Involving elderly research participants in the co-design of a future multi-generational cohort study

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Involving elderly research participants in the co-design of a future multi-generational cohort study

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Abstract

Objectives:
To report as a case study the process of involving study participants and how this impacted the co-design of a proposed multi-generational research study, using a novel standardised reporting tool.

Background:
An advert was sent to 14,268 participants of the ASPREE-XT observational cohort study to invite them to be involved in co-design and planning of a future multi-generational research study. Participants were aged 74 years and older from Australia, located in both metropolitan and rural locations.

Interventions:
We used participatory action research to involve elderly research participants in the co-design of a proposed multi-generational cohort study between 2017-2019 using newsletters, telephone interviews and an in-person workshop.

Primary and secondary outcome measures:
We used the ‘Standardised Data on Initiatives (STARDIT)’ Alpha Version to plan and report how participant involvement activities positively impacted the study design.

Results:
Fifty-nine ASPREE participants were interviewed by telephone and 18 participants attended a face-to-face workshop event. Involving participants positively impacted the proposed study design by improving the research objectives, developing protocols, influencing funding decisions and improving ethics applications. Learning points included the importance of maintaining the ideals of the ASPREE study (respect, quality and transparency); research participants’ preference for the option of receiving results (including genetic results); participants’ need for involvement in decisions about recruitment, data access, governance and other ethical issues; and the preference for different communication methods, including both face-to-face and online methods. The process was highly valued by all stakeholders, including research participants, study staff and lead investigators. Involvement of participants was described by the lead study investigator as “enormously helpful”.

Conclusions:

This case study provides evidence that including participants in the design of a research study positively impacted the study design, participants and researchers. Using a standardised reporting tool to describe the methods and impacts provides a way for learning from this case study to inform future research studies planning to involve people.

Plain English summary [required 250 words max]

Research which follows a population of people over time is a valuable way to learn about what contributes to both wellness and disease. Some studies of this kind look at multiple generations and are called multi-generational, longitudinal cohort studies. Such studies have had significant positive impacts on public health. However, such studies are challenging to establish and expensive to maintain.
It has been proposed that the existing ASPirin in Reducing Events in the Elderly Extension study (ASPREE-XT) would be a good basis for a future multigenerational research study (MGRS). ASPREE-XT participants are all over the age of 74.

There is evidence that involving potential participants in co-designing research can improve the quality of the study design, recruitment and acceptability of the research. A team of people including current a ASPREE-XT participant were involved in co-designing how potential participants would be involved in the co-design process. An advert was sent to 14,268 participants of the ASPREE-XT observational cohort study to invite them to be involved in the process of planning a future multi-generational research study.

Fifty-nine ASPREE-XT participants were interviewed by telephone and 18 attended a face-to-face workshop event. We used the ‘Standardised Data on Initiatives (STARDIT)’ Alpha Version to plan and report how participant involvement activities positively impacted the study design. Involving participants positively impacted the proposed study design by improving the research objectives, developing protocols, influencing funding decisions and improving ethics applications. This case study provides evidence that including participants in the design of a research study positively impacted the study design, participants and researchers.
Introduction

The ASPirin in Reducing Events in the Elderly (ASPREE) trial (2010-2018), a randomised controlled trial for aspirin in healthy older people, recruited 19,114 participants from Australia and the USA. The median age at recruitment was 74 years. The ASPREE Healthy Ageing Biobank is a sub-study which has collected biospecimens from over 15,000 ASPREE participants, alongside detailed medical records, lifestyle, cognitive function and physical testing data. ASPREE researchers are conducting various types of genomics and biomarker research. ASPREE had a remarkably high retention rate, with only 1.2% of participants withdrawing from the study and 90% still attending annual visits after an average of 4.5 years follow-up.

ASPREE XT is a follow up observational study that was established in 2018 to continue to collect data from ASPREE participants for another 5 years. ASPREE-XT participants are over the age of 74 (defined as elderly) and Australian participants are located in both metropolitan and rural locations. Multiple stakeholders, including a participant advisor, proposed a new multi-generational research study (MGRS) which could recruit direct descendants of ASPREE-XT participants, as part of a longitudinal observational study alongside their relatives for two generations or more. MGRS are challenging to establish and expensive to maintain, yet their value to medical and epidemiological research is significant. ASPREE-XT has been proposed as a platform for a multi-generational study. Recruiting ASPREE-XT participants and their descendants to a MGRS would provide a large starting population by international standards.

Previous multi-generational, longitudinal cohort studies have had significant positive impacts on public health. Examples include the Framingham Heart Study, Lothian Birth Cohort and Dubbo Osteoporosis Epidemiology Study, all of which had substantially lower starting
populations than ASPREE-XT. ASPREE-XT provides a rare opportunity for such a study in Australia, already combining high-quality medical record data with genomic data on a large number of elderly Australians. Additionally, the cohort is already well engaged, with all surveyed participants supportive of a proposed MGRS.

The aim of this study was to report as a case study the processes of involving ASPREE-XT participants in the co-design of a proposed multi-generational research study, and how this impacted study design. The research objectives were to involve potential participants in the co-design of a new MGRS; plan the process in a standardised way; identify themes and preferences; and evaluate then report the process using a standardised reporting tool.

ASPREE-XT’s unique focus on healthy ageing adds substantial value to public health, medical, epidemiological and geriatric research. Previously collected genomic (including epigenetic) data, combined with detailed medical data and ongoing cognitive assessment, allow that a MGRS with the ASPREE-XT cohort would be of considerable value to science for decades to come, for example in dementia research. The proposed study would examine health outcomes in a large, well-monitored cohort; and provide data to help inform our understanding about the genetic and environmental determinants of health and disease, across multiple generations. However, this is only possible if people choose to participate.

Many clinical research studies are underpowered due to poor recruitment and retention. Involving participants and the public in research design has been shown to improve the recruitment, quality and relevance of research. The concept of involvement is research being done ‘with’ people rather than ‘on’ them. Involving the public, patients, research
participants and other stakeholders in actively contributing to the research process can lead to a range of positive impacts and outcomes. In human genomics research, the need to involve the public and other stakeholders is a crucial aspect of responsible research practice. The term ‘stakeholder’ here means anyone who has a ‘stake’ in the research, in particular those with important knowledge, expertise or views that should be taken into account. This includes ASPREE-XT participants, study staff and academic research investigators and the wider public. At the earliest stage of the research cycle (the conceptual stage), some current ASPREE-XT participants were invited to be involved in the co-design of a new MGRS. In this paper we aim to outline how people were involved in the co-design process in order to appraise the methodology and inform future best-practice.

**Terminology**

We have used consistent language in this paper to describe concepts such as ‘involvement’. To aid readers, Table 1 provides definitions of important terms used consistently throughout this paper.
### Table 1. Definitions of terms

**Involvement** – The words ‘involvement’ or ‘being involved’ describe the concept of people being ‘involved’ in research. This is when research is carried out ‘with’ people rather than ‘on’ them\(^{20}\). ‘Involvement’ can also be defined as when other people aside from the research team, such as the public, patients, research participants and other stakeholders, actively contribute to the research process\(^{21}\). It is the ‘active involvement’ in shaping and guiding research, rather than only providing data\(^{25-27}\).

**Participant** – a person who participated in the process of sharing views and perspectives about the proposed MGRS, including sharing views about preferences for any future involvement. The term ‘ASPREE-XT participants’ will be used when specifically referring to participants from the existing study.

**Participant advisor** – before inviting people to become participants, it was necessary to involve a small number of participants to help advise and plan the process. These participants were chosen from an existing reference group of ASPREE-XT participants.

**Stakeholder** – this term includes anyone who has a ‘stake’ in the research, in particular those who have important knowledge, views or perspectives that should be taken into account\(^{22,24}\). In this paper it refers to participants, participant advisors and ASPREE-XT study team members (including researchers, ASPREE-XT participant assessors and lead investigators).

**Study team** – this process was guided by the study team, who consisted of academic researchers, ASPREE-XT participant assessors, a participant advisor and a lead investigator.
The process – this term will be used to describe the process of involving ASPREE-XT participants by inviting them to share views and perspectives about a potential future MGRS. This process includes the co-creation of this case study with participant advisors.

**Participatory action research (PAR)** - this term describes a number of related approaches, including forms of action research which embrace a participatory philosophy and include ‘co-design’ and ‘co-production’ of research \(^{28(p1)}\).
Materials and Methods

Study design

A participatory action research (PAR) method was chosen to guide the process with co-design and reporting guided by a number of frameworks. PAR is an umbrella term which describes a number of related approaches, including forms of action research which embrace a participatory philosophy and include ‘co-design’ and ‘co-production’ of research. During the study design, we applied this co-design process where researchers and other relevant stakeholders (including research participants) “work together, sharing power and responsibility from the start to the end of the project”, including knowledge generation and translation. Figure 1 summarises the process we used (Fig 1). The ASPREE participant advisor was an integral member of the study team, through each stage.

Guiding Frameworks

The process was guided by a number of international PAR methodology frameworks, including the International Collaboration for Participatory Health Research. The PAR method was also informed by an international review of involvement in genomic research carried out by some members of the study team. Learning from this review informed the subsequent development and application of an Alpha version of ‘Standardised Data on Initiatives’ (STARDIT). STARDIT includes a tool to map people’s preferences for involvement in a standardised way, including mapping views on who should be involved and how. STARDIT was then used to guide co-design of the process, and to subsequently report how people were involved, using standardised data. An Alpha version of the STARDIT framework was also used.
in parallel with the thematic analysis to organise data into pre-defined ‘super-categories’ which allow consistent comparison with other data using this reporting framework\textsuperscript{13}.

We used a case study research methodology to record and describe the process of involving participants in the co-design of a proposed MGRS. Case study research methodology is a form of empirical inquiry and this study is presented as an instrumental case study, where the purpose is to understand the particular case and to provide data that could produce useful generalisations by using inferences from the data\textsuperscript{40(p109)}.

The selection of the population for this case study was informed by a number of factors which were appraised by the study team, including ethical, pragmatic and population considerations\textsuperscript{41,42}. One of the study team members (PL) is an investigator with the ASPREE-XT study, which was used as a starting point by the study team to explore the appropriateness of this case study.

**Data collection and reporting**

The data collection and reporting was informed by a number of frameworks for reporting involvement in research\textsuperscript{13,22,30,31,38,43–45}. Qualitative and quantitative data was collected from multiple sources, including meeting notes, audio recordings, documents, emails and surveys. Data sources were analysed using a number of methods including thematic analysis. Further information about how data sources were analysed is in ‘Table 2. Summary of data analysis’.

**Stages of research**

**Fig 1. Process timeline of the participant involvement**

[see file ‘Figure 1’ – ‘Process timeline of the participant involvement’]
Stage 1: Planning

The study team held four meetings to co-design the involvement activities. One participant advisor was involved in a number of tasks including reviewing and improving the written information, telephone interview questions, and the facilitation plan for the event.

Stage 2: Recruitment and telephone interviews

An advert was placed in a newsletter to 14,268 ASPREE participants. The process we used for recruiting and involving ASPREE-XT participants has been divided into four chronological stages, summarised above in Figure 1 (Fig 1).

After the recruitment and consent process, participants were interviewed by telephone. A summary of the questions and responses from the telephone interviews is shown in Table 3. For a list of all questions asked, see Additional file 2. Participants were asked about their willingness to provide feedback throughout the study, and to be involved in study design, as well as preferences for modes of communication. The definition of involvement below used in the script was co-designed with participants for subsequent interviewees, with versions 1 through 1.3 piloted with 15 participants.

Pre-question script: ‘Traditionally, research studies have been designed and conducted only by researchers, and people invited to participate. We are challenging this idea of researchers being the only experts.’

Question: Would you be willing to provide feedback and share your views and perspectives by commenting throughout the research process?’
Twenty relevant interviews were transcribed, coded and categorised (JN), with relevant interviews identified by two investigators independently (JN, PL). To reduce any unconscious selection bias, a sample of over 10% of the interviews was selected at random.

**Stage 3: Event**

Eighteen participants attended a four-hour workshop event in central Melbourne, led by JN, who is an experienced facilitator. The event was co-designed by the study team, and was informed by interview data and international best-practices for involvement events. The event included an introduction to the proposed MGRS by the lead ASPREE-XT genomics researcher (PL); a plain-English introduction to genomics by an expert in genomics who is also an ASPREE-XT participant (BH); a summary of the telephone interview results by the interviewer (MS); and an interactive session which included open questions about the types of information participants would like returned and recruitment of family members (JN).

The final session included a presentation and interactive discussion about involvement in research, led by the event facilitator (JN). This session explored preferences about how people would like to be involved, with open and closed questions. Questions included preferences about tasks and modes of communication.

Throughout the event, participants shared their views on a range of issues through interactive discussions, voting (by show of hands) and anonymous written feedback.
Stage 4: Evaluation and analysis

Table 2. Summary of data collection and analysis

| Data source description                                      | Analysis method                        |
|-------------------------------------------------------------|----------------------------------------|
| **Meetings** – including meeting notes, recordings and      | Qualitative (content analysis)         |
| relevant documents                                          |                                        |
| **Telephone interviews** – including recordings, notes      | Qualitative (thematic analysis,       |
| made by interviewer, summary documents and related          | content analysis)                      |
| emails                                                      | Quantitative (number of responses)     |
| **Event** – including audio and video recordings, interviews,| Qualitative (content analysis)         |
| written notes and feedback forms                            | Quantitative (number of responses)     |
| **Study team surveys** – responses to open ended questions  | Qualitative (content analysis)         |
| by email                                                    |                                        |
| **Other data** – this included reflexive research diaries,  | Qualitative (thematic analysis)        |
| relevant emails, financial and other relevant documents     | Quantitative (analysis of cost)        |

Evaluation and reflection

All members of the study team (except JN) were surveyed six months after the face-to-face event in order to integrate the valuable views and perspectives of those involved in co-designing and delivering the process. Design of surveys was informed by frameworks for planning and reporting public involvement\textsuperscript{30,31}. The study team were asked 11 questions (available in Additional file File 1) and the data from the four interviews was coded and categorised (JN) using STARDIT\textsuperscript{13}. 
Data analysis

The stages of qualitative data analysis included data mapping and familiarisation; transcription; coding; searching for themes; reviewing themes with study team members; labelling and summarising themes; and reporting the findings. In order to enhance validity of the analysis, two authors (JN, MS) independently analysed the data thematically, which was then ‘member checked’ by a third author (PL). Standardised categories (STARDIT) were used during content analysis of the data in order to facilitate comparison with other research projects. More information about the data sources and a STARDIT report available is in Additional file 1.

Results

The results are presented in three sections. Section one provides results from the stages outlined in Figure 1 (Fig 1), section two shares the perspectives of ASPREE-XT participants and other stakeholders, and in section three we describe how participant involvement impacted study design in seven ways. We quote participants directly, assigning each a unique number. Figure 2 summarises the entire process, results and impacts (Fig 2). For quantitative data from the telephone interviews, see Additional File 3.

Section 1: Results from stages 1-4

Results from Stage 1: Planning

The input of the participant advisor during the early co-design stage had clear benefits, in particular in identifying the best approaches for engaging the broader ASPREE population; improving the wording of participant information resources and improving question design for the interviews.

Results from Stage 2: Recruitment and telephone interviews
After reading the advert in the newsletter, 76 ASPREE-XT participants agreed to participate in the process. We interviewed 59 people by telephone, with the remaining 17 not responding to a number of follow-up calls.

Interview participants expressed a strong interest in receiving results from genetic research. The strongest preference was for genetic results of medical significance (rather than ancestry or drug-response).

All telephone interview participants expressed willingness to provide feedback throughout the process of planning the proposed MGRS. While most participants understood the concept of involvement, 9 of the 20 analysed expressed self-doubt about how they could be involved. Of 20 analysed telephone interviews, five participants seemed unclear about what tasks they could be involved in, suggesting the concept of participant involvement was new to them. Six participants sought clarity about what was expected of them when being involved, or discussed the skills and knowledge required. Further, participants stated that the goals and processes of involvement needed to be clear in order to avoid “spending too much time” and were willing to be involved as long as the task had purpose and was not “just for the sake of chatting” (P1).

During the telephone interview stage, the interviewer initially “struggled to explain” the concept of involvement (as distinct from participation in an active study) (MS). Co-designing and making changes to the language used in the telephone interviews appeared to improve participant understanding of the concept of involvement. Changes included providing a clear definition and pre-question script and including actual examples of involvement (such as overseeing ethical decisions about data access), thereby avoiding jargonistic abstract concepts such as ‘involvement’.
| Questions (closed)                                                                 | Results                  | % of interview participants |
|---------------------------------------------------------------------------------|--------------------------|-----------------------------|
| 1. Do you think participants like yourself should be involved in helping design research projects, or just researchers? | Just researchers         | 39                          |
|                                                                                 | Participants involved    | 46                          |
|                                                                                 | Unsure                   | 11                          |
| 2. Would you be willing to provide feedback and share your views and perspectives by commenting throughout the research process? | Yes                      | 100                         |
|                                                                                 | No                       | 0                           |
| 3. Would you like to be more involved in study design rather than just participating? | Yes                      | 65                          |
|                                                                                 | No                       | 12                          |
|                                                                                 | Unsure                   | 23                          |
| 4. Would you be more or less likely to participate in research if participants were involved in design | More likely              | 25                          |
|                                                                                 | Less likely              | 0                           |
|                                                                                 | Unsure                   | 13                          |
|                                                                                 | Wouldn’t influence        | 13                          |
|                                                                                 | participation            |                             |
| 5. What is your preferred mode of communication for being involved               | Face to face event       | 15                          |
|                                                                                 | Post                     | 31                          |
|                                                                                 | Online discussion        | 3                           |
|                                                                                 | Online questionnaire     | 15                          |
| 6. If genetic research results were made available, which types of genetic testing would you be interested in? * | Medical                  | 97                          |
|                                                                                 | Ancestry                 | 97                          |
|                                                                                 | Drug response            | 94                          |
7. If only one option for genetic testing was available, which one would you prefer?

| Option               | Participants |
|----------------------|--------------|
| Medical              | 65           |
| Ancestry             | 12           |
| Drug response        | 6            |
| No preference        | 18           |

*Note that percentages indicate the number of participants who responded with that response. On question six, participants could select multiple answers, so totals do not add up to 100%.

**Results from Stage 3: Event**

Eighteen of the invited 76 participants attended the face-to-face event. When specific tasks were discussed at the event, 10 participants expressed interest in being involved in recruitment and communication tasks, 7 in data access decisions, and 11 in ethical decisions. When surveyed after the event, the study team reported that involving participants positively impacted the proposed study design by improving the research objectives, developing protocols, influencing funding decisions and improving ethics applications. Videos of the event discussions and interviews with participants will be shared in the public domain.

**Results from Stage 4: Evaluation and reflection**

Participant feedback from the workshop event was analysed, with participants reporting it as a positive experience, one describing it as “brilliant” (P7). When surveyed, all members of the study team reported that the event achieved its intended aims and was a positive experience. No negative impacts were reported from any participants or study team members at any stage of the process.
One study team member described that through involving people, their perceptions “significantly changed” as they learned how participants could have “valid, interesting, and often novel ideas that researchers may not have considered”. The lead investigator (PL) noted:

“What I learnt is that sometimes the researchers’ assumptions about certain things can be off or even incorrect, and that researchers can miss critical points that are important to participants and the study”.

During the process, both participants and study team members reported changed views about the value of involvement in research, demonstrating ‘transformative learning’ and co-construction of knowledge.

A number of significant learning points were identified by the study team when responding to the question “do you have any advice to other researchers planning involvement for their research”. Significant learning points are summarised in Table 4.

**Table 4: Summary of learning points from study team**

| Summary of learning points from study team |
|-------------------------------------------|
| 1. Fund and prioritise involvement, make it a requirement |
| 2. Ethics processes take time, but can improve plans |
| 3. Know your audience – don’t make assumptions |
| 4. Value diversity in experience and knowledge |
| 5. A supportive team improves the experience for all |

**Cost and value**
The entire process of involving people was estimated to cost $10,000 AUD, including staff time, catering and event venue hire. The value of the process was summarised by the lead investigator who stated “I learnt a lot from the process and am very glad we made the effort”.

Section 2: Perspectives of ASPREE participants and other stakeholders

Recurring themes were identified by the study team from the data sources analysed, and seven specific impacts were reported as a result of involving participants (summarised below). The analysis is divided into two sections: (1) participant views about the proposed study, (2) stakeholder views about involvement in the proposed study.

1: Participant views about the proposed study

Participants were very positive about the proposed MGRS, with one stating ‘I think it sounds very good’[P20]. Altruism was a primary motivator for participation, with participants suggesting that outcomes of a MGRS could include benefits to themselves (personally and for their families); improving healthcare more generally; and the potential for saving lives, preventing diseases, improving quality of life, and improving future research.

Participant views on types of information they would like returned from genomic research and how the study should be funded (commercial funding versus public funding) were diverse and did not always align with the study team’s prior expectations. For example, two thirds of event participants wanted access to their own genomic data, which was described by the lead investigator as ‘very at odds with the current system’ (PL).

2: Stakeholder views about participant involvement in the proposed study
Participants were supportive about being involved, with all participants supportive of being involved by providing feedback throughout the research process (100%, 32/32), with a typical participant response being ‘I’d be happy to be involved’ [P3]. Views about enablers were shared in three of the 20 interviews coded, by all 18 of the event participants and all study team members surveyed. Views about barriers were shared in eight of the interviews coded and by half of the study team surveys. Further data is categorised in Table 5. Additional mapping of preferences for involvement using the STARDIT-PM tool can be found in Additional File 1.
Table 5: Enablers and barriers for involving participants

The lead investigator stated “the feedback has been so valuable” and that it will be “built into the design” of future research. However, the study team also identified barriers to involving people which exist for researchers. One study team member reported that at the start of the project they were “unsure how people without a science or health background” could be involved. They reported a personal shift during the involvement process from not understanding how participants could be involved and being concerned about “asking too much” to believing that, “with adequate resources (financial, training, time...), people can be involved in all aspects of genomics research”.

Other barriers identified by the study team included delays in obtaining ethical approvals for involving people and the cost of involvement in both time and money. One study team member reflected in the follow-up survey that they had “worried too much” about the time-burden of involving participants. The concern of “not putting further pressure” on participants was a theme
in email communication between the study team when making decisions to limit contact with
participants throughout the process of involving them.

Survey data from the study team also suggests that adequate funding, a supportive team,
involving participants in the very earliest stage of research planning and co-designing inclusive
methods of involvement all contributed to the impacts reported.

**Section 3: Impacts on study design from stakeholder involvement**

Involving stakeholders in the co-design process impacted the study in seven specific impacts
ways. By asking for participants’ views on aspects of the proposed study design, the study team
gained insight into participant preferences and opinions. While there was diversity in views, the
process allowed the study team to improve aspects of the study design. These impacts are
summarised in Table 6 ‘Summary of impacts on study design’
Table 6 ‘Summary of impacts on study design’

| Impact on planned research                      | Summary of impact                                                                                                                                 |
|-------------------------------------------------|---------------------------------------------------------------------------------------------------------------------------------------------------|
| 1: Recruitment and sample collection            | Recruitment and consent for the MGRS will occur online wherever possible, and saliva samples (rather than blood) will sent by post to be used as biospecimens for DNA analysis. |
| 2: Participant communication                    | A short video and ‘information pack’, which will explain the MGRS study, will be created to assist with recruiting family members.                  |
| 3: Participant involvement in governance        | Participants will be invited to be involved in overseeing governance, including funding decisions.                                                  |
| 4: Data access                                  | Study participants should be involved in controlling data access decisions and policies.                                                             |
| 5: Communication and ways of involving participants | Participants will be included on study advisory groups, including for study recruitment and communication, data                                     |
access and ethical oversight using multiple communication modes.

| 6: Provide feedback to participants about the research | Participants will be informed about the impact of the research, and how their involvement has affected the design and management of the study. |
|------------------------------------------------------|----------------------------------------------------------------------------------------------------------------------------------|
| 7: Create learning and development opportunities     | Learning and development opportunities will be created for potential participants, researchers and other stakeholders.           |
Impact 1: Recruitment and sample collection approach

A discussion about participants’ adult children being ‘time-poor’ highlighted the importance of a study design which minimised the time burden on younger generations. Only a third of participants thought their children would be willing to do a blood test. This allowed the research team to make a more informed decision about ‘trade-offs’ between the data that can be collected via blood or saliva, versus the potential effect on recruitment.

Impact 2: Participant communication

Most participants reported willingness to be involved in recruiting family members to a new study, if given appropriate information and supporting documents. Relatively inexpensive information resources, such as explanatory videos, could have a significant impact on the success of the recruitment and the study as a whole. One participant also advised that information produced for participants by researchers can be confusing, and that laypeople can help simplifying it.

Impact 3: Participant involvement in governance

Event participants unanimously agreed that they should be involved in all aspects of the research, whereas 11 of participants from the 20 analysed interviews thought that they should be involved...
in study design. One interview participant felt only researchers should be involved as they are “the qualified people” (P8). Five of the interview participants expressed the view that non-researchers are required in research as they provide an important alternative perspective.

Participants had mixed views about commercial organisations funding research. Four event participants were against it, some were ambivalent, and the majority indicated no objections. One participant was concerned about the risks posed by involvement of commercial organisations with opaque “vested interests”, asking “what is in it for them?” (P2), referring to individuals and organisations with real or perceived conflicting, competing or commercial interests. Participants suggested that involving people in governance (including funding and ethical oversight) was a way of mitigating this risk.

Study team members reflected that public funding would be preferable to commercial funding, as the responses at the event suggested that a commercially-funded study might negatively impact recruitment. As a result, the study team altered the proposed design to involve participants in governance, oversight and funding decisions.
Impact 4: Data access

Participants shared the view that they would like different kinds of genetic information returned from the research (see Table 3), including personal medical, ancestry and pharmacogenomic results. Two-thirds of event participants wanted access to their own genomic data, and had mixed views about who else should have access. All event participants stated they were comfortable with their data being held by academics, and none were comfortable with data being held by a commercial company. However, one participant suggested not “ruling private companies out completely” from research (P9).

General practitioners (GPs) were generally trusted to access and interpret genomic data, but participants felt GPs should not have access to data that they did not. All but two event participants agreed they should exclusively control access to their own data, with those disagreeing mentioning cognitive decline as a reason for a co-managed access model.
Some participants had concerns about themselves or their biological relatives (especially offspring) finding out information they “might not want to know” (P6), or having it imposed on them.

**Impact 5: Communication and ways of involving participants**

Preferences for communication mode differed between interview participants and event participants. Interview participants preferred questionnaires via post (30%, 18/50), and face-to-face events (25%, 15/59), over online questionnaires (17%, 10/59) and online discussions (8%, 5/59). Event participants suggested face-to-face meetings were helpful but only when there was an “occasional need” (P3). Participant responses also suggested that limiting face-to-face events where possible (in favour of telecommunication) may mean involvement is more inclusive.

Event participants felt certain tasks (such as reviewing information) could be done “more online” (P3). Participants spontaneously suggested using online, moderated forums and suggested that these should be hosted by trusted institutions (such as universities) rather than commercial organisations, as some ‘don’t trust’ social media companies [P17].

**Impact 6: Provide feedback to participants about the research**
Event participants stated that keeping people informed about what has been learned from the study is a good way of keeping people engaged in the study and improving retention. For example, ASPREE-XT sends a regular newsletter to participants. Participants also stated that they would like to be informed about when their involvement has made a difference.

**Impact 7: Create learning and development opportunities**

Participants often stated their willingness but also their uncertainty about how they could be involved. Learning and development opportunities were identified as an e of involvement by both a participant advisor.

**Fig 2. Co-designing multi-generational genomics research**

[See file – ‘Fig 2’ - Co-designing multi-generational genomics research]

**Discussion**

The participatory action research method gave insights into participants’ preferences that measurably impacted on the proposed study design. The improvement of the interview design
using the co-design process illustrates the value of a flexible and iterative approach to involvement in a study.

Participants’ preference for being informed about both the study outcomes and the outcomes of their involvement is supported by other studies which suggest that communicating about the research regularly and sharing results may improve retention. This is particularly relevant for those planning involvement in genomics research, which may span decades. Such regular communication should also be combined with learning and development opportunities, which could also help facilitate participant involvement by ensuring that people understand the values which motivate the tasks they are being asked to be involved in.

Participant views about data storage and access aligns with a 2015 survey by Genetic Alliance Australia, which indicated people mostly trusted universities and research institutes to use personal genetic information for research, with commercial companies least trusted. Event participants’ unanimous preference for having access to their own data and a general trust for GPs to access and interpret genomic data also aligns with findings from other studies.
Participants’ concerns about unintentional disclosure of data to biological relatives who ‘might not want to know’ certain information highlights the challenges of asking participants about information preferences, and recognises that this ethical decision extends beyond individuals [P6]. This issue is common to almost all ongoing and proposed genomic research studies and should be urgently addressed by contemporary individualist ethical frameworks 58.

Study strengths

By asking participants their preferences, the study team gained useful insights to inform the design of the proposed study. Participants preference for being involved in decision making about funding sources, data management and ownership, and what information to share with participants will help ensure any future study design aligns with participants’ values, ensuring the design is culturally safe and culturally competent 60. Similarly, the participants preference for being involved in reviewing participant information aligns with other research which suggests that involving potential participants in reviewing information can help improve recruitment 61.
The effective involvement of ‘stakeholders’ also includes involving all relevant staff and health professionals at all levels of an initiative, who may have important knowledge or perspectives that senior research staff do not.

The ‘transformative learning’ during the process reported from both study participants and the study team was an important impact captured by the participatory action research (PAR) method\textsuperscript{39,50}. The process showed that it was valuable to create regular involvement opportunities for each stakeholder. Reporting this process in a standardised way using ‘Standardised Data on Initiatives’ (STARDIT) meant that impacts such as transformative learning could be reported and that this case study can be compared to similar studies in the future \textsuperscript{[49]}.

**Study limitations**

Because the 76 participants who responded to the original advertisement were self-selected volunteers, our findings may not necessarily be accurate for the whole ASPREE-XT cohort. Our sample may reflect a sub-set of individuals who feel more strongly supportive of a proposed MGRS than the cohort average. However, the data from this process is still useful and valid\textsuperscript{57}. 
The number of interviews which were transcribed (20) and analysed was high for a case study. As there is no agreement in case study literature on whether to code all or some of the data, our methodology balanced exhaustive examination of the data with the capacity bias imposed by time constraints. Therefore, data transcribed and coded from the interviews does not include all the data collected in the interviews.

While the mixture of written and verbal feedback at the event ensured a range of ways to give feedback, the voting process (which involved people raising their hands in front of everyone) may have given different results if it was an anonymous ballot. Future research should explore and compare preferences for different methods of voting.

While the process of involving people described here did not exclude people based on language, the original ASPREE study required a certain level of English language skills in order for people to participate in some of the cognitive assessments and public events. The process described here may therefore have excluded people who cannot read English or do not feel confident speaking in English, such as people who speak English as a second language.
Conclusions

This case study provides evidence that including participants in the design of a research study positively impacted the study design. As many research studies are negatively impacted owing to poor recruitment and retention, such evidence is increasingly important for informing involvement in future studies. The process of involving ASPREE-XT participants in the design of a new MGRS was highly valued by stakeholders, and was positively impactful for both participants and the study team. The lead investigator stated “the feedback has been so valuable” and that it will be “built into the design” of future research. Learning from the case study suggests that adequate funding, a supportive team, involving participants in the very earliest stage of research planning and co-designing inclusive methods of involvement all contributed to the impacts reported.

The process of involving people can be viewed as a learning experience for both the participants involved and study team members. The process changed participant and study team members’ views about the value of involvement. For example, one a study team member stated they ‘learned that participants have valid, interesting, and often novel ideas that researchers may not
have considered’ and their ‘perceptions of involving participants in planning medical research
have significantly changed’, which can be viewed as an impact of ‘transformative learning’.

Using a standardised reporting tool to describe the methods and impacts provides a way for
learning from this case study to inform future research studies planning to involve people,
including studies beyond the discipline of public health genomics.
Declarations

Ethics approval and consent to participate

The Ethics Committee of the Alfred Hospital (Melbourne, Australia) granted approval for this project, which was considered Low Risk Review and approved on 16/04/2018.

Project No: 180/18

Project Title: ASPREE Genomics Engagement: Planning of Future Multi-Generational and Family Studies

Principal Researcher: Dr Paul Lacaze

Participants gave informed consent to participate, with further consent being granted by participants for sharing of recordings from the process.

Consent for publication
Availability of data and materials

All relevant data has been anonymised and shared in the additional files. Monash University is storing all raw data according to the relevant ethics policies, and invites requests for more detailed data. STARDIT reports will be added to databases in the future. Additional videos of the events will be shared in the public domain.

Competing interests

The authors declare that they have no competing interests.

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Authors' contributions
JN wrote the manuscript and analysed the data, facilitated face to face events, designed and carried out the evaluation of the process and the reporting (including STARDIT and GRIPP2 reports). MS edited the manuscript, conducted telephone interviews, ran sessions at the face to face event analysed data and checked data analysis. DA advised on methodology and edited the manuscript. BH was a participant advisor during the process, ran sessions at the face to face event, advised on the process and edited the manuscript. JT advised on the process and edited the manuscript. PL checked data analysis and edited the manuscript and ran sessions at the face to face event. All authors helped co-design the process, read and approved the final manuscript.

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References

1. McNeil JJ, Nelson MR, Woods RL, et al. Effect of Aspirin on All-Cause Mortality in the Healthy Elderly. *N Engl J Med*. 2018;379(16):1519-1528. doi:10.1056/NEJMoa1803955

2. McNeil JJ, Wolfe R, Woods RL, et al. Effect of Aspirin on Cardiovascular Events and Bleeding in the Healthy Elderly. *N Engl J Med*. 2018;379(16):1509-1518. doi:10.1056/NEJMoa1805819

3. McNeil JJ, Woods RL, Nelson MR, et al. Effect of Aspirin on Disability-free Survival in the Healthy Elderly. *N Engl J Med*. 2018;379(16):1499-1508. doi:10.1056/NEJMoa1800722

4. National Institutes of Health. Daily low-dose aspirin found to have no effect on healthy life span in older people. 2018. https://aspree.org/aus/wp-content/uploads/sites/2/NIH-PRESS-RELEASE-ASPREE-NEJM09.2018.pdf. Accessed July 16, 2019.

5. Lacaze P, Woods R, Zoungas S, McNeil J. The genomic potential of the Aspirin in Reducing Events in the Elderly and Statins in Reducing Events in the Elderly studies. *Intern Med J*. 2017;47(4):461-463. doi:10.1111/imj.13384

6. Lockery JE, Collyer TA, Abhayaratna WP, et al. Recruiting general practice patients for
large clinical trials: lessons from the Aspirin in Reducing Events in the Elderly (ASPREE) study. *Med J Aust.* 2019;210(4):168-173. doi:10.5694/mja2.12060

7. ASPREE. Protocol - ASPREE XT: A study of the long term effects of aspirin and other factors on the health of older persons. 2018. https://aspree.org/aus/wp-content/uploads/sites/2/ASPREE-XT-Protocol_Version_1_January2018.pdf. Accessed May 1, 2019.

8. WHO. Proposed working definition of an older person in Africa for the MDS Project. WHO. https://www.who.int/healthinfo/survey/ageingdefnolder/en/. Published 2016. Accessed July 21, 2019.

9. Mahmood SS, Levy D, Vasan RS, Wang TJ. The Framingham Heart Study and the epidemiology of cardiovascular disease: a historical perspective. *Lancet (London, England)*. 2014;383(9921):999-1008. doi:10.1016/S0140-6736(13)61752-3

10. Deary IJ, Gow AJ, Pattie A, Starr JM. Cohort Profile: The Lothian Birth Cohorts of 1921 and 1936. *Int J Epidemiol.* 2012;41(6):1576-1584. doi:10.1093/ije/dyr197

11. Simons LA, McCallum J, Simons J, et al. The Dubbo study: an Australian prospective community study of the health of elderly. *Aust N Z J Med.* 1990;20(6):783-789. doi:10.1111/j.1445-5994.1990.tb00423.x
12. Lacaze P, Woods R, Zoungas S, McNeil J. The genomic potential of the Aspirin in Reducing Events in the Elderly and Statins in Reducing Events in the Elderly studies. *Intern Med J.* 2017;47(4):461-463. doi:10.1111/imj.13384

13. Nunn JS, Shafee T, Chang S, et al. Standardised Data on Initiatives - STARDIT: Alpha Version. 2019. doi:10.31219/OSF.IO/5Q47H

14. Ryan J, Fransquet P, Wrigglesworth J, Lacaze P. Phenotypic Heterogeneity in Dementia: A Challenge for Epidemiology and Biomarker Studies. *Front Public Heal.* 2018;6:181. doi:10.3389/fpubh.2018.00181

15. Al-Shahi Salman R, Beller E, Kagan J, et al. Increasing value and reducing waste in biomedical research regulation and management. *Lancet (London, England).* 2014;383(9912):176-185. doi:10.1016/S0140-6736(13)62297-7

16. Crocker JC, Ricci-Cabello I, Parker A, et al. Impact of patient and public involvement on enrolment and retention in clinical trials: systematic review and meta-analysis. *BMJ.* 2018;363:k4738. doi:10.1136/BMJ.K4738

17. Gillies K, Chalmers I, Glasziou P, Elbourne D, Elliott J, Treweek S. Reducing research waste by promoting informed responses to invitations to participate in clinical trials. *Trials.* 2019;20(1):613. doi:10.1186/s13063-019-3704-x
18. Liabo K, Stewart R. Involvement in research without compromising research quality. J Heal Serv Res Policy. 2012;17(4):248-251. doi:10.1258/jhsrp.2012.011086

19. Nunn JS, Tiller J, Fransquet PD, Lacaze P. Public Involvement in Global Genomics Research: A Scoping Review. Front Public Heal. 2019;7:79. doi:10.3389/FPUBH.2019.00079

20. National Institute for Health Research. Briefing note eight: Ways that people can be involved in the research cycle. National Institute for Health Research. http://web.archive.org/web/20170605035051/http://www.invo.org.uk/posttyperesource/where-and-how-to-involve-in-the-research-cycle/. Published 2017. Accessed June 5, 2017.

21. National Institute for Health Research. Patient and public involvement in health and social care research. 2014. https://www.nihr.ac.uk/about-us/CCF/funding/how-we-can-help-you/RDS-PPI-Handbook-2014-v8-FINAL.pdf. Accessed February 2, 2018.

22. Burton H, Adams M, Bunton R, et al. Developing stakeholder involvement for introducing public health genomics into public policy. Public Health Genomics. 2009;12(1):11-19. doi:https://dx.doi.org/10.1159/000153426

23. Lemke AA, Harris-Wai JN. Stakeholder engagement in policy development: challenges and opportunities for human genomics. 2015. doi:10.1038/gim.2015.8
24. Deverka PA, Lavallee DC, Desai PJ, et al. Stakeholder participation in comparative effectiveness research: defining a framework for effective engagement. *J Comp Eff Res*. 2012;1(2):181-194. doi:10.2217/cer.12.7

25. Oliver S, Clarke-Jones L, Rees R, et al. Involving consumers in research and development agenda setting for the NHS: Developing an evidence-based approach. *Health Technol Assess (Rocky)*. 2004;8(15):1-148, III-IV. doi:10.3310/hta8150

26. Staniszewska S, Adebajo A, Barber R, et al. Developing the evidence base of patient and public involvement in health and social care research: the case for measuring impact. *Int J Consum Stud*. 2011;35(6):628-632. doi:10.1111/j.1470-6431.2011.01020.x

27. International Association for Public Participation. Participation Spectrum. https://www.iap2.org.au/Tenant/C0000004/00000001/files/IAP2_Public_Participation_Spectrum.pdf. Published 2014. Accessed July 12, 2018.

28. Macaulay AC. Participatory research: What is the history? Has the purpose changed? *Fam Pract*. 2016;351(3):cmw117. doi:10.1093/fampra/cmw117

29. Pandya-Wood R, Barron DS, Elliott J. A framework for public involvement at the design stage of NHS health and social care research: time to develop ethically conscious standards. *Res Involv Engagem*. 2017;3(1):6. doi:10.1186/s40900-017-0058-y
30. Collins M. PiiAF The Public Involvement Impact Assessment Framework Guidance. http://piiaf.org.uk/documents/piiaf-guidance-jan14.pdf. Published 2014. Accessed October 4, 2017.

31. Staniszewska S, Brett J, Simera I, et al. GRIPP2 reporting checklists: tools to improve reporting of patient and public involvement in research. 2017. doi:10.1186/s40900-017-0062-2

32. O’Brien BC, Harris IB, Beckman TJ, Reed DA, Cook DA. Standards for Reporting Qualitative Research. *Acad Med.* 2014;89(9):1245-1251. doi:10.1097/ACM.0000000000000388

33. Tong A, Sainsbury P, Craig J. Consolidated criteria for reporting qualitative research (COREQ): a 32-item checklist for interviews and focus groups. *Int J Qual Heal Care.* 2007;19(6):349-357. doi:10.1093/intqhc/mzm042

34. INVOLVE. *Guidance on Co-Producing a Research Project.*; 2018. http://www.invo.org.uk/wp-content/uploads/2018/03/Copro_Guidance_Mar18.pdf. Accessed March 14, 2018.

35. INVOLVE. *Co-Production in Action: Number One.* Southampton; 2019. https://www.invo.org.uk/wp-content/uploads/2019/07/Copro_In_Action_2019.pdf.
36. Prikken I, Burall S. Doing Public Dialogue. 2012;(July). https://www.involve.org.uk/wp-content/uploads/2012/01/120727-RCUK-Resource-FINAL.pdf. Accessed March 19, 2018.

37. INVOLVE. Guide on Co-Producing a Research Project.; 2018. https://www.invo.org.uk/posttypepublication/guidance-on-co-producing-a-research-project/. Accessed May 8, 2019.

38. INVOLVE. National Standards for Public Involvement.; 2018. http://www.invo.org.uk/posttypepublication/national-standards-for-public-involvement/. Accessed January 29, 2019.

39. International Collaboration for Participatory Health Research (ICPHR). Position Paper 1: What Is Participatory Health Research? Version: May 2013.; 2013. http://www.icphr.org/uploads/2/0/3/9/20399575/ichpr_position_paper_1_definition_version_may_2013.pdf. Accessed June 13, 2017.

40. Stake R. The Art of Case Study Research.; 1995. https://us.sagepub.com/en-us/nam/the-art-of-case-study-research/book4954. Accessed April 10, 2018.

41. Gerring J, McDermott R. An Experimental Template for Case Study Research. Am J Pol Sci. 2007;51(3):688-701. doi:10.1111/j.1540-5907.2007.00275.x
42. Seawright J, Gerring J. Case selection techniques in case study research: A menu of qualitative and quantitative options. *Polit Res Q*. 2008. doi:10.1177/1065912907313077

43. Staniszewska S, Brett J, Mockford C, Barber R. The GRIPP checklist: Strengthening the quality of patient and public involvement reporting in research. 2011;4:391-399. doi:10.1017/S0266462311000481

44. Concannon TW, Meissner P, Grunbaum JA, et al. A new taxonomy for stakeholder engagement in patient-centered outcomes research. *J Gen Intern Med*. 2012;27(8):985-991. doi:10.1007/s11606-012-2037-1

45. Bowen GA. Document Analysis as a Qualitative Research Method. *Qual Res J*. 2009;9(2):27-40. doi:10.3316/qrj0902027

46. Saldaña J. *The Coding Manual for Qualitative Researchers (2nd Ed.)*. SAGE Publications; 2013. doi:10.1017/CBO9781107415324.004

47. Nunn JS, National Institute for Health Research, Support MC. *Building Research Partnerships*. 2012. https://learnzone.org.uk/downloads/Building Research Partnerships - 2013 Report - Macmillan NIHR CRN.pdf. Accessed July 21, 2019.

48. Guest G, MacQueen K, Namey E. *Applied Thematic Analysis*. 2455 Teller Road, Thousand Oaks California 91320 United States: SAGE Publications, Inc.; 2012.
49. Hyett N, Kenny A, Dickson-Swift V. Methodology or method? A critical review of qualitative case study reports. *Int J Qual Stud Health Well-being*. 2014;9(1).
doi:10.3402/qhw.v9.23606

50. Kemmis S, Nixon R, McTaggart R. *The Action Research Planner: Doing Critical Participatory Action Research.*; 2014. doi:10.1007/978-981-4560-67-2

51. Cook T, Abma T, Gibbs L, et al. *Position Paper No. 3: Impact in Participatory Health Research.*; 2020.
http://www.icphr.org/uploads/2/0/3/9/20399575/icphr_position_paper_3_impact_-march_2020__1_.pdf. Accessed May 24, 2020.

52. Cotterell P, Morris C, Harlow G, Beresford P. Making user involvement effective: lessons from cancer care. 2009. https://eprints.soton.ac.uk/184013/. Accessed July 28, 2017.

53. Gamble C, Dudley L, Allam A, et al. Patient and public involvement in the early stages of clinical trial development: a systematic cohort investigation. *BMJ Open*. 2014;4(7):e005234. doi:10.1136/bmjopen-2014-005234

54. Centre for Research in Public Health and Community Care. *Guidance for Researchers: Feedback from Researchers to PPI Contributors.*; 2018. https://www.clahrc-
55. Gradinger F, Britten N, Wyatt K, et al. Values associated with public involvement in health and social care research: a narrative review. *Health Expect.* 2015;18(5):661-675. doi:10.1111/hex.12158

56. Genetic Alliance Australia Ltd. *Australian Patients’ and Families’ Perspectives on Genome Sequencing.*; 2016. http://www.geneticalliance.org.au/cmsAdmin/uploads/genetic-alliance-genome-sequencing-report-final.pdf.

57. Metcalfe SA, Hickerton C, Savard J, et al. Australians’ perspectives on support around use of personal genomic testing: Findings from the Genioz study. *Eur J Med Genet.* November 2018. doi:10.1016/J.EJMG.2018.11.002

58. Bilkey GA, Burns BL, Coles EP, et al. Genomic Testing for Human Health and Disease Across the Life Cycle: Applications and Ethical, Legal, and Social Challenges. *Front Public Heal.* 2019;7:40. doi:10.3389/fpubh.2019.00040

59. Etchegary H, Wilson B, Rahman P, Simmonds C, Pullman D. Public interest in whole genome sequencing and information needs: an online survey study. *Per Med.* June
Beaton A, Hudson M, Milne M, et al. Engaging Māori in biobanking and genomic research: a model for biobanks to guide culturally informed governance, operational, and community engagement activities. *Genet Med*. 2017;19(3):345-351. doi:10.1038/gim.2016.111

Ennis L, Wykes T, Callaghan P, et al. Impact of patient involvement in mental health research: longitudinal study. *Br J Psychiatry*. 2013;203(5):381-386. doi:10.1192/bjp.bp.112.119818

Guest G, Bunce A, Johnson L. How Many Interviews Are Enough? *Field methods*. 2006;18(1):59-82. doi:10.1177/1525822X05279903

Seidman I. *Interviewing as Qualitative Research: A Guide for Researchers in Education and the Social Sciences*. Teachers College Press; 2006.

https://books.google.com.au/books?id=pk1Rmq-Y15QC&dq=Seidman,+2006),&lr=&source=gbs_navlinks_s. Accessed February 18, 2019.

ASPREE. ASPREE Protocol. 2014. https://aspree.org/usa/wp-content/uploads/sites/3/2014/04/ASPREE-Protocol-Version-9_-Nov2014_FINAL.pdf.
Accessed July 16, 2019.
Additional files

Additional File 1 - Data and Analysis - Detailed description of the data sources in this case study, how they were analysed, including STARDIT reports.

Additional File 2 - Telephone Questionnaire - Version Comparison - A comparison of the versions of the telephone questionnaire used.

Additional File 3 - Quantitative results - A summary of the quantitative results

Additional File 4 - GRIPP2 report

Additional File 5 - STARDIT report
### Table 5: Enablers and barriers for involving participants

| Enablers | Quotations | Barriers | Quotations |
|----------|------------|----------|------------|
| Financial remuneration for people’s time; financial support for travel and accommodation | Running a business ‘limits me and my time’ [P15] | Living in rural areas and travel logistics a barrier to participation in face-to-face events | ‘I’d like to be involved online rather than face to face because of travel difficulties’ [P6] |
| Learning and development opportunities for participants and researchers | ‘if you tell me what would be useful, I could do it’ [P13] | Poor explanation of abstract concepts such as ‘involvement’, which can be jargonistic | ‘there seemed to be confusion, or a lack of understanding of what this ‘involvement’ would actually look like’ (MS) |
| Small groups at events gave more people a chance to share perspectives | ‘Small group’ discussions at the event ‘worked well’ | Self-doubt about their skills or knowledge mean they don’t think they could be involved | ‘I’d probably ask a stupid question’ [P2] |
| Early notice of events | ‘give me enough notice’ [P10] | Some people not comfortable being part of a face-to-face group | ‘I get very uncomfortable in a group of people’ [P12] |
| Clear information about timings and | Ensure ‘people are advised’ what’s going to happen at the workshop’ [P8] | Lack of clarity about expected time commitments | Being involved ‘depends on what’s involved and time’ [P1] |
|-----------------------------------|-------------------------------------------------|-----------------------------------------------|--------------------------------------------------------|
| time commitments, frequency of involvement and available support | ‘What’s the endpoint – what’s the goal?’ [P11] | Unclear about what tasks they could be involved in | ‘I don’t know how I could but willing to help’ [P13] |
| Clear information about purpose and expectations of involvement, feeling their involvement has consequences | Independent facilitator when working in groups (either face-to-face or online) | I’d feel more comfortable if I had someone who was facilitating [P14] | Face to face discussions ‘dominated’ by more confident or knowledgeable people |
| | | | ‘Participants from professional backgrounds to some extent dominated some of the discussions’ (Study team) |
| Short events ensure people do not get fatigued | ‘any longer and I think fatigue would have dampened the enthusiasm’ (Study team) | People feel they have limited time, are busy with work or social commitments | ‘I don’t have a lot of time left in life’ [P1] |
| Having access and literacy in using computers and online tools | ‘if I could negotiate [online discussions] I’d be happy to do that’ [P15] | Lacking access, literacy or trust in using computers, smartphones or | ‘I’m hopeless with computers’ [P16], ‘I don’t have internet’ [P12] |
online tools (including social media)

| A selection of flexible communication modes (such as face to face and facilitated online discussion forums) | ‘I’d be happy to be involved – more online but if there was an occasional need to come into the city I’d be happy to do that’ [P3] | Only one mode of communication, such as expecting people to travel to events | ‘online is often easier’, face to face only ‘as long as it’s not too far’ [P22], ‘travel distance is an issue’ [P20] |
|---------------------------------------------------------|---------------------------------------------------------------------------------|------------------------------------------------------------------|-----------------------------------------------------------------------|
| Involving people in research ethics and governance | ‘ethics is the difference between right and wrong – you know what’s right and you don’t do what’s wrong’ [P18] | A ‘researchers know best’ attitude that doesn’t value the process of involving people (MS) | Researchers ‘don’t see the forest for the trees’ [P19] |
Figures

Figure 1

Process timeline of the participant involvement
Co-designing a multi-generational genomics study

**ASPREE**

The ASPirin in Reducing Events in the Elderly (ASPREE) trial has **over 19K participants**

ASPREE has **DNA** from **over 15K participants**, with **90%** still attending annual visits 5 years on

**Stage 1: Planning**

Multi-generational research study proposed by stakeholders

Stakeholders **co-designed involvement and ethics**

**Stage 2: Interviews**

Newsletter sent to **14,268 participants** inviting them to get involved in **co-designing study**

Telephone interviews with 59 participants are **analysed, data informs event design**

**Stage 3: Event**

18 participants attend **co-designed event** to explore views about study and preferences for involvement

**Stage 4: Evaluation**

Study team interviewed and data from the entire 18 month process is **co-analysed with stakeholders**

**Involving participants resulted in multiple impacts**

**Changes** to the proposed **study design** included:

- **Recruitment done online, saliva collected by post** instead of blood samples
- Participants control data access
- **No data access for profit**

Involve participants **online and in-person**

**Figure 2**

Co-designing multi-generational genomics research

**Supplementary Files**

This is a list of supplementary files associated with this preprint. Click to download.
- AdditionalFile1DataandAnalysis.pdf
- AdditionalFile2TelephoneQuestionnaire.xlsx
- AdditionalFile3QuantitativeResults.xlsx
- AdditionalFile4GRIPP2.pdf
- AdditionalFile5STARDITAlphaVersionReport.pdf