## Association between stressful life events and psychotic experiences in adolescence: evidence for gene–environment correlations

Sania Shakoor, Helena M. S. Zavos, Claire M. A. Haworth, Phillip McGuire, Alastair G. Cardno, Daniel Freeman and Angelica Ronald

#### Background

Stressful life events (SLEs) are associated with psychotic experiences. SLEs might act as an environmental risk factor, but may also share a genetic propensity with psychotic experiences.

#### Aims

To estimate the extent to which genetic and environmental factors influence the relationship between SLEs and psychotic experiences.

#### Method

Self- and parent reports from a community-based twin sample (4830 16-year-old pairs) were analysed using structural equation model fitting.

#### Results

SLEs correlated with positive psychotic experiences (r = 0.12-0.14, all P < 0.001). Modest heritability was shown for psychotic experiences (25–57%) and dependent SLEs (32%). Genetic influences explained the majority of the

modest covariation between dependent SLEs and paranoia and cognitive disorganisation (bivariate heritabilities 74–86%). The relationship between SLEs and hallucinations and grandiosity was explained by both genetic and common environmental effects.

#### Conclusions

Further to dependent SLEs being an environmental risk factor, individuals may have an underlying genetic propensity increasing their risk of dependent SLEs and positive psychotic experiences.

#### Declaration of interest None.

#### **Copyright and usage**

© The Royal College of Psychiatrists 2016. This is an open access article distributed under the terms of the Creative Commons Attribution (CC BY) licence.

Studies investigating the aetiology of adolescent psychotic experiences report modest heritability estimates ranging between 33 and 58%, with the remaining variances attributable to environmental influences.<sup>1-3</sup> Population-based studies of children and adolescents have found that stress-provoking life experiences such as trauma and victimisation are predictive of psychotic experiences.<sup>4</sup> It is thus reasonable to hypothesise that the same may be true for other stressful life events (SLEs). SLEs are defined as events that require individuals to readjust or experience a change in life.<sup>5</sup> Literature on SLEs has made a distinction between dependent life events which are typically reliant on an individual's behaviour (such as breaking up with a boy/girlfriend), and independent life events where an individual usually has no control on the occurrence of the event (such as death of a friend or relative).6 The relationship between SLEs and psychotic experiences has been explored within the adult population,<sup>7</sup> with estimates of a fourfold increased risk of psychotic experiences among adults who experienced two SLEs and a sixfold increased risk of psychotic experiences among adults who reported six or more SLEs.8 Less, however, is known about the relationship between SLEs and psychotic experiences in adolescents. In one study, researchers found that young adolescents who had more than three SLEs were more likely to experience psychotic experiences.9 In another, researchers found that over a 3-yearperiod, adolescents with a larger number of SLEs had the highest risk of persistent auditory hallucinations.<sup>10</sup> These observations support the notion that SLEs in general, as well as trauma and victimisation, also contribute towards their risk of psychotic experiences. SLEs are often considered as an index of 'environmental risk', yet their heritability has been estimated on

average as 28%,11 31% for 'dependent' SLEs and 17% for 'independent' SLEs.<sup>11</sup> Since dependent SLEs are more influenced by an individual's behaviour than independent SLEs, they may share a genetic propensity with other heritable behaviours such as psychotic experiences. It is thus feasible to suggest that SLEs are not solely an environmental risk factor for psychotic experiences, but rather that SLEs and psychotic experiences co-occur because of a shared genetic propensity. This possibility needs investigation because the implications for clinical prevention and intervention strategies differ depending on the degree to which the association is driven by genes and the environment. For example if SLEs co-occur with psychotic experiences because of underlying shared genetic influences,<sup>11</sup> this would indicate the need for future research prevention and intervention strategies to investigate other heritable correlates of psychotic experiences and SLEs such as underlying personality traits.<sup>12–14</sup>

The heritability of 'environmental' factors such as SLEs is indicative of gene–environment correlation, whereby genetic factors may in part influence an individual's exposure to specific environments which in turn results in the environmental factors themselves being partly heritable (a gene–environment correlation (rGE)).<sup>15</sup> Assuming an absence of rGE by investigating 'environmental' risk factors outside of the context of genetic influences may provide a biased estimation of the magnitude of effect an environmental factor has on traits such as psychotic experiences. The investigation of rGE contributes to our understanding of environmental risk factors by showing that experiences are in part a result of genetic influences,<sup>16</sup> thus targeting environmental risk factors alone may not be beneficial. In addition to exploring environmental effects directly, this research area demonstrates that focusing attention on underlying pathways through which genetic propensities influence behaviours and traits will also be fruitful. Although to our knowledge no other studies have investigated the genetic and environmental overlap between SLEs and psychotic experiences among adolescents, there is some evidence to suggest that there is a modest degree of genetic overlap between SLEs and depression.<sup>17</sup> The considerable comorbidity between psychotic experiences and depression<sup>18</sup> lends support to the hypothesis that some degree of genetic overlap will also be observed between SLEs and psychotic experiences. This is the first study to utilise data from an adolescent twin sample to investigate the genetic and environmental influences contributing to the associations between SLEs and psychotic experiences, as well as the first to assess SLEs in relation to dimensional scales of self-reported psychotic experiences. Our aims were twofold, first to examine whether dependent and independent SLEs are associated with specific psychotic experiences in adolescence, and second, to estimate the extent to which genetic and environmental factors influence the association between dependent SLEs and psychotic experiences.

#### Method

#### Participants

The Longitudinal Experiences And Perceptions (LEAP) study<sup>19</sup> is part of the Twins Early Development Study (TEDS) which comprises a community sample of monozygotic (MZ) and dizygotic (DZ) twins born in England and Wales between 1994 and 1996.20 10 874 families from TEDS were invited to take part in the LEAP study. Parent reports for 5076 (46.7%) families and twin reports for 5059 (46.5%) pairs were obtained. Adolescents who participated in the LEAP project had a mean age of 16.32 years. Individuals were excluded (n = 327 families) if they did not provide consent at first contact (when TEDS was started), if they had a severe medical disorder, had experienced severe perinatal complications or if their zygosity was unknown. Exclusions for medical disorders included individuals with cystic fibrosis, cerebral palsy, fragile-X syndrome, autism spectrum disorder and those with chromosomal abnormalities such as Down syndrome. After exclusions, the sample reported on in this study comprised 4830 families (44.84% male, 35.94% MZ twin pairs). Comparing the participating and non-participating samples, 94% v. 91% were White respectively, and 16% v. 12% had mothers with one or more A-levels (UK advanced educational qualification) as their highest qualification respectively.

#### Measures

#### SLES

We assessed SLEs using 20 items from the Coddington Life Events Record.<sup>5</sup> Parents and adolescents were asked to report on SLEs that had occurred in the past year, by responding 'Yes' (1) or 'No' (0) to items such as 'death of a close friend or relative'. Parent and adolescent reports were combined to capture all occurrences of SLEs. This was done using an either/or approach, as simple combination rules work as well, if not better than, more complicated ones.<sup>21,22</sup> An SLE was scored as 'Yes' (1) if either adolescent or parent had reported it. In line with the literature on SLEs,<sup>6,23,24</sup> a distinction was made between dependent and independent life events. The dependent SLEs scale was the sum of ten items that assessed life events that occur or are potentially likely to arise as a consequence of one's behaviour (i.e. breaking up

with a boy/girlfriend). The independent SLEs scale was the sum of ten items that assessed life events that occur or are likely to arise independent of one's behaviour (i.e. death of a friend or relative).

#### Psychotic experiences

Psychotic experiences were assessed using the Specific Psychotic Experiences Questionnaire (SPEQ).<sup>19</sup> SPEQ assesses specific psychotic experiences as quantitative traits and includes five selfreport subscales: paranoia (15 items), hallucinations (9 items), cognitive disorganisation (11 items), grandiosity (8 items), anhedonia (10 items) and one parent-rated subscale: parent-rated negative symptoms (10 items). SPEQ items were derived for the most part from existing scales that were adapted in order for them to be suitable for adolescents.<sup>19</sup> The subscales were derived from principal component analysis and show good-to-excellent internal consistency (r=0.77-0.93) and test-retest reliability across a 9-month interval (r = 0.65 - 0.74) in this sample. In terms of validity, expert clinical opinion was obtained on the suitability of each item as a measure of adolescent psychotic experiences to ensure content validity.<sup>19</sup> Furthermore, levels of agreement between scores on SPEQ and the PLIKS (a known measure of psychosis-like symptoms)<sup>25</sup> showed that adolescents who reported 'definitely' having any psychosis-like symptoms on the PLIKS had significantly more psychotic experiences on all the SPEQ subscales (with the exception of anhedonia) when compared with those who did not report any definite psychosis-like symptoms (all significant at P<0.001). Positive and cognitive subscales of psychotic experiences showed significant positive correlations with the PLIKS quantitative score (hallucinations r = 0.60, paranoia r = 0.48, cognitive disorganisation r = 0.41, grandiosity r = 0.27, all P < 0.001).<sup>19,25</sup> Furthermore, for paranoia, cognitive disorganisation, grandiosity and parent-rated negative symptoms SPEQ subscales, individuals who reported a family history of psychosis, as measured by having a first- or second-degree relative with schizophrenia or bipolar disorder, scored higher than individuals without a family history of psychosis (all P < 0.05). Further information on the measure can be found in Ronald et al.<sup>19</sup>

#### **Statistical analyses**

All analyses were performed using Stata 12 and Open MX. Open MX uses the method of maximum-likelihood estimation and is widely used for analysing genetically sensitive data.<sup>26</sup> In line with standard behavioural genetics procedure, the effects of gender and age were regressed out, and analyses were conducted using standardised residuals.<sup>27</sup> Scales of SLEs and psychotic experiences were transformed using square root transformation techniques to reduce skewness and kurtosis and to ensure that the assumption of having a normal distribution was met for genetic modelling (online Table DS1).

#### The twin design

The twin design involves MZ and DZ twin pairs to determine the extent to which variation in a single phenotype or covariation between phenotypes are attributable to genetic and environmental influences. Within-pair similarities for MZ and DZ twin pairs were examined separately to establish the role of genetic and environmental influences based on the notion that: (a) MZ twin pairs share 100% of their segregating DNA code and DZ twin pairs share on average 50%; (b) MZ and DZ twin pairs share environmental factors common to both twins in the same family ('common environment'); and (c) exposure to environmental factors that are experienced differently or are specific to the

individual ('unique environment') contribute towards differences within twin pairs.  $^{\rm 16}$ 

#### Twin analyses

Structural equation modelling techniques were employed to establish the relative importance of additive genetic (A), common environment (C) and unique environmental influences (E) contributing to a phenotype.<sup>16</sup> This technique further extends to bivariate analyses, by exploring the covariation between phenotypes. The relative contributions of genetic and environmental factors to the association between SLEs and psychotic experiences are referred to as bivariate heritability (biva<sup>2</sup>), bivariate common environment (bivc<sup>2</sup>) and bivariate unique environment (bive<sup>2</sup>). Estimates of covariance between SLEs and psychotic experiences were also used to calculate genetic correlations  $(r_a)$ , common environment correlations  $(r_c)$  and unique environment correlations  $(r_e)$ , which indexed the extent to which the same set of genes or environments influence both phenotypes.<sup>28</sup> The relative fit of different models were compared with a saturated model (which provides a full description of the data) to establish the best fitting model for the data.<sup>29</sup> Parameter estimates were then calculated with confidence intervals using the maximum-likelihood method. The best fitting models were selected based on the lowest Akaike's information criterion (AIC) values. In instances where the AIC values were similar across models (i.e. ACE dropped  $r_{a \text{ and}}$  ACE dropped  $r_{c}$ ), resulting in the relative influences being difficult to distinguish, the full ACE model was chosen as being the most parsimonious.

#### rGE

Further to distinguishing genetic and environmental influences contributing to phenotypic variances and covariances, the twin design also allows for the investigation of *rGE*. Univariate twin models were used to test whether genetic factors influence an 'environmental' measure such as SLEs. A genetic influence on an environmental measure would be indicative of *rGE*. Bivariate twin models were also used. Findings suggested *rGE* if genetic factors mediated the association between environmental measures (for example SLEs) and traits (for example psychotic experiences).<sup>16</sup>

#### Results

#### **Phenotypic analyses**

Analyses of variance illustrated significant mean effects of gender on psychotic experiences (Table 1). Females reported higher levels of paranoia, hallucinations and cognitive disorganisation, in contrast to males who reported higher levels of grandiosity, anhedonia and had more parent-rated negative symptoms. Females also reported more dependent SLEs than males. No main effect for gender was present for independent SLEs. A main effect for zygosity was observed for paranoia, hallucinations, cognitive disorganisation and parent-rated negative symptoms, whereby DZ twins reported higher levels in comparison to MZ twins. However, the combined effect of gender and zygosity on the means was small ( $R^2 = 0.00-0.06$ ).

Phenotypic correlations between SLEs and psychotic experiences are presented in Table 2. Dependent and independent SLEs in adolescence were modestly associated with increased levels of positive psychotic experiences: paranoia, hallucinations, cognitive disorganisation and grandiosity (r=0.12–0.14, all P<0.001). Correlations with negative psychotic experiences were low for dependent SLEs (anhedonia r=-0.04, P<0.05, parent-rated

Table 1 Means, standard deviations and analysis of variance by gender	viations and an	alysis of variand		and zygosity for psychotic experiences and stressful life events (SLEs)	ychotic experi	ences and stre	essful life eve	nts (SLES)				
			Mean (s.d.)									
Psychotic experiences	Total	Male	Female	Monozygotic twins	Dizygotic twins	Score, range Cronbach, $\boldsymbol{\alpha}$		Gender, P	Zygosity, P	ANOVA Gender × zygosity	$R^{2}$	и
Paranoia	12.17 (10.62)	11.75 (10.42)	12.50 (10.77)	11.79 (10.46)	12.38 (10.70)	0-71	0.93	< 0.01	0.01	0.45	0.00	4777
Hallucinations	4.65 (6.00)	4.30 (5.77)	4.94 (6.16)	4.47 (5.91)	4.76 (6.05)	0-45	0.87	< 0.01	0.01	0.53	0.01	4785
Cognitive disorganisation	3.96 (2.85)	3.40 (2.72)	4.41 (2.87)	3.86 (2.82)	4.01 (2.86)	0-11	0.73	< 0.01	0.01	0.66	0.03	4778
Grandiosity	5.32 (4.42)	5.82 (4.56)	4.91 (4.27)	5.26 (4.35)	5.35 (4.46)	0-24	0.85	< 0.01	0.56	0.96	0.01	4781
Anhedonia	17.33 (7.93)	19.50 (7.98)	15.58 (7.44)	17.07 (7.96)	17.48 (7.91)	0-50	0.78	< 0.01	0.44	0.85	0.06	4781
Parent-rated negative symptoms	2.81 (3.89)	3.17 (4.10)	2.52 (3.69)	2.64 (3.57)	2.91 (4.06)	0-30	0.85	< 0.01	0.03	0.02	0.01	4792
Dependent SLEs	1.68 (1.22)	1.66 (1.29)	1.70 (1.16)	1.63 (1.18)	1.71 (1.25)	0-10	0.41	0.01	0.06	0.16	0.00	4782
Independent SLEs	1.58 (1.40)	1.58 (1.42)	1.57 (1.38)	1.53 (1.35)	1.61 (1.43)	0-10	0.42	0.58	0.06	0.02	0.00	4784
a. Means and standard deviations reported prior to transformation, analyses of variances were performed using one random member of each twin pair. Gender P-value associated with the effect of gender on the means; zygosity P-value associated with the of zygosity P-value associated with the effect of gender and zygosity P-value associated with the effect of gender and zygosity P-value associated with the effect of gender and zygosity P-value associated with the effect of gender and zygosity P-value associated with the effect of gender and zygosity P-value associated with the effect of gender and zygosity P-value associated with the effects of the interaction between gender and zygosity on the means and R <sup>2</sup> is the proportion of the total variance explained by gender and zygosity, <i>n</i> is the number of one random selected individual from each twin pair.	ed prior to transform. sity P-value associate	lation, analyses of var	iances were performe the interaction betwee	ed using one random r en gender and zygosity	member of each twi y on the means and	n pair. Gender <i>P</i> -va. $R^2$ is the proportion	lue associated wit n of the total varia	th the effect of ance explained	gender on the m by gender and zy	ormed using one random member of each twin pair. Gender <i>P</i> -value associated with the effect of gender on the means; zygosity <i>P</i> -value associated with the effect stween gender and zygosity on the means and $R^2$ is the proportion of the total variance explained by gender and zygosity; <i>n</i> is the number of one randomly	sociated with of one randor	the effect nly

		Stressfi	Il life events	
	Dependent stressful life	e events	Independent stressful	life events
Psychotic experiences	r (95% CI)	п	r (95 Cl)	п
Paranoia	0.14 (0.11 to 0.17)	4732	0.09 (0.06 to 0.12)	4734
Hallucinations	0.14 (0.11 to 0.16)	4740	0.12 (0.09 to 0.15)	4742
Cognitive disorganisation	0.14 (0.11 to 0.16)	4733	0.10 (0.07 to 0.13)	4735
Grandiosity	0.12 (0.10 to 0.15)	4736	0.06 (0.04 to 0.09)	4738
Anhedonia	-0.04 (-0.06 to -0.01)	4736	-0.03 (-0.06 to 0.00)	4738
Parent-rated negative symptoms	0.04 (0.01 to 0.06)	4773	0.08 (0.05 to 0.11)	4775

negative symptoms r = 0.04, P < 0.05) and independent SLEs (anhedonia r = -0.03, P < 0.10).

The prevalence of SLE and mean scores on specific psychotic experiences scales for individuals with each type of SLE are reported in online Tables DS2–8. For example, the largest effect sizes for paranoia (Cohen's d=0.48) and anhedonia (Cohen's d=0.34) were observed among adolescents who experienced the SLE 'becoming involved in drugs'. Those who reported 'being responsible for a road accident' had the largest effect size for cognitive disorganisation (Cohen's d=0.60). Adolescents who experienced 'suspension from school/college' had the largest effect size for hallucinations (Cohen's d=0.36), grandiosity (Cohen's d=0.28) and parent-rated negative symptoms (Cohen's d=0.50).

We did not perform behaviour genetic twin analyses on the independent SLEs measure because these events were family-wide and experienced by both twins within a twin pair. It was therefore not possible to partition variance into genetic and environmental influences. Behaviour genetic analysis of anhedonia and parent-rated negative symptoms with dependent SLEs were not assessed, as phenotypic correlations were considered to be too small (r = -0.04 and r = 0.04 respectively) to be decomposed into genetic and environmental influences.

#### Behaviour genetic analyses

For both psychotic experiences and SLEs, univariate twin correlations (Table 3) were indicative of genetic influences (A), because MZ correlations were consistently larger than DZ correlations. As the DZ correlations were greater than half of MZ correlations, this suggested some common environmental (C) influence. Furthermore, as MZ correlations were less than unity, this implied a moderate unique environmental effect (E).

Univariate model fitting analyses confirmed initial observations from the twin correlations by showing that genetic (A: 0.25–0.57) and unique environmental (E: 0.17–0.57) factors contributed the most to variances observed in psychotic experiences and dependent SLEs (online Table DS9). All univariate ACE models did not provide a significantly worse fit compared with the saturated models. C could be dropped from the models for paranoia, cognitive disorganisation and anhedonia, and explained small amounts of the variance (0.11–0.26) for the remaining scales.

Bivariate cross-twin cross-trait correlations (Table 3) provided an insight into the extent to which the covariance between dependent SLEs and psychotic experiences was explained by genetic and environmental influences. Collectively, MZ cross-twin cross-trait correlations were larger than DZ cross-twin cross-trait correlations, which is indicative of a genetic influence on the phenotypic associations between SLEs and psychotic experiences. DZ cross-twin cross-trait correlations were somewhat greater than half of MZ cross-twin cross-trait correlations thus implying a modest common environmental effect. Where MZ cross-twin cross-trait correlations were less than the phenotypic correlations between SLEs and psychotic experiences, correlations were suggestive of a unique environmental influence on the covariation.

Results from the bivariate correlated factors solution (online Table DS10) showed that for the association between dependent SLEs and paranoia and cognitive disorganisation scales, the ACE correlated factors solution with dropped  $r_{\rm c}$  fitted the data best based on the AIC fit index. Analyses (Table 4) demonstrated that the relationship between dependent SLEs and paranoia was almost completely explained by genetic influences ( $biva^2 = 0.86$ ), with the remaining covariance explained by unique environment. Genetic correlation indicated that a moderate degree of genetic influences overlapped between the two phenotypes ( $r_a = 0.33$ ). Furthermore, a small proportion of unique environmental overlap between dependent SLEs and paranoia was also found ( $r_e = 0.04$ ). Analyses investigating the association between dependent SLEs and cognitive disorganisation showed a similar pattern whereby high bivariate heritability was found ( $biva^2 = 0.74$ ). The remaining covariance was explained by unique environment ( $bive^2 = 0.26$ ). Genetic and unique environment correlations showed that there was modest genetic  $(r_a = 0.21)$  and unique environmental  $(r_e = 0.05)$  overlap between dependent SLEs and cognitive disorganisation.

Bivariate analyses further showed that for the association between dependent SLEs and hallucinations and SLEs and grandiosity, the ACE correlated factors solution fitted the data best

Table 3 Univariate twin and	cross-trait cross-t	win correlations <sup>a</sup>
	ICC (9	5% CI)
	Monozygotic twins	Dizygotic twins
Univariate twin correlations, psychotic experiences		
Paranoia Hallucinations Cognitive disorganisation	0.52 (0.49 to 0.56) 0.43 (0.39 to 0.47) 0.45 (0.41 to 0.48)	0.29 (0.24 to 0.34) 0.31 (0.26 to 0.35) 0.23 (0.18 to 0.28)
Grandiosity Dependent SLEs	0.48 (0.44 to 0.52) 0.52 (0.48 to 0.55)	0.28 (0.23 to 0.32) 0.34 (0.30 to 0.39)
Cross-trait cross-twin correlation, psychotic experiences and dependent SLEs		
Paranoia Hallucinations Cognitive disorganisation Grandiosity	0.13 (0.08 to 0.17) 0.06 (0.02 to 0.11) 0.07 (0.03 to 0.12) 0.13 (0.06 to 0.15)	
SLEs, stressful life events. a. Correlations were performed using o Intraclass correlations (ICC) using transf scales.		

			Dependent stre	ssful life events		
	<i>biva</i> <sup>2</sup> (95% Cl)	<i>bivc</i> <sup>2</sup> (95% CI)	<i>bive</i> <sup>2</sup> (95% CI)	r <sub>a</sub> (95% Cl)	r <sub>c</sub> (95% Cl)	<i>r<sub>e</sub></i> (95% Cl)
Paranoia	0.86 (0.72 to 1.00)	-	0.14 (-0.01 to 0.30)	0.33 (0.24 to 0.45)	-	0.04 (0.01 to 0.09)
Hallucinations	0.44 (-0.22 to 1.00)	0.39 (-0.18 to 0.96)	0.17 (-0.04 to 0.37)	0.18 (-0.09 to 0.46)	0.25 (-0.12 to 0.67)	0.04 (-0.01 to 0.08
Cognitive disorganisation	0.74 (0.52 to 0.94)	-	0.26 (0.06 to 0.48)	0.21 (0.13 to 0.31)	-	0.05 (0.01 to 0.10)
Grandiosity	0.42 (-0.09 to 0.94)	0.35 (-0.09 to 0.79)	0.23 (0.08 to 0.38)	0.19 (-0.04 to 0.42)	0.38 (-0.12 to 1.00)	0.07 (0.02 to 0.11)

and unique environment respectively. Bivariate genetic ( $r_a$ ), common environment ( $r_c$ ) and unique environment ( $r_c$ ) correlations indicate the genetic and environment average between psychotic symptoms and stressful life events.

(online Table DS10). Both genetic and common environmental influences appeared to explain part of the covariance between dependent SLEs and hallucinations, and SLEs and grandiosity (as indicated by the *biva*<sup>2</sup> and *bivc*<sup>2</sup> values in Table 4), and both genetic and common environmental influences had some overlapping influences across SLEs and these psychotic experiences (as indicated by the  $r_a$  and  $r_c$  values) but notably the confidence intervals all overlapped with zero. This meant it was not possible to differentiate the relative role of genetic and common environmental influences on the covariance, suggesting they may both play a role. The association between dependent SLEs and grandiosity was also influenced by a modest degree of unique environmental effects (*bive*<sup>2</sup> = 0.23, Table 4).

#### Discussion

Using a community sample of 16-year-old twins, this study showed that SLEs were correlated with positive psychotic experiences (paranoia, hallucinations, cognitive disorganisation, grandiosity) and weakly correlated with negative psychotic experiences. Shared genetic influences explained a substantial proportion of the covariation between paranoia, cognitive disorganisation and dependent SLEs. For hallucinations, and grandiosity, both genes and environment explained some of the covariation with SLEs.

### Are stressful life events associated with psychotic experiences in adolescence?

In our sample of adolescents, females reported more positive psychotic experiences (with the exception of grandiosity) and males reported more grandiosity, anhedonia and had more parent-rated negative symptoms. These findings are similar to those from other cohort-based studies,<sup>30</sup> and suggest that there may be continuity in gender differences in psychotic experiences among the general population and those with schizophrenia, where males report severer negative symptoms than females.<sup>3</sup> In keeping with previous studies,<sup>9,10</sup> having an increased number of dependent and independent SLEs was associated with higher levels of psychotic experiences. This association was stronger for positive (paranoia, hallucinations, cognitive disorganisation, grandiosity) than negative psychotic experiences. Among SLEs, 'becoming involved in drugs', 'suspension from school/college' and 'being responsible for a road accident' were associated with the highest levels of positive psychotic experiences. This specificity of life events is of interest as it is consistent with the association between substance use and psychotic experiences among adolescents.<sup>32</sup> It also highlights that other correlates such as 'suspension from school' may also be of relevance for understanding positive psychotic experiences in adolescence. Collectively, the modest associations reported in this study show that not all

adolescents who experience SLEs have psychotic experiences, and vice versa. Experiencing a number of SLEs or specific SLEs such as 'becoming involved in drugs' may therefore be a trigger for having elevated levels of positive psychotic experiences.

The association between SLEs and psychotic experiences is consistent with cognitive psychological theories of the development of psychotic experiences,<sup>33</sup> which suggests that exposure to 'triggering events' are particularly damaging in individuals predisposed to disruptions in their cognitive processes. This disruption in cognitive processes in turn may contribute to the risk for psychotic experiences. For example, experiencing an increased number of SLEs may lead individuals to develop cognitive biases that result in viewing their environment to be hostile and threatening. This feeling that 'the world is out to get me' may trigger psychotic experiences such as paranoia. Our results inform these models by showing that part of the explanation for individuals having SLEs that co-occur with psychotic experiences is an underlying genetic propensity for both SLEs and psychotic experiences. As we could not examine the temporal relationship between SLEs and psychotic experiences in the present study, it is also possible that adolescents with psychotic experiences may be more likely to have SLEs. For example, experiencing paranoia may lead to being suspicious of others and result in SLEs such as breaking up with a boyfriend or girlfriend. However, evidence from a number of studies has shown life stress (i.e. SLEs) to be a risk factor for psychotic experiences and psychosis among adults and adolescents,<sup>8,10,34,35</sup> thus supporting the role of SLEs as a catalyst for psychotic experiences such as paranoia.

## To what extent do genetic and environmental factors influence the associations between SLEs and psychotic experiences?

In line with previous research among adolescents within the general population, dependent SLEs and psychotic experiences were in part heritable,<sup>1,2,11,36</sup> with the remaining variance largely attributable to unique environmental factors. Our findings extend those of previous studies by showing that the relationship between dependent SLEs and psychotic experiences (paranoia and cognitive disorganisation) was almost completely explained by genetic influences. Our findings also provide support for the concept of gene-environment correlation. This can be in one of three ways: active, evocative or passive.11 Gene-environment correlations could be 'active', whereby the genetic propensity that leads individuals to seek out situations resulting in dependent SLEs is the same genetic influence that increases the risk for psychotic experiences (paranoia and cognitive disorganisation). Alternatively, it could be 'evocative', whereby dependent SLEs, which are partly genetically influenced, result in environments or incite behaviours from others that result in elevated levels of

paranoia and cognitive disorganisation. Finally, gene–environment correlations may be 'passive', whereby genetic factors that increase the likelihood of dependent SLEs on the part of the parent are shared with adolescents through the environments parents raise them in, and in turn are associated with psychotic experiences. Focusing on 'environmental' factors in isolation may not therefore be an optimal research strategy. Examining factors through which a genetic vulnerability for having dependent SLEs and psychotic experiences are translating into behaviours (such as home environment or parenting), may help in identifying underlying mechanisms contributing towards psychotic experiences and SLEs among adolescents. For the relationship between dependent SLEs and hallucinations, and grandiosity, both genetic and common environmental influences appeared to play a role but their relative role was not clear.

#### Limitations and strengths

The study's cross-sectional design did not make it possible to test for temporal priority. Therefore, although interpretations were in the direction of SLEs leading to psychotic experiences, it is possible that psychotic experiences may have altered individuals' behaviours resulting in the SLEs being reported here. Furthermore, as participants were asked to report on their psychotic experiences from the past month and SLEs from the past 12 months, there may be recall bias, whereby SLEs were more difficult to remember given that the reporting period was more distal. Second, we used self-reports of paranoia, hallucinations, cognitive disorganisation, grandiosity and anhedonia. This work could be replicated using in-depth interviews and reports from other informants. Third, we observed modest correlations between SLEs and psychotic experiences. Estimates of bivariate heritability and environmental influences reported in this study are therefore explaining a small proportion of variance within psychotic experiences.

This study also has a number of strengths. It is the first to investigate psychotic experiences and SLE among a large community sample of adolescents, an age just prior to the modal age at onset of psychotic disorders such as schizophrenia. Furthermore, the genetically informative study design allowed the relationships to be decomposed into genetic and environmental influences. In contrast to other studies that have focused on a specific type of psychotic experience (i.e. hallucinations<sup>37</sup>), this study included multiple informant reports of specific psychotic experiences, which were measured as dimensions and included both positive and negative psychotic experiences. Adolescents reported on paranoia, hallucinations, cognitive disorganisation, grandiosity and anhedonia and parents reported on negative symptoms.

#### Implications

Our work underlines the importance of viewing certain environmental risk factors within the context of genetics. It highlights the importance of not always categorising risk factors as either environmental or genetic as they may be a combination of the two. Our finding of a shared genetic propensity between SLEs and paranoia and cognitive disorganisation could help research and interventions focus on other types of (heritable) behaviours shown developmentally earlier (i.e. impulsivity), which may jointly increase the risk of psychotic experiences and dependent SLEs. Moreover, as DNA does not change throughout the life course, a shared genetic propensity between SLEs and psychotic experiences would imply that clinical intervention should take into account the continued vulnerability of individuals with psychotic experiences to have dependent SLEs. Further research is needed, but the results are suggestive that focusing on and dampening the effects of common environmental risks that contribute towards SLEs might decrease the risk of psychotic experiences such as hallucinations in vulnerable individuals.

SLEs are associated with positive psychotic experiences in adolescence. This is via a shared genetic propensity in addition to the more recognised mechanism of shared environment risk. An accurate understanding of the mechanisms by which risk factors increase the risk for psychotic experiences is imperative for improving intervention and prevention strategies in adolescence.

Sania Shakoor, PhD, Centre for Brain and Cognitive Development, Department of Psychological Sciences, Birkbeck, University of London, London; Helena M. S. Zavos, PhD, King's College London, MRC Social, Genetic & Developmental Psychiatry Centre, Institute of Psychiatry, Psychology & Neuroscience, London; Claire M. A. Haworth, Department of Psychology, University of Warwick, Coventry; Phillip McGuire, PhD, MD, FRCPsych, King's College London, Department of Psychosis Studies, Institute of Psychiatry, Psychology & Neuroscience, London; Alastair G. Cardno, PhD, MMedSc, FRCPsych, MBChB, Academic Unit of Psychiatry and Behavioural Sciences, University of Leeds, Leeds; Daniel Freeman, PhD, DClinPsy, CPsychol, FBPsS, Department of Psychiatry, University of Oxford, Oxford; Angelica Ronald, PhD, Centre for Brain and Cognitive Development, Department of Psychological Sciences, Birkbeck, University of London, London, UK

**Correspondence:** Sania Shakoor, Centre for Brain and Cognitive Development, Department of Psychological Sciences, Birkbeck, University of London, 32 Torrington Square, London WC1E 7JL, UK. Email: sania.shakoor@googlemail.com

First received 16 Oct 2014, final revision 13 Aug 2015, accepted 8 Oct 2015

#### Funding

This project was funded by the Medical Research Council (G1100559 to A.R.; G0901245 and previously G0500079 to Robert Plomin). D.F. was supported by the Medical Research Council (G0902308) and C.M.A.H. by the British Academy.

#### **Acknowledgements**

The authors thank the participants of TEDS for making this research possible. Thank you also to Andrew McMillan, Francesca Lewis, Louise Webster, Neil Harvey and Rachel Ogden, to Peter McGuffin for help planning the study, and to Robert Plomin for the TEDS collaboration.

#### References

- Hur YM, Cherny SS, Sham PC. Heritability of hallucinations in adolescent twins. *Psychiatry Res* 2012; **199**: 98–101.
- 2 Ericson M, Tuvblad C, Raine A, Young-Wolff K, Baker L. Heritability and longitudinal stability of schizotypal traits during adolescence. *Behav Genet* 2011; 41: 499–511.
- **3** Zavos HMS, Freeman D, Haworth CMA, McGuire P, Plomin R, Cardno AG, et al. Consistent etiology of severe, frequent psychotic experiences and milder, less frequent manifestations: a twin study of specific psychotic experiences in adolescence. *JAMA Psychiatry* 2014; **71**: 1049–57.
- 4 Arseneault L, Cannon M, Fisher HL, Caspi A. Childhood trauma and children's emerging psychotic symptoms: a genetically sensitive longitudinal cohort study. Am J Psychiatry 2011; 168: 65–72.
- 5 Coddington RD. The significance of life events as etiological factors in the diseases of children: II: a study of a normal population. J Psychosom Res 1972; 16: 205–13.
- 6 Brown GW, Harris TO. Social Origins of Depression: A Study of Psychiatric Disorder in Women. Tavistock, 1978.
- 7 Johns LC, Cannon M, Singleton N, Murray RM, Farrell M, Brugha T, et al. Prevalence and correlates of self-reported psychotic symptoms in the British population. Br J Psychiatry 2004; 185: 298–305.
- 8 Wiles NJ, Zammit S, Bebbington P, Singleton N, Meltzer H, Lewis G. Self-reported psychotic symptoms in the general population: results from the longitudinal study of the British National Psychiatric Morbidity Survey. Br J Psychiatry 2006; 188: 519–26.
- 9 Bartels-Velthuis AA, van de Willige G, Jenner JA, Wiersma D, van Os J. Auditory hallucinations in childhood: associations with adversity and delusional ideation. *Psychol Med* 2012; 42: 583–93.

- 10 Escher S, Romme M, Buiks A, Delespaul P, Van Os J. Independent course of childhood auditory hallucinations: a sequential 3-year follow-up study. *Br J Psychiatry* 2002; 181: 10–8.
- 11 Kendler KS, Baker JH. Genetic influences on measures of the environment: a systematic review. *Psychol Med* 2007; 37: 615–26.
- 12 Loehlin JC. The differential heritability of personality item clusters. *Behav Genet* 2012; 42: 500– 7.
- 13 Boyette L, Korver-Nieberg N, Verweij K, Meijer C, Dingemans P, Cahn W, et al. Associations between the Five-Factor Model personality traits and psychotic experiences in patients with psychotic disorders, their siblings and controls. *Psychiatry Res* 2013; 210: 491–7.
- 14 Kardum I, Krapic N. Personality traits, stressful life events, and coping styles in early adolescence. *Pers Individ Dif* 2001; 30: 503–15.
- 15 Jaffee SR, Price TS. Gene-environment correlations: a review of the evidence and implications for prevention of mental illness. *Mol Psychiatry* 2007; 12: 432–42.
- 16 Plomin R, Defries J, Knopik VS, Neiderhiser JM. Behavioral Genetics (6th edn). Worth Publishers, 2013.
- 17 Boardman J, Alexander K, Stallings M. Stressful life events and depression among adolescent twin pairs. *Biodemography Soc Biol* 2011; 57: 53–66.
- 18 Varghese D, Scott J, Welham J, Bor W, Najman J, O'Callaghan M, et al. Psychotic-like experiences in major depression and anxiety disorders: a population-based survey in young adults. *Schizophr Bull* 2011; 37: 389–93.
- 19 Ronald A, Sieradzka D, Cardno A, Haworth C, McGuire P, Freeman D. Characterization of psychotic experiences in adolescence using the Specific Psychotic Experiences Questionnaire (SPEQ): findings from a study of 5000 16-year-olds. *Schizophr Bull* 2014; 40: 868–77.
- 20 Haworth CMA, Davis OSP, Plomin R. Twins Early Development Study (TEDS): a genetically sensitive investigation of cognitive and behavioral development from childhood to young adulthood. *Twin Res Hum Genet* 2013; 16: 117–25.
- 21 Bird HR, Gould MS, Staghezza B. Aggregating data from multiple informants in child pychiatry epidemiological research. J Am Acad Child Adolesc Psychol 1992; 31: 78–85.
- 22 Piacentini JC, Cohen P, Cohen J. Combining discrepant diagnostic information from multiple sources: are complex algorithms better than simple ones? J Abnorm Child Psychol 1992; 20: 51–63.
- 23 Silberg J, Rutter M, Neale M, Eaves L. Genetic moderation of environmental risk for depression and anxiety in adolescent girls. *Br J Psychiatry* 2001; 179: 116–21.

- 24 Zavos H, Wong CC, Barclay NL, Keers R, Mill J, Rijsdijk F, et al. Anxiety sensitivity in adolescence and young adulthood: The role of stressful life events, 5HTTLPR. *Depress Anxiety* 2012; 29: 400–8.
- 25 Zammit S, Owen MJ, Evans J, Heron J, Lewis G. Cannabis, COMT and psychotic experiences. *Br J Psychiatry* 2011; **199**: 380–5.
- 26 Boker S, Neale M, Maes H, Wilde M, Spiegel M, Brick T, et al. OpenMx: an open source extended structural equation modeling frame-work. *Psychometrika* 2011; 76: 306–17.
- 27 Eaves L, Eysenck HJ, Martin NG. Genes, Culture, and Personality: An Empirical Approach. Academic Press, 1989.
- 28 Neale MC, Cardon LR. *Methodology for Genetic Studies of Twins and Families*. Springer Netherlands, 1992.
- 29 Rijsdijk FV, Sham PC. Analytic approaches to twin data using structural equation models. *Brief Bioinform* 2002; **3**: 119–33.
- 30 Wigman JTW, Vollebergh WAM, Raaijmakers QAW, ledema J, van Dorsselaer S, Ormel J, et al. The structure of the extended psychosis phenotype in early adolescence: a cross-sample replication. *Schizophr Bull* 2011; 37: 850–60.
- 31 Verdoux H, van Os J. Psychotic symptoms in non-clinical populations and the continuum of psychosis. *Schizophr Res* 2002; 54: 59–65.
- 32 Hides L, Lubman DI, Buckby J, Yuen HP, Cosgrave E, Baker K, et al. The association between early cannabis use and psychotic-like experiences in a community adolescent sample. *Schizophr Res* 2009; 112: 130–5.
- 33 Garety PA, Kuipers E, Fowler D, Freeman D, Bebbington PE. A cognitive model of the positive symptoms of psychosis. *Psychol Med* 2001; 31: 189–95.
- 34 Tessner KD, Mittal V, Walker EF. Longitudinal study of stressful life events and daily stressors among adolescents at high risk for psychotic disorders. *Schizophr Bull* 2011; 37: 432–41.
- **35** Van Winkel R, Stefanis NC, Myin-Germeys I. Psychosocial stress and psychosis. A review of the neurobiological mechanisms and the evidence for gene-stress interaction. *Schizophr Bull* 2008; **34**: 1095–105.
- 36 Polanczyk G, Moffitt TE, Arseneault L, Cannon M, Ambler A, Keefe RSE, et al. Etiological and clinical features of childhood psychotic symptoms. Arch Gen Psychiatry 2010; 67: 328–38.
- 37 Shevlin M, Dorahy M, Adamson G. Childhood traumas and hallucinations: an analysis of the National Comorbidity Survey. J Psychiatr Res 2007; 41: 222–8.



Data supplement to Shakoor et al. Association between stressful life events and psychotic experiences in adolescence: evidence for gene-environment correlations. Br J Psychiatry doi: 10.1192/bjp.bp.114.159079

Table DS1: Distribution parameters for psychotic experiences and stressful life events

	Skew	Kurtosis
Paranoia	0.17	3.07
Hallucinations	0.55	2.74
Cognitive	0.44	2.31
Grandiosity	-0.08	2.79
Anhedonia	-0.48	3.10
Parent-rated negative symptoms	0.56	2.72
Dependent SLEs	-0.58	3.17
Independent SLEs	-0.30	2.25

Note: SLEs= stressful life events.

Table DS2: Prevalence of stressful life events (SLEs)

	Prevalence: In pas	t year
Stressful Life Events	Yes % (N)	No % (N)
Dependent SLEs		
Becoming involved with drugs	5.44 (259)	94.56 (4,499)
Being sent away from home	3.78 (180)	96.22 (4,582)
Breaking up with a boyfriend/girlfriend	26.41 (1,257)	73.59 (3,503)
Suspension from school/college	2.85 (136)	97.15 (4,638)
Failing an important exam	21.09 (1,006)	78.91 (3,764)
Being responsible for a road accident	0.71 (34)	99.29 (4,738)
Getting pregnant or fathering pregnancy	1.07 (51)	98.03 (4,709)
Outstanding personal achievement	68.13 (3,237)	31.87 (1,514)
Becoming a member of a church	4.62 (220)	95.38 (4,537)
Beginning to date	35.18 (1,665)	64.82 (3,068)
Independent SLEs		
Loss of a job by their father or mother	12.86 (611)	87.14 (4,139)
Marital separation of their parents	3.97 (188)	96.03 (4,551)
The death of a close friend or relative	27.10 (1,293)	72.90 (3,479)
Being hospitalized for illness or injury	9.92 (471)	90.08 (4,276)
Hospitalization of their brother or sister	10.05 (478)	89.95 (4,277)
Remarriage of a parent to a stepparent	2.29 (109)	97.71 (4,661)
Hospitalization of a parent	11.97 (571)	88.03 (4,198)
Major decrease in parental income	15.67 (748)	84.33 (4,026)
Decrease in arguments between parents	34.17 (1,571)	65.83 (3,026)
Moving to a new school or college	31.19 (1,488)	68.81 (3,283)

Table DS3: Descriptive for stressful life events by psychotic experiences: paranoia

	Occurren	ce of SLE				
	No	Yes	-			
	Mean (SD)	Mean (SD)	T value	df	P value	Effect size (d)
Paranoia						
Dependent SLEs						
Becoming involved with drugs	11.87 (10.40)	17.40 (12.56)	-8.01	282.59	> 0.00*	0.48
Being sent away from home	12.06 (10.54)	14.79 (11.57)	-3.70	190.82	> 0.00*	0.25
Breaking up with a boyfriend/girlfriend	11.35 (10.05)	14.42 (11.66)	-9.19	2175.61	> 0.00*	0.28
Suspension from school/college	12.09 (10.51)	15.34 (13.28)	-2.84	138.53	0.01	0.27
Failing an important exam	11.66 (10.28)	14.23 (11.65)	-6.20	1500.06	> 0.00*	0.23
Being responsible for a road accident	12.14 (10.58)	16.18 (11.61)	-2.27	33.51	0.03	0.36
Getting pregnant or fathering pregnancy	12.15 (10.61)	12.65 (10.16)	-0.65	51.32	0.52	0.05
Outstanding personal achievement	12.75 (11.34)	11.89 (10.24)	1.53	2700.79	0.13	0.08
Becoming a member of a church	12.11 (10.58)	12.84 (10.58)	-1.07	238.68	0.28	0.07
Beginning to date	11.51 (10.28)	13.48 (11.10)	-6.36	3323.21	> 0.00*	0.18
Independent SLEs						
Loss of a job by their father or mother	11.99 (10.58)	13.55 (10.59)	-3.77	796.45	> 0.00*	0.15
Marital separation of their parents	12.13 (10.57)	13.47 (11.21)	-1.48	198.26	0.14	0.12
The death of a close friend or relative	11.94 (10.51)	12.77 (10.86)	-2.29	2255.40	0.02	0.08
Being hospitalized for illness or injury	12.01 (10.46)	13.46 (11.74)	-2.38	561.56	0.02	0.13
Hospitalization of their brother or sister	11.97 (10.42)	14.20 (12.06)	-3.69	566.73	> 0.00*	0.20
Remarriage of a parent to a stepparent	12.15 (10.60)	13.16 (10.47)	-1.16	112.32	0.25	0.10
Hospitalization of a parent	12.14 (10.64)	12.30 (10.38)	-0.59	739.45	0.56	0.02
Major decrease in parental income	11.85 (10.37)	13.91 (11.57)	-4.64	1009.67	> 0.00*	0.19
Decrease in arguments between parents	11.94 (10.53)	12.82 (10.84)	-2.76	3135.68	0.01	0.08
Moving to a new school or college	12.08 (10.40)	12.36 (11.05)	-0.01	2700.55	1.00	0.03

Note: Significant at Bonferroni corrected value of 0.0004. df= Welch's degrees of freedom to adjust for uneven group sizes

	Occurre	nce of SLE				
	No	Yes				
	Mean (SD)	Mean (SD)	T value	df	P value	Effect size (d)
Hallucinations						
Dependent SLEs						
Becoming involved with drugs	4.51 (5.85)	7.03 (7.65)	-5.33	277.15	> 0.00*	0.37
Being sent away from home	4.59 (5.91)	6.49 (8.03)	-3.44	188.51	> 0.00*	0.27
Breaking up with a boyfriend/girlfriend	4.27 (5.65)	5.70 (6.75)	-6.56	2038.33	> 0.00*	0.23
Suspension from school/college	4.58 (5.88)	7.21 (8.58)	-3.69	137.75	> 0.00*	0.36
Failing an important exam	4.33 (5.70)	5.90 (6.92)	-6.87	1481.27	> 0.00*	0.25
Being responsible for a road accident	4.64 (5.98)	6.97 (6.89)	-2.15	33.44	0.04	0.36
Getting pregnant or fathering pregnancy	4.64 (5.98)	6.24 (6.88)	-1.68	60.97	0.10	0.25
Outstanding personal achievement	4.93 (6.48)	4.51 (5.75)	1.08	2746.06	0.28	0.07
Becoming a member of a church	4.65 (6.03)	4.66 (5.32)	-0.43	240.18	0.67	0.01
Beginning to date	4.28 (5.57)	5.42 (6.71)	-5.69	3144.81	> 0.00*	0.18
Independent SLEs						
Loss of a job by their father or mother	4.55 (5.94)	5.37 (6.28)	-3.39	780.91	> 0.00*	0.13
Marital separation of their parents	4.61 (5.96)	5.54 (6.51)	-1.86	197.78	0.06	0.15
The death of a close friend or relative	4.51 (5.88)	5.04 (6.29)	-2.82	2238.77	0.00	0.09
Being hospitalized for illness or injury	4.58 (5.90)	5.38 (6.81)	-2.44	562.67	0.02	0.13
Hospitalization of their brother or sister	4.52 (5.84)	6.00 (7.14)	-4.81	569.29	> 0.00*	0.23
Remarriage of a parent to a stepparent	4.63 (5.97)	5.85 (6.94)	-1.93	111.54	0.06	0.19
Hospitalization of a parent	4.60 (5.96)	5.01 (6.21)	-1.49	721.29	0.14	0.07
Major decrease in parental income	4.46 (5.88)	5.71 (6.46)	-5.15	996.25	> 0.00*	0.20
Decrease in arguments between parents	4.32 (5.73)	5.31 (6.40)	-5.78	3053.55	> 0.00*	0.16
Moving to a new school or college	4.51 (5.80)	5.01 (6.40)	-2.44	2754.33	0.01	0.08

Table DS4: Descriptive for stressful life events by psychotic experiences: hallucinations

Note: Significant at Bonferroni corrected value of 0.0004. df= Welch's degrees of freedom to adjust for uneven group sizes

Table DS5: Descriptive for stressful life events by psychotic experiences: cognitive disorganisation

	Occurre	nce of SLE				
	No	Yes	_			
	Mean (SD)	Mean (SD)	T value	df	P value	Effect size (d)
Cognitive Disorganisation						
Dependent SLEs						
Becoming involved with drugs	3.89 (2.82)	5.11 (3.09)	-6.17	278.85	> 0.00*	0.41
Being sent away from home	3.95 (2.85)	4.35 (2.90)	-1.82	190.75	0.07	0.14
Breaking up with a boyfriend/girlfriend	3.72 (2.78)	4.62 (2.95)	-9.36	2095.15	> 0.00*	0.22
Suspension from school/college	3.93 (2.89)	5.03 (3.03)	-4.16	140.01	> 0.00*	0.27
Failing an important exam	3.72 (2.77)	4.84 (2.95)	-10.77	1503.12	> 0.00*	0.39
Being responsible for a road accident	3.95 (2.85)	5.51 (2.33)	-3.89	33.77	> 0.00*	0.60
Getting pregnant or fathering pregnancy	3.96 (2.85)	4.42 (2.85)	-1.15	51.15	0.26	0.16
Outstanding personal achievement	4.39 (2.94)	3.75 (2.78)	7.01	2768.46	> 0.00*	0.22
Becoming a member of a church	3.95 (2.85)	3.95 (2.77)	0.01	240.17	0.99	0.00
Beginning to date	3.76 (2.81)	4.32 (2.88)	-6.39	3286.56	> 0.00*	0.20
Independent SLEs						
Loss of a job by their father or mother	3.91 (2.84)	4.29 (2.87)	-3.02	787.99	0.00	0.13
Marital separation of their parents	3.94 (2.85)	4.40 (2.94)	-2.08	198.59	0.04	0.16
The death of a close friend or relative	3.90 (2.84)	4.10 (2.87)	-2.13	2269.26	0.03	0.07
Being hospitalized for illness or injury	3.92 (2.82)	4.33 (3.01)	-2.86	561.14	0.00	0.14
Hospitalization of their brother or sister	3.90 (2.83)	4.48 (2.91)	-4.10	576.86	> 0.00*	0.20
Remarriage of a parent to a stepparent	3.93 (2.84)	5.03 (2.88)	-3.92	112.02	> 0.00*	0.38
Hospitalization of a parent	3.95 (2.85)	4.00 (2.82)	-0.37	734.98	0.71	0.02
Major decrease in parental income	3.87 (2.84)	4.45 (2.83)	-5.19	1033.23	> 0.00*	0.20
Decrease in arguments between parents	3.93 (2.85)	4.04 (2.83)	-1.24	3166.91	0.22	0.04
Moving to a new school or college	3.86 (2.81)	4.17 (2.91)	-3.38	2756.96	> 0.00*	0.11

Note: \* Significant at Bonferroni corrected value of 0.0004 df= Welch's degrees of freedom to adjust for uneven groupsizes

	Occurre	nce of SLE				
	No	Yes	_			
	Mean (SD)	Mean (SD)	T value	df	P value	Effect size (d)
Grandiosity						
Dependent SLEs						
Becoming involved with drugs	5.31 (4.43)	5.52 (4.36)	-1.20	286.21	0.23	0.05
Being sent away from home	5.28 (4.42)	6.08 (4.32)	-3.38	195.36	> 0.00*	0.18
Breaking up with a boyfriend/girlfriend	5.26 (4.44)	5.48 (4.37)	-1.93	2249.16	0.05	0.05
Suspension from school/college	5.29 (4.41)	6.59 (4.89)	-3.05	139.50	0.00	0.28
Failing an important exam	5.24 (4.39)	5.62 (4.56)	-2.25	1557.59	0.02	0.08
Being responsible for a road accident	5.31 (4.42)	6.41 (4.48)	-1.47	33.52	0.15	0.25
Getting pregnant or fathering pregnancy	5.31 (4.41)	6.47 (5.15)	-1.46	50.98	0.15	0.24
Outstanding personal achievement	4.40 (3.98)	5.76 (4.55)	-10.59	2894.92	> 0.00*	0.32
Becoming a member of a church	5.26 (4.38)	6.77 (5.01)	-4.85	239.20	> 0.00*	0.32
Beginning to date	5.18 (4.39)	5.62 (4.49)	-3.57	3396.39	> 0.00*	0.10
Independent SLEs						
Loss of a job by their father or mother	5.28 (4.38)	5.58 (4.63)	-1.58	791.44	0.12	0.07
Marital separation of their parents	5.31 (4.40)	5.62 (4.64)	-0.63	198.24	0.53	0.07
The death of a close friend or relative	5.27 (4.37)	5.45 (4.56)	-0.87	2241.01	0.38	0.04
Being hospitalized for illness or injury	5.27 (4.43)	5.68 (4.39)	-2.06	577.97	0.04	0.09
Hospitalization of their brother or sister	5.29 (4.44)	5.67 (4.30)	-2.19	592.86	0.03	0.09
Remarriage of a parent to a stepparent	5.34 (4.43)	4.74 (4.25)	1.40	112.29	0.16	0.14
Hospitalization of a parent	5.29 (4.34)	5.60 (4.73)	-1.41	727.94	0.16	0.07
Major decrease in parental income	5.27 (4.36)	5.60 (4.78)	-1.37	1006.20	0.17	0.07
Decrease in arguments between parents	5.07 (4.33)	5.79 (4.50)	-6.25	3287.44	> 0.00*	0.16
Moving to a new school or college	5.30 (4.37)	5.37 (4.51)	-0.12	2779.98	0.90	0.02

Table DS6: Descriptive for stressful life events by psychotic experiences: grandiosity

Note: Significant at Bonferroni corrected value of 0.0004. df= Welch's degrees of freedom to adjust for uneven groupsizes

Table DS7: Descriptive for stressful life events by psychotic experiences: anhedonia

	Occurren	nce of SLE				
	No SLE	Yes SLE	-			
	Mean (SD)	Mean (SD)	T value	df	P value	Effect size (d)
Anhedonia						
Dependent SLEs						
Becoming involved with drugs	17.19 (7.84)	20.08 (8.94)	-5.07	276.99	> 0.00*	0.34
Being sent away from home	17.27 (7.89)	19.06 (8.76)	-2.68	188.55	0.01	0.21
Breaking up with a boyfriend/girlfriend	17.39 (7.88)	17.12 (8.06)	1.01	2160.95	0.31	0.03
Suspension from school/college	17.29 (7.86)	18.88 (9.66)	-1.88	137.18	0.06	0.18
Failing an important exam	17.21 (7.78)	17.86 (8.45)	-2.20	1477.91	0.03	0.08
Being responsible for a road accident	17.33 (7.91)	17.44 (10.07)	-0.06	33.31	0.95	0.01
Getting pregnant or fathering pregnancy	17.32 (7.88)	15.73 (8.71)	1.30	50.94	0.20	0.19
Outstanding personal achievement	18.48 (8.24)	16.81 (7.72)	6.60	2742.19	> 0.00*	0.21
Becoming a member of a church	17.35 (7.93)	16.80 (7.56)	1.04	240.97	0.30	0.07
Beginning to date	17.74 (7.94)	16.63 (7.82)	4.62	3414.87	> 0.00*	0.14
Independent SLEs						
Loss of a job by their father or mother	17.31 (7.92)	17.41 (7.97)	-0.30	789.41	0.77	0.01
Marital separation of their parents	17.27 (7.90)	19.03 (8.44)	-2.79	1.96.42	0.01	0.22
The death of a close friend or relative	17.46 (7.96)	17.08 (7.86)	1.45	2323.03	0.15	0.05
Being hospitalized for illness or injury	17.29 (7.90)	17.76 (8.23)	-1.17	563.92	0.24	0.06
Hospitalization of their brother or sister	17.39 (7.95)	16.82 (7.80)	1.52	587.50	0.13	0.07
Remarriage of a parent to a stepparent	17.34 (7.88)	18.09 (9.48)	-0.85	110.55	0.40	0.09
Hospitalization of a parent	17.37 (7.91)	17.21 (8.09)	0.45	725.34	0.65	0.02
Major decrease in parental income	17.31 (7.92)	17.39 (7.98)	-0.23	1024.55	0.82	0.01
Decrease in arguments between parents	17.61 (8.04)	16.75 (7.65)	3.52	3294.29	> 0.00*	0.11
Moving to a new school or college	17.45 (7.83)	17.05 (8.16)	1.57	2740.49	0.12	0.05

Note: Significant at Bonferroni corrected value of 0.0004. df= Welch's degrees of freedom to adjust for uneven group sizes

Table DS8: Descriptive for stressful life events by psychotic experiences: parent-rated negative symptoms

	Occurre	nce of SLE				
	No SLE	Yes SLE				
	Mean (SD)	Mean (SD)	T value	Df	P value	Effect size (d)
Parent-rated negative Symptoms						
Dependent SLEs						
Becoming involved with drugs	2.76 (3.84)	3.83 (4.58)	-4.46	283.29	> 0.00*	0.25
Being sent away from home	2.75 (3.80)	4.46 (5.27)	-4.74	186.77	> 0.00*	0.37
Breaking up with a boyfriend/girlfriend	2.73 (3.81)	3.05 (4.09)	-2.11	2128.50	0.04	0.08
Suspension from school/college	2.75 (3.82)	5.06 (5.32)	-6.03	141.60	> 0.00*	0.50
Failing an important exam	2.58 (3.69)	3.71 (4.47)	-8.24	1488.04	> 0.00*	0.28
Being responsible for a road accident	2.81 (3.88)	4.14 (3.98)	-2.30	33.49	0.03	0.34
Getting pregnant or fathering pregnancy	2.81 (3.89)	3.94 (4.36)	-2.13	49.98	0.04	0.27
Outstanding personal achievement	3.67 (4.51)	2.40 (3.48)	10.81	2717.64	> 0.00*	0.32
Becoming a member of a church	2.81 (3.90)	2.85 (3.82)	-0.16	238.36	0.87	0.01
Beginning to date	2.84 (3.93)	2.78 (3.82)	0.55	3406.58	0.58	0.02
Independent SLEs						
Loss of a job by their father or mother	2.76 (3.85)	3.13 (4.05)	-2.55	786.78	0.01	0.09
Marital separation of their parents	2.77 (3.86)	3.46 (4.06)	-2.47	200.46	0.01	0.17
The death of a close friend or relative	2.71 (3.77)	3.08 (4.18)	-2.58	2201.27	0.01	0.09
Being hospitalized for illness or injury	2.75 (3.83)	3.36 (4.29)	-3.11	562.45	0.00	0.15
Hospitalization of their brother or sister	2.79 (3.88)	3.06 (3.98)	-1.61	580.63	0.11	0.07
Remarriage of a parent to a stepparent	2.80 (3.90)	3.32 (3.62)	-2.03	111.29	0.04	0.13
Hospitalization of a parent	2.79 (3.89)	3.00 (3.88)	-1.99	736.83	0.05	0.14
Major decrease in parental income	2.72 (3.80)	3.37 (4.33)	-4.01	998.44	> 0.00*	0.16
Decrease in arguments between parents	2.79 (3.89)	2.88 (3.91)	-0.75	3138.43	0.45	0.02
Moving to a new school or college	2.74 (3.76)	3.01 (4.18)	-1.53	2709.39	0.13	0.08

Note: \*Significant at Bonferroni corrected value of 0.0004. df= Welch's degrees of freedom to adjust for uneven group sizes

	Model F	it								
	Compared with saturated model							Parameter estimates: proportion of variance		
	Model	-2LL	df	LRT	Δdf	AIC	р	explained by genetic and en A (CI) C (CI)	<u>E (CI)</u>	
Paranoia	Sat	23525.91	6527	-	<u> </u>	-	p			
1 aranola				-	-	-	-	F -		
	ACE	23529.68	6533	3.77	6	-8.23	.71	.45 (.34, .54) .07 (.00, .16)	.48 (.45, .52)	
	CE	23598.74	6534	72.83	7	58.83	< 0.1		-	
	*AE	23531.56	6534	5.64	7	-8.36	0.58	.52 (.49, .55) -	.48 (.45, .51)	
Hallucinations	Sat	22198.45	6537	-	-	-	-	-		
	*ACE	22199.49	6543	1.04	6	-10.96	.98	.25 (.14, .37) .18 (.08, .27)	.57 (.53, .61)	
	CE	22219.57	6544	21.12	7	7.12	< 0.1		-	
	AE	22211.86	6544	13.41	7	-0.59	0.06		-	
Cognitive	Sat	31571.04	6528	-	-	-	-		-	
disorganisation	ACE	31580.54	6534	9.50	6	-2.50	.15	.44 (.32, .48) .01 (.00, .11)	.55 (.52, .59)	
	CE	31637.18	6535	66.14	7	52.14	<. 01		-	
	*AE	31580.58	6535	9.54	7	-4.46	.22	.45 (.42, .48) -	.55 (.52, .58)	
Grandiosity	Sat	18442.47	6531	-	-	-	-		-	
	*ACE	18447.28	6537	4.81	6	-7.19	.57	.36 (.25, .47) .11 (.01, .20)	.53 (.50, .57)	
	CE	18488.48	6538	41.20	7	27.20	<. 01		-	
	AE	18451.65	6538	4.37	7	-9.63	0.04		-	
Anhedonia	Sat	44554.62	6531	-	-	-	-		-	
	ACE	44560.00	6537	5.39	6	-6.61	.49	.47 (.36, .51) .01 (.00, .10)	.52 (.49, .56)	
	CE	44628.46	6538	73.84	7	59.84	<. 01		-	
	*AE	44560.04	6538	0.04	7	-13.96	0.61	.48 (.45, .51) -	.52 (.48, .55)	
Parent-rated	Sat	17410.51	6512	-	-	-	-	-		
negative	*ACE	17416.10	6518	5.59	6	-6.41	.47	.57 (.50, .64) .26 (.19, .32)	.17 (.16, .18)	
symptoms	CE	17810.00	6519	399.5	7	385.50	<. 01		-	
symptoms				0						
	AE	17465.87	6519	55.37	7	41.37	<. 01		-	
Dependent SLE	sSat	10380.92	6500	-	-	-	-		-	
1	*ACE	10387.40	6506	6.48	6	-5.52	.37	.32 (.22, .43) .19 (.10, .28)	.49 (.46, .52)	
	CE	10424.84	6507	43.92	7	29.92	<. 01	- · · · ·	-	
	AE	10403.11	6507	22.19	7	8.19	<. 01		_	

Table DS9 Fit statistics and parameter estimates for best fitting univariate models

Note: Sat = saturated model; ACE = full model testing genetic, common and unique environmental influences; AE = model testing genetic and unique environment influences; CE = model testing common and unique environmental influences; 2LL = negative 2 log likelihood; df = degrees of freedom; LRT = likelihood ratio  $\chi^2$  test comparing the -2LL fit of each model to the -2LL fit of the saturated model;  $\Delta df$  = difference in degrees of freedom comparing each model to the saturated model; AIC = Akaike's Information Criterion (lower values reflect a better fit); p = p-value. \*Best fitting model.

	Model fit, compared with saturated model						
	Model	-2LL	df	LRT	$\Delta df$	AIC	Р
Paranoia	Saturated	35150.03	13019	-	-	-	-
	ACE	35177.92	13036	27.89	17	-6.11	0.05
	CE	35286.12	13039	136.10	20	96.10	< 0.01
	AE	35194.87	13039	44.84	20	4.84	< 0.01
	E	36556.99	13042	1406.97	23	1360.97	< 0.01
	ACE dropped <i>ra</i>	35184.18	13037	34.15	18	-3.85	0.01
	*ACE dropped <i>rc</i>	35178.73	13037	28.70	18	-7.30	0.05
	ACE dropped ra & rc	35249.87	13038	99.84	19	61.84	< 0.01
Hallucinations	Saturated	35413.78	13029	-	-	-	-
	*ACE	35432.21	13046	18.43	17	-15.57	0.36
	CE	35490.70	13049	76.92	20	36.92	< 0.01
	AE	35459.80	13049	46.02	20	6.02	< 0.01
	E	36644.26	13052	1230.48	23	1184.48	< 0.01
	ACE dropped ra	35433.93	13047	20.16	18	-15.84	0.32
	ACE dropped $r_c$	35434.07	13047	20.29	18	-15.71	0.32
	ACE dropped ra & rc	35476.56	13048	62.78	19	24.78	< 0.01
Cognitive disorganisation	Saturated	35248.56	13020	-	-	-	-
	ACE	35269.67	13037	21.11	17	-12.89	0.22
	CE	35364.09	13040	115.52	20	75.52	< 0.01
	AE	35285.26	13040	36.70	20	16.70	0.01
	E	36450.41	13043	1201.85	20	1161.85	< 0.01
	ACE dropped ra	35271.57	13038	23.00	18	-13.00	0.19
	*ACE dropped $r_c$	35270.06	13038	21.50	18	-14.50	0.25
	ACE dropped <i>ra</i> & <i>rc</i>	35295.90	13039	47.34	19	9.34	< 0.01
Grandiosity	Saturated	35213.41	13023	-	-	-	-
-	*ACE	35232.93	13040	19.52	17	-14.48	0.30
	CE	35311.78	13043	98.36	20	58.36	< 0.01
	AE	35252.84	13043	39.42	20	-0.58	< 0.01
	Е	36501.75	13046	1288.34	20	1248.34	< 0.01
	ACE dropped <i>ra</i>	35235.63	13041	22.21	18	-13.79	0.22
	ACE dropped <i>rc</i>	35235.38	13041	21.96	18	-14.04	0.23
	ACE dropped <i>ra</i> & <i>rc</i>		13042	85.54	19	47.54	< 0.01

Table DS10: Fit statistics for best fitting bivariate models

Note: Sat= saturated model, ACE= full model testing genetic, common and unique environmental influences; AE= model testing genetic and unique environmental influences; CE= model testing common and unique environmental influences; ACE dropped ra= full model testing genetic, common and unique environmental influences with genetic correlation fixed to 0; ACE dropped rc= full model testing genetic, common and unique environmental influences with common environmental correlation fixed to 0; ACE dropped ra and rc= full model testing genetic, common and unique environmental influences with genetic and common environmental correlations fixed to 0; 2LL = negative 2 log likelihood; df = degrees of freedom; LRT = likelihood ratio  $\chi^2$  test comparing the -2LL fit of each model to the -2LL fit of the saturated model;  $\Delta df$  = difference in degrees of freedom comparing each model to the saturated model; AIC = Akaike's Information Criterion (lower values reflect a better fit); p = p-value. \*Best fitting model.





# Association between stressful life events and psychotic experiences in adolescence: evidence for gene–environment correlations

Sania Shakoor, Helena M. S. Zavos, Claire M. A. Haworth, Phillip McGuire, Alastair G. Cardno, Daniel Freeman and Angelica Ronald *BJP* 2016, 208:532-538.

Access the most recent version at DOI: 10.1192/bjp.bp.114.159079

Supplementary Material	Supplementary material can be found at: http://bjp.rcpsych.org/content/suppl/2016/03/22/bjp.bp.114.159079.DC1.html
References	This article cites 33 articles, 10 of which you can access for free at: http://bjp.rcpsych.org/content/208/6/532#BIBL
Reprints/ permissions	To obtain reprints or permission to reproduce material from this paper, please write to <b>permissions@rcpsych.ac.uk</b>
You can respond to this article at	/letters/submit/bjprcpsych;208/6/532
Downloaded from	http://bjp.rcpsych.org/ on November 24, 2016 Published by The Royal College of Psychiatrists

To subscribe to *The British Journal of Psychiatry* go to: http://bjp.rcpsych.org/site/subscriptions/