

# **The Genetic and Environmental Determinants of the Association Between Brain Abnormalities and Schizophrenia: The Schizophrenia Twins and Relatives Consortium**

## ***Supplemental Information***

### **Recruitment**

The Dutch sample contributed three cohorts. The discordant twin sample (MZ and DZ) have been described previously (1,2). The control twins (1,3,4) were recruited from the twin sample of the Department of Psychiatry at the University Medical Center Utrecht, the Netherlands and from the population based Netherlands Twin Register (5). Healthy control twins from the University Medical Center Utrecht bipolar twin study were also added (6). The Finnish twins were drawn from a twin cohort comprised of all of the same-sex twins born in Finland from 1940 through 1957 in which both members of each pair were alive and residing in Finland as of 1967 (7). In the United Kingdom, probands were referred to the study from across the country by their consulting psychiatrists. Control twins were recruited from the Institute of Psychiatry Volunteer Twin Register and by national media advertisements. The German twin sample was collected as part of the Heidelberg-Jena twin study on schizophrenia. Twins from all sites, except Finland, were recruited through advertisements and in collaboration with treating hospitals from the surrounding areas.

### **Assessment**

All subjects underwent extensive psychiatric evaluation using available hospital records and structured psychiatric interviews (Utrecht: the Comprehensive Assessment of Symptoms and History interview (8), the Schedule for Affective Disorders and Schizophrenia: Lifetime Version (9), the Structured Interviews for DSM-IV (10), the Family Interview for Genetic Studies (11), and a medical history inventory; Helsinki: Structured Clinical Interview for DSM-III-R Disorders (SCID), Patient or Nonpatient Edition respectively (12) and Structured Clinical Interview for DSM-III-R Personality Disorders (13); London:

Schedule for Affective Disorders and Schizophrenia—Lifetime Version (9); Jena: Schedules for Clinical Assessment in Neuropsychiatry (14) and Family History Research Diagnostic Criteria (15)). At all sites psychiatric diagnosis was established according to DSM-IV criteria. The diagnostic interviews were conducted by trained and experienced psychologists and psychiatrists. In the case of doubt consensus was reached with a senior psychiatrist. For this study we excluded individuals with schizoaffective disorder. Exclusion criteria for all were the presence of significant medical or neurological illnesses including migraine, epilepsy, hypertension, cardiac disease, diabetes mellitus, endocrine disorders, cerebrovascular disease, alcohol or other drug dependence, significant past head trauma, or IQ below 80. Zygosity was based on DNA polymorphisms. Control twins were excluded in case of an axis I and/or schizophrenia-spectrum diagnosis.

## **Consent**

The studies were approved by their respective ethics committees. All subjects gave written informed consent to participate in the study after a full explanation of the study aims and procedures.

**Table S1.** Scanner type and acquisition protocol used at each study site

<b>Site</b>	<b>Scanner</b>	<b>Protocol/orientation</b>	<b>Voxel dimensions (mm)</b>	<b>TE (msec)</b>	<b>TR (msec)</b>	<b>Flip angle</b>
Utrecht	Philips NT 1.5T	3D-FFE/coronal	1x1x1.2	4.6	30	30°
London Maudsley	GE Signa 1.5T	3D-SPGR/coronal	0.781x0.781x1.5	5	35	35°
London St Georges	GE Signa 1.5T	3D-SPGR/coronal	0.781x0.781x1.5	5	35	35°
Jena	Philips ACS II 1.5T	3D-FFE/sagittal	1x1x1	5	13	25°
Helsinki	Siemens Magnetom Impact 1.0T	MPRAGE/sagittal	1x1x1.2	4.4	11.4	12°

**Table S2.** Standardized estimates (with 95% confidence interval) of the bivariate AE genetic models for schizophrenia and relevant MRI brain volume.

	$h^2_{BV}$	$e^2_{BV}$	$r_g$	$r_e$	$r_{ph}$
Cerebrum	<b>.76</b> (.69 / .82)	<b>.24</b> (.18 / .31)	<b>-.21</b> (-.31 / -.10)	<b>-.37</b> (-.57 / -.14)	<b>-.22</b> (-.30 / -.14)
Cerebral white	<b>.73</b> (.65 / .80)	<b>.27</b> (.20 / .35)	<b>-.20</b> (-.31 / -.09)	-.07 (-.30 / .16)	<b>-.17</b> (-.25 / -.09)
Lateral ventricles	<b>.73</b> (.64 / .80)	<b>.27</b> (.20 / .36)	.05 (-.08 / .19)	<b>.42</b> (.14 / .65)	.10 (.00 / .20)
3 <sup>rd</sup> ventricle	<b>.76</b> (.67 / .82)	<b>.25</b> (.18 / .33)	<b>.20</b> (.07 / .33)	<b>.22</b> (-.08 / .49)	<b>.18</b> (.08 / .28)
Temporal cortical gray	<b>.55</b> (.41 / .66)	<b>.45</b> (.34 / .59)	.03 (-.12 / .19)	<b>-.34</b> (-.58 / -.08)	-.04 (-.01 / .06)
Parietal cortical gray	<b>.68</b> (.57 / .76)	<b>.32</b> (.24 / .44)	.09 (-.05 / .23)	<b>-.25</b> (-.50 / .01)	.03 (-.07 / .13)

$h^2$ ,  $e^2$  = standardized additive genetic and non-shared environmental variance components;  $r_g$ ,  $r_e$  = genetic, and non-shared environmental correlation;  $r_{ph}$  = total phenotypic correlation. Confidence intervals not including zero indicate significance (bold). Fixed (genetic) model for schizophrenia used:  $h^2 = .81$ ,  $c^2 = .11$ ,  $e^2 = .08$  and prevalence of 1%. A = additive genetic effects; E = unique environmental effects, including measurement error.

## Supplemental References

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