Validity of self reports in a cohort of Swedish adolescent smokers and smokeless tobacco (snus) users

A Post, H Gilljam, I Rosendahl, L Meurling, S Bremberg, M R Galanti

Objective: To validate self reports of cigarette and smokeless tobacco (snus) use in a prospective cohort of adolescents.

Design: A cross sectional analysis of a cohort sub-sample.

Setting: County of Stockholm, Sweden.

Subjects: 520 adolescents in the final grade of junior high school (mean age 15.0 years).

Main outcome measure: Concordance between self reported tobacco use and saliva cotinine concentration.

Results: Using a cut point of 5 ng/ml saliva cotinine to discriminate active tobacco use, there was a 98% concordance between self reported non-use in the past month and cotinine concentration. The sensitivity of the questionnaire compared to the saliva cotinine test, used as the gold standard, was 90% and the specificity 93%. One hundred and fifteen out of 520 subjects (22%) reported monthly tobacco use. Among these, 67% (46/69) of the exclusive cigarette smokers, 82% (23/28) of exclusive snus users, and 94% (15/16) of mixed users (cigarettes + snus) had cotinine concentrations above 5 ng/ml. Among subjects reporting daily use 95% (64/67) had salvia cotinine concentrations above the cut point. Exclusive current cigarette users were more likely to be classified discordantly by questionnaire and cotinine test compared to snus users (odds ratio 3.2, 95% confidence interval 1.2 to 8.6).

Conclusion: This study confirms the reliability of adolescents’ self reported tobacco use. In a context of low exposure to environmental tobacco smoke a cut off for saliva cotinine of 5 ng/ml reliably discriminated tobacco users from non-users. Irregular use of tobacco in this age group probably explains the discrepancy between self reported use and cotinine concentrations.

Tobacco use and exposure to ETS

Lifetime use of tobacco was investigated by the following questions: “Did you ever try smoking a cigarette, even a puff/ever try snus?” (Yes/No). “Did you smoke a whole cigarette in your life?” (Yes/No). Current tobacco use was assessed by the questions “Do you smoke/use snus at present?” with alternatives “Not at all”, “Less than once a month”, “Every month, but less than one cigarette per week”, “Each week, but less than one cigarette per day” and “At least one cigarette every day”. A reported frequency of tobacco use of...
at least once a month was considered as current use. The following question was used to estimate recentness of use: “When did you last smoke/use snus?” with seven response alternatives ranging from “today” to “more than one month ago”. Recent use (that is, use in the window of detectable cotinine concentration) was categorised as any use in the week before the survey. Regular users (that is, subjects reporting at least weekly use) were asked “How many cigarettes do you usually smoke/how frequently do you use snus during one week?” to assess intensity of use.

In order to rule out exposure to environmental tobacco smoke the following question was asked: “During the last week, approximately how many hours have you spent indoors in a place where someone smoked or had recently smoked?” The alternatives for estimated time of exposure were 0, 1–2, 3–4, 5–7, 8–14, and > 14 hours for each of the following environments: (1) In your home; (2) In someone else’s home; (3) During school hours; (4) In a café, restaurant, pub, discotheque etc; (5) In a car; (6) Elsewhere.

Collection of saliva specimens
Five months before the survey the selected adolescents and their parents received a letter explaining the purpose, but not the time, of the test. The free choice and the confidentiality of the results were underlined. Consent was assumed in all cases of no explicit refusal. The questionnaire survey was conducted at school, using the same procedure as in previous years. At the time of the survey, the students were not reminded of the coming test. Nevertheless, the samples were taken either later the same day or, with few exceptions, in the next few days. The median time between questionnaire and saliva sampling was 2.0 days (range 0–21 days).

Saliva was chosen as the medium because the non-invasive procedure is more acceptable in this age group. A team consisting of dental hygienists assisted by the school nurse collected the saliva specimens. The pupils were asked to rinse the mouth with water (in order to avoid possible contamination with snus) and to chew on a roll of cotton, which was then spat directly into a test tube, marked with an individual study code. The tubes were brought to the laboratory the next working day, centrifuged, and stored at −80°C until analysis. The school nurses took saliva samples on later occasions from students who were absent from school during the test day. No feedback was given to the study subjects or to the schools concerning the survey or the test results.

Saliva cotinine assay
To identify tobacco users among adults a cut point of 15 ng cotinine/ml saliva is customary.4 Owing to the low intensity of smoking in this age group, and the low exposure to environmental smoke in Swedish homes, a cut off at 5 ng/ml was chosen as a threshold for active tobacco use (N Benowitz, personal communication with MR Galanti).3 Values of less than 5 ng/ml were thus interpreted as no tobacco use in the preceding seven days, or low level exposure to passive smoking only. Cotinine in saliva has a half life of 16–20 hours with approximately 100 hours from intake to the chosen cut off value, and was determined by gas chromatography-mass spectrometry (GC-MS), using the method of Curvall et al3 with cotinine-d3 as an internal standard. LOD (limit of detection) was 0.2 ng/ml and LOQ (limit of quantitation) 2 ng/ml. Precision of the method was determined by running controls at three levels during three consecutive days. Calibration curves were constructed each day. Within-day precision was determined to coefficient of variation (CV) of: 4.7% at 5 ng/ml (n = 5); 4.8% at 100 ng/ml (n = 5); and 2.9% at 300 ng/ml (n = 5). Between-day precision was determined to 4.53% at 5 ng/ml (n = 15); 5.82% at 100 ng/ml (n = 15); and 4.1% at 300 ng/ml (n = 15). Calibration curves had always correlation constants better than 0.998 and varied marginally in slope between days.

Statistical methods
For the purpose of assessing reliability of self reports, we primarily conducted a descriptive analysis by comparing the expected proportion of cotinine-negative among self reported non-users (expected 100%) and of cotinine-positive among self reported current users.

We also analysed the same proportions separately by type of tobacco used. In tabular data, the departure of the observed from the expected distribution was formally tested by means of χ2 statistics. We also analysed three categorical predictors of discordance between self reported tobacco use and classification on the cotinine test, namely type of product (cigarettes only or snus, alone or in combination with cigarettes), intensity of use (no weekly use, 1–30 or more than 30 cigarettes/snus dips per week); and recentness of use (within one day, 2–7 days, or > 7 days before the survey). As measures of association odds ratios (OR) and their corresponding 95% confidence intervals (CI) were computed by ordinary logistic regression. This analysis was conducted both among all subjects and after restriction to current users. The level for the significance was conventionally set at 5% (p ≤ 0.05).

All data analyses were conducted using the SPSS 10.0.5 for Windows (SPSS Inc, Chicago, Illinois, USA).

RESULTS
The characteristics of the study sample are shown in table 1. At the time of the survey, 386 subjects (74.2%) had at least tried tobacco, 115 (22.1%) were current users (reported use at least monthly), and 102 (19.7%) were recent users (use in the week before the survey). The sensitivity and specificity of the questionnaire compared to the saliva cotinine test (gold standard) were 90% and 93%, respectively, at the cut off of 5 ng/ml. The overall concordance between the two measures was 93%. At the most commonly adopted cut off, 15 ng/ml, the questionnaire had a sensitivity of 93% and a specificity of 92%. Table 2 shows the cross classification of the cotinine test by self reported use. Ninety-eight per cent of subjects reporting no use or less than monthly use had a saliva cotinine concentration above 5 ng/ml (false negative according to the questionnaire), seven reported exposures to environmental smoke and three subjects reported recent use (past week). Among daily users about 96% had concentrations above cut-off. Among subjects reporting weekly, but not daily, use, every second adolescent had a cotinine concentration below the cut off (false positives). A comparison between the two different sets of

Table 1 Characteristics of the study sample

<table>
<thead>
<tr>
<th>Characteristics of the study sample</th>
<th>Number</th>
<th>%</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Sex</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Boys</td>
<td>255</td>
<td>49.0</td>
</tr>
<tr>
<td>Girls</td>
<td>265</td>
<td>51.0</td>
</tr>
<tr>
<td><strong>Parental education index</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Compulsory school</td>
<td>61</td>
<td>11.8</td>
</tr>
<tr>
<td>College</td>
<td>202</td>
<td>39.2</td>
</tr>
<tr>
<td>University</td>
<td>252</td>
<td>48.5</td>
</tr>
<tr>
<td><strong>Lifetime use of tobacco</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>None</td>
<td>134</td>
<td>25.8</td>
</tr>
<tr>
<td>Snus only</td>
<td>54</td>
<td>10.4</td>
</tr>
<tr>
<td>Smoking only</td>
<td>114</td>
<td>21.9</td>
</tr>
<tr>
<td>Snus and smoking</td>
<td>218</td>
<td>41.9</td>
</tr>
</tbody>
</table>

*At baseline for details see Tobacco Control 2003;12:74–8.

Validity of self-reports in adolescents


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questions concerning tobacco use revealed that “non- current” users also reported being “non-recent” users in 389/404 (96%) of cases (data not shown).

Among current users 69 (61.0%) were exclusive smokers, 28 (24.8%) used only snus, and 16 (14.2%) used both types of tobacco (two subjects had incomplete data). The tobacco consumption reported by regular smokers (mean 47 cigarettes/week), snus users (mean 31 pinches/week), and mixed users (mean 66 cigarettes or pinches/week) correlated significantly to saliva cotinine concentration (Spearman $r = 0.42$; $p < 0.01$) (data not shown).

The distribution of saliva cotinine concentration among current users is shown in fig 1. The median concentration was significantly different between exclusive smokers, exclusive users of snus, and mixed users. Concordantly, there was a significant difference in the proportions classified as positive by the cotinine test among self reported users (table 3). In fact, one third of the smokers had cotinine concentrations below the threshold, while this was found in nearly one out of six snus users and for only one in 16 mixed users.

In analyses using logistic regression, current users had a higher likelihood of discordance between self reports and cotinine tests, compared to non-users. Predictors of discordance were also: recentness (use not occurring the very same day of the survey); use of a moderate amount of tobacco per week (1–30 pieces—that is, cigarettes and snus dips) compared to heavier use (more than 30 tobacco pieces per week), and patterns of product use. Exclusive cigarette users were more likely to be classified discordantly by the questionnaire versus cotinine test compared to snus and mixed users (OR 3.2; 95% CI 1.2 to 8.6), even after adjustment for quantity or recentness of use. The adjusted estimates, however, were imprecise because of the paucity of the observations.

DISCUSSION

In this sample of Swedish 15 year olds there was a high overall correspondence between self reported tobacco use and use assessed by saliva cotinine. Indeed, the correspondence was higher for self reported non-users than for users. Overall, the correlation between the biological marker and the reported patterns of use (frequency, recentness, and intensity) was very good. A considerable discrepancy was found among subjects reporting non-daily tobacco use: half of the weekly users, for example, were classified as non-users by the biological test. The discrepancy is likely explained by the very unstable patterns of tobacco use in this age group, for which recent consumption does not imply intensive, nor regular use.10 In fact, infrequent and low level users in this cohort were less likely to be consistently classified by the two instruments than non-users or heavy/frequent users. It should also be noted that patterns of inhalation might vary substantially in this age group, not only between subjects, but also within subjects on different occasions.11 Our study also shows that recent and habitual use probably capture the same conceptual definition of intensity of use—in fact, virtually all of “non-current” users were also “non-recent” users. More self reported smokers than snus users or combined users fell below the cut-off value of the biological marker, and were therefore classified as non-users. To our knowledge this comparison has not been reported before. Since there is no reason to believe that smokers, more than snus users, would intentionally exaggerate their tobacco consumption, this difference may be explained in terms of frequency/intensity of use. However, in this study the type of tobacco used remained a predictor of discordance between self report and biological classification, even after adjustment for quantity or recentness of use, indicating that factors connected to topography of use and/or uptake of nicotine also play a role. For example, the exposure to nicotine from snus (a pinch of tobacco put under the lip and left in place for approximately 30 minutes) is likely to be more prolonged than that occurring after cigarette smoking. Although not a primary interest in this study, this finding deserves further

### Table 2

<table>
<thead>
<tr>
<th>Saliva cotinine (ng/ml)</th>
<th>Current self reported tobacco use</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Not at all or less than every month</td>
</tr>
<tr>
<td></td>
<td>Number</td>
</tr>
<tr>
<td>&lt; 5</td>
<td>395.0</td>
</tr>
<tr>
<td>≥ 5</td>
<td>9.0</td>
</tr>
<tr>
<td>Total</td>
<td>404.0</td>
</tr>
</tbody>
</table>

$\chi^2 p = 0.000.$

### Table 3

<table>
<thead>
<tr>
<th>Saliva cotinine (ng/ml)</th>
<th>Only cigarettes</th>
<th>Only snus</th>
<th>Both cigarettes and snus</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Number</td>
<td>%</td>
<td>Number</td>
</tr>
<tr>
<td>&lt; 5</td>
<td>23.0</td>
<td>5</td>
<td>17.9</td>
</tr>
<tr>
<td>≥ 5</td>
<td>46.0</td>
<td>23</td>
<td>82.1</td>
</tr>
<tr>
<td>Total</td>
<td>69.0</td>
<td>100</td>
<td>100</td>
</tr>
</tbody>
</table>

$\chi^2 p = 0.045.$

![Figure 1](https://www.tobaccocontrol.com)

**Figure 1** Quartiles, mean, and range of cotinine concentrations among current (at least monthly) users of the different tobacco products. Kruskall-Wallis test for difference between medians $p < 0.001$. *Extreme value."
What this paper adds

In general, the validity of adolescent self-reported smoking has been found to be accurate whereas little is known about smokeless tobacco users. Discordant results between the biological marker and reports could be explained by, for example, fear of breaking confidentiality, unstable patterns of tobacco use, lack of specificity of questionnaires, or by the biological marker examined.

This is the first biochemically verified study of self reports among Swedish adolescent smokers and smokeless tobacco (snus) users. Using a cut point of only 5 ng/ml saliva cotinine to identify active tobacco use, there was a 98% concordance between self reported non-use and cotinine concentration. More self reported smokers than snus users or combined users fell below the cut off value of the biological marker. The type of tobacco use remained a predictor of discordance between self report and biological classification, even after adjustment for quantity or recentness of use.

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attention because of its obvious implication for the natural history of nicotine dependence among young snus users.

This study has some weaknesses. First, it was not possible to standardize the time interval between self report and cotinine test, since the test procedure had to be coordinated with the annual survey. This may have caused some occasional users to be classified as non-users by the biological marker. At the time of the survey the study subjects were almost certainly not aware that the test was soon going to be performed, because the information received some months earlier was not repeated, and was never announced at the time of the test. This makes the possibility of behavioural or report modification unlikely. This possibility cannot be ruled out for school absentee tested at a later date, but very few subjects fell into this group. Because of the relatively low prevalence of regular tobacco use among the Swedish youths there were few frequent users, a fact that hampered a more refined analysis of the reliability of the reported intensity of use.

Strengths of the study included the very high participation rate, the detailed history of both cigarette smoking and smokeless tobacco use, as well as the very sensitive assay employed for the test. The set of questions tackling tobacco use in the annual questionnaire seems to classify correctly the different tobacco patterns typical of adolescence. This is, to our knowledge, the first study of biochemical verified self reports of tobacco use among Swedish adolescents and one of few internationally published covering smokeless tobacco use.13

Above all this study shows that adolescents give valid reports of tobacco use. Our results also suggest that the high reliability already observed in other youth studies is not affected by strongly negative societal attitudes towards tobacco use, cigarette smoking in particular.13 A cut off of 5 ng/ml discriminated tobacco users from non-users in this study without appreciable loss in sensitivity and specificity compared to the most commonly adopted cut off of 15 ng/ml.

The generally low exposure to passive smoking in Sweden, especially when children and adolescents are concerned, might also have contributed to the very good correspondence between gold standard and self reports at a very low cut off.14 The question of cost effectiveness of biological validation of tobacco use in future adolescents’ studies should therefore be considered.

ACKNOWLEDGEMENTS

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REFERENCES