Childhood Brain Tumors, Residential Insecticide Exposure, and Pesticide Metabolism Genes

Susan Searles Nielsen, Roberta Mckean-Cowdin, Federico M. Farin, Elizabeth A. Holly, Susan Preston-Martin, and Beth A. Mueller

Public Health Sciences Division, Fred Hutchinson Cancer Research Center, Seattle, Washington, USA; Norris Comprehensive Cancer Center/Department of Preventive Medicine, Keck School of Medicine, University of Southern California, Los Angeles, California, USA; Center for Ecogenetics and Environmental Health, Department of Environmental and Occupational Health Sciences, University of Washington, Seattle, Washington, USA; Department of Epidemiology and Biostatistics, School of Medicine, University of California, San Francisco, San Francisco, California, USA; Department of Epidemiology, School of Public Health and Community Medicine, University of Washington, Seattle, Washington, USA

BACKGROUND: Insecticides that target the nervous system may play a role in the development of childhood brain tumors (CBTs). Constitutive genetic variation affects metabolism of these chemicals.

METHODS: We analyzed population-based case–control data to examine whether CBT is associated with the functional genetic polymorphisms PON1−108T, PON1Q192R, PON1L55M, BCHEA397T, FMO1C−956A, FMOEL158E, ALDH3A1313A and GSTT1 (null). DNA was obtained from newborn screening archives for 201 cases and 285 controls, ≤ 10 years of age, and born in California or Washington State between 1978 and 1990. Conception-to-diagnosis home insecticide treatment history was ascertained by interview.

RESULTS: We observed no biologically plausible main effects for any of the metabolic polymorphisms with CBT risk. However, we observed strong interactions between genotype and insecticide exposure during childhood. Among exposed children, CBT risk increased per PON1−108T allele (odds ratio OR = 1.8; 95% confidence interval CI, 1.1–3.0) and FMO1C−956A (6) allele (OR = 2.7; 95% CI, 1.2–5.9), whereas among children never exposed, CBT risk was not increased (PON1: OR = 0.7; 95% CI, 0.5–1.0, interaction p = 0.005; FMO1: OR = 1.0; 95% CI, 0.6–1.6, interaction p = 0.009). We observed a similar but statistically nonsignificant interaction between childhood exposure and BCHEA397T (interaction p = 0.08). These interactions were present among both Hispanic and non-Hispanic white children.

CONCLUSION: Based on known effects of these variants, these results suggest that exposure in childhood to organophosphorus and perhaps to carbamate insecticides in combination with a reduced ability to detoxify them may be associated with CBT. Confirmation in other studies is required.

KEY WORDS: acetylcholinesterase inhibition, childhood cancer, children, gene–environment interaction, insecticides, pesticides, xenobiotic metabolism. Environ Health Perspect 118:144–149 (2010). doi:10.1289/ehp.0901226 available via http://dx.doi.org/ [Online 5 October 2009]

Both environmental exposure and genes affect childhood brain tumor (CBT) development. Ionizing radiation to the head and selected heritable syndromes are established risk factors (Fisher et al. 2007). However, these account for only a small proportion of CBT cases. Several epidemiologic studies suggest pesticides might be associated with CBT (Infante-Rivard and Weichenthal 2007), but most relied on retrospective questionnaire data with little detail. One study that considered the type and timing of pesticide exposure observed an increased risk of CBT with prenatal exposure to flea/tick products, but not herbicides, fungicides, or mollusccides (Pogoda and Preston-Martin 1997). The specificity of this finding is interesting because the major classes of insecticides—organophosphorus (OP), carbamate, organochlorine (OC), and pyrethrin/pyrethroid—readily cross the blood–brain barrier and target the nervous system, whereas pesticides aimed at plants and fungi inherently rely on different mechanisms of action.

Because constitutive genetic variation influences insecticide metabolism, we previously examined whether CBT is associated with two single nucleotide polymorphisms (SNPs) in the gene that codes for paraoxonase (PON1) (Searles Nielsen et al. 2005). We observed no association between CBT and the single coding region SNP PON1Q192R, but a strong dose–response relationship between CBT and PON1−108T: a promoter-region SNP associated with enzyme levels. CBT risk was increased among children who presumably had a reduced ability to detoxify chlorpyrifos and diazinon, the most common residential insecticides for many years, and this association was restricted to children whose homes had been chemically treated for insects. Here we examine these SNPs and six additional genetic polymorphisms that affect insecticide metabolism using an expanded number of cases and controls born in Washington State and California.

Materials and Methods

Participant selection and specimen retrieval. This analysis includes children enrolled in the U.S. West Coast CBT Study, a population-based case–control study described previously (Preston-Martin et al. 1996). Briefly, those cases were children diagnosed from 1984 through 1991 with a primary tumor of the brain, cranial nerves, or meninges [International Classification of Diseases–Onology (ICD-O) (World Health Organization 1976) codes 191.0–192.1] and who were identified through the Surveillance, Epidemiology and End Results registries in the Seattle–Puget Sound region of Washington, the San Francisco–Oakland area of California, and Los Angeles County, California. Control children living in the same counties were identified by random digit dialing. Children with a biological mother who spoke English or Spanish in a home with a telephone were eligible. After informed consent, interviews were completed with the mothers of 75% of controls who met these criteria (88% screened) and who were invited to participate while being frequency matched to cases by age and sex (1:1 in Los Angeles and 2:1 elsewhere) and 73% of cases for whom physician permission was received (97% of all cases) and who met the above criteria (or potentially met them but who were not located, 13% of all cases).

We identified those participating children for whom a dried blood spot (DBS) from newborn screening might still be archived at the Washington State Department of Health.
Child brain tumors and insecticide metabolism

(born in Washington in 1978–1990) or the California Department of Public Health (born in California in 1982–1990). As detailed elsewhere (Searles Nielsen et al. 2008), DBS were located and anonymized for 88% of children for whom a specimen was sought in Washington, including 66 (94%) cases and 137 (86%) controls included in both the present and earlier (Searles Nielsen et al. 2005) CBT–PON1 analyses. Similar specimen collection methods were used in California, where DBS for 86% of sought children were located, including 26 (93%) cases and 50 (75%) controls from San Francisco and 110 (92%) cases and 99 (85%) controls from Los Angeles. Institutional review board approvals from all relevant agencies were obtained before the study began.

Genotyping. The Functional Genomics Core Laboratory of the Center for Ecogenetics and Environmental Health at the University of Washington in Seattle, which was unaware of case status, obtained DNA using commercially available kits (QIAamp DNA Mini Kit for Washington DBS (Searles Nielsen et al. 2005) and the Repli-g Kit (Paynter et al. 2006) for California DBS; Qiagen, Valencia, CA), determined glutathione S-transferase theta 1 (GSTT1) null status (Kelada et al. 2003), and used custom TaqMan Detection System–based assays by–Design service (Applied Biosystems, Inc., Foster City, CA) for seven functional pesticide metabolism SNPs (Table 1) and PON2S311C (rs7493). We included PON2S311C to investigate whether CBT–PON1 associations might be a result of PON1’s generic antioxidant capabilities, because PON2 does not metabolize OPs (Draganov et al. 2005). Fragment lengths required for the TaqMan assays ranged from 84 to 202 bp (base pairs) and from 215 to 480 bp for the GSTT1 assay. Negative controls (no DNA) and sequencing-verified positive controls were included in each batch of analyses. Results were verified by sequencing as needed, and approximately 10% of all samples were resequenced. For the Washington samples, blind duplicate or quadruplicate specimens for 6% of cases and 6% of controls were assayed for all nine polymorphisms [205 of 207 (99.0%) pairs agreed], and the PON1C–108T assay results were confirmed using a different TaqMan assay (100% concordance). Complete genotyping data were available for all but two cases and one control.

Table 1. Functional effect and hypothesized “high-risk” alleles in genetic pesticide metabolism polymorphisms.

| Enzyme                      | Function                                  | Expression                                                                 | Polymorphism       | Hypothesized “high-risk” allele |
|-----------------------------|-------------------------------------------|---------------------------------------------------------------------------|--------------------|---------------------------------|
| Paraoxonase (PON1)          | Hydrolyzes environmentally and CYP-      | In blood and liver, expressed during the fetal period; increases with    | PON1C–108T (rs705379), | T: Reduced PON1 levels in neonates (63% lower in those with 2 vs. 0 variants) (Chen et al. 2003) and adults (Brophy et al. 2001; Levieu and James 2000) |
|                             | activated intermediates (AChE-inhibiting | gestational age and after birth; adult levels reached at 6–25 months of | adjacent to Sp1 binding site in the promoter region |                                  |
|                             | oxons) of selected OP insecticides (e.g., | age                        | PON1G120R (rs862),    |                                  |
|                             | chlorpyrifos, diazinon, parathion) such   |                            | amino acid substitution near the catalytic center |                                  |
|                             | that they cannot inhibit AChE            |                            | PON1L358M (rs854560), |                                  |
| Butyrylcholinesterase (BuChE)| Sequesters all OP and carbamate           |                              | amino acid substitution |                                  |
|                             | insecticides through stoichiometric      |                            | BChE632T (rs1803274), | T ("K variant"): 30–40% reduction in BuChE activity (Babaoglu et al. 2004; Bartels et al. 1992; Maitzler et al. 2009) |
|                             | (1:1) binding such that they cannot inhibit |                            | amino acid substitution |                                  |
| Flavin-containing           | Metabolizes some OP insecticides (e.g.,   | FM01: highest in the embryo during extensive brain development, and     | FM01G926A (rs12720462), in a promoter region | A (*6): Eliminates Y11 binding, 2- to 3-fold loss of promoter activity (Hines et al. 2003) |
| monoxygenase (FM01 and FM03)| phorate, terbufos, fenitrothion,           | high throughout the fetal period; decreases in the liver and brain      |                                  |                                  |
|                             | fonofos) and some carbamate               | after birth; also present in the small intestine and lung (at levels    |                                  |                                  |
|                             | insecticides (e.g., aldicarb and         | greater than liver after birth)                                        |                                  |                                  |
|                             | methiocarb) oxidizes the thioether sulfur  |                                  |                                  |                                  |
|                             | to form a sulfoxide; does not form oxons  |                                  |                                  |                                  |
| Aldehyde                   | Metabolizes a product of permethrin       | FM03: Not present in the postembryonic fetal period; in brain by 1–2 years of age | FM03S158K (rs2266782), amino acid substitution | K: One-third activity (Lattard et al. 2003) |
| dehydrogenase 3A1 (ALDH3A1)| (pyrethroid insecticide)                  |                                  |                                  |                                  |
| Glutathione                 | Metabolizes OC insecticides (DDT,         | In brain, stomach, and lung                                             | ALDH3A1S154A (rs887241), amino acid substitution | A: May reduce enzyme activity (Satomichi et al. 2003) |
| S-transferase theta 1      | (GSTT1)                                    |                                  | GSTT1 null                     |                                  |
| (GSTT1)                     | (applied to methylparathion, EPN,         |                                  |                                  |                                  |
|                             | triazine herbicides (atrazine), and       | In brain, liver (including during the fetal period), lung, small intestine, and blood (erythrocytes) |                                  |                                  |
|                             | chloroacetanilide herbicides (alachlor)   |                                  |                                  |                                  |

Abbreviations: DDT, dichlorodiphenyltrichloroethane; EPN, ethyl p-nitrophenyl phenylphosphonothioate.
ratios (OR) and 95% confidence intervals (CIs) of CBT in relation to each polymorphism. The GSTT1 assay provided dichotomous (any/no GSTT1) results. For all other polymorphisms we checked Hardy–Weinberg equilibrium by exact chi-square test [butyrylcholinesterase (BCHE) and flavin-containing mono-oxygenase 1 (FMO1) SNPs, with a minor allele frequency of ≤ 0.20] or Pearson’s chi-square test (other SNPs) and, unless noted, modeled genotype linearly (coded the variable as 0, 1, or 2 hypothesized “high-risk” alleles) (Table 1). Likelihood ratio tests confirmed that single linear variables were appropriate. We adjusted all models for study center, sex, diagnosis/reference age, and race/ethnicity [African American (either parent African American), Hispanic (not African American, and either parent Hispanic), white (both parents non-Hispanic white), Asian/other]. We excluded five cases and two controls with unknown race/ethnicity.

We conducted haplotype analyses for the four PON SNPs and the two FMO SNPs. We inferred haplotypes (PHASE software, version 2.1; Stephens and Donnelly 2003) while accounting for distance between SNPs, and using 100 additional control DBS (anonymous children born in Washington in 1980–1991). We modeled haplotype linearly while including children for whom both alleles were sufficiently frequent (> 1%) and estimated with > 80% probability.

To examine the potential for gene–insecticide interaction, we estimated separate CBT-genotype ORs for a) unexposed children (never exposed during pregnancy or childhood), b) children exposed during pregnancy, and c) children exposed during childhood. We formally assessed interaction on a multiplicative scale in logistic regression; reported p-values are from a single product term (genotype multiplied by dichotomous exposure). Prenatal and childhood insecticide exposure were correlated, so, when possible, we stratified prenatal exposure models by childhood exposure, and vice versa. We attempted to confirm all observed interactions in case–only gene–environment models (Khoury and Flanders 1996), because these would not be influenced by the composition of our control group or their reporting of insecticide use.

We checked whether our main gene and gene–insecticide CBT ORs were consistent across racial/ethnic groups, age, and study center. For these comparisons we dichotomized diagnosis/reference age at the median (3 years), combined centers (California, Washington), and examined stratum-specific estimates for sufficiently large racial/ethnic groups (Hispanic, non-Hispanic white). We also considered CBT histologic subtype. Included among the present sample were ICD-O histology codes 9380, 9382, 9400, 9401, 9420, 9421 (astroglial tumors, n = 96, 48% cases), 9470, 9471, 9473 (medulloblastoma/primitive neuroectodermal tumors, n = 55, 27% cases), and 9391–9393 (ependymoma, n = 25, 12% cases).

Results
All children were ≤ 10 years of age at diagnosis/reference, and most were < 5 years of age (Table 2). Proportionally more cases than controls were Hispanic or nonwhite. Only three cases and three controls had a heritable syndrome that predisposes to brain tumor, or a first-degree relative with a history of brain tumor. Farm residence and maternal prenatal agricultural occupation also were uncommon (2–4% cases, 1–2% controls).

Residential insecticide exposure. During pregnancy, proportionally more mothers of cases (27%) than controls (21%) reported treatment of the home for termites, fleas, ants, cockroaches, silverfish, or other pests (Table 2); we did not observe this difference in Washington State, where treatment was less prevalent than in California (data not shown). In contrast, treatment of the home for insects during childhood was more common among controls (33%) than among cases (23%) (Table 2), a difference observed in all study centers (data not shown). Among children in the pesticide follow-up study in Los Angeles, any residential insecticide use was prevalent both during pregnancy and childhood (≥ 70% of cases and controls). In general, use of flea/tick products was more common among cases than among controls, and the reverse for nuisance pests.

CBT and pesticide metabolism polymorphisms. Genotype frequencies were in Hardy–Weinberg equilibrium for each racial/ethnic group (all p-values > 0.05). Overall, we observed no marked differences between cases and controls for any polymorphism [see Supplemental Material, Table 1 (available online at doi:10.1289/ehp.0901226.S1 via http://dx.doi.org)]. Any potential heterogeneity in the CBT–genotype ORs by race/ethnicity was not statistically significant (all interaction p-values > 0.24). Main effect ORs for all racial/ethnic groups combined (and

Table 2. Characteristics of children with and without brain tumors, West Coast Childhood Brain Tumor Study, children with genotyping data and born in California or Washington State in 1978–1990 [no. (%)].

| Characteristic | Cases (n = 201) | Controls (n = 285) |
|---------------|----------------|-------------------|
| Study center  |                |                   |
| Los Angeles   | 110 (55)       | 99 (36)           |
| San Francisco | 25 (12)        | 50 (18)           |
| Seattle       | 66 (33)        | 136 (48)          |
| Birth year    |                |                   |
| 1978–1984     | 99 (49)        | 141 (49)          |
| 1985–1990     | 102 (51)       | 144 (51)          |
| Age (years)   |                |                   |
| < 5           | 167 (83)       | 222 (78)          |
| 5–10          | 34 (17)        | 63 (22)           |
| Child’s race/ethnicity | | |
| White         | 105 (54)       | 192 (68)          |
| Hispanic      | 62 (32)        | 61 (22)           |
| African American | 14 (7) | 13 (5)           |
| Asian/other   | 15 (8)         | 17 (6)            |
| Male          | 121 (60)       | 168 (59)          |
| Brain tumor in first-degree relative, or personal/family history of Li-Fraumeni syndrome, neurolentromatosis, or tuberous sclerosis | | |
| Farm residence during pregnancy/childhood | 9 (4) | 5 (2) |
| Maternal agricultural occupation in pregnancy | 4 (2) | 4 (1) |
| Chemical treatment of home for insect pests | | |
| During pregnancy | 55 (27) | 60 (21) |
| During childhood, up to diagnosis/reference | 46 (23) | 94 (33) |
| Insecticides for home, yard, garden, pets, or lice | | |
| During pregnancy | 61 (30) | 40 (17) |
| Fleas or ticks | 33 (41) | 19 (28) |
| Nuisance pests | 43 (57) | 39 (60) |
| During childhood | 60 (77) | 46 (70) |
| Fleas or ticks | 33 (41) | 22 (32) |
| Nuisance pests | 41 (51) | 40 (59) |

*All study participants for whom a usable DBS was obtained from newborn screening archives in California or Washington State. • African American: either parent African American; Hispanic: either parent Hispanic and neither parent African American; white: both parents non-Hispanic white; percentages exclude five cases and two controls with non-Hispanic white mothers and for whom paternal race was unknown. • Termites, fleas, ants, cockroaches, silverfish, or other pests; percentages exclude participants for whom prenatal (one control) or childhood (one case, one control) insecticide exposure was unknown. • From 1 month before conception until birth. • Between birth and diagnosis (cases) or comparable reference date (controls). • Based on children also participating in a pesticide follow-up study in Los Angeles only (Pogoda and Preston-Martin 1991) and with respective exposure data (76–80 cases and 65–88 controls). • Ants, cockroaches, and other nuisance pests; does not include termites or fleas.
adjusted for this factor) were close to null, with the possible exception of BCHE*A597T and FMO3*E158K (Table 3). FMO haplotype analyses suggested that any increased CBT risk in relation to the FMO3*158K allele was restricted to children with two FMO1*9536C alleles, but 95% CIs were wide (data not shown).

**Genotype–insecticide interactions.** We observed statistically significant interactions between insecticide treatment of the home during childhood and two promoter region pesticide metabolism SNPs (interaction p = 0.005 for PON1*108T7, 0.009 for FMO1*9536A; Table 3). We also observed an interaction for the coding region SNP BCHE*A597T of borderline significance (interaction p = 0.08). ORs per “high-risk” (hypothesized poor detoxification) allele (PON1*108T7, FMO1*9536A, and BCHE*A597T) were greater among children whose homes had been treated during childhood than among children whose homes never had been treated. These interactions were present among Non-Hispanic white children (interaction p = 0.11 for PON1*108T7, 0.04 for FMO1*9536A, 0.04 for BCHE*A597T) and Hispanic children (interaction p = 0.13 for PON1*108T7, 0.12 for FMO1*9536A, 0.16 for BCHE*A597T; Table 4). We observed the interactions between childhood insecticide exposure and PON1*108T7 and FMO1*9536A with or without exposure during pregnancy (all interaction p = 0.01–0.06; see Supplemental Material, Table 2 [doi:10.1289/ehp.0901226.S1]). These interactions also appeared independent of nearby SNPs, because we observed the insecticide–FMO1*9536A across FMO3*9158K genotypes, and the insecticide–PON1*108T7 interaction when modeling PON1–PON2 as a haplotype. The PON1*108T7 interaction appeared to vary by age at diagnosis/refererence: Among children < 3 years of age at diagnosis/reference, the OR per PON1*108T allele was 2.4 (95% CI, 1.0–5.7) if exposed to insecticides and 0.5 (95% CI, 0.3–0.7) if unexposed (interaction p = 0.001), and among older children 1.4 (95% CI, 0.7–2.7) if exposed and 1.2 (95% CI, 0.7–2.0) if unexposed (interaction p = 0.06; data not shown). This did not appear to be a result of the correlation between age and birth year or between prenatal and childhood exposure.

When we stratified genotype ORs by home insecticide treatment during pregnancy, we observed variability between exposed and unexposed children for some of the pesticide metabolism polymorphisms (Table 3). For example, the GSTT1 null genotype was associated with a reduced risk of CBT only among the exposed children, and the “high-risk” FMO3*158K allele was associated with an increased risk of CBT only among the unexposed children. However, none of the possible interactions between genotype and prenatal insecticide exposure was statistically significant (each interaction p > 0.15). Statistically significant or borderline interactions were suggested only in modestly sized subgroups involving PON1*108T7 and FMO1*9536A (data not shown).

Among children from Los Angeles with supplementary pesticide data, we observed possible interactions between both prenatal and childhood insecticide exposure and BCHE*A597T (genotype dichotomized; interaction p = 0.05–0.06 for any residential insecticide, 0.05–0.13 for flea/tick products, and 0.06–0.16 for products for nuisance pests such as ants and cockroaches; data not shown). The “high-risk” BCHE*A597T allele was associated with increased CBT risk only among insecticide-exposed individuals.

Even within the larger sample, our ability to consider histologic tumor type was quite limited. Nonetheless, the interactions between insecticide treatment of the home during childhood and each of the three SNPs (PON1*108T7, FMO1*9536A, and BCHE*A597T) remained when we focused on our largest subgroup, astroglial tumors. However, these interactions were not strictly specific to this tumor type.

We were unable to formally confirm any interactions using case-only models because, among controls, genotype and exposure were not independent. Otherwise, these models supported all reported interactions.

**Discussion**

We attempted to build on prior studies of CBT and pesticide exposure by considering individual differences in the metabolism of insecticides that target the nervous system. We a priori designated a “high-risk” allele for each pesticide metabolism polymorphism.

### Table 3. Risk of CBT and functional pesticide metabolism polymorphisms and PON2, overall and by home insecticide treatment, West Coast Childhood Brain Tumor Study [OR (95% CI)]

| Pesticide metabolism polymorphism | Pesticide treatment of the home for insect pestsa | All children | Chemical treatment of the home for insect pestsb | Never | Ever in pregnancy | Ever in childhood |
|-----------------------------------|-----------------------------------------------|-------------|-----------------------------------------------|-------|------------------|------------------|
|                                   | (196 ca/283 co)c | (116 ca/152 co)d | (53 ca/60 co)c | (46 ca/94 co)d |
| PON1*108T7                        | 0.9 (0.7–1.2) | 0.7 (0.5–1.0) | 1.2 (0.7–2.0) | 1.8 (1.1–3.0)* |
| PON1*108R7                        | 1.0 (0.7–1.3) | 1.0 (0.7–1.4) | 0.8 (0.5–1.4) | 0.9 (0.5–1.6) |
| PON1*108M                         | 1.0 (0.8–1.3) | 1.1 (0.8–1.6) | 0.7 (0.4–1.3) | 1.0 (0.6–1.7) |
| PON1*108C                         | 0.9 (0.7–1.2) | 0.9 (0.6–1.3) | 0.6 (0.3–1.3) | 1.1 (0.6–1.9) |
| BCHE*A597T                       | 0.7 (0.5–1.0)* | 0.6 (0.4–1.0)* | 0.9 (0.4–1.9) | 1.3 (0.6–2.8) |
| FMO1*9536A                        | 1.1 (0.7–1.6) | 1.0 (0.6–1.6) | 0.9 (0.4–2.0) | 2.7 (1.2–5.9)* |
| FMO3*158K                         | 1.2 (0.9–1.7) | 1.4 (1.0–2.0) | 1.0 (0.5–1.8) | 1.1 (0.7–2.0) |
| ALDH1A1*288A                      | 1.1 (0.8–1.4) | 1.1 (0.8–1.6) | 1.6 (0.9–2.8) | 1.0 (0.6–1.7) |
| GSTT1 (null)                      | 0.8 (0.5–1.3) | 0.5 (0.3–1.0) | 0.3 (0.1–1.0) | 0.4 (0.1–1.3) |

**Abbreviations:** ca, cases; co, control.

**Based on maternal report of chemical treatment of the home for termites, fleas, ants, cockroaches, silverfish, or “other” pests by a professional, the mother, or someone else.** OR and 95% CI, for GSTT1 null versus non-null or for all other polymorphisms per “high-risk” allele [PON1*108T7, PON1*108R7, PON1*108M, BCHE*A597T, FMO1*9536A (*6), FMO3*158K, ALDH1A1*288A; see Table 1], or PON2*311C, adjusted for race/ethnicity (African American, Hispanic, non-Hispanic white, Asian/other), study center, sex, and age at diagnosis/refererence (continuous). *Excludes five cases and two controls with unknown race/ethnicity. **p = 0.005.

### Table 4. Risk of CBT and PON1*108T7, FMO1*9536A, and BCHE*A597T, by home insecticide treatment during childhood and child’s race/ethnicity, West Coast Childhood Brain Tumor Study [OR (95% CI)]

| Pesticide metabolism polymorphism |
|-----------------------------------|
| PON1*108T7 (per T allele)         |
| Non-Hispanic white (105 ca/191 co) | 1.5 (0.8–2.8) | 0.8 (0.6–1.3) |
| Hispanic (62 ca/61 co)             | 1.6 (0.5–5.7) | 0.6 (0.3–1.0) |
| FMO1*9536A (AA/AC vs. CC)         |
| Non-Hispanic white (105 ca/191 co) | 3.0 (1.1–8.3) | 0.7 (0.3–1.7) |
| Hispanic (62 ca/61 co)             | 3.3 (0.5–23.8) | 0.6 (0.3–1.3) |
| BCHE*A597T (TT/AT vs. AA)         |
| Non-Hispanic white (105 ca/191 co) | 1.2 (0.5–3.1) | 0.5 (0.3–1.0) |
| Hispanic (62 ca/61 co)             | 2.9 (0.2–34.3) | 0.5 (0.2–1.2) |

**Abbreviations:** ca, cases; co, control.

*Hispanic: neither parent African American and one or both parents Hispanic; non-Hispanic white: both parents non-Hispanic white. **Data are OR (95% CI) obtained from a single model for each racial/ethnic group containing variables for genotype, childhood exposure (any vs. none), gene–exposure product term, study center, sex, and age at diagnosis/reference (continuous). **Based on maternal report, chemical treatment of the home for termites, fleas, ants, cockroaches, silverfish or “other” pests by a professional, the mother, or someone else, between birth and diagnosis/reference. **Data are for 20 cases/23 controls exposed during childhood and 77 cases/118 controls unexposed during childhood (including 9 cases/11 controls with prenatal exposure); excludes one control without exposure information. **Data are for 12 cases/12 controls exposed during childhood and 50 cases/48 controls unexposed during childhood (including 14 cases/17 controls with prenatal exposure).
polymorphism based on the expected functional impact with respect to acetylcholinesterase (AChE) inhibition (OP and carbamate insecticides; PON1, BCHE, FMO1, FMO3, and GSTT1 polymorphisms) and ion channel stimulation (OC and pyrethroid insecticides; aldehyde dehydrogenase 3A1 [ALDH3A1] and GSTT1 polymorphisms). Although some insecticides metabolized by these enzymes are ubiquitous in the environment or diet, ORs for CBT in relation to the hypothesized “high-risk” allele for the nine polymorphisms were close to the null, or fluctuated equally above and below the null. However, we observed interactions between genotype and chemical treatment of the home for insects during childhood for three functional SNPs located on different chromosomes: PON1C–108T, FMO1C–9536A, and BCHE539T. The direction of these interactions was consistent and biologically plausible. Moreover, they were present in each of our two largest racial/ethnic groups. PON1C–108T and BCHE539T variants are respectively associated with reduced in vivo activity of PON1 (Brophy et al. 2001; Chen et al. 2003; Leviev and James 2000) and the butyrylcholinesterase enzyme (BuChE) (Babaoglu et al. 2004; Bartels et al. 1992; Maetzel et al. 2009). Both neutralize AChE inhibitors: PON1 hydrolyzes selected OPs, notably chlorpyrifos and diazinon (Furlong 2007), and BuChE sequisters all OP and carbamate insecticides (Cokrągaj 2003). In vitro studies suggest that FMO1C–9536A materially reduces promoter activity (Hines et al. 2003). Its product, flavin-containing monooxygenase 1 (FMO1), oxidizes the thioether sulfur of some OP and carbamate insecticides (Hajjar and Hodgson 1980), and for some substrates (e.g., fenithion; Furnes and Schlenk 2004) the resulting sulfoxide is a weaker AChE inhibitor than is its parent compound. FMO1 does not appear to oxidize other sulfur atoms in OP insecticides (Hajjar and Hodgson 1980) (activate the parent compound to its oxon). Thus, our results are consistent with the possibility that children with a reduced ability to metabolize OP and perhaps carbamate insecticides might be at increased risk of CBT when sufficiently exposed. The apparent specificity of the results to AChE inhibitors is interesting but in part reflects our selection of polymorphisms. Also, even if our results suggest a biological impact of the SNP’s and insecticides, it is unknown whether this is a result of AChE inhibition per se or to some other effect of AChE-inhibiting insecticides used residentially during the study period. For example, chlorpyrifos and diazinon induce neurotoxic effects in neonatal rats, even when administered at levels insufficient to inhibit AChE (Slotkin et al. 2008).

The consistency of results across the three SNPs for which we observed an interaction with childhood insecticide exposure is compelling but nevertheless could represent chance associations. Our results were based on modest numbers, and these SNPs have not been studied in independent samples of brain tumor patients, making the probability of false positives high (Wacholder et al. 2004). The interaction involving FMO1C–9536A must be interpreted especially cautiously. Whether the net effect of FMO1 would be protective may depend on the insecticide (Buronfosse et al. 1995; Furnes and Schlenk 2004; Levi and Hodgson 1988). Although children are exposed to FMO1 insecticide substrates, including disulphoton used residually outdoors, we have not identified an FMO1-metabolized insecticide registered for residential use indoors. The interaction with home insecticide exposure in childhood is also puzzling because FMO1 enzyme levels in the brain and liver drop substantially after birth (Kokouritsaki et al. 2002; Zhang and Cashman 2006). Still, FMO1 is not absent from these sites and is expressed at greater levels in the lung and small intestine (Zhang and Cashman 2006), presumably relevant to inhaled and hand-to-mouth exposure, respectively.

The presence of interactions between genotype and insecticide exposure occurring during childhood, but generally not during pregnancy, deserves further comment. During prenatal development, maternal enzymes serve as a first line of defense against exogenous exposures, and without maternal biospecimens we were unable to directly examine the effect of this. Also, perhaps fetal expression of some enzymes is too low, regardless of genotype, to alter insecticide dose sufficiently to protect the brain; here again, maternal enzymes may be important. Our data do not suggest a lack of effect of insecticide exposure during this potentially sensitive period, but rather a lack of synergism with fetal genotype.

We did not observe interactions for other PON or FMO SNPs. None were in the promoter region of their respective genes. Also, the effect of the PON1Q192R amino acid change is dependent on the substrate, and the R isoform may be protective for chlorpyrifos but not diazinon (Davies et al. 1996; Li et al. 2000; Mutch et al. 2007). FMO1 metabolizes insecticides better than FMO3 (Furnes and Schlenk 2005; Leoni et al. 2008; Usmani et al. 2004). Perhaps more important, given our results for FMO3C158K, this coding region SNP is in linkage disequilibrium with promoter region polymorphisms that confer opposing effects on FMO3 enzyme activity (Phillips and Shephard 2008).

The childhood insecticide–PON1C–108T interaction was confined to children < 3 years of age. This polymorphism has a greater effect on PON1 levels in neonates than in adults (Chen et al. 2003), and adult levels are reached before 3 years of age (Cole et al. 2003). In addition, by this age diet is the main source of chlorpyrifos (Buck et al. 2001; Clayton et al. 2003), so in older children dietary exposure to chlorpyrifos and diazinon may have overwhelmed any interaction between PON1C–108T and residential exposure.

Since the time when the children in our study may have been exposed to home insecticides, chlorpyrifos and diazinon have been phased out of residential use in the United States. Nonetheless, children remain exposed to these and other AChE inhibitors not only via the diet but also potentially via drift from use in agricultural areas, on golf courses, and for mosquito control. In the home, OP and carbamate insecticides remain, for example, in topical treatments for lice (malathion) and flea collars (tetrachlorvinphos, carbaryl, propoxur). Therefore, the present study may have had an increased ability to observe the reported interactions because of the greater residential use of AChE inhibitors, yet our results remain relevant.

Another strength of our study is the use of archived DBS, available for participants regardless of survival status. This makes it unlikely that a relationship between genotype and responsiveness to treatment could underlie the observed interactions. Other opportunities for selection bias were present, including during specimen collection (Searles Nielsen et al. 2008). Although it is therefore difficult to rule out bias in main effects, gene–environment interactions are generally unaffected by selection bias (Morigoto et al. 2003). Further, despite the potential for differential reporting of past exposures, this more likely attenuated than caused the interactions we report (Garcia-Closas et al. 1999).

To date, this is the largest study of CBT and genetic polymorphisms. Studies with more participants are needed to clarify the reported associations. Inclusion of additional polymorphisms in FMO3 and BCHE, especially those in the promoter region, would be worthwhile. These have been less studied than coding region polymorphisms in relation to cancer, but they appeared to be critical here. Objective measurement of specific insecticides in environmental or biological specimens, and detailed interview data on the timing of exposure (e.g., during spermatogenesis, by pregnancy trimester, and by childhood age) also would be important. Although our results most strongly indicated the importance of exposures during early childhood, it is likely that other periods are also important, notably prenatal development. In studies that do consider exposures before birth, it would be useful to assess parents’ genotypes and levels of selected enzymes, including PON1 and FMO1 that are relatively stable over time in adults.

148

VOLUME 111 | NUMBER 11 | January 2010 • Environmental Health Perspectives
REFERENCES

Babaoglu MO, Ocaci T, Bayar B, Kayaalp SO, Bazkurt A. 2004. Frequency and enzyme activity of the butyrylcholinesterase K-variant in a Turkish population. Eur J Clin Pharmacol 59:875–877.

Bartels CF, Jensen FS, Rockridge D, van der Spek AF, Rubinstein HM, Lubrano T, et al. 1992. DNA mutation associated with the human butyrylcholinesterase K-variant and its linkage to the atypical variant mutation and other polymorphic sites. Am J Hum Genet 50:1096–1103.

Brophy VH, Jamps LA, Clendenning JB, McKinstry LA, Jarvik GP, Furlong CE. 2001. Effects of 5′ regulatory-region polymorphisms on paraoxonase-1 (PON1) expression. Am J Hum Genet 68:1428–1436.

Buick RJ, Dzikmayak H, Xue J, Zartner VG, Hamstrom K. 2001. Modeled estimates of chlorpyrifos exposure and dose for the Minnesota and Arizona NHEXAS populations. J Expo Anal Environ Epidemiol 11:253–268.

Buronfosse T, Moroni P, Benoiti E, Riviere JL. 1995. Stereoselective sulfonation of the pesticide methiocarb by flavin-containing monooxygenase and cytochrome P450-dependent monooxygenases of rat liver microsomes. Anticholinesterase activity of the two sulfoxide enantiomers. J Biochem Toxicol 10:179–189.

Chen J, Kumar M, Chan W, Berkowitz G, Wetmur JG. 2003. Increased influence of genetic variation on PON1 activity for detoxifying organophosphorus compounds. J Biochem Mol Toxicol 21:197–205.

Cole TB, Jampsa RL, Walter BJ, Arndt TL, Richter RJ, Shih DM, McCarver DG. 2003. Genetic variability at the human FMO1 locus: significance of a basal promoter yin yang 1 element polymorphism (FMO76). J Pharmacol Exp Ther 306:1319–1328.

Infante-Rivard C, Weichenthal S. 2007. Pesticides and childhood cancer: an update of Zahm and Ward’s 1998 review. J Toxicol Environ Health B Crit Rev 10:81–99.

Kelada SN, Stapleton PL, Farin FM, Bammler TK, Eaton DL, Smith-Weller T, et al. 2002. Glutathione S-transferase M1, T1, and P1 polymorphisms and Parkinson’s disease. Neurosci Lett 357:5–8.

Khoury MJ, Flanders WD. 1996. Nontraditional epidemiologic approaches in the analysis of gene-environment interaction: case-control studies with no controls! Am J Epidemiol 144:207–213.

Koukouritaki SB, Simpson P, Yeung AE, Rettie AE, Hines RN. 2000. Stereoselective sulfoxidation of the pesticide methiocarb by flavin-containing monooxygenase and cytochrome P450-dependent monooxygenases of rat liver microsomes. Pharmacogenomics 1:329–341.

Lattard V, Zhang J, Tran Q, Furnes B, Schlenk D, Cashman JR. 2003. Two new polymorphisms of the PON1 gene in Caucasian and African-American populations: comparative genetic and functional studies. Drug Metab Dispos 31:854–860.

Levi C, Buratti FM, Testai E. 2008. The participation of human hepatic flavin-containing monooxygenases and aldehyde oxidase in the biotransformation of the insecticide fenthion. Toxicol Appl Pharmacol 233:342–352.

Levi PE, Hodgson E. 1988. Stereospecificity in the oxidation of phorate and paraoxon by purified FAD-dependent monoxygenase and cytochrome P-450 isoenzymes. Xenobiotica 18:29–39.

Leviev I, Deskin S, James RW. 2001. Decreased stability of the M54 isoform of paraoxonase as a contributory factor to variations in human serum paraoxonase concentrations. J Lipid Res 42:528–535.

Leviev I, James RW. 2000. Promoter polymorphisms of human paraoxonase PON1 gene and serum paraoxonase activities and concentrations. Arterioscler Thromb Vasc Biol 20:516–521.

Li WF, Costa LO, Richter RJ, Hagen T, Shih DM, Twedt A, et al. 2000. Catalytic efficiency determines the in vivo efficacy of PON1 for detoxifying organophosphorus compounds. Pharmacogenetics 10:767–779.

Maelzer W, Keller S, Michelsen J, Koehler N, Stransky E, Becker C, et al. 2008. No differences of butyrylcholinesterase protein activity and allele frequency in Lewy body diseases. Neurobiol Dis 35:296–301.

Morimoto LM, White E, Newcomb PA. 2003. Selection bias in the assessment of genetic-environment interactions in case-control studies. Am J Epidemiol 158:259–263.

Mutlu E, Daly AK, Williams FM. 2007. The relationship between PON1 genotype and PON1–192 genotype in detoxification of three oxons by human liver. Drug Metab Dispos 35:315–320.

Paynter RA, Skibola DR, Skibola CF, Buffler PA, Wiemels JM, Smith MT. 2006. Accuracy of multiplexed Illumina platform-based single-nucleotide polymorphism genotyping compared between genomic and whole genome amplified DNA collected from multiple sources. Cancer Epidemiol Biomarkers Prev 15:2533–2538.

Phillips JR, Shepherd EA. 2008. Flavin-containing monooxygenases: mutations, disease and drug response. Trends Pharmacol Sci 29:294–301.

Pogoda JM, Preston-Martin S. 1997. Household pesticides and risk of pediatric brain tumors. Environ Health Perspect 105:1246–1250.

Preston-Martin S, Pogoda JM, Mueller BA, Holly EA, Lijinsky W, Davis RL. 1996. Maternal consumption of cured meats and vitamins in relation to pediatric brain tumors. Cancer Epidemiol Biomarkers Prev 5:399–405.

Saitomichi A, Nakajima Y, Takeuchi A, Takagaki Y, Saijenki G, Shibuya A. 2000. Primary structure of human hepatocellular carcinoma-associated aldehyde dehydrogenase. Biochem Biophys Acta 1481:328–334.

Searels Nielsen S, Mueller BA, De Roos AJ, Checkoway H. 2008. Newborn screening archives as a specimen source for epidemiologic studies: feasibility and potential for bias. Environ Health Perspect 116:1169–1173.

Searels Nielsen S, Mueller BA, De Roos AJ, Viernes HM, Farin FM, Checkoway H. 2005. Risk of brain tumors in children and susceptibility to organophosphorus insecticides: long-term follow-up of paraoxonase (PON1). Environ Health Perspect 113:909–913.

Slatkin TA, Seidler FJ, Fumagalli F. 2008. Targeting of neutrophilic factors, their receptors, and signaling pathways in the developmental neurotoxicity of organophosphates in vivo and in vitro. Brain Res Bull 76:424–438.

Stephens M, Donnelly P. 2003. A comparison of Bayesian methods for haplotype reconstruction from population genotype data. Am J Hum Genet 73:1162–1169.

Usmani KA, Karoly ED, Hodgson E, Rose RL. 2004. In vitro sulfoxidation of theophorin compounds by human cytochrome P450 and flavin-containing monooxygenase isoforms with particular reference to the CYFZC subfamily. Drug Metab Dispos 32:333–339.

Vasholdar S, Chanock S, Garcia-Closas M, El Ghormli L, Rothman N. 2004. Assessing the probability that a positive report is false: an approach for molecular epidemiologic studies. J Natl Cancer Inst 96:434–442.

World Health Organization. 1976. International Classification of Diseases for Oncology, 1st ed. Geneva:World Health Organization.

Zhang J, Cashman JR. 2006. Quantitative analysis of FMO2 gene mRNA levels in human tissues. Drug Metab Dispos 34:19–26.