Association of snuff use with chronic bronchitis among South African women: implications for tobacco harm reduction

O A Ayo-Yusuf,1 P S Reddy,2 B W van den Borne3

ABSTRACT

Objective: Nasal use of snuff is the predominant form of tobacco use among black South African women. This study examines the association between snuff use and chronic bronchitis (CB) among black South African women.

Design: The study investigated a nationally representative sample of 4464 black South African women ≥25 years old who participated in the 1998 South African Demographic and Health Survey. Data on participants’ tobacco use patterns, medical history and other relevant factors were obtained through an interviewer-administered questionnaire. Peak expiratory flow rates (PEFR) were also measured. Data analysis included χ2 statistics, Student t tests and multiple logistic regression analysis.

Outcome measure: CB, defined as reporting a productive cough for ≥3 months/year for at least 2 successive years.

Results: The prevalence of current snuff use was 16.1% (n = 719). Compared to non-users of snuff, snuff users were not only more likely to present with a history of tuberculosis (TB) (23.3% vs 15.9%; p = 0.06), but they were also more likely to present with CB (5.3% vs 2.8%; p < 0.01) and a lower PEFR (275 litres/min vs 293 litres/min; p < 0.01). Significant determinants of CB included snuff use > 8 times/day (odds ratio (OR) 2.86, 95% CI 1.17 to 7.02), a history of TB (OR 7.23, 95% CI 4.02 to 13.03), current smoking (OR 2.84, 95% CI 1.60 to 5.04) and exposure to smoky cooking fuels (OR 1.98, 95% CI 1.17 to 7.02), a history of TB (OR 7.23, 95% CI 4.02 to 13.03), current smoking (OR 2.84, 95% CI 1.60 to 5.04) and exposure to smoky cooking fuels (OR 1.98, 95% CI 1.17 to 7.02).

Conclusions: These data suggest that snuff use, in the form predominantly used in South Africa, increases the risk of CB. This challenges the idea that snuff may be a less harmful alternative to smoking in South Africa.

Both cigarette smoking and smokeless tobacco (SLT) or snuff use are common in South Africa, with an estimated national prevalence of 24.6% and 6.7% respectively in 1998.1 Unlike in most western countries, about 80% of snuff is taken nasally in South Africa. While fewer than 1% of South African men use snuff, among black South African women snuff is the predominant form of tobacco use, with a prevalence of 13.2% among women in 1998 compared to the 5.3% prevalence of smoking among women.1 Of particular concern is a recent report of the World Health Organization (WHO) global youth tobacco survey, which suggests that in 2002 the prevalence of snuff use among South African adolescents was 14.5%.3

There are traditional homemade and a limited range of commercial/industrialised SLT products on the South African market. The nicotine delivery potentials of the popular SLT products have been previously published.3 Snuff, available as fine powdered tobacco, is much less expensive than cigarettes in South Africa; a can of snuff typically costs 3 Rands (US$0.40) compared to 14 Rands (US$2.00) for a pack of cigarettes. The concentration of free base nicotine available for absorption by the users of South African SLT products previously tested was comparable and sometimes greater than that obtainable from commercial filtered cigarettes, depending on the heaviness of use. The most popular traditional and industrialised products on the South African market, typically with a moisture content of about 40%,3 are used both by oral and nasal application. The route of delivery therefore does not appear to be product-specific in South Africa, but may be dependent on cultural preferences as suggested by a recent study finding.4

Most of the existing literature on the health effects of snuff comes from western nations, particularly Sweden, where most users are Caucasian male snuff dippers. Since consistent association between snuff and major diseases is lacking (particularly in developed nations), snuff is widely considered to be considerably less harmful than cigarettes and snuff is therefore promoted as a reduced-harm product.5 Although the snuff products commonly used in South Africa may differ from those used in such western countries,6 in South Africa, too, snuff use is perceived by some adolescents to be a safer alternative to cigarettes.7 New Zealand is currently considering allowing the sale of nasal snuff as part of a harm-reduction strategy. The strategy has been proposed by some members of the public health community.8 Nasal snuff is also reported to be making its way back onto the English market as a result of the recent ban on public smoking.9

In fact, long-term use of certain types of snuff has been associated with adverse reproductive outcomes10 and the development of oropharyngeal and upper respiratory tract cancers.10-13 The results of recent large-scale studies have suggested that compared to non-use of any tobacco product, snuff use is associated with an increased risk of pancreatic cancer,14 higher mortality from all causes,15,16 and an increased risk of lung cancer mortality17 and lung cancer incidence among elderly American women.18 Snuff use and chronic bronchitis (CB) have also previously been reported as strong risk factors for lung cancer in a Moroccan population,19 but the association between snuff use and CB has not been previously reported.
None of the reviews of the earlier literature on the health effects of oral or nasal use of snuff has explored a link between snuff use and CB, despite the fact that even during nasal application snuff dust can conceivably be inhaled, and snuff has been suggested as a source of bacteria in CB. A more recent study has also questioned the microbiological safety of some finished snuff products on the South African market. The majority of users in South Africa use snuff with high nicotine-delivery capabilities, and nicotine has been suggested to be a potent bronchial gland stimulant. This study therefore sought to determine the association between snuff use and CB among black South African women.

**METHOD**

**Data source and study design**

Data for this study were obtained from black South African women aged 25 years and older (n = 4,464) who participated in the first (and so far the largest publicly available) South African Demographic and Health Survey (SADHS), a nationally representative, cross-sectional household questionnaire survey conducted between February and September 1998. Questionnaires were prepared in all 11 of South Africa’s official languages and translated to check for consistency of meaning. The 1998 SADHS used a stratified, two-stage probability sample design. The methods used to standardise data collection, and the interview and consent procedures have been previously published. The protocol for the SADHS was approved by the ethics committee of the South African Medical Research Council.

**CB risk-factor assessment and definitions**

The questionnaire was administered by trained fieldworkers. It inquired into socio-demographic characteristics and past medical history, including a past history of tuberculosis (TB) infection. An asset index, which is a measure of socioeconomic status, was derived from a composite score of five household items (electricity, television, telephone, refrigerator and washing machine) owned by respondents as identified in a factor analysis for inclusion. The respondents answered ‘yes’ (code 1) or ‘no’ (code 0) to each of the listed household items on the questionnaire. The scale can be considered to be very reliable, because it has a Cronbach alpha score of 0.80. The scores that were obtained were then ranked to classify the respondents into three socioeconomic categories.

Respondents who answered ‘yes’ to the question ‘Have you ever smoked tobacco?’ were categorised as ‘ever’ smokers. Those who said that they had never smoked but that they may have used snuff were classified as ‘never’ smokers. ‘Ever’ smokers were further categorised as ‘current regular smokers’ if they said that they currently smoke daily or occasionally, and as ‘ex-smokers’ if they said that they had previously smoked daily, but did not smoke at all when the survey was conducted. Respondents who said that smoking was a problem were asked about average daily frequency of snuff use. There was no question on the past history of snuff use in the 1998 SADHS. Other exposure and personal history variables included occupational exposure to dust/fumes/strong smells, domestic fuel exposure and domestic and work exposure to second-hand smoke.

Data on the symptomology of CB were based on a series of four standardised questions on chronic productive coughing, adapted from the internationally used British Medical Research Council Questionnaire. CB was defined as presenting with a history of a complex of symptoms including a chronic cough with phlegm every day for at least 3 months of the year for at least 2 successive years. As part of an effort to internally validate the questionnaire diagnosis of CB, each participant’s peak expiratory flow rate (PEFR) was tested at his/her home using a Tru-Zone mini-peak flow meter (Trudell Medical International, London, Ontario, Canada).

**Statistical analysis**

All statistical analyses were done using STATA Release 8 (Stata Corporation, College Station, Texas, USA) with appropriate adjustment made for selection probabilities and the complex sample design of the SADHS. Data were summarised as prevalence rates (%) with 95% confidence intervals. Statistical comparisons were made between the PEFR and CB prevalence among current and non-current snuff users. Group differences were assessed using χ² statistics and independent sample Student t tests. Multiple logistic regression models were constructed to determine the independent association between different intensities of snuff use and CB prevalence, while controlling for the influence of age, smoking duration and other covariates identified in both published literature and our bivariate analysis as being significantly associated with snuff use and CB. Guided by the previously reported frequency of snuff use, for the purpose of analyses, respondents were categorised as those who do not currently use snuff, those who use 1–8 times/day and those who use >8 times/day. As there were too few women snuff users currently smoking (n = 4) and also too few ex-smokers currently using snuff (n = 10), these individuals were grouped as smokers and ex-smokers respectively during subsequent statistical analyses. Sensitivity analysis showed that exclusion of these categories of snuff users did not change the parameter estimates obtained in this study. Statistical significance was set at p<0.05.

**RESULTS**

Of the study sample, 16.1% (n = 719) reported current snuff use and 3.2% (n = 142) reported CB. Subjects who met the questionnaire diagnostic criteria for CB had significantly lower mean PEFR values than those who did not (257 litres/min vs 292 litres/min; p<0.01). The mean difference was 55 litres/min (95% CI 38–71). The prevalence of both snuff use and of CB were lowest among the most educated respondents and those in the highest socioeconomic class, and the incidence of CB was highest among those who reported a past history of TB (table 1). Compared to those without a history of TB, respondents with a history of TB were also more likely to be current snuff users (23.3% vs 15.9%; p = 0.06).

Current snuff users and current smokers were more likely to report CB than non-users of tobacco. Compared to those not currently using snuff, current snuff users were also more likely to present with a lower PEFR (275 litres/min vs 293 litres/min; p<0.01). A history of domestic and/or occupational exposure to second-hand smoke was not significantly associated with CB (age-adjusted odds ratio (OR) 1.01, 95% CI 0.67–1.54), but occupational exposure to dust/fumes/strong smells and domestic exposure to smoking cooking fuels were strongly associated with both CB and snuff use in both the bivariate and multivariate analyses. Even after controlling for these potential confounders in a multivariate analysis, although the effect of snuff use was found to be grossly attenuated, a dose–response relationship remained between snuff use and CB (table 2). Snuff use of >8 times/day remained significantly associated with CB.
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### Table 1
Prevalence of snuff use and chronic bronchitis relative to socio-demographic and exposure characteristics

<table>
<thead>
<tr>
<th>Characteristics</th>
<th>Snuff use (%)*</th>
<th>Chronic bronchitis (%)*</th>
</tr>
</thead>
<tbody>
<tr>
<td>Prevalence (95% CI)</td>
<td>16.1 (14.6–17.6)</td>
<td>3.2 (2.6–3.8)</td>
</tr>
<tr>
<td>Asset index (tertiles) (n = 4447):</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Poorest</td>
<td>18.5</td>
<td>5.6</td>
</tr>
<tr>
<td>Middle</td>
<td>16.6</td>
<td>2.4</td>
</tr>
<tr>
<td>Richest</td>
<td>10.6</td>
<td>1.3</td>
</tr>
<tr>
<td>p Value</td>
<td>&lt;0.001</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>Education level (n = 4449):</td>
<td></td>
<td></td>
</tr>
<tr>
<td>None</td>
<td>2.45</td>
<td>5.6</td>
</tr>
<tr>
<td>1–7 years schooling</td>
<td>16.6</td>
<td>3.3</td>
</tr>
<tr>
<td>8–12 years schooling</td>
<td>11.5</td>
<td>1.8</td>
</tr>
<tr>
<td>&gt;12 years schooling</td>
<td>2.5</td>
<td>–</td>
</tr>
<tr>
<td>p Value</td>
<td>&lt;0.001</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>History of tuberculosis (n = 4451):</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Never</td>
<td>15.9</td>
<td>2.8</td>
</tr>
<tr>
<td>Ever</td>
<td>23.3</td>
<td>21.1</td>
</tr>
<tr>
<td>p Value</td>
<td>0.06</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>Cooking fuel used (n = 4455):</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Gas/electricity/paraffin</td>
<td>15.2</td>
<td>2.3</td>
</tr>
<tr>
<td>Wood/coal/dung (smoky)</td>
<td>19.1</td>
<td>6.2</td>
</tr>
<tr>
<td>p Value</td>
<td>0.02</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>Occupational exposure (n = 4455):</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Never</td>
<td>15.8</td>
<td>2.8</td>
</tr>
<tr>
<td>Ever</td>
<td>21.7</td>
<td>5.8</td>
</tr>
<tr>
<td>p Value</td>
<td>&lt;0.01</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>Smoking status (n = 4464):</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Never smoked</td>
<td>17.1</td>
<td>2.9</td>
</tr>
<tr>
<td>Ex-smoker</td>
<td>14.9</td>
<td>1.5</td>
</tr>
<tr>
<td>Current smoker</td>
<td>1.3</td>
<td>7.3</td>
</tr>
<tr>
<td>p Value</td>
<td>&lt;0.001</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>Snuff use status (n = 4464):</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Non-user</td>
<td>2.8</td>
<td></td>
</tr>
<tr>
<td>Use 1–8 times/day</td>
<td>4.3</td>
<td></td>
</tr>
<tr>
<td>Use &gt;8 times/day</td>
<td>11.7</td>
<td></td>
</tr>
<tr>
<td>p Value</td>
<td>&lt;0.01</td>
<td></td>
</tr>
</tbody>
</table>

Sample sizes for some variables do not sum to the total (4464) due to missing data.

*Weighted prevalence estimates.

(fully adjusted OR = 2.86, 95% CI 1.17–7.02) and so was current smoking (fully adjusted OR 2.84, CI 1.60–5.04), irrespective of the intensity of smoking. However, a non-linear dose–response was observed between the risk of CB and the reported number of cigarettes smoked per day (CPD) (table 2). The median CPD was four. Compared to never smokers, those who reported smoking 1–4 CPD had a higher risk of presenting with CB than those who reported smoking >4 CPD.

### DISCUSSION

This study demonstrates the first findings of a significant association between snuff or SLT use and CB. These findings also demonstrate common risk factors for snuff use and a diagnosis of CB, suggesting that snuff users in South Africa could be at a cumulative disadvantage for developing chronic airway diseases. Also significant is this study’s support of a previous report,23 which suggested that a history of TB infection was strongly associated with CB. In addition, our study suggests that snuff users are more likely to present with a history of TB. This observation calls for further investigation of the potential risk of snuff users spreading TB or being at high risk of contracting TB, especially in countries like South Africa which have a relatively high prevalence of TB. Snuff users and those suffering from CB are also more likely to be poor, uneducated women, who are more likely to cook with smoky fuels and work in conditions where they are exposed to dust, fumes and/or strong smells. Significantly, however, even after controlling for these potential confounders, this study shows that current daily heavy (>8 times/day) snuff users, but not less frequent users, had almost three times greater odds of developing CB when compared to non-users of snuff. By contrast, albeit consistent with the literature,23 those smoking even at low rates of 1–4 CPD as compared to never smokers were significantly more likely to present with a history of CB.

Although somewhat consistent with the findings of a similar study that reported the correlates of occurrence of CB among the general population of South African women,24 the unexpected finding that black South African women that reported smoking 1–4 CPD had higher odds of developing CB than those smoking >4 CPD may be related to compensatory changes in smoking behaviour. This view is supported by findings of a recent study that showed a similar unexpected non-linear relationship between changes in lung function and reduced smoking rates among smokers with early chronic obstructive pulmonary disease (COPD).24 It is indeed possible that heavy smokers that experienced early symptoms of COPD may have opted to reduce their smoking rates, but because of compensatory smoking (eg deeper and more frequent puffs) would effectively still be exposed to similar or even greater levels of cigarette toxins as before the reduction. Nonetheless, the unexpected observation of the larger impact of lower smoking rate on lung health in the current study may also be related to reporting bias. In a society where smoking by black South African women in particular is traditionally viewed as socially unacceptable, it is conceivable that heavier smokers may find it more socially desirable to report lower smoking rates.

Although it is still a possibility that uncontrolled residual confounders may explain the observed association of snuff use with CB, it is biologically plausible that nasal use of snuff could lead to CB. A previous report notes nasal symptoms associated with nasal mucosal changes in snuff users and these nasal symptoms in turn have been associated with increased occurrence of CB.31 Furthermore, this study’s finding would be consistent with that of a previous case study of CB, which suggests that nasal use of snuff could have been the source of the bacteria in CB.23

Contrary to a recent suggestion that smoke constituents other than nicotine may be responsible for increased pulmonary airway resistance,32 this study’s findings, supported by results from other studies,23 33 34 suggest that the nicotine in snuff could be associated with hyper-secretion of local bronchial glands and could therefore contribute to CB, especially in the presence of bacterial infection. Nicotine has also been shown to prolong the survival of neutrophils,35 the accumulation of which in the lungs has been implicated in the pathogenesis of CB.36 The current study’s findings may differ from those of other studies because the snuff products used in South Africa have a relatively high occurrence of CB.31 Furthermore, this study’s finding would be consistent with that of a previous case study of CB, which suggests that nasal use of snuff could have been the source of the bacteria in CB.23

The current study’s findings may differ from those of other studies because the snuff products used in South Africa have a relatively high prevalence of TB. Snuff users and those suffering from CB are also more likely to be poor, uneducated women, who are more likely to cook with smoky fuels and work in conditions where they are exposed to dust, fumes and/or strong smells. Significantly, however, even after controlling for these potential confounders, this study shows that current daily heavy (>8 times/day) snuff users, but not less frequent users, had almost three times greater odds of developing CB when compared to non-users of snuff. By contrast, albeit consistent with the literature, those smoking even at low rates of 1–4 CPD as compared to never smokers were significantly more likely to present with a history of CB.

Although somewhat consistent with the findings of a similar study that reported the correlates of occurrence of CB among the general population of South African women, the unexpected finding that black South African women that reported smoking 1–4 CPD had higher odds of developing CB than those smoking >4 CPD may be related to compensatory changes in smoking behaviour. This view is supported by findings of a recent study that showed a similar unexpected non-linear relationship between changes in lung function and reduced smoking rates among smokers with early chronic obstructive pulmonary disease (COPD). It is indeed possible that heavy smokers that experienced early symptoms of COPD may have opted to reduce their smoking rates, but because of compensatory smoking (eg deeper and more frequent puffs) would effectively still be exposed to similar or even greater levels of cigarette toxins as before the reduction. Nonetheless, the unexpected observation of the larger impact of lower smoking rate on lung health in the current study may also be related to reporting bias. In a society where smoking by black South African women in particular is traditionally viewed as socially unacceptable, it is conceivable that heavier smokers may find it more socially desirable to report lower smoking rates.

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Contrary to a recent suggestion that smoke constituents other than nicotine may be responsible for increased pulmonary airway resistance, this study’s findings, supported by results from other studies, suggest that the nicotine in snuff could be associated with hyper-secretion of local bronchial glands and could therefore contribute to CB, especially in the presence of bacterial infection. Nicotine has also been shown to prolong the survival of neutrophils, the accumulation of which in the lungs has been implicated in the pathogenesis of CB. The current study’s findings may differ from those of other studies because the snuff products used in South Africa have a relatively high pH and thus a higher nicotine-delivery capability than the SLT products commercially available elsewhere. It is however, pertinent to note that Bolinder et al also previously reported a significantly increased risk for reporting symptoms of COPD in a Swedish cohort of oral SLT users, as compared to non-users. Considering that similar respiratory problems have been noted among tobacco factory workers, although to a lesser degree, it is also possible that respirable snuff dust particles may...
considerably lower health risks compared to smoking. 6  
The South African market, have been reported to pose  
moist SLTs, some of which have recently been introduced onto  

This study’s finding that a current frequent snuff user’s risk  
weakens arguments that nasal snuff use may be sufficiently less  
harmful than smoking to be worth promoting as an alternative.  
Taking into account the risk-use equilibrium,27 promoting such  
a strategy in the studied population could be a potentially risky  
population experiment, given that the current number of black  
South African women snuff users is already almost three times  
higher than the number of smokers. Furthermore, given that  
any suggestion of the relative safety of any snuff product in a  
fairly uneducated population could inadvertently encourage  
continued or increased snuff use, such a harm-reduction  
strategy could potentially increase total tobacco-related disease  
in the studied population, especially since a significant decline in  
smoking rates has already been observed in this population due  
to current governmental policy.14 2 Such a strategy would also  
be problematic in parts of Africa and India where the nasal use  
of snuff is still relatively common and the relatively high cost of  
modern oral SLT products may be prohibitive. Considering that  
these third-world populations of snuff users may be larger in  
number than those of current SLT users in developed countries,  
the findings of this study suggest that future policy debates on  
SLT as a harm-reduction strategy need to be broadened to  
include the implications of such a strategy for populations  
outside developed countries.

Study limitations
Our study findings are subject to some limitations. Firstly, as  
mentioned above, the findings may not be generalisable to other  
SLT products. Secondly, the study relies on self-reporting of  
tobacco use and CB. The participants’ responses are therefore  
potentially subject to reporting bias, especially in respect of  
smoking, which is generally regarded as a social taboo among  
black South African women. It is therefore possible that such  
reporting bias may have weakened the association reported  
between smoking and CB. Furthermore, the responses to  
questions used for CB diagnosis may be subject to recall bias.  
Nevertheless, the criteria used for CB diagnosis are similar to  
those used in many other national surveys, with which our

<table>
<thead>
<tr>
<th>Table 2</th>
<th>Multiple logistic regression model for chronic bronchitis (n = 4425)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Crude odds ratio (95% CI)</td>
</tr>
<tr>
<td>Snuff use:</td>
<td></td>
</tr>
<tr>
<td>Never/none-current</td>
<td>1</td>
</tr>
<tr>
<td>1–8 times/day</td>
<td>1.56 (0.96 to 2.55)</td>
</tr>
<tr>
<td>&gt;8 times/day</td>
<td>4.63 (2.09 to 10.22)</td>
</tr>
<tr>
<td>Smoking status:</td>
<td></td>
</tr>
<tr>
<td>Never smoked</td>
<td>1</td>
</tr>
<tr>
<td>Ex-smoker</td>
<td>0.48 (0.12 to 1.99)</td>
</tr>
<tr>
<td>1–4 CPD</td>
<td>2.96 (1.43 to 6.11)</td>
</tr>
<tr>
<td>&gt;4 CPD</td>
<td>2.31 (1.16 to 4.59)</td>
</tr>
<tr>
<td>Asset (tertile) index:</td>
<td></td>
</tr>
<tr>
<td>Lowest rank (poorest)</td>
<td>1</td>
</tr>
<tr>
<td>Middle</td>
<td>0.44 (0.30 to 0.66)</td>
</tr>
<tr>
<td>Highest rank</td>
<td>0.24 (0.10 to 0.63)</td>
</tr>
<tr>
<td>Tuberculosis:</td>
<td></td>
</tr>
<tr>
<td>Never</td>
<td>1</td>
</tr>
<tr>
<td>Ever</td>
<td>9.41 (5.41 to 16.38)</td>
</tr>
<tr>
<td>Cooking fuel:</td>
<td></td>
</tr>
<tr>
<td>Gas/Electricity/Paraffin</td>
<td>1</td>
</tr>
<tr>
<td>Smoky fuels</td>
<td>2.82 (1.92 to 4.14)</td>
</tr>
<tr>
<td>Occupational exposure to irritants:</td>
<td></td>
</tr>
<tr>
<td>Never</td>
<td>1</td>
</tr>
<tr>
<td>Ever</td>
<td>2.12 (1.31 to 3.43)</td>
</tr>
</tbody>
</table>

*In addition to these covariates, the model also controlled for age,  
CPD, cigarettes per day.

...
results are thus comparable. Moreover, in our study, CB diagnosis was cross-validated with PEFR measurements. Thirdly, because this was a cross-sectional study, reverse causality is possible, although not likely. Accordingly, any inference about causality based on the current study should be made with caution. Snuff use must be confirmed as a true risk factor in longitudinal studies that take into account the temporal order of events. Fourthly, the SADHS did not include relevant information such as the exact form of SLT use, past history of use and the lifetime duration of use. Therefore, age (which is expected to correlate with years of use) was used as a covariate in all the analyses, irrespective of the level of statistical significance detected. Conceivably, if some of the study participants were in fact snuff-dippers or if a significant proportion of non-current users of snuff were ex-users of snuff, these may have attenuated rather than inflated the strength of the association reported for current snuff use in this study. Lastly, the data that were used are somewhat dated. However, considering that the biological relationship hypothesised between snuff and bronchitis is not time-dependent, this is not likely to influence the study findings significantly, except if the snuff products used before 1998 differed from those currently in use. However, the reported pH of a commonly used traditional snuff mixture and that of the most popular industrialised brand tested in the past (unpublished results) are similar to the pH of similar products tested more recently. This implies that the nicotine-delivery capability of these products are also similar. Thus the nicotine-delivery potential of snuff products currently in use is not likely to be considerably lower than those used prior to 1998.

Despite these limitations, this study provides useful information about the risk of SLT products used by populations outside Europe.

CONCLUSIONS

Although further investigation is needed on the exact biological mechanisms, this study demonstrates for the first time that snuff use, in the form predominantly used in South Africa, increases the risk for CB. This challenges the idea that snuff may be a much less harmful alternative to smoking in South Africa and in particular, among black South African women. The study findings also highlight the need for public health interventions that discourage the adoption of any form of tobacco use and promote tobacco use cessation, particularly among South African adolescents.

What this paper adds

Only limited information is available about health risks associated with the use of smokeless tobacco (SLT) products other than those currently available in Europe and North America. However, the harm-reduction debate extends to populations as far removed from such markets as South Africa and New Zealand, where nasal use of snuff is either currently the predominant form of SLT use or the form currently being advocated. This study provides useful information needed to improve the current basis for risk assessment of SLT use in populations outside of Europe. This study’s findings suggest that heavy snuff use is equally as likely to be associated with chronic bronchitis as smoking in the studied South African population, challenge any public health strategy that may want to promote a switch to SLT use as a public policy response for the reduction of tobacco-related diseases in South Africa.

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Competing interests: None.

REFERENCES


FUNDING AVAILABLE FOR RESEARCH PROJECTS

The Committee on Publication Ethics (COPE) has established a Grant Scheme to fund research in the field of publication ethics. The Scheme is designed to provide financial support to any member of COPE for a defined research project that is in the broad area of the organisation’s interests, and specifically in the area of ethical standards and practice in biomedical publishing. The project should have a specific goal and be intended to form the kernel of a future publication.

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An application form can be obtained by contacting Linda Gough, COPE administrator, at LGough@bmj.com or 020 7383 6602. For more information on COPE, see http://www.publicationethics.org.uk/

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