Motivational interviewing for modifying diabetes risk: a randomised controlled trial

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ABSTRACT

Background
Around 10–15% of adults aged over 40 years have pre-diabetes, which carries a high risk of progression to type 2 diabetes. Intensive lifestyle intervention reduces progression by as much as 58%. However, the cost and personnel requirements of these interventions are major obstacles to delivery in NHS primary care.

Aim
To assess the effectiveness of a low-cost intervention, delivered in primary care by non-NHS staff, to reduce the risk of diabetes through weight loss and physical activity.

Design of study
Pragmatic single-blind randomised controlled trial with researchers and statistician blinded to group allocation.

Setting
UK primary care.

Method
One-hundred and forty-one participants with a body mass index of 28 kg/m² or more, but without diabetes or heart disease, received either information leaflets or individual behavioural counselling using motivational interviewing techniques. The intervention was delivered by five counsellors recruited from the local community. The primary outcomes were the proportions of participants meeting predefined targets for weight loss (5%) and moderate physical activity (150 minutes/week) after 6 months.

Results
Using intention-to-treat analysis, more people in the intervention group achieved the weight-loss target (24% versus 7% for controls; odds ratio [OR] = 3.96; 95% confidence interval [CI] = 1.4 to 11.4; number needed to treat [NNT] = 6.1 (95% CI = 4 to 21)). The proportion achieving the physical activity target did not increase significantly (38% versus 28% for controls; OR = 1.6; 95% CI = 0.7 to 3.8).

Conclusion
Short-term weight loss, at a level which, if sustained, is clinically meaningful for reducing diabetes risk, is achievable in primary care, without excessive use of NHS monetary or personnel resources.

Keywords
behaviour therapy; diabetes mellitus; intervention studies; primary prevention; randomised controlled trial; weight loss.

INTRODUCTION

Recent large studies in the US, China, and Finland have shown that modest changes in physical activity and diet can have a large health impact in terms of preventing progression to type 2 diabetes. Around 15% of adults aged over 40 years have either impaired fasting glycaemia or impaired glucose tolerance (collectively known as pre-diabetes), and large numbers may be detected in UK primary care over the next few years, as initiatives for primary prevention of heart disease and (possibly) targeted diabetes screening are implemented.

The problem with the diabetes-prevention studies conducted to date is that to achieve the required behavioural changes has required very intensive interventions. The Finnish Diabetes Prevention Study involved seven individuals sessions with a nutritionist, quarterly review sessions, plus separate individual physical activity coaching and behavioural counselling sessions. The US programme involved 16 weekly individual sessions, followed by an individualised maintenance programme including further individual and group sessions. The intervention was delivered by...
registered dieticians, exercise physiologists, professional behavioural psychologists, and health educators. This intensity of professional-led intervention is not likely to be affordable in UK primary care, and the resources in terms of available staff time are also a major constraint. The need for evaluation of a range of more practical community-based options has been highlighted by key authors in this field. 4, 6, 7

One promising approach, which offers a well-defined and evidence-based model for behaviour change, is motivational interviewing. 8–11 Motivational interviewing focuses on exploring and making explicit the reasons someone might have to engage in a particular behaviour or course of action. Strongly patient-centred interviewing techniques are used to contrast the pros and cons of change, reduce sources of resistance, and identify possible barriers to change. Action plans are then made, which specifically ‘situate’ the planned change in the context of the person’s day-to-day life. The principles and techniques are easy to learn, and well-established training courses are available through a national network of accredited trainers. 12 The techniques can also be used successfully by non-health professionals. 13

The aim of this study therefore was to see whether changes in weight and physical activity similar to those obtained in the Finnish and US diabetes-prevention studies could be achieved with a simplified intervention, based on motivational interviewing, and delivered by trained health-promotion workers not currently working for the NHS.

METHOD

Design

A single-blind, pragmatic, randomised controlled trial was conducted in a community setting, with participants allocated to receive either intervention or ‘information only’ in addition to any usual care being received. ‘Usual care’ means that the care of the participant was not interfered with in any way outside of the study setting, although changes in medication or new diagnosis of illness during the course of the study were monitored.

Intervention

Five ‘health-promotion counsellors’ were trained to deliver motivational interviewing in one-to-one individual consultations. They worked part-time and were selected to have prior experience in promoting physical activity or healthy diet with adults, and not to be current NHS employees. Two were ex-NHS staff (health visitor, rehabilitation nurse) and three were postgraduate students in sports and health science. The training included 2 days on the assessment of existing diet and physical activity, and setting appropriate, individually tailored, diet and activity targets. Dietary recommendations were based on reducing overall calorific intake and portion size, reducing overall fat intake, reducing saturated fat content, and increasing fibre intake. This included aiming to increase ingestion of wholegrain products, vegetables, fruit, low-fat milk, low-fat meat, soft margarines, and vegetable oils rich in monounsaturated fat. Physical activity recommendations focused on increasing overall physical activity within the context of the individual’s existing life, rather than specifying a predetermined activity regime. Training in supporting behaviour change consisted of a 2-day course in motivational interviewing, provided by an accredited motivational interviewing trainer. 12

Participants in the intervention group received up to 11 individual sessions over a 6 month period. A mixture of one-to-one contacts (median 8) and telephone contacts (median 1.5) was received with a mean 34 minutes per contact. Action plans were made and checked/modified at subsequent contacts using relapse-management/relapse-prevention techniques, and targets were increased gradually to build/reinforce confidence over time. Participants were also encouraged to self-monitor their own weight, activity, and energy levels. A specific aim of the intervention was to encourage participants to develop sustainable cognitive and behavioural skills for managing their diet and physical activity. The fixed-term nature of the intervention was emphasised, and the length of appointments tapered off to allow a planned withdrawal of support and to minimise dependence on the therapist. Interventions were delivered in a consulting room at a local community hospital in Devon. Intervention fidelity was checked by analysing transcripts of interviews conducted by the counsellors with an actor who responded based on a given scenario. One behavioural scientist and one nurse consultant with experience of using motivational interviewing techniques marked the transcripts, using the Behaviour Change Counselling Index (BECCI). 14 The standard achieved was deemed (in the opinion of the assessors) to be satisfactory for all counsellors. Attendance records indicated that 96% of the...
intervention group received at least three sessions of treatment, and 79% received six or more sessions. Reviews of motivational interviewing suggest that at least two sessions and at least 60 minutes of contact time are required for benefit. Hence, the majority of intervention participants received a therapeutic dose of treatment.

Control
Control group participants received a standardised information pack promoting similar diet and physical activity recommendations to those above. Specifically, British Heart Foundation health-promotion materials were used; the NHS Smoking Cessation Service ‘Green Book’; and locally produced information on physical activity and local ‘walk and talk’ activities. At the end of the study (post data collection), controls received two individual sessions with the study counsellors.

Participants
The study recruited adults (age ≥18 years) identified by their GP as having a body mass index (BMI) of 28 kg/m² or more (a level clearly associated with increased diabetes risk) recorded in the last 10 years. Participants were excluded if they had existing diabetes, heart disease, or severe joint problems, if they felt they would be unable to engage in at least moderate physical activity, if the GP felt the patient to be unsuitable (for example, severe uncontrolled hypertension, dementia, or mental illness), or if they did not speak English fluently.

Measures
Outcomes were measured at baseline and 6 months after the first consultation. Baseline demographic data on age, sex, and level of education (primary, secondary, or degree level) were collected. Participants were also asked about the onset of new medical conditions, medication use, and attendance at other weight-loss or physical activity programmes during the course of the study. The primary outcomes were the proportion of patients achieving the prespecified target of 5% reduction in weight, and the proportion of patients achieving the UK government target of 150 minutes of moderate activity per week. Secondary outcomes were weight (kg) and waist circumference (cm), measured using appropriate guidelines. Physical activity levels were self-reported using the Modifiable Activity Questionnaire (MAQ).

Analysis
The primary analysis was an intention-to-treat analysis of the proportions achieving the specified targets, using logistic regression with group allocation and baseline values entered into the regression model (except for weight loss, where the baseline value is integral to the measure). Where follow-up data were missing, the baseline value was carried forward (that is, no change from baseline was assumed). Secondary outcomes were compared using analysis of covariance (ANCOVA; with baseline-dependent variable scores entered into the model). Further analyses assessed the sensitivity of both primary and secondary analyses to differences in baseline characteristics (age, sex, and education level) and to other factors that might affect weight (incidence of new illness, or medication that might affect weight, and use of other weight-loss programmes). Analyses were conducted using SPSS (version 14).

Sample size calculation
The sample size was calculated based on the analysis of primary outcomes (target weight and activity). Estimates of the likely proportions were taken from previous studies (43% achieving target weight loss and 36% physical activity in the intervention group versus 13% and 16% in the control group, using intention-to-treat basis). To have an 80% chance of detecting these differences, with a (two-sided) 0.05 significance level, 34 patients per group were required for weight, and 73 per group for physical activity.

Procedure
Participants were recruited between February and June 2006 from two GP surgeries in a semi-rural town in the UK. A list of eligible patients was identified by searching practice databases for people meeting the BMI, age, diabetes, and heart disease criteria. The patients’ GPs checked for additional exclusion criteria. Participants were invited by letter to attend group recruitment meetings at a local community hall, and the researchers attempted to confirm attendance by telephone contact. At the meetings, the study was explained in a brief presentation. Refreshments were served and participants could leave at this stage if desired. Written consent was taken individually, using well-spaced desks at one end of the hall. Participants completed baseline questionnaires and then their weight and waist were measured in private rooms.

Randomisation
Patients were randomly allocated to treatment groups using sealed envelopes, each containing either a control or an intervention slip. These were drawn from a box by a researcher, following completion of baseline measures. A batch of envelopes was prepared at each recruitment session by the chief investigator or the delegated representative, to match the number of participants attending.

Blinding
Researchers taking weight and waist measures were blind to group allocation (at baseline, measures were taken prior to randomisation, and at follow-up, patients went into a private room for measurement, which involved only a brief interaction with the researcher). Participants were, by necessity, unblinded. The statistician was blinded to group allocation by withholding the key to group labelling until after analysis.

RESULTS
Of the 247 eligible patients that could be contacted, 141 (57%) agreed to take part. At the 6-month follow-up point, 115 patients (82%) provided weight and waist circumference data, although only 88 (62%) provided physical activity questionnaires. The recruited sample was broadly similar to the eligible population (mean age 51.9 years, 58% female), being older by 2.7 years and with a slightly higher proportion of females (64%). The flow of participants through the study is shown in Figure 1. Table 1 shows the sample characteristics at baseline.

Table 2 shows the outcomes at 6 months. A significantly higher proportion in the intervention group achieved 5% weight loss (24% versus 7% for controls; odds ratio [OR] = 4.0; 95% confidence interval [CI] = 1.4 to 11.4). The number needed to treat (NNT) to achieve 5% weight loss was 6.1 (95% CI = 3.6 to 20.8). A higher proportion in the intervention group achieved the 150-minute physical activity target, although this difference was not significant (38% versus 28% for controls; OR = 1.6; 95% CI = 0.7 to 3.8).

Secondary analyses showed a significant difference in weight (adjusted mean difference 1.3 kg; 95% CI = 0.2 to 2.4), and a small, non-significant difference in waist circumference between the two groups (adjusted mean difference: 1.6 cm; 95% CI = -0.01 to 3.3, P = 0.051).

The results showed very little sensitivity to baseline demographic characteristics. A sensitivity analysis excluding six people who developed potentially confounding medical conditions during the study (two thyroid problems, one pre-diabetes with metformin treatment, one heart condition, two musculoskeletal problems), and eight who received alternative weight-loss interventions (one orlistat, seven slimming or gym-based programmes) slightly increased the odds ratio for achieving weight loss (OR = 4.5 (95% CI = 1.4 to 14.6). For those completing the study (per-protocol analysis), 29% in the intervention group achieved the weight-loss target (versus 9% in the control group), and 46% achieved the physical activity target (versus 34% of controls).

The costs for delivering the intervention were estimated to be £263 per patient, based on financial records of the direct costs associated with the project (salaries, supervision/management time, stationery costs) and estimates of indirect costs (consulting room, reception support and hospital overheads, telephone, secretarial support, and general practice costs for the identification and referral of patients) but not project set-up costs (for example, recruitment). Appendix 1

| Table 1. Baseline characteristics. |
|-------------------------------|-------------------|-------------------|
|                               | Intervention (n = 72) | Control (n = 69)  |
| Mean (SD) age, years          | 53.3 (12.3)         | 54.5 (11.5)       |
| Female sex, n (%)             | 46 (64)             | 44 (64)           |
| Secondary level education, n (%)| 45 (63)             | 45 (66)           |
| Degree level education, n (%) | 26 (36)             | 20 (29)           |
| Mean (SD) weight, kg          | 91.6 (13.3)         | 94.4 (14.2)       |
| Mean (SD) waist circumference, cm | 104.2 (11.3)      | 106.4 (10.5)      |
| Meeting physical activity target, n (%) | 24 (33.3)         | 20 (29.0)         |

*Did not attend follow-up weight/waist measurement session. LVCF = last value carried forward.
Table 2. Outcomes at 6 months.

<table>
<thead>
<tr>
<th></th>
<th>Intervention (n = 72)</th>
<th>Control (n = 69)</th>
<th>Size of effect/ significance</th>
</tr>
</thead>
<tbody>
<tr>
<td>Total (%) achieving 5% weight loss*</td>
<td>17 (23.6)</td>
<td>5 (7.2)</td>
<td>OR = 4.0; 95% CI = 1.4 to 11.4</td>
</tr>
<tr>
<td>Total (%) achieving 150 minutes activity*</td>
<td>27 (37.5)</td>
<td>19 (27.5)</td>
<td>OR = 1.6; 95% CI = 0.7 to 3.8</td>
</tr>
<tr>
<td>Number (%) achieving 150 minutes moderate activity, but not achieving this at baseline*</td>
<td>10 (13.9)</td>
<td>5 (7.2)</td>
<td>OR = 2.1; 95% CI = 0.7 to 6.4</td>
</tr>
<tr>
<td>Mean (SD) weight, kg</td>
<td>91.3 (13.7)</td>
<td>92.6 (15.0)</td>
<td>Mean difference 1.3; 95% CI = 0.2 to 2.4</td>
</tr>
<tr>
<td>Mean (SD) waist circumference, cm</td>
<td>102.6 (11.2)</td>
<td>104.1 (11.6)</td>
<td>Mean difference 1.6; 95% CI = -0.01 to 3.3</td>
</tr>
</tbody>
</table>

OR = odds ratio. *Analysis takes into account baseline value as an integral part of the measure.
**Analysis and means (but not proportions) adjusted for value of baseline dependent variable.

DISCUSSION

Summary of main findings

The study shows that personnel recruited from outside the existing health service, with brief training in motivational interviewing, can help people to achieve a 5% weight-loss target. The results suggest an additional small increase in the proportion achieving a clinically meaningful change in physical activity, although this was not significant.

Strengths and limitations of the study

The study’s main strengths were in its pragmatic nature and setting and broad inclusion criteria, which increase the likely generalisability of the results. The effect was estimated in a practice setting, with realistic dropout and non-adherence to treatment taken into account using intention-to-treat analysis. A high retention rate was achieved in both groups, with 88% of men and 76% of women completing the intervention. The relatively high degree of engagement and retention of men is worth noting. The collection of objective measures (weight and waist) was blinded to group allocation. However, costs were not rigorously estimated or analysed. The study had insufficient power to detect the (smaller than expected) changes in proportions achieving the physical activity target. This was compounded by the fact that the response rate for physical activity questionnaires was low. In addition, the self-report of physical activity has low sensitivity and limited validity (a tendency for over-estimation and low-to-moderate correlations with objective measures). Generalisability may also be limited by the self-selecting nature of the sample (around 57% agreed to take part), although it should be noted that this kind of self-selection would also apply to a GP-mediated offer of referral to a weight-loss intervention.

The practices involved were semi-rural with a broad population demographic, but with only limited representation from ethnic minority groups. A larger study would therefore be needed to establish generalisability to other UK practices.

A further limitation of this study is that it is not possible to say whether the effects found would persist beyond 6 months. It is known that the effects of more-intensive interventions, such as the Finnish Diabetes Prevention Study, can be largely maintained for as much as 7 years. However, less-intensive behavioural interventions are less effective and often lose their effects over time, and weight-loss interventions often suffer from high dropout. It could be hypothesised that the type of intervention used in the present study should provide good maintenance of weight loss, as motivational interviewing aims to equip people with the cognitive tools to make and sustain behaviour change. This is supported by review evidence that effect sizes in nine trials of motivational interviewing, involving 1500 people, did not significantly decrease between 20 and 67 weeks. However, the existence of longer-term, generalisable benefits for this specific application of motivational interviewing needs to be established in larger clinical trials.

Comparison with existing literature

This study’s findings are consistent with other trials of motivational interviewing, which have produced significant weight loss and physical activity increases.

The Finnish Diabetes Prevention Study suggests that weight is the dominant behavioural predictor of progression to diabetes. In their study, people with hyperglycaemia achieving an initial 5% weight loss were 55% less likely (hazard ratio 0.45) to develop type 2 diabetes within 7 years. In the present study, the difference in the proportions achieving 5% weight loss (17%) was less than that achieved in the Finnish trial (30%). One reason for this could be that the participants did not have pre-diabetes, which may increase motivation for behaviour change. However, it is likely that the reduced intensity of intervention was also a factor.

This study therefore suggests that in devising lower-cost, simpler interventions, there may be a trade off between effect size and cost. The costs per patient of the study (estimated £263) were well below those of the US and Finnish diabetes-prevention programmes (£179638 and £211639 respectively, when inflated to 2007 prices). A comparison with pharmaceutical approaches is also possible. The NNT for 2 years to achieve 5% weight loss using orlistat is 6 (95% CI = 4 to 14), with an estimated treatment cost (at 2007 prices) of £1759 per patient. However, it is worth...
noting that the present study did not set out to formally measure or compare cost-effectiveness, and these figures are presented mainly to contextualise this study as a lower-cost approach.

**Implications for future research and clinical practice**

Future research in this area should use less-burdensome and more objective measures of physical activity (for example, a simpler questionnaire and accelerometers). The longer-term effectiveness and cost-effectiveness of a range of well-defined intervention approaches, with different levels of intensity, needs to be established to inform policy decisions and future clinical practice in this area. Monitoring of intervention processes (both intervention delivery and intra-individual/psychological processes) and the relation of processes to outcomes is also essential for advancing the science and methods of behavioural intervention.21

The current study has shown that it is feasible to deliver a low-cost intervention, which produces meaningful changes in weight, at a level which, if sustained, can modify progression to diabetes. The intervention can be delivered in primary care, without creating an excessive drain on NHS monetary or personnel resources.

**Funding body**

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**Ethical approval**

North and East Devon Local Research Ethics Committee (05/Q2102/04)

**Competing interests**

The authors have stated that there are none.

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**REFERENCES**


### Appendix 1. Estimated intervention costs.

<table>
<thead>
<tr>
<th>Cost Item</th>
<th>Cost (£)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Total project worker time (inclusive of on-costs, supervision, and training time) 1015 hours @ £10.57</td>
<td>10 728.55</td>
</tr>
<tr>
<td>Management team/supervision time: 52 weeks at 2 hours per week @ £19.98/h (including on-costs)</td>
<td>2077.92</td>
</tr>
<tr>
<td>Room cost/reception cover (including hospital overheads at 100%)</td>
<td>2481.72</td>
</tr>
<tr>
<td>Stationery</td>
<td>135.00</td>
</tr>
<tr>
<td>Telephone (estimated)</td>
<td>500.00</td>
</tr>
<tr>
<td>Secretarial support: 52 hours (1/week) @ £9.80 (including on-costs)</td>
<td>509.60</td>
</tr>
<tr>
<td>Training (motivational interviewing), as charged</td>
<td>1500.00</td>
</tr>
<tr>
<td>Training (internal) 4 hours × 4 trainers × (average) £34.80/h (including on-costs)</td>
<td>556.80</td>
</tr>
<tr>
<td>GP costs (including on-costs) for identification and referral of patients (5 minutes administrator £14.60/hour and 5 minutes GP time £60.63/hour) ÷ 72 patients</td>
<td>451.38</td>
</tr>
</tbody>
</table>

**Total cost**  
18 940.97

**Cost per intervention**  
263.07