Evaluation of the digital diabetes prevention programme pilot: uncontrolled mixed-methods study protocol

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ABSTRACT

Introduction The prevalence of type 2 diabetes is rising steeply. National Health Service England (NHSE) is exploring the potential of a digital diabetes prevention programme (DDPP) and has commissioned a pilot with embedded evaluation.

Methods and analysis This study aims to determine whether, and if so, how, should NHSE implement a national DDPP, using a mixed-methods pretest and post-test design, underpinned by two theoretical frameworks: the Coventry, Aberdeen and London - Refined (CALO-RE) taxonomy of behavioural change techniques for the digital interventions and the Consolidated Framework for Implementation Research (CFIR) for implementation processes. In eight pilot areas across England, adults with non-diabetic hyperglycaemia (NDH) (glycated haemoglobin (HbA1c) 42–47 mmol/mol or fasting plasma glucose 5.5–6.9 mmol/L) and adults without NDH who are overweight (body mass index (BMI) >25 kg/m²) or obese (BMI >30 kg/m²) will be referred to one of five digitally delivered diabetes prevention interventions. The primary outcomes are reduction in HbA1c and weight (for people with NDH) and reduction in weight (for people who are overweight or obese) at 12 months. Secondary outcomes include use of the intervention, satisfaction, physical activity, patient activation and resources needed for successful implementation. Quantitative data will be collected at baseline, 6 months and 12 months by the digital intervention providers. Qualitative data will be collected through semistructured interviews with commissioners, providers, healthcare professionals and patients. Quantitative data will be analysed descriptively and using generalised linear models to determine whether changes in outcomes are associated with demographic and intervention factors. Qualitative data will be analysed using framework analysis, with data pertaining to implementation mapped onto the CFIR.

Ethics and dissemination The study has received ethical approval from the Public Health England Ethics and Research Governance Group (reference R&D 324). Dissemination will include a report to NHSE to inform future policy and publication in peer-reviewed journals.

INTRODUCTION

Diabetes is a national and international priority for health services, with a steeply rising prevalence. Globally, it affects over 400 million people or around 9% of the adult population, and of these, over 90% have type 2 diabetes mellitus (T2DM). There were an estimated 3.7 million people with diabetes in England in 2016–2017. People with diabetes are at risk of complications including cardiovascular disease, nephropathy, retinopathy and neuropathy. The costs of treating diabetes and its complications are estimated at 10% of the total National Health Service (NHS) budget, or some £10 billion per annum.

Risk factors for developing T2DM include diet, lack of physical activity, obesity, genetic factors and deprivation. For many people, T2DM is a preventable illness, with prevention focused on the modifiable factors of diet, physical activity and weight. There is high-quality international evidence that face-to-face programmes focusing on these three factors can reduce the rate of progression to T2DM in high-risk individuals. To be successful, these programmes appear to require intensive sustained input over a prolonged period of time. In light of this...
evidence NHS England (NHSE) launched Healthier You: The NHS Diabetes Prevention Programme (NHS DPP) in 2015, initially in seven demonstrator sites, and subsequently rolled out across England. All programmes within the NHS DPP must offer at least 16 hours of face-to-face contact, spread over 13 sessions, with the total programme lasting at least 9 months.16

NHSE is also considering a national digital diabetes prevention programme (DDPP), and in order to generate the evidence to inform future policy in this area, launched a pilot DDPP in 2017, called ‘Healthier You: NHS DPP digital stream’.17 The reasons cited by NHSE for exploring the potential of a digital DPP are twofold: first, digital delivery may overcome some of the challenges affecting face-to-face programmes; and second, systematic review evidence that digital health interventions can be effective in increasing physical activity, changing diets and promoting weight loss,18-27 all behaviours which are effective in preventing T2DM. Challenges affecting the population impact of face-to-face programmes include problems of acceptability, as their intensive nature may make it difficult for people who work or have other commitments in their lives to attend; and there may be perceived stigma in attending a programme aimed at prevention of T2DM. Finally, the face-to-face programmes are costly, particularly when implemented at scale and a digital programme could potentially be easier to deliver at scale and more cost-effective.

However, there is little evidence to support these potential advantages,28 and there are some well-documented challenges in the delivery of digital health interventions. Three of the most important of these challenges are problems with engagement and adherence; concerns around the ‘digital divide’; and well-documented problems with implementation. Many digital health interventions show low rates of initial uptake as well as high rates of subsequent attrition from the intervention, which limits their potential for population impact.29 Moreover, there is uncertainty as to how best to improve engagement and adherence—although there are data which suggest that a certain amount of human input, for example, in the form of supportive or coaching telephone calls, can improve engagement, the requirement for human input can impact on the scaleability and costs of digital health interventions.30-32 There is real concern that the ‘digital divide’ (the divide between those who do and do not make regular use of digital technology) will exacerbate health inequalities, as many of those with greatest health needs (older people, people with long-term health problems and those with low incomes) are also people who make less use of digital technology.33-37 Third, the challenges of successful implementation of digital health interventions are well known,38 39 with few examples of successful integration of digital health interventions into routine healthcare and considerable uncertainty as to how best to achieve such integration.

In the light of this potential, coupled with these major areas of uncertainty, NHSE commissioned a pilot DDPP with an associated evaluation to run alongside the national DPP. In the initial tender document, NHSE specified that the goal of the pilot and associated evaluation was to inform future policy in this area, and as such, the overarching aim of the evaluation was to determine: ‘Whether, and if so, how, should NHSE roll out a national DDPI at the end of the pilot?’ Specific areas of interest were around uptake, use and impact on weight and glycated haemoglobin (HbA1c); the likely impact of a digital programme on health inequalities; and how the interventions should be integrated into NHS workflows, including determining the resource requirements for successful implementation. Although a formal comparison of the effectiveness of different interventions was out of scope, NHSE was interested in whether specific features of different interventions appeared to be associated with variation in observed uptake, use or impact. The tender specified that a formal health economic analysis was out of scope. The evaluation is due to report in 2020.

The specific objectives of the evaluation can be grouped into three areas: uptake, use and impact of the interventions; the extent to which uptake, use and impact vary by socioeconomic status (SES) as an indicator of likely impact on health inequalities; and factors relating to implementation.

Objectives pertaining to uptake, use and impact:
1. Determine uptake and use of the digital diabetes prevention interventions (DDPI) by people referred to them through the DDPP.
2. Determine the effects of DDPI on people referred to them through the DDPP.
3. Explore the extent to which these benefits vary according to differences in key features of the selected DDPI.
4. Explore user views about the acceptability of DDPI, including perceptions relating to use/non-use and impact on relevant behaviours.

Objectives pertaining to health inequalities:
5. Explore the extent to which uptake, use and effects vary by SES.

Objectives pertaining to implementation:
6. Describe the various implementation strategies applied in the eight demonstrator sites.
7. Determine the costs associated with implementing and delivering a DDPP, from an NHS perspective.
8. Explore commissioner, healthcare professional and provider views about key factors influencing implementation, uptake and impact of the DDPI.
9. Summarise and synthesise these data in a report which can be used to inform the policy decisions about whether, and if so, how, to roll out a DDPP across England.

METHODS

Design

Mixed-methods pretest and post-test design, underpinned by theory.
Theoretical frameworks
This evaluation will be underpinned by two theoretical frameworks: one pertaining to the effectiveness of the DDPI and one pertaining to the implementation processes.

Understanding the likely and observed effectiveness of the selected DDPI will be promoted by applying the Coventry, Aberdeen and London - Refined (CALORE) taxonomy of behaviour change techniques and describing interventions using the Template for Intervention Description and Replication (TiDIER) framework (online supplementary appendix 1).

To help with describing and understanding the implementation processes, we will use the Consolidated Framework for Implementation Research (CFIR). This specifies that the key constructs which determine whether an intervention is successfully implemented or not are: intervention characteristics; outer setting; inner setting; individual characteristics and the implementation process (for details see online supplementary appendix 2).

Patient and public involvement
The board overseeing the NSHE programme in diabetes and diabetes prevention is made up of a triumvirate of NHSE, Public Health England (PHE) and Diabetes UK (DUK). DUK is the largest charity representing the voice of people with diabetes in the UK. Patient and public involvement (PPI) in this study was, therefore, provided by DUK, through their membership of the board. This board determined how the DDPP should be piloted and evaluated, what the requirements for the evaluation were, including overall design (numbers of demonstrator sites, DDPI, pretest and post-test design, primary outcomes and duration of study). The board also provides oversight of the conduct and progress of the study and will receive the reports of the study. Hence, there was PPI input into determining the research questions, outcome measures, study design and dissemination.

Setting
Eight demonstrator sites were selected by NHSE in parallel with the selection of the evaluator team, who had no input into site selection. Sites volunteered to participate in the digital pilot, and were selected to achieve a range of geographies and demographics, including rural, semirural, urban and metropolitan areas, with widely varying proportions of people from black and minority ethnic backgrounds, SES and pre-existing levels of digital readiness and engagement with diabetes prevention.

Populations and participants
There were three populations specified by NHSE: (1) adults with non-diabetic hyperglycaemia (NDH) defined as having had an Hba1c measurement of 42–47 mmol/mol, or a fasting glucose measurement of 5.5–6.9 mmol/L, in the 12 months prior to referral; (2) adults who are overweight (body mass index (BMI) of 25 to <30 kg/m²) without NDH and (3) adults who are obese (BMI of 30 kg/m² or over) without NDH. It is the responsibility of the eight demonstrator sites to determine how they will identify people who fell into these three populations, how general practitioner (GPs) and patients will be informed about the programme, and how patients will be referred to a DDPI. In most sites, patients will be referred to the interventions by their healthcare professionals, usually their GPs. In some areas, identification and referral may be undertaken by alternative providers contracted to undertake assessment of cardiovascular risk as part of the NHS Health Check Programme, which includes a two-stage assessment that aims to identify NDH and undiagnosed T2DM. Referring healthcare professionals are responsible for discussing the referral with patients, to ensure patients understand their diagnosis, the type of intervention they are being referred to and the expected benefits. Referring healthcare professionals are responsible for sending the patient’s name and contact details to the relevant DDPI provider; that provider is then responsible for contacting the patient and onboarding them to the intervention. These processes are identical to those used in the national face-to-face DPP, with the only difference being that the provider is offering a digital, rather than a face-to-face intervention.

Interventions
A total of five DDPI were selected following a multistage selection process undertaken by an independent assessor (Our Mobile Health) under contract to NHSE. Following widespread advertising of the opportunity, 84 companies registered an interest and underwent initial screening against six criteria. These were: (1) that the intervention supported behavioural change; (2) was available by June 2017; (3) was localised for the English market; (4) was underpinned by an evidence-based approach; (5) did not require any further integration with existing health information technology (IT) systems before launching; (6) had a pricing system in place (although cost itself was not a criterion). Thirty providers met these criteria and progressed to a self-assessment exercise which explored eight criteria: safety; privacy and security; pricing; evidence based or indicators of effectiveness; usability and accessibility; technical stability; change management and interoperability. The self-assessment questionnaire can be found at https://developer.nhs.uk/daq. This led to a shortlist of 14 interventions, which were reviewed by subject matter experts including behaviour change theory experts, clinical safety officers, GPs, diabetologists, diabetes specialist nurses and dieticians. The final sample was selected to vary on factors known to be important in influencing uptake, use and effectiveness of digital health interventions, namely: the delivery platform (smartphone vs not); the amount of human interaction to promote uptake; and inclusion (or not) of wearables. Within this, interventions with better pre-existing evidence for uptake, use and effectiveness as well as capabilities and infrastructure to implement at scale, were prioritised.
All five interventions focused on dietary intake, weight loss and physical activity, and all interventions set personalised goals and provided feedback on progress towards these goals.

Intervention A (Buddi Nuijer) is a smartphone app, which links to a wristband for monitoring physical activity. The participants log their eating habits and activities via the app, receive three motivational messages from the app per day and complete a total of 22 learning modules over the course of the 12-month programme. Onboarding is done by an initial phone call and email for registration. A customer services team is available for those who have technical difficulties, but apart from that, the service is entirely digital.

Intervention B (Hitachi) is smartphone (Android and Apple), tablet and desktop compatible. The solution provides a hybrid of digital and non-digital interactions with a website and a series of phone calls with an experienced health advisor, whose advice follows NHS guidelines. Participants and health advisors set an action plan at the start of the programme. Participants then self-report their outcomes and log their lifestyle on the website to understand their progress against key milestones and objectives. This information goes back to the health advisor team who prompt on guidance and can interact with participants. Health advisors contact participants monthly for the first 6 months and have one more step-down call at the 8-month mark. Pedometers and scales can be provided on request. Onboarding is done through an initial brief phone call, at which point a longer goal setting telephone call is arranged.

Intervention C (Liva) is smartphone (Android and Apple), tablet and desktop compatible. There is an initial face-to-face appointment with a health coach for onboarding and goal setting, followed by 26 coaching sessions with the same coach, delivered weekly for the first 12 weeks and then tapering off in frequency. There is an online peer support group.

Intervention D (Ourpath) is smartphone (Android and Apple), tablet and desktop compatible and includes wireless weighing scales and a wearable activity tracker. Participants are entered into a peer group of up to 10 other people with similar goals who live locally. Groups interact by group messaging, and group targets are set as well as individual ones. The programme has three stages: the ‘Core’ programme with daily education content received through the app for the first 6 weeks, the ‘Sustain’ programme with weekly education content until the 6-month mark and the final 6 months where the user will have completed all the education modules but still have access to the dietician and group support. Onboarding is done through two phone calls—an initial introductory one, and a second one to set the participant up with the programme and group.

Intervention E (Oviva) is an app (Android and Apple compatible) with supporting material (learning materials, podcasts, recipes) delivered through an online portal. For patients without a smartphone, the content can be delivered via phone calls. The app allows users to track their activity, weight, and food and drink intake (using a photo food diary). The programme is a mix of digital and non-digital interactions with a series of phone calls accompanying the app. The programme is more intense at the start with a weekly phone call in the first 8 weeks to cover the 16-topic curriculum, tapering to a monthly phone call thereafter. The phone calls are all conducted by the same dietician who is a specialist diabetes dietician with at least 2 years experience. Onboarding is done by the dietician in the first phone call.

The interventions will be described according to the TiDIER Framework for describing complex interventions and the CALO-RE Behaviour Change Technique taxonomy.

**Outcomes**

The primary outcomes were prespecified by NHS England, and were change between baseline and 12 months in Hba1c for the population with NDH and weight for all three populations. Changes over 6 months are considered secondary outcomes.

Secondary and explanatory outcomes have been selected according to our pathway of action model, which posits that the DDPI achieves its goal of reducing a user’s risk of diabetes by promoting behavioural change, specifically, promoting dietary change and an increase in physical activity. Taken together these behavioural changes result in reduced Hba1c and reduced weight. To achieve these changes requires the user to: register with the intervention, use the intervention, initiate behaviour change and sustain behaviour change. Effects at each stage will be moderated by intervention factors and by patient factors (see figure 1). The context and implementation process will also affect the overall population uptake and impact.

Secondary outcomes are listed in table 1 and include intervention uptake, use and satisfaction; behavioural and clinical outcomes; and costs related to implementation, although it should be noted that a formal health economic analysis was excluded from the brief. Data will be collected at baseline, 6 months and 12 months.

Measures used are the Friends and Family Test (FFT) to measure satisfaction with care, the International Physical Activity Questionnaire (IPAQ) to measure physical activity and the 13-item Patient Activation Measure (PAM-13).

Demographic characteristics are collected at baseline only and will be used as explanatory factors. Demographic data to be collected includes age (date of birth), gender, ethnicity, postal code (to be used for determining SES by mapping against the Index of Multiple Deprivation (IMD)) and highest level of education attained.

**Qualitative and explanatory outcomes**

Reasons for observed differences in quantitative outcomes will be explored through qualitative interviews. These will be undertaken with a range of stakeholders, including commissioners/leads for the DDPP in selected...
local health economies (LHE); healthcare professionals (GPs, practice nurses, diabetes nurses, healthcare assistants, practice managers); digital diabetes intervention providers; the implementation teams charged with implementing the DDPP in the selected LHE and patients.

Interviewees will be selected through purposive sampling, aiming for variation across geographical area, digital diabetes intervention provider, disciplinary backgrounds and areas of high and low uptake. Recruitment of interviewees will continue until (1) all digital diabetes intervention providers and representatives from each demonstrator site have been interviewed and (2) data saturation is achieved.

Interviews will be conducted using semi-structured topic guides, by trained interviewers, working for RSM. The topic guides will vary according to the background of the stakeholder, with the main areas covered summarised in table 2. Interviews will be undertaken in waves, with wave 1 taking place during initial implementation and set-up, wave 2 once the programme is well established and wave 3 towards the end of the pilot.

Data collection
Referral forms will include basic demographic and clinical data, including HbA1c and weight recorded in the previous 3 months to confirm eligibility.

Quantitative data
The DDPI providers are responsible for collecting all quantitative data. Baseline data will be obtained from referral forms and supplemented with data obtained during onboarding interviews with patients. Follow-up self-report data (FFT, IPAQ and PAM-13) will be collected online. HbA1c and weights will be measured by the DDPI providers or by patients’ general practices, with the method and site of measurement recorded.

HbA1c measurements may be done on either venous or capillary blood samples, using either registered NHS laboratories or validated point-of-care testing kits. Whichever measurement process is used at baseline should be used at follow-up. Weights will be recorded using calibrated scales, with patients wearing light indoor clothing.

Data analysis
Quantitative data
The baseline characteristics of the three cohorts (NDH, overweight and obese) will be summarised with respect to sociodemographic characteristics, intervention uptake, behavioural and clinical outcomes and economic outcomes. Continuous data will be summarised in terms of the mean, SD and number of observations or, where skewed, median and IQR. Binary/categorical data will be summarised in terms of frequency counts and percentages. Descriptive statistics will also be used to explore differences in baseline characteristics across the eight demonstrator sites and five DDPIs.

The primary analysis will be based on participants with complete data. For continuous outcomes, the overall effectiveness of the programme will be assessed in pre–post analyses by comparing the mean outcomes in each cohort from baseline to 6 months, and from baseline to 12 months (presented with 95% CIs for the estimated change in outcomes). The statistical significance of any changes will be assessed using a paired t-test.
categorical outcomes, pre–post analyses will be conducted using McNemar’s test. Multivariable generalised linear models will be used to determine whether changes in outcomes are associated with demographic factors, adjusting for baseline outcome scores. Where necessary, continuous outcomes will be transformed to ensure good regression model fit. Exploratory analysis of the influence of potential mediators will be conducted by adding variables relating to usage and features of the DDPI to the regression models. The potential for clustering effects by demonstrator site will be considered by the inclusion of random effects for the demonstrator site in the generalised linear models. Three-level models accounting for clustering by GP practice within demonstrator sites will also be explored.

Reasons for missing data will be documented and the baseline characteristics of those with and without missing data compared. Although the primary analysis will be based on participants with complete data, we will undertake sensitivity analyses using various imputation models. The potential for bias due to non-random attrition will be addressed by fitting a propensity score model to account for drop-out on the basis of baseline characteristics and then using inverse probability weighting based on the propensity score to fit the treatment effectiveness model. No formal adjustment for multiple significance testing will be applied.

### Sample size

Target referral and registration numbers were preset by NHSE as part of the tender at 3500 registrations for the NDH population and 1500 for the overweight/obese population. We estimated minimum detectable effect sizes at 90% power and a 5% significance level for the key research questions, given these fixed sample sizes. Assuming a 25% completion rate (at 12 months), it will be possible to detect standardised effect sizes of $d=0.11$ and $d=0.17$ when assessing overall effectiveness.

<table>
<thead>
<tr>
<th>Table 1 Quantitative outcomes</th>
<th>Population</th>
<th>Time point for collection</th>
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<tbody>
<tr>
<td>Primary Outcome</td>
<td>HbA1c</td>
<td>Weight</td>
</tr>
<tr>
<td>NDH</td>
<td>Baseline</td>
<td>6 months</td>
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<tr>
<td>Overweight/obese</td>
<td>X</td>
<td>X</td>
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<tr>
<th>Secondary outcomes</th>
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<tr>
<td>Intervention factors</td>
</tr>
<tr>
<td>Amount of human support planned and delivered (coaching, phone calls, emails).</td>
</tr>
<tr>
<td>Numbers referred</td>
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<tr>
<td>Numbers registered</td>
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<tr>
<td>Numbers who start to use the intervention</td>
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<tr>
<td>Numbers who complete the intervention</td>
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<tr>
<td>Usage data for each user</td>
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<tr>
<td>Friends and Families Test</td>
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<tr>
<th>Behavioural and clinical outcomes</th>
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<tbody>
<tr>
<td>Height for calculation of BMI</td>
</tr>
<tr>
<td>Physical activity (IPAQ)</td>
</tr>
<tr>
<td>Patient activation (PAM-13).</td>
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</tbody>
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<tr>
<th>Economic outcomes</th>
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<tbody>
<tr>
<td>Cost of the digital diabetes prevention intervention</td>
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<tr>
<td>Types of staff involved in implementation in each LHE</td>
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<tr>
<td>Time spent by each member of staff on implementation of the DDPP (estimated)</td>
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<tr>
<td>Additional costs</td>
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</table>
### Table 2  Qualitative and explanatory outcomes

<table>
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<tr>
<th>Stakeholder group</th>
<th>Main areas of topic guide</th>
<th>Time point for collection</th>
</tr>
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</table>
| Local health economies (LHE) (commissioners, diabetes leads, health service managers) | About the LHE: ► Geography, demography and priorities  
► Culture and organisational style  
► Rationale for engaging with Digital Diabetes Prevention Programme (DDPP) (hopes, expectations, fears)  
About the digital diabetes prevention intervention (DDPI) selected  
► How and why this DDPI was selected  
► Views of the selected intervention  
About the implementation plan  
► Describe the implementation plan  
► Reflections on progress, strengths, weaknesses, amendments proposed or made  
Resources required  
► Types and numbers of staff involved  
► Time per staff member (estimated)  
► Other costs/resources  
Overall lessons learnt | X  
X  
X  

| Healthcare professionals | Local geography, demography and clinical priorities  
Understanding and prioritisation of DDPP  
Views about DDPI in use in local area  
Views about potential benefits / harms of DDPP, including impact on health inequalities  
Views about implementation process locally  
Overall lessons for future national delivery | X  
X  
X  

| Digital Diabetes Programme Intervention Providers | Describe the intervention  
Describe the evidence base for the intervention  
Onboarding process  
Views on how the implementation is going in participating LHE  
Explanations and reflections on reasons for successes/challenges in implementation  
Views on whether and how this programme could be scaled up nationally  
Observed usage and impact of intervention on patients and reasons for these  
Overall lessons learnt | X  

| Patients | Knowledge about diabetes and its prevention  
Relative prioritisation of diabetes prevention  
Experience of DDPP including identification, referral, onboarding to DDPI, use of DDPI  
Reasons for use/non use of DDPI  
Impact of DDPI on lifestyle and health behaviours  
Preferences for digital versus face to face  
Suggestions for improvement  
Overall views about the programme | X  

in the NDH and overweight/obese groups respectively, assuming clustering is ignorable. This compares favourably with a weighted mean effect size of d=0.22 (95% CI 0.20 to 0.23) estimated in a meta-analysis by Johnson et al\textsuperscript{(9)} for behaviour change interventions targeting eating and physical activity. Further power analysis allowing for clustering effects by demonstrator site (with an intraclass correlation coefficient (ICC) of 0.02 based on a median estimate of 0.0185 in a study of ICCs in adults with diabetes in primary care practices\textsuperscript{(5)}) gave minimum detectable effect sizes of d=0.18 and 0.22 in the NDH and overweight/obese groups, respectively, assuming a 25% completion rate at 12 months. For the purpose of analysis, completion is defined as obtaining data on weight and HbA1c at 12 months.

Qualitative data
Interviews will be recorded, transcribed verbatim and anonymised prior to analysis. Transcripts will be analysed using framework analysis\textsuperscript{52} which is well suited to policy-relevant research, with specific questions and a priori issues. The five steps of framework analysis are (1) familiarisation, (2) identifying a thematic framework; (3) indexing; (4) charting and (5) mapping and interpretation. Familiarisation will be achieved by reading and rereading transcripts, with an a priori framework based on the CFIR used to index and chart the data. Data that cannot be coded using CFIR will be noted. Mapping and interpretation will take place in multidisciplinary data clinics where interpretations can be proposed, discussed and refined.

Ethics, research governance and data security
PHE is the sponsor for this research.

Data security and information governance
Data will be handled according to the principles of the General Data Protection Regulation, the European Union framework for data protection which became law in the UK in May 2018.

Quantitative data
The DDPI providers will be responsible for obtaining and pseudonymising the quantitative data. No personally identifiable data will be handled by RSM. Postal code mapping for IMD will be undertaken by the LHE (with support from RSM).

Qualitative data
1. Patients. Patients will be invited to participate in the interviews via the digital diabetes prevention providers and/or the LHE and will opt in to providing fully informed consent for interviews.
2. Healthcare professionals, commissioners and providers. RSM has the contact details for these individuals as RSM is also responsible for the implementation of the DDPI. Written informed consent will be obtained prior to undertaking interviews.

Interview tapes will be stored securely on RSM servers. Only anonymised transcripts will be shared with the evaluation team outside RSM.

Dissemination
Contracted outputs include a 12-month report to the funder, based on 6-month follow-up data, and a final report, based on 12-month follow-up data. The findings will inform the scale of future provision of digital approaches within the NHS DPP.

Academic dissemination will also be undertaken in the form of conference presentations and publications in peer-reviewed journals. These presentations and publications will require advance approval from NHS England. Approval will not be unreasonably withheld, but academic dissemination may have to be delayed till after major policy decisions have been taken and made public.

Contributors
EM and JI developed the initial outline evaluation plan in response to the NHSE tender; this was developed, refined and operationalised by EM, KD, WH and AL. WH led on the statistical analysis plan. JV chairs the NHSE Programme Board which developed and oversaw the tender. EM wrote the first draft of the protocol paper; all authors have read and commented on drafts and have approved the final version. PPI involvement was provided by Diabetes UK membership of the Board overseeing this work.

Funding
This work was supported by NHS England, as part of the overall contract for implementation and evaluation of the Digital Diabetes Prevention Programme. EM is part funded by the NIHR School for Primary Care Research and the NIHR Collaboration for Leadership in Applied Health Research and Care, North Thames. AL is funded by the HEE Deanery (North Thames).

Competing interests
JI is a partner and KD a management consultant at RSM. RSM hold the contract from NHSE to implement and evaluate the Digital Diabetes Prevention Programme. EM and WH receive consultancy fees for their work on the evaluation. EM is managing director of a not-for-profit Community Interest Company, HeLP-Digital, which exists to disseminate a digital diabetes self-management programme, HeLP-Diabetes, across the NHS. JV is the National Clinical Director for Diabetes and Obesity at NHS England.

Patient consent for publication
Not required.

Ethics approval
Ethical approval has been granted by the Public Health England Research Ethics and Governance Group, reference R&D 324.

Provenance and peer review
Not commissioned; externally peer reviewed.

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