Comorbid Problems in ADHD: Degree of Association, Shared Endophenotypes, and Formation of Distinct Subtypes. Implications for a Future DSM

Submitted as:

Chapter 10

Abstract

We aimed to assess which comorbid problems (oppositional defiant behaviors, anxiety, autistic traits, motor coordination problems, and reading problems) were most associated with Attention-Deficit/Hyperactivity Disorder (ADHD); to determine whether these comorbid problems shared a neuropsychological endophenotype with ADHD; and to determine whether an ADHD-endophenotype supported the hypothesis that ADHD with comorbid problems is a qualitatively different phenotype than ADHD without comorbid problems. An aggregated ADHD-endophenotype was based on ten neuropsychological tasks administered to 816 children from ADHD- and control-families. Additional data on comorbid problems were gathered using questionnaires. Results indicated that oppositional defiant behaviors appeared the most important comorbid problems of ADHD, followed by autistic traits, and then followed by motor coordination problems, anxiety, and reading problems. The ADHD-endophenotype was related to the comorbid problems and cross-related to comorbid problems in siblings, suggesting a shared etiology of ADHD and comorbid problems. ADHD in co-occurrence with comorbid problems did not appear to be a distinct subtype of ADHD.
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Introduction

Attention-Deficit/Hyperactivity Disorder (ADHD) (American Psychiatric Association [APA], 1994) is frequently associated with a range of other psychiatric and neurological disorders. It is estimated that around 60-100% of patients with ADHD also exhibit one or more comorbid disorders (Gillberg et al., 2004) that often continue into adulthood (Biederman, 2004; Kessler et al., 2006). Around 42-90% of patients meet criteria for Oppositional Defiant Disorder (ODD) and/or Conduct Disorder (CD) (Angold et al., 1999; Bauermeister et al., 2007; Cunningham & Boyle 2002; Gillberg et al., 2004; Jensen et al., 1997), disorders characterized by externalizing behavioral problems, such as aggressive behavior, difficulty with authority (ODD) or lying, stealing, and vandalism (CD). Furthermore, around 13-51% of ADHD patients suffer from internalizing disorders, such as anxiety or depression (Angold et al., 1999; Bauermeister et al., 2007; Gillberg et al., 2004; Jensen et al., 1997). Currently, the Diagnostic and Statistical Manual for Mental Disorders (DSM) IV (APA 1994) rules out a diagnosis of autistic disorder with ADHD. Nevertheless, a large percentage (65-80%) of children with ADHD portrays symptoms in the autistic spectrum (Clark et al., 1999; Gillberg et al., 2004). Other disorders frequently observed in patients with ADHD, are dyslexia (25-40%), motor coordination problems (50%), dyscalculia (10-60%), sleep disorders (25-50%), and enuresis and/or encopresis (30%) (Bhatia et al., 1991; Gillberg et al., 2004; Owens 2005; Willcutt et al., 2005b). ADHD patients with comorbid problems compared to ADHD patients without comorbid problems appear to have a more severe form of ADHD, are often more impaired in their daily functioning, and have a poorer long term prognosis (Bauermeister et al., 2007; Biederman et al., 1996; Connor et al., 2003; Gillberg et al., 2004). This may have implications for diagnosis and treatment of ADHD, like broad assessments covering multiple childhood psychiatric disorders and not only ADHD and interventions that also address the comorbid problems (Biederman et al 1991; Gillberg et al., 2004; Hechtman et al., 2005; Jensen et al., 2001).

Clearly, comorbid problems are an important aspect of ADHD. However, the word ‘comorbid’ has various meanings. For example, it may refer to one disorder leading to another, it may refer to two (or more) clearly separable and independent disorders occurring together, or it can refer to two (or more) disorders that share a common underlying etiology (Angold et al., 1999; Biederman et al., 1991; Caron & Rutter 1991; Gillberg et al., 2004). Understanding how comorbidity arises may inform our understanding of the development of psychopathology (Angold et al., 1999). In light of the development of a future DSM, studying the degree and
nature of the association between ADHD and its comorbid problems deserves further attention (Jensen et al., 1997).

Therefore, the first aim of our study was to investigate the degree of association between ADHD and several comorbid problems (oppositional defiant behaviors, anxiety, autistic traits, motor coordination problems, and reading problems). It was analyzed which comorbid problem was the most important co-occurring with ADHD. To assess this, we used questionnaires. Questionnaires generally correlate strongly with structured interviews (Biederman et al., 2005; Conners 1996) and may aptly reflect the underlying continuous distribution of traits (Reich et al., 1975). Using questionnaires, it was possible to estimate the degree of phenotypic association between ADHD and its comorbid problems.

The second aim of our study was to clarify whether ADHD and comorbid problems are 'merely' associated with each other at a phenotypic level or whether they partly arise from the same underlying vulnerability traits. In previous studies (Rommelse et al., 2007a, b, c, d, 2008a), we have identified neuropsychological endophenotypes for ADHD: heritable deficits that formed underlying, vulnerability traits for ADHD. Characteristic of endophenotypes is that they are also present in at risk non-affected family-members of ADHD patients and show resemblance between family-members (Waldman, 2005). These endophenotypes may shed light on the nature of the association between ADHD and its comorbid problems: if the ADHD-endophenotype also relates to the comorbid problems (while correcting for ADHD), it is likely that both disorders (partly) relate to the same underlying neuropsychological substrate and are not merely phenotypically associated with one another. Further evidence for this hypothesis may be obtained, when these neuropsychological endophenotypes cross-correlate with comorbid problems in siblings (i.e. neuropsychological deficits of a child relate to comorbid problems in his/her siblings), suggesting similar familial influences give rise to ADHD and comorbid problems. These correlations and sibling cross-correlations may indicate whether or not ADHD and comorbid problems partly arise from the same underlying neuropsychological vulnerability traits. The neuropsychological tasks used in this study comprise both executive/cognitive and motor functions and, therefore, may aptly reflect broad neuropsychological functioning.

The third aim of our study was to investigate whether the specific combination of ADHD with a comorbid problem may be seen as forming a distinct phenotype (Banaschewski et al., 2005; Biederman et al., 1991; Caron & Rutter 1991; Jensen et al., 2001) and not merely as 'more of the same of both disorders'. If the combination of both disorders is different from the additive effect of both problem domains, the interaction term will relate significantly to the ADHD-endophenotype, indicating that the combination of both disorders differs qualitatively from the
individual syndromes. If, however, the combination of both problem domains is similar to the additive effect of both disorders, the interaction term will not relate significantly to the ADHD-endophenotype, indicating that the combination of both disorders differs only quantitatively from the individual syndromes.

**Methods**

**Participants**

Participants were recruited in the Dutch part of an international multicenter ADHD genetic study (IMAGE) that aims to identify genes that increase the risk for ADHD using QTL linkage and association strategies (Brookes et al., 2006). Families with at least one child with the combined subtype of ADHD (proband) and at least one additional sibling (regardless of possible ADHD-status) participated. Proband were clinically referred to specialist centres. Additional control families were recruited from primary and high schools from the same geographical regions as the participating ADHD-families. Controls and their first degree relatives were required to have no formal or suspected ADHD diagnosis. A total of 238 ADHD-families and 147 control-families fulfilled inclusion and exclusion criteria. Within the ADHD-families, 238 probands (all with combined subtype ADHD), 112 affected siblings (64 with combined subtype, 28 with inattentive subtype and 20 with hyperactive-impulsive subtype) and 195 non-affected siblings participated. Control-families consisted of 271 children. For 51 control children, no additional control sibling could be recruited for the study. The groups did not differ in age, but the group of probands and affected siblings had a larger percentage of males than the groups of non-affected siblings and controls (probands: $M$ age = 12.0 (2.5), % males = 84.5; affected siblings: $M$ age = 12.0 (3.4), % males = 56.3; non-affected siblings: $M$ age = 11.5 (3.6), % males = 45.1; controls: $M$ age = 11.6 (3.2), % males = 40.6). All children were between the ages of 5 and 19 years and were of European Caucasian descent. Participants were excluded, if they had an IQ < 70, a diagnosis of autism, epilepsy, brain disorders or known genetic disorders, such as Down syndrome or Fragile-X-syndrome.

The exact screening procedures and measures for ADHD phenotyping have been described previously (Brookes et al., 2006). Briefly, screening questionnaires (parent and teacher Conners’ long version rating scales [Conners, 1996] and parent and teacher Strengths and Difficulties Questionnaires [Goodman, 1997]) were used to identify children with ADHD symptoms. Scores were considered clinical if $T$-scores were obtained $\geq 63$ on at least one
Conners’ ADHD-subscale (DSM-IV Inattention, DSM-IV Hyperactive-Impulsive, and DSM-IV ADHD Total) or scores > 90th percentile on the SDQ-hyperactivity scale. Additionally, a semistructured interview, the Parental Account of Children’s Symptoms (PACS) (Taylor, 1986), was administered for children scoring clinically on any of the questionnaire scales of interest. For diagnostic purposes, questionnaire data and the PACS were subjected to a standardised algorithm to derive each of the 18 DSM-IV ADHD symptoms, providing operational definitions for each behavioral symptom (see for a description Rommelse et al., 2007a). The section on autistic traits of the PACS was administered to exclude possible cases with autistic disorder, if a clinical score (raw score ≥ 15) was obtained on the Social Communication Questionnaire (SCQ; Berument et al., 1999). Concerning control children, the Conners’ long version for both parents and teachers was completed and control children were required to obtain non-clinical scores on all scales measuring ADHD related symptomatology. A measure of ADHD for further analyses was operationalized by averaging the scaled Conners’ N-subscale (ADHD Total) for the parent and teacher rating.

This sample of children allowed for analyses on continuously distributed data on ADHD and comorbid problems, since they represented the whole range of possible scores on the Conners’ ADHD scales (T-scores from 40 to 90: control children had T-scores up till 62, probands had T-scores of 63 and over, and siblings of probands had scores in the whole range) and since no disorders were excluded in the screening, except a full diagnosis of autism.

Measures
Data on Comorbid Problems
Oppositional Defiant Behaviors
The Conners’ Long Version filled out by the parents and teachers was used to get an indication of oppositional defiant behaviors (Subscale A) (Conners, 1996). Subscale A consisted of ten items that were rated on a 4-point scale (0 = not true at all; 3 = very much true). Higher scores were indicative of more oppositional defiant behaviors. Reliability and validity of the Conners’ have been established (Conners, 1996). A measure of oppositional defiant behaviors was operationalized by averaging the scaled Conners’ A-subscale (Oppositional) for the parent and teacher rating.

Anxiety
The Conners’ Long Version was used to obtain a measure of anxiety (Subscale D), both as observed by parents and teachers (Conners, 1996). Subscale D consisted of eight items that were
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rated on a 4-point scale (0 = not true at all; 3 = very much true). Higher scores were indicative of more symptoms of anxiety. A measure of anxiety was operationalized by averaging the scaled Conners’ D-subscale (Anxious-Shy) for the parent and teacher rating.

**Autistic Traits**

Ratings on symptoms in the autism spectrum were obtained using the Dutch 49-item Children’s Social Behavior Questionnaire (CSBQ) (Hartman et al., 2006). Parents were asked to rate the behaviors of their child on a 3-point scale (0 = not; 2 = clearly/often). The questionnaire consists of several subscales: Unadjusted behavior, Tendency to withdrawal, Orientation problems, Difficulty understanding, Stereotype movements, and Anxiety for changes. The reliability and validity of this questionnaire have been established (Hartman et al., 2006). The total sum score was used as a measure of symptoms in the autism spectrum, with higher scores indicative of more severe autistic like behaviors.

**Motor Coordination Problems**

Motor coordination problems were rated by parents and teachers. Parents filled out the Developmental Coordination Disorder Questionnaire (DCD-Q) (Wilson et al., 2000). The DCD-Q consists of 17 items that were rated on a 5-point scale (1 = not at all like this child; 5 = extremely like this child). The total score was used as an indication of motor coordination problems, with a low score reflective of more severe motor coordination problems. The reliability and validity of this questionnaire have been confirmed (Wilson et al., 2000). Teachers rated motor coordination problems on the 17-item Groninger Motor Observation-scale (GMO) (Kalverboer & Van Dellen, 1990), using a 4-point scale (1 = not applicable; 4 = applicable). Items related both to fine and gross motor coordination. The total score was used as indication of motor coordination problems, with higher scores reflecting more severe problems. The reliability and validity of this questionnaire have been established (Kalverboer & Van Dellen, 1990). A measure of motor coordination problems was operationalized by averaging the standardized total scores for the parent and teacher rating.

**Reading Problems**

Reading problems were assessed with the 6-item questionnaire of Willcutt and colleagues filled out by the parents (Willcutt et al., 2008). A 5-point scale was used (1 = never/not at all; 5 = always/a great deal). A higher score reflected more severe reading problems. The total score of
this questionnaire is known to correlate between .61 and .71 with validated measures of reading problems (Willcutt et al., 2008) and was used as measure of reading problems.

**Neuropsychological Tasks**

The ten experimental tasks described in this study have been fully described elsewhere (Rommelse et al., 2007a, b, c, d, 2008a). A short description of each task is given below. Based on previous results (Rommelse et al., 2007a, b, c, d, 2008a), the task variable that showed the most optimal result in the endophenotypic analyses in the five previous studies was chosen for the current analyses. These task variables were significantly associated with ADHD (i.e. affected children performed more poorly than control children) and were significantly correlated between siblings.

**Stop Task**

The Stop Task was used to measure speed of inhibition of an ongoing response (Rommelse et al., 2008a). Subjects were presented two types of trials: go-trials and stop-trials. Go-trials consisted of the presentation of a go-stimulus (drawing of a plane) that was either pointing to the right or to the left. Children were instructed to press a response button that corresponded to the direction of the stimulus as quickly and as accurately as possible. Stop-trials were identical to the go-trials but in addition a stop-signal was presented (drawing of a cross that was superimposed on the plane). Children were required to withhold their response to the stop-signal. Go stimuli were displayed for 1000 ms, preceded by a 500 ms fixation point. Stop signals were displayed for 1000 ms minus delay time. Inter-trial intervals were 3000 ms. The delay between the go- and stop-signal was dynamically varied so that it could be estimated when the child successfully inhibited 50% of the stop-trials, and unsuccessfully inhibited the other 50%. At this point, the go-process and stop-process were of equal duration, which made it possible to estimate the latency of the stop-process: the Stop Signal Reaction Time (SSRT). A total of 2 practice blocks and 4 experimental blocks were administered, each consisting of 60 trials. The first practice block consisted of only go-trials. The second practice block and the 4 experimental blocks consisted of 75% go-trials and 25% stop-trials. Go- and stop-trials were pseudo-randomly presented. Task administration took about 15 minutes. Based on previous results (Rommelse et al., 2008a), the dependent measure was the SSRT, which showed endophenotypic-like group differences and correlated between siblings.
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Shifting Attentional Set

Shifting Attentional Set was designed to measure the accuracy of motor inhibition and cognitive flexibility (Rommelse et al., 2007b). The task consisted of three blocks of which the first block was designed to acquire a baseline of the accuracy of responding with which the performance on the second (motor inhibition) and third (cognitive flexibility) block could be compared. In all blocks, trials consisted of a horizontal bar with ten grey squares presented permanently at the centre of the screen. From trial to trial, a coloured square moved across the bar in a random direction (either one square to the right or to the left). Responses were required to be initiated between 150 to 5000 ms after a square moved one position, otherwise a trial was replaced. The task was self-paced with post-response intervals of 250 ms. In the first block, the moving square was coloured green, and compatible responses were required. Children were instructed to press a response button as quickly and as accurately as possible that corresponded to the direction in which the stimulus moved. In the second block, the moving square was coloured red, and incompatible responses were required. The suppression of the automatic compatible response, in order to generate a non-automatic incompatible response, was hypothesized as requiring inhibitory control. In the third block, the colour of the moving square alternated randomly between green and red, and both compatible and incompatible responses were required. Thus, both the direction and the colour of the square were unpredictable. This mixture of both compatible and incompatible trials was hypothesized as requiring high levels of cognitive flexibility in addition to inhibitory control. The first and second block consisted of 10 practice trials and 40 experimental trials. The third block consisted of 16 practice trials and 80 experimental trials. Administration took about 10 to 15 minutes. The dependent measure was the percentage of errors across blocks measuring overall inaccuracy on cognitive taxing tasks, which was the best indicator of endophenotypic vulnerabilities (Rommelse et al., 2007b).

Time Reproduction

The Timetest Application Version 1.0 was used to measure the accuracy of time reproduction (Rommelse et al., 2007a). Stimuli consisted of temporal intervals with different durations (4, 8, 12, 16, 20 s) that had to be reproduced as accurately as possible. The task was administered first in the visual modality (light bulb) and thereafter in the auditory modality (tone). Children were not informed about the length of the intervals. In both modalities, 3 practice and 20 experimental trials were administered. The five interval lengths were randomly presented four times. Task administration for both modalities required 15 minutes. Based on previous results, the main dependent measure was the precision of the reproduction (operationalized as the absolute
discrepancy between the response length and the stimulus length) averaged across trials and modalities, which was abnormal in children with ADHD and their non-affected siblings and correlated between siblings (Rommelse et al., 2007a).

**Visuo-Spatial Sequencing**

The Visuo-Spatial Sequencing task was used to measure accuracy of visuo-spatial working memory (Rommelse et al., 2008a). Stimuli consisted of nine circles symmetrically organized in a square (3 by 3). On each trial, a sequence of circles was pointed at by a computer-driven hand. Subjects were instructed to replicate the exact same sequence of circles, by pointing to them with the small, self-driven hand. There were no time constrictions. One practice trial and 24 experimental trials were presented. Every succeeding trial increased in difficulty level: an increase in the number of circles required to be remembered and/or an increase in the complexity of the spatial pattern (i.e. the trial consisted of circles that were spatially further removed from one another instead of being close to one another), hence manipulating working memory demands. Task administration took about 7 minutes. Based on previous results, the total number of correct targets in the correct order was used as dependent measure reflecting endophenotypic-like group differences and correlating between siblings (Rommelse et al., 2008a).

**Digit Span**

The Digit Span backwards of the WISC-III and WAIS-III was used to obtain an indication of verbal working memory (Rommelse et al., 2008a). The task consisted of repeating a sequence of numbers in the opposite order. Children were instructed to reproduce sequences as accurately as possible. One digit was added to the sequence if a child reproduced the sequence successfully. Two practice trials with a 2 digit sequence and (dependent on the child’s performance) a maximum of 8 experimental sequences were administered. Dependent measure was the maximum Digit Span backwards, which proved useful as endophenotypic candidate (Rommelse et al., 2008a).

**Pursuit**

This task was designed to measure precision of motor control under continuous adaptation (Rommelse et al., 2007c). The stimulus consisted of a randomly moving target (asterisk) that was required to be ‘caught’ by moving a mouse cursor on top of the asterisk. The target moved at a constant speed of 10 mm/s. Children were instructed to ‘catch’ the randomly moving target as precisely as possible. One practice (13 s) and one experimental session (60 s) were
administered for both hands separately. Administration took about 5 minutes. The dependent measure was the precision (mean distance in mm between target and cursor calculated per second and averaged across the 60 s experimental session) of the left hand. Previous results have shown that mainly the performance of the left hand was most strongly associated with ADHD (Rommelse et al., 2007c).

**Tracking**
This task aimed to measure precision of motor control without continuous adaptation required (Rommelse et al., 2007c). The stimulus consisted of an inner and outer circle (radius 7.5 and 8.5 cm, respectively). Children were instructed to trace an invisible midline (radius 8 cm) between the inner and outer circle as quickly and precisely as possible with a mouse cursor. One practice and one experimental session were administered for both hands separately (clockwise with the right hand and counter clockwise with the left hand). Administration took about 3 minutes. The dependent measure was the precision (mean distance to midline in mm averaged across 60 equal parts of the circle) of the left hand. Previous results have shown that precision of the left hand showed endophenotypic-like characteristics (Rommelse et al., 2007c).

**Tapping**
This task measured variability of self-generated motor output (Rommelse et al., 2007d). Children were required to tap as frequently as possible within a certain time period. During tapping, the number of taps was continuously counted and displayed on the screen. One practice session (5 s) and one experimental session (18 s) were administered for both hands separately. The task was first practised and executed with the index finger of the non-preferred hand, thereafter practised and executed with the index finger of the preferred hand. Administration took about 3 minutes. The dependent measure was the variability (SD of intertap intervals in ms) averaged across hands. Previous results have shown that this measure correlates between siblings (Rommelse et al., 2007d).

**Baseline Speed**
This task was designed to measure variability on a simple reaction time task (Rommelse et al., 2007d). Stimuli consisted of a fixation cross in the centre of a computer screen that changed unpredictably into a white square. Immediately following the response, the white square changed back into the fixation cross. The time interval between a response and the emergence of the next white square varied randomly between 500 to 2500 ms in order to prevent anticipation strategies.
Subjects were required to press a key as quickly as possible when the white square appeared. A practice session (10 trials) and an experimental session (32 trials) were administered for both hands separately. The task was first practised and executed with the index finger of the non-preferred hand, thereafter practised and executed with the index finger of the preferred hand. Administration took about 5 minutes. Dependent measure was the variability (SD of reaction times in ms) of responses averaged across hands. Previous results have shown that this measure was associated with ADHD and correlates between siblings (Rommelse et al., 2007d).

Motor Timing

This task was designed to measure variability of motor timing (Rommelse et al., 2007d). In this task a 1 s interval had to be produced. The start of the interval was announced by a tone (80 db, 50 ms). After the subject’s response, visual feedback was given, indicating whether the response was correct, too short or too long. A response was regarded as correct, if it fell between the lower and upper boundary set by a dynamic tracking algorithm. Boundaries were set at 500 to 1500 ms at the beginning of the task. If the response fell within these boundaries, the boundaries for the subsequent trial were narrowed by 100 ms. Likewise, the boundaries of the subsequent trial were widened with 100 ms, if the response on the previous trial fell outside those boundaries. Subjects were instructed to produce as accurately as possible the 1 s interval. Twenty practice trials and 80 experimental trials were administered. Both sessions were preceded by presenting 10 times a cartoon figure for exactly 1 s on the screen to demonstrate the duration of 1 s. Administration took about 8 minutes. The dependent measure was the variability (SD of productions in ms). Previous results have shown this measure to be a viable endophenotypic candidate (Rommelse et al., 2007d).

Procedure

Administration of the neuropsychological tasks in children with ADHD and their siblings took place at the VU University Amsterdam or at the Radboud University Nijmegen Medical Centre and was conducted simultaneously for all children in a family. Psychostimulants were discontinued for at least 48 hours before testing took place (Pelham et al., 1999). Children were motivated with small breaks. At the end of the session, a gift worth approximately € 4.- was given. Control children were tested in a similar way in a quiet room at their school. The study had medical-ethical approval.
Data Analyses

Measures with less than 5% missing data were subjected to expectation maximization to replace the missing data (Tabachnick & Fidell, 2001). Percentages of missing data of the Stop Task (9.2%) and GMO (8.3%) were slightly too large to replace. All measures were subjected to a Van der Waerden transformation (Lehmann, 1975) to normalize the measures and to depict all measure on the same scale (z-scores) (see Figure 10.1). Some measures were mirrored, so that the scores of all variables would imply the same: Higher scores are indicative of poor performance or reflect more severe comorbid problems. In order to obtain a reduced number of neuropsychological variables and a more robust neuropsychological construct, a principal component analysis was performed on the ten neuropsychological task variables. All ten task measures were related to one major component, explaining 47% of the task variance. Additional components did not have an eigenvalue greater than 1 and each of the additional components explained 10% or less additional variance. Therefore, the following results report only on the main factorial component, which is labelled ‘the ADHD-endophenotype’.

Analyses were performed in SAGE (Statistical Analysis for Genetic Epidemiology version 5.4, 2007), which corrects for relatedness of measurements since more than one child per family participated. The following two terms were used: correlation (referring to a correlation between two variables in the same subject), and sibling cross-correlation (referring to a correlation between siblings for two different variables). Correction for multiple comparisons was applied using the False Discovery Rate (FDR) controlling procedure with a q-value of 0.05 (Benjamini & Hochberg, 1995). Correlations were interpreted according to Cohen’s guidelines (Cohen, 1988): 0.10-0.29 (small/modest), 0.30-0.49 (medium/moderate), and 0.50-1.0 (large/strong).

Concerning the first research aim, correlations were calculated between ADHD and the comorbid problems. Correlation coefficients were compared using dependent correlation two-sided t-tests (Chen & Popovich, 2002). Concerning the second research aim, correlations and sibling cross-correlations were calculated between the ADHD-endophenotype and the measures of comorbid problems in order to examine whether the ADHD-endophenotype was related to comorbid problems. The partial correlations and sibling cross-correlations were calculated, allowing investigation of whether or not the association between the ADHD-endophenotype and the comorbid problems was present independent of ADHD. Partial correlations between endophenotype and comorbid problems were calculated by adjusting for the correlations endophenotype-ADHD and ADHD-comorbid problem. Partial sibling cross-correlations between endophenotype and comorbid problems were calculated by correcting for the sibling cross-
correlations endophenotype-ADHD and ADHD-comorbid problem. Regarding the third research aim, correlations between the ADHD-endophenotype and the interaction score between ADHD and comorbid problem were calculated to test whether an endophenotype related to a qualitatively different phenotype.

**Results**

**Degree of Association between ADHD and Comorbid Problems**

Correlations indicated a strong association between ADHD and oppositional defiant behaviors ($r = .72$, $p < .001$) and autistic traits ($r = .68$, $p < .001$). A medium association was found with motor coordination problems ($r = .51$, $p < .001$), anxiety ($r = .45$, $p < .001$), and reading problems ($r = .38$, $p < .001$). The correlation between ADHD and oppositional behaviors was nominally significantly stronger than the correlation between ADHD and autistic traits ($t = 2.22$, $p = .03$), and significantly stronger than the correlation between ADHD and motor coordination problems, anxiety, and reading problems ($t = 8.49$, 10.73, and 11.99, respectively, $p < .001$). The correlation between ADHD and autistic traits was stronger than the correlations of ADHD with motor coordination problems, anxiety, and reading problems ($t = 7.28$, 8.87, and 10.86, respectively, $p < .001$). Motor coordination problems were significantly more correlated with ADHD compared to reading problems ($t = 4.04$, $p = .001$), but not compared to anxiety ($t = 1.92$, $p = .06$). The difference between the correlations of anxiety and reading problems with ADHD did not survive correction for multiple testing ($t = 1.98$, $p = .05$).

**Relation between the ADHD-Endophenotype and Comorbid Problems**

The ADHD-endophenotype was modestly to moderately correlated with all comorbid problems. Following adjustment for ADHD, the correlations between the ADHD-endophenotype and oppositional defiant behaviors and anxiety did not survive statistical correction (see Table 10.1). Correlations between the ADHD-endophenotype and autistic traits, motor coordination problems and reading problems survived adjustment for ADHD. The ADHD-endophenotype cross-correlated modestly to the comorbid problems (except anxiety) in siblings, also after adjustment for ADHD.
Figure 10.1 Normalized and standardized measures of ADHD, comorbid problems, and the ADHD-endophenotype (aggregated score of ten neuropsychological task variables).
Figure 10.1 Continued Normalized and standardized measures of ADHD, comorbid problems, and the ADHD-endophenotype (aggregated score of ten neuropsychological task variables).
Table 10.1 Correlations and sibling cross-correlations between the ADHD-endophenotype and comorbid problems

<table>
<thead>
<tr>
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<th>Correlations $^a$</th>
<th>Sibling cross-correlations $^b$</th>
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<tbody>
<tr>
<td></td>
<td>$r \backslash r_p \text{,}^\gamma$</td>
<td>$r \backslash r_p \text{,}^\delta$</td>
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<tr>
<td>Oppositional defiant behavior</td>
<td>.17 \ .03</td>
<td>.12 \ .11</td>
</tr>
<tr>
<td>Anxiety</td>
<td>.12 \ .03</td>
<td>.05 \ .04</td>
</tr>
<tr>
<td>Autistic traits</td>
<td>.33 \ .26</td>
<td>.21 \ .20</td>
</tr>
<tr>
<td>Motor coordination problems</td>
<td>.41 \ .36</td>
<td>.14 \ .13</td>
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<tr>
<td>Reading problems</td>
<td>.37 \ .32</td>
<td>.11 \ .32</td>
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Note.

$^a$ Correlation endophenotype-comorbid problem measured in the same subject based on 816 children (affected children, non-affected siblings, and controls) corrected for the non-independency of children.

$^b$ Correlation endophenotype-comorbid problem measured in two different subjects (siblings) based on 540 sibling-pairs (including affected children, non-affected siblings, and controls) corrected for the non-independency of sibling pairs.

$^\gamma r_p = $ Partial correlation endophenotype-comorbid problem corrected for the correlations endophenotype-ADHD and ADHD-comorbid problem.

$^\delta r_p = $ Partial sibling cross-correlation endophenotype-comorbid problem corrected for the sibling cross-correlations endophenotype-ADHD and ADHD-comorbid problem.

**Bold** correlations are significant after correction for multiple testing.

ADHD-Endophenotype in Relation to ADHD with Comorbid Problems

In order to test whether the ADHD-endophenotype was related to a qualitatively different phenotype, we analysed whether the ADHD-endophenotype correlated with the interaction term between ADHD and the comorbid problem. If the combination of both problem domains is similar to the additive effect of both disorders, the interaction term will not relate significantly to the ADHD-endophenotype, indicating the combination of both disorders differs only quantitatively from the individual problem domains. If, however, the combination of both disorders is different from the additive effect of both disorders, the interaction term will relate significantly to the ADHD-endophenotype, indicating the combination of both disorders differs qualitatively from the individual problem domains. The ADHD-endophenotype did not significantly correlate with any of the interaction terms (with correlations ranging from -.03 to .04 and $p$-values ranging from .44 to .91).
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Discussion

The aim of this study was threefold. First, to examine the degree of association between ADHD and comorbid problems (oppositional defiant behaviors, anxiety, autistic traits, motor coordination problems, and reading problems) in order to determine which comorbid problems were most important in relation to ADHD. Second, to examine whether an aggregated score of previously investigated candidate neuropsychological ADHD-endophenotypes was related to comorbid problems (with and without adjusting for their interdependence on ADHD). Third, to determine whether the ADHD-endophenotype related to qualitatively different phenotypes if ADHD co-occurred with comorbid problems.

With respect to the first aim of our study, the various comorbid problems were all significantly associated with ADHD. That is, having a more severe form of ADHD was related to having more severe oppositional defiant behaviors, higher levels of anxiety, more autistic traits, and more severe motor coordination and reading problems, which concurs with previous studies reporting on various disorders in combination with ADHD (Angold et al., 1999; Bauermeister et al., 2007; Gillberg et al., 2004). Oppositional defiant behaviors and autistic traits were most strongly correlated with ADHD severity compared to the other investigated comorbid problems. The former finding is well established in literature (Angold et al., 1999; Bauermeister et al., 2007; Gillberg et al., 2004; Jensen et al., 1997). Since oppositional defiant behaviors with ADHD may significantly influence the type of treatment that can be best prescribed (Hechtman et al., 2005; Jensen et al., 2001), it seems important to assess oppositional defiant behavior in ADHD-patients. The latter finding (ADHD in combination with autistic traits) is still controversial, since the DSM-IV does not support a diagnostic combination of ADHD and autistic disorder (Corbett & Constantine, 2006; Goldstein & Schwebach, 2004). This may need to be reconsidered when developing the future DSM, since our findings and numerous previous findings suggest that ADHD is frequently associated with characteristics of autistic disorder (Corbett & Constantine, 2006; Goldstein & Schwebach, 2004; Leyfer et al., 2006). Furthermore, ADHD correlated moderately with motor coordination problems and reading problems, although the correlation between ADHD and motor coordination problems appeared stronger than the correlation between ADHD and reading problems. Since both motor coordination problems and reading problems can significantly interfere with academic functioning in a child that is already vulnerable to academic failure because of ADHD symptoms (Rasmussen & Gillberg, 2000), it is important to determine vulnerabilities of motor coordination and reading in children in ADHD assessment. Anxiety was moderately correlated with ADHD, to a comparable degree as motor
coordination problems and reading problems. Anxiety as internalizing disorder may be an easily overlooked comorbid problem in children with ADHD given the externalizing nature of ADHD symptoms. However, since comorbid anxiety may influence response to methylphenidate (Pliszka, 1989; Tannock et al., 1995) and is predictive of a range of psychiatric disorders in adolescence (Bittner et al., 2007), assessment of anxiety in children with ADHD is important (Angold et al., 1999; Bauermeister et al., 2007; Gillberg et al., 2004).

Second, comorbid problems were not merely associated phenotypically, but an aggregated score of previously investigated candidate neuropsychological ADHD-endophenotypes (Rommelse et al., 2007a, b, c, d, 2008a) was also modestly to moderately related to these disorders. The severity of the comorbid problems could be predicted to some degree by the endophenotypic dysfunction. It may be argued, however, that these correlations may be explained by the mutual dependence of the ADHD-endophenotype and the comorbid problems with ADHD. However, even when this dependence was controlled for, almost all associations between the ADHD-endophenotype and comorbid problems remained significant (except for oppositional defiant behaviors and anxiety), suggesting that the ADHD-endophenotype is related independently of ADHD to the majority of comorbid disorders. These findings may pinpoint to a shared underlying general neuropsychological dysfunction that may give rise to both ADHD and several associated domains. Further support for this hypothesis was found, when the sibling cross-correlations between the ADHD-endophenotype and the comorbid problems were calculated and were almost all found to be significant (except for anxiety). This suggests that most comorbid problems may arise from similar neuropsychological endophenotypes that are related to ADHD. It appears that at least a part of the relationship between ADHD and comorbid problems relies on common neuropsychological endophenotypes that may have multiple behavioral consequences, pleiotrophy (Banaschewski et al., 2005). The chain of events leading to ADHD symptoms does not appear to be independent of the sequence of events leading to other domains. Atypical brain development may be the basis of developmental disorders, resulting in frequent co-occurrence of multiple disorders in the same child (Kaplan et al., 2001). No such evidence was found for anxiety in relation to ADHD, however, suggesting the occurrence of anxiety disorders in children with ADHD may be merely phenotypical.

Results for the third aim of our study revealed that the ADHD-endophenotype did not relate to the interaction term between ‘ADHD scores’ and ‘comorbid scores’. This suggested that larger endophenotypic dysfunctions resulted in ‘more of the same’ (a more severe form of ADHD and comorbid problem) but not in a phenotypically distinct subtype (Banaschewski et al.,
2005; Biederman et al., 1991; Caron & Rutter, 1991; Jensen et al., 2001). Based on these results, it does not appear necessary to define new diagnostic categories in a future DSM entailing ADHD with specific comorbid problems.

Several limitations of this study warrant consideration. First of all, questionnaires were used to gather information on comorbid problems. Although questionnaires may more aptly reflect the underlying continuously distributed nature of disorders, they do not provide sufficient information to diagnose a child according to DSM-IV criteria. Additional measures that might have allowed us to diagnose all children for comorbid problems would have added to the study findings. A second limitation that should be noted, is that our sample was biased towards having ADHD (or being biologically related to someone who does) and biased towards not having ADHD. Although the whole range of ADHD severity and severity of comorbid problems was represented in our sample, the current findings on the associations between the comorbid problems may, therefore, not be applicable to the occurrence of these symptoms in the general population. Third, the ADHD-endophenotype used in this study was defined as being a single component derived from measures, previously shown to be candidate ADHD-endophenotypes (Rommelse et al., 2007a, b, c, d, 2008a). It is unclear which neuropsychological process or processes are likely to be tapped by this overall factor or how this factor could account for ADHD or the comorbid problems. However, it may be hypothesized that this factor represents general cognitive functioning in combination with variability of reacting. This factor appears to be a powerful endophenotypic construct and we hope to clarify in future studies what genetic and neurobiological factors may underlie this construct. A fourth limitation is related to the fact that children with a full autistic disorder were excluded from the study, since these children can not be diagnosed with ADHD according to the DSM-IV. Our findings on autistic traits are thus limited to children with sub clinical autistic disorder.

Understanding the presence of comorbidity between psychiatric conditions offers a mean of correcting and validating psychiatric nosology (Angold et al., 1999). The co-occurrence of ADHD and comorbid problems is evident and arises partly from shared neuropsychological endophenotypic dysfunctions, suggesting these symptoms can not be viewed, diagnosed or treated independently of one another. This may have implications for the theoretical background on which the development of a future DSM as well as for the diagnostic and treatment procedures utilized in daily clinical practice. With respect to the latter, these findings suggest always assessing possible comorbid conditions in addition to ADHD, even when symptoms of ADHD appear the most prominent. It appears, however, that new diagnostic categories in a future DSM entailing ADHD with specific comorbid problems are not necessary.