N-acetyltransferase 1 polymorphism increases cotinine levels in Caucasian children exposed to secondhand smoke: the CCAAPS birth cohort

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Cotinine is a proxy for secondhand smoke (SHS) exposure. Genetic variation along nicotine and cotinine metabolic pathways may alter the internal cotinine dose, leading to misinterpretations of exposure-health outcome associations. Caucasian children with available SHS exposure and hair cotinine data were genotyped for metabolism-related genes. SHS-exposed children had 2.4-fold higher hair cotinine (0.14 ± 0.22 ng mg⁻¹) than unexposed children (0.06 ± 0.05 ng mg⁻¹, P < 0.001). SHS-exposed children carrying the NAT1 minor allele had twofold higher hair cotinine (0.18 ± 0.12 ng mg⁻¹ for heterozygotes and 0.17 ± 0.08 ng mg⁻¹ for homozygotes) compared with major allele homozygotes (0.09 ± 0.02 ng mg⁻¹, P = 0.0009), even after adjustment for SHS dose. These findings support that NAT1 has a role in the metabolic pathway of nicotine/cotinine and/or their metabolites. The increased cotinine levels observed for those carrying the minor allele may lead to SHS exposure misclassification in studies utilizing cotinine as a biomarker. Additional studies are required to identify functional single-nucleotide polymorphism(s) (SNP(s)) in NAT1 and elucidate the biological consequences of the mutation(s).

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INTRODUCTION

Secondhand smoke (SHS) exposures have been estimated to cause between 150 000 and 300 000 lower respiratory tract infections and exacerbate 400 000–1 000 000 cases of childhood asthma annually.1,2 Exposures also cause acute and recurring middle ear disease, cough, phlegm, wheeze and breathlessness, and may be responsible for induction of new cases of asthma in children and adolescents.3 Despite these health risks, 35% of children in the United States live in homes where residents or visitors smoke on a regular basis4 and 50–75% of children have detectable levels of serum cotinine, indicating systemic exposure to SHS.1,5

Nicotine and cotinine measures serve as a proxy for systemic exposure to over 7000 chemicals present in cigarette smoke.6 Although biomarkers of SHS are routinely used to determine systemic exposures, there is great deal of interindividual variability in these measurements, even in pharmacokinetic studies utilizing labeled nicotine and cotinine where exposure levels are identical.7–10 Some of these variations are because of known factors, such as race, gender, age and genetic variation.11,12

Nicotine is mainly metabolized through hepatic oxidation, and 70 and 80% is converted to cotinine through a two-step process,11,13 mediated by a cytochrome p450 (CYP) system and catalyzed by a cytoplasmic aldehyde oxidase (AOX1).14–15 The CYP system consists of genes including CYP2A6, CYP2B6, CYP2D616–19 and CYP2A13.12 CYP2A6 is responsible for ~90% of the oxidation of nicotine to cotinine.8,20,21 Other liver enzymes, including UDP-glucuronosyltransferases, and flavin-containing monooxygenases are involved in the metabolism of nicotine and cotinine to additional metabolites.8,11,12 The genes that regulate the metabolism of nicotine are rich in polymorphisms, and many alter the conversion of nicotine to cotinine and their metabolites. Genetic variation in the CYP2A6 gene is well documented to cause decreased or absent nicotine metabolism in adults. Smokers and adults dosed with labeled nicotine that carry variant CYP2A6 alleles have decreased nicotine metabolism and urinary cotinine levels.22–24 No studies, however, have evaluated the effect of polymorphisms in these genes and other metabolism-related genes with cotinine levels in SHS-exposed children.

The objective of this study is to determine how polymorphisms in genes along the nicotine and xenobiotic metabolism pathways contribute to cotinine level variability, with respect to parental-reported SHS exposures. We genotyped Caucasian children participating in the Cincinnati Childhood Allergy and Air Pollution Study on a custom single-nucleotide polymorphism (SNP) chip for 176 tagging and functional variants in 12 genes known to be involved in the metabolism of nicotine, cotinine or other constituents present in SHS.

MATERIALS AND METHODS

Study population and SHS exposure determination

Caucasian children were participants in the Cincinnati Childhood Allergy and Air Pollution Study, a birth cohort of 762 infants born to atopic parent(s).25 The study was limited to Caucasians because of the known
differences in nicotine metabolism in Caucasians compared with African-Americans. Infants were identified by birth records and eligible parents had at least one allergy symptom and were skin prick test-positive to at least one aeroallergen. Children were examined annually at ages 1, 2, 3, 4 and 7 years of age for allergic disease development. SHS exposure histories were obtained by utilizing a panel of four questions. Briefly, parents were asked about the smoking habits of the mother, the number of smokers living in the child’s home, the number of hours per day the child is around SHS at any location and SHS exposures in the car. Children whose parents answered positively to any of these four questions at the age 2 or 4 clinical examinations were defined as being exposed to SHS, whereas those answering no to all were considered unexposed.

Hair cotinine quantification
Hair samples were obtained by cutting two samples of ~ 20 strands from the root end of the hair in the occipital region in the scalp. Hair samples from the 2 and 4 years of age visit were adjusted for weight and analyzed for cotinine level by radioimmunoassay at the Hospital for Sick Children in Toronto, Ontario, Canada; the limit of detection was 0.02 ng mg⁻¹ (see ref. 27). The average hair cotinine level from the age 2 and 4 samples was used. If a child had only one sample available from either of those ages, that value was used.

Gene and SNP selection, DNA isolation and genotyping
Genes were selected for inclusion on a custom Illumina GoldenGate Assay (http://www.illumina.com) based on function and relevance to nicotine metabolism. Genes implicated in nicotine metabolism (AOX1, CYP1A1, CYP1B1, CYP2A6, CYP2A13, CYP2B6, CYP2D6, flavin-containing monoxygenase 3, UDP-glucuronosyltransferase 2B10) as well as those having functions in xenobiotic metabolism (AHR, CYP1A1, NAT1, NAT2) were included. Tagging SNPs in the CEU and YRI populations with an R-square cutoff of 0.8 and minor allele frequency ≥ 10% which captured the common genetic variation, were selected using HapMap (http://hapmap.ncbi.nlm.nih.gov/). Functional SNPs in the CYP2A6 gene were also included regardless of minor allele frequency. Table 1 describes the number of SNPs selected in each gene as well as the gene name, function and associated disorders.

DNA was isolated from buccal cells collected via cytology brush or saliva using the Zymo Research Genomic DNA II Kit (Orange, CA, USA) or Oragene DNA Kit (DNA Genotek Inc., Ontario, Canada). Genotypes were assigned using BeadStudio’s genotyping module (BeadStudio v3.2, San Diego, CA, USA). A total of 176 SNPs were considered for analyses.

Statistical analyses
Hair cotinine values were log-transformed. SNPs were excluded (n = 59) that failed Hardy–Weinberg equilibrium in the non-allergic control group (P < 0.0001), had minor allele frequencies below 10% (except two SNPs in CYP2A6) or had missing call rates greater than 10%. Principal component analyses were performed using the 100 included ancestry-informative markers in EIGENSTRAT to account for potential population stratification; λ = 1 indicated no adjustment needed. Using PLINK, associations with hair cotinine were tested adjusting for sex using the additive linear regression model. To address multiple testing, we determined the average pairwise LD (as measured by r²) for all SNP combinations and calculated the Bonferroni correction using Simple Interactive Statistical Analyses Software (http://www.quantitativeskills.com/sisa/). Associations were therefore considered significant at 0.000067. Linkage disequilibrium between SNPs of interest was calculated using JMP Genomics 6.0 (SAS, Cary, NC, USA).

RESULTS
Demographics of subjects
Of the 431 Caucasian children genotyped on the Illumina array, 51 who did not have available hair cotinine levels and/or had > 20% of their total SNP calls missing were excluded, yielding 380 children for these analyses (Table 2). Children with parental-reported SHS exposure had 2.4-fold higher hair cotinine levels (0.139 ± 0.216 ng mg⁻¹) compared with children with no parental-reported exposures (0.058 ± 0.051 ng mg⁻¹, P < 0.0001; Table 2). The proportion of males did not differ between groups (Table 2).

Genetic associations with hair cotinine level
Genetic associations of the flavin-containing monoxygenase 3, CYP1B1, AOX1, UDP-glucuronosyltransferase 2B10, AHR, NAT1, NAT2, CYP2A6, CYP2B6, CYP2A13 and CYP2D6 SNPs were evaluated and adjusted for sex (Figure 1a). After consideration for multiple comparisons and stratification for reported SHS exposures, one tagging SNP in NAT1 (rs13253389) was significantly associated with hair cotinine level in children with parental-reported SHS exposures (P = 0.0006, Figure 1b, Table 3). Other SNPs in the NAT1 gene were associated with hair cotinine level at a nominal level but were not significant after multiple testing considerations (Table 3). Children without reported SHS exposures exhibited no significant genetic associations (Figure 1c).

The LD-plot of the NAT1 SNPs tested is shown in Figure 2. The significant tagging SNP in NAT1 is an A → G polymorphism with a minor allele frequency of 34%. Children whose parents reported SHS exposures and were heterozygous or homogygous for the minor ‘G’ allele had twofold higher hair cotinine levels (0.178 and 0.175 ng mg⁻¹, respectively) than subjects who were homozygous for the major ‘A’ allele (0.089 ng mg⁻¹, P = 0.0009; Figure 3). Common genetic variation at this locus did not alter hair cotinine levels of children with no reported SHS exposures.

To ensure that subjects carrying the AG or GG genotypes did not have higher hair cotinine levels because of higher SHS exposures, we evaluated the proportion of subjects whose parents reported SHS exposure by NAT1 rs13253389 genotype. There were no significant differences in the proportion of children with reported SHS exposures for subjects that were AA (35.6%), AG (41.8%) or GG (35.6%), P = 0.45. In addition, we evaluated the association between the NAT1 SNP and hair cotinine level in children exposed to SHS adjusted for sex and the SHS dose, defined by the average number of cigarettes reportedly smoked by all household members at ages 2 and 4. Even after adjustment, we found that the NAT1 rs13253389 SNP still independently and significantly contributed to hair cotinine level (P = 0.012, data not shown).

DISCUSSION
Our data support that the NAT1 gene has a role in the metabolic pathway of nicotine to its metabolites. Children whose parents indicate that they are exposed to SHS that carry the minor ‘G’ allele of the rs13253389 SNP had twofold higher hair cotinine levels than those who were homozygous for the major ‘A’ allele. This association remained significant after adjustment for SHS exposure level, indicating that this NAT1 SNP independently contributes to hair cotinine level. In contrast, children whose parents indicated no SHS exposures had no differences in hair cotinine levels by genotype. These results collectively suggest that NAT1, either directly or indirectly, regulates the metabolism of nicotine or cotinine to its downstream metabolites. To our knowledge, this is the first report of an association between the NAT1 gene and hair cotinine level.

NAT1 rs13253389 is an intronic SNP located 4305 base pairs downstream of the transcription start site. Whereas it is possible that this tagging SNP has a direct role in the phenotype observed, it is likely that this SNP is in linkage disequilibrium with the causal variant. The NAT1 gene is part of the family of arylamine N-acetyltransferases that are xenobiotic-conjugating enzymes. These enzymes are responsible for the acetylation of arylamines and arylhydrazines, O-acetylation of N-aryldihydroxylamines, and N, O acetyl transfer of N-hydroxamic acid. Genetic polymorphisms of both the NAT1 and NAT2 genes have been associated with slow and fast acetylator phenotypes. Whereas the metabolic pathways of nicotine and cotinine have been extensively characterized, NAT1 has not yet been implicated in nicotine or cotinine metabolism.
## Table 1. Selected genes, functions and associated disorders

| Total SNPs | SNPs after exclusions | Gene name | Full gene name | Chr. | Reported processes and function* | Reported associated disorders* |
|------------|-----------------------|-----------|----------------|------|----------------------------------|------------------------------|
| 18         | 12                    | FMO3      | Flavin-containing monooxygenase 3 | 1q24.3 | Intrinsic to ER membrane, flavin adenine dinucleotide binding, monooxygenase activity, NADP binding | Trimethylaminuria, metabolic disorder, familial adenomatous polyposis, adenoma |
| 10         | 7                     | CYP1B1    | Cytochrome P450, family 1, subfamily B, polypeptide 1 | 2p22.2 | Angiogenesis, arachidonic acid and cellular aromatic compound metabolic process, endothelial cell migration and adhesion, epoxide and omega-hydroxylation activity, redox process, response to toxins, steroid and xenobiotic metabolism, visual perception, ER membrane, aromatase, monooxygenase, oxidoreductase and electron carrier activity, heme, oxygen and iron binding | Breast cancer, congenital glaucoma tumors, cancer, glaucoma, prostate cancer, endometrial cancer, endometriosis, colorectal cancer, glaucoma open-angle |
| 46         | 37                    | AOX1      | Aldehyde oxidase 1 | 2p33.1 | Aldehyde oxidase and electron carrier activity, metal ion and NAD binding | None |
| 3          | 7                     | UGT2B10   | UDP-glucuronosyltransferase 2 family, polypeptide B10 | 4q13.2 | Lipid metabolism, ER membrane, glucuronosyltransferase activity | Breast cancer, tumors, cancer, lung cancer, shock, adenocarcinoma |
| 17         | 15                    | AHR       | Aryl hydrocarbon receptor | 7p21.1 | Apoptosis, blood vessel development, cell cycle, intracellular receptor signaling pathway, regulation of transcription from RNA polymerase II, prostate gland development, regulation of B-cell proliferation and transcription, response to stress and xenobiotics, xenobiotic metabolism, cytosolic AHR complex, DNA, transcription factor and protein binding | Lung, mammary tumor, retinoblastoma, inflammation, endometriosis |
| 19         | 10                    | NAT1      | N-acetyltransferase 1 | 8p22 | Small molecule and xenobiotic metabolism, cytosol, acetyltransferase activity | Cancer, breast cancer, bladder cancer, colorectal cancer, polyps, tumors, lung cancer, prostate cancer, pancreatic cancer, colon cancer |
| 15         | 14                    | NAT2      | N-acetyltransferase 2 | 8p22 | Small molecule and xenobiotic metabolism, cytosol, acetyltransferase activity | Bladder cancer, cancer, breast cancer, colorectal cancer, lung cancer, tumors, prostate cancer, polyps, colon cancer, endometriosis |
| 1          | 0                     | CYP1A1    | Cytochrome P450, family 1, subfamily A, polypeptide 1 | 15q24.1 | 9-cis-retinoic acid biosynthetic process, aging, amine, coumarin, insecticide, porphyrin and flavenoid metabolism, camera-type eye development, cell proliferation, dibenzop-dioxin catabolism, digestive tract development, embryo development, hepatocyte differentiation, hydrogen peroxide biosynthesis, parturition, antibiotic, food, herbicide, hyperoxia liposaccharide, nematode, organic cyclic compound, virus, vitamin A, wounding and arsenic response, demethylase, electron carrier, flavonoid 3'-monooxygenase and oxidoreductase activity, enzyme, heme and iron ion binding | Lung cancer, breast cancer, cancer, tumors, prostate cancer, ahh inducibility, squamous cell carcinoma, oral cancer, colorectal cancer, breast carcinoma |
| 7          | 2                     | CYP2A6    | Cytochrome P450, family 2, subfamily A, polypeptide 6 | 19q13.2 | Coumarin metabolism and catabolism, exogenous drug, steroid, xenobiotic and small molecule metabolism, cytoplasm microtubule, ER membrane, coumarin 7-hydroxylation, oxidoreductase and electron carrier activity, enzyme, heme and iron ion binding | Lung cancer, cancer, tumors, adenocarcinoma, smoking habit, liver diseases, colorectal cancer, adenoma, renal disease, carcinoma |
| 30         | 13                    | CYP2B6    | Cytochrome P450, family 2, subfamily B, polypeptide 6 | 19q13.2 | Electron carrier, monooxygenase and oxidoreductase activity, heme and iron binding, incorporation or reduction of molecular oxygen | Tumors, prostate cancer, cancer, inflammation, breast cancer, immunodeficiency, hypertrophy, virus infection, metastasis, necrosis |
| 5          | 3                     | CYP2A13   | Cytochrome P450, family 2, subfamily A, polypeptide 13 | 19q13.2 | Xenobiotic and small molecule metabolism, ER membrane, aromatase and electron carrier activity, heme and iron binding | Cancer, lung cancer |
| 1          | 1                     | CYP2D6    | Cytochrome P450, family 2, subfamily D, polypeptide 6 | 22q13.2 | Electron carrier and oxidoreductase activity, heme and iron binding, incorporation or reduction of molecular oxygen | Breast cancer, Parkinson disease, schizophrenia, lung cancer, autoimmune hepatitis, cancer, hepatitis c, Alzheimer's disease, bladder cancer, tumors |

Abbreviations: AOX1, aldehyde oxidase 1; ER, endoplasmic reticulum; FMO, flavin-containing monooxygenase. *Obtained from Gene Ontology website (www.geneontology.org). *Reported disease associations were obtained from the top 10 Novoseek disease relationship hits (number of articles in which both the gene's symbol or description and the disease appear) from GeneCards (www.genecards.org).
Nicotine is metabolized to six primary metabolites, and cotinine is quantitatively the most important. Cotinine is also further broken down to six primary metabolites. The majority of variations in the clearance of nicotine via the cotinine pathway is genetically influenced (heritability estimate 60.8%). After taking into account the effect of variation in CYP2A6, the heritability estimates are reduced but remain substantial at 51.8%. Thus, a large proportion of the genetic variance in nicotine metabolism remains, suggesting the involvement of additional genetic factors that are yet to be identified. There are many potential mechanisms by which NAT1 could, in part, modulate cotinine levels (Figure 4). NAT1 might promote the breakdown of nicotine to an as yet unidentified metabolite and the SNP (or a causal SNP in linkage disequilibrium with this SNP) might reduce the breakdown of nicotine, resulting in a compensatory increase in cotinine formation (Figure 4b (1)). Alternatively, NAT1 may regulate cotinine metabolism with the SNP preventing cotinine formation.

| Variable               | Parental report of SHS | No parental report of SHS |
|------------------------|------------------------|---------------------------|
| Total children (n)     | 160                    | 271                       |
| Exclusions¹ (n)        | 14                     | 37                        |
| Children after exclusions (n) | 146                   | 234                       |
| Percent male           | 54.8%                  | 54.3%                     |
| Mean (s.d.) ng mg⁻¹ hair cotinine² | 0.139 (0.216)   | 0.058 (0.051)             |

Abbreviations: CCAAPS, Cincinnati Childhood Allergy and Air Pollution Study; SHS, secondhand smoke; SNP, single-nucleotide polymorphism; UGT, UDP-glucuronosyltransferase. ¹Individuals without an available hair cotinine level or more than 20% of their total SNPs missing were excluded. ²Hair cotinine was significantly higher in children whose parent(s) reported SHS exposure (P < 0.0001).

Figure 1. Association of NAT1 SNPs with early hair cotinine level in CCAAPS Caucasian children. (a) All children; (b) SHS-exposed children; (c) SHS-unexposed children.
breakdown resulting in cotinine accumulation (Figure 4b (2)). As nicotine metabolism is also affected by the intake of certain foods such as menthol and grapefruit juice, $^{11}$ NAT1 could have a role in regulating the breakdown of one of these foods and thereby impact cotinine levels. Finally, NAT1 might regulate pathways upstream of the CYP system or other known enzymes that metabolize nicotine, indirectly affecting cotinine levels.

Cotinine is regarded as the biologic gold standard of SHS assessment. $^{5}$ Clinically, these observations are important because they highlight the NAT1 gene as a source of variability in cotinine measurement among Caucasian children. In our study, Caucasian children exposed to SHS who carry the NAT1 minor ‘G’ allele had increased cotinine levels that were not because of increased exposures, potentially leading to exposure misclassification. This is important because exposure misclassification can produce spurious differences between groups or mask true associations, and decrease power to detect true effects. $^{37}$ As biomarker variability leading to exposure misclassification has a considerable impact on all smoke exposure-related studies, sources of variability and limitations in biomarker measurement, including genetics, needs consideration.

These findings may also have an impact on the study of nicotine dependence as it is possible that the increased cotinine levels are because of an increased metabolism of nicotine. People who metabolize nicotine more quickly might smoke more cigarettes per day compared with slower metabolizers. $^{11}$ Genetic variations in other genes, such as CYP2A6, have also been reported to cause increased or decreased cotinine levels $^{22,24}$ and associate with smoking behavior. $^{18,35}$ However, the allele frequencies of these polymorphisms are low (0–8%), with a combined frequency

### Table 3. Beta of genetic associations of SNPs in the NAT1 gene with hair cotinine level in Caucasian CCAAPS children

| SNP          | Major/minor alleles | MAF  | All subjects (n = 380) | SHS exposed (n = 146) | SHS unexposed (n = 234) |
|--------------|---------------------|------|------------------------|-----------------------|--------------------------|
|              | Beta                | P-value | Beta                | P-value                  | Beta                   | P-value                      |
| rs7017402    | G/A                 | 0.123 | 0.172                 | 0.089                  | 0.466                   | 0.0052                      |
| rs4921580    | C/G                 | 0.126 | 0.204                 | 0.046                  | 0.492                   | 0.0056                      |
| rs4921581    | A/G                 | 0.333 | 0.147                 | 0.034                  | 0.403                   | 0.0010                      |
| rs13253389   | A/G                 | 0.344 | 0.132                 | 0.058                  | 0.419                   | 0.0006                      |
| rs17693103   | G/T                 | 0.179 | 0.059                 | 0.512                  | 0.325                   | 0.0417                      |
| rs6586714    | G/A                 | 0.086 | 0.054                 | 0.659                  | 0.152                   | 0.5067                      |
| rs4921880    | A/T                 | 0.227 | 0.128                 | 0.109                  | 0.366                   | 0.0086                      |
| rs11777998   | G/C                 | 0.123 | 0.286                 | 0.006                  | 0.547                   | 0.0008                      |
| rs7003890    | T/C                 | 0.442 | 0.120                 | 0.085                  | 0.123                   | 0.3240                      |
| rs8190845    | G/A                 | 0.126 | −0.081                | 0.431                  | −0.115                  | 0.5728                      |

Abbreviations: CCAAPS, Cincinnati Childhood Allergy and Air Pollution Study; MAF, minor allele frequency; SHS, secondhand smoke; SNP, single-nucleotide polymorphism. Bold indicates significance at the LD-adjusted Bonferroni level.

Figure 2. LD Plot of SNPs in the NAT1 Gene.

Figure 3. Association of NAT1 SNPs with early hair cotinine level in CCAAPS Caucasian children.

uptstream of the CYP system or other known enzymes that metabolize nicotine, indirectly affecting cotinine levels.

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of 9.1% in whites\(^{23}\); therefore, it is not surprising that we did not find any associations with the functional CYP2A6 SNPs in our population, given we were not powered to detect associations with minor allele frequencies < 10%.

As the objective of this study was to examine how genetic variability in genes involved in nicotine and xenobiotic metabolism affects children’s hair cotinine levels, we utilized parental report of SHS to stratify our exposure groups. Thus, it is possible that the carriers of the NAT1 minor allele(s) had greater SHS exposures because of parental underreporting, which contributed to the observed increased cotinine levels. To minimize this likelihood, parental-reported SHS exposures were defined by a four-question panel designed to provide the parent(s) an opportunity to report exposures without divulging personal habits.\(^{26}\) Consequently, there were no significant differences in the proportion of children with parental-reported SHS exposures by genotype. In addition, NAT1 rs13253389 remained significant in the multivariate model even after adjustment for the total number of cigarettes smoked per day by all smokers living in the child’s home, indicating that this SNP independently contributes to children’s hair cotinine level.

In conclusion, our data support that NAT1 has a role in the metabolic pathways from nicotine to cotinine and/or their associated metabolites, leading to increased cotinine levels. These findings are clinically relevant because genetic variability in NAT1 can affect cotinine biomarker levels independent of SHS exposure levels. Additional studies are required to identify functional SNP(s) in NAT1 and elucidate the biological consequences of the mutation(s).

CONFLICT OF INTEREST
The authors declare no conflict of interest.

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