The Prenatal Environment in Twin Studies: A Review on Chorionicity

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Received: 5 October 2015 / Accepted: 1 January 2016 / Published online: 5 March 2016 © The Author(s) 2016. This article is published with open access at Springerlink.com

Abstract A literature search was conducted to identify articles examining the association of chorionicity (e.g., whether twins share a single chorion and thus placenta or have separate chorions/placentas) and genetics, psychiatry/behavior, and neurological manifestations in humans twins and higher-order multiples. The main aim was to assess how frequently chorionicity has been examined in relation to heritability estimates, and to assess which phenotypes may be most sensitive to, or affected by, bias in heritability estimates because of chorionicity. Consistent with the theory that some chorionicity effects could lead to overestimation and others to underestimation of heritability, there were instances of each across the many phenotypes reviewed. However, firm conclusions should not be drawn since some of the outcomes were only examined in one or few studies and often sample sizes were small. While the evidence for bias due to chorionicity was mixed or null for many outcomes, results do, however, consistently suggest that heritability estimates are underestimated for measures of birth weight and early growth when chorionicity is not taken into account.

Keywords Chorionicity · Genetics · Heritability · Prenatal environment · Twins

Introduction

Twin studies have long been used to estimate the unique contributions of genetic and environmental influences on variation in human traits. One assumption of the quantitative genetic theory underlying twin studies is the equal environments assumption, which states that the exposure to environmental events that create resemblance between co-twins for the trait under study is equal for monozygotic (MZ) and dizygotic (DZ) twin pairs (Loehlin and Nichols 1976; Scarr and Carter-Saltzman 1979). The prenatal environment is a specific and crucial environmental influence on many human traits (Barker 1990), and while twins and higher-order multiples share the womb, the prenatal environment may not be equal for both twins in a pair, or for other higher-order multiples. Thus, the prenatal environment cannot necessarily be considered as an environmental factor creating resemblance in children sharing the
womb at the same time. How twins experience the prenatal environment depends, in part, on chorionicity, i.e., whether twins share a single chorion (monochorionic, MC) or have separate chorions (dichorionic, DC). Monozygotic (MZ) twins can be mono- or dichorionic, whereas dizygotic twins are dichorionic.

In this review, we first introduce the concepts of the chorion, amnion, and placenta. Next, we discuss how chorionicity may shape the prenatal environment of twins and higher-order multiples and aim to summarize the types of outcomes that have been linked to chorionicity. Finally we review and summarize studies which have examined the influence of chorionicity on twin-based heritability estimates in order to draw conclusions about whether chorionicity introduces bias and, if there is bias, whether this bias affects phenotypes in a consistent manner.

**Chorionicity**

The chorion is the outer-most fetal membrane that contains the amnion/amniotic sac. The amnion is the thin inner-most fetal membrane that protects the embryo/fetus and contains amniotic fluid. The chorion connects the amnion, amniotic sac, and the fetus to the placenta and contributes to placental development. Thus, if twins share a chorion (e.g., are monochorionic or MC) they will share a single placenta, whereas twins with separate chorions (e.g., dichorionic or DC twins) develop individual placentas. DZ twins are dichorionic, since they form from two separately fertilized eggs, although very rare exceptions have been described in the literature (e.g., Souter et al. 2003). Figure 1a, b provides an illustration of the multiple ways twins can share the chorion and amnion. Figures 2, 3, 4 show ultrasound images of monochorionic (Fig. 2), dichorionic (Fig. 3), and trichorionic triplets (Fig. 4). Generally, it is thought that the timing of division of the blastocyst/embryo determines amnionicity and chorionicity (Hall 2003; De Paepe 2015), such that later cleavage (e.g., between 4 and 13 days) leads to MC twins and earlier cleavage (e.g., before 4 days) leads to DC twins. Later cleavage (e.g., 8–13 days) may lead to monoamniotic twins and earlier cleavage (e.g. before 8 days) to diamniotic twins. However, what determines whether and when a fertilized egg splits, and if the resulting MZ twins (or triplets or other higher order multiples) will develop separate chorions, are questions for which very little empirical data are available (Knopman et al. 2014; Herranz 2015).

**Prevalence**

Epidemiological data indicate that the MZ twin prevalence is fairly consistent at around 4 per 1000 maternities worldwide (Tong et al. 1997). DZ twinning rates differ around the globe and over time (e.g., increasing with maternal age and as artificial reproductive techniques have become more widely available and used; Hoekstra et al. 2008). For example, among Caucasian populations (e.g., United States, Europe, Australia), total twinning rates were estimated at 15–16 per 1000 in 2003 (Hoekstra et al. 2008), whereas Asian countries had lower rates at about 9 per 1000 (Smits and Monden 2011; Hoekstra et al. 2008).
African populations have higher twinning rates, of about 12–18 in sub-Saharan countries and over 18 per 1000 in central African countries (Smits and Monden 2011). Thus, in Caucasian and sub-Saharan African populations, MZ twins comprise ~26 % of all twins, whereas in Asian populations, MZ twins represent over half of all twins, and in central African populations MZ twins represent less than 5 % of all twins. Given that heritability estimates are specific to the population being studied, differences in the prevalence of MZ and DZ twins in different populations will likely affect the extent to which chorionicity might affect heritability estimates in these populations.

Of all MZ twin pairs, about two-thirds (70–74 %) are monochorionic (MZ-MC) and one-third (35–30 %) are dichorionic (MZ-DC) (Hall 2003). However, 1–2 % of MZ twin pairs are monoamniotic (Hall 2003) although this percentage varies by sample. Given the low prevalence of monoamniotic twins this review focuses on the potential effects of chorionicity rather than amnionicity. For Caucasian populations (where most twin research has been done) about 17 % of all twin pairs are MZ-MC, ~9 % are MZ-DC, and ~74 % are DZ-DC. However, the proportion of MZ-MC, MZ-DC, and DZ-DC twins in any given study varies widely and is not always reported (Petterson et al. 1998).

**Determination**

A large body of literature has examined appropriate ways to determine chorionicity. Prospectively, chorionicity is best determined via ultrasound. Determining chorionicity is highly accurate (96 %) by ultrasound in the first trimester, though still accurate (80 %) in the second (e.g., see Audibert and Gagnon 2011 for review; Machin 2004). Placental pathology examination also provides a direct assessment of chorionicity shortly after birth (De Paepe...
2015). Retrospective self-report determination of chorion type, for example by asking twin participants “how many placentas” there were at birth, has been suggested to be unreliable: 60 % accurate for MZ and 37 % accurate for DZ twins (Derom et al. 2003). Some studies have also tried to use dermatoglyphics to retrospectively determine chorionicity (e.g., Davis et al. 1995; Reed et al. 1991, 1978, 2002; Melnick and Myrianthopoulos 1979; Steinman 2001). Placental pathology examination and ultrasound appear to be the most reliable methods of determining chorionicity; thus, for the remainder of this paper we focus on studies which employed one of these two methods.

**Placental function**

The MC placenta functions like a single placenta, although a single placenta was not designed to support the growth of two fetuses. Therefore, MC placentation has a profoundly different biology than DC placentation. The greatest danger associated with MC placentation is related to the structure of blood vessels. One twin usually has better placement and therefore receives more of the nutrients. Inter-fetal vascular connections also form vascular anastomoses (i.e., the joining of two blood vessels) and connect the circulation of one twin to the circulation of the other, so in some pregnancies, there is direct blood sharing of MC twins. These inter-fetal vascular connections rarely form in DC twin pairs (Machin and Bamforth 1996; Phillips 1993).

Unequal placental sharing is a major cause of fetal growth discordance in MZ twins (Chang 2008; Cleary-Goldman and D’Alton 2008; Nikkels et al. 2008). For example, specific reductions in five amino acids have been shown to explain discordant growth in MZ twins, suggesting that the inter-twin distribution of blood and nutrients accounts for within-pair differences in birth weight, as opposed to more general placental dysfunction (Bajoria et al. 2001). Extreme discordant growth due to unequal placental sharing can result in twin-to-twin transfusion (TTTS) syndrome, a severe pregnancy complication unique to MC twin pairs where there is also direct blood sharing (occurring in 5–30 % of MC twin pairs; Haverkamp et al. 2001; Phillips 1993). The imbalanced blood flow and twin-to-twin transfusion has been reported to influence MZ twin resemblance for birth weight (see Foley et al. 2000 for review, and supplemental Table 1). These findings result in a difference in MC and DC twins for some birth outcomes including birth weight discordance, as MC twins are more likely to have higher birth weight discordance than DC twins who do not share a placenta.

The placenta also functions as a barrier, allowing small molecules (e.g., gases, nutrients, waste material, antibodies) to pass between mothers and children through passive transport (Page 1993; Schneider 1991). Other small molecules that may have an effect of fetal development (e.g., some maternal hormones like cortisol; bacteria; teratogens such as illicit drugs) can also be diffused through the placenta (van der Aa et al. 1998; Page 1993). Thus, the composition of the placenta and efficiency of transport between mother and child can affect fetal development. The placenta also functions as an endocrine organ (Melmed et al. 2012), synthesizing a large array of hormones (e.g., sex steroids and protein hormones) and cytokines that play a key role in fetal development (and maternal endocrine function). There are individual differences in hormone production, and sharing a placenta may lead to similarities in MC twins that are related to the levels and changes in placental hormone production relative to DC twins. Sharing a placenta in this case may lead to more similar in utero environments for MC twins relative to DC twins. However, endocrine function is, to some extent, linked to the vascular system, and the amount of pathogen, infection, nutrient, and gas and waste diffusion may also be linked to the proportion of the placenta dedicated to each child (Melmed et al. 2012). The potential impact of diffusion and endocrine function on similarity and differences of MC versus DC twins has not, to our knowledge, been investigated and is potentially an important area for future research. Thus, while some placental mechanisms (diffusion and endocrine function) may lead to more similar whereas others (unequal sharing of the vascular system) may lead to more different in utero environments, these mechanisms are linked and so the reality is less clear-cut.

**Chorionicity and heritability**

Because of the placental mechanisms leading to similarities and differences of the in utero environments for twins of different types, chorionicity may bias the heritability estimates found in twin studies (see Table 1). The potential challenge that chorionicity plays in the validity of twin studies is not a new concept (Price 1950), and has been highlighted in a number of studies (Derom et al. 2001; Foley et al. 2000; Munsinger 1977; O’Brien and Hay 1987; Phelps et al. 1997; Prescott et al. 1999; Price 1950). The prenatal environment could be more similar for MC twins relative to DC twins because of the shared chorion, or less similar because of the vascular and placental sharing inequalities often observed in MC but not DC pregnancies. Vascular differences found in MC twins often lead to differences in intrauterine growth of the twins, and thus MC twins can appear quite dissimilar especially early in life. If zygosity is only determined via questionnaire, MC twins may be misclassified as DZ twins, which would bias results of twin studies (Machin 2001, 2009). Even with correct classification, if MC twins are more dissimilar because of unequal placental sharing, then heritability estimates may...
be underestimated because MZ twins would have a lower correlation, closer to that of DZ twin pairs (Price 1950). That is, the subset of MZ-DC twins may be more similar to DZ-DC and less similar to MZ-MC twins in their sibling correlations. This would, in turn, affect the intra-class correlations for MZ and DZ pairs (e.g., reduce the contrast) and downwardly bias the estimates of heritability. Further, MZ twins often have poorer outcomes than DC twins (see review below and Supplemental Table). This may lead to mean-level or variance differences in the outcomes between MC and DC twins due to a possible violation of the equal environments assumption, which could also bias heritability estimates. For example, if in a pair of MC twins, one of the twins is at increased risk for a particular outcome (e.g., through limited blood supply because of TTTS), then the prenatal environment is not ‘shared’ although the MC status is considered ‘shared’.

However, if sharing a placenta makes twins more similar because of similar intrauterine environments (e.g., passive transport), then the potential bias could indeed operate in the opposite direction, leading to overestimation of genetic influences (Phillips 1993). For example, MC pairs may be more likely to experience the same environmental exposures and pathogens, including infections and substance use exposure (Prescott et al. 1999). The crux of understanding how chorionicity may influence heritability estimates lies in understanding whether the prenatal environment is more or less similar for MC twins, and for which outcomes chorionicity matters for twin similarity.

This ‘chorionicity debate’ led to the proposal for chorion-control studies, where MZ-MC twins are compared with MZ-DC twins on a specific trait, or multiple traits, and a call for including chorionicity in classical twin studies (Phelps et al. 1997). However, methodological challenges have made the examination of the potential role of chorionicity difficult and largely theoretical; as noted above, a reliable assessment of chorionicity ideally requires placental pathology examination or prenatal ultrasound. As there is an increasing interest in simultaneously examining prenatal and genetic influences as exemplified in this special issue of Behavior Genetics, it is important to revisit the question of whether chorionicity may influence outcome variables assessed in twin studies and whether such influence could bias heritability estimates from studies that include predominantly twins.

### Method

#### Medical library database search

The purpose of the literature search was to identify articles examining associations of chorionicity and genetics, psychiatry/behavior, and neurological manifestations in humans (twins/multiples). We searched PubMed (yielding 2111 articles after deleting duplicates), Embase, 1947 to present, OvidSP (yielding 1455 articles after deleting duplicates), and PsycINFO 1806 to Present (yielding 138 articles after deleting duplicates). The entire search strategy, including all search terms for each database, is included in Appendix. A variety of search terms were used (both text words [tw] and the PubMed search also included Medical Subject Heading terms [MeSH]), including but not limited to variants of multiple birth (e.g., multiple birth, twin), chorionicity (e.g., chorion, monochori*, dichori*, placentation), genetics (e.g., genetic*, epigenetic*, gene, genes, genotype), intelligence (e.g., intelligence, IQ), psychiatry/behavior (e.g., psychology, psychiatry*, mental, psychology*, behavior, neuropsych), neurological manifestations (e.g., neuromorbidity, neurologic*), and
concordance/discordance (e.g., twin, discordan*, concor-
dan*). In Embase, twin concordance and discordance was
searched in combination with the outcome separately
because of poor representation of chorionicity in the bib-
liographic records. Animal studies were excluded in all
searches. We did not filter by language or date of pub-
cation. After duplicates from the multiple searches were
excluded, there were a total of 2920 unique articles.

Selecting relevant articles

Each of the abstracts of the 2920 articles were read and
judged for relevance to chorionicity and genetics/behavior/
psychiatry (e.g., identifying sources which examine the
association of chorionicity with behavioral outcomes). Full
texts were also searched for “chor” to aid with determining
whether articles were relevant. Case studies and non-em-
pirical articles were excluded from the final selections. We
also excluded studies that used retrospective report of
chorionicity as well as other alternative proxies for chori-
onicity (e.g., birth weight discordance, handedness, mir-
roring). At the end of this culling, 307 articles were
identified as potentially relevant.

These 307 articles were further classified into back-
ground/review articles (n = 68), studies that compare the
prevalence of various outcomes stratified by chorionicity
(reviewed below and in the Supplementary Table, n = 134), studies that examined chorionicity effects in the
context of behavioral genetic designs (n = 38), epigenetic
studies (n = 5), and irrelevant studies (e.g., not examining
chorionicity directly, or conference abstracts which may be
preliminary and not peer reviewed, vetted findings, n = 62).
This sorting was done by reading the abstracts and
articles to the depth required to make a decision. Of primary
interest for the current review were the studies that exam-
ined chorionicity effects in the context of behavioral genetic
designs. These studies were reviewed in detail in order to
conclude whether chorionicity may bias results of heri-
tability estimates for the diverse outcomes studied. We did
not restrict our search based on outcomes during this phase.

Results

Chorionicity and prevalence of birth outcomes
and human traits

A very large body of literature has examined whether there
are prevalence differences in various birth, perinatal, and
other outcomes based on chorionicity (see Supplementary
Table for a summary of the 134 articles reviewed). The
best-characterized outcomes influenced by chorionicity
include immediate pregnancy and birth outcomes rather
than longer term growth and psychiatric outcomes. We
highlight the most consistent findings here (see Supple-
mentary Table for details and exceptions). Most studies
found that MC pregnancy infers higher risk of mortality
than DC pregnancies (see Supplementary Table), but
effects are not always consistent (e.g., Baghdadi et al.
2003; Lenis-Cordoba et al. 2013). Fetal growth has also
been robustly linked with chorionicity. For example, birth
weight discordance occurs more frequently in MC twins
than DC twins (although this effect is not found in every
study). Further, MC twins generally have lower birth
weight (especially the smaller twin), lower birth weight
after adjusting for gestational age (Ananth et al. 1998;
Shrim et al. 2010), and shorter crown-rump length.
Intrauterine growth restriction is more prevalent in MC
twins than DC twins. However, fetal growth velocity has
not been shown to differ for MC versus DC twins (Smith
et al. 2001; Taylor et al. 1998). A host of obstetric and
perinatal complications have also been examined exten-
sively in relation to chorionicity. Most studies have found
that DC twins are born at older gestational ages than MC
twins, and experience fewer morbidities (e.g., patent ductus
arteriosus, sepsis, vision and auditory loss, congenital
malformations, anemia, intracranial lesions). In general,
MC pregnancies are riskier than DC pregnancies.

In contrast to pregnancy and birth outcomes, associa-
tions of chorionicity and cognitive, psychiatric, and
behavioral outcomes are not as frequently studied or as
consistent. The limited literature hints that MC twins have
worse cerebral white matter outcomes than DC twins. For
example, MC twins have higher cerebral white matter
lesions (Adegbite et al. 2005) and a higher incidence of
antenatal necrosis of cerebral white matter (Bejar et al.
1990) than DC twins. However, another study showed no
differences in clinical neurologic indicators of perinatal
asphyxia (van Steenis et al. 2014). In terms of cognitive
performance, results are mixed. One study suggested that
MC twins have higher rates of pathological nonverbal
performance and learning disabilities (Einaudi et al. 2008),
whereas other studies showed no difference in mental
development indexes (e.g., on the Bayley; Welch et al.
1978; Steingass et al. 2013). Studies examining cerebral
palsy are inconsistent, with some suggesting that MC twins
are at a higher risk (Burguet et al. 1995, 1999), but others
finding no difference in prevalence of cerebral palsy in MC
versus DC twins (Steingass et al. 2013; Hack et al. 2009),
or that the association was attenuated when controlling on
other perinatal factors (Livinec et al. 2005).

Chorionicity and behavioral genetic designs

We identified 38 articles that examined chorionicity within
a behavioral genetic design. Of these, one was excluded
because no full text was available in English. An additional seven were excluded because chorionicity was not determined via placental pathology or ultrasound. We organized the resulting 30 studies into the following outcome-based categories (although some studies have multiple outcomes across multiple categories): birth weight and early growth, screening/vaccination, handedness, anthropomorphic measures, cognitive/brain measures, and behavioral measures. Reviewed studies are presented in Table 2.

Eight studies examined chorionicity effects on intra-pair associations/differences and/or included chorionicity in classical twin models decomposing the variance in a phenotype into additive genetic (A), common environmental (C), and non-shared environmental (E) influences (e.g., ACE models) in regard to birth weight and early growth patterns. Across these studies, generally it was found that MC twins grew more slowly, were less variable, and less correlated for birth weight than DC twins, and that including chorionicity yielded attenuated, more precise heritability estimates (Buzard et al. 1983; Vlieitten et al. 1989; Gielen et al. 2008; Touwsager et al. 2010; Welch et al. 1978; Mukherjee et al. 2009; Spitz et al. 1996; Loos et al. 2001a). Although effects were not always significant (e.g., trend-level; Buzard et al. 1983), the evidence does point to biased heritability estimates in studies of birth weight; where, without accounting for chorionicity, heritability is underestimated.

One study examined screening for trisomy 21 and one examined responses to vaccination (Wojdemann et al. 2006; Gupta et al. 2008). Neither study found evidence of a chorionicity effect on twin similarity. Two studies examined handedness (Carlier et al. 1996; Melnick and Myrianthropoulos 1979). Neither found any effects of chorionicity on twin similarity.

Eleven studies measured various anthropometric measures. Chorionicity effects varied with outcome and over time. For example, MZ-DC twins were more discordant for cholesterol levels from cord blood than MZ-MC twins (Corey et al. 1976). There were significant chorionicity effects when modeled explicitly for height at age 4 years, explaining a small percentage of variance (4 %), but not for weight (Hur and Shin 2008). One study suggested that MZ-MC twins were more discordant than MZ-DC twins for height at 8–12 years (Spitz et al. 1996), however another found that there were no differences in the concordance of MZ-MC and MZ-DC twins for height in at 10–16 years (Gutknecht et al. 1999). MZ-MC twins were more discordant than MZ-DC twins for weight and BMI throughout childhood and adolescence (Gutknecht et al. 1999; Spitz et al. 1996; Mukherjee et al. 2009). There was also some evidence that MZ-MC twins were more similar than MZ-DC twins for saccadic eye movements in adolescence (Blekher et al. 1998). In adults, there were no differences in the twin similarity of various obesity-related measures (or very small effects; Loos et al. 2001a), lung measures, or conventional and ambulatory blood pressure (Loos et al. 2001a; van den Borst et al. 2012; Souren et al. 2007; Fagard et al. 2003). The only significant chorionicity effect on twin similarity found in adults was for fasting fibrinogen: MZ-DC twins were more similar than MZ-MC twins (Loos et al. 2001b). In sum, chorionicity appears to maintain an effect on twin similarity for a variety of anthropometric measures even after birth, but these effects seem to dissipate in later adolescence and adulthood. However the directions of effects varied for each measure. Based on the limited evidence provided here, heritability estimates may be overestimated for cord blood cholesterol, saccadic eye movements, and height at age 4 years. However, heritability estimates may be underestimated for height at 8–12 years, weight and BMI in childhood and adolescence, and fasting fibrinogen in adults.

Eight studies examined cognitive and brain-based measures, and findings were generally mixed. Studies very early in life (e.g., from in utero to 1 year) found no significant effects of chorionicity on twin similarity for head circumference, intracranial volume (Mukherjee et al. 2009), or anterior fontanelle development (Melnick et al. 1980). In toddlerhood, there were no chorionicity effects on twin similarity for the Bayley Mental Development scores (Welch et al. 1978). In childhood, there was evidence of two populations of MZ twins with regard to variation in IQ, as MZ-MC twins differed from DZ twins but MZ-DC twins did not (Melnick et al. 1978), suggesting considerable influence of the prenatal environment on IQ. However, another study showed that there were no differences in twin similarity based on chorionicity for the McCarthy Scales of Children’s Abilities (Sokol et al. 1995). Also in childhood, one study found that MC twins were more similar for arithmetic and vocabulary (with chorionicity explaining 14 and 10 % of the total variance respectively; Jacobs et al. 2001), whereas another found no effect of chorionicity on twin similarity for vocabulary (Spitz et al. 1996). MZ-MC twins were more similar than MZ-DC twins for measures of personality in one study (Sokol et al. 1995), whereas another study found null findings for measures of personality (Gutknecht et al. 1999) in childhood. Some studies found relatively few significant effects of chorionicity on twin similarity (relative to the number of tests examined, e.g., Gutknecht et al. 1999; Spitz et al. 1996). There was only one replicated finding: MZ-MC twins were more similar than MZ-DC twins for the block design but not for vocabulary in children and adults (Spitz et al. 1996; Rose et al. 1981). One reason for the mixed findings in the literature likely is the small sample sizes used to investigate these effects. Nonetheless, there is evidence that chorionicity may have an effect on twin similarity for some cognitive measures, particularly...
### Table 2 Reviewed studies examining chorionicity with a behavioral genetic design

| Reference | N (twin pairs) | Location | Placenta determination | Age | Comparison | Analysis | Outcome | Effect | Other notes |
|-----------|---------------|----------|------------------------|-----|------------|---------|---------|--------|-------------|
| Birth weight (BWT), early growth | | | | | | | | | |
| Buzzard et al. (1983) | 52 MZMC 40 MZDC 72 DZDC | Canada | Placental pathology | 20–41 years | MC versus DC and MZ versus DZ | Intra-pair differences | BWT | MC = DC; MZ = DZ | Trends for intra-pair differences in chorionicity and chorionicity by sex interaction |
| Vlietinck et al. (1989) | 762 MZ 1093 DZ | Belgium | Examination of placenta | Birth | MZMC versus MZDC versus DZ | Extended ACE model with chorionicity included | BWT | rMC < rDC (MC also less variable than DC) | EFPTS |
| Gielen et al. (2008) | 4252 twin pairs | Belgium | Examination of placenta | Birth | MZ versus DZ, chorionicity a covariate | ACE models, chorionicity a covariate | BWT | | EFPTS |
| Touwslager et al. (2010) | 522 individuals | Belgium | Examination of placenta | 0–2 years | MZMC versus MZDC versus DZ | Intra-pair growth correlations | Growth during infancy | rMZMC = rMZDC at 0-1, 0-6, or 6-12 months; rMZMC > rMZDC at 12-24 months; EFPTS; MZMC grew more slowly than MZDC |
| Misc. screening/vaccination | | | | | | | | | |
| Wojdemann et al. (2006) | 31 MC 150 DC | Denmark | Ultrasound | Birth | MC versus DC | ICCs | Nachal transluency | rMC = rDC | |
| Gupta et al. (2008) | 117 DC 16 MC | India | Examination of placenta | Newborns | MC and MZ versus DZ | Intra-pair agreement | BCG vaccine reaction | rMC = rDC; rMZ = rDZ | |
| Handedness | | | | | | | | | |
| Carlier et al. (1996) | 20 MZMC 24 MZDC 24 DZ | France | Examination of placenta | 8–12 years | MZMC versus MZDC versus DZ | Comparison of difference between groups on the average absolute twin difference; if chorion effect not significant, all MZs pooled for MZ/DZ comparison | Manual performance, direction, laterality (handedness) | No chorionicity effects | rMZ = rDZ in classic twin design (chorionicity not controlled) |
| Melnick and Myrianthopoulos (1979) | 117 MZMC 56 MZDC | USA | Sex, nine blood groups, and gross and microscopic examination of placenta | Birth through 7 years | MZMC versus MZDC | Comparing concordance | Finger ridge count and right-left asymmetry of ridge count and congenital anomaly (e.g., differentiation of body parts, tissue differentiation or dysplasias) | rMZDC < rMZMC for total ridge count. No chorionicity effects for congenital anomalies, other assessments of dermatoglyphics | NCPP twin population |
| Anthropomorphic measures | | | | | | | | | |
| Corey et al. (1976) | 30 MZMC 22 MZDC | USA | Examination of fetal membranes | Newborns | MZMC versus MZDC | General linear model for twin data, applied to MC and DC twins | Cholesterol from cord blood | Female MZ > male MZ for cord blood cholesterol | |
| Reference                  | N (twin pairs) | Location | Placenta determination | Age               | Comparison | Analysis | Outcome | Effect | Other notes |
|---------------------------|---------------|----------|------------------------|-------------------|------------|----------|---------|--------|-------------|
| Hur and Shin (2008)       | 81 MCMZ 47 DCMZ 457 DZ | S. Korea | Examination of placenta | 2–9 years (mean = 4 years) | MZC versus MZDC versus DZ | ACE models including chorion effects | Height, weight, BMI | A + C sig. chorion effects for height sig but only 4 %, not sig for weight/BMI | follows up a traditional MZ/DZ study showing sig. genetic influence |
| Blekher et al. (1998)     | 17 MZMC 16 MZDC | USA | Examination of placenta | 9–18 years (mean 13.2 years) | MZMC versus DC | ICCs | Saccadic eye movements: accuracy, slope, velocity at 15 degree saccade | MZMC > MZDC for reaction times; not phasic component of saccadic command. | EFPTS; regular twin study run subsequently |
| Souren et al. (2007)      | 240 MZ 138 DZ | Belgium | Examination of placenta | Adults | MZMC versus MZDC | ICCs | Body mass, BMI, fat mass, waist-to-hip ratio, sum of four skinfold thickness, lean body mass, fasting glucose, fasting insulin, insulin resistance and beta cell function, and insulin-like growth factor binding protein-1 levels, total, LDL, HDL, total:HDL ratio for cholesterol, triacylglycerol, NEFA and leptin levels | MZMC = MZDC | EFPTS; regular twin study run subsequently |
| van den Borst et al. (2012)| 165 MZ 71 DZ 102 single twins | Belgium | Examination of placenta | Adults | MZMC versus MZDC | ICCs | Lung measures: forced expiratory volume in 1 s and forced vital capacity | MZMC = MZDC | EFPTS; regular twin study run subsequently |
| Fagar et al. (2003)       | 125 DZ 97 MZDC 128 MZMC | Belgium | Examination of placenta | 18–34 years | MZMC versus MZDC versus DZ | Intra-pair correlations; ACE models fit to MZ/ DZ; MZDC/ DZ; MZMC/ DZ | Conventional and ambulatory blood pressure | MZMC = MZDC | EFPTS |
| Loos et al. (2001a)       | 280 MZMC 212 MZDC 311 DZ | Belgium | Examination of placenta | 18–34 years | MZMC versus MZDC versus DZ | Intra-pair concordance | Abdominal obesity | Contribution of zygosity and chorionicity low (< 1.7 %) for adult weight. /DC > /MC for BWT only | EFPTS |
| Loos et al. (2001b)       | 67 MC 56 DC | Belgium | Examination of placenta | M = 25 years | MZMC versus MZDC versus DZ | ICCs | Fasting fibrinogen | MZMC < MZDC | Suggested chorionicity effect would change heritability |
| Cognitive and brain measures | Mukherjee et al. (2009) | USA | Placental pathology, or ultrasound if pathology not performed or unavailable | Prenatal and birth | MZMC versus MZDC versus DZ | Comparing concordance | Head circumference and weight at 22 weeks, 32 weeks, birth, and intracranial volume | MZMC = MZDC | head circumference and weight; Sig. group difference on ICV (DZ > MZMC and MZDC) |
| Reference                  | N (twin pairs) | Location    | Placenta determination | Age          | Comparison                | Analysis          | Outcome                                                                 | Effect                                                                 | Other notes                                      |
|---------------------------|----------------|-------------|-------------------------|--------------|---------------------------|------------------|--------------------------------------------------------------------------|------------------------------------------------------------------------|-------------------------------------------------|
| Welch et al. (1978)       | 20 MZMC, 12 MZDC | USA         | Gross and microscopic placenta examination | 18 months    | MZMC vs MZDC              | Differences in within-pair mean squares | BWT, Bayley mental development scores | $\Delta$DC > $\Delta$MC for BWT; not significant (in opposite direction) for Bayley | Positive correlation of BWT and Bayley |
| Melnick et al. (1978)     | 86 MZ, 173 DZ   | USA         | Sex, nine blood groups, and gross and microscopic examination of placentae | 7 years      | MZMC vs MZDC and MZDC vs DZ | Heritability estimates | IQ | heterogeneity between MZM/DZ total variances, but not MZDC/DZ. Estimated genetic variance comparing MZM/DZ not sig diff from 0, but MZDC/DZ was. | NCPP twin population, also split by race |
| Sokol et al. (1995)       | 23 MZMC, 21 MZDC | USA         | Gross and microscopic examination of placentae | ~6 years     | MZMC vs MZDC              | Intra-pair differences | McCarthy Scales of Children’s Abilities and Personality Inventory for Children | $\Delta$MZMC > $\Delta$MZDC for 3/4 Personality Inventory for Children factor scales, 8/12 clinical scales, and 2/4 validity/screening scales. No differences on 6 McCarthy scales. | |
| Antoniou et al. (2010)    | 663 twin pairs  | Belgium     | Examination of placenta  | 7–15 years, $M = 10.4$ years | MZ vs DZ on placenta | Bivariate ACE models | BWT-IQ and cord knots-IQ associations | No chorionicity effect | |
| Spitz et al. (1996)       | 20 MZMC, 24 MZDC, 24 DZ | France | Ultrasound and microscopic examination at delivery | 8–12 years | MZMC vs MZDC              | Intra-class correlations | Anthropometric measures at birth and assessment; Cognitive battery (Vocabulary, block design from WISC-R, K-ABC, perception and mental rotation | $\Delta$MC < $\Delta$DC for BWT, child weight, height, BMI; $\Delta$MC > $\Delta$DC block design. No chorion effect for vocabulary, K-ABC, perception, or mental rotation | |
| Jacobs et al. (2001)      | 161 MZMC, 82 MZDC, 188 DZ | Belgium | Examination of placenta  | 9–11 years   | MZMC vs MZDC and MZDC vs DZ | ACE models including chorion effects | WISC-R | $\Delta$MZMC > $\Delta$MZDC on arithmetic (chorionicity = 14 % of total variance) and vocabulary (chorionicity = 10 %) | EPPTS; High heritability for almost all subscales and IQ scores. |
| Gutknecht et al. (1999)   | 20 MZMC, 23 MZDC | France      | Examination of placenta  | 10–16 years  | MZMC vs MZDC              | Comparison of average within-pair difference between groups | Weight, height, BMI; WISC-III; figurative reasoning; schools’ standardized exams; Personality and behavioral measures | $\Delta$MC > $\Delta$DC, weight, BMI; 1/6 WISC-III measures; figurative reasoning, 1/5 personality/behavioral variables (most null) | Lack of power; but found consistent direction of effects in even null findings |
Table 2 continued

| Reference            | N (twin pairs) | Location | Placenta determination | Age            | Comparison                          | Analysis                                                | Outcome                                      | Effect                                                                 | Other notes                                                                 |
|----------------------|----------------|----------|------------------------|----------------|-------------------------------------|---------------------------------------------------------|---------------------------------------------|------------------------------------------------------------------------|---------------------------------------------------------------------------|
| Rose et al. (1981)   | 17 MZMC, 15 MZDC, 28 DZ | Canada   | Examination of placenta | Adults         | MZMC versus MZDC versus DZ         | Mean differences, ICCs, heritability estimates (from several contrasts) | WAIS vocabulary and block designs             | rMZMC > rMZDC (~ 2x) for block design only; vocabulary: rMZMC = rMZDC > rDZ |                                                                           |
| Melnick et al. (1980) | 94 MZ, 187 DZ    | USA      | Examination of placenta | 4 months and 1 year | MZ versus DZ, checked chorion effects | Within-pair mean square estimates of genetic variance | Anterior fontanelle development               | No chorionicity effects at 4 months or 1 year                           | NCPP twin population, also split by race                                  |
| Chen et al. (1990)   | 44 MZ, 18 DZ     | China    | Examination of placenta | 6 months       | MZ versus DZ and MZMC versus MZDC  | Intra-pair correlations                                 | Temperament                                  | rMZMC = rMZDC                                                          | Found genetic influences                                                   |
| Hur (2007)           | 56 MZMC, 34 DCMZ, 316 DZ | S. Korea | Examination of placenta | 2–9 years (M = 4 years) | MZMC versus MZDC versus DZ         | ACE models including chorion effects                  | Prosocial behaviors                           | No chorionicity or shared environment effects.                           | Twin study estimates good                                                 |
| Riese et al. (1999)  | 48 MZMC, 29 MZDC | USA      | Examination of placenta | Neonates before released from hospital | MZMC versus MZDC                          | ICCs                                                   | Temperament                                  | rMZMC = rMZDC                                                          |                                                                           |
| Wichers et al. (2002)| 202 MZMC, 125 MZDC, 425 DZ | Belgium | Examination of placenta | 6–17 years     | MZMC versus MZDC versus DZ         | ICCs, ACE model                                       | CBCL total problems score                     | rMZDC = rMZMC > rDZ. Chorionicity effect was 0 in the full ACE model | EFPTS                                                                    |

Described effects are not necessarily comprehensive. Findings were interpreted by comparing MC and DC similarity (e.g., ICCs)—with MC and DC similarity denoted by r (e.g., the within-twin pair correlation).

a N number of twin pairs unless otherwise noted, MZMC monozygotic monochorionic twins, MZDC monozygotic dichorionic twins, DZDC Dizygotic dichorionic twins, BWT birth weight, MC monochorionic, DC dichorionic, MZ monozygotic, DZ dizygotic, EFPTS East Flanders Prospective Twin Survey, ICC intra-class correlation, BCG Bacille Calmette-Guerin vaccine, NCPP NINCDS Collaborative Perinatal Project, BMI body mass index, LDL low density lipoprotein, HDL high density lipoprotein, NEFA non-esterified fatty acids, ICV intracranial volume, BWT birth weight, WISC Wechsler Intelligence Scale for Children, K-ABC Kaufmann Assessment Battery for Children, WAIS Wechsler Adult Intelligence Scale, CBCL Child Behavior Checklist.
during childhood. When effects were found, MC twins were generally more similar on the cognitive or personality assessment than DC twins were, suggesting that for some cognitive measures heritability estimates may be overestimated when not accounting for chorionicity.

We identified four studies that examined other behavioral phenotypes. For measures of temperament in very early childhood, MC twin similarity was equal to DC twin similarity (Chen et al. 1990; Riese 1999). Similarly, there was no chorionicity effect on twin similarity for prosocial behavior or Child Behavior Checklist (CBCL) total problems in childhood and adolescence (Hur 2007; Wichers et al. 2002). Thus, it is unlikely that chorionicity biases heritability estimates of toddler temperament and child and adolescent prosocial or problem behavior, although the studies were quite small and few in number.

Discussion

We presented the state of the literature on twin chorionicity in relation to a series of human outcome traits, and addressed the question of to what extent chorionicity differences in MZ twins may influence heritability estimates. We found a large body of literature on the effects of chorionicity on health and behavioral outcomes and a much smaller, but notable body of literature (30 articles in total) that examined chorionicity in relation to twin similarity, which could be used to draw tentative conclusions about whether chorionicity may bias heritability estimates. With only three studies from Asian populations and no studies from African populations, we were unable to draw even tentative conclusions about whether potential chorionicity biases may differ in populations with different twinning rates and MZ-MC/MZ-DC/DZ-DC proportions.

Consistent with the theory that some chorionicity effects could lead to overestimation and others to underestimation of heritability, there were instances of each across the many phenotypes considered here. However, firm conclusions should not be drawn since some of the outcomes were only examined in one or few studies and often sample sizes were small. In this same issue, van Beijsterveldt et al. (2015), using a sample of over 9000 twin pairs, report on chorionicity and heritability estimates on 66 phenotypes, including weight, height, motor milestones, child problem behaviors, cognitive function, wellbeing and personality. For only a few traits, within-pair similarity differed between MC-MZ and DC-MZ pairs. For traits influenced by birth weight, such as weight in young children MC twins were more discordant for 5 out of 13 measures. For traits where blood supply is important, MC-MZ twins were more concordant than DC-MZ for 3 traits. van Beijsterveldt et al. conclude that “the influence on the MZ twin correlation of the intra-uterine prenatal environment, as measured by sharing a chorion type, is small and limited to a few phenotypes”.

In our review, we also see that the most robust findings for chorionicity biasing heritability estimates were for birth weight (Vlietinck et al. 1989; Gielen et al. 2008; Touwslager et al. 2010; see Buzzard et al. 1983 for trend effect). This may be due to differences in placental sharing and vascularization between MZ-MC co-twins, which would reduce MC twin similarity and subsequently underestimate heritability of BW (see Table 1). That chorionicity could lead to underestimates of heritability for birth weight is interesting because despite the low heritability estimates from twin studies for birth weight, recent genome-wide association studies for this phenotype yielded significant hits (Horikoshi et al. 2013; Freathy et al. 2010).

Chorionicity may continue to effect heritability estimates of anthropometric traits later in life, but here effects are attenuated and less consistent. For example, heritability of weight and BMI are likely to be underestimated in childhood and adolescence (Gutknecht et al. 1999; Spitz et al. 1996; Mukherjee et al. 2009), while findings for height are inconsistent (Hur and Shin 2008; Spitz et al. 1996; Gutknecht et al. 1999). By adulthood, chorionicity did not appear to bias heritability estimates for the majority of studied anthropomorphic measures (e.g., various obesity-related measures, lung measures, or conventional and ambulatory blood pressure (Loos et al. 2001a; van den Borst et al. 2012; Souren et al. 2007; Fagard et al. 2003), however, chorionicity had an effect on fasting fibrogen (Loos et al. 2001b). It is important to note that specific outcomes have not been studied systematically. Therefore, it is unclear to what extent chronicity affects specific anthropometric outcomes across development.

Similarly, the effect of chorionicity on cognitive and personality measures in childhood and adolescence was mixed, although when effects were found they pointed to overestimation of heritability estimates. In measures of early brain and cognitive development, chronicity appeared to play no role (Mukherjee et al. 2009; Melnick et al. 1980; Welch et al. 1978). Chorionicity also appeared to play no role in the twin similarity for trisomy 21, vaccination responses, handedness, toddler temperament, or child and adolescent prosocial or problem behavior. One study found evidence that heritability of was overestimated without accounting for chorionicity (Davis and Phelps 1995; Davis et al. 1995); however, this finding has yet to be replicated.

Taken together, chorionicity biases heritability estimates for some outcomes at some points in during development. It is unclear for which outcomes heritability estimates are likely to be biased in a meaningful or measurable way. This review suggests that outcomes that are related to birth weight are more likely to be influenced by chorionicity. There is also qualitative evidence to suggest that chorionicity effects on heritability may be relatively greater for early compared to
later developmental outcomes, as was observed with anthropometric traits. With the exception of measures of birth weight and early growth, this review did not find evidence of any replicated effects of chorionicity on the heritability of human traits. Given the wide range of outcomes measured and small sample sizes it is unclear whether chronicity has a measurable effect on behavioral and cognitive measures. It thus would seem that concerns about heritability estimates based on the classical twin design, which relies on the equal environment assumption, are unwarranted when considering the prenatal environment.

**Funding** This work supported by NIH Grants: DA023134 (Knopik); a 2011 NARSAD Distinguished Investigator Grant [Boomsma: NARSAD (18633)]. Dr. Marceau is supported by T32 DA016184 and T32 MH019927.

**Compliance with Ethical Standards**

**Conflict of Interest** Kristine Marceau, Minni T. B. McMaster, Taylor F. Smith, Joost G. Daams, Catharina E. M. van Beijsterveldt, Dorret I. Boomsma, and Valerie S. Knopik declare that they have no conflicts of interest.

**Human and Animal Rights and Informed Consent** This review article does not contain any studies with human participants or animals performed by any of the authors. Each study reviewed in this article is assumed to have been conducted in accordance with the ethical standards of the institutional and/or national research committee and with the 1964 Helsinki declaration and its later amendments or comparable ethical standards. For this type of review, formal consent is not required.

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Appendix: search strategy

Appendix 1. Search strategy

Multiples (covered by pubmed search term "multiple birth offspring"):
exp triplets/ or exp twins/ or gemellus or exp triplet pregnancy/ or exp twin pregnancy/ or exp multiple pregnancy/
Specified on:
psychiatry/behavior/intelligence/genetics

CONCEPT 1 [multiples]
("Multiple Birth Offspring"[Mesh] OR Multiple Birth*[tw] OR Sextuplet*[tw] OR quadruplet*[tw] OR quintuplet*[tw] OR triplet*[tw] OR twins*[tw] OR twin*[tw] OR gemell*[tw] OR "Pregnancy, Multiple"[Mesh] OR multiple pregnanc*[tw] OR quadruplet pregnan*[tw] OR quintuplet pregnan*[tw] OR triplet pregnan*[tw] OR twin pregnan*[tw])

CONCEPT 2 compare different forms of chorionicity (monochorial dichorial, etc):
("Chorion"[Mesh] OR chorion*[tw] OR monochori*[tw] OR dichori*[tw] OR ("Embryonic and Fetal Development"[Mesh] OR fetal development[tw] OR embryo* development[tw]) AND outcome*[tw])

Probably too broad:
"Chorion"[Mesh] OR chorion*[tw] OR monochori*[tw] OR dichori*[tw] OR "Placentation"[Mesh] OR placentat*[tw]

Exclusion of animal studies:
NOT ("Animals"[Mesh] NOT "Humans"[Mesh])

Validation set PubMed:
14749653[uid] OR 9610996[uid] OR 16946215[uid] OR 10438438[uid] OR 6682287[uid] OR 18482623[uid] OR 988747[uid] OR 23355123[uid] OR 11084545[uid] OR 23101489[uid] OR 21727159[uid] OR 11665320[uid] OR 11360946[uid] OR 9822493[uid] OR 21830245[uid] OR 9505178[uid] OR 7487842[uid] OR 12044201 [UID]

PubMed 20140922:
(("Multiple Birth Offspring"[Mesh] OR Multiple Birth*[tw] OR Sextuplet*[tw] OR quadruplet*[tw] OR quintuplet*[tw] OR triplet*[tw] OR twins*[tw] OR twin*[tw] OR gemell*[tw] OR "Pregnancy, Multiple"[Mesh] OR multiple pregnanc*[tw] OR quadruplet pregnan*[tw] OR quintuplet pregnan*[tw] OR triplet pregnan*[tw] OR twin pregnan*[tw]) AND ("Chorion"[Mesh] OR chorion*[tw] OR monochori*[tw] OR dichori*[tw] OR (("Embryonic and Fetal Development"[Mesh] OR fetal development[tw] OR embryo* development[tw]) AND outcome*[tw])) AND outcome*[tw])
NOT ("Animals"[Mesh] NOT "Humans"[Mesh])

Added later:
(twin discordan*[tw] OR twin concordan*[tw] OR ((twin[tiab] or twins[tiab]) AND (concordan*[tiab] OR discordan*[tiab])) and outcome*[tw])

Specifications:
Genetics:
genetic*[tw] OR epigenetic*[tw] OR gene*[tw] OR genes*[tw] OR intelligence*[tw] OR iq*[tw] OR genotyp*[tw] OR genotyp*[tw] OR gene*[tw] OR genes*[tw] OR intelligence*[tw] OR iq*[tw] OR genotyp*[tw] OR genotyp*[tw] OR "genetics"[Subheading] OR "Genetic Techniques"[Mesh] OR Genetics*[Mesh] OR "Congenital, Hereditary, and Neonatal Diseases and Abnormalities"[Mesh] OR "Genetic Phenomena"[Mesh]

Psychiatry/behavior:
"Psychiatry and Psychology Category"[Mesh] OR "psychology"[Subheading] OR psychiatr*[tw] OR mental*[tw] OR psycholog*[tw] OR behavior*[tw] OR neuropsych*[tw]

"Neurologic manifestations":
nervous system diseases[mh] OR neuromorbidity[tw] OR neurologic*[tw]

Final search, PubMed 20150119 (20150119: 2111 hits / after deleting double records):
(("Multiple Birth Offspring"[Mesh] OR Multiple Birth*[tw] OR Sextuplet*[tw] OR quadruplet*[tw] OR quintuplet*[tw] OR triplet*[tw] OR twins*[tw] OR twin*[tw] OR gemell*[tw] OR "Pregnancy, Multiple"[Mesh] OR multiple pregnanc*[tw] OR quadruplet pregnan*[tw] OR quintuplet pregnan*[tw] OR triplet pregnan*[tw] OR twin pregnan*[tw]) AND ("Chorion"[Mesh] OR chorion*[tw] OR monochori*[tw] OR dichori*[tw] OR (("Embryonic and Fetal Development"[Mesh] OR fetal development[tw] OR embryo* development[tw]) AND outcome*[tw])) AND outcome*[tw])
NOT ("Animals"[Mesh] NOT "Humans"[Mesh])
Embase 1947 to Present, OvidSP, 20150119 (20150119: 1455 hits/after deleting double records):
exp multiple pregnancy/ or exp twins/
(multiple birth or multiple offspring or Sextuplet* OR quadruplet* OR quintuplet* OR triplet* OR twins
OR twin OR gemell* OR multiple pregnane* OR quadruplet pregnan* OR quintuplet pregnan* OR triplet
pregnan* OR twin pregnan*).ab,kw,ti
or/1-2
chorion/
(chorion* OR monochori* OR dichori*).ab,kw,ti
embryo development/ or fetus development/
((embryo* OR fetus OR fetal) ADJ3 development).ab,kw,ti
6 or 7
outcome?.mp
8 and 9
4 or 5 or 10
3 and 11
twin concordance/ or twin discordance/
((discordan* or concordan*) adj3 twin?).ab,kw,ti.
9 and (13 or 14)
12 or 15
gene/ or genetics/ OR genetic procedures/ or congenital disorder/ OR heridity/
(genetic* OR epigenetic* OR gene? OR intelligence OR iq OR genotyp* geno typ* OR phenotyp* OR pheno typ*).ab,kw,ti
exp psychiatry/ or exp psychology/
(psychiatr* OR mental OR psycholog* OR behavior* OR neuropsych*).ab,kw,ti
exp neurologic disease/
(neuromorbiditiy OR neurologic*).ab,kw,ti
or/17-22
16 and 23
(animal/ or animal experiment/ or animal model/ or nonhuman/ or rat/ or mouse/ or (rat or rats or mouse
or mice).ti.) not human/
24 not 25..dedup 26

Validation set (19 records):
("23355123" OR "2013110881" OR "21830245" OR "2011467101" OR "2008220392" OR
"2006431534" OR "20040456200" OR "2002202403" OR "2001184521" OR "200413754" OR
"1999283616" OR "1998397679" OR "1998179819" OR "1998106571" OR "1995299640" OR
"1983121977" OR "0977184937" OR "2001343682" OR "2013644154").an

Comment: twin (dis/con)cordance in combination with outcome searched for separately because of poor
representation of chrorionicity in bibliographic records.

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PsycINFO 1806 to Present, 20150119 (138 hits)
exp multiple births/
(multiple birth or multiple offspring or Sextuplet* OR quadruplet* OR quintuplet* OR triplet* OR twins
OR twin OR gemell* OR multiple pregnane* OR quadruplet pregnan* OR quintuplet pregnan* OR triplet
pregnan* OR twin pregnan*).ab,id,ti
or/1-2
(chorion* OR monochori* OR dichori*).ab,id,ti
prenatal development/
((embryo* OR fetus OR fetal) ADJ3 development).ab,id,ti
5 or 6
outcome?.mp
7 and 8
4 or 9
3 and 10
((discordan* or concordan*) adj3 twin?).ab,id,ti.
8 and 12
11 or 13..dedup15
(1996-16528-001 or 2002-01801-013).an
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