Mixed methods protocol for a realist evaluation of electronic personal health records design features and use to support medication adherence (ePHRma)

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ABSTRACT

Background National Health Service policy suggests that increasing usage of electronic personal health records (PHR) by patients will result in cost savings and improved public health. Medication adherence means that patients take their prescribed medication as agreed with their doctors. Some of the claimed benefits of PHRs are decreasing healthcare costs and improving medication adherence and patient outcomes.

Methods This is a mixed methods convergent study, primarily qualitative. The qualitative and quantitative data collection and analysis will occur in parallel, and then be synthesised. We are interviewing and surveying adults with long-term conditions to identify what are the most important and useful features of their current PHR. The data collection comprises patient demographics, the Medication Adherence Questionnaire, the personality scale Big Five Inventory-2 Extra-Short Form and the WHO Quality of Life-BREF scale. Qualitative data will be analysed using the Framework method.

Ethics We have received a favourable ethical opinion from the Health Research Authority/Research Ethics Committee.

BACKGROUND

The WHO reported that medication adherence in patients with long-term conditions averages to 50% in developed countries. Based on the results of a 2017 US survey, roughly 40% of patients who are chronically ill were interested in using technology to assist them with medication, diagnosis, test results and managing their condition in their home environment. National Health Service (NHS) England policies published over the past decade such as Personalised Health and Care 2020 (PHC2020) and the Five Year Forward View specify that the NHS needs to use information and communication technologies (ICT) to reduce healthcare costs and improve healthcare outcomes. The most recent NHS policy document, The NHS Long Term Plan, focuses on ‘personalised healthcare’ to improve quality of life and public health and aspires that over the next 5 years with the introduction of further ICT, outpatient visits will drop by one-third. The aim is that people with long-term conditions such as diabetes, respiratory or renal problems will have further access to technology designed to help them manage their condition, for example, continuous glucose monitoring for all pregnant patients with type 1 diabetes.

Recent literature suggests that there is limited evidence that the use of health applications can improve patient adherence to prescribed medication, and the quality of the evidence is often questionable. The impact of ICT on cost, quality and safety of healthcare remains questionable, based on the conflicting evidence of more ‘optimistic’ versus more cautious studies.

Medication adherence is ‘the extent to which a person’s behaviour—taking medication, following a diet, and/or executing lifestyle changes, corresponds with agreed recommendations from a healthcare provider’ and is considered a well-known challenge in healthcare, acknowledged by PHC2020. The ABC taxonomy is selected as the conceptual framework for medication adherence in this study, since it is well cited and it is considered more comprehensive than the WHO Five interacting dimensions that affect adherence, which will still inform this study as a secondary source.

It is widely suggested in the literature that medication adherence should be measured in conjunction with quality of life in order to define whether a patient’s health is better. Health-related quality of life (HRQoL) is the ‘subjective assessment of the impact of disease and treatment across the physical, psychological, social and somatic domains of functioning and
well-being. Literature also suggests that patient personality traits often affect medication adherence. The Five-Factor Model is an established taxonomy of personality traits. According to it, personality can be described in terms of five basic personality trait dimensions: agreeableness, conscientiousness, extraversion, neuroticism and openness to experience.

The NHS standards for commissioning personal health records (PHR) provide guidance on good practice for the development of PHRs in England, but they do not provide enough evidence on how the PHR standards impact public health, nor on what design features should a PHR include nor evidence on how these features impact health outcomes. PHRs are ‘online systems that include collections of patients’ healthcare and medical data, which utilise health informatics standards to enable patients to share, organize and manage these data according to their own views’. Some of the many claimed benefits of PHRs are the ability of PHR to improve patient outcomes, decrease healthcare costs, allow patients the ability to self-manage their health, empower patients and improve medication adherence.

### Aim and objectives

The aim of this study is to determine how best computerised PHR features should be designed to help patients get the full benefit of their prescribed medication. It could be hypothesised that health and information technology literacy may be important factors in identifying these essential PHR design features. Health literacy is ‘the degree to which individuals have the capacity to obtain, process, and understand basic health information and services needed to make appropriate health decisions’. The Health Education England defines digital literacy as ‘the capabilities that fit someone for living, learning, working, participating and thriving in a digital society’.

### Primary objective

Identify the essential design features of PHRs to improve medication adherence in adults with long-term conditions.

### Secondary objectives

- Identify how patient and disease-specific factors mediate the impact of PHRs.
  - Patient specific: personality traits and sociodemographics.
  - Disease specific: progression, severity, intervention type, polypharmacy.
- Develop a theoretical model that describes the interaction between the PHR design features and the patient and disease-specific factors, to help determine what works for whom in what circumstances.

### METHODS

In the absence of a comprehensive checklist for our mixed methods study design, we have adopted the Strengthening the Reporting of Observational Studies in Epidemiology (STROBE) statement given that our study is observational. The STROBE statement covers cohort, case-control and cross-sectional studies and we will follow the relevant statement’s sections about the cohort (for our qualitative data) and cross-sectional (for our quantitative data) studies.

### Study design

This study will follow the mixed methods research methodology, based on a pragmatist perspective, which is considered appropriate in the literature for medical informatics. The convergent mixed methods design will be used. The convergent design (or parallel or concurrent) is parallel and both qualitative and quantitative data are gathered at the same time, are analysed separately and then synthesised. The purpose of this design is to gather complementary data for the topic and aims to mitigate the pitfalls of solely quantitative or qualitative approaches, by illustrating quantitative and qualitative results together and examining relationships between variables. The results can then be merged and compared to answer the study’s aims and objectives. In this study, the qualitative data have been identified as dominant, as the area is largely under-researched and complex. The quantitative data will supplement and indicate the extent with which findings from the qualitative work are present. Figure 1 illustrates the different stages of the research and a more detailed explanation follows.

This study’s evaluation is based on the Medical Research Council guidance on developing and evaluating complex interventions that emphasises the importance of assessing the effectiveness and the outcomes of complex interventions contextually in interventions that are currently implemented and cannot be reversed.

This study will use the realist evaluation framework as its overarching theoretical approach to identify the optimal configurations of context, mechanism and outcomes in the use of PHRs for medicine optimisation. A realist evaluation includes a contextual evaluation (in what circumstances), a process evaluation (what works for whom) and outcomes evaluation (how). This holistic evaluation approach makes realist evaluation framework an ideal candidate to deal with the uncertainty and the complexity of interventions in medical informatics.

The evaluation will begin with the construction of an initial programme theory (table 1), informed by the literature review, which identifies the outcomes that are derived from the intervention and what mechanisms are in place to generate these outcomes as well as what context will affect them. The context–mechanism–outcome (CMO) configuration will be used as analytical approach, to identify mechanisms and contextual factors that are associated with the variation in outcomes. The study will iteratively revise the programme theory complying with the realist evaluation principles.

### Sampling and recruitment

Based on the study’s intention to directly relate the data sets, as well as to avoid influencing our ability to converge...
the results, we will use the same individuals to participate in both the quantitative and the qualitative strands of the study. The primary data of this study are qualitative; thus the sampling strategy is based on the qualitative component of the study, which will be merged with the quantitative data strand. A purposive convenient sample of participants will be recruited from volunteers through a variety of methods. The ‘gold standard’ for a purposive sample is to achieve saturation, which is impossible to predict. Therefore, based on the aims and objectives of this study and the type of sampling scheme selected, a typical recommendation is medium sample size, roughly 30–40 participants to achieve theoretical saturation.

Posters will be displayed in general practitioner surgeries, pharmacies in Hampshire and University of Portsmouth communal areas. Recruitment flyers will be electronically distributed by social media and via groups and forums of patients who are chronically ill, such as Diabetes UK. A project website will be created that will also make available the flyer and information sheets. Furthermore, more volunteers will be recruited via charities and other universities and colleges in South East England.

Participants

An electronic consent form and demographic information including their age and gender will be collected prior to the interviews and survey. The recruitment will occur in two stages. The first stage will not specify a long-term condition; rather we will interview anyone who complies with the inclusion criteria and consents. When we reach 15 (40%) of the intended participants, we will select the three majority long-term conditions, as was advised by the initial patient and public involvement (PPI) focus group. The participant recruitment will finish in August 2019.

Inclusion criteria

- Participant is willing and able to provide informed consent for participation in the study.
- Male or female, aged 18 years or above.
- Participant has a long-term condition and is treated outside hospital.
- Participant is able to self-administer his/her medication.
- Participant is frequently (at least once a week) using a PHR to manage in full or partially his/her medication.
- Participant is able to communicate freely.

Figure 1  Study design. BFI-2-XS, Big Five Inventory-2 Extra-Short Form; MAQ, Medication Adherence Questionnaire; WHOQOL-BREF, WHO Quality of Life-BREF.
Table 1  Initial programme theory (acronyms explained and further details provided in the Measures section)

Programme theory
The use of PHR can improve medication adherence in adults with chronic conditions.

Context—C
What conditions do we need for an intervention to trigger mechanisms to produce particular outcome patterns?
► Adults with long-term conditions
► Patient-specific factors (demographic characteristics, health and digital literacy)
► Disease-specific factors (comorbidities, duration of condition, severity)
► Patient personality traits: BFI-2-XS

Mechanism—M
What leads an intervention to have a particular outcome in a specific context?
► PHR design features that are used
► PHR type (tethered or stand-alone)
► PHR problems

Outcome pattern—O
What effects do the causal mechanisms have when triggered in a specific context?
► PHR benefits
► Medication adherence: MAQ
► Health-related quality of life: WHOQOL-BREF
► PHR design features that would be nice to use
► Perceived medication adherence and health-related quality of life

BFI-2-XS, Big Five Inventory-2 Extra-Short Form; MAQ, Medication Adherence Questionnaire; PHR, personal health record; WHOQOL-BREF, WHO Quality of Life-BREF.

Exclusion criteria
► Participants who are pregnant or terminally ill or considered vulnerable adults and patients with cancer.
► Adults with medically serious problems that are not classified as long-term conditions.
► Patients require assistance with taking their medication.
► Patients unable to communicate or unable to self-manage their medication.
► Inpatients or patients living in care homes.

Measures
This study will gather qualitative and quantitative data from the same population during an online/telephone interview, and will collect extra quantitative data via an online survey.

Quantitative data
The Medication Adherence Questionnaire (MAQ) scale will be used to measure medication adherence, which is a free-to-use four-item scale and has sensitivity (81%), specificity (44%), and Cronbach’s alpha reliability of 0.61. The MAQ scale was preferred over the more reliable and widely used 8-item Morisky Medication Adherence Scale (MMAS-8) or the 13-item Self-Efficacy for Appropriate Medication Use Scale. Both instruments were regarded as too long by our PPI group, additionally MMAS-8 is a licence product and this study is not resourced for this expenditure.

The WHO Quality of Life-BREF (WHOQOL-BREF) scale will be used to measure HRQoL, which is a free licence-based valuable scale for measuring quality of life in multiple chronic conditions, scoring 0.92 internal consistency reliability and has validity similar to the 36-Item Short Form Survey (SF-36) scale. The WHOQOL-BREF was preferred over other well-cited measures of HRQoL which include the SF-36, EuroQol EQ-5D, and the WHOQOL-100, purely for brevity following the PPI advice.

The shortest available form of the Big Five Inventory (BFI) questionnaire that is used to measure personality traits, namely the Big Five Inventory-2 Extra-Short Form (BFI-2-XS), was selected for this study. Although the BFI-2-XS has a lower psychometric scope than the complete BFI questionnaire, it appears to be adequate for this study based on PPI advice.

The sociodemographic information that we intend to gather is age group, gender, education, domestic status, health literacy, health and digital literacy as high, medium or low, based on further scales, based on the PPI advice regarding potential participation fatigue. We will simply infer the degree of health and digital literacy as high, medium or low, based on generic categorisations in the literature. Health literacy will be inferred from the questions 1, 4 and 5, and digital literacy from the questions 1 and 5 of the interview (see online supplementary file 1), as well as the employment and education status of the participant.

The guideline that we use to infer the health and digital literacy is provided in the online supplementary file 2.

Qualitative data
Qualitative dimensions of the concepts of effectiveness of PHR design features with medication adherence will be assessed with a semistructured interview (see online supplementary file 1). Questions that infer the medication adherence and quality of life are included in the interview, to strengthen our methodological and analytical approach.

Analysis
The qualitative data analysis will use the Framework method, which will enable themes to be developed inductively (from the interviews) but also incorporating themes identified in the literature review. The Framework method is systematic, thorough, data driven but also flexible and enables visual representation of the data. There are step-by-step guides to the application of the
Framework method and this study intends to follow the guidelines of Gale et al.24 25

We will use NVivo V.12 for coding. To achieve coding inter-rater reliability, a subset of files (25%) will be coded by an independent researcher and then the themes will be compared and the inter-rater reliability score will be calculated.24 After coding the 50% of the transcripts, a framework will be developed in an iterative manner, to apply to the rest of the transcripts. The framework will include codes that are grouped together and are clearly defined. The interpretation of the data will occur in conjunction with the quantitative strand during data merging.

We will use Microsoft Excel 2016 for descriptive statistics; namely mean, minimum, maximum, SD and coefficient of variance.

Joint displays will be used in a fashion similar to Vaughan Dickson et al.,26 so as the quantitative and qualitative results will be analysed thematically based on the CMO configuration to identify the essential design features of PHRs to improve medication adherence in adults with long-term conditions, taking into consideration the different personality types, adherence levels and quality of life opinions of the participants.

Patient and public involvement
A PPI focus group with eight participants took place at the University of Portsmouth in June 2018. This group’s suggestions are taken into consideration by the research team. The study protocol has been also reviewed and approved by this PPI group. PPI will be involved throughout this research, in order to minimise bias, increase transparency and ensure approval.

Limitations
This study is limited to the design features of a PHR and not the overall design. There are no explicit questions in the interview guide regarding usability or interface design. Another limitation is that the sample size of this study is limited to 30–40 participants. The dominant data of this study are the qualitative and it is currently acceptable in broad literature to interview approximately 40 people.34 35 Due to time constraints a higher number would be unrealistic.

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Contributors
EA drafted the manuscript, and is the guarantor of the protocol. PJS revised the manuscript multiple times for methodological and intellectual content. HH revised the manuscript twice for methodological, conceptual and intellectual content from a pharmacological perspective.

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Competing interests
None declared.

Patient consent for publication
Not required.

Ethics approval
This research has been approved by the NHS North East—Newcastle & North Tyneside 2 Research Ethics Committee (10 September 2018). The findings will be published in peer-reviewed journals and presented at national and international scientific conferences.


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