**REVIEW**

Effect of smokeless tobacco (snus) on smoking and public health in Sweden

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Objective: To review the evidence on the effects of moist smokeless tobacco (snus) on smoking and ill health in Sweden.

Method: Narrative review of published papers and other data sources (for example, conference abstracts and internet based information) on snus use, use of other tobacco products, and changes in health status in Sweden.

Results: Snus is manufactured and stored in a manner that causes it to deliver lower concentrations of some harmful chemicals than other tobacco products, although it can deliver high doses of nicotine. It is dependence forming, but does not appear to cause cancer or respiratory diseases. It may cause a slight increase in cardiovascular risks and is likely to be harmful to the unborn fetus, although these risks are lower than those caused by smoking. There has been a larger drop in male daily smoking (from 40% in 1976 to 15% in 2002) than female daily smoking (34% in 1976 to 20% in 2002) in Sweden, with a substantial proportion (around 30%) of male ex-smokers using snus when quitting smoking. Over the same time period, rates of lung cancer and myocardial infarction have dropped significantly faster among Swedish men than women and remain at low levels as compared with other developed countries with a long history of tobacco use.

Conclusions: Snus availability in Sweden appears to have contributed to the unusually low rates of smoking among Swedish men by helping them transfer to a notably less harmful form of nicotine dependence.

In recent times the tobacco industry has been active in developing and marketing new products that might be perceived as less harmful to health than typical cigarettes. At the same time, there has been an increasingly vigorous debate within the public health community over the most appropriate response to the new products being developed by the industry. In this debate, public health advocates have been mindful of the historical precedents set by previous tobacco industry attempts to introduce new product lines that have been perceived as less harmful. It is now clear that so called “light” cigarettes were widely believed to be less harmful (and continue to be by the majority of consumers) but in fact are no less deadly than standard cigarettes. The introduction and marketing of these products may well have had a serious adverse effect on public health by duping hundreds of millions of smokers into the belief that they could continue to smoke at reduced risk.

In the current debate over tobacco harm reduction, some have cited the “Swedish experience” as an example of tobacco specific nitrosamines (TSNAs). It has become clear that different selection and curing methods can affect the levels of nitrates and hence TSNAs present in the raw tobacco before processing. Over recent decades snus manufacturers have selected tobacco blends that have been air and sun cured (dried), while US moist snuff products tend to include blends high in fire cured tobacco.

After curing, raw cured tobacco is cut into small strips, dried, ground, and sifted before processing. In Sweden, by tradition, snus production has included a process in which the tobacco is heat treated with steam for 24–36 hours (reaching temperatures of approximately 100°C). Ingredients added are: 45–60% water, 1.5–3.5% sodium chloride, 1.2–3.5% sodium bicarbonate, and less than 1% flavouring. It is claimed that the heating process kills bacteria, producing a relatively sterile product. The product is then packaged in cans and refrigerated during storage. In Sweden the product is also kept in refrigerators by the retailers. One study examined levels of carcinogenic TSNAs in snus kept at temperatures ranging from −20°C to +23°C for 20 weeks. This exposure to a variety of temperatures over time did not produce a significant increase in concentrations of TSNAs, suggesting that the exposure to heat during manufacturing is unlikely to be of concern.

**MANUFACTURING**

Snus both contains and delivers a number of harmful substances, including cancer-causing tobacco specific nitrosamines (TSNAs). It has become clear that different selection and curing methods can affect the levels of nitrates and hence TSNAs present in the raw tobacco before processing. Over recent decades snus manufacturers have selected tobacco blends that have been air and sun cured (dried), while US moist snuff products tend to include blends high in fire cured tobacco.

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manufacturing may itself have prevented microbial activation of nitrates.10
This manufacturing process contrasts with that traditionally used in the USA, in which the product is fermented (rather than being subject to high temperatures), allowing the continued formation of TSNAS. In addition, North American smokeless tobacco is not typically stored in refrigerators. One study found that nitrite and TSNA levels increased significantly in US snuff stored at 37°C for four weeks.11
Although different products vary in their pH levels, snus typically has a pH in the range 7.8–8.5.12–13 This is important because only nicotine in the free-base form is rapidly absorbed through the mucosal membrane, and the proportion of free-base nicotine available from tobacco is determined by the pH level. For example Brunnemann and Hoffmann compared two brands and found that one brand with a pH of 5.84 had only 1% of the nicotine in the free-base form and another brand with a pH of 7.99 had 59% of the nicotine available in free-base form for absorption.14 Another study found that a leading Swedish snus brand had a higher pH (and therefore probably more efficient nicotine delivery) than five comparison brands of US smokeless tobacco.15

Delivery of harmful substances
Possibly as a result of the differences in manufacturing and storing procedures, snus has been claimed to contain lower levels of some harmful substances than many of the brands marketed in different countries. The total TSNA concentration varied greatly among the US brands from 4.1 to 128 (mg/g dry tobacco). There is little evidence to support claims that TSNA levels have consistently dropped over the past decade in North American snuff (for example, Copenhagen brand in 1994 had a measured TSNA level of 17.2 and in 2000 it was 13.8). Snus brands selected in Sweden from 1990, 1991, and 1994 had a measured TSNA level of 17.2 and in 2000 it was 11.2

Table 2

<table>
<thead>
<tr>
<th>Toxin</th>
<th>Limit</th>
</tr>
</thead>
<tbody>
<tr>
<td>Nitrate</td>
<td>3.5 mg/kg</td>
</tr>
<tr>
<td>Tobacco specific nitroamines (TSNA)</td>
<td>5 mg/kg</td>
</tr>
<tr>
<td>N-Nitrosodimethyamine (NDMA)</td>
<td>5 µg/kg</td>
</tr>
<tr>
<td>Benz(a)pyrene (BaP)</td>
<td>10 µg/kg</td>
</tr>
<tr>
<td>Cadmium</td>
<td>0.5 mg/kg</td>
</tr>
<tr>
<td>Lead</td>
<td>1.0 mg/kg</td>
</tr>
<tr>
<td>Arsenic</td>
<td>0.25 mg/kg</td>
</tr>
<tr>
<td>Nickel</td>
<td>2.25 mg/kg</td>
</tr>
<tr>
<td>Chromium</td>
<td>1.5 mg/kg</td>
</tr>
</tbody>
</table>

mg/kg, thousandth gram per kilogram product (based on Snus with 50% water content); µg/kg, millionth gram per kilogram product (based on Snus with 50% water content); double the limits for dry weight equivalents.

Table 1

<table>
<thead>
<tr>
<th>Country and brand (year sampled)</th>
<th>Manufacturer</th>
<th>Nicotine (mg/g)</th>
<th>NNN (µg/g)</th>
<th>NNN (µg/g)</th>
<th>Total TSNA (µg/g)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sweden</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Three brands* (1990–91)</td>
<td>Swed Match</td>
<td>1.4 – 2.1</td>
<td>5.2 – 5.7</td>
<td>9.2 – 11.2</td>
<td></td>
</tr>
<tr>
<td>Elton Snus (2000)</td>
<td>Swed Match</td>
<td>0.5</td>
<td>1.1</td>
<td>2.8</td>
<td></td>
</tr>
<tr>
<td>Sudan (Toomabak)</td>
<td></td>
<td>32.2 – 102.4</td>
<td>630 – 7870</td>
<td>830 – 3805</td>
<td></td>
</tr>
<tr>
<td>5 Samples* (1990)</td>
<td></td>
<td>8.4 – 26.0</td>
<td>1140 – 2790</td>
<td>420 – 1550</td>
<td></td>
</tr>
<tr>
<td>3 Samples* (1993)</td>
<td></td>
<td>188 – 362</td>
<td>241 – 369</td>
<td></td>
<td></td>
</tr>
<tr>
<td>USA</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>2 Samples* (1991)</td>
<td></td>
<td>18.6 – 20.6</td>
<td>0.5 – 0.8</td>
<td>4.8 – 9.0</td>
<td></td>
</tr>
<tr>
<td>1 Sample* (1992)</td>
<td></td>
<td>16.7</td>
<td>0.6</td>
<td>5.6</td>
<td></td>
</tr>
<tr>
<td>Copenhagen (1994)</td>
<td>USSTC</td>
<td>12 (11.7 – 11.3)</td>
<td>1.9 (1.3 – 2.5)</td>
<td>8.7 (10.1 – 7.3)</td>
<td>17.2 (20.2 – 14.2)</td>
</tr>
<tr>
<td>Skol, Original Fine cut(1994)</td>
<td>USSTC</td>
<td>11.9 (13.4 – 10.7)</td>
<td>1.3 (1.4 – 1.2)</td>
<td>8.2 (9.5 – 6.9)</td>
<td>14.9 (17.4 – 12.4)</td>
</tr>
<tr>
<td>Skol Bands Straight(1994)</td>
<td>USSTC</td>
<td>10.1 (10.9 – 9.3)</td>
<td>0.9 (1.2 – 0.6)</td>
<td>5.1 (6.1 – 4.1)</td>
<td>8.2 (9.9 – 6.5)</td>
</tr>
<tr>
<td>Kodak: Wintergreen(1994)</td>
<td>Canwood</td>
<td>10.9 (10.1 – 11.7)</td>
<td>0.6 (0.8 – 0.4)</td>
<td>6.3 (7.4 – 5.2)</td>
<td>11.0 (13.4 – 8.6)</td>
</tr>
<tr>
<td>Hawken Wintergreen(1994)</td>
<td>Canwood</td>
<td>3.2 (3.4 – 3.0)</td>
<td>0.2 (244 – 16)</td>
<td>3.1 (3.4 – 2.8)</td>
<td>4.1 (4.5 – 3.7)</td>
</tr>
<tr>
<td>Skoll (2000)</td>
<td>USSTC</td>
<td>4.3</td>
<td>20.8</td>
<td>64.0</td>
<td></td>
</tr>
<tr>
<td>Copenhagen(2000)</td>
<td>USSTC</td>
<td>3.4</td>
<td>14.3</td>
<td>41.1</td>
<td></td>
</tr>
<tr>
<td>Timber Wolf(2000)</td>
<td>Swed. Match</td>
<td>0.95</td>
<td>3.0</td>
<td>7.5</td>
<td></td>
</tr>
<tr>
<td>Silver Creek(2000)</td>
<td>Swisher</td>
<td>17.8</td>
<td>41.4</td>
<td>127.9</td>
<td></td>
</tr>
</tbody>
</table>

Table 1.* All available pertinent data from referenced studies is reported. Blank cells indicate that data were not provided in the referenced study.

*Brand not specified in study.
Figure 1: Venous blood concentrations in nanograms of nicotine per millilitre (ng/ml) of plasma as a function of time for various nicotine delivery systems; all plasma nicotine concentrations have been reconfigured such that the pre-absorption level starts at 0 ng/ml (that is, to take out the baseline differences). Cigarette, and 2 mg nicotine gum, adapted from Russell et al, page 100. Swedish snus plasma nicotine concentrations in 10 Swedish snus users from a single 2 g pinch of loose snus adapted from Holm et al.

about twice as high as the nicotine concentrations typically obtained from nicotine replacement therapy. The nicotine levels shown in fig 1 are from users of loose snus and it is possible that some other brands (particularly portion packed products) or those with a lower pH may give different levels.

IS SNUS HARMFUL TO HEALTH AND IS IT LESS HARMFUL TO AN INDIVIDUAL USER THAN CIGARETTES?

Many of the smokeless tobacco users participating in the older epidemiological studies discussed below may have been exposed to products delivering higher quantities of harmful substances than current versions of these products.

Nicotine dependence

Given the pattern of nicotine absorption described above there can be no doubt that snus is dependence forming in much the same way as other forms of tobacco consumption. There is some evidence that the dependence potential of nicotine and other psychoactive drugs is related to their speed of delivery to the brain and so one would expect snus and other non-inhaled forms of nicotine delivery to be proportionately less addictive than inhaled tobacco smoke. However, there is clear evidence that users of products with snus-like nicotine delivery profiles develop cravings and nicotine withdrawal symptoms when attempting to abstain, and find it difficult to quit. While snus probably does not produce strongly nicotine dependence than smoking, it has just minimal, if any, advantages over cigarettes or other smokeless tobacco products in terms of its lower potential to induce dependence. In fact, its high nicotine delivery and hence dependence potential (relative to most other non-smoked delivery modalities) may be a critical factor enabling it to compete with the more rapidly absorbed nicotine from smoked tobacco.

Oral cancer

One of the biggest concerns about the use of smokeless tobacco stems from the relatively large body of evidence from a number of countries showing that oral tobacco use can cause cancer of the mouth, head, and neck. With regards to its use in India, the 2001 US Institute of Medicine (IOM) report stated, “A large number of studies in India, including cohort, case-control, and intervention studies, support an association between oral cancer and smokeless tobacco, and these studies are consistent, strong, coherent and temporally plausible” (p 427). The IOM report stated that toombak users in Sudan also have a much higher relative risk (RR) of oral cancer than non-users and that “In spite of conflicting US data, it can be concluded that snuff use in the United States also increases the risk of oropharyngeal cancers” (p428). In contrast, there is consistent evidence from two case–control studies in Sweden showing no increased risk of cancer of the head, neck, or mouth among snus users.

Schildt and colleagues investigated whether snus leads to increased risk of oral cancer by comparing various risk factors in 410 cases of oral cancer and 410 matched controls identified during the period 1980–89. Ninety six per cent of the identified cases and 91% of identified controls participated in the study (leaving full data from 354 matched pairs) and 20% of the overall sample were current or ex snus users. Univariate analyses found significant increased risk of oral cancer as a result of smoking (odds ratio (OR) 1.8 for active smokers), and alcohol (OR 1.9 for beer drinkers) but no increased risk for active snus use (OR 0.7, 95% confidence interval (CI) 0.4 to 1.1). The authors concluded: “Our results do not support any association between use of oral snuff and oral cancer.”

Levin and colleagues conducted a similarly designed study, identifying cases of head and neck cancer in two regions of Sweden between 1988 and 1991 and matched controls. Interviews were conducted with high proportions of identified cases (90%) and controls (85%). This study found significantly increased risks of head and neck cancers associated with alcohol use and smoking, but no increased risk associated with former or current snus use. The RR for head and neck cancer among snus users as compared with non-snus users, after adjusting for age, region, alcohol, and smoking was 1.0 (95% CI 0.6 to 1.6). Similarly there were no significant relations between duration of snus use or lifetime consumption and head/neck cancer.

A recent systematic review of the health effects of smokeless tobacco concluded: “Chewing betel quid and tobacco is associated with a substantial risk of oral cancers in India. Most recent studies from the US and Scandinavia are not statistically significant, but moderate positive associations cannot be ruled out due to lack of statistical power.” Snus causes a number of non-malignant oral diseases, including oral lesions and dental caries. However, it appears as though the lesions produced by snus are reversible and disappear if snus use ceases.

Other cancers

Ye and colleagues conducted a case control study (504 cases and 1164 controls) examining the effects of smoking, alcohol, and snus use on gastric cancer in Sweden. They found a significant dose and duration related increase in gastric cancer risk with smoking, but no effect of snus or alcohol. They concluded that “smoking, but not the oral use of tobacco in the form of moist snuff, is positively associated with risk of gastric cancer”.

Lagergen et al conducted a case–control study designed to test the association between smoking, snus, and alcohol use, and cancer of the oesophagus and gastric cardia in Sweden. Combined smoking and alcohol use was strongly associated with oesophageal squamous cell carcinoma (OSCC) (OR 23.1 for heavy users compared with never users), but snus use was not significantly associated with any of the cancer sites under study in multivariate analyses. There was some indication of a possible link between snus use and OSCC in that the odds ratio was 2.0 for use for over 25 years versus never snus use.
although because of the relatively small size of this sub-sample (n = 14 cases) this was not significant (95% CI 0.9 to 4.1). The authors concluded: “we found no statistically significant association between snuff dipping and risk of any of the studied tumors.”

It remains possible, but unlikely, that a carcinogenic effect of snus only emerges after very long term use. Bolinder and colleagues found a non-significant RR of death from cancer of 1.1 for snus users compared with never tobacco users (95% CI 0.9 to 1.4) in a prospective study of Swedish construction workers that included a relatively large sample, many of whom had used snus for over 40 years. The RR for cancer death was 1.0 (compared with non-tobacco users) for those over 40 years old (RR 1.0, 95% CI 0.7 to 1.4). This study found a significantly increased all cause mortality in snus users compared with never tobacco users, largely from elevated cardiovascular mortality. The RR for lung cancer among snus users was 0.8, compared with never tobacco users compared with never tobacco users, largely from elevated cardiovascular mortality. The RR for lung cancer among snus users was 0.8, compared with never tobacco users (RR 1.4, 95% CI 1.2 to 1.6). This excess risk was significantly increased for smokers of at least 15 cigarettes per day (95% CI 1.2 to 1.6).

Overall, the results of the five large studies examining snus in relation to cancer are consistent in finding no increased cancer risk among snus users. All of the Swedish studies of the relation between snus and cancer were robust enough to detect significant effects for tobacco smoking (often involving very large effect sizes), and the studies of oral cancer were also able to detect significant relations with alcohol use. The lack of relation with snus is therefore unlikely to be caused by methodological problems such as low statistical power.

Cardiovascular disease

Bolinder and colleagues conducted a series of epidemiological and clinical studies examining the effects of long term snus use on health, focusing on cardiovascular risk factors and myocardial infarction. Their first report focused on a cross sectional study of almost 98 000 Swedish construction workers undergoing health examinations in 1971–4, including over 5000 exclusive snus users. This study found an increased prevalence of circulatory and respiratory symptoms among snus users and heavy smokers as compared to non-tobacco users, and an increased prevalence of hypertension in snus users compared to non-tobacco users. Surprisingly this study found the lowest effects for tobacco smoking (often involving very large effect sizes), and the studies of oral cancer were also able to detect significant relations with alcohol use. The lack of relation with snus is therefore unlikely to be caused by methodological problems such as low statistical power.

Bolinder’s second study examined the relation between tobacco use and cardiovascular mortality in a larger sample (n = 135 036) of Swedish male construction workers recruited at a health examination in 1971–4 and followed up 12 years later.

This study found that snus users had a significantly higher risk of dying from a cardiovascular event than never tobacco users (RR 1.4, 95% CI 1.2 to 1.6). This excess risk was comparable to that of ex-smokers who had quit in the past five years, but smaller than heavy smokers (RR 1.9 compared with never tobacco users). The analyses in this study adjusted for age and region of origin, and (for at least some analyses, although it was not always stated) also adjusted for body mass index, blood pressure, diabetes, and heart problems at the time of entering the study. Alcohol consumption and cholesterol were not measured and so could not be controlled for.

Subsequent studies focused on a smaller sample of Swedish firemen (around 140, split approximately equally between snus users, smokers, and non-tobacco users). These studies found that snus use did not influence exercise capacity, or play a major role in the atherosclerotic process (both of which were adversely affected by smoking). However, they replicated the previous finding of higher daytime (but not night time) heart rate and blood pressure among both snus users and smokers compared to non-tobacco users. Overall, these studies by Bolinder and colleagues are suggestive of an increased cardiovascular risk from snus use, that is probably mediated by nicotine’s sympathetic stimulant effects, and is of a smaller magnitude than the excess cardiovascular risks caused by smoking. It was suggested that snus’ effects on blood pressure may be related to its sodium content (1.3–3.5% sodium chloride and 1.5–3.5% sodium bicarbonate).

However, two subsequent case–control studies by Huhtasaari and colleagues did not find a significantly increased risk of myocardial infarction among snus users as compared to non-tobacco users. Both of these studies were based on data collected in northern Sweden as part of the World Health Organization MONICA (multinational monitoring of trends and determinants in cardiovascular diseases) project. In both reports, the cases and controls were identified in the 1990s.

Huhtasaari and colleagues found an age adjusted OR for myocardial infarction (MI) of 0.89 (95% CI 0.62 to 1.29) for snus use versus no tobacco use, whereas smoking significantly increased risk of an MI (OR 1.87, 95% CI 1.40 to 2.48). In multi-variate analyses smoking remained significantly associated with MI, whereas snus use was not.

Huhtasaari subsequently conducted a larger study than the one reported in 1992, and included more detailed tobacco use histories and closer matching of cases and controls (matched for sex, date of birth, and area of residence). This study found (after adjustment for multiple cardiovascular risk factors) that cigarette smoking significantly increased risk of an MI (OR 3.53, 95% CI 2.48 to 5.03), whereas snus use significantly reduced the risk (OR 0.58, 95% CI 0.35 to 0.94) compared with men who never became regular tobacco users. When the analysis focused only on fatal cases, there was a tendency towards increased risk in snus users, but this was not significant (OR 1.5, 95% CI 0.45 to 5.03).

There is no clear explanation for the difference in results between the Bolinder and Huhtasaari studies, although the different study populations, time periods covered, and outcomes measured (sudden death versus non-fatal MI) may have contributed. The similar magnitude of effect for fatal cardiovascular events found in these studies is suggestive of a slightly increased risk overall. On the other hand it remains possible that the effect of snus in the Bolinder study was caused by some unmeasured (and therefore uncontrolled) confounding factor, with dietary habits and alcohol consumption being examples of baseline variables not measured in that study. This possibility is supported by a recent report of the effects of smokeless tobacco in the USA, based on analyses of the First National Health and Nutrition Examination Survey epidemiologic followup study (NHANES-1) data. This study had 96% follow up of the original 14 407 participants and 98% identification of death certificates for the 4604 decedents by 1992. Male smokeless tobacco users were found to have moderately increased risks of some disorders, but all of these excess risks disappeared when variables such as race and poverty were controlled for. For example, the crude hazard ratios for male smokeless users versus non-tobacco users were 1.5 and 2.1 for circulatory and respiratory diseases before adjustment, but after adjustment for confounders these hazard ratios became 1.0 and 0.9. One potentially serious flaw with this study is that pipe and cigar users were included in the “non-tobacco user” comparison group, seriously undermining confidence.
in their conclusion that US smokeless tobacco users have similar mortality outcomes to non-tobacco users. We cite this paper as an example of the changes in outcomes that can result from controlling for baseline variables, rather than as evidence of the safety of US smokeless tobacco.

Bolinder et al’s first study found snus users to be at excess risk of a number of respiratory symptoms. For example, the OR for ‘cough in the morning’ for snus users versus never tobacco users was 2.1 (95% CI 1.8 to 2.4), as compared with an OR of 7.9 for smokers versus never tobacco users. It is not easy to think of a plausible mechanistic pathway whereby exclusive snus use might cause respiratory symptoms. This study excluded all those who reported mixed use of snus and cigarettes or reported being an ex-snuser (n = 59,864 excluded). However, the increased respiratory symptoms suggest the possibility that some of those reporting exclusive snus use were actually occasional or ex-smokers. Passive smoke exposure is another possible confounding factor that could potentially contribute to these findings. This study was initially funded by a health insurance group with the purpose of examining factors affecting sick leave and disability pensions. Some participants may have under-reported their recent or ex-smoking due to their belief that it either was not related to their pension or that it was not serious enough to affect their future benefits.

In reviewing the evidence from a range of clinical and experimental studies, Benowitz concluded: “Overall, the epidemiologic and experimental data suggest that nicotine absorbed from smokeless tobacco, nicotine gum or transdermal nicotine is not a significant risk factor for accelerating coronary artery disease or causing acute cardiovascular events.” This conclusion is supported by a recent case-control study that examined risk factors for stroke among Swedish men. In multivariate analyses, controlling for other risk factors, smoking was related to increased risk of stroke (OR 1.74) whereas snus use was not (OR 0.87, 95% CI 0.41 to 1.83).

Given the inconsistencies in the results of these studies, it remains possible that snus users have a slightly increased cardiovascular risk as compared to never tobacco users, even after controlling for other confounding factors. However, all of the large studies of the effects of tobacco use on cardiovascular disease in Sweden are in agreement that “the use of smokeless tobacco (with snus being the most studied variant) involves a much lower risk for adverse cardiovascular effects than smoking does”.

Respiratory diseases
A Pubmed search did not identify any studies that specifically examined the effect of snus on respiratory diseases; similarly the IOM report did not address the effects of smokeless tobacco on respiratory illnesses. The reason for this is presumably that there is no plausible causal mechanism whereby smokeless tobacco could cause respiratory disease. A recent study of mortality in US smokeless users reported no increased risk of respiratory diseases in smokeless users. This contrasts heavily with the effect of continued smoking on chronic obstructive pulmonary disease, with 50% of elderly Swedish smokers developing the condition as compared with less than 20% of never smokers.

Diabetes
Bolinder found that smokers had significantly higher fasting blood glucose values than never tobacco users whereas snus users were not significantly different from never users. Eliasson and colleagues found that neither smoking nor snus use was associated with changed glucose tolerance or insulin concentrations. However, a more recent study by Persson found an increased risk of (asymptomatic) type 2 diabetes among both heavy smokers (25+ cigarettes per day) and heavy snus users (3+ cans per week), with significant odds ratios of 2.7 and 2.6, respectively, for these two groups as compared with non-tobacco users. It should be noted that this study specifically recruited men over 35 years old, 50% of whom had a family history of diabetes. The effects of snus on risks for diabetes are unclear and it may be that any effects are restricted to heavy users and/or those with a family history of diabetes.

Pregnancy
A Pubmed search did not identify any studies that had specifically examined the effects of snus use during pregnancy. However, given that animal studies have implicated nicotine as a cause of some of the widely known adverse effects of tobacco exposure during pregnancy (on both the health of the mother and healthy development of the fetus), it follows that snus use during pregnancy is likely to incur some of the risks associated with smoking during pregnancy. The preliminary results of one study (as yet unpublished) have been presented at a conference earlier this year. The study examined data from the Swedish Birth Register for women who delivered babies during 1999–2000. The study compared 789 snus users to 11,242 cigarette smokers and 11,500 women not using any tobacco. Smokers gave birth to babies weighing an average of 206 g (7.3 ounces) less than non-tobacco users. Snus users gave birth to babies weighing an average of 40 g (1.4 ounces) less than tobacco users. Snus users were more likely than both smokers and non-tobacco users to deliver prematurely (perhaps partially explaining the slightly lower birth weight), and were more likely than both smokers and non-tobacco users to suffer pre-eclampsia. Clearly, the full results of this study and additional studies on this topic are required before coming to conclusions, particularly given the possibilities for confounding variables to cause small sized effects. However, given the known risks of nicotine in pregnancy, and the preliminary results of this unpublished study, it seems likely that snus use can cause adverse health effects in pregnancy and should not be promoted as safe for use in pregnancy. It would be a particular cause for concern were there to be evidence of increased snus use among women of reproductive age, without an equal or greater reduction in smoking in that group. Given that smoking during early pregnancy in Sweden has already declined from 31% in 1983 to 12% in 2000, it could be argued that the potential for snus to have a “positive” impact on smoking in pregnancy has similarly shrunk. It would seem as though Swedish women are on a positive trend towards tobacco-free pregnancies without snus, and that it would be best kept that way.

The Pattern of Nicotine Use in Sweden over the Past Century
Total consumption of snus and cigarettes in Sweden have changed dramatically over the past century, with the most pronounced changes occurring over the past 20 years when cigarette consumption has reduced significantly and at the same time snus consumption has risen significantly. Figure 2 provides only a crude snapshot of overall sales, that hides sex-specific changes and changes in the size of the population. Adult (over 14) cigarette consumption went from approximately 0.2 kg/person in the 1920s to 1.1 kg/person in 1970 and then down to 0.6 kg/person at the end of the 20th century. Across the same time points Snus consumption fell from 1.4 kg/person to 0.4 kg/person and then has increased again to 0.9 kg/person by 2000. Figure 2 also serves as a reminder of some of the other factors affecting cigarette consumption; the large drop in cigarette sales in 1997 was probably related to an 18% price increase in January of that
the population in Sweden has increased by approximately 60% from 1916 to 2000. The same research group has recently published a prospective follow up study of over 70% of the participants in 1986, 1990, and 1994 who were successfully followed up in 1999 (n = 1651). This study found a continuing trend away from smoking among men in northern Sweden, moving to a smoking prevalence around 10% in those followed up in 1999. Of those men who were smokers (no snus use) in the 1986–94 surveys, 39% had quit smoking by 1999, one third of whom had switched to snus use. Among women who were smokers at the baseline surveys, 30% had quit by 1999, only 10% of whom had switched to snus. This study concluded: “use of snus played a major role in the decline of smoking rates amongst men in northern Sweden. The evolution from smoking to snus use occurred in the absence of a specific public health policy encouraging such a transition.”

It should be noted that fig 4 (consistent with fig 3) shows that while cigarette smoking has fallen dramatically among Swedish men, overall tobacco use has not. Some may view this as a failure of tobacco control (compared with some other countries). We view changes in tobacco caused disease as the decisive factor when evaluating the effects of tobacco control, and as discussed below, these changes have been very positive for Swedish men. It could also be argued that this reduction in male smoking may have occurred without snus. Here we regard the comparison with Swedish women (little snus use, smaller smoking reduction, smaller health improvement) and the characteristics of male ex-smokers (large proportion switching to snus when quitting smoking) as strongly suggestive of snus having a direct effect on the changes in male smoking and health.

WHAT HAVE BEEN THE NET EFFECTS OF SNUS ON PUBLIC HEALTH IN SWEDEN

The reductions in male smoking prevalence that have occurred in Sweden over the past 25 years have been the largest of any developed nation in the world. At the same time, Swedish men have also experienced a notable reduction in the incidence of the major smoking caused diseases. To exemplify this, fig 6 shows the pattern of changes in lung cancer incidence in Sweden and its near neighbour, Norway, from 1960 to 1999. Since the mid 1970s there has been a

A more detailed picture of the likely role of snus in smoking cessation in Sweden can be gained by examining the prevalence of ex-smoking among ever smokers by history of snus use and by sex in the Rodu study. As shown in fig 5, a higher proportion of male than female ever smokers had quit, and most of these had also used snus. The data from this study provide strong support for the role of snus in promoting smoking cessation among Swedish men (fig 5).

One recent study has specifically examined whether snus use appears to have directly influenced smoking rates in northern Sweden. This study used the dataset developed for the northern Sweden component of the WHO MONICA study. This involved collection of data from four representative population based surveys conducted in 1986, 1990, 1994, and 1999, including detailed questions on tobacco use among approximately 1500 adults at each time point. This study found stable prevalence of “all tobacco use” among men (at around 40%) over the 13 year period, but with male smoking decreasing from 23% to 14% and snus use increasing from 22% to 30%, as the proportion of snus using ex-smokers increased from 9% to 14%. In women, smoking prevalence remained stable from 1986 to 1994 at 27% then dropped to 22% in 1999 when snus use rose from 2% to 6% (fig 4).

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pronounced reduction in the incidence of lung cancer in Swedish men, as compared with Swedish women, and both men and women in Norway.

As of 2000, Sweden has a lower standardised rate of male lung cancer incidence than any comparable developed nation in the world. Sweden also has a low rate of oral cancer by international standards and this low rate has been falling over the past 20 years while snus use has increased. Sweden’s low rates of both lung and oral cancer are shown in fig 7.
Interestingly Swedish men have also had a significant improvement in cardiovascular health over the same period. For example, Rosen and colleagues studied trends in heart attacks in Sweden over the years 1987 to 1995 (amounting to 360,000 separate heart attacks in total). They found a 22% drop in heart attacks in men aged 30–64 years during that period, roughly double the decline among same aged women over the same period (fig 8).

It is noteworthy that these improvements in tobacco caused illnesses have occurred primarily in men, despite a stable consumption of tobacco among men during that time period. The main factor that has changed is that many Swedish men have switched from smoked tobacco to snus. Of course one cannot state with absolute certainty that if snus had not been available in Sweden that just as many men would have quit smoking either without assistance or perhaps by switching to nicotine replacement therapy. However, the pattern of sex differences in smoking cessation and snus use within Sweden, together with the between-country differences in smoking prevalence changes and health changes (comparing Sweden with other similar countries that have lower snus use, such as Norway), strongly suggests that a significant portion of the health improvement among Swedish men over the past 20 years has been due to a large proportion quitting smoking or never starting to smoke, but using snus instead.

**IS SNUS A ‘’GATEWAY’’ TO SMOKING OR A PATHWAY FROM SMOKING IN SWEDEN?**

It has been argued that smokeless tobacco could become a “gateway” product, hooking young people on nicotine from a cheaper and more easily concealed product, before they more easily move on to yet more addictive and harmful products such as cigarettes. For many reasons, the evidence from Sweden is not supportive of such a view. Firstly, if snus was acting to attract young people towards smoking one might expect the only country in Europe with a sizable snus market to have had the worst record for reducing smoking prevalence rather than the best. Secondly, when one examines the sex differences in tobacco use patterns, if snus was attracting young men towards smoking, one would expect the change in smoking prevalence to have been worse for men than for women, whereas it has been significantly better (that is, smoking prevalence has fallen more for men than for women).
than for women in Sweden). Looking only at daily smoking prevalence among 16 year olds in Sweden, this has remained remarkably stable at around 11% for boys and 16% for girls for the past 20 years. Again this is not consistent with the idea that snus is acting as a gateway to smoking among boys.

Thirdly, when one looks at the pattern of changes in tobacco use among Swedish men, the proportion of current smokers who are ex snus users is consistently smaller than the proportion of current snus users who are ex smokers (4% vs 14% of the adult male population in 1999, with only 3% current users of both snus and smoked tobacco, in the Rodu et al 2002 study).

A study recently presented by Ramstrom examined smoking status in Sweden by snus use, using data from a representative sample (n = 6700) of the Swedish population aged 16–79 years collected in 2001–2. In the sample of men, 15% were daily smokers and 20% were daily snus users (19% vs 2% among women). Among 2879 men, 468 (16%) were primary daily snus users (that is, they started daily snus without having previously started smoking). Twenty per cent of this group subsequently became daily smokers, compared with 45% of men who were not primary snus users. This suggests that snus use is protective against smoking rather than a gateway towards it. It is possible that this pattern of results could be caused by a combination of age and cohort effects. However, when we examined this issue we found lower rates of smoking onset among primary snus users in both older (born 1922–56) and younger (born 1957–1985) cohorts. Among those men who ever became daily smokers, 71% with a history of snus use quit smoking completely, compared with 54% of those with no snus history. Of those men who have quit smoking completely after having used snus as a cessation aid, 75% are currently daily snus users and 25% have quit snus use as well. Of all those men who quit smoking and mentioned the use of a single smoking cessation aid, 62% stated that they used snus as a cessation aid, compared with 38% who mentioned using nicotine replacement therapy. Again this is more consistent with snus being a pathway from smoking.

A study recently reported by Gilljam and Galanti also suggests that snus has primarily been a pathway from smoking among Swedish men. Their study consisted of a survey of approximately 1000 current smokers and 1000 ex-smokers (all men aged 25–55 years). Twenty-nine per cent of the ex-smokers had used snus to quit, and smoking cessation was significantly more likely among men who had used snus as compared with men who had not. Among those who used snus, 28% gave “health concerns” as their primary reason for snus use (for example, to help quit smoking or because it was less dangerous than smoking).

A recent study of tobacco use among young people in Sweden reported a larger prevalence of combined snus and cigarette use than reported in adult studies. This study was based on a 1998 survey targeting all 15–16 year old children in Stockholm (the capital city). Only 1.3% of girls reported snus use so this paper focused on boys (n = 6287): 14.3% were cigarette users, 5.7% were snus users, and 13.8% used both. Thus the majority (71%) of male snus users at that age were also smoking tobacco, although it should be noted that these percentages include people using these tobacco products less than daily. This study also highlighted the fact that at this age the young people had not yet established a stable profile of tobacco use, and that tobacco use, and particularly combined smoking and snus use, was linked with a number of other problem behaviours. Thus the likelihood of being a current snus user was several times higher among boys who reported having been drunk (OR 9.6), or used illicit drugs (OR 2.4) compared with those who did not. The authors of this study concluded: “smokeless tobacco use in adolescence does not substitute cigarette smoking and can be an indicator of a drug- and risk-seeking lifestyle.” It therefore seems unlikely that either form of tobacco use is a “gateway” to the other, but rather that both are markers of risk taking behaviour in adolescents. Smoking cessation aid, 62% stated that they used snus as a cessation aid, 75% are currently daily snus users and 20% use among Swedish men. Their study consisted of a survey of approximately 1000 current smokers and 1000 ex-smokers. Among 2879 men, 468 (16%) were primary daily snus users (that is, they started daily snus use at age 15 will subsequently quit smoking and will transfer to exclusive snus use. However, given the high frequency of combined snus and cigarette use in this study, it is clear that the pattern of transitional and combined use of different tobacco products among young people should continue to be closely monitored. One study in Finland examined the effect of tobacco use on prevalence and some negative consequences, including 12% of existing snuff users switching to smoking.

A recent report on smokeless tobacco use in the USA and Sweden concluded that at least 77% of US smokeless users and 83% of Swedish snus users appear to be “non-gateway” users in that their snuff use did not lead to smoking or their smoking preceded their snuff use.

Overall, the patterns of tobacco use in Sweden suggest that those who start using snus are less likely to become smokers, and that among people who become smokers, their chances of quitting smoking are higher if they use snus.

**WHAT ARE THE LESSONS FROM THE SWEDISH EXPERIENCE?**

None of the studies reviewed in this paper were randomised controlled trials and so no specific causal relations can be inferred from any individual study. Both within and outside Sweden, smoking is primarily influenced by factors other than availability of smokeless tobacco (for example, real price of cigarettes, health education, smoke-free air policies, industry marketing, etc). That having been said, we feel that the analysis of the change in patterns of tobacco use and health outcomes over time described here, including the comparison between countries and between sexes within Sweden, is suggestive of a positive rather than a negative net effect of snus use on tobacco smoking and hence on public health in Sweden. A key component of the evidence on this is the differential smoking quit rate between men and women. Most of the other important background factors affecting cigarette consumption (for example, price) would be expected to have similar effects on men and women and so...
they are unlikely to account for the sex differences in quitting. One important reason for male smoking cessation rates being higher is that many more Swedish men than women use snus.

We do not assume that these same benefits would automatically transfer to other countries, or even that they will remain constant in Sweden. Most countries of the world have very limited (or no) regulation of tobacco ingredients, or marketing. There may be little to stop a company from introducing a product that delivers significantly higher quantities of toxins than snus, and directing the marketing at young people, or even young non-smoking women of childbearing age. In such a scenario it is perfectly possible that snus or other smokeless products would have a negative effect on public health. However, in Sweden we have a concrete example in which availability of a less harmful tobacco product has probably worked to produce a net improvement in health in that country.

The main conclusions that can be drawn from the Swedish Experience are as follows:

- Implementation of stronger, evidence based regulation of tobacco products is necessary to avoid unintended public health consequences from both tobacco availability and tobacco bans in Sweden and worldwide.5 71

- Significant proportions of smokers are capable of transferring their nicotine dependence from an ultra-fast nicotine delivery product (a cigarette) to a medium rate nicotine delivery product (snus) so long as it delivers comparable amounts of nicotine, and so long as it is competitive on price, accessibility, and long term availability. This suggests that were a comparable non-tobacco pharmaceutical product (for example, a high dose nicotine gum) to become available and be able to compete on an even (or advantageous) basis, it may also have similar effects in helping a significant proportion of the smoking population transfer to a safer product. Unfortunately pharmaceutical nicotine replacement therapy (NRT) is currently regulated as part of a different regulatory system (along with medicines) that puts it at a competitive disadvantage as compared with tobacco products.6 They total elimination of most of the toxins in the nicotine delivery product (as in NRT) is clearly preferable to the marginal or unverified reductions in toxin delivery that are typically achieved by tobacco products.

- It appears to be extremely unlikely that nicotine is capable of stimulating cancer under normal use conditions. The media regularly issues scare stories about nicotine replacement products potentially causing cancer, usually stemming from media coverage of laboratory studies in animals or test tubes that interpret their findings as implying that nicotine may cause cancer in humans. In Sweden large numbers of snus users are consuming large quantities of nicotine, absorbed at a single part of the body (the mouth), along with significant concentrations of other carcinogens, for most of their adult life without evidence of increased cancer risk. The epidemiology of snus and cancer in Sweden does not support the view that nicotine itself is a risk factor for cancer.27–33

- Snus is certainly not harmless. It can cause reversible lesions in the mouth, it most likely causes harmful effects to the unborn fetus when used by a pregnant woman, and long term use may contribute to cardiovascular disease (although most of the available evidence suggests that cardiovascular risks are not increased by snus).

- Snus is clearly less harmful to the individual user than smoked tobacco, and also less harmful than the types of smokeless tobacco used in some other parts of the world, notably Sudan and India.29 The manufacturers of snus have voluntarily set fairly sensible toxicity standards for their product in order to reduce health risks as much as technologically possible. These or more thorough standards should now be applied across the industry and across countries. It could be argued that these same standards should also be applied to the other tobacco products (chewing tobacco, cigars, and pipe tobacco) that Swedish Match also produces and sells, and that a similar set of standards should apply to all nicotine delivery products.

In accepting that we now have smokeless tobacco products available that are less harmful than the dominant products (cigarettes), public health professionals and policymakers need to decide whether to focus our effort on restricting access to the most harmful products (smoked tobacco products), or focus much time, energy, and legislation on restricting access to the least harmful products, that under some circumstances can produce a net public health benefit.

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