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Randomised controlled trial of smoking cessation intervention after admission for coronary heart disease

Petter Quist-Paulsen, Frode Gallefoss

Abstract

Objective To determine whether a nurse led smoking cessation intervention affects smoking cessation rates in patients admitted for coronary heart disease.

Design Randomised controlled trial.

Setting Cardiac ward of a general hospital, Norway.

Participants 240 smokers aged under 76 years admitted for myocardial infarction, unstable angina, or cardiac bypass surgery, 118 were randomly assigned to the intervention and 122 to usual care (control group).

Intervention The intervention was based on a booklet and focused on fear arousal and prevention of relapses. The intervention was delivered by cardiac nurses without special training. The intervention was initiated in hospital, and the participants were contacted regularly for at least five months.

Main outcome measure Smoking cessation rates at 12 months determined by self report and biochemical verification.

Results 12 months after admission to hospital, 57% (n = 57/100) of patients in the intervention group and 37% (n = 44/118) in the control group had quit smoking (absolute risk reduction 20%, 95% confidence interval 6% to 33%). The number needed to treat to get one additional person who would quit was 5 (3 to 16). Assuming all dropouts relapsed at 12 months, the smoking cessation rates were 50% in the intervention group and 37% in the control group (absolute risk reduction 13%, 9% to 20%).

Conclusion A smoking cessation programme delivered by cardiac nurses without special training, significantly reduced smoking rates in patients 12 months after admission to hospital for coronary heart disease.

Introduction

Smoking cessation after myocardial infarction is associated with a 50% reduction in mortality after three to five years. Reduced mortality is apparent after a few months and increases with time. After a coronary event, 30-45% of patients stop smoking spontaneously. Randomised investigations of smoking cessation after admission to hospital for coronary heart disease have obtained mixed results. Studies of interventions to change lifestyle, where helping patients to quit smoking was only part of the intervention, have not shown any statistically significant effects on smoking cessation rates.

In studies addressing only smoking cessation, those with brief interventions have been ineffective. Three of five trials with longer interventions (4-6 months) have shown increased smoking cessation rates. Only one of these, however, verified that patients had quit smoking by biochemical means.

This study used a complicated psychological approach, with specially trained nurses, and 29% of participants in the usual care group were lost to follow up.

We aimed to determine whether a nurse led smoking cessation intervention with emphasis on fear arousal affected smoking cessation rates after 12 months among patients admitted for coronary heart disease.

Methods

We invited to participate in our study all patients admitted to Vest-Agder Hospital, Kristiansand, Norway for myocardial infarction, unstable angina, or care after coronary bypass surgery performed at other hospitals. Eligible patients had to be under 76 years of age and daily smokers until the start of their present coronary symptoms. Patients who had undergone bypass surgery had to have been daily smokers until they received the date for surgery, and the cause had to be symptomatic coronary artery disease. Patients had to be sufficiently recovered to reliably receive the intervention, had to be able to read Norwegian, and had to live in Vest-Agder or Aust-Agder county. We excluded patients with serious illnesses associated with short life expectancies (cancer, chronic obstructive lung disease, renal or liver failure), serious psychiatric problems, alcoholism, and dementia.

We aimed to detect a 20% difference between the two groups. With a power of 80% ($\beta = 0.2$) and an $\alpha$ of less than 0.05, 98 patients were needed in each group ($\chi^2$, two tailed test, Sample Power version 1, SPSS, Chicago). To allow for dropouts, we decided to enrol 250 patients, and we estimated an inclusion period of two years.

Randomisation and intervention

The nurses recruited patients two to four days after admission. After providing written informed consent, the participants answered a baseline questionnaire and were randomly allocated to usual care (control group) or intervention. The nurses were given a serially numbered sealed envelope from a secretary who was otherwise uninvolved in the study. Randomisation was in blocks of varying sizes.

Doctors were not involved in the programme. Our intention was that all participants should receive doctors’ usual messages for quitting smoking. The doctors were not informed of the patients’ inclusion in the trial or of their randomisation status.

Control group

Patients were offered group sessions twice a week with the nurses, in which the importance of smoking cessation was mentioned. At sometime during these sessions a video was shown and a booklet handed out that contained general information on...
coronary heart disease and advice on quitting smoking. The control group received no further specific instructions on how to stop smoking.

**Intervention group**

One of three nurses consulted the patients once or twice during their hospital stay. The intervention was based on a 17 page booklet specially produced for the trial. This booklet emphasised the health benefits of quitting smoking after a coronary event. Two illustrations showed the differences in mortality between those who continued smoking after myocardial infarction or unstable angina and those who stopped. One of the illustrations was a bar chart showing a 60% risk reduction for death after five years of quitting, and the other was a linear chart showing that after 13 years 18% of patients who continued smoking were alive compared with 63% of those who had quit. On the basis of these figures, the participants were told that they most probably would have another heart attack if they continued smoking (fear arousal message).

The booklet also contained information on how to prevent relapse, how to stop smoking for those who had not stopped or had relapsed, and how to use nicotine replacements. Also explained was how to identify and cope with high risk situations for relapse, with action plans.

The patients were advised not to smoke during their hospital stay. Those with strong withdrawal urges were encouraged to use nicotine replacements (gum or patch). Spouses who smoked were also advised to quit.

The nurses contacted participants by telephone two days, one week, three weeks, three months, and five months after discharge. Those with special needs were telephoned monthly thereafter. At six weeks all participants in the intervention group had a consultation at the outpatient clinic with one of the cardiac nurses. The outpatient contacts included prevention of relapses and positive feedback (for example, telling patients who were still not smoking that they already had less chance of another heart attack). The health benefits of quitting were repeated and, if necessary, a fear arousal message given. Those who continued smoking or relapsed were offered additional support and advice.

**Outcome measures**

The patients were asked to return after 12 months. Those missing the appointment were given another one by letter. If they missed that one, they were telephoned and given an appointment. A home visit was offered to patients who failed to attend.

Smokers who stated that they were still smoking were classified as smokers and those who claimed they had quit and had a nicotine metabolite concentration in urine < 2.0 mmol/mol creatinine were classified as non-smokers. This limit was as validated at the university hospital in Sahlgren, Sweden, where urinary analyses were performed. The hospital used a radioimmunoassay method with antibodies labelled with iodine-125 (Double Antibody Nicotine Metabolite; Diagnostic Products, CA) with cotinine as standard for calibration, and calibration values 0.6-85.5 μmol/l.

**Statistical methods**

The χ² test was used to assess the effect of intervention. We calculated the number needed to treat, with confidence intervals. Simple and multiple logistic regression models were used to test the relation between baseline characteristics and outcome measures. We used SPSS for Windows (version 11) for all analyses.

### Table 1

<table>
<thead>
<tr>
<th>Characteristic</th>
<th>Intervention group (n=118)</th>
<th>Control group (n=122)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Mean (SD) age (years)</td>
<td>57 (7)</td>
<td>57 (7)</td>
</tr>
<tr>
<td>Married or living with partner</td>
<td>90 (76)</td>
<td>94 (77)</td>
</tr>
<tr>
<td>Employed</td>
<td>67 (57)</td>
<td>52 (43)</td>
</tr>
<tr>
<td>No education after primary school</td>
<td>46 (39)</td>
<td>33 (27)</td>
</tr>
<tr>
<td>Retried</td>
<td>21 (16)</td>
<td>35 (29)</td>
</tr>
<tr>
<td>Alcohol consumption &gt;1 unit a day</td>
<td>5 (4)</td>
<td>8 (7)</td>
</tr>
<tr>
<td>Previously no coronary artery disease</td>
<td>93 (79)</td>
<td>85 (70)</td>
</tr>
<tr>
<td>Myocardial infarction</td>
<td>91 (77)</td>
<td>85 (69)</td>
</tr>
<tr>
<td>Bypass surgery</td>
<td>10 (8)</td>
<td>18 (15)</td>
</tr>
<tr>
<td>Unstable angina</td>
<td>17 (14)</td>
<td>19 (16)</td>
</tr>
<tr>
<td>Mean (SD) no of days in hospital</td>
<td>6.9 (4.4)</td>
<td>6.7 (3.4)</td>
</tr>
<tr>
<td>Mean (SD) no of days in intensive care unit</td>
<td>2.7 (2.0)</td>
<td>2.9 (2.1)</td>
</tr>
<tr>
<td>Mean (SD) years of smoking</td>
<td>38.3 (13.9)</td>
<td>37.6 (11.5)</td>
</tr>
<tr>
<td>Mean (SD) no of cigarettes a day</td>
<td>143.5 (57)</td>
<td>156.6 (83.3)</td>
</tr>
<tr>
<td>Mean (SD) no of previous attempts to quit</td>
<td>2.3 (3.1)</td>
<td>2.3 (3.0)</td>
</tr>
<tr>
<td>Smoked in 24 hours before admission</td>
<td>101 (87)</td>
<td>114 (93)</td>
</tr>
<tr>
<td>Spouse who smokes</td>
<td>45 (36)</td>
<td>51 (42)</td>
</tr>
</tbody>
</table>
In both groups, 21% of patients reported smoking while in hospital. Despite the doctors’ claims that they gave firm and unequivocal advice about stopping smoking, only 59% (64/108) of patients in the intervention group and 58% (70/120) in the control group remembered receiving such a message.

**Intervention data**

Patients in the intervention group had an average of 1.6 (SD 0.7) consultations as inpatients and 1.6 (SD 1.5) as outpatients. They also received a mean of 8.5 (SD 3.2) telephone calls. Most patients (85%) received more telephone calls than the intended minimum of five. The mean total time devoted to each patient was 147 minutes (SD 50), including time to fill in questionnaires. Thirty six per cent (36/100) of the participants in the intervention group and 28% (33/118) in the control group used nicotine replacements (without a statistically significant difference).

**Smoking cessation rates**

Six of the patients who stated that they were non-smokers at 12 months had nicotine metabolite concentrations above the reference limit (three in each group), and three refused to provide a urine sample (one in the intervention group and two in the control group). All these patients were classified as smokers. The validated smoking cessation rates at 12 months were therefore 57% (57/100) in the intervention group and 37% (44/118) in the control group (absolute risk reduction 20%, 95% confidence interval 6% to 33%; table 2). The number needed to treat to get one additional patient to quit was 5 (3 to 16). The groups showed similar smoking cessation rates while in hospital and at six weeks’ follow up (table 2). No biochemical validations were made at this stage.

Of the 22 patients lost to follow up, seven died or changed address. Seven of the remaining 15 said they were non-smokers (not validated biochemically) and seven said they were smokers at the time of withdrawal. Assuming that these 15 patients returned to smoking at 12 months, in an intention to treat analysis the smoking cessation rates were 50% (57/114) in the intervention group and 37% (44/119) in the control group (absolute risk reduction 13%, 0% to 26%, number needed to treat 8, 4 to 250).

**Predictors of outcome**

In a simple logistic regression model, positive predictors of abstinence at 12 months were the number of days spent in the intensive care unit, myocardial infarction as reason for admission, no previous coronary heart disease, and being employed. In a multiple logistic regression analysis of predictors of abstinence, the intervention versus control group and absence versus occurrence of previous coronary heart disease showed statistically significant positive associations with cessation.

Only 9% (7/80) of the patients who smoked while in hospital or at six weeks’ follow up were abstinent at 12 months (three in the intervention group and four in the control group).

**Discussion**

A nurse led smoking cessation intervention with at least five months’ follow up increased smoking cessation rates among patients admitted to hospital for coronary heart disease. Further intervention had little impact in those patients who smoked while in hospital or at six weeks’ follow up. Since there were no differences in smoking cessation rates between the groups at six weeks, we speculate that a long intervention period was an important factor. It is also possible that the initial intervention provided such a strong motivation for abstinence that it prevented later relapses.

Our dropout rate of less than 10% was lower than in comparable studies. Only one patient in the control group, excluding those who died or changed address, was lost to follow up. The dropout rate in the intervention group was higher than in the control group. This may have been a result of the intervention itself.

Several reasons could explain why there were no differences in smoking cessation rates between the two groups at six weeks. Firstly, our medical department is small (110 beds), and it was therefore not possible to segregate the two groups. The control group could have been contaminated if patients in the intervention group passed along relevant messages or the booklet. We do not know to what extent this happened, but the study nurses did notice some communication about smoking cessation between the patients. Secondly, we did not emphasise to which group patients were allocated. Thus, some patients in the control group believed they received a smoking cessation programme, which could have motivated them to abstain. Thirdly, staff members knew that a smoking cessation trial was being conducted, and this may have focused their attention on the smoking intervention. It is possible that the control group received a better standard of advice on smoking cessation than usual.

Only six out of 10 patients remembered receiving advice to stop smoking from a doctor, and even though around three quarters of the patients stopped smoking during their hospital stay, it is possible that this could have been increased further by including doctors in the programme. Specially trained nurses may also have prevented more relapses than nurses without special skills in smoking cessation. However, these methods may not be feasible in routine care. Most smoking cessation programmes use a group format, which may be more effective. But several weeks of enrolment would be needed to gather enough patients for such a programme, even in a large hospital, and by this time patients’ motivation for quitting has probably diminished.

We believe that our study is the largest to date addressing only smoking cessation, with a long intervention period. The applicability of the results was strengthened by the low dropout rate and by the inclusion of most patients who smoked, regardless of previous coronary heart disease. Further, the principles of the intervention, focusing on fear arousal and relapse prevention, were simple. Assuming a time commitment of 2.4 hours for each patient, a nurse working part time has the capacity to conduct individual smoking cessation programmes on eight patients a week. This intervention is probably cost effective compared with other medical treatments in preventing cardiac events and deaths, and we suggest that similar programmes should be provided as part of routine care in wards dealing with cardiac conditions.

We thank the cardiac nurses Tone Baeck, Eva Boroye, and Anne Kari Kjellesvik for delivering the intervention and collecting data.

**Contributors:** PQ-P and FG contributed to the planning and conduction of the study, the analysis and interpretation of the data, and writing the paper.
They will act as guarantors for the paper. Kjetil Drangsholt, Pal Friis, Finn Tore Gjestvang, and Anders Wahlstedt contributed to the planning of the study.

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Ethical approval: The study was approved by the regional ethics committee.


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