# Association of Birth Weight With Polymorphisms in the *IGF2*, *H19*, and *IGF2R* Genes

RONALD M. ADKINS, GRANT SOMES, JOHN C. MORRISON, JAMES B. HILL, ERIN M. WATSON, EVERETT F. MAGANN, AND JULIA KRUSHKAL

Departments of Pediatrics [R.M.A.] and Preventive Medicine [G.S., J.K.], University of Tennessee Health Science Center, Memphis, Tennessee 38103; Department of Obstetrics and Gynecology [J.C.M.], University of Mississippi Medical Center, Jackson, Mississippi 39216; Department of Obstetrics and Gynecology [J.B.H., E.M.W., E.F.M.], Naval Medical Center Portsmouth, Portsmouth, Virginia 23708

ABSTRACT: There is a substantial genetic component for birth weight variation. We tested 18 single nucleotide polymorphisms (SNPs) in the IGF2, H19, and IGF2R genes for associations with birth weight variation in 342 mother-newborn pairs (birth weight 2.1-4.7 kg at term) and 527 parent-newborn trios (birth weight 2.1–5.1 kg) across three localities. SNPs in the IGF2R (rs8191754; maternal genotype), IGF2 (rs3741205; newborn genotype), and 5' region of the H19 (rs2067051, rs2251375, and rs4929984) genes were associated with birth weight. Detailed analyses to distinguish direct maternal, direct newborn, and parent of origin effects for the most strongly associated H19 SNP (rs4929984) determined that the association of maternal genotype with newborn birth weight was due to parent of origin effects not direct maternal effects. That SNP is located near the CTCF binding sites that influence expression of the maternally imprinted IGF2 and paternally imprinted H19 locus, and there are statistically significant and independent opposite effects of the same rs4929984 allele, depending on the parent from which it was inherited. (*Pediatr Res* 68: 429-434, 2010)

Term, low birth weight infants are second only to premature infants in their rates of morbidity and mortality (1). As adults, individuals born small for GA (SGA) are at increased risk of multiple chronic illnesses (2). The link between reduced birth weight and adult illness might be explained by uteroplacental insufficiency that alters organ function and hormonal milieu to make the individual more susceptible to disease (3). Alternatively, genetic factors may exist that both reduce fetal growth and increase predisposition to disease (4).

Birth weight variation has a substantial genetic component. The likelihood of an SGA individual giving birth to an SGA child is  $3.5\times$  for men,  $4.7\times$  for women, or  $16.3\times$  if both parents were SGA, implying a multiplicative effect (5). Total heritability for birth weight is 42% to 45%, with a slightly stronger maternal *versus* paternal component (6,7). In addi-

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Correspondence: Ronald Adkins, Ph.D., Department of Pediatrics, University of Tennessee Health Science Center, 308 West Patient Tower, 50 North Dunlap, Memphis, TN 38103; e-mail: radkins1@uthsc.edu

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tion, SGA births both cluster in families and recur in successive generations (8,9).

IGF2 its receptor (IGF2R), and H19 are strong candidate genes for influencing birth weight variation. The maternal allele of IGF2 is imprinted in the fetus, meaning that the maternally transmitted allele is mostly silenced. Mouse knockouts of the paternal Igf2 allele are growth retarded, whereas fetuses overexpressing Igf2 exhibit overgrowth (10,11). In humans, the fetal growth restriction of Silver-Russell and fetal overgrowth of Beckwith-Wiedemann Syndromes (BWS) are associated with depressed versus elevated levels of IGF-II (12,13). H19 is an untranslated gene  $\sim$ 100 Kb from IGF2 for which the RNA transcript seems to be the functional product. H19 is growth inhibitory (14) and functionally antagonistic to *IGF2*. The paternal allele of *H19* is imprinted, and the binding of the CTCF protein to an imprinting control region (ICR) upstream of H19, regulated to a great extent by DNA methylation, reduces the IGF2 gene's access to a key enhancer located 3' of H19 (15,16). Finally, IGF2R exhibits inconsistent paternal imprinting in humans and functions mainly to remove IGF-II from the circulation (17–19). Correspondingly, mouse knockouts of Igf2-r exhibit fetal overgrowth (11). Numerous studies have shown association between birth weight or adult body mass and variation in the IGF2, H19, and IGF2R genes, although replication has been problematic (20-28).

By using three independent populations, we tested the association between single nucleotide polymorphisms (SNPs) in the *IGF2*, *H19*, and *IGF2R* genes and birth weight. Although some previously associated SNPs failed to replicate in our cohorts, we identify at least one SNP in each gene that associates with birth weight when present in the maternal or newborn genome. We also make the novel observation that maternal *versus* paternal transmission of the same *H19* allele has an opposite effect on birth weight.

Abbreviations: BWS, Beckwith-Wiedemann Syndrome; DMR, differentially methylated region; ICR, imprinting control region; IGF2AS, IGF2 antisense; IGF2R, IGF2 receptor; MBMI, maternal BMI; MBMI27, maternal BMI in excess of 27; SGA, small for GA; SNP, single nucleotide polymorphism

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### **METHODS**

Subjects. From 2003 to 2005, maternal blood, umbilical cord blood, and placental tissue were obtained from healthy pairs of mothers and newborns (Table 1) during labor and delivery at the University of Mississippi Medical Center (Jackson, MS; n = 104 pairs) and the Regional Medical Center (Memphis, TN; n = 123). From 2007 to 2008, healthy women (n = 115) were recruited during prenatal care into a longitudinal cohort study of cognitive development in Memphis, TN. Maternal blood was obtained from these women during prenatal care, and umbilical cord blood was obtained at delivery. Finally, from 2008 to 2009, maternal and umbilical cord blood was collected from mothers and newborns (n = 527) and mouthwash saliva (Listerine Cool Mint) from fathers (n = 502) at the Naval Medical Center Portsmouth, VA. All newborns were healthy at the time of delivery. Inclusion criteria were singleton pregnancy, >36-wk gestation, 18 to 40 y old. Exclusion criteria eliminated many additional factors influencing birth weight and included, among others, diabetes (type 1, type 2, and gestational), hypertension/vascular disease, preeclampsia, autoimmune disease, infectious disease (i.e. hepatitis, HIV), sickle-cell disease, uterine infection, illicit drug use, smoking, and birth defects. The Institutional Review Boards of the University of Tennessee Health Science Center, University of Mississippi Medical School, and Naval Medical Center at Portsmouth approved this study, and informed consent was obtained from all mothers and fathers.

Genotyping. SNPs were selected based on previously published associations with birth weight or body mass-related phenotypes or for the purpose of tagging haplotypes (Table 2; Fig. 1). Haplotype tagging SNPs were selected using SNPbrowser software version 3.5 (Applied Biosystems, Inc., Carlsbad, CA) and data from the International HapMap Project with the criteria  $r^2 \ge 0.8$  with untyped SNPs and minor allele frequency >20% in the Yoruban population (or Caucasian, if Yoruban data unavailable). IGF2R is a large gene, and haplotypes were tagged for only the 5′ one third of the gene. All genotyping used TaqMan reagents (Applied Biosystems, Inc.). Genotyping was ambiguous for H19 SNP rs2251375, and this was verified by either direct DNA sequencing or allele-specific PCR. Only 203 mother-newborn pairs were genotyped for rs2251375, all African-American. Genotyping was attempted for IGF2 SNP rs3741211, but allelic discrimination was poor, and this site is not reported.

Statistical analysis. All statistical analyses were performed using Stata SE version 10.1 (StataCorp, College Station, TX). Before inclusion of genetic data, a nongenetic model was constructed of potential confounders or effect modifiers, including GA (nearest week), maternal age, parity, newborn gender, maternal BMI (MBMI), and pregnancy weight gain. Each variable was plotted against birth weight to determine how best to present it. The relationship between MBMI and birth weight is nonlinear with birth weight increasing up to an MBMI of 27 and then decreasing beyond that. Therefore, this variable was represented as a linear spline of MBMI and MBMI27 (coded as 0 when MBMI  $\leq$ 27 and MBMI27 when MBMI  $\geq$ 27). The final model was identified by iteratively removing the least significant variable via "backward selection" until all remaining variables were significant ( $p \leq 0.05$ ). The final model included GA, newborn gender, MBMI, and MBMI27.

Association between birth weight and each SNP (0, 1, or, 2 copies of the minor allele) was tested in the combined Memphis and Jackson populations by addition of the genetic data to the nongenetic multiple linear regression

**Table 1.** Characteristics of the participants

		his, TN, and kson, MS	Portsmouth, VA		
Variable	Median	Range	Median	Range	
Race (maternal)					
African-American (N)	286		94		
Caucasian (N)	56		406		
Other $(N)$	0		27		
Mothers					
Age (yr)	22	17 to 40	26	18 to 44	
Parity	2	1 to 11	2	1 to 6	
Weight gain (kg)	15.3	-4.1 to 47.6	15.4	0 to 37.2	
BMI (kg/m <sup>2</sup> )	24.9	16.1 to 56.7	24.7	15.4 to 45.2	
Newborns					
GA (wk)	39	36 to 42	39	36 to 42	
Female (%)	44		49		
Weight (kg)	3.3	2.1 to 4.7	3.4	2.1 to 5.1	
Z	-0.1	-2.24 to $3.09$	0.0	-2.74 to $3.48$	

model. Differences in African and European allele frequencies (admixture) can produce false-positive associations (29,30). To adjust for admixture, we included an estimate of the proportion of African genomic ancestry in each newborn as a covariable, calculated by the program Admixmap (31). These estimates were based on European and African allele frequencies and used the candidate gene SNPs, 50 SNPs from 13 other protein-coding loci (32–34) and 12 unlinked ancestry informative SNPs (minor allele frequency difference >80% between Yoruban and European).

Whether an additive, recessive, or dominant genetic model best fits the data are unknown a priori. Performing all three tests at each SNP introduces a multiple testing burden. Although Bonferroni correction is too stringent because of nonindependence among SNPs in a small genomic region, permutation can determine empirical significance levels without a loss of power (35). We tested all three genetic models and calculated p values by permuting the dependent variable (birth weight) 100,000 times and determining the rank of the original regression t statistic (36). For the most strongly associated SNP (rs4929984), we genotyped the family trios in our replication sample from Portsmouth, VA, and used the polytomous logistic regression approach of Kistner et al. (37) to distinguish between direct maternal and parent of origin genetic effects. Because that method does not allow the inclusion of covariables, birth weight Z scores (standard deviations from the average birth weight for race, gender, and GA using national statistics for 2003) were used as the dependent variable. Some possibility for confounding exists because of a negative correlation between African-American-specific birth weight Z scores and genetically estimated proportions of African genomic ancestry (34). Hardy-Weinberg genotypic expectations were tested via an exact test (command genhwi), and measures of linkage disequilibrium were calculated using the pwld command (authored by David Clayton, Cambridge Institute for Medical Research) in Stata.

## **RESULTS**

General characteristics of the population. Overall characteristics of the mothers and newborns (Table 1) were similar among recruitment sites. Birth weights covered a broad range, with 2% to 4% (n=13 Memphis and Jackson, n=13 Portsmouth) SGA ( $\leq 10$ th percentile for GA, gender, and ethnicity) and 4% to 9% (n=12 Memphis and Jackson, n=47 Portsmouth) large for GA (LGA;  $\geq 90$ th percentile). Likely, the proportion of SGA and LGA newborns was cumulatively < 20% because of our selection of only healthy mothers and newborns, a decision intended to enrich for the genetic component of normal birth weight variation.

Genetic variants. IGF2 SNP rs680 deviated (p = 0.01) from Hardy-Weinberg genotypic expectations among African-American newborns, with a deficit of eight heterozygotes and nearly equal excesses of both homozygotes. Given that 34 statistical tests were performed, this could be a result of type I error.

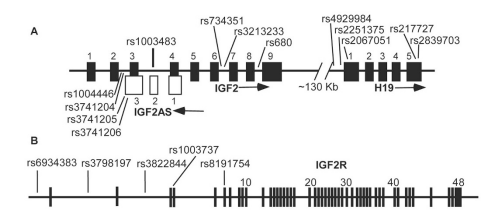
In Memphis and Jackson, four SNPs (IGF2R rs8191754 and H19 rs2067051, rs2251375, and rs4929984) among mothers and one SNP (IGF2 rs3741205) among newborns showed significant association with birth weight (Table 2). Linkage disequilibrium is high among the H19 SNPs ( $D' \ge 0.97$ ;  $R^2 =$ 0.20-0.72) but low between the H19 and IGF2 SNPs  $(D' \le 0.27; R^2 = 0)$ . Maternal genotypes of rs4929984 significantly (p = 0.04) associated with birth weight, but neither paternal nor newborn genotypes were associated (p > 0.27), among the Caucasian family trios from our replication population from Portsmouth, VA, who should exhibit few effects of admixture. By using polytomous logistic regression, which is insensitive to population stratification, in all Portsmouth trios, parent of origin (p = 0.016), but not direct maternal genetic association (p = 0.396), was significant for rs4929984. In regression (Table 3) of birth weight on GA, gender, MBMI,

Table 2. Statistical	analyses	of SNPs in	Momphis and	Lackson	nonulations	N =	3421
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	Association with birth weight $(p)$								
	Minor allele frequency* Maternal					Fetal			
	African-American	Caucasian	Additive	Dominant	Recessive	Additive	Dominant	Recessive	
IGF2R									
rs6934383	0.36	0.18	0.8452	0.8122	0.9152	0.9916	0.9243	0.9612	
rs3798197	0.41	0.26	0.2263	0.3740	0.2938	0.7921	0.5283	0.9276	
rs3822844	0.44	0.30	0.8446	0.6223	0.8994	0.8848	0.3191	0.5468	
rs1003737	0.48	0.54	0.6813	0.4419	0.9567	0.9426	0.8541	0.7725	
rs8191754	0.13	0.07	0.0462	0.9911	0.0283	0.7728	0.4536	0.9065	
IGF2-H19									
rs2839703	0.12	0.42	0.2200	0.6911	0.2025	0.1672	0.8648	0.0800	
rs217727	0.08	0.13	0.1734	0.4075	0.2073	0.1856	0.2289	0.2670	
rs2067051	0.09	0.28	0.0318	0.5225	0.0305	0.9735	0.1093	0.5073	
rs2251375	0.38	_	0.0140	0.1689	0.0172	0.3672	0.7188	0.3177	
rs4929984	0.31	0.48	0.0013	0.0009	0.0276	0.2268	0.5223	0.2307	
rs680	0.07†	0.21	0.3523	0.1610	0.5963	0.6281	0.3878	0.8071	
rs3213233	0.10	0	0.9765	0.7003	0.8837	0.8319	0.5404	0.9467	
rs734351	0.25	0.30	0.7531	0.8741	0.6393	0.6327	0.3547	0.9209	
rs1003483	0.16	0.50	0.6032	0.9680	0.5427	0.1579	0.2132	0.2805	
rs3741206	0.28	0.45	0.1436	0.9632	0.0621	0.7680	0.9763	0.6904	
rs3741205	0.30	0.26	0.1754	0.2712	0.2668	0.0383	0.2944	0.0385	
rs3741204	0.43	0.30	0.7660	0.8057	0.8123	0.1842	0.2582	0.2966	
rs1004446	0.44	0.31	0.9567	0.9054	0.8566	0.1475	0.4209	0.1332	

<sup>\*</sup> Among newborns.

<sup>†</sup> Test of Hardy-Weinberg genotypic expectations, p < 0.05 among newborns.



**Figure 1.** *A) IGF2, IGF2AS*, and *H19* genes. *B) IGF2R* locus. The locations of genotyped SNPs are shown. Although not to scale, the relative spacing of exons is represented. *Arrows* indicate the direction of transcription.

**Table 3.** Multiple linear regression of birth weight (g) on parental origin of alleles at rs4929984 in the Memphis, Jackson, and Portsmouth African-American and Caucasian participants

	All $(N = 686)$		Primiparous $(N = 228)$		Multiparous $(N = 458)$	
Variable	β	p	β	p	β	p
GA (wk)	124	< 0.001	127	< 0.001	135	< 0.001
Ethnicity (African-American/Caucasian)	249	< 0.001	226	< 0.001	265	< 0.001
Gender (male/female)	-119	< 0.001	-76	0.135	-147	< 0.001
MBMI	30	< 0.001	39	< 0.001	26	< 0.001
MBMI27	-33	< 0.001	-50	0.004	-29	0.005
Paternally transmitted allele (C/A)	78	0.012	178	0.001	33	0.392
Maternally transmitted allele (C/A)	-89	0.007	-86	0.136	-84	0.039

MBMI27, ethnicity, and alleles of inferred parental origin in only the African-American and Caucasian participants from all locations, both the maternally (p = 0.007) and paternally transmitted (p = 0.012) alleles were significant, indicating independent associations with both alleles. Maternal transmission of the A allele was associated with an 89 g decrease in birth weight, whereas paternal transmission was associated

with a 78 g increase. This implies opposite associations with the same allele, depending on whether it was paternally or maternally transmitted. We can assume that the paternal genotype does not directly affect fetal growth. However, direct maternal, direct fetal, and parent-of-origin effects (both maternal and paternal alleles) may be operating singly or in any combination, making it very difficult to isolate one effect from

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**Table 4.** Effect of the paternally transmitted allele of rs4929984 when maternal genotype and her transmitted allele are held constant

Maternal genotype (Z)	Paternal allele	Z (SD)	<i>N</i> *	p†
CC (0.02)	С	-0.05 (0.06)	197	0.09
	A	0.11 (0.08)	127	
AA (-0.19)	C	-0.32(0.09)	79	0.05
	A	-0.02(0.12)	65	

<sup>\*</sup> Memphis, Jackson, and Portsmouth combined.

the others. One approach is to eliminate variation in some of these factors while testing for the others. Therefore, to better understand the possibility of opposing parent-of-origin associations, we considered only homozygous mothers from all localities, which holds constant both the maternal genotype and the maternally transmitted allele (Table 4). Newborns whose mothers were CC (maternally transmitted C allele) versus AA had significantly higher birth weight Z scores (n = 468; t = 2.25; two-sided p = 0.025), as would be expected (Table 2). Within each category of homozygous mother, newborns who received an A allele from their father had a higher average Z score. The significance of this difference was marginal, likely because of small sample sizes within these narrow subcategories.

## DISCUSSION

IGF2R. The association of IGF2R with birth weight is inconsistent. Among 884 Japanese mothers and newborns, rs8191754 CC homozygotes had the highest birth weight (p =0.02), with heterozygotes intermediate between homozygotes (23). In contrast, among 97 Greek newborns, neither neonatal rs8191754 genotypes nor parental origin of alleles were associated with birth weight (24). We found association between maternal, but not newborn, rs8191754 genotypes and birth weight. In our case, GG homozygotes were heaviest (3425 g), but this was based on only five individuals and is probably unreliable. A contrast (t test, Ho:  $BW_{CC} \neq BW_{CG}$ ; p = 0.04) between the genotypes CC (3301 g; n = 242) and CG (3186 g; n = 79) should be more accurate and is consistent with the Japanese pattern. Although consistent between Japanese and our cohort, the association of rs8191754 with birth weight remains equivocal given the conflicting association with neonatal versus maternal genotypes and the modest effect size estimated.

*IGF2.* We surveyed eight SNPs in the *IGF2/IGF2AS* (*IGF2* antisense transcript) genes, some of which have shown association with birth weight, fetal growth disorders, or mature body mass. Polymorphism rs680 is associated with birth weight among Japanese (23) but not among Brazilians or Belgians (21,38), and with adult male BMI among some British cohorts (20,22,27,28), but not others (38). In Memphis and Jackson, we found no association between birth weight and either maternal or newborn rs680 genotypes or in analyses of individual newborn genders (p > 0.08).

Polymorphism rs3741205 is the defining SNP of a CAGA haplotype (rs1004446, rs3741204, rs3741205, and rs3741206)

in a differentially methylated region (DMR) termed DMR0 in intron 2 of IGF2 that is enriched in patients with sporadic BWS (39). BWS is characterized by pre- and postnatal overgrowth and is commonly associated with altered methylation of DMRs near H19 and KCNQ1 (500 Kb from IGF2) and their loss of imprinting. Among Japanese, inferred paternal inheritance of IGF2 haplotype CTG at rs1003484, rs3741211, and rs3741206 was associated with a higher frequency of SGA births (25). We observed a significant association between birth weight and newborn genotypes at rs3741205 for both an additive and recessive (TT versus GT and GG) model. The presence of a G allele is associated with an increase of 74 g after adjustment for covariables. Unlike among the Japanese population, we find no significant association with inheritance of the inferred paternal allele (p > 0.2 in all subjects and)within newborn genders).

The CAGA haplotype more frequent among patients with BWS was associated with loss of methylation of an exon 10 DMR (KvDMR1) in the maternally imprinted *KCNQ1* gene (39). Lack of maternal methylation of KvDMR1 permits expression of an antisense transcript (*KCNQ10T1*), which reduces expression of the *KCNQ1* and *CDKN1C* genes, and this is observed in about half of BWS cases. Genetic variation in the region of *IGF2* DMR0 may be associated with changes in DNA methylation at KvDMR1. Alternatively, given its proximity to the gene, this variant may influence the expression of *IGF2AS*, whose function is presently unclear.

H19 region. Among British mothers and mother-newborn pairs, rs217727 in the paternally imprinted H19 gene exhibited independent associations of maternal and newborn rs217727 genotypes with birth weight and a further association of maternal rs217727 genotypes with umbilical cord IGF-II levels (26). There was no evidence of a parent of origin effect. We found no birth weight association with either maternal or newborn genotypes of rs217727. The minor allele of rs217727 is rare (8-13%; five newborn and three maternal TT homozygotes) among our subjects, reducing statistical power in regression analyses. Nevertheless, restricting attention to CC (268 mothers and 275 newborns) and CT (59 mothers and 51 newborns) individuals, the relationship to birth weight remains nonsignificant (p > 0.3), despite a 45 g (newborn, adjusted for covariables) or 55 g (maternal) increase in birth weight among CT heterozygotes.

In Memphis and Jackson, we found strong association between birth weight and maternal genotypes at rs4929984 and weaker association with rs2067051 and rs2251375. Linkage disequilibrium is high (D'=0.94-1) among those SNPs, meaning that the association with each site is not independent and indicating 1) only one of the SNPs is functionally important, 2) they are functionally important as a haplotype, or 3) all three are in linkage disequilibrium with a functional genetic variant that was not surveyed.

The association of the *H19* rs4929984 polymorphism was replicated in the Portsmouth family trios. Polytomous logistic regression (37) is capable of disentangling direct maternal, direct fetal, and parent of origin associations and reveals that the association is due to parent of origin, not to direct association with maternal genotypes. Analyses of the effect of the

<sup>†</sup> Two-sided t test.

maternally and paternally transmitted alleles at rs4929984 indicate that maternal inheritance of the A allele results in an average 89 g decrease (adjusted for covariables; Table 3), whereas paternal inheritance of the same allele results in an average 78 g increase in birth weight. These associations remain if only the paternally or maternally transmitted alleles are included in the regression model, but with smaller effect sizes (maternal allele, 74 g, p = 0.02; paternal allele, 60 g, p =0.05), consistent with the possibility of mutual confounding because of opposite associations with the same allele depending on parental origin. This may be why paternal genotypes are not directly associated with birth weight in the Virginia population. The maternally transmitted allele may confound the ability to detect an association with paternal genotype because of its opposite and slightly larger effect. Analyses within maternal genotypic classes (Table 4) that eliminate variation because of the maternal genotype and maternally transmitted allele illustrate that the paternal allele has a consistent association in the opposite direction of the maternal allele.

The association of rs217727 with birth weight (26) was strongest for the first born child, suggesting a contribution from *in utero* growth constraint in first pregnancies. To determine whether a similar effect exists among our participants, we distinguished between first and subsequent pregnancies (Table 3). Among multiparous women, the maternally transmitted allele had a significant effect but not among primiparous women. However, the estimated effect size of the maternal allele was similar within both groups, and the nonsignificant result may be because of lack of statistical power in the comparatively small subset (n = 228). Interestingly, the paternally transmitted allele was significantly associated only among primiparous mothers, suggesting a pattern similar to that previously observed for maternal genotypes of rs217727.

A functional basis for the opposite direction of association of parentally transmitted alleles of rs4929984 requires explanation. The imprinting of both IGF2 and H19 is under the control of an ICR with seven CTCF binding sites that resides 500 to 5,200 nucleotides upstream of *H19*. The significantly associated SNPs near H19 flank the ICR (rs4929984 upstream; rs2251375 and rs2067051 downstream). Therefore, it is unlikely that these SNPs directly disrupt the insulator function of CTCF binding. However, these SNPs may be in linkage disequilibrium with variants that alter CTCF binding or affect the methylation of the ICR and its ability to block IGF2 access to an enhancer 3' of H19. Alternatively, these SNPs, or ones in linkage disequilibrium with them, may affect the level of H19 transcription or stability. Given the competitive use of the H19 ICR by IGF2 and H19 and the activity of those loci on different parental chromosomes, it is quite feasible that the same genetic variant could promote fetal growth when on the paternally inherited chromosome and retard fetal growth when on the maternally inherited chromosome. Interestingly, the pattern of birth weight variation that we have observed with respect to rs4929984 closely resembles the "bipolar" (homozygotes similar, with one type of heterozygote smaller and the other larger) dominance pattern of imprinting for postnatal growth found at some genomic regions in an intercross of the large *versus* small inbred mouse lines (40).

In summary, we identified polymorphisms in *IGF2R*, *IGF2*, and near *H19* that are associated with birth weight. Although the *H19* association was replicated in a second cohort, the sample sizes were small, and these results should be validated in larger, independent populations. Recently, a large genomewide association meta-analysis (41) was performed that identified two strong candidates for birth weight association, neither of which were in the loci studied here. However, that study focused entirely on the child's genotype and would not be expected to identify the associations we primarily observed with maternal genotypes.

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