions: adjacent to frontal horns (frontal caps), adjacent to the wall of the lateral ventricles (bands) and adjacent to the occipital horns (occipital caps). The total vascular lesion load was calculated by adding the region-specific scores (range 0–9). In the present study, only participants rated 0–1 were included.

Image acquisition and processing

All participants underwent the same imaging protocol, which included standard clinical sequences (Fluid Attenuated Inversion Recovery (FLAIR), Proton Density-T2-weighted) and a whole-brain high-resolution T1-weighted sequence obtained in the sagittal plane using a modified driven equilibrium Fourier transform sequence (TE/TR = 2.4/7.92 ms, flip angle: 15°, voxel size: 1 mm³) using a 3 T Allegra MR imager (Siemens, Erlangen, Germany) with a standard quadrature head coil. All planar sequence acquisitions were obtained in the plane of the anterior commissure–posterior commissure line. Particular care was taken to center the subject in the head coil and to restrain the subject’s movements with cushions and adhesive medical tape. MRI-based quantification of cerebral volumes was performed using Freesurfer (v.4.0.5) software package (http://surfer.nmr.mgh.harvard.edu). Freesurfer includes a sophisticated automated segmentation algorithm, which delineates gross brain anatomy into a series of cortical and subcortical labels. The stream consists of five different stages, fully described elsewhere.1,13,14 Initially, the MRI volumes were registered to the Talairach space and the output images were intensity normalized. At this stage, manual intervention is needed to visualize and edit areas of skull and the areas of cortex or cerebellum that should be corrected. After skull stripping, the output brain mask was labeled using a probabilistic atlas,15 and a complex algorithm combining information on image intensity, probabilistic atlas location and the local spatial relationships between structures.16–18 For the purpose of this study, calculated volumes (in mm³, subsequently converted to ml) for these labels were summed up to derive estimates of total gray and white matter volume as well as total brain volume. The Freesurfer software and its documentation can be downloaded from http://surfer.nmr.mgh.harvard.edu.

Discovery sample data

Summary results including risk variants, their P-values and associated odds ratios from the recent international collaborative GWAS on 9394 SCZ cases and 12 462 healthy controls were collected from the Psychiatric GWAS Consortium. Relevant information on the methods used by the consortium is described elsewhere.1

Genetic analysis of the target sample

Genotyping of the Italian target sample was performed using a semi-custom Affymetrix myDesign genotyping array (Affymetrix, Santa Clara, CA, USA), based on a CEU (Caucasian residents of European ancestry from UT, USA) marker backbone including 518 722 SNPs, and a custom marker set including 102 537 SNPs. The array was designed using the Affymetrix Design center (www. affymetrix.com), applying diverse selection criteria.19 Genotyping was performed by Affymetrix on a GeneTitan platform. Several quality control steps were applied (SNP call rate >97%, Fisher’s linear discriminant, heterozygous cluster strength offset, homozygote ratio offset). In a subsequent step, some SNPs were filtered out based on minor allele frequency <0.02 or if the χ²-test for Hardy–Weinberg equilibrium was <1×10⁻⁶.19 For the present study, SNPs on chromosomes X and Y and mitochondrial DNA were excluded, leaving 574 505 SNPs for analyses. This SNP set was then used to calculate multidimensional scaling components in order to control for population stratification. Similarly, the inbreeding coefficient was calculated by making use of a pruned version of this SNP set. MDS components and inbreeding coefficients were calculated using PLINK.20

Derivation of polygenic scores

Polygenic SCZ risk scores. Markers directly genotyped in the target sample were used for the generation of this score. As described in the Psychiatric Genomics Consortium guidelines regarding polygenic risk profile analyses (http://pgc.unc.edu), all SNPs in the extended major histocompatibility complex region were removed at this stage, except one representing the best hit in this region. PSS were calculated following the methods described by Purcell et al. In brief, sets of SNPs with P-values below different cutoffs (0.0001; 0.0005; 0.001; 0.005; 0.01; 0.05; 0.1; 0.2; 0.3; 0.4) in the discovery sample were defined. In order to identify polygenic effects

Table 1. Variance explained (R²) and P-values of the association of PSS (based on different P-value thresholds) and of PWM (based on the complete or partial SNP set) with total brain, white matter and gray matter volumes in the healthy target sample (N = 122)

| No. SNPs | Total brain | White matter | Gray matter |
|----------|-------------|--------------|-------------|
|          | R² change | P-value | R² change | P-value | R² change | P-value |
| PSS (polygenic SCZ risk score) | | | | | | |
| P < 0.0001 | 106 | –0.007 | 0.613 | –0.008 | 0.740 | –0.007 | 0.590 |
| P < 0.0005 | 311 | –0.009 | 0.928 | –0.010 | 0.903 | –0.009 | 0.801 |
| P < 0.001 | 482 | 0.007 | 0.180 | –0.009 | 0.776 | 0.023 | 0.054 |
| P < 0.005 | 1480 | –0.006 | 0.559 | –0.009 | 0.983 | 0.000 | 0.320 |
| P < 0.01 | 2528 | –0.005 | 0.507 | –0.006 | 0.558 | –0.005 | 0.570 |
| P < 0.05 | 8555 | –0.006 | 0.582 | –0.007 | 0.613 | –0.008 | 0.710 |
| P < 0.1 | 14491 | –0.009 | 0.819 | –0.008 | 0.678 | –0.009 | 0.999 |
| P < 0.2 | 24670 | –0.008 | 0.689 | –0.009 | 0.856 | –0.007 | 0.624 |
| P < 0.3 | 33660 | –0.010 | 0.635 | –0.010 | 0.941 | –0.005 | 0.489 |
| P < 0.4 | 41398 | –0.007 | 0.661 | –0.007 | 0.941 | –0.003 | 0.445 |
| PWM (polygenic white matter score) | | | | | | |
| Complete SNP set | 185 | –0.008 | 0.719 | –0.009 | 0.744 | –0.001 | 0.333 |
| Partial set | 142 | –0.002 | 0.391 | –0.009 | 0.872 | 0.006 | 0.187 |

Abbreviations: SCZ, schizophrenia; SNP, single-nucleotide polymorphism. R² refers to R² adjusted for the number of predictors in the model.
due to independent SNPs in linkage equilibrium, each SNP set was pruned based on a pairwise $r^2$ threshold of 0.25 and a sliding window of 50 SNPs shifting 5 SNPs at each step (see Table 1 for the total number of SNPs finally included under each $P$-value threshold). For each subject in the target sample, a PSS was calculated for the different $P$-value thresholds. For each SNP, the number of risk variants (0, 1, 2) an individual carries was multiplied by the logarithm of the odds ratio for that particular variant. Both pruning and scoring were performed using PLINK.20

Polygenic white matter scores. A total of 66 out of the 186 SNPs that constitute the PWM7 were directly genotyped in the Axiom array. The remaining 120 SNPs were imputed using IMPUTE v2.212. Imputation was performed with the 1000 Genomes Project Phase 1 (March 2012) reference panel, and best-guess genotypes were used for polygenic scoring. One of the SNPs (rs9880959) could not be imputed. Mean ± s.d. imputation quality score was 0.906 ± 0.127. First, PWM including the complete set of 185 SNPs was calculated following the previously described methods. Second, an alternative PWM was calculated using a partial set of 142 SNPs, including only those directly genotyped and those with imputation quality score $>0.9$ (Figure 1).

Statistical analysis
Standardized values of total brain, white matter and gray matter volumes were obtained after correction for age, gender and intracranial volume. These corrected brain dimensions were used as dependent variables in a linear regression model. Ten multidimensional scaling components, the inbreeding coefficient and the number of SNPs used to calculate the polygenic scores were selected as covariates of potential relevance. $R^2$ values derived from a model including all of these covariates were subtracted from $R^2$ values from a model including covariates plus the respective polygenic score. The difference between the $R^2$ adjusted for the number of predictors in the model represents the increase in the variance explained attributable to the polygenic score. All these calculations as well as sign tests were carried out with SPSS 17.0 (IBM-Deutschland GmbH, Munich, Germany).22 PLINK20 was used for testing association of each of the 185 SNPs belonging to PWM with the different brain volumetric variables. QUANTO software23 was used for power calculations in the target sample.

RESULTS
The target sample comprised a total of 122 Italian subjects (56 males, 66 females), aged $45.0 ± 17.0$ (mean ± s.d.) years (range 20–80), with an education level of $14.4 ± 3.7$ (mean ± s.d.) years (range 5–24 years). Mean volumes (± s.d.) (ml) were $978.1 ± 112.7$, $407.1 ± 55.7$ and $551.5 ± 65.5$ for total brain, white matter and gray matter, respectively.

PSS were not associated ($P>0.05$ in all comparisons) with any of the brain dimensions in the present sample of healthy subjects, irrespective of the $P$-value cutoffs selected for analysis (Table 1).

**Figure 1.** Scheme of source and number of single nucleotide polymorphisms (SNPs) used to calculate polygenic white matter scores (PWM).

![Diagram](image_url)
group by removing individuals that were >55 years old failed to provide even the least signal of an association between PSS or PWM and total brain or white matter dimensions.

The original study\(^7\) used the Illumina HumanHap550 beadchip (Illumina) for SNP genotyping, whereas we used semi-custom Axiom (Affymetrix). As both arrays are based on very similar principles regarding linkage disequilibrium thresholds for genome-wide coverage, have a similar number of markers and are identical regarding about 30\% of the SNPs, this technological difference does not seem to account for non-reproducibility of the original publication.

SNP imputation quality was nearly perfect in 76 SNPs, whereas 43 SNPs did not reach the highest quality threshold. However, as described in the Materials and Methods section, the overall mean quality score was >0.9. Nevertheless, with a total of 142/185 SNPs, at least an approximation of the PWM results should have been reached.

The scanner in which MRI sequences were acquired is different in the original study (1.5 \(T\)) compared with the present work (3 \(T\)). In fact, 3 \(T\) scanners increase baseline magnetization, leading to an about twofold increase in signal-to-noise ratio, which, in turn, improves accuracy and reproducibility of tissue classification results and thus sensitivity of volumetry regarding morphometric differences.\(^4\) Therefore, also this difference should not be critical for our failure to replicate the previous results. Moreover, the deep methodological (software) differences between the two segmentation processes used in the original\(^7\) versus our study might account for the mismatch between the two samples. In this regard, it has to be pointed out that the method used here is nowadays the gold standard in brain segmentation, whereas the method employed by Terwisscha van Scheltinga et al.\(^7\) is based on older software.

Taken together, even with the limitations of our replication sample discussed above (and always bearing in mind the possibility of a winner’s curse effect in the original study), at least a ‘signal in the right direction’ would have to be expected if the PSS or PWM associations with brain dimensions were of general validity. In light of the obvious interest of studying effects of polygenic risk scores on specific subphenotypes of relevance for complex psychiatric disorders, our results admonish that replication studies are absolutely essential for this kind of analyses. This all the more as polygenic settings of interest cannot easily be explored in animal models to confirm their specific importance. It remains to be established whether by including even larger numbers of individuals in case–control GWAS, the heterogeneity problem of SCZ or other mental diseases (and of health) will be solved. Although much more labor intense than GWAS, large-scale phenotype-based genetic association studies will be pivotal for further investigating the genotype contribution to complex disease phenotypes, thereby extending and complementing the GWAS efforts.

CONFLICT OF INTEREST

The authors declare no conflict of interest.

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