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The relationship between ADHD and key cognitive phenotypes is not mediated by shared familial effects with IQ


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**Background.** Twin and sibling studies have identified specific cognitive phenotypes that may mediate the association between genes and the clinical symptoms of attention deficit hyperactivity disorder (ADHD). ADHD is also associated with lower IQ scores. We aimed to investigate whether the familial association between measures of cognitive performance and the clinical diagnosis of ADHD is mediated through shared familial influences with IQ.

**Method.** Multivariate familial models were run on data from 1265 individuals aged 6–18 years, comprising 920 participants from ADHD sibling pairs and 345 control participants. Cognitive assessments included a four-choice reaction time (RT) task, a go/no-go task, a choice–delay task and an IQ assessment. The analyses focused on the cognitive variables of mean RT (MRT), RT variability (RTV), commission errors (CE), omission errors (OE) and choice impulsivity (CI).

**Results.** Significant familial association (rF) was confirmed between cognitive performance and both ADHD (rF = 0.41–0.71) and IQ (rF = −0.25 to −0.49). The association between ADHD and cognitive performance was largely independent (80–87%) of any contribution from etiological factors shared with IQ. The exception was for CI, where 49% of the overlap could be accounted for by the familial variance underlying IQ.

**Conclusions.** The aetiological factors underlying lower IQ in ADHD seem to be distinct from those between ADHD and RT/error measures. This suggests that lower IQ does not account for the key cognitive impairments observed in ADHD. The results have implications for molecular genetic studies designed to identify genes involved in ADHD.

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**Key words:** ADHD, cognitive, heritability, IQ, intermediate phenotype.

**Introduction**

Research on attention deficit hyperactivity disorder (ADHD) has identified specific cognitive measures, such as reaction time (RT) performance and commission errors on go/no-go tasks, as potential intermediate phenotypes that may mediate the association between genes and behavioral symptoms (Kuntsi et al. 2006; Rommelse, 2008; Jester et al. 2009). ADHD is also associated with lower IQ, and this association has been shown to be due largely to shared genetic influences.
(Kuntsi et al. 2004; Polderman et al. 2006). Yet it remains unclear to what extent impairment in general cognitive function can explain the observed associations with the other cognitive indices. Here we investigate, using a genetic model-fitting approach, the role of IQ in relation to cognitive impairments that are known to be associated with ADHD and share familial (genetic) influences with the clinical disorder.

Previous research has evaluated the suitability of cognitive performance measures as potential intermediate phenotypes using five main criteria (Gottesman & Shields, 1973; Gottesman & Gould, 2003). Two of the initial criteria are (1) that the cognitive performance measures are associated with the clinical disorder and (2) that the cognitive performance measures share overlapping genetic influences with the disorder or symptoms of the disorder in the general population. Until recently, ADHD research has mainly used a proband–sibling design to nominate potential intermediate phenotypes. This approach compares the means of cognitive performance measures in affected ADHD probands, unaffected siblings of probands and controls. Shared familial influences between the cognitive measure and the disorder are implied when the sibling mean is significantly different from the control group mean, in the direction of the proband mean. Although this method can provide an estimate of the size of the familial effects (Andreou et al. 2007), it cannot be used to investigate the extent to which multiple cognitive measures share the same familial effects.

An alternative approach is to use structural equation modeling (SEM), which provides estimates of the size of shared familial influences between the experimental measure and the clinical disorder and also allows comparison between two or more potential intermediate phenotypes. SEM approaches in twin studies have found little or no evidence for shared environmental effects on either ADHD or the associated cognitive variables (Burt, 2009; Wood et al. 2009 b), so it can be assumed that the familial effects are genetic in origin (Andreou et al. 2007). The multivariate SEM approach to the analysis of putative intermediate phenotypes will allow us to describe the underlying familial architecture and thus the degree to which cognitive variables share etiological influences with each other and with the clinical phenotype. These results will also facilitate reducing the number of intermediate phenotype measures to take forward into genetic mapping studies, where multiple testing is a major problem.

ADHD is associated with impairments on executive function tasks, especially those measuring RT, response inhibition (indexed by commission errors) and sustained attention (indexed by omission errors) (Willcutt et al. 2005; Klein et al. 2006; Johnson et al. 2009; Kuntsi et al. 2009; Wood et al. 2009 b). A strong association has emerged between ADHD and RT variability (RTV) (Klein et al. 2006; Rommelse et al. 2008; Kuntsi et al. 2009; Wood et al. 2009 b). In our own research, using a large proband–sibling and control sample, we previously showed an association with combined-type ADHD on subsets of the current sample for commission and omission errors on a go/no-go task (Uebel et al. 2010), in addition to mean RT (MRT) and RTV on the go/no-go and a four-choice RT tasks (Andreou et al. 2007; Uebel et al. 2010). We also demonstrated an association with ‘choice impulsivity’ (CI; preference for smaller-immediate rewards, incorporating ‘delay aversion’; Marco et al. 2009). Using identical tasks, similar findings emerged in a large general population twin sample (ages 7–10) for the RT variables commission errors† (Kuntsi et al. 2009) and CI (Paloyelis et al. 2009).

In the proband–sibling and control sample we observed improvements in RT mean and variability under incentive or combined fast/incentive conditions that was greater in cases than controls, suggesting an important role for motivational or energetic factors on the processes that underlie the response time measures (Andreou et al. 2007; Uebel et al. 2010). By contrast, case-control differences in omission and commission errors were not altered under the different conditions, suggesting a potentially different underlying cognitive process that was not influenced by motivational or energetic factors for these variables (Uebel et al. 2010).

Using the population twin sample we estimated the heritability of MRT and RTV to be around 50–60% (Wood et al. 2009 b). Furthermore, the estimates increased to around 70% when corrected for measured test–retest unreliability (Kuntsi et al. 2006), nearing the average ‘broad sense’ heritability for ADHD of 70% (Burt, 2009). Quantifying results from other studies that report shared familial variance between RT data and ADHD (Nigg et al. 2004; Bidwell et al. 2007), the genetic correlation between the RT variables and ADHD symptom scores was estimated at around 0.7 (Wood et al. 2009 b), indicating that approximately 70% of the genetic influences on ADHD also influence RT performance, and that the familial variance in sibling studies represents largely genetic influence. Previous analyses on a subset of the present ADHD-proband and control sibling-pair sample similarly indicated that 58–70% of the covariation between ADHD and RT variables was due to shared familial influences (Andreou et al. 2007). In other analyses,

† Omission errors were not investigated because of the small number of such errors made in this general population sample.
performance on the stop signal RT from the stop task (Schachar et al. 2005; Waldman et al. 2006; Bidwell et al. 2007; Rommelse et al. 2008) and commission errors on the continuous performance task (Bidwell et al. 2007) also indicated shared familial variance with ADHD, as indicated by mean scores in unaffected siblings or parents of ADHD-probands that were significantly different from those of controls. Using the go/no-go task, twin data indicated heritability estimates of up to 45% for error data (Kuntsi et al. 2006) and ADHD–unaffected sibling–control means comparisons further suggested shared familial variance with ADHD, assumed to be largely genetic, as above (Slaats-Willemsen et al. 2003; Andreou et al. 2007; Uebel et al. 2010).

ADHD is also associated with lower IQ and twin data indicate that this is also mainly the result of shared genetic influences (Kuntsi et al. 2004; Polderman et al. 2006). An important clinical question therefore is whether lower general cognitive ability, as indexed by lower IQ, can explain some or all of the more specific cognitive performance deficits associated with ADHD. One investigation in ADHD sibling pairs suggested independent familial segregation of executive functioning and IQ in ADHD families (Rommelse et al. 2008), which concurred with results using SEM on the twin sample (7–10 years). Most of the genetic covariance (66–82%) between RT variables and ADHD symptom scores was due to genetic factors that are not shared with IQ, with 92–95% of the overall phenotypic covariance arising independently of etiological (genetic and environmental) factors shared with IQ (Wood et al. 2009b). Establishing whether this translates to a clinical sample is a key aim in the current analyses.

To address this question in a more clinically relevant sample, we now extend our previous IQ-related model-fitting analyses on the twin sample to a large clinical sample of ADHD probands, their siblings and a control sibling-pair sample, and further extend the analysis to additional cognitive variables. Using familial multivariate model fitting, we aimed to investigate whether the familial association between five measures of cognitive performance (MRT, RTV, OE, CE and CI) and a clinical diagnosis of ADHD is mediated through shared familial influences with IQ. A measure of CI was included in light of recent findings that suggest that (unlike the RT data findings) covariation between ADHD and reward preference may, at least in part, be explained by the covariation between ADHD and IQ (Bitsakou et al. 2009; Marco et al. 2009). An additional aim is to examine whether there is justification for aggregating across measures of the same cognitive index, gained either from different tasks (RT variables) or from different conditions of the same task (accuracy variables). Such aggregation across measures is likely to be beneficial for future genetic analyses, as psychometrically robust variables are created (Kuntsi et al. 2006) and the overall number of variables is reduced.

### Method

#### Sample

**ADHD probands and siblings**

Participants were recruited from eight specialist clinics in seven European countries (Belgium, Germany, Ireland, Israel, Spain, Switzerland and the UK), through the International Multicenter ADHD Genetics (IMAGE) project (see Chen et al. 2008 for a detailed description of ascertainment and diagnostic procedures). All participants were of European Caucasian descent and aged 6–18. All probands had a clinical diagnosis of combined subtype ADHD (ADHD-CT) and had one or more full siblings and biological parents available for ascertainment of clinical information and DNA. Siblings within the same age range as the ADHD probands were included in the study and were therefore unselected for ADHD status. Exclusion criteria applying to both probands and siblings included IQ < 70, autism, epilepsy, general learning difficulties, brain disorders and any genetic or medical disorder associated with externalizing behaviors that might mimic ADHD. Where families had more than two siblings, the ADHD index cases were matched to only one of the siblings, to maintain a simple proband-sibling structure for all families included in this analysis. Sibling selection was based, first, on gender and, second, on nearest age to the index proband.

**Control sample**

The control group was recruited from primary (ages 6–11 years) and secondary (ages 12–18 years) schools in the UK, Germany and Spain, aiming for an age and sex match with the clinical sample. The same exclusion criteria were applied as for the clinical sample. In addition, one child subsequently withdrew after testing and three were excluded for having an IQ < 70. A further 10 controls were excluded for having both parent and teacher Conners’ DSM-IV ADHD subscale T scores > 63, to exclude potential, undiagnosed ADHD cases.

**Final sample**

The ADHD proband and sibling sample consisted of 920 individuals and the control sample of 345 individuals. The final total sample therefore consisted of...
1265 individuals, which comprised 580 complete sibling pairs and 105 singletons. Of the 1265 individuals, 524 with ADHD-CT were classified as affected, 16 who met criteria for the hyperactive-impulsive or inattentive subtypes were classified as a ‘subthreshold group’ (who met criteria for the hyperactive-impulsive or inattentive subtypes), and a further 664 individuals were unaffected siblings and controls.

An additional 61 participants had cognitive data, but no clinical data, and their affection status was coded as missing. Ethical approval was obtained from local ethical review boards.

**Procedure**

ADHD probands and their siblings were invited to the research centre for the cognitive assessments and for the parent interview. A minimum of a 48-h stimulant-medication-free period was required for cognitive testing. Patients on non-stimulant medications, such as atomoxetine, were excluded from the study. Controls and their siblings were either invited to the research centre or assessed in schools. Children were given short breaks as required and the total length of the test sessions, including breaks, was approximately 2.5–3 h.

**Measures**

**Diagnosis**

The Parental Account of Child Symptoms (PACS) interview (Taylor et al. 1986) was conducted with the parents to derive the 18 DSM-IV symptoms for ADHD index cases plus siblings who were thought, on the basis of parents’ descriptions of behavior or Conners’ scores ≥ 65, to have ADHD. Situational pervasiveness was defined as some symptoms occurring within two or more different situations from the PACS, in addition to the presence of one or more symptoms scoring 2 or more from the DSM-IV ADHD subscale of the teacher-rated Conners’ (Conners et al. 1998). Impairment criteria were based on severity of symptoms identified in the PACS. Across the IMAGE sites a mean κ coefficient of 0.88 and an average agreement of 96.6% were obtained for ADHD diagnostic categories (Asherson et al. 2008).

**Cognitive tasks**

*Wechsler Intelligence Scales for Children, Third Edition*/*Wechsler Adult Intelligence Scale, Third Edition*. The vocabulary, similarities, picture completion, and block design subtests from the Wechsler Intelligence Scales for Children (WISC-III; Wechsler, 1991; Sattler, 1992), or the Wechsler Adult Intelligence Scale (for those over 16; WAIS-III; Wechsler, 1997) were used to obtain an estimate of the child’s IQ.

The go/no-go task. On each trial of the go/no-go task (Borger & van der Meere, 2000; Kuntsi et al. 2005), one of two possible stimuli appeared for 300 ms in the middle of the computer screen. The child was instructed to respond only to the ‘go’ stimuli and to react as quickly as possible, but to maintain a high level of accuracy. The proportion of ‘go’ stimuli to ‘no-go’ stimuli was 4:1. The children performed the task under three conditions (slow, fast and incentive; see Uebel et al., 2010), matched for length of time on task. Here we present data from the slow condition, with an inter-stimulus interval (ISI) of 8 s and consisting of 72 trials, and the fast condition, with an ISI of 1 s and consisting of 462 trials. The order of presentation of the slow and fast conditions varied randomly across children. The variables obtained from the task are the MRT, standard deviation (SD) of the RT, or RTV, the commission error (CE) and the omission error (OE).

The fast task. The baseline condition of the fast task (Kuntsi et al. 2006; Andreou et al. 2007), with a fore period of 8 s and consisting of 72 trials, followed a standard warned four-choice RT. A warning signal (four empty circles, arranged side by side) first appeared on the screen. At the end of the fore period (presentation interval for the warning signal), the circle designated as the target signal for that trial was filled (colored) in. The child was asked to make a compatible choice by pressing the response key that corresponded directly in position to the location of the target stimulus. Following a response, the stimuli disappeared from the screen and a fixed inter-trial interval of 2.5 s followed. Speed and accuracy were emphasized equally. If the child did not respond within 10 s, the trial terminated. A comparison condition with a fast event rate (1 s) and incentives followed the baseline condition (further details in Andreou et al. 2007). The variables obtained from the task are MRT and RTV; here reported for the baseline condition.

The Maudsley index of childhood delay aversion. Two conditions, each with 20 trials, were administered (Kuntsi et al. 2006; Marco et al. 2009). In each trial, the child had a choice between a smaller-immediate reward (one point involving a 2-s pre-reward delay) and a larger-delayed reward (two points involving a 30-s pre-reward delay). In the no post-reward delay condition, choosing the small reward led immediately to the next trial, reducing the overall length of the condition. In the post-reward delay condition, choosing the small reward led to a delay period of 30 s, and
choosing the large reward led to a delay period of 2 s before the next trial; therefore, the overall delay was constant and independent of the choice made. The order of the two conditions was chosen randomly for each participant. Here, we report data for CI; the percentage of choices for the larger reward in the no post-reward delay condition (reverse scored), which showed the greatest association with ADHD (Marco et al. 2009).

Selection of cognitive task variables for model-fitting analyses. RT data were available from the go/no-go and fast tasks: MRT and RTV were obtained from baseline (slow) conditions, where a strong association with ADHD is observed (Andreuou et al. 2007; Kuntsi et al. 2009; Uebel et al. 2010). CE and OE data were available from the go/no-go task: here we use data obtained from slow and fast conditions only, which showed the strongest associations with ADHD (Kuntsi et al. 2009; Uebel et al. 2010). CI data were obtained from the no post-reward delay condition of the choice-delay task, as this reflects the strongest association with ADHD from this task over and above ‘delay aversion’ (Marco et al. 2009; Paloyelis et al. 2009).

Analyses

Familial structural equation models

The SEM program Mx (Neale et al. 2006) was used to conduct the genetic analyses and to estimate phenotypic correlations. To account for the selected nature of the sample, the selection variable (ADHD status) is included in all models with its parameters fixed. This necessitated ordinal data analysis with the age- and sex-regressed residual scores of the cognitive variables ordinalized into five equal-sized categories, because the Mx program cannot include both ordinal and continuous data in the same analysis. Ordinal data analysis assumes the combination of ordered categories to reflect measurements of an underlying multivariate normal distribution of the traits, with one or more thresholds for each liability distribution to distinguish between the ordered categories. The threshold for ADHD status was fixed to a z value of 1.64 to give a population prevalence of 5%, and its parameters fixed to expected population estimates, with the familiality of ADHD fixed to 80% (sibling correlation of 0.40; see Rijsdijk et al. 2005 for further explanation and validation of this approach).

Phenotypic correlations

Sibling correlations are estimated from a phenotypic correlation model specified in a Gaussian decomposition to give maximum likelihood correlations between the phenotypic variance in each measure for each sibling, and to allow additional constraints. In addition to the constraints outlined above, further constraints reflect the assumptions of the familial model: that phenotypic correlations across traits are the same across siblings and that cross-trait cross-sibling correlations are independent of sibling status (first- or second-born).

Genetic models: Cholesky decomposition (Fig. 1)

Using the information that siblings reared together share, on average, 50% of their segregating alleles, multivariate models use cross-trait cross-sibling correlations to decompose the covariance between traits into familial [F; 50–100% of additive genetic (A) + 100% common environmental (C)] influences, and individual-specific environmental (E) influences, which include possible measurement error. Without knowing the underlying ratio of A:C influences for each variable, it is not possible to specify a variance/covariance structure that accurately estimates the amount of variance due to A+C influences, and as we are here focusing on shared variance, overall percentages for variance due to F and E parameters for each variable are not presented (although estimates are available in Fig. 1).

A triangular, or Cholesky, decomposition is imposed on the data, which allows an estimation of the extent to which traits share common F and E influences. Although the ordering of variables in the Cholesky is often arbitrary for computational reasons, in the multivariate models we assigned IQ to be the first measured variable, to allow an estimation of the extent to which the covariance between cognitive data and ADHD was independent of risk factors shared with IQ. Because of the computational intensity of ordinal data analysis, 95% confidence intervals are not available. However, the significance of parameters in the main model (Fig. 1) were tested by dropping, in turn, each parameter and comparing the $\chi^2$ of the reduced model to that of the full model with a 1-df test of freedom at the $p < 0.05$ level. A significant result indicates that the model was a worse fit without this parameter, and thus the parameter was significant with an $\alpha$ level of 0.05.

Results

Group differences between ADHD-CT probands, siblings of probands and controls existed for gender and parent and teacher ratings of ADHD behaviors; and between probands and controls, and siblings and controls (but not probands and siblings) for IQ and age (Table 1). The use of definition variables in Mx was not possible because of the computational intensity of the
integration in ordinal data analysis. Accordingly, the data were regressed for age and gender prior to the familial modeling and the age- and sex-corrected residuals used. IQ and ADHD status were included as measured variables.

**Multivariate familial models across ADHD and MRT, RTV, CE or OE**

To examine whether cognitive variables across similar (theoretically related) tasks, or across different conditions of the same task, reflect similar etiological influences, models were run across two sets of data for each cognitive index (ADHD was also included to correct for ascertainment bias). The similar phenotypic and cross-sibling correlations from the constrained, phenotypic model indicate that shared familial effects underlie task (for MRT and RTV) or condition (for CE and OE) level covariance (Table 2). This is reflected in the high familial correlations between task- or condition-level data on the same cognitive construct of between $r^F = 0.69$ and $0.83$ (Table 2).

**Multivariate familial models across IQ, ADHD, CI and mean MRT, RTV, CE or OE scores (Fig. 1)**

The correlations between ADHD and IQ were $-0.20$ at the phenotypic level and $-0.17$ at the familial level.

Given the results outlined above, the extent to which etiology of any overlap between cognitive indices and ADHD was independent of etiology shared with IQ was examined using mean scores across the measures of MRT, RTV, CE or OE, using a Cholesky decomposition (Table 3). By summing the contribution of $F$ and $E$ factors that contribute to the covariation between cognitive indices and ADHD that do not influence the population variance in IQ, and taking them as a percentage of the total covariance, we obtain the percentage of the covariation that is independent of shared etiological influences with IQ.

Etiological ($F/E$) correlations for mean scores with ADHD were as expected from task- or condition-specific measures (Table 2). The overlap between ADHD and the cognitive indices was largely independent of any shared etiology between ADHD and IQ. Between 73% and 81% of the familial influences that were shared between ADHD and the cognitive indices were independent of those shared with IQ. The exception was CI, which was lower at 62%, indicating a greater degree of overlap with the familial influences shared between ADHD and IQ. The percentage of the covariation with ADHD that was independent of any shared etiological ($F + E$)
Data from a large ADHD and control sibling-pair sample showed that the association between ADHD and several cognitive measures (MRT, RTV, CE and OE) is largely (80–87%) independent of etiological influences shared with IQ. This confirms and extends previous model-fitting findings on a general population twin sample (Wood et al. 2009b), and also previous findings on separate clinical samples using different analytical techniques (Rommelse et al. 2008; Jester et al. 2009). The evidence is therefore accumulating that the relationship between ADHD and key cognitive phenotypes is not mediated by shared familial effects with IQ. This suggests that several distinct processes are involved and that impairments in general cognitive ability are unlikely to explain the specific deficits seen in ADHD.

**Table 1. Group means (and standard deviations) for sample characteristics and cognitive variables**

<table>
<thead>
<tr>
<th></th>
<th>ADHD probands</th>
<th>Siblings of ADHD probands</th>
<th>Controls</th>
</tr>
</thead>
<tbody>
<tr>
<td>Male (%)abc</td>
<td>89.01</td>
<td>49.78</td>
<td>70.43</td>
</tr>
<tr>
<td>Age (years)ac</td>
<td>11.45 (2.73)</td>
<td>11.38 (2.96)</td>
<td>12.07 (2.47)</td>
</tr>
<tr>
<td>IQac</td>
<td>102.02 (15.44)</td>
<td>103.43 (13.59)</td>
<td>108.91 (13.71)</td>
</tr>
<tr>
<td>Parent-rated Conners’ DSM-IV ADHD subscaleabc</td>
<td>78.87 (8.51)</td>
<td>54.80 (13.62)</td>
<td>52.20 (10.83)</td>
</tr>
<tr>
<td>Teacher-rated Conners’ DSM-IV ADHD subscaleabc</td>
<td>71.20 (10.70)</td>
<td>56.54 (12.41)</td>
<td>50.32 (9.17)</td>
</tr>
<tr>
<td>Mean RT</td>
<td>645.70 (233.85)</td>
<td>538.97 (184.81)</td>
<td>495.26 (233.85)</td>
</tr>
<tr>
<td>Fast task (baseline condition)abc</td>
<td>924.01 (352.18)</td>
<td>879.75 (401.17)</td>
<td>672.08 (208.34)</td>
</tr>
<tr>
<td>Go/no-go task (slow condition)abc</td>
<td>455.39 (343.55)</td>
<td>357.82 (323.58)</td>
<td>202.58 (178.50)</td>
</tr>
<tr>
<td>Mean RT variability</td>
<td>312.79 (221.37)</td>
<td>225.48 (169.37)</td>
<td>143.54 (178.4)</td>
</tr>
<tr>
<td>Go/no-go task (slow condition)abc</td>
<td>52.84 (23.57)</td>
<td>43.48 (24.79)</td>
<td>37.64 (22.53)</td>
</tr>
<tr>
<td>Commission errors</td>
<td>53.92 (17.89)</td>
<td>44.39 (18.97)</td>
<td>41.28 (17.84)</td>
</tr>
<tr>
<td>Omission errors</td>
<td>13.04 (14.39)</td>
<td>8.15 (10.93)</td>
<td>3.56 (5.47)</td>
</tr>
<tr>
<td>Choice impulsivityac</td>
<td>72.22 (32.72)</td>
<td>76.65 (29.23)</td>
<td>86.43 (23.75)</td>
</tr>
</tbody>
</table>

ADHD, Attention deficit hyperactivity disorder; RT, reaction time.

* Significant differences between probands and controls (p < 0.05).

**Table 2. Maximum likelihood phenotypic, cross-sibling and familial correlations for cross-task or cross-condition data from constrained phenotypic models across ADHD (used for ascertainment correction) and cognitive variables**

<table>
<thead>
<tr>
<th></th>
<th>Phenotypic correlation</th>
<th>Cross-sibling correlation</th>
<th>Familial correlation</th>
</tr>
</thead>
<tbody>
<tr>
<td>Mean RTa</td>
<td>0.52</td>
<td>0.19</td>
<td>0.69</td>
</tr>
<tr>
<td>RT variabilitya</td>
<td>0.49</td>
<td>0.20</td>
<td>0.70</td>
</tr>
<tr>
<td>Commission errorsb</td>
<td>0.59</td>
<td>0.16</td>
<td>0.74</td>
</tr>
<tr>
<td>Omission errorsb</td>
<td>0.50</td>
<td>0.20</td>
<td>0.83</td>
</tr>
</tbody>
</table>

ADHD, Attention deficit hyperactivity disorder; RT, reaction time.

Choice impulsivity is not included as the variable is based on only one condition/task.

influences with IQ was 85% for MRT, 87% for RTV, 84% for CE, 80% for OE and 61% for CI.

**Discussion**

Data from a large ADHD and control sibling-pair sample showed that the association between ADHD and several cognitive measures (MRT, RTV, CE and OE) is largely (80–87%) independent of etiological influences shared with IQ. This confirms and extends previous model-fitting findings on a general population twin sample (Wood et al. 2009b), and also previous findings on separate clinical samples using different analytical techniques (Rommelse et al. 2008; Jester et al. 2009). The evidence is therefore accumulating that the relationship between ADHD and key cognitive phenotypes is not mediated by shared familial effects with IQ. This suggests that several distinct processes are involved and that impairments in general cognitive ability are unlikely to explain the specific deficits seen in ADHD.

For individual cognitive measures, the high familial correlations (0.69–0.83) obtained across conditions or tasks indicate that they are largely measuring the same underlying liability. These results, on familial sharing, indicate that performance seems to be relatively stable across task and condition, when focusing on the cognitive measures that are associated with ADHD. These results support the aggregation of data across the variables examined here for future genetic mapping analyses of the common genetic influences that span the various measures. They also suggest that the individual cognitive measures are indexing the same
unitary construct across these two tasks, providing support for combining datasets for meta-analytic studies, where the data are gathered using similar paradigms. This is important for genetic mapping studies because replication of preliminary findings and pooling of data to reach genome-wide levels of significance are essential to confirm the identity of true genetic associations. Although these results are promising, caution must be advised in considering the exact task parameters. For example, for RTV we have shown, using the current sample (Andreou et al. 2007; Uebel et al. 2009), how the strength of association with ADHD depends crucially on task condition parameters, such as event rate and incentives.

Our results across tasks and conditions show a striking similarity with results in a younger, general population twin study (Wood et al. 2009b). An example is the comparability of the genetic correlations between ADHD symptom scores and RTV in the fast and go/no-go tasks in the twin study (~0.6–0.7) and the familial correlations in the current study (~0.6–0.8). In addition to suggesting that the familial covariance is largely genetic, these findings emphasize the robustness of the methods and findings, which replicate not only across tasks and samples but also across definitions of ADHD (diagnosis versus a continuum of symptoms in the general population), supporting the conceptualization of ADHD as the extreme of a continuously distributed trait. Future analyses will aim to extend this work and examine whether there are separate pathways between the RT and error variables to account, for example, for bottom-up influences from subcortical arousal structures and brief reductions in the top-down control of sustained attention and inhibition (Halperin & Schulz, 2006; Johnson et al. 2007, 2008; Halperin et al. 2008; O’Connell et al. 2008; Loo et al. 2009; Kuntsi et al. in press). The current data emphasize that these processes do not arise out of pathways shared with the more generalized deficit of lowered IQ.

The familial sharing between ADHD and CI was lower (with a familial correlation of ~0.14) than that found for the other cognitive variables. The percentage of the covariation with ADHD that was independent of shared etiological influences with IQ was also lower, at 61%, indicating that CI and IQ are more closely related constructs at the etiological level. Research investigating whether there are separate and dissociable mechanisms, underpinned by different neural circuitry (Sonuga-Barke, 2005), may clarify the role of CI in ADHD symptomatology. Overall, the evidence in support of CI as an intermediate familial phenotype in ADHD is less strong than for the other cognitive variables investigated here, but it is unclear at present whether this reflects, at least in part, psychometric properties of the particular measure used in this study (in particular, ceiling effects; see Kuntsi et al. 2006) and should therefore be further investigated using alternative measures of this construct.

The current analyses add to the emerging understanding of the genetic architecture of the cognitive and energetic processes that underlie the symptoms of ADHD. For the first time, a clinical sample has been used to quantify that the familial influences ADHD shares with IQ are largely separable from those that ADHD shares with the other key cognitive indices associated with the disorder. The aetiological factors that give rise to lower IQ in ADHD seem to be largely distinct from those that give rise to the association of ADHD with RT variables, CE and OE. Lower IQ does not seem to be a general explanation for the impairments in these specific cognitive domains.

### Table 3. Etiological correlations from correlated factors solutions of Cholesky models estimating the etiological influences across IQ, ADHD status, and cognitive variables

<table>
<thead>
<tr>
<th>Phenotypic correlations</th>
<th>Cross-sibling correlations</th>
<th>Familial correlations</th>
<th>Individual-specific correlations</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>ADHD</td>
<td>IQ</td>
<td>ADHD</td>
</tr>
<tr>
<td>Mean RT*</td>
<td>0.42</td>
<td>−0.24</td>
<td>0.22</td>
</tr>
<tr>
<td>RT variability*</td>
<td>0.47</td>
<td>−0.25</td>
<td>0.23</td>
</tr>
<tr>
<td>Commission errors*</td>
<td>0.24</td>
<td>−0.16</td>
<td>0.12</td>
</tr>
<tr>
<td>Omission errors*</td>
<td>0.33</td>
<td>−0.23</td>
<td>0.17</td>
</tr>
<tr>
<td>Choice impulsivity</td>
<td>−0.16</td>
<td>0.30</td>
<td>−0.03</td>
</tr>
</tbody>
</table>

ADHD, Attention deficit hyperactivity disorder; RT, reaction time.

*Mean across fast task and slow condition of the go/no-go task.

*Mean across slow and fast conditions of the go/no-go task.

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Declaration of Interest

None.

References


Johnson KA, Kelly SP, Bellgrove MA, Barry E, Cox M, Gill M, Robertson IH (2007). Response variability
in attention deficit hyperactivity disorder: evidence for neuropsychological heterogeneity. *Neuropsychologia* 45, 630–638.


Rommelse NN (2008). Endophenotypes in the genetic research of ADHD over the last decade: have they lived up to their expectations? *Expert Reviews in Neurotherapy* 8, 1425–1429.


Waldman ID, Nigg JT, Gizer IR, Park L, Rapley MD, Friderici K (2006). The adrenergic receptor alpha-2A gene (ADRA2A) and neuropsychological executive functions as putative endophenotypes for childhood
ADHD. Cognitive, Affective and Behavioral Neuroscience 6, 18–30.


Wood AC, Asherson P, van der Meere JJ, Kuntsi J (2009b). Separation of genetic influences on attention deficit hyperactivity disorder symptoms and reaction time performance from those on IQ. *Psychological Medicine*. Published online: 15 September 2009. doi:10.1017/S003329170999119X.