Gene-Environment Interplay in the Association Between Pubertal Timing and Delinquency in Adolescent Girls

K. Paige Harden and Jane Mendle
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Gene-Environment Interplay in the Association Between Pubertal Timing and Delinquency in Adolescent Girls

K. Paige Harden
University of Texas

Jane Mendle
University of Oregon

Early pubertal timing places girls at elevated risk for a breadth of negative outcomes, including involvement in delinquent behavior. While previous developmental research has emphasized the unique social challenges faced by early maturing girls, this relation is complicated by genetic influences for both delinquent behavior and pubertal timing, which are seldom controlled for in existing research. The current study uses genetically informed data on 924 female-female twin and sibling pairs drawn from the National Longitudinal Study of Adolescent Health to (1) disentangle biological versus environmental mechanisms for the effects of early pubertal timing and (2) test for gene-environment interactions. Results indicate that early pubertal timing influences girls’ delinquency through a complex interplay between biological risk and environmental experiences. Genes related to earlier age at menarche and higher perceived development significantly predict increased involvement in both nonviolent and violent delinquency. Moreover, after accounting for this genetic association between pubertal timing and delinquency, the impact of nonshared environmental influences on delinquency are significantly moderated by pubertal timing, such that the nonshared environment is most important among early maturing girls. This interaction effect is particularly evident for nonviolent delinquency. Overall, results suggest early maturing girls are vulnerable to an interaction between genetic and environmental risks for delinquent behavior.

Keywords: delinquency, puberty, behavioral genetics, twins, gene-environment interaction
Potential Biological Mechanisms for the Effects of Early Pubertal Timing

In order to specify more precisely the pathways by which early pubertal maturation confers risk for antisocial behavior, it is necessary to acknowledge the importance of genetic influences on girls’ pubertal timing. While it is not yet fully known why some girls begin puberty earlier than others, girls’ “inmate developmental clock” appears to be under strong genetic control. Behavior genetic studies report that correlations for age at menarche are consistently higher among MZ than among DZ twins (Doughty & Rodgers, 2000; Meyer, Eaves, Heath, & Martin, 1990; Rowe, 2000; Treloar & Martin, 1990), with approximately 61–68% of the variance in menarche accounted for by genetic effects. Likewise, estimates for the heritability of secondary sexual characteristics, such as breast development or skin changes, range from .50 (Ge, Natsuaki, Neiderhiser, & Reiss, 2007) to .88 (Mustanski, Viken, Kaprio, Pulkkinen, & Rose, 2004). Molecular genetic studies have also identified specific genes involved in pubertal timing. Most notably, the GPR54 gene has been shown to influence the initial secretion of the gonadotropin releasing hormone (GnRH), which is necessary for gonadal maturation (Seminara et al., 2003). However, understanding of the genetic basis of pubertal timing is incomplete. As Sisk and Foster (2004) describe, “Identification of master regulatory genes directing the unique maturational components of the first and most important transition to fertility remains an unsolved part of the puberty mystery” (p. 1042).

In addition, it is well-established that antisocial behavior is heritable (e.g., Arsenault et al., 2003; D’Onofrio et al., 2007; Scourfield, Van den Bree, Martin, & McGuffin, 2004; Slutske et al., 1997; Young et al., 2002; for reviews see Miles & Carey, 1997; Raine, 2002; Rhee & Walman, 2002; Rowe, 2001). Although there have been few empirical tests of whether genes related to early pubertal timing overlap with those for delinquency (Mendle et al., 2007), common genetic influences have been implicated in the association between delinquency and fertility relevant outcomes strongly correlated with pubertal timing (Udry, 1979), including age of first sexual intercourse (Harden, Mendle, Hill, Turkeheimer, & Emery, 2008), risky sexual behavior (Verweij, Zeitsch, Bailey, & Martin, 2009), and adolescent childbearing (Harden et al., 2007). In addition, shorter alleles of the X-linked androgen receptor (AR) gene have been associated with aggression and impulsivity in males and with earlier ages of physical maturation in females (Comings et al., 2002).

Emerging research on adolescent brain development has suggested a novel explanation for why genetically influenced differences in pubertal timing may be important for girls’ delinquency. In addition to a cascade of endocrine events that culminate in reproductive maturity, puberty also involves a cascade of neural changes, a “second period of structural reorganization and plasticity in the brain” (Blakemore, Burnett, & Dahl, 2010). One key change initiated by the hormonal events at puberty is the remodeling of neural circuits involved in reward-motivated behavior, particularly in the striatum, nucleus accumbens and in dopaminergic pathways to the prefrontal cortex (Blakemore et al., 2010; Forbes & Dahl, 2010; Kuhn et al., 2010). Female pubertal hormones (most notably, estradiol and progesterone) have significant effects on the anatomy of dopamine systems relevant for reward-motivated behavior (Kuhn et al., 2010). Moreover, pubertal development, independent of chronological age, has been found to be associated with changes in activity in the striatum and medial and ventrolateral prefrontal cortex in response to monetary rewards and social threats (Forbes, Phillips, Ryan, & Dahl, in press; Forbes et al., 2010). This is congruent with findings that pubertal development and pubertal hormones, independent of age, are linked to changes in the personality trait of sensation-seeking, which has been conceptually linked to responsiveness to rewards and novelty (Martin, Kelly, Rayens, Brogl, Brenzel, Smith, & Omar, 2002; Martin, Logan, Leukefeld, Milich, Omar, & Clayton, 2001; Steinberg, Albert, Cauffman, Banich, Graham, Woolard, 2008; Zucker- man, Buchsbaum, & Murphy, 1980).

Because the timing of pubertal development is strongly influenced by genes, early maturing girls can thus be thought of as having genetic propensities for early neurobiological change. Moreover, for early maturing girls, these puberty-linked neurobiological changes are asynchronous with age-based neurobiological changes in cortical structures that underlie behavioral inhibition and effortful control. As articulated by the dual systems model of adolescent brain development (Casey, Getz, & Galvan, 2008; Somerville, Jones, & Casey, 2010; Steinberg, 2008), this “maturity gap” between the development of subcortical versus cortical brain regions results in an elevated propensity for adolescents—particularly early maturing adolescents—to engage in delinquent behavior, and in risk-taking behavior more generally.
The Social Challenges of Early Pubertal Timing

In contrast, the prevailing view in the developmental literature emphasizes the unique environmental dilemmas posed by early pubertal timing. The most widely accepted theoretical explanation for the relation between pubertal timing and delinquency, as well as a breadth of other psychosocial problems more common among girls with early pubertal timing, is the maturation disparity hypothesis (reviewed in Ge & Natsuaki, 2009). This theory stresses that a girl’s changing physical appearance during puberty creates new social experiences that may promote delinquent behavior, including increased autonomy from parents, increased romantic and sexual attention from males, and increased affiliation with older peer groups (Caspi et al., 1993; Ge, Brody, Conger, Simons, & Murry, 2002). Early matures are believed to be particularly vulnerable to the psychosocial risks posed by these new experiences, because they are forced to navigate the changing social landscape with fewer emotional or cognitive resources than girls who reach the same developmental milestones at a later chronological age. Moreover, theories specific to adolescent delinquency, most notably Moffitt’s (1993) taxonomy of adolescent-limited delinquency, emphasize the lack of rights and privileges granted by adults to physically mature adolescents. Caught between a mature physical appearance and an immature social status, adolescents are believed to engage in rebellious activities designed to demonstrate independence (i.e., running away from home, lying to parents, truancy), imitate adult actions (i.e., sexual activity, drinking), or showcase lack of concern for adult-enforced social rules (i.e., larceny, property damage; Moffitt, 1993; Moffitt et al., 1996; Moffitt, Caspi, Rutter, & Silva, 2001).

In addition to experiencing heightened environmental stress during the pubertal transition, early maturing girls are also more likely to have experienced prior environmental stress in early childhood. In particular, early childhood stressors such as father absence (Ellis, 2004); child maltreatment (Bergen, Bukowski, & Karasov, 2003; Turner, Runz, & Galambos, 1999; Wise, Palmer, Rothman, & Rosenberg, 2009); harsh parenting (Belsky et al., 2007); and poverty (Obeidallah et al., 2000) are all correlated with a comparatively precocious timing of development. While these associations have been interpreted in light of Belsky, Steinberg, and Draper’s (1991) evolutionary hypothesis, it is important to note that this line of research has been strongly criticized by behavioral geneticists for interpreting correlations between parent characteristics and child outcomes as due to purely environmental mechanisms, without considering the role of genetic transmission from parent to child (Rowe, 2002). In our previous genetically informed research on this topic, we have presented evidence that early family structure (father absence and step-fathering) are associated with accelerated pubertal and sexual development via genetic transmission rather than an environmental effect (Mendle et al., 2006, 2009). Nevertheless, regardless of their causal status, adverse home environments are certainly correlated with early pubertal development, and these early environmental stressors may partially account for the elevated rates of antisocial behavior problems seen in early maturing girls.

Integrating Biological and Social Risk:
Gene-Environment Interaction

Early pubertal timing is not only associated with increased exposure to environmental risks for delinquency, but early maturing girls may also be more sensitive to environmental risks. In support of this, social context has been shown to be a critical determinant of resiliency versus vulnerability for early developers. For example, Caspi, Lynam, Moffitt, and Silva (1993) found that early maturing girls were at elevated risk for delinquency only if they attended mixed-sex schools versus same-sex schools. They attributed this interaction effect to greater exposure to male peers and male romantic partners. Likewise, Natsuaki, Biehl, & Ge (2009) found that engagement in a romantic relationship was associated with greater psychological distress for early matures rather than their peers. Finally, pubertal timing has been found to interact with characteristics of the parent–child relationship and with neighborhood characteristics, with early maturing girls showing the highest involvement in delinquent behavior when maternal nurturance and parental monitoring were low (Mrug et al., 2008) and when concentrated neighborhood disadvantage was high (Obeidallah, Brennan, Brooks-Gunn, & Earls, 2004).

Considering the biological and environmental risks faced by early maturing girls, it is possible that early pubertal timing may increase delinquent behavior via a heightened sensitivity to environmental context, also known as gene × environment interaction (G × E). Only one previous study has tested this hypothesis. Burt, McGue, DeMarte, Krueger, & Iacono (2006) found compelling evidence for G × E interaction in the association between timing of menarche and conduct disorder symptoms: Among girls with an age of menarche of 11 years or earlier, only 8% of the variance in conduct disorder symptoms could be attributed to genes and the remaining 92% was attributable to environmental influences. In contrast, for girls who experienced an average menarcheal timing (12–13 years), the majority of the variance in conduct disorder symptoms (67%) was attributable to genes and only 33% attributable to the environment. These findings are consistent with more general observations from the behavior genetics literature that heritability is not static; rather, genetic propensities may be suppressed or augmented in certain social circumstances (e.g., Harden, Hill, Emery, & Turkheimer, 2008; Harden, Turkheimer, & Loehlin, 2007).

Sensitivity to the unique environmental challenges of puberty may be particularly pertinent for nonviolent, rule-breaking forms of delinquent behavior (such as property crime). In contrast, violent or aggressive delinquency differs behaviorally, developmentally, and etiologically from rule-breaking (Achenbach, 1991, 2001; Barker et al., 2009; Eley, Lichtenstein, & Stevenson, 1999; Loeb, Burke, & Pardini, 2009; Moffitt, 1993; Tackett, Krueger, Sawyer, & Graetz, 2003; Tuvblad, Eley, & Lichtenstein, 2005). Adolescents with histories of violent behavior show poor executive functioning, verbal processing, and neuropsychological impairment (Barker et al., 2007; Déry, Toupin, Paucé, Mercier, & Fortin, 1999), while results from neuroimaging studies confirm that adolescents with high levels of violent behavior display reduced activation in the frontal and temporal cortices, compared to normal controls, when watching pain inflicted on another person (Decety, Michalska, Akitsuiki, & Lahey, 2009). Similar decrements have been observed in adults with histories of physical aggression.
(e.g., Siever, 2008; Volkow et al., 1995). There deficits are believed to be rooted in heritable predispositions (Moffitt, 1993), and a broad array of candidate genes have been identified as playing a role in violent behavior (Brunner et al., 1993; Davridge et al., 2004; Flory et al., 2007; Meyer-Lindentag et al., 2006; Mik et al., 2007). In contrast, nonviolent forms of delinquency are developmentally transient (Stanger et al., 1997), show less genetic stability (Eley, Lichtenstein, & Moffitt, 2003; Burt & Neiderhiser, 2009) and are more strongly influenced by environmental factors (Burt, 2009).

Measurement of Pubertal Timing

Lastly, the relative importance of biological versus environmental pathways for the relation between pubertal timing and delinquent behavior may depend, in part, on how the construct of pubertal timing is measured and operationalized. Typically, assessing pubertal stage using a physical examination by a trained physician or nurse has been considered the “gold standard” of objective measurement; however, physical exams are infrequently used, particularly in large-scale epidemiological research. Adolescent self-report of age at menarche is a far more common method for measuring pubertal timing, because of its relative ease of use. Test–retest reliability coefficients for self-report of menarche have been found to be good (.61–.81; Dorn, Dahl, Woodward, & Biro, 2006). Moreover, adults’ retrospective reports of menarche are quite accurate as compared to childhood medical records, even after spans of up to 40 years (Bean, Leeper, Wallace, Sherman, & Jagger, 1979; Casey et al., 1991).

A third strategy, which was developed as an alternative to physical exam, is to query adolescents regarding changes in breast size and other secondary sex characteristics (perceived development; e.g., Biehl, Natsuaki, & Ge, 2007; Natsuaki, Biehl, & Ge, 2007; Graber, Lewinsohn, Seeley, & Brooks-Gunn, 1997; Graber, Seeley, Brooks-Gunn, & Lewinsohn, 2004; Wichstrom, 2000). Measures of perceived pubertal development have only modest agreement with more objective measures of pubertal timing, such as menarcheal age or Tanner stages as determined by physical exam (Dorn et al., 2006). Nevertheless, girls’ perceptions of themselves as physically mature, regardless of the accuracy of those perceptions, may be salient for their risk for engaging in delinquent behavior. In particular, girls who perceive themselves as more advanced in their pubertal development may also perceive a wider “maturity gap” until they gain full adult status (Moffitt, 1993), which may provoke additional delinquent behavior. A limited number of studies have evaluated the relative impact of objectively early maturation versus girls’ perceptions of themselves as early maturers (Brooks-Gunn and Warren, 1985; Brooks-Gunn, Attie, Burrow, Rosso, & Warren, 1989; Michael & Eccles, 2003; Rierdan et al., 1988). These studies have consistently found that girls who define themselves as early maturers—even if their development is objectively on-time or even late—are more likely to exhibit adverse developmental outcomes. Thus, both “objective” early pubertal timing (e.g., menarche) and girls’ “subjective” perceptions of early pubertal development may result in increased antisocial behavior, but no previous research has examined both objective and perceived pubertal timing while also attempting to disentangle biological versus social processes.

Goals of the Current Study

Collectively, previous research indicates that, while the relation between pubertal timing and girls’ delinquent behavior is well-established, the underlying mechanisms remain to be elucidated. The present study uses a behavioral genetic approach to this question. We utilize two different measures of pubertal timing, perceived development and menarcheal age, to predict both nonviolent and violent delinquency in a sample of sister dyads from the National Longitudinal Study of Adolescent Health (Add Health). The utility of the behavioral genetic approach is twofold. First, comparisons between biological relatives allow us to discriminate between genetic and environmental pathways of risk: If Twin A matures earlier than Twin B, does she also show higher levels of delinquent behavior? A significant within-twin pair association, which controls for all genetic factors shared by twins, would indicate that environmental differences (rather than genetic differences) in pubertal timing are important pathways of risk for delinquency. Second, recent developments in statistical methods for twin data allow us to test hypothesis about gene-by-environment interaction: Are environmental or genetic influences on delinquency stronger among early maturing girls?

Method

Participants

Data are drawn from the National Longitudinal Study of Adolescent Health (Add Health; Udry, 2003), a study of health behaviors in a nationally representative study of adolescents. The Add Health study used stratified random sampling of U.S. high schools (79% of targeted schools agreed to participate). From the participating schools, 90,118 students completed a confidential in-school survey during the 1994–1995 academic year. School rosters were then used to randomly select a sample of over 20,000 adolescents (10,480 females; 10,264 males) to complete a comprehensive, 90-min, in-home interview between April and December 1995. Participants ranged in age from 11 to 21 years, M = 16 years, 25th percentile = 14 years, 75th = 17 years. There have been three follow-up interviews with the Add Health participants: Wave II in 1996, Wave III in August, 2001-2002, and Wave IV in 2007–2008.

The Add Health study deliberately oversampled sibling pairs, even if one member of the pair did not attend a high school in the original probability sample. The current study only uses female-female sibling pairs; thus, our subsample includes 1848 females from 924 sibling pairs: 145 monozygotic (MZ) twin pairs, 116 dizygotic (DZ) twin pairs, 369 full sibling pairs (FS), 117 half-sibling pairs (HS), 112 pairs of cousins raised in the same household (CO), and 65 nonbiologically related pairs (e.g., step-siblings or adoptive siblings; NR). Twin zygosity was determined primarily on the basis of self-report and responses to four questionnaire items concerning similarity of appearance and frequency of being confused for one’s twin. Similar questionnaires have been utilized widely in twin research and have been repeatedly cross-validated with zygosity determinations based on DNA (e.g., Loehlin & Nichols, 1976; Spitz et al., 1996). Race/ethnicity were classified as either White, N = 984, 53.2%; African American, N = 493, 26.7%; Hispanic, N = 245, 13.3%; or Other, including Asian American and Native American, N = 126, 6.8%. The mean age was 16.12 years, SD = 1.67 years.
Measures

Pubertal timing. In the in-home interview for both waves I and II, participants were asked whether they had ever had a menstrual period, and if so, which month and year it first occurred. At Wave III, participants were asked, “How old were you when you got your period for the first time?” From these waves of data, age at menarche was constructed as follows: If the participant reported menarche occurring before Wave I, the Wave I report of age at menarche was used (86.8% of participants). If the participant reported menarche occurring between Waves I and II, the Wave II report of age at menarche was used (8.2% of participants). Lastly, if participants reported getting menarche between Waves II and III, the Wave III report was used (2.7%). Menarche data was missing for 23 individuals (1.2% of the sample). The mean age of menarche in the sample was 12.23 years, SD = 1.42, Range: 7.0 years – 19.0 years. African American girls, on average, experienced menarche at a significantly younger age (10.9 years – 19.0 years. African American girls, on average, experienced menarche at a significantly younger age (12.23 years, SD = 1.44, Age = 1.43, p < .01; based on a mixed effects model that accounted for clustering of girls within sibling pairs) than White girls, M = 12.34, SD = 1.44. Hispanic girls also trended toward younger mean ages at menarche, M = 12.14, SD = 1.37, although this difference was not significant, p = .08, in the sibling pairs subsample.

In addition, at Wave 1, participants’ perceptions of their level of pubertal development were assessed using two items about breast size [1 = “My breasts are about the same size as when I was in grade school” to 5 = “My breasts are a whole lot bigger than when I was in grade school; they are as developed as a grown woman’s breasts”] and body curviness [1 = “My body is about as curvy as when I was in grade school” to 5 = “My body is a whole lot more curvy then when I was in grade school”]. Higher scores on these items thus represent a higher perceived pubertal status; to calculate a measure of subjective pubertal timing, we calculated the deviation of each participant’s score from the mean level of development reported by adolescents of the same age and standardized this deviation score, M = 0, SD = 1. Thus, higher scores reflect more advanced perceived development relative to other same-aged adolescents. Age at menarche was measured using a group fight, shot or stabbed someone. Following Guo, Roettger, & Cai (2008), responses to violent delinquency items were also on a 0–3 scale, except for two items (shooting or stabbing someone; pulling a knife or gun on someone). These items were scored as, 0 = Never and 3 = One or more times. Both delinquency scales showed good internal reliability, α = .92 for nonviolent and .77 for violent. We calculated sum scores for both delinquency scales, M nonviolent = 3.27, SD = 4.13; M violent = 2.29, SD = 3.39, and then standardized them, M = 0, SD = 1, by age in years, so that higher scores could be interpreted as more delinquent behavior relative to other same-aged adolescents. Age at menarche was significantly and negatively correlated with both nonviolent delinquency, r = -.09, 95% CI = -.18, -.05, and violent delinquency, r = -.11; 95% CI = -.15, -.02. Perceived development was also significantly correlated with both forms of delinquency, nonviolent: r = .16; 95% CI = .10, .22; violent: r = .11, 95% CI = .04, .17.

Sibling correlations for perceived development, age at menarche, violent, and nonviolent delinquency, by type of sibling pair, are summarized Table 1.

Analytic Plan

Data were analyzed using structural equation modeling in Mplus (Muthen & Muthen, 1998–2007), using Full Information Maximum Likelihood (FIML) to account for missing data. Absolute model fit was assessed using the CLI and the RMSEA. Nested models were compared using differences in log-likelihood, which are distributed as chi-squares.

Table 1
Sibling Pair Correlations and Zero-Order Correlations Among Variables

<table>
<thead>
<tr>
<th>Sibling Pair Correlations</th>
<th>Non-violent delinquency</th>
<th>Violent delinquency</th>
<th>Age at menarche</th>
<th>Perceived development</th>
</tr>
</thead>
<tbody>
<tr>
<td>MZ Twins</td>
<td>.46 (N = 140)</td>
<td>.54 (N = 140)</td>
<td>.61 (N = 139)</td>
<td>.38 (N = 137)</td>
</tr>
<tr>
<td>DZ Twins</td>
<td>.44 (N = 113)</td>
<td>.38 (N = 113)</td>
<td>.31 (N = 111)</td>
<td>.29 (N = 109)</td>
</tr>
<tr>
<td>Full Siblings</td>
<td>.30 (N = 365)</td>
<td>.46 (N = 365)</td>
<td>.29 (N = 362)</td>
<td>.17 (N = 355)</td>
</tr>
<tr>
<td>Half Siblings</td>
<td>.35 (N = 109)</td>
<td>.30 (N = 110)</td>
<td>.18 (N = 109)</td>
<td>.08 (N = 109)</td>
</tr>
<tr>
<td>Cousins</td>
<td>.26 (N = 65)</td>
<td>.22 (N = 65)</td>
<td>.22 (N = 65)</td>
<td>.04 (N = 64)</td>
</tr>
<tr>
<td>Nonbiological Siblings</td>
<td>.20 (N = 111)</td>
<td>.08 (N = 111)</td>
<td>.08 (N = 110)</td>
<td>.04 (N = 107)</td>
</tr>
</tbody>
</table>

Note. MZ = Monozygotic; DZ = Dizygotic. Sample sizes refer to number of pairs. Significant correlations (p < .05) are in bold font.
Data were analyzed in two steps. First, we fit bivariate twin models that tested whether the associations between pubertal timing, as measured by age at menarche and perceived development, and violent and nonviolent delinquency were due to environmental versus genetic pathways. An example of this model, for nonviolent delinquency, is shown in Figure 1. The twin model (see Neale & Cardon, 1992 for full details) typically decomposes variation in a given phenotype into three latent factors: variance due to additive genes ($A$), variance due to environmental influences that make twins and siblings similar to each other, also known as the shared environment ($C$), and variance due to environmental influences that make twins and siblings different from each other, also known as the nonshared environment, plus measurement error ($E$). The correlation between the additive genetic factors for the first and second sibling in each pair are fixed according to genetic theory: 1.0 for MZ twins, 0.5 for DZ twins and full siblings, 0.25 for half siblings, 0.125 for cousins, and 0 for nonbiologically related siblings.

Previous analyses of the Add Health data, which combined menarche and perceived development as indicators of a single construct, found that shared environmental influences on pubertal timing are negligible (Ge et al., 2007). Our reanalysis of the data found that shared environmental influences on menarche were nonsignificant and could be dropped from the model without loss of fit. ACE model: $\chi^2 = 25.41$ (df = 26; $p = .50$); AE model: $\chi^2 = 26.98$ (df = 27; $p = .47$); $\Delta \chi^2 = 1.57$, $\Delta df = 1$, $p = .21$. In contrast, a model without additive genetic influences on menarche did fit significantly worse, CE model: $\chi^2 = 59.14$ (df = 27, $p < .01$); $\Delta \chi^2 = 33.7$, $\Delta df = 1$, $p < .01$. Similarly, shared environmental influences on perceived development were also nonsignificant and could be dropped from the model without loss of fit. ACE model: $\chi^2 = 25.95$ (df = 26; $p = .47$); AE model: $\chi^2 = 26.32$ (df = 27; $p = .50$); $\Delta \chi^2 = 0.37$, $\Delta df = 1$, $p = .54$, while a model without additive genetic influences on menarche did fit significantly worse, CE model: $\chi^2 = 35.16$ (df = 27; $p = .13$); $\Delta \chi^2 = 9.21$, $\Delta df = 1$, $p < .001$. Consequently, the models in the current paper only include genetic and nonshared environmental influences on each measure of pubertal timing.

In the first multivariate model, delinquency was regressed on the $A$ and $E$ factors of each measure of pubertal timing (parameters labeled $b_A$ and $b_E$). The regression on $A$ tests whether higher involvement in delinquent behavior is predicted by genes related to pubertal timing. In contrast, the regression on $E$ essentially compares the following within sibling pairs: Does the twin who has earlier pubertal timing than her co-twin also demonstrate higher delinquency?

Second, we fit bivariate interaction models that tested whether, after controlling for the genetic and environmental associations between pubertal timing and delinquency, early pubertal timing significantly moderated the residual genetic and environmental variance in delinquency. An example of this model, for menarche and nonviolent delinquency, is illustrated in Figure 2 (only one twin per pair shown). This model is identical in form to the bivariate twin model, described previously, except that the paths from the $A$, $C$, and $E$ components of delinquency are moderated by pubertal timing (either age at menarche or perceived development). A significant interaction effect on path from the $E$ component of delinquency would indicate that girls with early pubertal timing are more sensitive to environmental influences.

**Results**

**Is the Association Between Early Pubertal Timing and Higher Delinquency Due to Genetic or Environmental Pathways?**

**Mean comparisons.** In order to illustrate the association between perceived development and delinquency, we first divided the sample into *higher* versus *lower* perceived pubertal development (based on a mean split), and calculated the mean levels of nonviolent and violent delinquency for each group. These mean comparisons are shown in Figure 3. The first set of bars in each plot (labeled *between families comparison*) shows the mean levels of delinquency for adolescents from sibling pairs where both adolescents had higher perceived development versus adolescents from sibling pairs where both adolescents had lower perceived development. These results are consistent with the extant literature on pubertal timing: On average, adolescents who perceive them-
selves to be more mature, relative to their peers’ self-perceptions, show higher levels of both violent and nonviolent delinquency. The second set of bars in each plot (labeled within family comparison) shows the mean levels of delinquency for siblings who are discordant for perceived development. In contrast to the mean difference evident in the between-families comparison, siblings who differ in their perceived pubertal development do not appreciably differ in their involvement in violent or nonviolent delinquency. Because there is no shared environmental variance, this suggests that genetic factors shared by siblings may account for the association between perceived development and delinquency.

Next, we divided the sample into earlier age at menarche (less than 12 years old) and later age at menarche (12 years old or older) groups, and calculated a similar set of between-family and within-family comparisons (illustrated in Figure 4). Again, results from the between-family comparison were consistent with previous research on pubertal timing and delinquency: Girls with an earlier age at menarche demonstrated, on average, higher levels of delinquency. However, this association was sharply attenuated when comparing within sibling pairs who were discordant for earlier versus later age at menarche.

Twin models. Results from the bivariate twin models are summarized in Table 2. Overall, the fit of the bivariate models was fair to good (RMSEAs <0.07). Genetic influences accounted for approximately 41% of the variance in perceived development. Genetic influences for more advanced perceived development, in turn, significantly predicted higher levels of both nonviolent, $b_A = .237$, and violent delinquency, $b_A = .174$. Similarly, genetic influences accounted for 61% of the variance in menarcheal age, and genes related to earlier age at menarche significantly predicted higher levels of nonviolent delinquency, $b_A = -1.09$, but not violent delinquency. After controlling for these genetic associations, the environmental paths between perceived development and delinquency, and between age at menarche and delinquency, were not significant. Overall, results from the bivariate twin models suggest that biological differences in pubertal timing—rather than environmental mechanisms—are responsible for the main effects of pubertal timing on delinquency.

Does Early Pubertal Timing Result in Increased Sensitivity to Environmental Risks?

Results from the best-fitting interaction models are summarized in Table 3. Across all models, a consistent pattern emerged for nonshared environmental influences to be stronger for girls with earlier pubertal timing. For models using perceived development, the $e'$ parameters were significant and positive, $+.110$ for nonviolent delinquency and $+.065$ for violent delinquency, indicating that girls who rated their pubertal development as advanced relative to their peers were more sensitive to the nonshared environment. Similarly, for models using age at menarche, the $e'$ parameters were significant and negative, $-.093$ for nonviolent delinquency and $-.100$ for violent delinquency, indicating that girls with earlier menarche were more sensitive to the nonshared environment.

For models of nonviolent delinquency, there were significant interactions between pubertal timing and genes, such that genetic effects were less influential for girls with higher perceived development, $a' = -.082$, and earlier ages at menarche, $a' = .124$. In addition, there was a significant interaction between age at menarche and perceived development as continuous variables.

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1 Previous research has used various cut-offs to define “early” pubertal maturation; we chose 12 years as the cut-off because it was close to the mean menarcheal age in the full sample, thus yielding groups of approximately equal size. This choice is consistent with classifications used by numerous previous studies (e.g., Caspi & Moffitt, 1991; Deardorff, Gonzales, Christopher, Roosa, & Millsap, 2005; Stattin & Magnusson, 1990). Ultimately, any dichotomization of menarche into “early” versus “late” is to some degree arbitrary. Therefore, our subsequent behavioral genetic analyses (main effects models and interaction models) examine both menarche and perceived development as continuous variables.
arche and the shared environmental component of nonviolent delinquency, with higher shared environmental influences in girls with an early age at menarche, \( c' = -0.115 \). A similar interaction with the shared environment was evident for perceived development, \( c' = 0.057 \); although this parameter was not significantly different than zero, nested model comparisons indicated that this interaction could not be dropped from the model without a significant decrease in model fit, \( \Delta \chi^2 = 251.96, p < .001 \). Overall, interaction models of nonviolent delinquency indicated that early maturing girls were more sensitive to shared and non-shared environmental influences, and less influenced by genes. Results from the full interaction models for nonviolent delinquency are illustrated in Figure 5.

For models of violent delinquency, the pattern of results for the genetic and shared environmental interactions was less clear. The interaction between perceived development and genes was not significantly different than zero, \( a' = 0.024 \), although this parameter could not be dropped from the model without a significant loss in model fit, \( \Delta \chi^2 = 208.38, p < .001 \). The interaction between perceived development and shared environment was also not significant, and nested model comparisons indicated that the \( c' \) parameter could be dropped from the model, \( \Delta \chi^2 = 0.12, p = .73 \). The only clear result from the interaction model of perceived development and violent delinquency, then, was the significant \( e' \) parameter, indicating that early maturing girls were more sensitive to nonshared environmental influences. Results from the best-fitting interaction model (with \( c' \) fixed to zero) for perceived development and violent delinquency are illustrated in the left side of Figure 6. In contrast, for menarche and violent delinquency, both the genetic, \( a' = 0.063 \), and shared environmental, \( c' = 0.115 \), interactions were significant, but in the opposite direction as observed for nonviolent delinquency. That is, genetic influences

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Table 2

<table>
<thead>
<tr>
<th>Model</th>
<th>Perceived → Non-violent</th>
<th>Menarche → Non-violent</th>
<th>Perceived → Violent</th>
<th>Menarche → Violent</th>
</tr>
</thead>
<tbody>
<tr>
<td>Parameters From Bivariate Twin Models of Pubertal Timing and Delinquency</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Indices of Model Fit</th>
<th>Perceived → Non-violent</th>
<th>Menarche → Non-violent</th>
<th>Perceived → Violent</th>
<th>Menarche → Violent</th>
</tr>
</thead>
<tbody>
<tr>
<td>( \chi^2 (df, P) )</td>
<td>111.73 (73, .002)</td>
<td>105.80 (73, .007)</td>
<td>118.78 (73, &lt;.001)</td>
<td>125.04 (73, .001)</td>
</tr>
<tr>
<td>RMSEA</td>
<td>.059</td>
<td>.054</td>
<td>.064</td>
<td>.068</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Variance in Pubertal Timing</th>
<th>Perceived → Non-violent</th>
<th>Menarche → Non-violent</th>
<th>Perceived → Violent</th>
<th>Menarche → Violent</th>
</tr>
</thead>
<tbody>
<tr>
<td>Var (A)</td>
<td>1.33 (.20)*</td>
<td>1.24 (.10)*</td>
<td>1.33 (.20)*</td>
<td>1.24 (.10)*</td>
</tr>
<tr>
<td>Var (E)</td>
<td>1.93 (.18)*</td>
<td>0.78 (.08)*</td>
<td>1.93 (.18)*</td>
<td>0.78 (.08)*</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Regression Parameters</th>
<th>Perceived → Non-violent</th>
<th>Menarche → Non-violent</th>
<th>Perceived → Violent</th>
<th>Menarche → Violent</th>
</tr>
</thead>
<tbody>
<tr>
<td>Genetic path (b_a)</td>
<td>0.237 (.063)*</td>
<td>-0.109 (.043)*</td>
<td>0.174 (.059)*</td>
<td>-0.036 (.041)</td>
</tr>
<tr>
<td>Environmental path (b_e)</td>
<td>-0.057 (.033)</td>
<td>.049 (.054)</td>
<td>-0.045 (.030)</td>
<td>-0.017 (.048)</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Variance in Delinquency</th>
<th>Perceived → Non-violent</th>
<th>Menarche → Non-violent</th>
<th>Perceived → Violent</th>
<th>Menarche → Violent</th>
</tr>
</thead>
<tbody>
<tr>
<td>Var (A)</td>
<td>.278 (.098)*</td>
<td>.323 (.091)*</td>
<td>.373 (.076)*</td>
<td>.409 (.071)*</td>
</tr>
<tr>
<td>Var (C)</td>
<td>.164 (.053)*</td>
<td>.181 (.053)*</td>
<td>.188 (.048)*</td>
<td>.193 (.048)*</td>
</tr>
<tr>
<td>Var (E)</td>
<td>.472 (.051)*</td>
<td>.473 (.052)*</td>
<td>.386 (.039)*</td>
<td>.389 (.039)*</td>
</tr>
</tbody>
</table>

Note.  Standard errors are in parentheses.
* Parameter significantly different from zero at \( p < .05 \).
were higher, and shared environmental influences were lower for early maturing girls. Results from the full interaction model for menarche and violent delinquency are illustrated in the right side of Figure 6.

### Discussion

Early pubertal timing puts girls at risk for a breadth of emotional and behavioral problems, including heightened involvement in delinquent behavior during adolescence and criminal behavior in adulthood. Because pubertal development involves an orchestrated set of changes across physiological, cognitive, and social domains, it has been challenging for researchers to identify the specific aspects of early pubertal timing that are most salient for delinquent behavior. Specifically, this study addressed the question of whether genetic differences accounted for the increased delinquent behavior seen in early maturing girls. Using nationally representative data from over 900 adolescent sister pairs of varying degrees of genetic relatedness, our behavioral genetic analyses indicated that a common set of genes predisposed girls toward early pubertal maturation and resulted in elevated involvement in both nonviolent and violent forms of delinquency. This result was consistent across measures of pubertal timing, including both objective reports of menarcheal age and girls’ self-reports of pubertal development. After accounting for these common genetic influences, there were no significant environmentally mediated paths between pubertal timing and delinquency. That is, sisters who differed in their pubertal timing did not significantly differ in their delinquency. Overall, our results challenge a purely socioenvironmental account

### Table 3

Unstandardized Parameter Estimates From Best-Fitting Interaction Models of Pubertal Timing and Delinquency

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Perceived Non-violent</th>
<th>Menarche Non-violent</th>
<th>Perceived Violent</th>
<th>Menarche Violent</th>
</tr>
</thead>
<tbody>
<tr>
<td>Full Model Fit (-2LL)</td>
<td>11734.15</td>
<td>11181.84</td>
<td>11693.26</td>
<td>11138.5</td>
</tr>
<tr>
<td><strong>Regression Parameters</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Genetic path (bA)</td>
<td>0.232 (.065)*</td>
<td>-0.093 (.043)</td>
<td>0.187 (.061)*</td>
<td>-0.018 (.039)</td>
</tr>
<tr>
<td>Environmental path (bE)</td>
<td>-0.041 (.034)</td>
<td>0.036 (.054)</td>
<td>-0.055 (.031)</td>
<td>-0.026 (.044)</td>
</tr>
<tr>
<td><strong>Variance in Delinquency</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Main effect of genes (a0)</td>
<td>0.380 (.160)*</td>
<td>-1.140 (.864)</td>
<td>0.565 (.071)*</td>
<td>1.352 (.382)*</td>
</tr>
<tr>
<td>Gene × puberty interaction (a’)</td>
<td>-0.082 (.030)*</td>
<td>0.124 (.056)*</td>
<td>0.024 (.026)</td>
<td>-0.063 (.030)*</td>
</tr>
<tr>
<td>Main effect of shared environment (c0)</td>
<td>0.455 (.074)*</td>
<td>1.877 (.600)*</td>
<td>0.443 (.055)</td>
<td>-0.946 (.334)</td>
</tr>
<tr>
<td>Shared environment × puberty interaction (c’)</td>
<td>0.057 (.030)</td>
<td>-0.115 (.054)*</td>
<td>[0]††</td>
<td>0.115 (.025)*</td>
</tr>
<tr>
<td>Main effect of non-shared environment (e0)</td>
<td>0.713 (.036)</td>
<td>1.863 (.226)*</td>
<td>0.636 (.032)*</td>
<td>1.846 (.172)*</td>
</tr>
<tr>
<td>Non-shared environment × puberty interaction (e’)</td>
<td>0.110 (.016)*</td>
<td>-0.093 (.016)^</td>
<td>0.065 (.019)*</td>
<td>-0.100 (.013)*</td>
</tr>
</tbody>
</table>

* Parameter significantly different from zero at p < .05. † Variance in delinquency due to genes is calculated as (a0 + a’menarche)^2. Thus, within the observed range of menarcheal age (±2 SDs from mean ~9 to 14 years), the genetic variance is positive. †† Nested model comparisons indicated that this interaction parameter could be fixed to zero without significant loss of model fit.

Figure 5. Additive genetic, shared environmental, and nonshared environmental influences on nonviolent delinquency are moderated by perceived development and age at menarche.
of why early maturing girls show increased involvement in delinquent behavior, and suggest that the genetic differences between early and late maturing girls hold more salience for their behavioral problems than environmental differences.

At the same time, results from our interaction models indicated that early pubertal timing moderated the residual genetic and environmental influences on delinquency. For both measures of pubertal timing, and for both violent and nonviolent forms of delinquency, the magnitude of nonshared environmental influence on delinquency was highest for early maturing girls. That is, after accounting for the main effects of pubertal timing on delinquency, there was greater within-family environmental variability in early maturing sister pairs than in later matures. Thus, there are two pathways of influence between early pubertal timing and delinquency: Genetic factors related to early maturation directly influence delinquent behavior, and early maturation also increases sensitivity to other environmental influences on delinquency. The interaction between pubertal timing and nonshared environmental influence was more marked for nonviolent than for violent delinquent behavior. This is consistent with previous theoretical and empirical reports indicating that nonviolent forms of delinquent behavior are more substantially influenced by environmental risks, whereas violent delinquency is more consistent across contexts and across the life course (Burt, 2009; Burt & Neiderhiser, 2009; Eley et al., 1999, 2003; Loeber et al., 2009; Moffitt, 1993; Tackett et al., 2003; Tuvblad et al., 2005).

Although the behavior genetics models used in the current analyses are capable of discriminating between genetic and environmental pathways, identifying the specific genes responsible for the observed association remains an important question for future research. One possibility is the androgen receptor gene (AR). Comings et al. (2002) found that the short allele of the GGC repeat polymorphism was associated with both earlier menarche in girls and higher aggression and impulsivity in males. (Jorm, Christensen, Rodgers, Jacob, and Easteal (2004) failed to replicate the association between AR and fathers’ parenting behaviors, but did not present data regarding the association between AR and menarche.) It is important to note, however, that the effect sizes associated with individual genetic polymorphisms are typically very small, and no single gene is likely to account fully for the association between early pubertal timing and delinquency. Moreover, it is difficult to speculate about specific risk genotypes when the genetic origins of individual differences in pubertal timing remain largely unknown.

In addition to genetic main effects, we also found that early maturing girls were more sensitive to environmental influences. This may reflect the impact of early pubertal maturation on brain development. In particular, pubertal maturation is linked to changes in dopaminergic neural circuits involved in responsiveness to emotional and motivational stimuli (Blakemore et al., 2010; Forbes & Dahl, 2010; Kuhn et al., 2010), and pubertal timing has been shown to predict neural activity in the striatum and prefrontal cortex in response to rewards and social threats (Forbes, et al., in press; Forbes et al., 2010). These changes in responsiveness to rewards and emotions precede age-based neural changes in cortical structures that underlie behavioral inhibition and effortful control (Casey et al., 2008; Somerville et al., 2010; Steinberg, 2008). Thus, from a developmental neuroscience perspective, early maturing girls are expected to be particularly attuned to the rewards inherent in risky environmental situations, and this process may underlie the greater nonshared environmental influence evident in the current results.

Moreover, for nonviolent delinquent behavior, genetic factors independent of pubertal timing were less influential for early maturing girls. This result is consistent with previous research by Burt et al. (2006), which showed substantial heritability (>60%) for conduct disorder symptoms in girls with on-time menarche, and very weak genetic influence (<10%) for girls with menarche prior to age 11. More broadly, the amplification of genetic variance among later maturing girls can be conceptualized in terms of

![Figure 6. Additive genetic, shared environmental, and nonshared environmental influences on violent delinquency are moderated by perceived development and age at menarche.](image-url)
the “social push” hypothesis (Raine & Venables, 1981): Biological vulnerabilities are more important when the environment is relatively benign, and they are less important in the presence of social stressors that predispose to antisocial behavior. For example, previous research has found that psychophysiological variables, such as low resting heart rate and reduced electrodermal conditioning, are stronger predictors of antisocial behavior among children with more advantaged and harmonious home environments (reviewed in Raine, 2002). Thus, among later maturing girls who do not experience the social challenges of early pubertal timing, genetic factors may be more important for differentiating who goes on to show delinquent behavior and who does not.

Limitations and Future Directions

The current study is limited by several features of the Add Health data, and it is worth considering the impact of those limitations on our results. First, Add Health participants were drawn from a broad age range at the initial assessment wave. To correct for this age heterogeneity, we have standardized all variables by age, consistent with previous analyses of pubertal timing variables in Add Health (Ge et al., 2007). Moreover, the mean age of participants (age 16 years) is older than is typical for studies of pubertal timing; indeed, given the mean menarcheal age is 12.2 years, the average participant was likely to be post-pubertal. Thus, the current analyses may have failed to detect environmentally mediated effects of pubertal timing that were specific to the early adolescent transition and that did not persist into mid-adolescence.

Second, the Add Health survey used an abbreviated self-report measure of pubertal development. Dorn et al. (2006) have noted that agreement between different measurement modalities for pubertal timing (e.g., physician exam vs. self-report) is low to moderate; thus, a different pattern of results may have been evident if pubertal timing had been measured at an earlier chronological age or by a measurement strategy other than self-report. In addition, menarche occurs relatively late in the process of puberty; given that children also differ in their rate of pubertal timing (pubertal tempo), menarcheal age may misrepresent the pubertal experiences of girls who begin puberty relatively early but mature relatively slowly (Mendle, Harden, Brooks-Gunn, & Graber, 2010). To our knowledge, no previous study has combined a genetically informative sample with repeated assessments of pubertal development by multiple modalities (e.g., self-report and physical exam). While such a design would, of course, be quite resource-intensive, it also holds potential to shed new light on the mechanisms by which early pubertal timing impacts girls’ psychosocial development.

Third, the current data are silent regarding girls’ history of behavior problems in childhood; thus, it remains unclear whether the impact of pubertal timing is consistent across childhood-versus-adolescent-onset trajectories of antisocial behavior. According to Moffitt’s (1993, 2003) taxonomy of “adolescent-limited” versus “life-course persistent” antisocial behavior, individuals with a history of childhood behavioral problems that persist into adolescence (and subsequently into adulthood) are more likely to be characterized by a constellation of severe risk factors, including adverse family environments, genetic vulnerabilities, and neuropsychological deficits. While some authors have argued that this trajectory is not applicable for females, given the low prevalence of female externalizing problems in childhood (e.g., D’Unger, Land, & McCall, 2002; Silverthorn & Frick, 1999; Silverthorn, Frick, & Reynolds, 2001), a review of longitudinal research on female antisocial behavior by Fontaine et al. (2009) concluded that a very small proportion of females (1–2% of the population) do show an “early starter” or “childhood-onset” trajectory of antisocial behavior. Given the rarity of this trajectory, and the community nature of the Add Health sample, the current results are probably descriptive of the more common, adolescent-limited trajectory of delinquency, which has been estimated to characterize up to 25% of adolescent girls (Fontaine et al., 2009). To our knowledge, however, no study has specifically examined whether the sequelae of early pubertal timing are moderated by history of childhood behavior problems; this is a fruitful question for future research.

Fourth, the Add Health study is characterized by a high degree of racial and ethnic diversity within a genetically informed sample. In contrast, virtually all of the previous genetically informed research on pubertal timing and its effects on psychopathology have been conducted using highly homogeneous samples of Northern Europeans (e.g., Dick, Rose, Viken, & Kaprio, 2000) or European Americans (e.g., Burt et al., 2006; Culbert, Burt, McGue, Iacono, & Klump, 2009). Thus one major contribution of the current study is its representation of African American, Hispanic/Latina, and Asian American girls in behavioral genetic research. At the same time, the number of sibling pairs in Add Health was too small to conduct analyses (particularly tests of G × E interaction) separately by race/ethnicity and still maintain adequate power. Although some authors have found that the developmental correlates of early pubertal timing differ across race/ethnic groups (e.g., Cavanagh, 2004), Lynne et al. (2007) reported that early maturation predicts delinquent and aggressive behavior in Hispanic/Latina and African American adolescents, as well as in White adolescents. Currently, it remains unknown whether environmental mechanisms are more or less important in racial and ethnic minorities.

Finally, the Add Health study does not include measures of relational aggression (i.e., behavior that is intended to harm others through damaging their peer relationships or social standing; Crick & Grotepeter, 1995), but instead focuses on only overt or physical manifestations of aggressive behavior. Girls are increasingly more likely to engage in relational aggression, relative to boys, as they transition from childhood to adolescence (Archer, 2004), and this increase may be linked to pubertal development. Very few studies have examined whether relational forms of aggression are also more prevalent in early maturing girls. However, the little extant research on this topic has found that both pubertal stage and early pubertal timing are associated with higher relational aggression, particularly for girls who experienced less positive parenting (Hemphill et al., 2010; Mrug et al., 2008). The only behavior genetic study of relational aggression in adolescents found that, like physical aggression, it is substantially heritable (Tackett, Waldman, & Lahey, 2009), but it remains unclear whether the mechanisms underlying the association between pubertal timing and relational aggression are the same as other forms of antisocial behavior.
Conclusions

Overall, this study provides evidence for a complex interplay between genetic and environmental risk in the association between early pubertal timing and adolescent girls’ involvement in violent and nonviolent delinquent behavior. Genes predisposing girls to earlier pubertal timing also increased vulnerability to delinquent behavior, and this common set of genetic influences underlie the common genetic influences, and to determine the mechanisms underlying early maturing girls’ greater sensitivity to environmental influences on delinquency.

References


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