Stability of the Nicotine Metabolite Ratio in *ad Libitum* and Reducing Smokers

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Abstract

Background: The ratio of two nicotine metabolites, cotinine and trans-3'-hydroxycotinine (3-HC), has been validated as a method of phenotyping the activity of the liver enzyme cytochrome P450 (CYP) 2A6 and, thus, the rate of nicotine metabolism. Our objective was to evaluate the correlates and stability of the 3-HC to cotinine ratio in *ad libitum* and reducing smokers, using nicotine replacement therapy (NRT), over a period of months.

Methods: Smokers (*n* = 123, 94% Caucasian) participated in a smoking reduction study, where one-third of the sample smoked *ad libitum* for 8 weeks (Waitlist phase), before joining the rest of the participants for 12 weeks of cigarette reduction (Reduction phase) using NRT. Urinary nicotine, cotinine, and 3-HC were measured at each visit.

Results: The baseline 3-HC to cotinine ratio was significantly but weakly correlated with cigarettes per day (*r* = 0.19), BMI (*r* = −0.27), and waking at night to smoke (*r* = 0.23). As assessed by repeated measure ANOVA, the 3-HC to cotinine ratio was stable in the Waitlist phase [coefficient of variation for 3 to 4 measurements, 38% (range, 5-110%)], whereas minor variation was noted in the Reduction phase [coefficient of variation for 3-5 measurements, 35% (range, 10-107%)].

Conclusions: In nonreducing *ad libitum* smokers, the 3-HC to cotinine ratio was generally stable, whereas during smoking reduction using NRT, some small variation was detected. Although the current findings are suggestive of the stability of the 3-HC to cotinine ratio in a predominantly Caucasian sample smoking freely or reducing smoking with NRT, additional research is needed in more diverse populations. (Cancer Epidemiol Biomarkers Prev 2008;17(6):1396–400)

Individual differences in nicotine metabolism have been shown to influence the developmental spectrum of nicotine dependence from acquisition to maintenance to cessation (1). Nicotine is primarily metabolized by the liver enzyme cytochrome P450 (CYP) 2A6 to cotinine. Accordingly, one approach to estimating the rate of nicotine metabolism has been to genotype CYP2A6. Although genotypic data have been informative in explaining individual differences in smoking behavior, such data may be an imperfect index of enzyme activity, due to the presence of exogenous (e.g., menthol) or endogenous (e.g., female sex hormones) chemicals that can induce or inhibit CYP2A6 (2, 3). Thus, phenotypic assessment of CYP2A6 activity and nicotine clearance is also important.

Benowitz et al. (4) suggested measuring the ratio of two nicotine metabolites, cotinine and trans-3'-hydroxycotinine (3-HC), as a measure of CYP2A6 activity. This ratio has been used to phenotype CYP2A6 activity because CYP2A6 is the primary enzyme mediating nicotine metabolism, and CYP2A6 also catalyzes the conversion of cotinine to 3-HC. Hence, the 3-HC to cotinine ratio is a surrogate measure of CYP2A6 activity and nicotine clearance. A growing body of research has validated the 3-HC to cotinine ratio, showing it to be moderately correlated with both CYP2A6 genotype and the rate of nicotine metabolism (1, 3, 5-9).

Although the 3-HC to cotinine ratio seems to correlate well with independent measures of nicotine clearance, limited data on its stability over time within individuals are available. Lea et al. (10) examined the stability of the 3-HC to cotinine ratio in 6 smokers, sampling morning and evening over a 7-day period. The authors did not observe diurnal differences nor substantial variation over the sample week. However, the stability of the 3-HC to cotinine ratio has not been evaluated over a period of weeks or months, or during changes in nicotine intake. The purpose of this report was to evaluate the stability of the 3-HC to cotinine ratio over a period of 5 months in a well-defined population evaluated in the context of a within-subject study of nicotine replacement therapy (NRT)-facilitated smoking reduction (11, 12). First, we examined convergent validity of the 3-HC to cotinine ratio in our sample by assessing correlations between the ratio and baseline demographic, smoking, and smoking history variables. Second, we assessed whether the 3-HC to cotinine ratio was stable during extended *ad libitum* smoking and smoking reduction with NRT.
Materials and Methods

Subjects. The current analytic sample was drawn from a larger study of scheduled smoking reduction using NRT (11, 12). Before beginning this study, all subjects provided written, informed consent approved by the University of Minnesota Institutional Review Board. The sample was composed of a total of 123 subjects (52% female; 94.3% Caucasian); the mean age was 45.6 y (SD, 10.4; range, 20.0-68.0). The mean body mass index (BMI) was 27.0 kg/m² (SD, 6.1). Among females (n = 64), 7.8% reported using hormone birth control (e.g., orthotrycyclin). Baseline smoking characteristics were as follows: mean smoking rate, 26.4 cigarettes per day (SD, 7.2; range, 15.0-50.0); and years of cigarette use, 16.6 (SD, 12.1; range, 1.0-50.0). Most smoked regular or medium cigarettes (92.5%) that were not mentholated (92.5%). In terms of nicotine dependence, the average Fagerstrom Test of Nicotine Dependence (13) score was 5.8 (SD, 1.5; range, 2.0-9.0), with 45.5% of subjects smoking their first cigarette within 5 min of awakening. In past quit attempts, the mean number of Diagnostic and Statistical Manual of Mental Disorders 4th edition nicotine withdrawal symptoms experienced in a previous quit attempt was 3.9 (SD, 1.9). The mean longest quit attempt was 348 d (range, 1-4,015).

Procedures. The study consisted of two consecutive phases: (a) Waitlist ad libitum smoking (8 wk); and (b) Reduction with NRT (12 wk). One-third of the subjects completed the 8-wk, Waitlist ad libitum smoking phase before joining the remaining sample in a 12-wk Reduction phase that consisted of 6 wk of scheduled reduction followed by 6 wk of cigarette reduction maintenance. In the scheduled reduction phase, smokers were expected to reduce their daily cigarette use in 3, consecutive 2-wk stages (as a percentage of baseline smoking, cigarettes per day): (a) weeks 1 to 2, 25% reduction; (b) weeks 3 to 4, 50% reduction; (c) weeks 5 to 6, 75% reduction; and (d) weeks 7 to 12, maintain 75% reduction. Evidence of reduction was seen in decreases in a number of tobacco-related biomarkers, including 4-(methylNtritosamo)-1-(3-pyridyl)-1-butanol (11). To assist cigarette-smoking reduction, participants were provided brief behavioral treatment and NRT (i.e., 4 mg Nicorette gum as well as 14 and 21 mg Nicoderm CQ patches). After scheduled reduction, subjects attempted to maintain their reduction levels or to further decrease cigarette consumption for an additional 6 wk while continuing NRT use.

Biochemical and Subjective Measurements. At each clinic visit, urine samples were collected for assessment of nicotine, cotinine, and 3-HC levels. The large majority of samples were collected in the morning hours (median, 10:15 a.m.); a recent report suggests that sampling time during the day does not influence the 3-HC to cotinine ratio (10). The urinary levels of cotinine, nicotine, and 3-HC were determined by gas chromatography/mass spectrometry analysis as previously described (12). The reported drug or metabolite concentrations represent the totals of free and glucuronidated forms (c.f., 4). Urine concentrations were normalized to urinary creatinine concentrations (i.e., nanomol compound per milligram creatinine). Carbon monoxide was measured in expired breath samples at each clinic visit using a Bedfont Micro Smokerlyzer (Bedfont). The daily cigarette smoking rate was determined by averaging self-recorded smoking on daily diary cards (which captured the date and time each cigarette was smoked). Self-reported demographic characteristics and smoking history variables were collected at baseline by questionnaires (13).

Data Analysis. All analyses were conducted with the Statistical Analysis System Version 9.1.3 (14). Available sample sizes differed because of attrition and occasional missing data (the maximum and minimum sample sizes were 123 at baseline and 47 in the Waitlist phase). P values of <0.05 were considered statistically significant, based on two-tailed tests, unless otherwise specified. Spearman rank-order correlations were used to assess univariate associations. Correlates of the 3-HC to cotinine ratio were subsequently evaluated in a full-rank multiple linear regression model, using log-normalized ratios as the dependent variable. Stability of the 3-HC to cotinine ratio was examined in two ways. First, the coefficient of variation or CV [(SD/mean) × 100] was computed to indicate the relative percentage of variability around the mean (c.f., 10). Second, we used repeated measure ANOVA models to assess for changes in log-normalized 3-HC to cotinine ratios over time. Type I error rate was controlled using a Tukey adjustment.

Results

Baseline Biochemical Levels. Median baseline biochemical levels were as follows: expired air carbon monoxide, 20 parts-per-million (range, 5-55); total urine nicotine, 8.7 nmol/mg creatinine (range, 1.1-46.8); total urine cotinine level, 24.6 nmol/mg creatinine (range, 4.9-90.4); and total trans-3’-hydroxycotinine, 39.8 nmol/mg creatinine (range, 1.8-167). The distribution of the 3-HC to cotinine ratio followed a log-normal distribution with a median of 2.0 (range, 0.1-11.3).

Correlations Between Baseline Sample Characteristics and 3-HC to Cotinine Ratio. Spearman rank-order correlations were computed between demographic, metabolic, and nicotine dependence variables and the 3-HC to cotinine ratio (measured at baseline, before the Waitlist or Reduction phases). Available sample sizes ranged from 123 to 64 (for the birth control analysis restricted to females). The 3-HC to cotinine ratio was positively associated with daily cigarette smoking rate (r = 0.19; P < 0.05). Higher 3-HC to cotinine ratios were associated with lower BMI (r = −0.27; P < 0.01). Those reporting waking at night to smoke tended to have higher 3-HC to cotinine ratios (r = 0.23; P < 0.05). The association between 3-HC and 3-HC to cotinine was strong (r = 0.68, P < 0.0001).

We further evaluated the average 3-HC to cotinine ratio during weeks 2, 6, and 8 of the Waitlist phase with the 3 baseline variables that were associated with the baseline 3-HC to cotinine ratio. We found that in this reduced sample (n = 47), the correlations were similar in direction, although the magnitudes were somewhat larger for daily cigarette smoking rate (r = 0.38; P < 0.01) and waking at night to smoke (r = −0.49; P < 0.001) but not BMI (r = −0.21; not significant).

Based on the foregoing univariate analyses, we modeled log-normalized 3-HC to cotinine as a function of the predictor’s cigarettes per day, BMI, and waking to
smoke, with age and sex included as covariates. All three predictors remained significant: BMI, $\beta = -0.03$, $t = -2.27$, $P = 0.03$; cigarettes per day, $\beta = 0.02$, $t = 2.07$, $P = 0.04$; and waking to smoke, $\beta = 0.36$, $t = 2.14$, $P = 0.03$. However, only 9% of the total variability in the 3-HC to cotinine ratios was accounted for by the model.

**Stability of Nicotine Metabolism.** The stability of nicotine metabolism was first examined through calculating the coefficient of variation for each subject, during the Waitlist and Reduction phases. In the 8-week Waitlist phase (3–4 observations), subjects showed an average within-subject variation as follows: total cotinine, 32% (range, 2–103%); 3-HC, 35% (range, 5–72%); and the 3-HC ratio, 38% (range, 5–110%). In the 12-week Reduction phase (3–5 observations), subjects showed an average within-subject variation as follows: total cotinine, 39% (range, 8–148%); 3-HC, 40% (range, 1–180%); and the 3-HC to cotinine ratio, 35% (range, 10–108%).

To formally assess the stability of the 3-HC to cotinine ratio, we evaluated repeated measures within subject models for the Reduction and Waitlist phases individually, using log-normalized 3-HC to cotinine ratios. From each model, we back-transformed estimated least square means and associated 95% confidence intervals to estimated geometric means (see Fig. 1). In the Waitlist phase, the ratio remained stable (time: $F = 0.73$; $P = 0.53$; see Fig. 1A). In the Reduction phase, slight variation was observed (time: $F = 2.94$; $P = 0.02$). This effect was primarily driven by a small increase in the 3-HC to cotinine ratio, with significant increases observed at week 4.

**Figure 1.** Temporal stability of the 3-HC to cotinine ratio in those in the Reduction phase (A) and Waitlist phase (B). Values are geometric means and 95% confidence intervals, back transformed from log 3-HC to cotinine. *, week 4 significantly greater than week 1.
the metabolic ratio from week 1 to 4 \((t = 2.93; P = 0.029; \text{see Fig. 1B})\). No other significant differences between time points in the Reduction phase were detected.

**Discussion**

In our univariate and multivariate analyses, cigarettes per day, waking to smoke at night, and BMI were associated with the 3-HC to cotinine ratio. The association between faster nicotine metabolism and greater daily smoking has been noted before (4). This relationship may arise because smokers seek to maintain satisfactory blood concentrations, and therefore, increased nicotine clearance leads to increased smoking. Concerning the association between faster nicotine metabolism and rising at night to smoke, Reider et al. (15) have described a clinical phenomenon, “nocturnal sleep-disturbing nicotine craving”, which involves the smoker waking with a need to smoke a cigarette to fall back to sleep. From sleep onset to awakening, plasma nicotine levels decrease as a function of nicotine metabolism, although nicotine clearance is somewhat lower at night (16). Our present observation of faster nicotine metabolism in those rising to smoke is consistent with nocturnal sleep-disturbing nicotine craving. In the case of BMI, the 3-HC to cotinine ratio tended to decline with increased body mass, indicating slower nicotine metabolism. Relatively little work has been published on the effects of obesity on nicotine pharmacokinetics (17), and further pharmacokinetic studies are needed. These results lend some evidence of convergent validity the 3-HC to cotinine ratio.

Given the increasing importance of characterizing nicotine metabolism in smokers for research as well as potentially for treatment, a rapid measure of CYP2A6 activity will be highly valuable. The 3-HC to cotinine ratio has shown good concordance with other measures of nicotine clearance, including the CYP2A6 genotype and controlled laboratory tests of nicotine clearance. Levi and coworkers (7, 8) have shown that the predictive relationship of the 3-HC to cotinine ratio to nicotine clearance remains consistent under different patterns and quantity of smoking, permitting spot sampling. An important question concerns the long-term stability of the 3-HC to cotinine ratio. Following smokers for 7 days, Lea et al. (10) observed stability in the 3-HC to cotinine ratio within time of day and across the week (CV, 26%). In our sample, we extended the work of Lea and colleagues (10) to evaluate the stability of the 3-HC to cotinine ratio in both ad libitum and NRT-assisted reducing smokers over a period of months. We observed similar CVs in both the ad libitum Waitlist smokers (CV, 38%) and in those undergoing NRT-assisted smoking reduction (CV, 35%), however with some range of variation. We found that the variation in cotinine and 3-HC were similar to the 3-HC to cotinine ratio. Some evidence of a small increase in the 3-HC to cotinine ratio in NRT-assisted reducing smokers in the first month of reduction, indicates higher nicotine clearance. However, factors related to this transient increased nicotine clearance are unknown. Overall, our findings further support the use of a single assessment of nicotine metabolism through the 3-HC to cotinine ratio.

One limitation of the current study is that it relied solely on urine to assess tobacco biomarkers. Although the agreement between urine, saliva, and plasma matrices is generally good (18, 19), the current report would have been strengthened by including saliva or plasma measurements. A second limitation of this study was that 94.3% of participants were Caucasian. In addition, few participants were exposed to exogenous chemicals known to induce metabolism of nicotine (i.e., 7.5% smoked mentholated cigarettes, 7.8% of females used hormone birth control). The effects of racial and ethnic differences as well as exogenous chemicals on nicotine metabolism have been noted, and evaluation of the stability of the 3-HC to cotinine ratio in more diverse samples is still needed. A third limitation was that due to concurrent use of two forms of NRT in the reduction phase, we were unable to meaningfully evaluate the role of the 3-HC to cotinine ratio in smoking reduction because participants were able to compensate for nicotine not obtained from cigarettes with medicinal nicotine. Given recent interest in reduced smoking as an intervention for smokers unwilling or unable to quit, further evaluation of the role of nicotine metabolism in smoking reduction is warranted.

In conclusion, the 3-HC to cotinine ratio seems to be stable under conditions of extended ad libitum smoking in a mostly Caucasian sample. Some modest variation in the 3-HC to cotinine ratio was seen during NRT-facilitated reduction. Although the evidence of the temporal stability of the 3-HC to cotinine ratio is accumulating, additional research in more racially diverse samples is needed to strongly recommend spot sampling of cotinine and 3-HC as an approach to quickly achieve a useful estimate of nicotine clearance.

**Disclosure of Potential Conflicts of Interest**

No potential conflicts of interest were disclosed.

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**References**

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