**Large multicentre pilot randomised controlled trial testing a low-cost, tailored, self-help smoking cessation text message intervention for pregnant smokers (MiQuit).**

**Running head:** Randomised controlled trial of MiQuit

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**Declaration of competing interest:** None to declare**ABSTRACT**

**Aims:** To estimate the effectiveness of pregnancy smoking cessation support delivered by SMS text message and key parameters needed to plan a definitive trial.

**Design:** Multicentre, parallel-group, single-blinded, individual randomised controlled trial

**Setting:** 16 antenatal clinics in England.

**Participants:** 407 participants were randomised to the intervention (n=203) or usual care (n=204). Eligible women were <25 weeks gestation, smoked at least 1 daily cigarette (> 5 pre-pregnancy), were able to receive and understand English SMS texts and were not already using text-based cessation support.

**Intervention:** All participants received a smoking cessation leaflet; intervention participants also received a 12-week programme of individually-tailored, automated, interactive, self-help smoking cessation text messages (MiQuit).

**Outcome Measurements:** Seven smoking outcomes including validated continuous abstinence from 4 weeks post-randomisation until 36 weeks gestation, design parameters for a future trial and cost-per-quitter.

**Findings:** Using the validated, continuous abstinence outcome, 5.4% (11/203) of MiQuit participants were abstinent versus 2.0% (4/204) of usual care participants (odds ratio [OR] 2.7, 95% confidence interval [CI] 0.93 to 9.35). The Bayes Factor for this outcome was 2.23. Completeness of follow up at 36 weeks gestation was similar in both groups; provision of self-report smoking data was 64% (MiQuit) and 65% (usual care) and abstinence validation rates were 56% (MiQuit) and 61% (usual care). The incremental cost-per-quitter was £133.53 (95% CI -£395.78 to £843.62).

**Conclusions:** There was some evidence, though not conclusive, that a text messaging programme may increase cessation rates in pregnant smokers when provided alongside routine NHS cessation care.

**Keywords:** Smoking cessation, pregnancy, self-help, randomised controlled trial, SMS text messaging, mHealth

**INTRODUCTION**

Smoking in pregnancy is strongly associated with pregnancy complications including miscarriage,1 spontaneous preterm birth,2 small for gestational age,2 and stillbirth.3 4 Smoking in pregnancy also perpetuates health inequalities; rates are around five times higher in the most deprived women compared with the least deprived5-7 and children born to smokers have an increased risk of becoming smokers themselves.8 9 Systematic review evidence shows that behavioural smoking cessation interventions reduce the risks of preterm birth and low birthweight by around 18%.10

Structured self-help support helps pregnant smokers to stop.11 12 Mobile phone text messaging is a simple way of providing self-help support and is effective for non-pregnant smokers.13 However, many aspects of ‘generic’ text messaging cessation systems are unlikely to be appropriate in pregnancy. Available generic programmes make no mention of pregnancy,13 which for most pregnant smokers is the main reason they try quitting,14 and effective behavioural support for pregnant smokers is typically strongly pregnancy-orientated.10 Consequently, pregnant smokers may find much of the behavioural support delivered by generic programmes irrelevant, reducing its impact and perhaps even being counterproductive.15 Even more importantly, available ‘generic’ programmes provide some advice and support that potentially could be harmful in pregnancy. For example, use of nicotine replacement therapy (NRT) is encouraged without consideration of pregnancy-specific risks, and generic advice on keeping fit and weight gain after quitting are quite different from what might be appropriate in pregnancy.

To maximise the potential of self-help support for helping pregnant smokers to stop, we have developed an individually-tailored SMS text messaging intervention specifically for pregnant smokers, called MiQuit. This process followed the Medical Research Council framework for developing and evaluating complex interventions16 and was informed by extensive qualitative work with pregnant smokers.17 MiQuit can be used by all pregnant smokers as the support it provides is tailored to a woman’s level of motivation to quit. A randomised controlled trial (RCT) (N=207) demonstrated that randomisation to MiQuit or routine care is feasible, that women find MiQuit highly acceptable and that MiQuit is likely to encourage cessation until at least mid-pregnancy.18 This feasibility trial provided the best available estimate for MiQuit efficacy, albeit for a relatively brief cessation period; we believed cessation at the end of pregnancy would be a more appropriate outcome for a definitive trial as this would result in maximal benefits for the fetus. As MiQuit is a cheap intervention with potential for wide dissemination, we anticipated that even a 1-2% absolute effect on smoking cessation in pregnancy could prove clinically important and cost effective and the imprecise efficacy estimate we had obtained suggested that an impact of this size was potentially attainable. Consequently, we planned a full trial to detect such an effect on smoking cessation until the end of pregnancy and estimated this could require 3-4000 participants. This large, pilot RCT was conducted to investigate the feasibility of undertaking a much larger multi-centre RCT in UK National Health Service (NHS) settings to determine whether or not MiQuit can impact on cessation throughout pregnancy. The current trial would also provide estimates of effectiveness and cost effectiveness, with the latter enabling comparisons with other cessation interventions.

**METHODS**

**Design**

This was a multicentre, two-arm, parallel group, single blind, individually randomised controlled trial.

**Study population**

Participants were recruited from 16 English NHS hospital antenatal clinics between February and September 2014. They were aged 16 and over, less than 25 weeks pregnant, had smoked at least five cigarettes daily before pregnancy and at least one per day at enrolment, able to understand written English and owned a mobile phone with text messaging functionality. Participants already using text message-based smoking cessation support were excluded.

**Study protocol and interventions**

The study protocol was approved by Nottingham 1 Research Ethics Committee (Ref.:13/EM/0427) and subsequently published.19

Usual care
Participants were given a standard NHS booklet on smoking cessation for mums-to-be (appendix 1) and could access smoking cessation information, advice or support for stopping smoking offered as part of routine antenatal care.

Intervention

Two days after enrolment, in addition to the booklet and usual care, intervention participants started to receive MiQuit; an automated 12-week advice and support programme for quitting smoking in pregnancy delivered by SMS text message. MiQuit objectives are informed by Social Cognitive Theory,20 Perspectives on Change Theory (Borland, 2000, unpublished work), the Elaboration Likelihood Model of Persuasion21 and several additional cognitive determinants of quitting smoking in pregnancy.18 It uses 14 participant characteristics to individually-tailor support.22 Tailoring characteristics include gestation, motivation to quit, the hardest situation to avoid smoking, cessation self-efficacy, cigarette dependence and partner’s smoking status. 'Push' support (i.e. automated support sent to participants’ phones) is delivered according to a delivery schedule (0, 1 or 2 daily texts). Push message frequency is highest in the first 4 weeks. Push support includes motivational messages, advice about quit attempt preparation, managing cravings and withdrawal, dealing with trigger situations and preventing lapses, information about fetal development and how smoking affects this (see appendix 2 for example messages and tailoring variables). Users can alter support frequency by texting the keywords MORE or LESS, and are encouraged to set and send a quit date to MiQuit to enable them to receive additional support orientated around when their quit attempt begins. At 3 and 7 weeks into the programme, users are asked to respond to texts asking about smoking in the previous 3 days, so that subsequent support is further tailored to smoking behaviour.22 Additionally, system users can 'pull' on-demand support for combatting cravings or temptation to smoke by texting HELP and seek advice on returning to abstinence after a lapse by texting SLIP. Alternatively, texting QUIZ provides a multiple choice message trivia game designed to distract users from smoking. Support can be discontinued by texting STOP. More detailed information about the development and structure of the intervention can be found elsewhere18 22

Enrolment, randomisation and blinding
Research midwives (RMs) identified potential participants in antenatal clinics via their clinic notes or a screening questionnaire, and interested women were provided with participant information sheets. RMs sought written consent, but if time was insufficient, contact details were requested instead and verbal consent was sought later in a phone call from the RM or a researcher from the trial coordination team. Next, baseline data were collected and, after this was entered onto a web-based database, participants were individually randomised to usual care or the MiQuit intervention in a 1:1 ratio using the Nottingham Clinical Trials Unit web-based system with both the RM or researcher and the participant remaining masked to allocation. Randomisation used a computer generated pseudo-random code with random permuted blocks of randomly varying size, and stratification was by study site and gestation (<16 weeks vs. ≥16 weeks). Following randomisation, unblinded trial team members sent arm-specific information packs to participants, which included the usual care booklet. Those dispatching packs were not involved in collecting follow-up data. Trial staff involved in follow-up remained unaware of participants’ treatments until questions on the intervention were asked at the end of the study, after smoking outcome data had been collected.

Data collection

Baseline data included contact details, age, highest qualification, postcode to enable matching to Index of Multiple Deprivation (IMD) scores,23 ethnicity (based on UK Census categories), gestation, pre-pregnancy smoking rate, heaviness of smoking index,24 strength and frequency of urges to smoke,25 whether a quit date had been set, intention to quit,18 number of births beyond 24 weeks, partner’s (significant other’s) smoking status and health status using EQ-5D.26

Four weeks after randomisation participants were contacted to complete a questionnaire assessing smoking status over the past 7 days; we used text messages to notify them to expect a telephone call and if after several attempts the call was unsuccessful, we posted and emailed a link to the questionnaire. At 36 weeks gestation participants were similarly contacted and asked about smoking behaviour since 4 weeks post-randomisation and in the past 7 days, quit attempts lasting at least 24 hours and use of smoking cessation support. MiQuit arm participants were also asked their views on the intervention. Where 7-day complete abstinence from smoking was reported, we immediately attempted to biochemically validate this with exhaled-breath Carbon Monoxide (CO) readings and/or saliva samples tested for cotinine, with samples or readings collected at hospital or home visits. If face-to-face collection was not successful, postal saliva sample packs were used. Before samples were donated, participants were asked either verbally or by questionnaire about smoking status and use of nicotine replacement therapies (NRT) or e-cigarettes.

To encourage engagement, participants received a £5 shopping voucher for providing data at each of the first three contacts (i.e. £15 maximum); a £10 voucher was also provided after validation visits. Participants were informed of how to withdraw from data collection via postcard, telephone, text, or email.

**Outcomes**

Future trial design parameters
We monitored monthly rates of recruitment, outcome ascertainment rates, and estimated the validated abstinence rate in both trial arms combined. We aimed to enrol 400 participants in 12 months. The key smoking outcome for a future trial is described below (#1).

Smoking

Smoking measures were: 1) self-reported abstinence from 4 weeks post-randomisation until late pregnancy collected at late pregnancy follow up (approximately 36 weeks gestation), with no more than 5 cigarettes in total between the two time points,27 biochemically validated at the later time; 2) as 1 but self-report only; 3) self-reported 7-day point prevalence abstinence at late pregnancy; 4) as 3 but biochemically validated; 5) self-reported 7-day point prevalence abstinence at 4 weeks post-randomisation; 6) self-reported 7-day point prevalence abstinence at both 4 weeks post-randomisation and late pregnancy; 7) as 6 but biochemically validated in late pregnancy.

We stated a priori that we anticipated that outcome #1, continuous abstinence from 4 weeks post-randomisation until 36 weeks gestation, would be most appropriate for a future RCT to definitively assess MiQuit efficacy.19 We had concerns about the viability of using this outcome, so a key objective was to ascertain its feasibility of measurement. Where participants reported abstinence but were using NRT or e-cigarettes, CO readings alone were used for validation (cut point of <9 ppm). Otherwise, a saliva cotinine reading of <10 ng/ml was also required.28 Where data from only one validation method were available, a value below the relevant cut-point was considered sufficient. Saliva was analysed by ABS Laboratories Ltd, Hertfordshire.

Economic
As the usual care and intervention groups both had access to standard NHS smoking cessation and antenatal care, it was assumed that both groups had equal cost, therefore the only additional cost would be for delivering MiQuit. Costs included were the text messages and the annual running cost. These were based on historical costs incurred. Costs were calculated at 2014-2015 price per year from a NHS and Personal Social Services perspective.

Sample size
The sample size was justified primarily on the basis of how precisely key parameters for the design of a definitive RCT could be estimated. With 400 participants (200 per group), we could estimate the overall recruitment rate to within +/-1%, outcome ascertainment rates per treatment group to within +/-4%, and combined quit rates for both groups to within +/-3%. Precision estimates for detecting between-group differences in quit rates were calculated for ranges of treatment effects (i.e. odds ratio [OR]) and usual care group quit rates;19 for example, these showed that if a 5% usual care group quit rate occurred in late pregnancy, with 400 participants the trial would estimate an OR of 1.8 (as noted in a previous review)12 with 80% confidence intervals (CIs) of 1.06 to 3.05).19

**Statistical analysis**

A statistical analysis plan was agreed with the Trial Steering Committee and published with the trial protocol.19 Recruitment and outcome ascertainment rates were estimated with 95% CIs. For each treatment group, and for both groups combined, abstinence rates for each outcome were estimated with 95% Wilson CIs. Chi-squared tests (Fisher’s exact tests in cases with small expected frequencies) were performed to assess the association between smoking outcomes and treatment group. Firth (penalised) logistic regression models29 were then used to estimate odds ratios with 95% profile CIs30 to compare smoking outcomes between treatment groups, adjusting for factors used to stratify the randomisation via their inclusion as fixed covariates in each model (trial site, gestation at randomisation). Three additional models for all seven smoking outcomes were carried out, each adjusting for one of three baseline variables commonly associated with smoking in pregnancy (heaviness of smoking, partner’s smoking status and education),31 32 with likelihood ratio tests assessing whether these improved model prediction. Where convergence of a model could not be achieved due to low event rates within small centre sites, these centres were merged to overcome the issue.

An intention-to-treat (ITT) analysis was used, with all participants analysed within the treatment group to which they were randomised and, where missing outcome data, were assumed smoking.27 Participants who withdrew from the study due to miscarriage/stillbirth were included in the analyses and classed as smoking. Where validation of abstinence was required, participants not providing a breath or saliva sample were classed as smoking. Complete case sensitivity analyses were performed on all smoking outcomes.

The number of quit attempts since baseline was compared between groups using a Mann-Whitney U test. Participants’ views on the MiQuit intervention were reported using percentages with 95% Wilson score CIs. Analyses were carried out in Stata, Version 12.

After undertaking the planned analyses, we decided to generate a Bayes Factor from smoking outcome #1, using an online calculator33 with an expected effect size of OR 1.83 taken from a relevant systematic review.12 We used a conservative approach for estimation using a half normal distribution, where the mode at 0 indicated no intervention effect, and the standard deviation equal to the expected effect size.

# Economic analysis

# The main outcome was the incremental cost per additional quitter, calculated by dividing the average incremental cost per participant by the number of additional quitters derived from smoking outcome #1. Confidence intervals were generated using bootstrapping with 1,000 iterations.34

# RESULTS

Over 7 months, we assessed 1181 pregnant smokers for eligibility and 407 were recruited into the study; 203 were randomised to MiQuit and 204 to usual care. There was marked variation in recruitment between the 16 sites (median 12 participants, IQR 34), with one recruiting no participants. Figure 1 shows participant flow and reasons for exclusion. At 4 weeks, 295 (72%) participants provided smoking outcome data (68% MiQuit, 77% usual care). Further attrition in late-pregnancy was fairly minimal, with 261 (64%) participants providing these outcome data (64% MiQuit, 65% usual care). 230 (57%) provided smoking outcome data at both time points (55% MiQuit, 58% usual care) and 254 (62%) gave data used for smoking outcome #1 on abstinence between 4 weeks and late pregnancy (61% MiQuit, 64% usual care). We obtained validation samples for 37/64 (58%) of participants who reported abstinence at 36 weeks gestation (56% MiQuit, 61% usual care); with two (3.1%) and 15 (23%) participants providing only CO or cotinine readings respectively.

*Figure 1 here*

Table 1 shows baseline participant characteristics by trial arm; mean age was 26.5 (SD 5.7) years, 92% were White, mean gestation at enrolment was 14.7 (SD 4.4) weeks, and 60% reported smoking within 30 minutes of waking. 74% were very or extremely determined to stop smoking and 40% felt very or extremely confident in stopping until their baby was born. Participants’ characteristics were similar in both groups apart from that women randomised into the usual care group were more likely to reside in the most deprived (e.g. lower income) areas and have a non-smoking partner.

*Table 1 here*

**Smoking outcomes**

Table 2 shows cessation rates across and within treatment groups and provides estimates for MiQuit’s effects. For smoking outcome #1, 15 participants were classified as abstinent; 11/203 (5.4%) were in the MiQuit group and 4/204 (2.0%) in the usual care group (adjusted OR 2.70, 95% CI 0.93 - 9.35). Estimated treatment effects for the remaining smoking outcomes also favoured MiQuit aiding smoking cessation, with ORs ranging from 1.03 to 3.28; those for self-reported abstinence at both 4 weeks post-randomisation and in late pregnancy (smoking outcome #6) reached statistical significance. Adjusting for heaviness of smoking, partner’s smoking status and education did not result in any meaningful changes to the findings (see supplementary Table S1). In a sensitivity analysis based on women with complete outcome data, the ORs were increased for six out of the seven smoking outcomes, including outcome #1 (OR 3.11, 95% CI 1.05 to 10.80) (Table S2). The number of quit attempts between baseline and late pregnancy did not differ significantly between treatment groups (MiQuit median 2 [IQR 1,3], N=124; usual care median 1 [IQR 0,3], N=130; Mann-Whitney U *p*=0.118). The Bayes Factor for outcome #1 was 2.23, meaning that the hypothesis that MiQuit is effective is more than twice as likely to be correct than the hypothesis that it is not effective. This represents ‘anecdotal evidence’ for MiQuit having an intervention effect.35

*Table 2 here*

**Use of NHS cessation support**

Overall use of ‘non-trial’ cessation support was similar in both arms (Table 3). When examining specific types of support, midwife discussion of smoking was reported by notably more usual care participants.

*Table 3 here*

**Participant evaluations of MiQuit**

Among all MiQuit participants, 27 (13%) discontinued support early (mean days into programme 24.1, SD 15.7) having texted STOP and 13 (6.4%) changed their message frequency to “less”, 11 (5.4%) to “more”, and 1 (0.5%) to “less” followed by “more”. Among those at late-pregnancy follow-up who answered the relevant questions, 3/123 (2.4%) reported receiving no text messages and, of the remaining 120, 97 (81%) reported reading all messages at least once. Messages relating to fetal development were most frequently rated (by 35%) as the most helpful. Table 4 shows that 62% rated the text messages as quite or extremely helpful but 14% considered them annoying. 81% would either ‘probably' or ‘definitely’ recommend MiQuit support to a friend or relative.

*Table 4 here*

**Economic analysis**

The per-participant cost of sending MiQuit texts was estimated to be £2.95; a mean of 84.1 texts per participant at 3.5p per text. The annual running cost of delivering MiQuit was £339 (£1.67 per participant) and included a virtual reply number (£99) and server/web hosting including domain name (£240). Thus, the total per-participant MiQuit cost was £4.62. From Table 2, row 1, the relevant incremental quit rate estimate was 3.46%, giving an incremental cost per additional quitter of £133.53 (95% CI -£395.78 to £843.62). The probability of MiQuit being cost-effective was 96.5% if a decision maker was willing to pay £10,000 to gain an additional quitter.

**DISCUSSION**

**Statement of principal findings**

This trial demonstrates the feasibility of recruiting pregnant smokers from multiple UK hospital antenatal settings to a trial of a text message cessation support intervention; we met our recruitment target 5 months earlier than expected. We also found that it was feasible to measure smoking cessation in participants who were not expected to set a quit date using a stringent outcome measure. Using this outcome, we found that 5.4% of women in the MiQuit group stopped smoking during pregnancy and 2.0% did in the control group and this almost reached statistical significance; it is likely, when tested on a larger scale, that MiQuit will prove to be both effective and cost effective for promoting smoking cessation throughout pregnancy.

**Findings in context**The efficacy estimate provided using outcome #1 data suggests that, compared with usual care, MiQuit may almost triple the odds of sustained smoking cessation, but this has limited precision. However, it is the best estimate yet produced for the likely efficacy of text messaging used for smoking cessation in pregnancy. It is also of a similar magnitude to efficacy estimates derived from ‘definitive’ trials of similar interventions used by non-pregnant smokers36 37 and to that from a smaller MiQuit trial.18 Additionally, our estimate for the likely cost-effectiveness of the intervention is encouraging; compared with other cessation interventions a cost-per-quitter of £134 is low. For example, although financial incentives for smoking cessation in pregnancy are highly effective38 and cost-effective,39 their cost-per-quitter is almost 10 times higher (£1,127). Similarly, MiQuit's cost-efficacy compares favourably with that of cessation support delivered by traditional UK smoking cessation services; the ‘cost-per-person-setting-a-quit-date’ within such services has recently been estimated as £202.40 However, as only 34% of those setting a quit date achieve longer-term abstinence, the cost-per-quitter,41 inflated accordingly, is probably closer to £600. Although the trial did not include a formal cost-utility analysis, it is highly likely that, if cessation is maintained in the longer term, the calculated ‘cost-per-quitter’ will translate into longer-term cost effectiveness. One can assume that ‘quitters’ gain 1.94 Quality-Adjusted Life Years (QALYs) across their lifetime,42 43 so by multiplying this value by the seven additional quitters generated by MiQuit the incremental QALYs would be 13.58, making the incremental cost per additional QALY £69.06 – even after inflating this figure to take into account relapse to smoking,44 this would remain securely within most accepted cost-effectiveness benchmarks. Finally, it is noteworthy that the ‘non-text-message’ costs of MiQuit are fixed and so ‘per-user’ costs fall as the numbers using the intervention increase. For example, if MiQuit was used by 2000 pregnant smokers annually, per-user ‘non-text-message’ costs would be around 20p, reducing the incremental cost per additional quitter to approximately £91.

Importantly, systems such as MiQuit could be particularly useful for the high proportion of pregnant smokers who currently do not access ‘traditional’ methods of support.45 46 For example, in the UK around 83% of pregnant smokers do not use support offered46 but, if encouraged, many of these may use text support.

**Strengths and limitations of the study**

A limitation is that this RCT did not have a specified primary outcome; however, although multiple cessation outcomes were used, we indicated a priori which was anticipated to be the most appropriate as a primary outcome (outcome #1).19 Consequently, as we demonstrated that outcome #1 was feasible to measure, it is reasonable to use these data to represent MiQuit’s likely treatment effect. However, caveats to the interpretation of non-primary RCT outcomes still apply. Additionally, completeness of follow up and biochemical validation rates were not optimal, potentially reducing statistical power. However, we conservatively assumed that women lost to follow up were still smoking27 and outcome ascertainment rates were slightly higher in the usual care group; both factors would tend to attenuate rather than inflate the observed intervention effect. Consistent with this observation, the complete case analyses showed stronger intervention effects for most smoking outcomes, including a statistically significant between-group difference for cessation outcome #1. As with many RCTs, a further limitation is the unknown generalisability of findings to all pregnant smokers. We did not systematically record data on the numbers or characteristics of pregnant smokers attending hospital units during trial recruitment, so we cannot say how representative the trial sample is; although, based on socio-economic characteristics and smoking rates at pre-pregnancy and baseline, the sample was generally representative of women who smoke in pregnancy and are recruited to trials.10 Ease of recruitment in antenatal care settings suggests there is a substantial cohort of pregnant smokers who would be likely to use MiQuit if offered this as part of routine care. Moreover, we have already shown that 3-4% of pregnant smokers will initiate MiQuit after receiving a one-page leaflet advertising this in their ‘antenatal booking pack’.22

A key strength is that this is the largest RCT to investigate the efficacy of text message-based, self-help cessation support which is appropriate for and can be safely followed by pregnant smokers. The study was conducted to the highest RCT standards; it employed remote randomisation, those enrolling participants were blind to treatment allocations and abstinence was biochemically validated. Additionally, researchers collecting outcome data were, where possible, blind to treatment allocations, so outcome ascertainment bias was minimised. Intervention fidelity was high, 98% of MiQuit recipients recalled receiving text message support, and similarities between adjusted and unadjusted analysis models imply that chance differences in groups’ baseline characteristics do not explain findings. Similarly, it seems unlikely that use of other ‘non-trial’ cessation support explains findings; use of such support was very similar in both groups except that usual care group women were more likely to report having discussed smoking with a midwife. Such support would be expected to increase cessation in the usual care group, reducing the apparent efficacy of MiQuit. Overall, therefore, it seems likely that differences between groups’ smoking rates are due to MiQuit and not to other factors.

**Conclusions**
MiQuit is likely to be an effective smoking cessation intervention and further evaluative research is needed. If further research is confirmatory, pregnancy-orientated text message systems like MiQuit could quickly and cheaply be made available alongside other first-line support options to help pregnant smokers to stop.

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# Author contributions

FN designed and developed the MiQuit intervention, contributed to the conception and design of the study and helped draft and edit the protocol. SC contributed to the conception and design of the study, drafted and edited the protocol and participated in the coordination of the trial. KF contributed to the drafting and editing of the protocol and coordinated the study as trial manager. JE performed the statistical analysis of the study. JLB contributed to the conception and design of the study, helped draft and edit the protocol and is the trial statistician. SS designed and developed the MiQuit intervention and contributed to the conception and design of the study. MJ performed the health economic analysis of the study. MU contributed to the conception and design of the study, and helped draft and edit the protocol. RW participated in the coordination of the trial. ML contributed to the design of the study and helped draft the protocol. AM contributed to the design of the study and was involved in the development of the statistical analysis. SP contributed to the design of the study and was involved in the development of the health economics analysis. TC conceived and designed the study, drafted and revised the protocol, and is the trial’s chief investigator. FN, KF, JE and TC drafted the manuscript and all authors read, commented on and approved the final manuscript.

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**Figure 1** Trial flow

**Excluded** (n= 774)

|  |  |
| --- | --- |
| 1. Smokes < 1 per day now | 44 |
| 2. Smoked < 5 per day pre-pregnancy | 6 |
| 3. In another text service | 4 |
| 4. Not able to understand English | 8 |
| 5. > 25 weeks gestation | 82 |
| 6. Unwilling to consent | 14 |
| 7. Not interested in study | 363 |
| 8. Other | 167 |
| 9. Unknown | 86 |

**Assessed for eligibility**

(n=1181)

**Randomised**

(n=407)

**Allocated to usual care arm** (n=204)

**Allocated to MiQuit arm**

(n=203)

**Did not complete 4 week FU** (n=47)

Lost to follow up (n=47)

Withdrew consent (n=0)

Fetal death (n=0)

**Did not complete 4 week FU** (n=65)

Lost to follow up (n=61)

Withdrew consent (n=2)

Fetal death (n=2)

**Abstinence data collected at 4-week follow up**

(n=157)

**Abstinence data collected at 4-week follow up**

(n=138)

**Did not complete 36-week gestation FU** (n=26)

Lost to follow up (n=23)

Fetal death (n=3)

**Did not complete 36-week gestation FU** (n=39)

Lost to follow up (n=34)

Fetal death (n=3)

Withdrew consent (n=1)

Unknown reason (n=1)

**Abstinence data collected at 36 weeks gestation**

(n=132)\*\*

**Abstinence data collected at 36 weeks gestation**

(n=129)\*

**Analysed**

(n=204)

**Analysed**

(n=203)

\*Includes 17 MiQuit participants without 4-week follow up data

\*\*Includes 14 usual care participants without 4-week follow up data

**Table 1** Baseline characteristics by treatment group

|  |  |  |
| --- | --- | --- |
|  | **MiQuit†**(N=203) | **Usual Care†**(N=204) |
| **Age (years)** Mean[SD]Median[1st Q, 3rd Q]Min, max | 26.6 (5.7)25.7 (22.1, 30.8)16.9, 40.0 | 26.4 (5.7)25.8 (21.9, 29.7)16.6, 41.3 |
| **Highest qualification** No formal qualification GCSE or similar A level or similar Degree or similar Declined to answer | 37 (18.2)117 (57.6)32 (15.8)16 (7.9)1 (0.5) | 44 (21.6)106 (52.0)37 (18.1)13 (6.4)4 (2.0) |
| **IMD score\***Quintile 1Quintile 2Quintile 3Quintile 4Quintile 5missing | 13 (6.4)16 (7.9)22 (10.8)53 (26.1)92 (45.3)7 (3.5) | 6 (2.9)13 (6.4)21 (10.3)50 (24.5)108 (52.9)6 (2.9) |
| **Ethnicity** White Indian Pakistani Bangladeshi Black Caribbean Black African Black(other)Chinese Other Asian (non-Chinese)Mixed Not given | 188 (92.6)0 (0)3 (1.5)0 (0)1 (0.5)2 (1.0)1 (0.5)0 (0)1 (0.5)6 (3.0)1 (0.5) | 185 (90.7)0 (0)2 (1.0)0 (0)4 (2.0)1 (0.5)1 (0.5)0 (0)0 (0)11 (5.4)0 (0) |
| **Gestation at randomisation (weeks)**Mean[SD] Median[1st Q, 3rd Q] Min, max | 14.6 (4.2)13 (12, 19)4, 23 | 14.7 (4.5)13 (12, 20)3, 24 |
| **Cigarettes per day before pregnancy** Mean[SD] Median[1st Q, 3rd Q] Min, max | 15.7 (6.7)15 (10, 20)5, 40 | 16.4 (6.6)15 (10, 20)5, 40 |
| **Cigarettes per day now** Mean[SD] Median[1st Q, 3rd Q] Min, max | 9.0 (5.9)8 (5, 10)1, 40 | 9.4 (6.1)10 (5, 10)1, 40 |
| **Time to first cigarette after waking**Within 5 minutes 6-30 minutes 31-59 minutes 1-2 hours More than 2 hours | 64 (31.5)56 (27.6)41 (20.2)22 (10.8)20 (9.9) | 64 (31.4)61 (29.9)31 (15.2)29 (14.2)19 (9.3) |
| **Frequency of urges to smoke in the past 24 hours** Not at all A little of the time Some of the time A lot of the time Almost all the time All the time | 3 (1.5)36 (17.7)94 (46.3)44 (21.7)16 (7.9)10 (4.9) | 8 (3.9)37 (18.1)88 (43.1)42 (20.6)18 (8.8)11 (5.4) |
| **Strength of urges to smoke in the past 24 hours** No urges Slight Moderate Strong Very strong Extremely strong | 4 (2.0)58 (28.6)78 (38.4)39 (19.2)15 (7.4)9 (4.4) | 6 (2.9)55 (27.0)95 (46.6)28 (13.7)14 (6.9)6 (2.9) |
| **Have you set a quit date?** No Yes | 193 (95.1)10 (4.9) | 192 (94.1)12 (5.9) |
| **Are you seriously planning to quit?**NoWithin the next 3 monthsWithin the next 30 daysWithin the next 2 weeks | 17 (8.4)68 (33.5)55 (27.1)63 (31.0) | 19 (9.3)57 (27.9)59 (28.9)69 (33.8) |
| **Number of births beyond 24 weeks** Mean[SD] Median[1st Q, 3rd Q] Min, max | 1.4 (1.5)1 (0, 2)0, 10 | 1.4 (1.4)1 (0, 2)0, 9 |
| **Parity** 0 births beyond 24 weeks 1 or more births beyond 24 weeks | 66 (32.5)137 (67.5) | 65 (31.9)139 (68.1) |
| **Partner/significant other’s smoking status** Smoker Non-smokerNot applicable (no partner) | 135 (66.5)34 (16.8)34 (16.8) | 128 (62.8)44 (21.6)32 (15.7) |

Data are N (%) unless specified.

\* Index of Multiple Deprivation, Office for National Statistics. Quintile 1 represents least deprivation

†Data was complete for all baseline variables other than Index of Multiple Deprivation (IMD) score (3.2% missing: no match to home postcode), highest qualification (1.2% missing) and ethnicity (0.25% missing). Similar proportions per trial arm were missing baseline data.

**Table 2** MiQuit treatment effect estimates on seven smoking outcomes

|  |  |  |  |  |  |  |
| --- | --- | --- | --- | --- | --- | --- |
| **Outcome** | **Measure** | **MiQuit\***N=203 (%) | **Usual Care\***N=204 (%) | **Total\***N=407 (%) | **P value\*\*** | **Adjusted Odds Ratio (95% CI)\*\*\*** |
| **Abstinence reported from 4 weeks post-randomisation until late pregnancy (smoking outcome #1) †** | Validated | 11 (5.42) | 4 (1.96) | 15 (3.69) | 0.064 | 2.70 (0.93-9.35) |
| Abstinence reported from 4 weeks post-randomisation until late pregnancy (smoking outcome #2) † | Self-report | 33 (16.26) | 33 (16.18) | 66 (16.22) | 0.983 | 1.03 (0.61-1.75) |
| 7-day point prevalence abstinence at late pregnancy (smoking outcome #3) | Self-report | 36 (17.73) | 28 (13.73) | 64 (15.72) | 0.267 | 1.34 (0.79-2.31) |
| 7-day point prevalence abstinence at late pregnancy (smoking outcome #4) | Validated | 15 (7.39) | 9 (4.41) | 24 (5.90) | 0.202 | 1.67 (0.72-4.03) |
| 7-day point prevalence abstinence at 4 weeks post-randomisation (smoking outcome #5) | Self-report | 15 (7.39) | 7 (3.43) | 22 (5.41) | 0.077 | 2.11 (0.89-5.46) |
| 7-day point prevalence abstinence at both 4 weeks post-randomisation and late pregnancy(smoking outcome #6) | Self-report | 13 (6.40) | 4 (1.96) | 17 (4.18) | 0.025 | 3.16 (1.14-10.69) |
| 7-day point prevalence abstinence at both 4 weeks post-randomisation and late pregnancy(smoking outcome #7) | Validated | 8 (3.94) | 2 (0.98) | 10 (2.46) | 0.062 | 3.28 (0.90-17.36) |

\* All smoking outcomes are calculated out of 407 participants in total (203 MiQuit, 204 usual care). Participants lost to follow up or with missing outcome data are assumed to be smoking.

\*\* Unadjusted, from a chi-squared test using a 2-sided p value (Fisher’s exact test p values were used in the case of small expected frequencies).

\*\*\* Model-based, adjusted by site and gestation at randomisation (95% profile confidence intervals reported).

† Russell Standard criterion (permits no more than 5 cigarettes in total). The criterion for all other smoking outcomes was total abstinence (‘not even a puff’).

**Table 3** Use of NHS and other cessation support during trial period

|  |  |  |
| --- | --- | --- |
| **Outcome\*** | **MiQuit**(N=124) | **Usual Care**(N=130) |
| Reported use of any stop smoking support N (%) | 83 (66.9) | 98 (75.4) |
| Reported use of different types of support N (%): |  |  |
|  | GP or nurse discussion | 20 (16.1) | 26 (20.0) |
|  | Midwife discussion | 45 (36.3) | 72 (55.4) |
|  | Stop smoking helpline | 5 (4.0) | 6 (4.6) |
|  | NHS Smokefree website | 16 (12.9) | 15 (11.5) |
|  | Other smoking cessation website | 7 (5.7) | 9 (6.9) |
|  | NRT | 26 (21.0) | 36 (27.7) |
|  | Individual NHS behavioural support | 9 (7.3) | 15 (11.5) |
|  | Group NHS behavioural support | 3 (2.4) | 3 (2.3) |

\* Outcomes are calculated out of 254 participants with response data at late pregnancy follow up (124 MiQuit, 130 usual care)

**Table 4** Interventionparticipant views of and preferences for the MiQuit intervention

|  |  |
| --- | --- |
|  | **MiQuit** |
| Reported receiving text messages (n=123) | 120 (97.6, 93.1-99.2) |
| Discontinued the support prematurely by texting ‘STOP’ (n=203) | 27 (13.3, 9.3-18.7) |
| Rated the text messages as ‘quite’ or ‘extremely’ helpful (n=120) | 74 (61.7, 52.7-69.9) |
| Rated the text messages as ‘quite or ‘extremely’ annoying (n=120) | 17 (14.2, 9.0-21.5) |
| Rated the number of text messages received as (n=120): |  |
|  | ‘far too many’ or ‘a little too many’ | 25 (20.8, 14.4-29.2) |
|  | ‘about right’ | 79 (65.8, 56.8-73.9) |
|  | ‘not enough’ or ‘not nearly enough’ | 16 (13.3, 8.3-20.8) |
| Would ‘probably’ or ‘definitely’ recommend the support (n=120) | 97 (80.8, 72.8-86.9) |

Data are N (%, 95% Wilson CI)