Smoking reduction with oral nicotine inhalers: double blind, randomised clinical trial of efficacy and safety

Chris T Bolliger, Jean-Pierre Zellweger, Tobias Danielsson, Xandra van Biljon, Annik Robidou, Åke Westin, André P Perruchoud, Urbain Säwe

Abstract

Objectives To determine whether use of an oral nicotine inhaler can result in long term reduction in smoking and whether concomitant use of nicotine replacement and smoking is safe.

Design Double blind, randomised, placebo controlled trial. Four month trial with a two year follow up.

Setting Two university hospital pulmonary clinics in Switzerland.

Participants 400 healthy volunteers, recruited through newspaper advertisements, willing to reduce their smoking but unable or unwilling to stop smoking immediately.

Intervention Active or placebo inhaler as needed for up to 18 months, with participants encouraged to limit their smoking as much as possible.

Main outcome measures Number of cigarettes smoked per day from week six to end point. Decrease verified by a measurement of exhaled carbon monoxide at each time point compared with measurement at baseline.

Results At four months sustained reduction of smoking was achieved in 52 (26%) participants in the active group and 18 (9%) in the placebo group (P < 0.001; Fisher’s test). Corresponding figures after two years were 19 (9.5%) and 6 (3.0%) (P = 0.012).

Conclusion Nicotine inhalers effectively and safely achieved sustained reduction in smoking over 24 months. Reduction with or without nicotine substitution may be a feasible first step towards smoking cessation in people not able or not willing to stop abruptly.

Introduction

The best way to prevent the detrimental health consequences of cigarette smoking is to quit, and efforts to date have focused on this strategy. 1 Many smokers, however, find it impossible to quit, even with help, because of their dependence on nicotine, which is a highly addictive psychoactive drug. 2 Nicotine replacement therapy is an established pharmacological aid to help smokers quit and has consistently been shown almost to double the abstinence rate, irrespective of the level of additional interventions. 3 Increasing experience with trials on smoking cessation, however, has shown that successful abstinence is usually obtained in smokers with low to moderate nicotine dependence, whereas heavily dependent smokers have the highest relapse rates. 4 Unfortunately this latter group has the highest cigarette consumption and is therefore at the highest risk of developing disease related to tobacco consumption.

Given that few smokers are ready to quit at any time, plus the fact that many smokers try to quit several times before succeeding, new treatment approaches are clearly needed. One such strategy could be to reduce tobacco consumption substantially in smokers who are unwilling or unable to quit right away. For such smokers, sustained reduction might reduce the known health risks by reducing tobacco exposure and may also move them towards the ultimate goal of quitting. 5 6 In a preliminary study Fagerström et al showed that short term smoking reduction with nicotine replacement therapy over a period of five weeks was possible and that the combination of reduced smoking with nicotine replacement therapy was well tolerated. 7 The efficacy and safety of nicotine replacement therapy in achieving sustained smoking reduction, however, has not yet been assessed. Another important issue is whether smoking reduction can increase motivation to quit in recalcitrant smokers.

Smoking cessation is no longer regarded as a dichotomous process (cessation or not) but rather as a continuum that entails several stages, as described by DiClemente and Prochaska. 8 There is empirical evidence to suggest that reduced smoking, also referred to as controlled smoking or harm reduction, is a therapeutic option for those smokers unable or unwilling to quit. Glasgow et al 9 and Hughes et al 10 found that smokers randomised to such an intervention were no less likely, and possibly even more likely, to quit smoking in the long term compared with smokers randomised to more conventional interventions. While not the first treatment of choice, reduced smoking might be considered for recalcitrant smokers unwilling to repeat traditional cessation attempts.
The oral nicotine inhaler, which is one of the newer nicotine replacement products, has been shown to be an effective aid for smoking cessation. Use of a nicotine inhaler by smokers unwilling or unable to stop smoking completely might be a good approach to reducing cigarette consumption as the inhaler imitates some aspects of cigarette smoking and contains nicotine. We tested the efficacy and safety of the nicotine inhaler in achieving sustained smoking reduction.

Methods

Study design

Smokers were recruited into this two centre, double blind, placebo controlled, randomised clinical trial through newspaper advertisements that asked for healthy smokers who were unwilling or unable to quit but were interested in reducing their smoking. All participants were given information about possible ways to achieve this goal. Smoking cessation was recommended as the ultimate goal throughout the study.

Participants

Participants in the trial had to be at least 18 years of age, smoke 15 or more cigarettes a day, have a carbon monoxide concentration in exhaled air ≥ 10 ppm, have smoked regularly for three or more years, have failed at least one serious attempt to quit within the past 12 months, want to reduce smoking as much as possible with the help of the nicotine inhaler, be prepared to adhere to the protocol, and be willing to provide informed consent. Exclusion criteria were current use of nicotine replacement therapy or any other behavioural or pharmacological smoking cessation or reduction programme, use of other nicotine-containing products, or any condition that might interfere with the study.

The ethics committees of the universities of Basle and Lausanne approved the study.

Treatment

Independent pharmacists dispensed either active or placebo inhalers according to a computer generated randomisation list. All smokers received information about the general implications of smoking and its effects on health. Participants were asked to reduce the number of cigarettes smoked daily as much as possible, and an initial reduction of 50% was suggested.

The active treatment comprised nicotine replacement through an inhalation device (Nicorette Inhaler, Pharmacia and Upjohn). The inhaler consists of a plastic mouthpiece into which a disposable cartridge containing 10 mg nicotine and 1 mg menthol is inserted. At room temperature the total available nicotine content is 4-5 mg per cartridge. The inhaler delivers about 13 µg of nicotine per puff (average puff volume of 50 ml), which means that about 80 puffs are required to obtain 1 mg nicotine. The placebo inhalers were identical in appearance and contained only menthol. Both treatment groups were allowed to use the inhalers as needed, with the recommendation to use between six and 12 cartridges over 24 hours. Participants were encouraged to decrease use of the inhaler after four months but were permitted to continue treatment for 18 of the 24 months in the study.

Assessment

After the initial telephone screening and baseline assessment participants were reassessed at the clinic after one, two, three, and six weeks and three, four, six, 12, 18, and 24 months. Counselling on smoking reduction was provided at each visit.

Admission criteria and demography, including the Fagerström test for nicotine dependence, reasons for reducing smoking, and medical and smoking history were assessed at baseline. At all key visits (baseline and months four, 12, and 24) we measured expired carbon monoxide concentrations, symptoms of withdrawal, smoking status, respiratory function, blood pressure, pulse, weight, plasma cotinine concentration, haematological variables, and concentrations of blood lipids and fibrinogen. We also assessed smoking status, intention to quit, compliance, concomitant medications, adverse events, and quality of life (with the SF-36 questionnaire). Reported adverse events and plasma cotinine concentrations served as a basis for the safety analysis of concomitant smoking and nicotine replacement therapy.

Measures of outcome

The primary efficacy measure (success) was defined as self reported reduction of daily cigarette smoking by at least 50% compared with baseline from week six to month four, the duration for which the study was powered. This reduction was verified by decreased carbon monoxide concentrations at week six and months three and four. Results up to 24 months are presented in this paper. Smoking cessation was defined as not smoking from week six and a carbon monoxide concentration < 10 ppm at all subsequent visits. Smoking reduction and cessation are also presented with verification of carbon monoxide concentrations at each time point (point prevalence).

Statistical analysis

To ensure an adequate number of participants we recruited 200 smokers into each group, giving a total of 400 participants. The primary analysis was an intention to treat analysis including all participants who were randomised and received medication. As in other studies of smoking cessation studies participants who dropped out were regarded as treatment failures. All statistical methods were two tailed, and P values of <0.05 were considered significant. Odds ratios together with corresponding confidence intervals were calculated when applicable, and Fisher’s test was used for categorical and binary variables. Confidence intervals of means were calculated for continuous variables when applicable, and differences within participants were tested for with Wilcoxon’s signed rank sum test.

Results

Baseline characteristics and rates of follow up

Table 1 shows the baseline characteristics of all enrolled participants. The only significant difference was that there were more women in the active...
P values (Wilcoxon’s rank sum test) for difference between intervention and placebo: *P=0.004; †P<0.001; ‡P=0.003; §P=0.01; ¶P=0.03; **P=0.02.

BMJ

treatment group. At each follow up visit there were more participants in the active treatment group than in the placebo group (figure).

Treatment compliance
Inhaler use decreased over time, as expected. Of participants present at week six, 222/368 (60%) used the inhaler every day. Corresponding figures after four, 12, and 18 months were 146/318 (46%), 39/331 (12%), and 30/289 (10%), respectively. Participants in the active treatment group used an average of 4.5 cartridges a day after two weeks and 2.6 a day after 18 months; they reduced their cigarette intake significantly more than participants in the placebo group and by eight in the placebo group and nine in the active treatment group, with corresponding figures for palpitation (1.13 to 15.6) and coughing (13 v 4; 1.1 to 10.6). The total numbers of adverse events relating to symptoms usually associated with nicotine (nausea, vomiting, and palpitation) were significantly more common in the active treatment group than the placebo group; throat irritation (14 v 4; 95% confidence interval for odds ratio 1.13 to 15.6) and coughing (13 v 4; 1.1 to 10.6). The sustained rates of complete abstinence were low in both groups, with no significant differences between groups. Four participants in the active treatment group and one in the placebo group were not smoking any cigarettes at four, 12, and 24 months. The point prevalence cessation rates, however, increased over time, and 38 participants were not smoking after 24 months (table 3).

Safety
Adverse events were reported by 114 participants in the placebo group (193 adverse events) and 113 participants in the active treatment group (227 adverse events). Fifty three serious events occurred, none of which was related to treatment. Two local symptoms were significantly more common in the active treatment group than the placebo group: throat irritation (14 v 4; 95% confidence interval for odds ratio 1.13 to 15.6) and coughing (13 v 4; 1.1 to 10.6). The total numbers of adverse events relating to symptoms usually associated with nicotine (nausea, vomiting, and palpitation) were evenly distributed between the groups; nausea or nausea and vomiting was reported by eight in the placebo group and nine in the active treatment group, with corresponding figures for palpitation of two and one, respectively.

Table 1 Baseline demographic characteristics, including smoking status and history. Values are expressed as means (SD); range

<table>
<thead>
<tr>
<th>Placebo (n=200)</th>
<th>Active (n=200)</th>
</tr>
</thead>
<tbody>
<tr>
<td>No of men</td>
<td>104</td>
</tr>
<tr>
<td>Age (years)</td>
<td>45.8 (10.5); 22-77</td>
</tr>
<tr>
<td>Weight (kg)</td>
<td>81.5 (12.3); 57-121</td>
</tr>
<tr>
<td>Age when started smoking (years)</td>
<td>17.1 (2.7); 11-35</td>
</tr>
<tr>
<td>No of cigarettes (per day)</td>
<td>39.3 (12.1); 15-70</td>
</tr>
<tr>
<td>CO as % of baseline value</td>
<td>27.1 (11.1); 10-61</td>
</tr>
<tr>
<td>FTND score</td>
<td>5.6 (2.0); 1-10</td>
</tr>
</tbody>
</table>

CO=carbon monoxide; FTND=Fagerström test for nicotine dependence.

Efficacy
Table 3 shows that the success rates from week six onwards were significantly higher for the active group at four, 12, and 24 months. The point prevalence reduction rates were also higher in the active treatment group, but the difference was significant only at four months (P < 0.001), with only a trend towards significance at 12 months (P = 0.085) and 24 months (P = 0.357).

The sustained rates of complete abstinence were low in both groups, with no significant differences between groups. Four participants in the active treatment group and one in the placebo group were not smoking any cigarettes at four, 12, and 24 months. The point prevalence cessation rates, however, increased over time, and 38 participants were not smoking after 24 months (table 3).

Safety
Adverse events were reported by 114 participants in the placebo group (193 adverse events) and 113 participants in the active treatment group (227 adverse events). Fifty three serious events occurred, none of which was related to treatment. Two local symptoms were significantly more common in the active treatment group than the placebo group: throat irritation (14 v 4; 1.13 to 15.6) and coughing (13 v 4; 1.1 to 10.6). The total numbers of adverse events relating to symptoms usually associated with nicotine (nausea, vomiting, and palpitation) were evenly distributed between the groups; nausea or nausea and vomiting was reported by eight in the placebo group and nine in the active treatment group, with corresponding figures for palpitation of two and one, respectively.

Table 2 Number of inhalers used, reduction in number of cigarettes smoked, and exhaled carbon monoxide concentration in participants using inhaler every day; active and placebo treatment groups. Values are expressed as means (SD); range

<table>
<thead>
<tr>
<th>Time point</th>
<th>Intervention group</th>
<th>Placebo group</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>No of inhalers/day</td>
<td>No of participants</td>
</tr>
<tr>
<td>1 week</td>
<td>169</td>
<td>4.3 (2.1); 1-12</td>
</tr>
<tr>
<td>2 weeks</td>
<td>168</td>
<td>4.5 (2.0); 1-10</td>
</tr>
<tr>
<td>6 weeks</td>
<td>117</td>
<td>4.3 (2.1); 1-12</td>
</tr>
<tr>
<td>4 months</td>
<td>84</td>
<td>3.9 (2.0); 1-10</td>
</tr>
<tr>
<td>12 months</td>
<td>27</td>
<td>3.5 (1.9); 1-7</td>
</tr>
<tr>
<td>18 months</td>
<td>22</td>
<td>2.6 (1.7); 0-6</td>
</tr>
</tbody>
</table>

P values (Wilcoxon’s rank sum test) for difference between intervention and placebo: *P<0.004; †P<0.001; ‡P=0.003; §P=0.01; ¶P=0.03; **P=0.02.
Table 3 Efficacy results measured as sustained and point prevalence reductions in smoking and point prevalence abstinence rates according to treatment with oral nicotine inhaler (active treatment) or placebo

<table>
<thead>
<tr>
<th>Definition</th>
<th>Time point (months)</th>
<th>No (%) with active treatment</th>
<th>No (%) with placebo</th>
<th>Odds ratio (95% CI)</th>
<th>P value (Fisher’s test)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sustained</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Reduction*</td>
<td>4</td>
<td>52 (26.0)</td>
<td>18 (9.0)</td>
<td>3.55 (2.04 to 6.19)</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td></td>
<td>12</td>
<td>26 (13.0)</td>
<td>8 (4.0)</td>
<td>3.59 (1.65 to 7.80)</td>
<td>0.002</td>
</tr>
<tr>
<td></td>
<td>24</td>
<td>19 (9.5)</td>
<td>6 (3.0)</td>
<td>3.39 (1.38 to 8.29)</td>
<td>0.012</td>
</tr>
<tr>
<td>Point prevalence</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Reduction†</td>
<td>4</td>
<td>83 (41.5)</td>
<td>44 (22.0)</td>
<td>2.52 (1.63 to 3.87)</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td></td>
<td>12</td>
<td>59 (29.5)</td>
<td>43 (21.5)</td>
<td>1.53 (0.97 to 2.40)</td>
<td>0.085</td>
</tr>
<tr>
<td></td>
<td>24</td>
<td>55 (27.5)</td>
<td>46 (23.0)</td>
<td>1.27 (0.81 to 2.00)</td>
<td>0.357</td>
</tr>
<tr>
<td>Abstinence‡</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Reduction*</td>
<td>4</td>
<td>13 (6.5)</td>
<td>12 (6.0)</td>
<td>1.36 (0.63 to 2.95)</td>
<td>0.557</td>
</tr>
<tr>
<td></td>
<td>12</td>
<td>16 (8.0)</td>
<td>15 (7.5)</td>
<td>1.26 (0.65 to 2.47)</td>
<td>0.609</td>
</tr>
<tr>
<td></td>
<td>24</td>
<td>21 (10.5)</td>
<td>17 (8.5)</td>
<td>1.26 (0.65 to 2.47)</td>
<td>0.609</td>
</tr>
</tbody>
</table>

*Sustained reduction in number of cigarettes smoked daily by at least 50% from week 6, verified by decreased carbon monoxide concentrations compared with baseline.
†Point prevalence reduction of cigarettes smoked daily by at least 50% at months 4, 12, and 24, verified by decreased carbon monoxide concentrations compared with baseline.
‡No cigarettes smoked, verified by carbon monoxide concentrations <10 ppm at months 4, 12, and 24.

Intention to quit
With regard to changes in participants’ interest to quit smoking no difference could be detected between reducers and non-reducers or active and placebo treatment.

Discussion
This trial confirmed that it is possible to achieve a sustained reduction in cigarette smoking in people unable or not willing to quit. Although the overall success rates were relatively small, active treatment with the nicotine inhaler was more effective in obtaining this reduction than placebo over the entire period of two years. The combination of smoking and using a nicotine inhaler was well tolerated.

Except for the fact that more women were randomised to active (114) than placebo (96) treatment, all baseline parameters in our study group were comparable to the cohorts found in smoking cessation studies. One notable difference, implied by the recruitment criteria, was that our participants were not prepared to stop smoking immediately, although over two thirds said they wanted to stop eventually, whereas all participants were motivated to stop in cessation studies.

As the study of smoking reduction is relatively new, there is no consensus regarding what level of reduction should be chosen to define success. We therefore arbitrarily selected a reduction of >50% in daily cigarette smoking, verified by a reduction in exhaled carbon monoxide concentrations compared with baseline. Smoking cessation has traditionally been defined as sustained abstinence combined with an expired carbon monoxide concentration of < 10 ppm.

Safety
No serious or important adverse events related to treatment occurred during the study period. Symptoms of possible nicotine overdose were evenly distributed between treatment groups. Throat irritation and coughing were the only symptoms significantly more common in the active treatment group, and this was expected on the basis of previous experience with the nicotine inhaler in smoking cessation studies.

Our finding, that the combination of reduced smoking and use of the nicotine inhaler was well tolerated, should allay the concern that concomitant use may lead to nicotine intoxication.

Concept of smoking reduction
One argument against smoking reduction is that encouragement of this strategy may give smokers an easy option and undermine efforts to stop smoking completely, but the converse may also apply. Smoking reduction may promote smoking cessation by allowing smokers to take control of their smoking gradually, and this seemed to be the case in one trial. Other supportive studies in which nicotine treatments were used to reduce smoking in smokers who were trying to quit suggest that nicotine treatments do suppress smoking behaviour and would thus benefit smokers trying to reduce smoking. For example, in the lung health study 60% of the participants reduced rather than stopped smoking, and 39% of these reduced their smoking by at least 50%. Similarly, a recent analysis of the 1410 people who smoked both at baseline and at two year follow up in the community intervention trial for smoking cessation (COMMIT), which involved participants in 22 US cities, reported that at two years 17% had decreased their smoking by 5-25%, 15% by 24-49%, and 8% by at least 50%. Reduced smoking at two years neither promoted nor undermined cessation at a later date. In our opinion, the fact that in our

What is already known on this topic
Smoking cessation has been the sole goal in studies on the treatment of tobacco dependence

Many smokers find it impossible to stop smoking without help because of their dependence on cigarettes

What this study adds
Smokers who are unwilling or unable to stop smoking abruptly can use nicotine replacement therapy to reduce the amount they smoke

Reduction in smoking was safely achieved with the aid of a nicotine inhaler
Effect of restrictions on smoking at home, at school, and in public places on teenage smoking: cross sectional study

Melanie A Wakefield, Frank J Chaloupka, Nancy J Kaufman, C Tracy Orleans, Dianne C Barker, Erin E Ruel

Abstract

Objective To determine the relation between extent of restrictions on smoking at home, at school, and in public places and smoking uptake and smoking prevalence among school students.

Design Cross sectional survey with merged records of extent of restrictions on smoking in public places.

Setting United States.

Participants 17 287 high school students.

Main outcome measures Five point scale of smoking uptake; 30 day smoking prevalence.

Results More restrictive arrangements on smoking at home were associated with a greater likelihood of being in an earlier stage of smoking uptake (P < 0.05) and lower 30 day prevalence (0.91 (0.83 to 0.99), P = 0.03). School smoking bans were related to a greater likelihood of being in an earlier stage of smoking uptake (0.89 (0.85 to 0.90), P < 0.05) and lower prevalence (0.86 (0.77 to 0.94), P < 0.001) only when the ban was strongly enforced, as measured by instances when teenagers perceived that most or all students obeyed the rule.

Conclusions These findings suggest that restrictions on smoking at home, more extensive bans on smoking in public places, and enforced bans on smoking at school may reduce teenage smoking.

Introduction

Restrictions on smoking at work and home are associated in adults with reduced daily smoking rate and increased cessation.1,2 As these types of smoking restrictions become more pervasive,3,4 smoking is being in an earlier stage of smoking uptake (P < 0.05) and lower 30 day prevalence (0.91 (0.83 to 0.99), P = 0.03). School smoking bans were related to a greater likelihood of being in an earlier stage of smoking uptake (0.89 (0.85 to 0.90), P < 0.05) and lower prevalence (0.86 (0.77 to 0.94), P < 0.001) only when the ban was strongly enforced, as measured by instances when teenagers perceived that most or all students obeyed the rule.