effect of obesity was negligible. It is surprising that the earlier reports from the Framingham Study* (based on two-year incidence estimates) as well as the 10-year results from the Seven Countries Study† failed to show a convincing relation between obesity and coronary heart disease. A recent report from Framingham,‡ however, based on 26-year incidence data, identified obesity as an important and independent risk factor (especially in women).

In the Whitehall Study mortality from causes other than coronary heart disease and cancer was increased by about 50% in the lowest quintile of body mass index, relative to the middle quintiles. This unexplained finding merits further study.

We are grateful for the co-operation of the Civil Service and their medical advisory service. The work was supported in part by a grant from the British Heart Foundation.

References


(Received 24 June 1982)

Randomised controlled trial of nicotine chewing-gum

M J JARVIS, MARTIN RAW, M A H RUSSELL, C FEYERABEND

Abstract

The effectiveness of 2 mg nicotine chewing-gum as an aid to stopping smoking was compared with a placebo containing 1 mg nicotine, but unbuffered, in a double-blind randomised trial. Of 58 subjects given the active gum, 27 (47%) were not smoking at one-year follow-up compared with 12 (21%) of the 58 subjects treated with placebo (p < 0.05). By the most stringent criterion of outcome, 18 (31%) subjects in the active treatment group and eight (14%) in the placebo group had not smoked at all from the start of treatment to follow-up at one year (p < 0.05).

Subjects receiving the active gum experienced less severe withdrawal symptoms and rated their gum as more helpful than did the placebo group. Minor side effects were common but only gastric symptoms were more frequent with the active gum. Subjects receiving active gum used it for longer than those receiving placebo but most stopped using it within six months and only four (7%) developed longer-term dependence. The number of gums used daily correlated significantly with pretreatment blood nicotine concentrations in the active
treatment group and with pretreatment cigarette consumption in the placebo group. A lower pretreatment blood nicotine value was the best predictor of success at one year (p < 0.001) but there was no significant relation to cigarette consumption, sex, and social class.

The results clearly confirm the usefulness of nicotine chewing-gum as an aid to stopping smoking and imply a definite role for nicotine in cigarette dependence and withdrawal. Successful use of the gum requires careful attention to subjects' expectations and clear instructions on how to use it.

Introduction

Many smokers give up smoking without any special help or treatment, but others have great difficulty and fail many times. The first smoking-cessation clinic was started in Stockholm in 1955. Since then there has been an intensive search for an effective treatment for dependent smokers. Simple support and encouragement, given individually or in groups, has a success rate of around 15-25% abstinent at one-year follow-up. Numerous other methods have been tried, including tranquillisers, lobeline, electric aversion therapy, rapid smoking, hypnosis, and, more recently, acupuncture. None of these methods, however, has been found to have a specific effect over and above the attention-placebo element inherent in any treatment.

We have reported encouraging results from the use of nicotine chewing-gum (Nicorette) in our smokers' clinic. In a comparative study the success rate of smokers who received the gum was 38% abstinent at one year of follow-up compared with only 14% of those who had had intensive psychological treatment. We now report the results of a randomised double-blind placebo-controlled trial of the gum with one-year follow-up and biochemical validation of reported abstinence from smoking.

Addiction Research Unit, Institute of Psychiatry, Maudsley Hospital, London SE5
M J JARVIS, BSc, MPhil, clinical psychologist
MARTIN RAW, BA, MPhil, clinical psychologist (now at St George's Hospital Medical School, London)
M A H RUSSELL, FRCP, FRCPsyCh, senior lecturer and consultant psychiatrist
Poisons Unit, New Cross Hospital, London SE14
C FEYERABEND, BSc, senior biochemist
Subjects and methods

The active gum contained 2 mg nicotine and was identical with the commercially available preparation. The placebo gum contained 1 mg nicotine and its biological availability was reduced by the lack of an alkaline buffer to promote absorption through the buccal mucosa.

The placebo was designed in this way to mimic the nicotine taste of the active gum without providing an effective pharmacological dose. In pretrial tests the placebo gum did produce appreciable plasma nicotine concentrations with excessive chewing (117 nmol/l (19 ng/ml) when chewed half-hourly for four hours). This suggests that anyone chewing 20 or more placebo gums a day would get a pharmacologically effective dose. The study could therefore be described as a dose-response study. Whatever terminology is preferred, there is no doubt that the “placebo” provided a fairly stringent test of the pharmacological role of nicotine in the efficacy of the active gum. Both active and placebo gums were packed identically and labelled as 2 mg Nicorette.

A total of 116 subjects were entered into the trial. All were cigarette smokers who attended the Maudsley Hospital smokers’ clinic for treatment between November 1979 and October 1980 and who agreed to participate in a trial of nicotine chewing gum. They were treated in groups of about 10, taken in order from the waiting list, each group being allocated at random to receive either the active or placebo gum. There were 12 groups in all, with each of two therapists treating three active-treatment and three placebo groups. Fifty-eight subjects were assigned to the active gum and 58 to the placebo. Therapists and subjects were blind to the allocation.

Before treatment all subjects completed questionnaires about their smoking and attended an assessment interview at which a blood sample was taken two minutes after finishing a cigarette to determine their baseline smoking values of plasma nicotine.10 Expired-air carbon monoxide11 or carboxyhaemoglobin12 values were also measured. Subjects assigned to the two treatments were well matched in demographic characteristics and pretreatment smoking habits, with the exception that those assigned to the active gum tended to be the heavier smokers (t = 2.1; p < 0.05; table I).

### TABLE I—Comparison of demographic and pretreatment smoking characteristics of subjects receiving active gum and placebo

<table>
<thead>
<tr>
<th></th>
<th>Active gum (n = 58)</th>
<th>Placebo gum (n = 58)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Mean age (years)</td>
<td>41.6</td>
<td>40.8</td>
</tr>
<tr>
<td>No (%) of men in group</td>
<td>58 (50.0)</td>
<td>56 (50.0)</td>
</tr>
<tr>
<td>No (%) of subjects in classes I and II</td>
<td>33 (56.9)</td>
<td>36 (62.0)</td>
</tr>
<tr>
<td>Mean No of cigarettes smoked daily</td>
<td>30.9</td>
<td>26.5</td>
</tr>
<tr>
<td>Mean plasma nicotine concentration (nmol/l)</td>
<td>199</td>
<td>247</td>
</tr>
<tr>
<td>Mean carboxyhaemoglobin value (%)</td>
<td>7.2</td>
<td>7.0</td>
</tr>
</tbody>
</table>

*p < 0.05. (No other differences statistically significant.)

Conversion: SI to traditional units—Nicotine: 1 nmol/l = 0.16 mg/ml.

All subjects were given the same instructions about the gum. They were told that it contained nicotine which would be absorbed through the lining of the mouth as it was chewed, and that it would reduce the craving for cigarettes and help to relieve other withdrawal symptoms. They were warned that it might take a few days to get used to the taste, and that they should not expect it to be a miracle cure that removed the necessity for them to work very hard at stopping. They were encouraged to stop smoking completely on the first day of treatment and told to chew a piece of gum whenever the desire to smoke was particularly strong. No restrictions were placed on the number of gums to be chewed each day. It was recommended that they should use the gum for at least three months before attempting to do without it.

Group meetings were held weekly for one hour for the first six weeks. Attendance was similar in those receiving the active and placebo gums, with 28 and 29 subjects respectively attending three or more meetings. Thereafter subjects attended as needed to collect gum and for follow-up at three months, six months, and one year. At the one year follow-up claims of abstinence were validated by measurement of expired air carbon monoxide concentrations. There were no cases of deception. In six subjects assigned to active gum and six assigned to placebo biochemical validation was not done but confirmatory reports were obtained from friends or relatives.

At each attendance subjects completed ratings of withdrawal symptoms, acceptability of their gum, and a check list of potential side effects. They were also given cards to record daily consumption of gum and cigarettes.

Statistical analyses were based on the binomial test, χ² test, and t test. Two-sided tests were used except for the analysis of outcome, where the one-sided hypothesis that the active gum was superior to the placebo was tested.

Results

Outcome

The figure shows the success rates of the active and placebo gums. A significant advantage for the active gum emerged as early as two weeks after the start of treatment, when 39 subjects assigned to the active gum were not smoking as compared with only 26 assigned to the placebo (Z = 2.24; p < 0.025). At one year 27 (47%) of those given the active gum were abstinent as compared with 12 (21%) of those given the placebo (see table II).

The results were unusual in that there were more subjects abstinent in the active-treatment group at one year than at six months, indicating that several subjects who had relapsed at the earlier point stopped smoking again before one year. Table II therefore gives the result of applying more stringent criteria of success to supplement the conventional analysis. Whichever criterion of success was applied the active gum was clearly more effective than the placebo.

### TABLE II—Success rates at one-year follow-up according to three different criteria. (Figures are percentages (%)) of successful subjects

<table>
<thead>
<tr>
<th>Criterion of success</th>
<th>Active gum (n = 58)</th>
<th>Placebo gum (n = 58)</th>
<th>Binomial test (Z)</th>
<th>t (10 df)*</th>
</tr>
</thead>
<tbody>
<tr>
<td>Abstinent at one year</td>
<td>27 (47)</td>
<td>12 (21)</td>
<td>2.75; p &lt; 0.01</td>
<td>2.31; p &lt; 0.025</td>
</tr>
<tr>
<td>Abstinent at end of initial treatment and at six months</td>
<td>22 (38)</td>
<td>9 (16)</td>
<td>2.52; p &lt; 0.01</td>
<td>2.38; p &lt; 0.025</td>
</tr>
<tr>
<td>No smoking at all from first week of treatment to one year of follow-up</td>
<td>18 (31)</td>
<td>8 (14)</td>
<td>2.00; p &lt; 0.025</td>
<td>1.92; p &lt; 0.05</td>
</tr>
</tbody>
</table>

Note: One subject receiving active gum was lost to follow-up and classified as a failure despite being abstinent from start of treatment to last contact at five months.

* Since treatment was randomised over groups, not individual subjects, group effects could exist which would invalidate use of binomial test. The more conservative t test allows for this possibility. t value derived from analysis of variance of angular transformation of group success rates.

Of those abstinent at the one-year follow-up, six in the active-treatment group and two in the placebo group were still using the gum. When for the sake of extreme stringency these subjects were excluded from the one-year abstainers, the active gum retained its clear-cut advantage over placebo with 21 (36%) abstinent at one year compared with 10 (17%) of the placebo group (Z = 2.10; p < 0.05). At a mean of 22 months after starting treatment, five of the six subjects given the active gum were still abstinent and three were still using the gum. Both groups given the placebo had relapsed to smoking.

The figure shows that the main effect of the active gum was to enable more subjects to stop smoking initially. There were also fewer relapses in the active-treatment group, although this failed to reach statistical significance. In the active-treatment group 12 of the 39 subjects abstinent at two weeks had relapsed to smoking at one year, compared with 14 of the 26 in the placebo group (χ²=2.57; df=1; NS).

### QUANTITY AND DURATION OF GUM USE

Use of the active gum was greater than the placebo at every stage of treatment and follow-up both in terms of the proportion of subjects who were using it and the number of pieces they were chewing a day. The differences, however, were statistically significant only at three and six months (table III). Of the seven subjects using active gum and three using placebo gum at the one-year follow-up, only four, all in the active group, had used it continuously throughout the year. The difference in point prevalence on the active gum was therefore 7%, and there were no such cases in the placebo group.

Among those who were not smoking at one month there was a
TABLE III—Comparison of MEDICAL assigned and asked active-treatment and placebo groups statistically significant.

<table>
<thead>
<tr>
<th>Time from start of treatment</th>
<th>Number of subjects using gum</th>
<th>Mean No of gums/day</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Active gum (n = 58)</td>
<td>Placebo gum (n = 58)</td>
</tr>
<tr>
<td>1 week</td>
<td>47 (81)</td>
<td>39 (67)</td>
</tr>
<tr>
<td>2 weeks</td>
<td>40 (66)</td>
<td>36 (62)</td>
</tr>
<tr>
<td>3 months</td>
<td>33 (57)</td>
<td>29 (50)</td>
</tr>
<tr>
<td>6 months</td>
<td>24 (41)</td>
<td>18 (31)</td>
</tr>
<tr>
<td>9 months</td>
<td>12 (21)</td>
<td>10 (17)</td>
</tr>
<tr>
<td>1 year</td>
<td>7 (12)</td>
<td>5 (9)</td>
</tr>
</tbody>
</table>

*p < 0.005. (No other differences between active and placebo groups statistically significant.)
† Number of gums daily are averages for those who were using it.

SIDE EFFECTS

Mild and transient symptoms were common with both the active and placebo gums (table IV). They were mainly non-specific effects of heavy chewing. Only in the case of hiccups, indigestion or stomach ache, and feeling sick was the incidence substantially higher and therefore interpretable as specific side effects of the active gum. In most cases these symptoms were mild and transient. In no cases were side effects cited as a cause for discontinuing the gum.

ACCEPTABILITY OF GUM

Ten subjects in the active-treatment group and 13 in the placebo group did not attend the first meeting after the start of treatment. We do not know whether this was due to dislike of the gum, disappointment with its efficacy, or other reasons. Among the remaining the active group rated the gum as significantly stronger (p < 0.05) and more helpful (p < 0.05) than the placebo group. Both groups rated the gum as moderately satisfying and slightly unpleasant tasting, and these ratings did not differ significantly between the two groups when averaged over the first six weeks of treatment. In the first week, however, the active group rated their gum as tasting more unpleasant than did the placebo group (p < 0.05), a difference which disappeared by the second week. By the second week the active group found the gum more satisfying than did the placebo group (p < 0.05).

WITHDRAWAL SYMPTOMS

Since the pattern of attendance at weekly treatment sessions did not differ between the active-treatment and placebo groups, each subject’s ratings of withdrawal symptoms over the first six weeks of treatment were averaged. Subjects who received the active gum experienced less severe withdrawal symptoms than the placebo group. They felt less irritable (p < 0.05), less sleepy (p < 0.01), and less hungry (p < 0.05). Differences between the groups in ratings of tensesness, feeling miserable, and difficulty doing without cigarettes also favoured the active gum but did not reach statistical significance. The active group, however, also rated themselves as less alert but this difference was not significant.

CHARACTERISTICS OF SUBJECTS AND OUTCOME

The relation of the variables listed in table I to outcome at the one-year follow-up was similar in the active-treatment and placebo groups, so that the two groups were combined to provide 39 successes and 77 failures. The successes tended to be older (mean ages 43 ± 38 years; p < 0.05) and to have lower pretreatment carboxyhaemoglobin values (mean 6.0% ± 7.6%; p < 0.01) and lower plasma nicotine concentrations (mean 172.0 nmol/l (27.9 ng/ml) v 229.3 nmol/l (37.2 ng/ml); p < 0.001). Their mean pretreatment daily cigarette consumption was also lower (26.3 ± 23.9) but this difference was not statistically significant. Social class and sex had no relation to outcome. The success rate among the men was 37% (19/52) compared with 31% (20/64) for the women (χ² = 0.36; NS).

Discussion

Treatment with the active nicotine chewing-gum achieved results that were substantially better than with the placebo. Success rates, based on three different criteria, were more than double those obtained with placebo. By the conventional criterion of smoking status at follow-up 47% of the subjects in the active-treatment group were abstinent at one year compared with 21% of the subjects given placebo. By the criterion of abstinence at the end of treatment and at six months and one year the success rates were 38% and 16% respectively, and when based on lapse-free abstinence throughout the year from the first week of treatment to the one-year follow-up
they were 31% and 14% respectively. The differences were statistically significant in all three cases and were of a similar order to our earlier study, which obtained a 38% success rate (conventionally defined) with nicotine chewing-gum compared with 14% for intensive psychological treatments. In both studies the active gum alone gave results that were well above the range (about 15-25%) reported for all other methods.

Our study was unusual in showing a better result at one year than at six months. This was because some subjects who relapsed returned for further treatment. This response contrasts with studies of other treatment methods, in which those who relapse tend to avoid returning even for follow-up. The tendency for those treated with nicotine gum to be more likely to return for a second course if they relapse may be a further reflection of its greater efficacy.

The rigorous design of our study with randomised allocation, double-blind placebo control, success rates based on all who started treatment, biochemical validation of reported abstinence, and the fact that the placebo provided nicotine to taste but with low biological availability all make it difficult to see how the gum could have achieved its effect other than by a specific action of the nicotine it provided. This further suggests an important role for nicotine in maintaining the habit of dependent smokers. That the active gum was significantly more effective in relieving withdrawal symptoms also supports the view that they may be caused partly by nicotine deprivation.

Another finding has possible implications for the role of nicotine in smoking. This is that the extent of gum use was significantly related to pretreatment plasma nicotine values but not to cigarette consumption in those given the active gum, whereas the reverse was true for those using the placebo. This suggests that the active gum was fulfilling a pharmacological need while the placebo may have been acting more as an oral substitute. The pharmacological role of nicotine is also supported by the fact that 7% of those assigned to receive the active gum developed some degree of dependence on it, while there were no instances of dependence on the placebo.

On the practical side, the active gum was apparently more effective in helping smokers to stop smoking during the first four weeks of treatment than in reducing the tendency to relapse thereafter. Other studies have found higher rates of long-term success among those who continued using the gum for longer periods. While it was realised that this may have been attributable to self-selection with people continuing to chew while they were successfully keeping off cigarettes but giving up the gum as soon as they relapsed to smoking, this finding was nevertheless used as a reason for suggesting that longer-term use of the gum might improve outcome. That our placebo group also showed a similar trend in this direction suggests that further study is needed to establish the optimal duration of gum use.

We emphasise that the high success rate achieved in this study was not necessarily due to the gum alone. The subjects had six group meetings with an experienced therapist. Above all they were given careful instructions on what they might realistically expect from the gum and how to use it correctly. This no doubt accounted for the lack of more than mild and transient side effects. That the active gum was initially the unpleasant to taste but subsequently became more satisfying points to the importance of encouraging subjects to persist with it for at least two weeks.

We conclude that after more than 20 years of unsuccessful research into all kinds of treatment methods for smokers, nicotine chewing-gum given to well-motivated smokers in a clinic setting is the first treatment to have been developed that has a specific effect over and above that attributable to an attention-placebo response. That it is also the first treatment to provide effective nicotine substitution has important implications for the role of nicotine in cigarette dependence.

We thank the Medical Research Council and Department of Health and Social Security for financial support; A B Leo for supplying the nicotine and placebo gumbals, and Vera Amato for secretarial help. Our colleagues Gill Devitt, John Stapleton, Steve Sutton, Colin Taylor, and Robert West gave helpful comments.

References


(Accepted 24 June 1982)