

Twin study confirms virtually identical prenatal alcohol exposures can lead to markedly different fetal alcohol spectrum disorder outcomesfetal genetics influences fetal vulnerability

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Abstract

Background: Risk of fetal alcohol spectrum disorder (FASD) is not based solely on the timing and level of prenatal alcohol exposure (PAE). The effects of teratogens can be modified by genetic differences in fetal susceptibility and resistance. This is best illustrated in twins.

Objective: To compare the prevalence and magnitude of pairwise discordance in FASD diagnoses across monozygotic twins, dizygotic twins, full-siblings, and half-siblings sharing a common birth mother.

Methods: Data from the Fetal Alcohol Syndrome Diagnostic & Prevention Network clinical database was used. Sibling pairs were matched on age and PAE, raised together, and diagnosed by the same University of Washington interdisciplinary team using the FASD 4-Digit Code. This design sought to assess and isolate the role of genetics on fetal vulnerability/resistance to the teratogenic effects of PAE by eliminating or minimizing pairwise discordance in PAE and other prenatal/postnatal risk factors.

Results: As genetic relatedness between siblings decreased from 100% to 50% to 50% to 25% across the four groups (9 monozygotic, 39 dizygotic, 27 full-sibling and 9 half-sibling pairs, respectively), the prevalence of pairwise discordance in FASD diagnoses increased from 0% to 44% to 59% to 78%. Despite virtually identical PAE, 4 pairs of dizygotic twins had FASD diagnoses at opposite ends of the fetal alcohol spectrum—Partial Fetal Alcohol Syndrome versus Neurobehavioral Disorder/Alcohol-Exposed.

Conclusion: Despite virtually identical PAE, fetuses can experience vastly different FASD outcomes. Thus, to protect all fetuses, especially the most genetically vulnerable, the only safe amount to drink is none at all.

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Introduction

The effects of teratogens can be modified by genetic differences in fetal susceptibility and resistance [1]. Fetal alcohol syndrome (FAS) is a permanent birth defect syndrome caused by maternal consumption of the teratogen alcohol

during pregnancy. FAS is characterized by growth deficiency, a unique cluster of minor facial anomalies and central nervous system (CNS) structural and/or functional abnormalities. Not all fetuses exposed to and damaged by prenatal alcohol exposure (PAE) have FAS. The physical, cognitive, and behavioral deficits observed among individuals with PAE are not dichotomous, that is either normal or clearly abnormal. Rather, the outcomes all range along separate continua from normal to clearly abnormal [2]. This full range of outcomes caused by PAE is called Fetal Alcohol Spectrum Disorders (FASD). Diagnoses like FAS, Partial FAS (PFAS), Static Encephalopathy/Alcohol Exposed (SE/AE) and Neurobehavioral Disorder/Alcohol Exposed (ND/AE) fall broadly under the umbrella of FASD [2,3].

Fetal risk of damage from PAE is not just dependent on the timing, frequency, and quantity of exposure. Fetal alcohol spectrum disorders (FASD) are caused by a complex interaction of genes and environment, and are regulated by both parental and fetal genes [4]. Fetal genetics influences a fetus' vulnerability to the teratogenic effects of PAE [5]. In a 1993 study of 16 monozygotic and dizygotic twin pairs, Streissguth and Dehaene [6] reported 100% pairwise concordance in FASD diagnoses among monozygotic twin pairs, while dizygotic twins were only 64% concordant. The outcomes of that study strongly suggested that genetic loci regulate susceptibility to, or resistance against FASD. This 1993 study was conducted prior to the creation of rigorous FASD diagnostic systems. Patients were diagnosed as FAS or Fetal Alcohol Effects. If two fetuses exposed to identical levels of alcohol can experience vastly different FASD outcomes, this would have important implications for public health messaging and the setting of exposure thresholds in FASD diagnostic guidelines.

The purpose of this study was to compare the prevalence and magnitude of pairwise discordance in FASD diagnoses across four groups of sibling pairs: monozygotic twins, dizygotic twins, fullsiblings, and half-siblings sharing a common birth mother. All sibling pairs had virtually identical or reportedly similar levels of PAE, were raised together, were diagnosed by the same interdisciplinary team using the 4-Digit Diagnostic Code and were identical or similar in age at the time of diagnosis. This sibling-pair design sought to more fully assess and isolate the role of genetics on fetal vulnerability to the teratogenic effects of PAE by eliminating or minimizing pairwise discordance in age, PAE and other prenatal and postnatal risk factors.



Specific aims

- To determine if the prevalence of FASD diagnostic discordance was higher among dizygotic twin pairs than among monozygotic twin pairs.
- 2. To determine if the prevalence of FASD diagnostic discordance increases as the proportion of genome shared between sibling-pairs decreases across the four study groups: monozygotic twins, dizygotic twins, full-siblings, and half-siblings sharing a common birth mother.
- 3. To document the greatest magnitude of FASD diagnostic discordance observed between twin pairs with virtually identical PAE. Can twins with virtually identical PAE present at opposite ends of the fetal alcohol spectrum?
- To estimate the proportion of phenotypic variance in FASD diagnoses due to genetic factors (heritability).

Methods

A retrospective study was conducted using data collected from twin and sibling pairs that received a FASD diagnostic evaluation at the University of Washington Fetal Alcohol Syndrome Diagnostic & Prevention Network (FASDPN).

FASD diagnostic method

FASD diagnoses were rendered using the FASD 4-Digit Diagnostic Code. It is described in full by Astley [2,7]. Briefly, the 4 digits of the FASD 4-Digit Code reflect the magnitude of expression of the 4 key diagnostic features of FASD, in the following order: 1) growth deficiency, 2) FAS facial phenotype, 3) CNS structural/functional abnormalities, and 4) PAE (Figure 1). The magnitude of expression of each feature is ranked independently on a 4-point Likert scale, with 1 reflecting complete absence of the FASD feature and 4 reflecting a strong "classic" presence of the FASD feature. Each Likert rank is specifically case defined. There are a total of 102 4-Digit Codes that fall broadly under the umbrella of FASD (Figure1 A-D).

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		3		4	3		4						
	4	H and W ≤ 3 %		All 3 features	Structural / Neurological Abnormalities	C	onfiri Hig	med h		4	~	A CONTRACT	
nk	3	H or V ≤ 3 %		2.5 features	Severe Dysfunction	С	onfiri	med		3		3	
Ra	2	H and V all els	V 1-2 features Mode Dysfu		Moderate Dysfunction	ι	Jnkno	own		2	~~	57	
	1	H and W > 10 %		No features	No Dysfunction	С	onfiri Abse	med ent		1	Y	T	
		Grow	⁄th	Face	CNS	А	lco	hol	4	A Lip-	Philtrum Guide I	Philtrum Guide	F
				Diagnosis		Gro	wth	FAS Fac	e	CNS		Alcohol	
1.	FA	S		Fetal Alcoho	Syndrome	gro	wth	face		severe		alc	
2. PFAS Partial FA			FAS			face		severe		alc			
3.	SE	AE	Sta	atic Encephalopa	athy / Alc Exposed					severe		alc	
4. ND/AE			Neu	robehavioral Dis	order / Alc Exposed						moderate	alc	

Figure 1: FASD 4-Digit Diagnostic Code. A) Abbreviated case-definitions for the fetal alcohol spectrum disorder (FASD) 4-Digit Code [2]. The 4-Digit Code 3434 is one of 12 4-Digit Codes that fall under the diagnostic category FAS. B) The Rank 4 FAS facial phenotype requires 3 features: I) palpebral fissure lengths 2 or more standard deviations below the mean; 2) a smooth philtrum (Rank 4 or 5 on the University of Washington Lip-Philtrum Guide); and 3) a thin upper lip (Rank 4 or 5 on the University of Washington Lip. Philtrum Guide). C and D) The 4-Digit Code produces four-diagnostic subgroups under the umbrella of FASD: FAS (Diagnostic Categories A, B); PFAS (Diagnostic Category C); SE/AE (Diagnostic Categories E,F); and ND/AE (Diagnostic Categories G,H). Abbreviations: CNS: central nervous system; H: height percentile; W: weight percentile.

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These four clinically codes cluster under meaningful FASD diagnoses: fetal alcohol syndrome (FAS) (4-Digit Code Diagnostic Categories A, B): Partial FAS (PFAS) (Diagnostic Static Encephalopathy/Alcohol-Category C); Exposed (SE/AE) (Diagnostic Categories E, F); and Neurobehavioral Disorder/Alcohol-Exposed (ND/AE) (Diagnostic Categories G, H) (Figure 1D). Individuals that did not meet criteria for one of these FASD diagnostic classifications were identified in this study as Not FASD/Alcohol-Exposed (Diagnostic Categories I, J).

Study groups

The sibling pairs were partitioned into four study groups: 1. monozygotic twins, 2. dizygotic twins, 3. full-siblings, and 4. half-siblings sharing a common birth mother. Monozygotic twin pairs share virtually 100% of their genome. Dizygotic twin pairs and full-sibling pairs share, on average, 50% of their genome. Half-sibling pairs with a common birth mother share, on average, 25% of their genome [8,9].

Data from all twin and sibling pairs that met the following inclusion criteria were used in this study:

- Sibling pairs were monozygotic twins, dizygotic twins, full-siblings, or half-siblings sharing the same birth mother.
- Both siblings received an FASD diagnostic evaluation at an FASDPN clinic by the same interdisciplinary team using the 2004 FASD 4-Digit Diagnostic Code.
- Siblings did not present with another genetic syndrome.
- Age at diagnosis could range from newborn to adult. Effort was made to select pairs that both fit into one of three age ranges at the time of diagnosis (0-3 years, 4-8 years, 9 or more years). This minimized the chance that FASD diagnostic contrasts between pairs may be due to one sibling being too young to fully assess or comparably assess brain function.
- All siblings had confirmed PAE. Twin pairs, by definition, had virtually identical PAE. Full sibling pairs and half-sibling pairs had to have concordant Alcohol Ranks (e.g., both siblings had to have Rank 3



alcohol exposure or both had to Rank 4 alcohol exposure).

- Effort was made to select siblings raised together who experienced identical or similar other prenatal and postnatal risk factors.
- Siblings could be of any gender or race.

Data set

All data collected during an FASD diagnostic evaluation at the FASDPN are entered into the FASDPN database with patient consent and University of Washington Human Subjects Division approval. Approximately 3,000 patients have been evaluated in the clinic to date. The data document patient demographics, PAE, all other reported prenatal and postnatal adverse exposures and events and measures of growth, facial features, and structural and/or functional brain abnormalities used to derive the FASD 4-Digit Code. These data are recorded on three standardized diagnostic forms: the New Patient Information Form, FASD Diagnostic Form, and FAS Facial Photographic Analysis Report posted on the FASDPN website www.FASDPN.org [2,8].

Key data used in this study included the patient's FASD 4-Digit Code, FASD diagnostic category (FAS, PFAS, SE/AE, ND/AE and Not FASD/AE), and their Growth, Face, CNS and Alcohol Ranks (Figure 1). The CNS Rank in the 4-Digit Code serves two purposes: 1) Ranks 1 through 4 document the probability of underlying CNS structural abnormality (Rank 1: unlikely; Rank 2: possible; Rank 3: probable; and Rank 4: definite). 2) Ranks 1 through 3 also document the magnitude of CNS dysfunction as measured using standardized neuropsychological tools (Rank 1: no dysfunction; Rank 2: moderate dysfunction; and Rank 3 severe dysfunction). The CNS functional Ranks 1-3 introduced by the 4-Digit Code were case-defined to predict increasing likelihood of structural CNS abnormality-a predictive correlation that was subsequently confirmed through magnetic resonance imaging [7]. To distinguish these two CNS measures in the current study, they are labeled CNS1-4 and CNS1-3. PAE is ranked by the 4-Digit Code on a 4-point Likert scale (Figure 1A). Only subjects with Rank 3 or Rank 4 PAE were enrolled in this study. An Alcohol Rank 4 is assigned when PAE is confirmed and reported to be high risk (generally high peak blood alcohol concentrations

delivered at least weekly in early pregnancy). An Alcohol Rank 3 is assigned when PAE is confirmed, but the amount reported is low to moderate risk (designated as PAE Rank 3b in this study) or the details on the amount and timing are unknown (designated as PAE Rank 3a in this study). The information used to generate the Alcohol Rank is presented on page 8 of the 4-Digit Code FASD Diagnostic Form [2]. In preparation for a FASD diagnostic evaluation, efforts are made to document the quantity, frequency and timing of maternal alcohol use before and during the index pregnancy. Although 99.5% of patients evaluated at the FASDPN clinic have confirmed PAE, only 30 to 40% have quantity, frequency and/or timing of exposure detailed in their records [10]. Recall error and reporting bias likely impact the accuracy of this more detailed information, therefore the more global measure of PAE "Alcohol Rank" was used as the primary measure of PAE risk in this study. In addition to the risk posed by PAE, measures of other prenatal and postnatal risks were also used in this study. Other prenatal risk factors documented in the FASDPN clinical database include poor prenatal care, pregnancy complications, presence of abnormalities. syndromes/genetic other and prenatal exposure to other substances (e.g., medications, tobacco, illicit drugs, and/or other teratogens). The 4-Digit Code ranks the magnitude of these other prenatal risks in a single composite measure labeled "Other Prenatal Risks Rank". Rank 1 equals no risk; Rank 2 equals unknown risk; Rank 3 equals some risk; and Rank 4 equals high risk. Rank 4 is assigned when there is exposure to another teratogen (e.g., Dilantin) or when another syndrome or genetic condition is present (e.g., Down syndrome, Fragile X, etc.). Rank 3 is assigned to all other prenatal risks. Postnatal risk factors documented in the FASDPN database include perinatal complications, number of home placements, physical and/or sexual abuse, neglect, and trauma. The 4-Digit Code ranks the magnitude of these other postnatal risks in a single composite measure labeled "Other Postnatal Risks Rank". Rank 1 equals no risk; Rank 2 equals unknown risk; Rank 3 equals some risk; and Rank 4 equals high risk. Rank 4 is used to note severe postnatal circumstances that have been shown to have a significant adverse effect on development in most instances. Examples include physical/sexual abuse, multiple home placements, trauma, and severe neglect) [2]. Rank 3 is used to note conditions akin



to those in Rank 4, but the circumstances are less severe.

Statistical analyses

The primary focus of the study was to compare the prevalence of discordant FASD diagnoses between sibling pairs across the 4 study groups. Descriptive statistics (means, SD, and proportions expressed as valid percentages (e.g., subjects with missing data are not included in the denominator)) were used to profile the demographic and clinical outcomes of the 4 study groups. The chi-square test with tests for linear trend was used to compare proportions between the study groups. One-way ANOVA was compare outcomes used to measured on a continuous scale between the study groups. Twotailed p-values less than 0.05 were interpreted as statistically significant.

Heritability is formally defined as the proportion of phenotypic variation that is caused by genotypic variation in a population. FASD is not a genetic disorder, but fetal genetics appears to modify the teratogenic effects of PAE [5,6]. Heritability has historically been estimated from studies of twins. Monozygotic twin pairs share essentially 100% of their genome. Dizygotic twin pairs share, on average, 50% of their genome. If a trait appears to be more similar in monozygotic twins than in dizygotic twins (when the twin pairs were raised together in the same environment), genetic factors likely play an important role in determining (or modifying) that trait. By comparing a trait in monozygotic twins versus dizygotic twins, one can calculate an estimate of its heritability. Heritability estimates range from 0% to 100%. A heritability close to 0% indicates that almost all of the variability in a trait is due to environmental factors (e.g., PAE and other prenatal and postnatal risk factors), with very little influence from genetic differences. A heritability estimate close to 100% indicates that almost all of the variability in a trait comes from genetic differences, with very little contribution from variability in environmental factors. When a phenotype is determined by a combination of genetic and environmental factors, heritability will be somewhere between 0% and 100%. Comparing discordance for monozygotic versus dizygotic twins allows an indirect estimate of the importance of genetic factors in producing the phenotype. Heritability estimates based on twin discordance studies can be simplistically viewed as:

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percent heritability = ((dizygotic discordance minus monozygotic discordance) / dizygotic discordance) *100 [9,11]. It is important to understand that heritability does not indicate what proportion of a trait is determined by genes and what proportion is determined by environment. A heritability of 80% does not mean that a trait is 80% caused by genetic factors; it means that 80% of the variability in the trait in a population is due to genetic differences. Heritability measures the fraction of variation between individuals in a population that is due to their genotypes.

Results

Demographic and clinical profiles of the four study groups

The study selection criteria generated 84 sibling pairs broken into four study groups (monozygotic twins, dizygotic twins, full siblings and half sibling sharing the same birth mother) with key factors that defined and differentiated the groups (Table 1).

Table 1. Key factors that defined and differentiated the 4 study groups.

		Study	Groups	
Features shared between sibling pairs	Monozygotic twins	Dizygotic twins	Full-siblings	Half-siblings
	9 pairs	39 pairs	27 pairs	9 pairs
Birth mother	identical	identical	identical	identical
Birth father	identical	identical	identical	different
Genome shared	~100%	~50%	~50%	~25%
Prenatal alcohol exposure	virtually identical	virtually identical	100% same Rank	100% same Rank
Other prenatal risks	virtually identical	virtually identical	100% same Rank	88% same Rank
Siblings raised together	100%	100%	96%	100%
Other postnatal risks	100% same Rank	87% same Rank	83% same Rank	75% same Rank
Matched in age within one of 3 age ranges (0-3; 4-8, 9+ years)	100%	100%	*93%	*89%

* 2 pairs of full-siblings and 1 pair of half-siblings had one sibling that was in a younger age category. In each of these 3 pairs, the younger sibling had the more severe FASD diagnostic outcome.

The demographic and FASD diagnostic profiles of the study sample (Table 2) were highly representative of the entire FASDPN clinic population (n = 3,000) from which it was selected [10]. The gender and age distributions were comparable between the 4 study groups. Race was 100% concordant across all 84 twin/sibling pairs. The number of full and half sibling pairs included in the study may appear smaller than one would expect from a patient population of 3,000. A number of factors inherent in the FASDPN clinical

dataset limited the number of full and half sibling pairs available for inclusion in the study. In general, 85% of the patients evaluated by the FASDPN are in foster/adoptive care—no longer living with their birth parents. Confirmation of full or half sibling status requires knowledge of both birth parents' names. This is typically available on only half of the FASDPN patient population. Of the 54% (n = 1,617) with birth parent names available, 8% (n = 129) were siblings (full or half). Seventy-two of the



129 full and half siblings met the inclusion criteria for the study.

The primary reason siblings failed to meet the study's inclusion criteria were they were too

different in age (e.g., infant vs adolescent) at the time of their FASD evaluation to draw valid conclusions regarding the concordance/discordance of their diagnoses.

Table 2. Comparison of demographic and FASD clinical profiles between the four study groups.

Demographic and	Mono	zygotic	Dizygoti	c	Full-siblings		Half-siblings		Total	
Clinical Characteristics	N = 18	N = 18 (9 pairs)		N = 78 (39 pairs)		(27 pairs)	N = 18 (9 pairs)		N = 168 (84 pairs)	
Gender (N pairs; valid %)										
female-female	4	44.4	10	25.6	7	25.9	2	22.2	23	27.4
male-male	5	55.6	10	25.6	11	40.7	4	44.4	30	35.7
Mixed gender	0	0.0	19	48.7	9	33.3	3	33.3	31	36.9
Overall proportion of female subjects	8/18	44.4	37/78	47.4	23/54	42.6	7/18	38.9	75/168	44.6
Race (N; valid %)										
Caucasian	4	22.2	32	41.0	32	59.3	14	77.8	82	48.8
African American	0	0.0	0	0.0	0	0.0	0	0.0	0	0.0
Native American	0	0.0	8	10.3	2	3.7	1	5.6	11	6.5
Hispanic	0	0.0	4	5.1	6	11.1	0	0.0	10	6.0
Mixed race	14	77.8	34	43.6	14	25.9	3	16.7	65	38.7
Age at Diagnosis (years) (N	N, valid ^o	%)								
0 - 3.9	4	22.2	30	38.5	17	31.5	2	11.1	53	31.5
4 - 8.9	12	66.7	28	35.9	22	40.7	9	50.0	71	42.3
9 - 19.7	2	11.1	20	25.6	15	27.8	7	38.9	44	26.2
Sibling pairs raised togeth	er (N pa	irs; valid %	⁄0)							
yes	9	100.0	39	100.0	26	96.3	9	100.0	83	98.8
Prenatal Alcohol Exposure	e: 4-Digi	t Alcohol I	Rank (N, va	alid %)						
Rank 3a: Exposure confirmed, amount unknown	8	44.5	30	38.5	26	48.1	6	33.3	70	41.7
Rank 3b: Exposure confirmed, amount low to moderate	4	22.2	0	0.0	4	7.3	0	0.0	8	4.8
Rank 4: Exposure confirmed and level high	6	33.3	48	61.5	24	44.4	12	66.7	90	53.5

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Rank 1: no risk	0	0.0	0	0.0	0	0.0	0	0.0	0	0.0		
Rank 2: unknown risk	2	11.1	8	10.3	0	0.0	2	11.1	12	7.1		
Rank 3: moderate risk	16	88.9	70	89.7	54	100.0	15	83.3	155	92.3		
Rank 4: high risk	0	0.0	0	0.0	0	0.0	1	5.6	1	0.6		
Postnatal Risks: 4-Digit Rank (N, valid %)												
Rank 1: no risk	0	0.0	2	2.6	1	1.9	0	0.0	3	1.8		
Rank 2: unknown risk	2	11.1	0	0.0	5	9.3	1	5.6	8	4.8		
Rank 3: moderate risk	6	33.3	37	47.4	31	57.4	4	22.2	78	46.4		
Rank 4: high risk	10	55.6	39	50.0	17	31.5	13	72.2	79	47.0		
FASD Diagnoses (N, valid	%)											
FAS	4	22.2	1	1.3	2	3.7	1	5.6	8	4.8		
PFAS	0	0.0	5	6.4	3	5.6	2	11.1	10	6.0		
SE/AE	8	44.5	15	19.2	12	22.2	5	27.8	41	24.4		
ND/AE	4	22.2	48	61.5	23	42.6	9	50.0	84	50.0		
Sentinel Physical Findings/AE	0	0.0	0	0.0	4	7.4	1	5.6	4	2.4		
Not FASD/AE	2	11.1	9	11.5	10	18.5	0	0.0	21	12.5		

Other Prenatal Risks: 4-Digit Rank (N, valid %)

The matching criteria used to select twin and sibling pairs effectively minimized pairwise discordance in PAE and other prenatal and postnatal risk factors (Table 3). By definition, PAE was 100% concordant (virtually identical) between the monozygotic and dizygotic twin pairs. The Other Prenatal Risk Rank was 100% concordant between the monozygotic, dizygotic and full-sibling pairs, and 87.5% concordant between the half-sibling pairs. The Postnatal Risk Rank was highly concordant across all 4 groups (monozygotic: 100%; dizygotic: 87.2%; full-sibling: 91.3% and half-sibling: 75.0%), but did decrease linearly as the proportion of genome shared between siblings decreased.



Table 3. Matching criteria effectively minimized pairwise discordance in PAE and other prenatal and postnatal risk factors.

	Monozyg	gotic	Dizygoti	c	Full-sibl	ings	Half-siblings	
Prenatal and Postnatal Risks	N = 18 (9	pairs)	N = 78 ((39 pairs)	N = 54 (2	27 pairs)	N = 18 (9	pairs)
	N Pairs	valid%	N Pairs	valid%	N Pairs	valid%	N Pairs	valid%
Pren	atal Alcoho	l Exposur	e (PAE): 4	-Digit Cod	le Rank			
Concordant Ranks between sibling pai	rs							
Overall	9	100.0	39	100.0	27	100.0	9	100.0
Rank 1: Confirmed absence of PAE	0	0.0	0	0.0	0	0.0	0	0.0
Rank 2: PAE unknown	0	0.0	0	0.0	0	0.0	0	0.0
Rank 3a: PAE confirmed, amount unknown	4	44.5	15	38.5	13	48.1	3	33.3
Rank 3b: PAE confirmed, amount low to moderate	2	22.2	0	0.0	2	7.3	0	0.0
Rank 4: PAE confirmed, amount high	3	33.3	24	61.5	12	48.6	6	66.7
Discordant Ranks between sibling pair	s							
Overall	0	0.0	0	0.0	0	0.0	0	0.0
	Other Pro	enatal Risl	ks: 4-Digit	Code Ran	k			
Concordant Ranks between sibling pai	rs							
Overall	9/9	100.0	39/39	100.0	27/27	100.0	8/9	88.9
Valid Overall (excluding pairs with unknown risk)	8/8	100.0	35/35	100.0	27/27	100.0	7/8	87.5
Rank 1: no risk	0	0.0	0	0.0	0	0.0	0	0.0
Rank 2: unknown risk	1	11.1	4	10.3	0	0.0	1	11.1
Rank 3: some risk	8	88.9	35	89.7	27	100.0	7	77.8
Rank 4: high risk	0	0.0	0	0.0	0	0.0	0	0.0
Discordant Ranks between sibling pair	s							
Overall	0/9	0.0	0/39	0.0	0/27	0.0	1/9	11.1
Valid Overall (excluding pairs with unknown risk)	0/9	0.0	0/39	0.0	0/27	0.0	1/8	12.5
Rank 4: high risk – Rank 3: some risk	0	0.0	0	0.0	0	0.0	1	11.1

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	Postnatal Risks: 4-Digit Code Rank									
Concordant Ranks between sibling pairs										
Overall	9/9	100.0	34/39	87.2	21/27	77.8	6/9	66.7		
Valid Overall (excluding pairs with unknown risk)	8/8	100.0	34/39	87.2	20/23	91.3	6/8	75.0		
Rank 1: no risk	0	0.0	1	2.6	0	0.0	0	0.0		
Rank 2: unknown risk	1	11.1	0	0.0	1	3.7	0	0.0		
Rank 3: some risk	3	33.3	16	41.0	13	48.1	0	0.0		
Rank 4: high risk	5	55.6	17	43.6	7	26.0	6	66.7		
Discordant Ranks between sibling pairs										
Overall	0/9	0.0	5/39	12.8	6/27	22.2	3/9	33.3		
Valid Overall (excluding pairs with unknown risk)	0/8	0.0	5/39	12.8	3/23	13.0	3/8	25.0		
Rank 3: some risk – Rank 2: unknown risk	0	0.0	0	0.0	2	7.4	1	11.1		
Rank 3: some risk – Rank 1: no risk	0	0.0	0	0.0	1	3.7	0	0.0		
Rank 3: some risk – Rank 4: high risk	0	0.0	5	12.8	2	7.4	2	22.2		
Rank 4: high risk – Rank 2: unknown risk	0	0.0	0	0.0	1	3.7	0	0.0		
Abbreviation: PAE: prenatal alcohol exposure										

Specific aims 1 and 2: Pairwise concordance/ discordance of FASD diagnostic outcomes

FASD diagnoses (FAS, PFAS, SE/AE, ND/AE, Not FASD/AE) were 100% concordant between monozygotic twin pairs, but only 56.4% concordant among dizygotic twin pairs (Table 4). This closely mirrored the proportion of the genome shared between twin pairs (monozygotic 100%; dizygotic 56%).

The prevalence of pairwise concordance in FASD diagnoses decreased significantly and linearly as the proportion of genome shared between siblings decreased across the 4 study groups (100% among 9 monozygotic twin pairs sharing 100% of their genome; 56.4% among 39 dizygotic twin pairs sharing 50% of their genome; 41.7% among 27 pairs of full-siblings sharing 50% of their genome; and 22.2% among 9 pairs of half-siblings sharing 25% of their genome) (Chi2 linear trend = 1.7, p = 0.001) (Table 4 and Figure 2). When looking at the sub components that define FASD (growth

and CNS deficiency, FAS facial phenotype, structural and/or functional abnormalities), the prevalence of pairwise discordance in the 4-Digit Rank for each of these components increased across the 4 groups as the proportion of genome shared between siblings decreased (Table 4 and Figure 2). It is interesting to note that the prevalence of pairwise discordance in the Face Rank was the only component that increased significantly and linearly as the proportion of genome shared between siblings across the four groups decreased. The Rank 4 FAS facial phenotype, as defined by the 4-Digit Code, is the only component of FASD that is confirmed to be specific to (caused only by) PAE [7]. Other prenatal and postnatal risks can impact growth and brain development, but only PAE can cause the FAS facial phenotype. Thus, even though the prevalence of pairwise discordance in the Postnatal Risk Rank significantly increased across the 4 groups (monozygotic: 0%; dizygotic: 12.8%; full-sibling: 16.7%; and half-sibling: 25.0%) (Table 3 and Figure 2), discordance in postnatal risk factors

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cannot influence the pairwise discordance observed in Face Ranks. It is also interesting to note that the proportion of twin and sibling pairs within each study group that have the same (concordant) Face Ranks closely matches the proportion of the genome shared between the twin and sibling pairs within each study group (Figure 2).

There was no evidence in this study population that gender influenced the severity or pairwise discordance of FASD diagnostic outcomes. The severity of the FASD diagnosis was comparable between males and females across the entire study sample (n = 168) (Chi2 = 5.4, p = 0.14).

Among the 19 dizygotic twins with discordant genders:

- The female had a more severe FASD outcome than the male in 31.6% of the pairs (6/19).
- The male had a more severe FASD outcome than the female in 26.3% of the pairs (5/19).
- The male and female had the same FASD outcome in 42.1% of the pairs (8/19).



Figure 2. Twin/Sibling Pairwise Concordance in FASD outcomes and prenatal/postnatal risks. Monozygotic twins, dizygotic twins, full siblings and half siblings share 100%, 50%, 50% and 25% of their genome respectively as depicted by the first set of bars. If fetal genetics is modifying the teratogenic impact of PAE, the pattern of pairwise concordance reflected in the bars for each FASD outcome will more closely resemble the pattern of bars for Genome Shared than the patterns of bars reflecting

pairwise concordance in Alcohol Rank, other Prenatal Risks or Postnatal Risks. The bar patterns across all FASD outcomes are far more reflective of the pattern of bars for Genome Shared than the pattern of bars for Alcohol Rank, Prenatal Risk Rank or Postnatal Risk Rank. Although the bar pattern for Postnatal Risk Rank resembles the bar pattern for Face Rank, discordance in postnatal risk factors cannot be contributing to discordance in Face Rank because only prenatal factors can impact facial morphology. Since the FAS facial phenotype, as defined by the 4-Digit Code, is so specific to (caused only by) PAE, the most compelling evidence supporting the role genetics plays in modifying the teratogenic impact of PAE is illustrated in how highly correlated the bar patterns are between Genome Shared and Face Rank and how poorly correlated the bar patterns are between Face Rank and Alcohol Rank (especially between monozygotic and dizygotic twins with virtually identical PAE).

Specific aims 1 and 2: Pairwise concordance/ discordance of FASD diagnostic outcomes

FASD diagnostic discordance: Since there was 100% diagnostic concordance between monozygotic twin pairs, this analysis focused on the 39 dizygotic twin pairs. Despite virtually identical PAE, 4 of the 39 dizygotic twin pairs had FASD diagnostic contrasts as large as PFAS vs ND/AE (Table 4). The sibling in each sibling pair with PFAS experienced severe CNS functional and/or structural abnormalities (i.e., the 3rd digit in

their 4-Digit Code was Rank 3 or 4) while their cotwin experienced low to moderate CNS dysfunction dizygotic twins (4344 *vs* 2324, 1343 *vs* 1123). The four twin pairs had virtually identical PAE, were raised together, and had reportedly comparable prenatal and postnatal experiences (e.g., prenatal tobacco exposure, prenatal exposure to illicit drugs, multiple home placements, neglect, and/or physical/ sexual abuse). In other words, their contrasts in FASD outcomes would appear to better-explained by their discordant genetic vulnerability to PAE, than their discordant environmental influences.

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 Table 4. Prevalence of FASD diagnostic concordance and discordance between twin and sibling pairs.

	Monozy	gotic	Dizygoti	c	Full-sibl	ings	Half-siblings	
Concordance and Discordance in FASD Outcomes between Twin and Sibling Pairs	N = 18 (9	9 pairs)	N = 78	(39 pairs)	N = 54 (27 pairs)		N = 18 (9 pairs)	
I win and Sidling Pairs	N Pairs	valid%	N Pairs	valid%	N Pairs	valid%	N Pairs	valid%
Pairwise FAS	E, ND/AE	, Not FASI	D/AE)					
Concordant outcomes between sibling	pairs							
Total concordant pairs	9	100.0	22	56.4	11	40.7	2	22.2
FAS-FAS	2	22.2	0	0.0	0	40.7	0	0.0
PFAS-PFAS	0	0.0	0	0.0	0	0.0	0	0.0
SE/AE-SE/AE	4	44.4	4	10.3	2	7.4	0	00
ND/AE-ND/AE	2	22.2	16	41.0	6	22.2	2	22.2
Not FASD/AE-Not FASD/AE	1	11.2	2	5.1	3	11.1	0	0.0
Discordant Ranks between sibling pair	rs							
*Total discordant pairs	0	0.0	17	43.6	16	59.3	7	77.8
FAS-PFAS	0	0.0	1	2.6	0	0.0.	0	0.0
FAS-SE/AE	0	0.0	0	0.0	0	0.0	1	11.1
# FAS-ND/AE	0	0.0	0	0.0	2	7.4	0	0.0
PFAS-SE/AE	0	0.0	0	0.0	1	3.7	1	11.1
# PFAS-ND/AE	0	0.0	4	10.3	2	7.4	1	11.1
SE/AE-ND/AE	0	0.0	7	17.5	5	18.5	4	44.4
SE/AE-Not FASD/AE	0	0.0	0	0.0	2	7.4	0	0.0
ND/AE-Not FASD/AE	0	0.0	5	12.8	2	7.4	0	0.0
Not FASD/AE-Not FASD/AE	0	0.0	0	0.0	2	7.4	0	0.0
	Pairwi	se FASD D	Diagnostic	Features				
Discordant outcomes between sibling	pairs							
Growth Ranks 1-4	***1	11.1	17	43.6	10	37.0	4	44.4
**Face Ranks 1-4: Total discordant pairs	0	0.0	10	25.6	8	29.6	5	55.6
Face Rank 1 vs 4	0	0.0	0	0.0	0	0.0	0	0.0
Face Rank 2 vs 4	0	0.0	1	2.6	1	3.7	2	22.2
Face Rank 3 vs 4	0	0.0	0	0.0	2	7.4	0	0.0

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CNS Ranks 1-4: probability of structural abnormality (none, possible, probable, definite)	***1	11.1	17	43.6	14	51.9	5	55.6
Alcohol Ranks 3-4	0	0.0	0	0.0	0	0.0	0	0.0
CNS Functional Ranks 1-3: (no, moderate, severe dysfunction)	0	0.0	17	43.6	10	37.0	5	55.6
Microcephaly (head circumference <= 3 rd percentile)	***1	11.1	2	5.1	5	18.5	3	33.3
Seizure disorder	0	0.0	4	10.3	1	3.7	1	11.1

Linear trend across 4 study groups: MH Chi²: *10.7, p 0.001; ** 5.1, p 0.02. *** One twin pair had discordant growth Ranks in their 4-Digit Codes (<u>3</u>244-<u>1</u>244). Another pair had discordant CNS structural Ranks (12<u>4</u>3-12<u>3</u>3) because only one twin presented with microcephaly. Their CNS functional Ranks, however, were both Rank 3. These contrasts in a single component of the 4-Digit Code did not result in discordant FASD diagnostic classifications. Both twin pairs had concordant diagnoses of SE/AE. #Large contrast in pairwise FASD diagnostic outcomes

Four full-sibling pairs and one half-sibling pair also experienced large contrasts in FASD diagnoses (FAS vs ND/AE: 4-Digit Codes: 3443 vs 1223 and 3343 vs 1123) and PFAS vs ND/AE: 4-Digit Codes: 2434 vs 2324; 1443 vs 1323 and 2324 vs 4344) (Table 4). And like the dizygotic twins described above, these five sibling pairs had the same Prenatal Alcohol Ranks, were raised together, and had reportedly comparable prenatal and postnatal experiences (e.g., prenatal tobacco exposure, prenatal exposure to illicit drugs, multiple home placements, neglect, and/or physical/sexual abuse). Once again, their contrasts in FASD outcomes would appear to better-explained by their discordant genetic vulnerability to PAE, than their discordant environmental influences.

FAS facial phenotype discordance: The 4-Digit Code ranks the magnitude of expression of the FAS facial phenotype on a 4-point Likert scale (Rank 1: absent; Rank 2: mild; Rank 3: moderate; Rank 4: severe) (Figure 1A). As the proportion of genome shared between siblings decreased from 100% to 50% to 50% to 25% across the four groups (9 monozygotic, 39 dizygotic, 27 full-sibling and 9 half-sibling pairs, respectively), the prevalence of pairwise discordance in the FAS Facial Rank increased from 0.0% to 25.6% to 29.6% to 55.6% (Table 4). The prevalence of concordance in Facial Rank across the four groups closely followed the proportion of genome shared between siblings across the four groups (Figure 2).

Since the Rank 4 FAS facial phenotype, as defined by the 4-Digit Code, is the only FASD physical feature confirmed to be highly specific to (caused only by) PAE [7] it is interesting to document the prevalence of pairwise discordance (if any) involving the Rank 4 FAS facial phenotype (Table 4).

Two of the nine monozygotic twin pairs presented with the Rank 4 FAS facial phenotype. Both twins in each pair presented with concordant Rank 4 faces. Three of the 39 dizygotic twin pairs presented with the Rank 4 FAS facial phenotype; two pairs presented with concordant Rank 4 facial phenotypes and one pair presented with discordant Face Ranks (Rank 4 vs Rank 2). One twin in the discordant dizygotic twin pair presented with PFAS and a Rank 4 FAS facial phenotype. All three facial features of FAS were present-short PFLs, smooth philtrum and thin upper lip (mean PFL z-score -3.5, philtrum smoothness Rank 5, upper lip thinness Rank 4). In contrast, the co-twin presented with ND/AE and a Rank 2 Facial Phenotype. Only one of the three FAS facial features was present-a thin upper lip (mean PFL z-score -1.8, philtrum smoothness Rank 3, upper lip thinness Rank 4).

The Rank 4 facial phenotype was also observed among full-siblings and half-siblings (Table 4). Three of the 27 full-sibling pairs presented with the Rank 4 facial phenotype—all three pairs were discordant (Face Ranks 2 vs 4 and Face Ranks 3 vs 4). The Rank 4 facial phenotype was also observed in two of the 9 half-sibling pairs—both pairs had discordant Face Ranks (Face Ranks 2 vs 4). Even though the Alcohol Ranks were concordant for each of these five full-sibling and half-sibling pairs, this does not ensure that the day-to-day level of PAE was identical between each sibling pair.

Specific aim 4: Heritability

Comparing pairwise discordance in FASD outcomes for monozygotic versus dizygotic twins allows an indirect estimate of the importance of genetic factors in modifying the teratogenic effects of PAE in this study sample. Percent heritability (((dizygotic discordance minus monozygotic discordance) / dizygotic discordance) *100) for different FASD outcomes in the current study were as follows:

FASD Diagnosis (FAS, PFAS, SE/AE, ND/AE, Not FASD/AE):

((0.436 - 0.00) / 0.436)*100 = 100%

FAS Facial Rank: ((0.256 - 0.0) / 0.256)*100 = 100%

Growth Rank:

((0.436 - 0.111)/0.436)*100 = 74.5%

CNS 1-4 Rank:

((0.436 - 0.111)/0.436)*100 = 74.5%

Monozygotic twin pairs in this study had virtually identical genomes and virtually identical PAE. Under those genetic-environmental conditions, their FASD diagnoses and FAS facial phenotype Ranks were identical (0% discordant). In contrast, the dizygotic twin pairs in this study had virtually identical PAE, but shared only 50% of their genomes. Under those genetic-environmental conditions, 43.6% of the twin pairs had discordant FASD diagnoses and 25.6% had discordant FAS phenotype Facial Ranks. Based on these discordance rates for the monozygotic and dizygotic twin pairs, heritability estimates for the FASD diagnosis and the FAS Facial Rank were both 100%. In other words, essentially all of the discordance observed between twin pairs for these two outcomes appears to be due to differences in their genotypes, not differences in their environmental risk factors.

Heritability estimates for the Growth Rank and the CNS 1-4 Rank were both 74.5%, signifying that the Growth and CNS Ranks were determined by a combination of genetic and environmental factors. Although the prenatal environmental factor PAE was virtually identical between the twin pairs, PAE was not the only environmental risk factor in this study that could adversely impact growth and CNS development. Other prenatal risk factors like prenatal exposure to tobacco and other illicit drugs



can impact growth and CNS development, but they, like PAE were virtually identical between the twin pairs. Thus, it is unlikely that these other prenatal risk factors explain the discordance in growth and CNS development observed between the twin pairs. On the other hand, postnatal environmental risk factors like neglect, abuse, and multiple home placements can adversely impact a child's growth and CNS development and did vary slightly between the dizygotic twin pairs-5 of the 39 dizygotic pairs had discordant Postnatal Risk Ranks (Table 3). But a much higher number of dizygotic pairs had discordant Growth Ranks (n = 17) and discordant CNS 1-4 Ranks (N = 17) (Table 4). More specifically, not all 5 dizygotic pairs with discordant Postnatal Risk Ranks had discordant Growth or CNS 1-4 Ranks. Only 2 of the 5 pairs had discordant Growth Ranks and only 3 of the 5 pairs had discordant CNS 1-4 Ranks. Stated another way, only 2 (12%) of the 17 dizygotic pairs with discordant Growth Ranks had discordant Postnatal Ranks, and only 3 (18%) of the 17 dizygotic pairs with discordant CNS1-4 Ranks had discordant Postnatal Ranks. Thus, as the heritability estimates suggest, variations in the Growth and CNS 1-4 Ranks appeared to be influenced by both genetic and environmental factors. Overall, the heritability estimates generated for the growth (74.5%), FAS face (100%) and CNS (74.5%) components of FASD are reflective of the fact that growth and CNS development are susceptible to a multitude of prenatal and postnatal environmental risk factors, whereas the FAS facial phenotype is highly specific to early prenatal exposure to alcohol.

If fetal genotype is modifying the teratogenic impact of PAE, the prevalence of pairwise concordance across FASD outcomes would more closely reflect the percent of genome shared between sibling pairs than the pairwise concordance of PAE and other prenatal and postnatal risk factors. This is illustrated graphically In Figure 2 and Table 4. Monozygotic twins, dizygotic twins, full siblings and half siblings share on average 100%, 50%, 50% and 25% of their genome respectively as depicted by the first set of bars. The pattern of pairwise concordance reflected in the bars for each FASD outcome more closely resemble the pattern of bars for Genome Shared than the patterns of bars reflecting pairwise concordance in Alcohol Rank, other Prenatal Risks, or Postnatal Risks. Since the FAS facial phenotype, as defined by the 4-Digit Code, is so highly specific to (caused only by) PAE, the most compelling evidence supporting the role

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genetics plays in modifying the teratogenic impact of PAE is illustrated in how highly correlated the bar patterns are between Genome Shared and Face Rank and how poorly correlated the bar patterns are between Face Rank and Alcohol Rank (especially between monozygotic and dizygotic twin pairs with virtually identical PAE).

Discussion

Fetal genotype modifies the teratogenic effects of PAE

The outcomes of this study provide conclusive evidence that fetal genotype can modify the teratogenic effects of PAE. When twin pairs with virtually identical PAE were genetically identical, their FASD diagnoses were identical. When twin pairs with virtually identical PAE were genetically different, their FASD diagnoses were often different (44% presented with discordant FASD diagnoses). And when their diagnoses were discordant, the magnitude of discordance was extreme 10% of the time. For example, four of the 39 pairs of dizygotic twins were born at opposite ends of the fetal alcohol spectrum (PFAS and ND/AE), despite virtually identical PAE. Finally, as the proportion of genome shared between siblings decreased from 100% to 50% to 50% to 25% across the four study groups (monozygotic, dizygotic, full-sibling and half-sibling pairs respectively), the prevalence of pairwise discordance in FASD diagnoses increased linearly from 0% to 44% to 59% to 78%.

The prevalence of pairwise concordance in FASD diagnoses observed in our 48 twin pairs (monozygotic 100%, dizygotic 56%) was comparable to the prevalence of concordance observed in 16 twin pairs (monozygotic 100%, dizygotic 64%) reported by Streissguth and DeHaene [6] back in 1993-the only other FASD twin group study published to date. The 16 twin pairs (5 monozygotic and 11 dizygotic) were born to alcohol-abusing mothers from two countries (the United States and France). The study population included 11 Caucasian and 5 Native American twin pairs ranging in age from 1.5 to 30 years. The study was conducted prior to the creation of rigorous, case-defined FASD diagnostic systems. Patients were diagnosed as FAS or Fetal Alcohol Effects (FAE) in accordance with gestalt approach to diagnosis published by Clarren and Smith in 1978 [12]. Thirty-nine percent had FAS, 19% had FAE and 42% were alcohol exposed but unaffected. The higher concordance observed in their dizygotic twin pairs will be due in part to the fact that concordance in a study using only two FASD diagnoses (FAS and FAE) will always be higher than concordance in a FASD diagnoses (FAS, PFAS, SE/AE, and ND/AE). Over the years, the outcomes of ten additional twin pairs with PAE have been published as single-case studies [13-20]. Formal diagnostic FASD evaluations were rarely conducted, but the clinical descriptions of the twin pairs were consistent with monozygotic twin pairs having more concordant outcomes than dizygotic twin pairs.

Similar to humans. evidence of genetic modification of FASD outcomes also come from animal studies. For example, a study by Debelak and Smith [21] examined 11 genetic strains of chick embryos following ethanol exposure during early neurulation and found that the strains could be classified into very sensitive, moderately sensitive, or insensitive to ethanol-induced apoptosis of cranial neural crest cells, which give rise to facial structures. Comprehensive reviews on the genetics of FASD are presented by Mead and Sarkar [4] and Eberhart and Parnell [5].

Discordance in the FAS facial phenotype

It is interesting to note the rather high prevalence of pairwise discordance (43.6%) in FASD diagnoses among dizygotic twins in this study, despite virtually identical PAE. It is clear that discordance in PAE does not explain their discordant FASD diagnoses, but are there factors other than PAE that may explain the discordance? FASD is characterized by growth deficiency, a specific cluster of minor facial anomalies, and abnormal CNS structure and/or function. Only the Rank 4 FAS facial phenotype, as defined by the 4-Digit Code, is confirmed to be specific to (caused only by) PAE [7]. The FAS facial phenotype is not caused by other prenatal and postnatal risk factors. In contrast, growth deficiency and CNS structural/ functional abnormalities can be caused by a multitude of other prenatal and postnatal risk factors. Despite efforts to minimize postnatal contrasts between the twin pairs in this study (Table 3 and Figure 1), some of the pairwise discordance in growth and CNS outcomes used to generate the FASD diagnoses is likely explained, in part, by discordant postnatal risk factors.

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Documenting the prevalence of discordance (if any) for the Rank 4 FAS facial phenotype is of particular interest because the Rank 4 FAS facial phenotype is so highly specific to PAE [7]. If identical PAE can result in discordant Rank 4 facial phenotypes between twin pairs, this would further strengthen the evidence that genes are modifying the teratogenic impact of PAE. Since the FAS facial phenotype requires PAE in a very narrow window of time (during the gastrulation period of fetal development) [22,23], only the pairwise discordance in Rank 4 facial phenotypes that would be meaningful (that could be validly interpreted) would be among monozygotic and dizygotic twin pairs-the only two groups where the timing of PAE can be confirmed to be virtually identical on a day-to-day basis between twin pairs.

Two of the nine monozygotic twin pairs presented with Rank 4 FAS facial phenotypes. Both twins in each pair presented with concordant Rank 4 faces. Three of the 39 dizygotic twin pairs presented with Rank 4 FAS facial phenotypes; two pairs presented with concordant Rank 4 faces and one pair presented with discordant Face Ranks (Rank 4 vs Rank 2). The twin pair with discordant Face Ranks presented as follows: Twin 1: PFAS, Face Rank 4, all three of the FAS facial features (mean PFL zscore -3.5, philtrum smoothness Rank 5, upper lip thinness Rank 4). Twin 2: ND/AE, Face Rank 2, only 1 of the 3 FAS facial features-a thin upper lip (mean PFL z-score -1.8, philtrum smoothness Rank 3, upper lip thinness Rank 4). The work by Das et al., [24], presented below, provides a compelling genetic explanation for why Face Ranks were always concordant among monozygotic twins, but occasionally discordant among dizygotic twins.

Das et al., [24] reported a significant geneenvironment interaction explaining variation in facial morphology associated with ethanol use in pregnancy. Genetic diversity in ethanol metabolizing enzymes occurs in the general population. Ethanol is metabolized to acetaldehyde by two enzyme systems: the microsomal ethanol oxidizing system and alcohol dehydrogenase (ADH) [25,26]. The presence of the ADH1B*3 allele has been found to be protective for offspring neurodevelopmental and growth outcome after maternal ethanol consumption in pregnancy [27]. In 2004, Das et al. [24] demonstrated that among African American women and their offspring, the presence of an ADH1B*3 allele was protective for the effects of maternal ethanol ingestion during



on infant facial pregnancy formation. The protective effect demonstrated was present with the allele present in only the mother, only the infant, or both the mother and the infant. Exposure to ethanol and absence of the ADH1B*3 allele in both the mother and infant resulted in significant reductions in three facial measurements obtained from infant facial photographs-palpebral fissure length, inner canthal distance and the distance from the bridge of the nose to the bottom of the upper lip. Based on the findings of Das et al., [24] one could speculate that discordant FAS Face Ranks could occur in dizygotic twins (as observed in our study) if the ADH1B*3 allele was absent in the mother and one twin, but present in the other twin. Based on the same line of reasoning, one would expect monozygotic twins to always present with concordant Face Ranks (as was observed in our study). Replication of the Das et al., study using a study population of monozygotic and dizygotic twins with PAE would greatly advance our understanding how the ADH1B*3 allele modifies the teratogenic impact of PAE.

Implications for public health messaging and setting FASD diagnostic exposure thresholds

Despite virtually identical PAE, 4/39 (10%) dizygotic twin pairs had FASD diagnostic contrasts as large as PFAS vs ND/AE. The four twin pairs had virtually identical PAE, were raised together, and had reportedly identical prenatal and postnatal experiences (e.g., prenatal tobacco, illicit drug exposure, home placements, physical/sexual abuse). In other words, their contrasts in FASD outcomes to better-explained by would appear their discordant genetic vulnerability to PAE, than their discordant environmental influences. These 4 twin pairs provide powerful evidence that what may be construed in public health messaging (and some FASD diagnostic guidelines [28-30] as a safe level of exposure for one fetus, may very well place another fetus at significant risk. Not only can the same level of PAE cause strikingly different outcomes in two fetuses, but PAE reportedly below the threshold of exposure required by some FASD diagnostic guidelines can result in full FAS (See Figure 2 in Astley, et al., [31]). Thus, as stated by the U.S. Centers for Disease Control and Prevention [32] "There is no guaranteed safe level of alcohol use at any time during pregnancy. Fetal alcohol spectrum disorders are completely preventable if a woman does not drink during pregnancy."

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Potential limitations

Zygosity classification: Twins were classified as monozygotic or dizygotic for this study based on clinical and social service records shared with the FASDPN clinic at the time of their FASD evaluation. It is unknown how many twin pairs had zygosity confirmed through DNA genotyping. While there remains a small chance that one or more twin pairs in this study have their zygosity misclassified, a study of 578 twin pairs conducted by the National Academy of Sciences found parent report of zygosity was confirmed accurate by DNA genotyping over 95% of the time [33]. When misclassification occurred, it was most likely to occur among monozygotic twins who were not strikingly similar in appearance and thus incorrectly classified as dizygotic. This direction of error would lead to more conservative estimates of heritability. In the current study, all monozygotic twins looked identical and all dizygotic twins were easily distinguished from one another.

Do twins share identical prenatal environments? Not necessarily. It is for that reason the prenatal environments and PAE shared between our 48 twin pairs is described throughout this study as virtually identical. How twins experience the prenatal environment depends, in part, on chorionicity, i.e., whether twins share а single chorion (monochorionic) or have separate chorions (dichorionic). Monozygotic twins can be mono- or dichorionic, whereas dizygotic twins are dichorionic [34]. The chorionicity of the 9 monozygotic twin pairs in the current study was unknown. The chorion is the outer-most fetal membrane that contains the amnion/amniotic sac. The amnion is the thin innerprotects most fetal membrane that 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) they will share a single placenta, twins with separate chorions (e.g., whereas dichorionic) develop individual placentas. Dizygotic twins are dichorionic, since they form from two eggs. Among Caucasian separately fertilized populations, total twinning rates were estimated at 15-16 per 1,000 in 2003 [35]. In Caucasian populations, monozygotic twins comprise ~26% of all twins. For Caucasian populations about 17% of all twin pairs are monozygotic-monochorionic, ~9% are monozygotic-dichorionic and ~74% are



dizygotic-dichorionic. All twins can be expected to have many kinds of in-utero differences, as placental flow in monozygotic twins such and the amount of microchimerism in dizygotic twins [36]. The greatest risk associated with monochorionic placentation is related to the structure of blood vessels. One twin typically has better placement and therefore receives more of the nutrients [34]. The placenta also functions as a barrier, allowing small molecules (e.g., gases, nutrients, waste material, antibodies) to pass between mother and child through passive transport [37,38]. Other small molecules that can impact fetal development (e.g., some maternal hormones like cortisol; bacteria; teratogens such as alcohol) can also be diffused through the placenta [37,39]. Unequal placental sharing is a major cause of fetal growth discordance in monozygotic twins [40]. It is interesting to note that one of the few occurrences of discordant outcomes between monozygotic twins in the current study was discordant growth in one twin pair (Growth Rank 3 versus Growth Rank 1). Both twins had concordant weight percentiles (ranging from the 20th to 40th percentiles) at 2 years of age. Height percentiles, birth and however, were significantly discordant. One twin was significantly shorter (1st and 10th percentiles at birth and 2 years of age) than the other (20th and 60th percentiles at birth and 2 years of age). All in all, while prenatal environments, including level of PAE, are not necessarily 100% identical between twin pairs, the near perfect match between percent of genome shared and FASD diagnostic concordance (monozygotic twins: 100% genome shared and 100% FASD concordance; dizygotic twins: 50% genome shared, 56% FASD concordance) suggests the prenatal environments and PAE levels in our 48 twin pairs were virtually identical.

Conclusion

Not all fetuses are equally vulnerable to the adverse effects of prenatal alcohol exposure. Risk is not just dependent on timing and level of exposure. Fetal genetics plays an important role. As demonstrated in this study, despite virtually identical prenatal alcohol exposures, two fetuses can experience vastly different FASD outcomes. So which fetus is genetically vulnerable? We currently have no way of knowing. Thus, to protect all fetuses, especially the most genetically vulnerable, the only safe amount to drink is none at all.

Ethical Approval

This study was approved by the University of Washington Human Subjects Division.

Author Contribution

All authors are members of the interdisciplinary FASD diagnostic team and participated in the interpretation and reporting of the study's outcomes. SJAH conducted the statistical analyses.

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