Developing research resources And minimum data set for Care Homes' Adoption and use (the DACHA study)

PLAIN ENGLISH SUMMARY

BACKGROUND

Approximately 420,000 people in England and Wales live in a care home, many with multiple health problems and other life-limiting conditions, including 70% with dementia. Care homes, commissioners and regulators collect and hold a lot of information about residents' medical history, needs, preferences, and care. At present, there is no consistent approach to how this is done with consent (including those with dementia) nor how it is used securely. Different people collect different information in different ways, making it difficult to understand the care needs of residents, how these are met as they move between organisations. Consequences include duplication of effort, some needs not being identified or reviewed, feelings of frustration and failures in communication. At a national level, it is difficult to plan for future needs of residents, required treatments and services, and to research their effectiveness.

A more consistent approach to how information about care home residents is recorded and handled is needed. An agreed data set would help researchers and practitioners enhance the care and quality of life for residents, families, and staff. RESEARCH QUESTIONS

- What is already known about how multiple health and social care organisations are working with and for care home residents by sharing systems of information (data)?
- How can existing and future collection and use of residents' data be optimised for planning and delivering care and research?
- What is the minimum amount of information that it is feasible and appropriate for care homes to collect routinely and how that can be linked to routinely held information by GPs and hospitals?

STUDY

First, the study will include a review of what outcomes are commonly measured in care home research, to help to identify potential items to include in an agreed data set, and help researchers planning future research. A second review of evaluations of intervention research studies in care homes will give information about what happens when undertaking research and using different outcome measures in the day to day setting of a care home. These findings will be compared with what care home residents, family members and care home staff involved in providing day-to-day care identify as information that is important to them.

We will bring together existing data, collected by various research teams throughout the UK. Often, teams have collected similar data about different care home residents, for example, medication records, or studies may use the same questionnaires to measure outcomes. By bringing these existing data together and asking new questions that had not been researched in the original studies, we can understand the care home population better. We already have agreement from five research teams that we can pool their data, and plan to ask more as new studies are published.

To address what is already known and what works when using residents' data to inform care, we will review the kind of resident data that are currently collected and what needs to be in place for them to be useful. We will also survey English care homes to discover what information they are already collecting about their residents. These findings will be compared with what care home residents, family members and care home staff involved in providing day-to-day care identify as information that is important. Next, we will test how to combine information which is already routinely collected for health and social care with new information to achieve consistent ways of organising data for the benefit of residents and researchers. We will consult with stakeholders to develop a schedule of questions and linked outcome measures that could be the basis of a national database. Working with IT specialists, we will test these questions and the feasibility of asking them in practice.

Throughout the study we will work with all interested parties (commissioners, care home owners, residents and their representatives, NHS staff and the regulator, researchers and data management and privacy experts) to develop a consensus on the minimum content of shared records, guidance on how to standardise the way it is recorded and outcomes measured and linked to existing routinely collected data. We will test the feasibility of using the shared care record in geographically dissimilar areas that are already working to integrate health and social care data. Throughout the study, we aim to create new ways of working and doing research in and with care homes, so that the outputs benefit not only researchers, but also residents.

SUMMARY OF RESEARCH (ABSTRACT)¹

BACKGROUND In England, long-term continuing care for older people is principally provided by care homes. Residents and staff rely on the NHS for medical care and access to specialist nursing and therapy services. The creation of Integrated Care Systems https://www.england.nhs.uk/integratedcare/integrated-care-systems/ in England and recognition of the role of social care as an essential part of care provision for this population underline the need to develop reciprocal systems of working between the NHS and care homes that optimise current provision and research on its effectiveness. Within ten years future demand for long term care will outstrip current capacity (1, 2). There is currently no agreed framework for collating, sharing and interpreting data collected by the stakeholder organisations involved in supporting care homes – this is a missed opportunity for co-ordination of care, service development and commissioning, as well as the conduct of research. Other countries have highly evolved minimum datasets which have shown utility in all of these areas.

AIMS

To establish what data need to be in place to support research, service development and uptake of innovation in care homes

To synthesise existing evidence and data sources with care home generated resident data to deliver a minimum data set (MDS) that is usable and authoritative for different user groups (residents, relatives, business, practitioners, academics, regulators and commissioners).

DESIGN A mixed method study drawing on design and implementation theories.

¹ Glossary of abbreviations included with reference list

- Work package (WP) 1 Evidence reviews on what improves the productivity of care homes research and key measures and outcomes that have been incorporated in UK care home research that could inform a MDS.
- WP 2 Creation of a care home trial repository, initially including trial data on 6000 residents, with the ability to add new trials beyond the project duration. This will allow secondary data analysis and test key resident characteristics and outcomes relevant to the development of a MDS
- WP 3 To establish what is known about the implementation and content of MDS relevant to UK systems of care. A realist review building on WP1 to understand how MDS work in different care home settings and the attributes and situations in which their use supports improved outcomes for residents, family, staff and organisations. A scoping review of the content of Minimum data sets relevant for English systems of care MDS and survey of existing care home generated data to inform the development of a MDS.
- WP 4 Mapping and characterisation of existing sources of data on care home residents to create resident datasets from routine NHS and Local Authority data in two integrated care sites (ICS) to link with data collected in study care homes. Combined with WP1-3 findings and national consultation this will enable critical appraisal of feasibility of inclusion of assessment and outcome measures in a care home generated MDS for testing
- WP 5 Pilot and test the MDS by collecting data at three time points on 300 residents of care homes across two integrated care systems (ICS). We will pilot an electronic prototype interface to collect the care home components of the MDS, alongside the protocol for integrating these with native data from NHS and social care databases to form the full dataset. Data from the MDS will be collected at baseline, 6 and 12 months to understand the impact of seasonality. Focus groups conducted in care homes at each time point will develop an understanding of implementation issues including the perceived utility of the MDS data to key stakeholders. Descriptive statistics will be used to understand the measurement attributes of the MDS alongside factor analysis support item reduction for the MDS.
- Consultation with stakeholders: This is embedded throughout study: A national expert group of up to 40 resident and relative representatives, care home owner and staff representatives, commissioners of health and social care to care homes, researchers and NHS providers and staff will convene at three points in the study. This will inform the planning of WP 1-4, develop a consensus on proposed MDS for testing, and develop recommendations for implementation. Ensuring this element of co-production is part of the project throughout will ensure outputs are fit for use and "shovel ready".

PROPOSED OUTPUTS

- i) Guidance for researchers and commissioners on health care implementation in care homes, resource on assessment tools and core outcomes in care home research;
- ii) A repository of care home RCTs to support secondary data analysis and inform MDS development that can support more research and useful findings for priority questions, without the expense and burden of involving care homes and residents in research
- iii) Evidence on what needs to be in place for MDS to support staff uptake, resident assessment care and improved outcomes
- iv) An implementation strategy for MDS that links NHS, social care and care home generated data
- v) A prototype MDS to inform commissioning, needs assessment and care delivery.

BENEFITS There is widespread interest in how to improve the uptake of best evidence and care for care home residents. Combining existing data with care home generated data in the development of the MDS could improve the quality of life of care home residents and their experience of care and reduce NHS and related costs. By working closely with resident representatives, the care home industry, NHS England, Local Authorities, commissioners and the regulator this study addresses policy objectives of integrated care for this group with a paradigm shift towards individual and care home level information being routinely shared and used to underpin research, innovation and intervention.

BACKGROUND AND RATIONALE

There are approximately 420,000 older people living in long term care settings (3, 4). They are among the most frail and vulnerable in society and rely on the NHS for access to medical care, specialist nursing, and therapy (5, 6). Care homes are a sector under pressure both financially and because of rising numbers of the oldest old (1, 2, 7) and this directly affects how the whole health and social care system performs. Care home providers and NHS partners need to understand who their residents are and what kind of services enable them to live and die well. Capacity to assess quality, monitor resource use and care costs, and evaluate and spread innovations is radically undermined by a lack of strategy and standardisation of how resident needs and service use are documented.

There is a growing abundance of data about the care home sector and the residents it serves from both service delivery and research data. These data are not, though, currently harnessed in an effective way to the benefit of the sector and the residents it serves.

Care homes compile data to plan and document care, to inform business plans, and as evidence for regulators, Local Authorities, and NHS commissioners. Data on residents are also generated by NHS practitioners (e.g. GPs, community nurses, therapists, and paramedics) when they visit care homes and when residents are admitted to hospital. Increasingly they are also generating resident specific data via the use of sensor technologies that monitor movement and vital signs (8). In England some work has been done to systematise resident data (9) and Scotland will have an integrated dataset (10) by early autumn. There are, however, no mechanised systems to aggregate this data at a care home level that captures the resident experience of care even in these pioneering areas.

From a research perspective, equivocal findings and underused data from robust RCTs conducted in care home environments are a further concern (11). Researchers require care home specific evidence on how intervention processes and organisational context affect uptake, and how best to collect and measure data (12, 13). Optimising existing data to maximise learning about residents from previous research (14) would help to target future expenditure, minimise risks of repeating identifiable failings in trial implementation, and maximise a return on investment that has already been made in research.

Consumers (residents and relatives) also want information about care homes (15) with value placed on indicators of residents' quality of life as well as satisfaction with services (15-17). More efficient use of existing data and the creation of Minimum Data Sets (MDS) could answer key service delivery and research questions about the way NHS services are delivered to care homes, the impact of different models of care, or which services are needed by which subgroups of residents and how resident data are shared (5, 18). There is a need to map current information provision and how it can be organised and standardised to inform and translate securely to IT systems that all parties want to use, and will be able to use, to support the planning and provision of care to care home residents.

Minimum datasets provide a comprehensive account of resident characteristics, resource use, and quality of care outcomes in key areas (e.g. functional ability, pain, and infections (19-24). MDS can also help to inform planning and evaluation of care and research from assessment on admission to end of life (25, 26). One of the most widely used MDS, the international Resident Assessment Instrument (interRAI), was developed in the late 1980s and implemented in North American nursing homes in 1996 (20, 27) and is now used in many countries internationally. Countries often use MDS because they are mandated and/or linked to reimbursement systems and quality monitoring (28, 29). What is less clear is how the priorities of social care providers, residents and families are addressed, how the resources required to sustain the system are established, and how use of MDS supports change in care practices and personalised care within individual care homes (30).

This submission responds to all parts of the commissioning brief.

Why the research is needed now

The recent NIHR themed review of care home research (11) concluded that research in care homes is relatively undeveloped. It recommended that future work is grounded in the reality of care home life, supporting partnership working between health professionals, staff, residents, and relatives in designing and delivering new approaches to care, and this proposal arises from such collaboration. The importance of building capacity in care home research and the need for a minimum data set to improve understanding of how care for residents is planned, delivered, and reviewed were identified as priorities. Discussions about the proposal with representatives from: care home chains, NHS England, Care Quality Commission (CQC), representatives from the Cabinet Office working group on social care measurement confirm that this work is both timely and urgently needed.

In England, the absence of a national mandate, lack of links with NHS data, and implementation challenges (31) have meant that MDS and data-driven approaches to resident assessment have been limited to single projects e.g.(32, 33). Routinely collected data on resident and care home characteristics (e.g. by GPs), RCT data from care home research, and studies of health and social care outcomes specific to social care (16) have the potential to inform a MDS specific to the national situation. The challenge is to establish systems of assessment and documentation that are evidence-based and usable by all those involved in receiving, providing, commissioning, and regulating care.

Test-bed initiatives supported by NHS England (Care home Vanguards), The Health Foundation, Nuffield Trust (34, 35) and NIHR-funded research (36, 37) have demonstrated the benefits for residents and staff of close working relationships between care homes and the NHS. The New Care Models Programme (36), by beginning to integrate NHS data sources, demonstrates that health and care professionals can access the necessary information to provide treatment and support. Research is needed that exploits these developments to improve the care of residents.

In July 2018, at a meeting of the National Care Home Research and Development Forum, 67 participants representing commissioners, family carers, regulators, care homes, and the NHS described local initiatives where health and social care services are sharing data e.g. to support safeguarding and risk reduction. There were also a few examples of shared access to cloud-based systems. This work with key stakeholders confirmed that a MDS is recognised as a needed resource when the care of frail older people requires the ability of different services and practitioners to work across health and social care organisations. (38, 39).

AIMS

To provide resources that support research and uptake of innovation in care homes.

To synthesise existing evidence and data sources with care-home-generated resident data to deliver a Minimum Data Set that is usable and authoritative for different user groups (residents, relatives, business, practitioners, academics, regulators, and commissioners).

Objectives

To establish evidence-based resources that can both improve the productivity of care home research and capture the key measures and outcomes used in UK care home research (MDS) (Work packages (WP) 1&2)

To develop an understanding of how a MDS could work in the UK context and develop a MDS that meets the needs and priorities of commissioners, providers, regulators, and recipients of care. (WP3 &5)

To demonstrate how the creation of administrative datasets on care home residents derived from routine health and social care data can complement care-home-generated resident data (WP 4&5)

To assess the feasibility of collecting data directly from care homes and matching this to native data to populate a complete MDS (WP 3 &5)

Demonstrate the utility of the matched MDS data to different stakeholders involved in commissioning, providing, receiving, and regulating care (WP 3-5)

RESEARCH PLAN AND METHODS

Design and Theoretical Framework

As an overarching approach to think through implementation issues (and theory) and how different stakeholders will inform the co-design of a MDS we have used the Double-Diamond design process (www.bsbd.org.uk/double-diamond-design-process) to structure the work packages (WPs) of this mixed methods study. There are four stages involved in this process:

- 1. **Discover** open up and explore the focus of the project and needs of knowledge users (WP1-3)
- 2. Define focus in on the important issues to tackle and define problems to be addressed. (WP1-3)
- 3. **Develop** collectively design and test potential solutions to the problem (corresponding to the national stakeholder consultation and WPs 4-5)
- 4. Deliver narrow in on a practical, working solution and implement it. (WP 4-5).

This approach to implementation is guided by the principles of integrated knowledge translation (IKT), that emphasizes collaboration between researchers and decision-makers (40) and by design-based approaches to co-production (41-44). Each of these is based in the idea that collaboration between those who produce research and those who use it will improve its quality, relevance, and usefulness. From IKT we take the broad framing of knowledge co-production based in an ongoing relationship between researchers and decision-makers or knowledge users (such as clinicians, managers, or policy-makers). This ensures the production of mutually beneficial research that both supports decision-making and the kind of group-level identity transformation needed to support the development and delivery of a MDS for health and social care (40, 45, 46).

National Consultation with expert reference group on Minimum Data Set development

In addition to Patient and Public Involvement and Engagement (PPIE), and to support the codesign approach throughout the study, we will convene a national expert consultation group meeting as four regional groups (North-East, West, and East of England, Yorkshire and Humber) organised in collaboration with the Collaboration for leadership in Applied Health Research and Care and the Academic Health Science Networks (CLAHRC/AHSN) national care home collaboration. This will involve ten representatives per region meeting three times during the study. Meeting 1 will review the findings of WP1-3 to validate/debate the findings and prioritise learning about implementation and outcomes of interest. Meeting 2 will review the proposed MDS developed from the combined evidence from WP 1-4. Meeting 3 will prioritise and agree the proposed MDS prior to implementation. This approach recognises the importance of emphasizing process and of ensuring we bring together all participants residents, families, staff, managers, clinicians, commissioners and regulators - in a way that enables us to collectively elicit and share experiences and perspectives and to engage in "collective making" as a form of knowledge mobilisation (42). We will work with the conceptualisation of stakeholder engagement proposed by Boaz and colleagues (41) and with their proposed 15 design principles for stakeholder engagement.

Work package 1 (WP) Research in and with care homes

This work responds to the research brief's identification of the need for evidence that can both improve the productivity of care home research and capture the key measures and outcomes used in UK care home research that could inform a MDS. The evidence reviews meet the call requirements as described in the Scope/Part 1:- 'Research must include a review of completed studies in the UK' and Scope/Part 2.5 'Within this topic there is the option to include research into systems for capturing and using quality of life, quality of care, relational and patient-centred care measures and how these might be used at scale for individual or care home measurement'.

Review One will provide an overview of resident reported outcomes and care quality relating to interventions or programme implementation in care homes, and any evidence of sensitivity to change. Attention will be paid to the use of functional and quality of life measures. Based on our prior work we anticipate considerable heterogeneity and knowledge gaps (e.g. in resident/family reported outcome measures). By identifying those outcomes measured in care home studies that are sensitive to change, clinically meaningful, and address the lived experience of the older people resident in care homes this review will both be a resource for researchers planning future work and inform the work of developing a MDS.

The review will include RCTs, pragmatic trials, natural experiments and implementation studies. It will focus on the range of interventions conducted in care homes since 2009 to reflect contemporary research practice. Included studies will be categorised by focus and outcome categories (e.g. cognitive ability, function, quality of life and care) and analysis will consider frequency of use across studies and data on the feasibility and utility of the outcome measures used.

Review 1 will systematically identify trials, outcome and contextual measures that have been conducted in care homes to directly inform WP2 and 3. It will collate a list of outcome measures and other contextual measures of relevance to resident outcomes that have been used in care home studies to date. Measures of relevance to quality of life, quality of care, relational and patient-centred care will be identified for consideration for possible inclusion in the MDS. The analysis will consider the frequency of a measure's use across studies and the feasibility and utility of a measure within an MDS, and any evidence of sensitivity to change. The data

collected will inform the prioritisation of datasets of most value to the MDS for linkage in WP4 and decisions on the content of the final MDS for piloting in WP5

Review Two will provide a synthesis of the evidence from process evaluations of complex interventions conducted in UK care homes. The process of delivering an intervention in a care home setting can mediate the validity of the outcome measures directly. A process evaluation describes how delivery is achieved, what is actually delivered, how the delivered intervention was reported as producing change, and what affected implementation and outcomes (47, 48). WP1 will synthesise implementation factors that have been argued within published process evaluations of care home research to have influenced study outcomes.

Evidence from review 2 will inform the pilot implementation of the MDS in WP5 and inform the interview topics for the assessment of potential barriers and facilitators to wider implementation in WP5. This will be complemented by an early deliverable; a guide to researchers, to support and improve the design, conduct, and impact of future health services research in the care home sector.

Review 2, by producing a synthesis of process evaluations of care home research, will provide further in-depth evidence of factors that influence resident outcomes that may not have been directly measured in research or data systems to date, and that could be of importance to care home research in the future that could be considered for inclusion within the MDS. This evidence synthesis will identify common factors from previously published studies and programmes that have reported a process evaluation that may affect outcomes, fidelity, and quality of implementation. Taking a human-factors approach to data organisation and analysis, it will examine the relationship between the organisational context of care homes involving a subset of embedded work systems: i) physical environment, ii) tools and technologies, iii) care tasks, and iv) the carer/resident. This fits with the design principles of stakeholder engagement (41) with care home staff that places the resident at the centre of the work system model.

Search strategy: Both reviews will use high quality evidence synthesis methods. Protocols will be registered on PROSPERO and conduct and reporting will follow the guidelines of the PRISMA statement where appropriate. Searches for relevant evidence will include databases of peer-reviewed literature (MEDLINE, EMBASE, CINAHL, Social Sciences Citation Index) and sources that include grey literature (SIGLE, Open Grey, Google Scholar. Studies for inclusion will be limited to English language and of relevance to the UK care home context. To assess quality of conduct and reporting of studies, studies will be assessed against the MRC recommended criteria for process evaluation (<u>https://mrc.ukri.org/documents/pdf/mrc-phsrn-process-evaluation-guidance-final/</u>).GRADE-CERQual will be used to assess the overall strength of the available evidence to guide recommendations (<u>www.cerqual.org/</u>).

A preliminary scoping of the literature for Review One identified 75-100 studies offering UK relevant evidence. A preliminary scoping exercise for Review Two identified 12 studies with a published process evaluation, and 14 ongoing process evaluations in a care home setting.

WP1 will identify a list of salient contextual factors that influence implementation of care interventions in a care home setting. This will also inform implementation of the MDS. The output will be guidance and resources for researchers and commissioners aiming to test and implement health and care interventions in care homes.

Work Package 2 Creation of an individual patient data (IPD) repository of UK care home trials

WP2 complements WP1 and responds to the research brief's identification of "the potential for secondary data analysis of existing research data to track residents' outcomes before and

after significant events or new approaches to providing health and social care". This WP will develop a repository of individual resident data from existing care home trials. Secondary analysis of existing trial data is low-risk, low-cost, and a priority for NIHR (49). Large RCTs conducted solely in care homes are a growing resource. While these RCTs focus on a variety of health/care topics (e.g. falls risk, medication management, nutrition, or infection) there is much overlap in outcome measures used, and information collected on both residents and the care home structure. Trials in care homes monitor participants regularly, for up to one year. Outcome measures, health resource use, and clinical events as well as care home characteristics can therefore be tracked over this period, allowing for longitudinal analysis. If individual patient data (IPD) from existing trials could be pooled, they would collectively provide a much larger and more useful dataset. Individual patient data (IPD) can be used for exploratory analysis to better understand this population, reduce duplication of effort, and refine future research questions (50-52).

To enable development of the repository, we will first complete a scoping review to identify potential care home trials for inclusion. Initial work has identified large trials conducted in UK care homes (see table below). We have secured the agreement of five lead investigators, who will work with us. Work-package coordinator (LI) has a working knowledge of one of these datasets (54), as trial health economist. Additional trials will be identified systematically through the reviews in WP1, and by contacting all trialists listed in the NIHR "Advancing Care" Themed Review (11) (44 studies featured), the CLAHRC National Work stream Report (55) (32 studies featured), and snowballing techniques. We have set up a database of potential studies for inclusion that monitors how IPD is requested and managed and that logs all contacts with trialists and respective ENRICH units.

A collaboration of trialists will make up the repository Steering Committee, to oversee sharing, combining and repurposing of their existing trial data. Together they will agree ground rules for the collaboration, including approval process for data requests. Based on conversations and confirmation of intention to participate, the repository will combine trial data for over 4200 residents from 254 care homes across the UK, with the potential to add seven more trials (representing 4166 residents from 384 care homes) as the repository develops.

| Trial | OPERA (56) | ChallengeDe mCare | DCM- EPIC(57) | CHIPPS (54) | ELECTRIC(58) |
|-----------------------|-------------------------|-------------------------|-------------------------|---------------------------|--|
| Data available | Now | Now | 2019 | 2021 | 2021 |
| PI | Underwood | Moniz-Cook | Surr | Wright | Booth |
| Location | England | England | England | England, Scotland & NI | England & Scotland |
| N participants | 1054 | 832 | 1020 | 880 | 500 |
| N care homes | 77 | 63 | 50 | 44 | 20 |
| Follow-up (months) | 12 | 12 | 16 | 6 | 4 |
| Treatment focus | Exercise for depression | People with Dementia | People with Dementia | Medication management | People with urinary Incontinence |

Initial trials to include in Repository

The repository will be set up as a parallel to (59, 60) similar collaborations (notably the VISTA stroke repository) and have infrastructure to host data securely and expertise to manage future trial inclusion and data access requests beyond the duration of DACHA.

NIHR 127234 DACHA study

Informed by WP1 and national consultation, key variables will be prioritised. Study characteristics (inclusion/exclusion, trial methodology, and intervention details), care home specific details (staff ratios, staff retention, bed size, case mix, ownership) and individual participant level data (demographics, outcome measures, resource use, and medications) will be included. Variable names will be standardized and data reorganized so that *pooled* individual patient data can be presented to future users in user-friendly datasets. We will report baseline characteristics about care homes and residents as derived from all pooled trial data, focusing on what data from trials may be appropriate to include in a MDS framework. Possible proxy measures for the outcomes listed in WP4 will be also be explored, such as using prescribing data to identify long term conditions. We will also evaluate how generalizable the repository data is, compared to alternative data sources. Specifically, we will compare point estimates from data derived in WP4 (data collected from administrative sources e.g. CQC) and WP5 (data prospectively collected within the MDS framework). This will test if there a difference in care home resident's participation in RCTs and observational studies and what can be derived from GP, hospital and social care

We will apply for HRA and CAG approval and complete a Data Protection Impact Assessment to demonstrate GDPR compliance for the trial repository. Each contributing trial will hold ethical approval to collect the primary data. Trialists who contribute their data for use in the DACHA study will complete a gatekeeper agreement, identifying them as the gatekeeper for their respective datasets. We will request that all data received will be fully de-personalised (such as converting 'date of birth' to 'age at randomisation'). Therefore, only *completely anonymised data* will be held in the repository.

The repository will host trials with a range of clinical focus – it is therefore likely that some measures will be unique to single trials. However, several key outcome measures – e.g. Barthel; EQ5D; DEMQol, MMSE, are used in almost all RCTs conducted in care homes (and will contribute to the research questions under examination. Additionally, clinical indicators such as hospitalisations, falls, and death rates are routinely reported.

Data will be compiled based on the availability of pre-specified outcome measures, in part identified through WP1 evidence reviews. The identification and critique of relevant outcome measures within existing trials will help inform the development of a prototype MDS (WP4). Where queries exist on submitted measures, these will be resolved through examination of the accompanying documents that support the datasets (e.g. papers, funders reports, data dictionaries), through review of existing, published assessment tools, and through correspondence with contributors to clarify queries on the assessment tools used, including versions, modifications and appropriate interpretation.

Dependent on adequate sample size, we would like to extend the analysis of the IPD data by performing mapping (crosswalks) between outcome measures

Finally, drawing on the findings from WPs 1&3, feedback from the national expert consultation group, and with PPIE input (see text box below) we will prioritise questions for future use of the repository data. For example can we use trial IPD to identify sub-populations of residents who are most likely to benefit from particular types of interventions?

The input of the consultation group will also be important at this stage in ensuring that the thinking around these issues remains oriented to the ultimate goal of successful implementation.

The repository will be a lasting legacy of the DACHA programme – a valuable source of highquality, anonymised, individual participants' data (IPD) to inform the development of future research, testing of hypotheses and optimisation of study design issues. Data will not be used to re-examine the original research questions set out by the primary studies, but anonymised data will be available to the wider research community for further examination in novel exploratory analyses.

National Expert Consultation Group Meeting 1: Recruitment of the 4 regional groups (North East, East and West of England, East Midlands and Yorkshire) will be organised through the Care Home Collaboration. We will purposively recruit local care home managers (2) Family members (2) Commissioners from NHS and LA (2), Local CQC regulator (1), GP, community therapist/ nurse (2), IT/software expert working with care homes (1). They will meet for half a day to review the findings of WP1-2 to validate/debate their relevance for a MDS. Using nominal group techniques the groups will prioritise learning about implementation and outcomes of interest. Researchers will facilitate and record the discussions to ensure all views are captured and where necessary follow up with telephone interviews for points of clarification or where there were polarised views within the group. Findings from the four groups will be grouped and synthesised to refine the MDS matrix development (see WP3) and fed back to participants for final comment.

Work package 3 (WP) Development and implementation of a MDS

This WP has 3 elements: (i) a realist review; (ii) mapping of care home generated data, and (iii) a consultation and priority setting exercise. The aims are to:

- Develop an understanding of how a MDS could work in the UK context,
- Identify care home generated data that could contribute to a MDS
- Generate a matrix of potential variables and their characteristics, and identify stakeholder priorities for inclusion in a MDS.

(i) Realist review: The purpose of the realist review in WP3 is to develop a theory-driven understanding of how internationally deployed MDS systems offer transferable learning and/or utility for the UK. Interrogating research on how MDS have been used by care home staff will increase understanding of how different contexts, assumptions and mechanisms affect its feasibility, and relevance to practice and for different groups of residents.

We will draw on realist methods (61) to develop a theory driven understanding of how internationally deployed MDS systems offer transferable learning and/or utility for the UK. The realist approach assumes that causal mechanisms are embedded within particular contexts and social processes, and that it is crucial to understand the relationship between these mechanisms and the effect that context has on their operationalization and outcome (62).

Established MDS aim to support comprehensive assessment of the resident, their needs and linked care planning (63). The Inter RAI (International Resident Assessment Instrument) is the most commonly used MDS. It consists of a data collection form, a set of assessment items with identical definitions, time frames for observation specific to the long term care setting, built in protocols for clinical and quality assessment, triggers that identify residents at risk, status and outcome measures and user manual. Research has demonstrated the value of a MDS to commissioners and service providers in enabling identification of care needs and residents at risk of ill health (20, 64-68). In North America, where its use is mandated, it is used in large scale research and cross national comparisons, for example in measuring

the quality of care in different facilities. Our NIHR OPTIMAL study on NHS working with care homes employed an adapted InterRAI. Whilst it was possible to identify resident characteristics associated with unplanned hospital admission, some measures were not appropriate for UK residents, staff found it added to the burden of record keeping plus, its use did not influence care decisions. Other studies identify the need to integrate the use of MDS into the routines of the care home to achieve a sustained impact on resident outcomes (69). There is also evidence that residents with worse health status are more likely to have missing data, it adds to work stress (affecting staff retention) and does not always capture what is important to residents and carers (27, 70-72)

To develop a theoretical understanding of how effective implementation of a MDS improves resident and organisational outcomes within the current care system the review will have three iterative stages: (1) Synthesis of review findings from WP1, linked work on integration of NHS, social care and CQC data, the use of Comprehensive Geriatric Assessment (73) in care homes and scoping of how MDS is represented as changing staff behaviours, practice and quality of care to achieve improved resident and service outcomes; (2) Development of likely 'Context-Mechanism-Outcome' (C-M-O) configurations to build an explanatory theory(or theories) of how a MDS might work in the UK that are tested in the evidence on MDS implementation; (3) analysis and synthesis of the programme theory (61, 74, 75).

Searches will be conducted from 2005 (to reflect the growing interest in MDS related research) purposive, iterative and including MEDLINE, EMBASE, CINAHL, Social Sciences Citation Index, sources of grey literature and Google Scholar and lateral searches.

The outcomes of interest will be informed by WP1 as well as resident identified priorities for care from the PPIE work (if different to the research evidence) but will include, resident outcomes, staff adherence to MDS processes and resources used. The review will consider recurring patterns of association between contexts and mechanisms (demi-regularities) detectable across studies. It is likely analysis will be informed by systems thinking and organisational theories of change that focus on the way that values and beliefs defined at a strategic level are embedded across the workforce (76, 77)

Findings from the review will be tested further in four group interviews with frontline staff in two areas (East of England, West of England) working with staff from Quantum Care (who use predominately paper based records supplemented with resident specific digital data from monitoring technology) and Somerset Care (who have used e records for over 15 years). The realist review findings will directly inform how the development and introduction of a MDS in WP5 is conceptualised and implemented.

(ii) Scoping of international MDS content and mapping of care home generated data for inclusion in MDS: To identify the full range of data variables, scales and outcome measures used in MDS across the world, we will conduct a rapid scoping review of international MDS research on resident assessment and care. Using the libraries generated by searches for WP1 and the realist review, this review will chart the variables/outcome measures used and care processes or experiences evaluated. Our focus will be on outcomes relevant to resident quality of life and the provision/monitoring of care, but not available from existing health and social care data (e.g. on pain, social engagement). It recognises that in addition to the programmes of work in specific topic areas (e.g. pain, depression) undertaken by the Inter-RAI consortium there is research that addresses other measures such as quality of end of life care, quality of life and testing the validity (or not) of oral health measures used within existing MDS.Research that identifies gaps in the assessment and care of older people is also relevant. For example, a review on the implementation of Comprehensive Geriatric Assessment (CGA) found that frailty had not

been considered as a determinant of outcomes among older people. A realist review of CGA by the team (CG, ALG) in care home settings signalled the importance of knowing when and who from the health and care home teams is involved in assessment and review of care.

We will use established methods (78, 79) and apply strict inclusion criteria (English language publications since 2005 identified by searches in WP1 and 2, observational or experimental study designs, data generated by MDS in long term care facilities for older adults). Data will be extracted and charted without quality assessment. This work will produce an accessible guide to the current content of international MDS, and relate each data item to the component of resident care or experience being measured. Of direct relevance to WP4 and 5, the content review will also be a freely available resource for researchers, supporting the development of international comparative studies and MDS development.

(iii) Mapping of care home generated data for potential inclusion in MDS: Our previous work with care homes has identified multiple data sources specific to the care home that are not synthesised, shared or electronically stored. Care home providers collect information to support care delivery and for reporting to commissioners and regulatory authorities. This includes technology generated data on resident vital signs and mobility, medication intake, preferences and risk assessments. This component of WP3 aims to map data collected by care home providers for their own use that has potential for inclusion in an MDS. We will compile a list of the content, form and use of care home generated data, and assess the willingness of care home providers to contribute data items to an MDS that is shared with external partners such as the NHS.

Participants: Working with a purposive sample of English care providers (110 or 20% of the 5500 organisations who run the 11300 care homes in the UK) that will include the five largest providers who are responsible for over 20% of beds. The sampling matrix will account for funding source for care homes (not/for profit) and residents (self-funder, local authority, NHS continuing care); size of organisation; bed numbers; nursing/residential status; CQC rating and geographical location. We would aim to achieve a response rate of 60%. An adequate response rate to ensure that we have captured the range of data collected by care homes would be between 40-50%.

Intervention: The findings of the realist and scoping reviews will be used to develop a semi structured telephone and online survey for care home providers and local regulators. The semi-structured telephone and online survey for care home providers and local regulators will collect objective data on what and how information (on care processes and delivery, resident characteristics, health status, functional ability and quality of life) is collected, collated and shared within care homes and provider organisations. Our experience of working with commissioners and care providers in this area, points to recent, rapid growth in new data collection initiatives in care homes. For example, use of the National Early Warning Score in care homes is now encouraged in some regions, with NHS support for digital data collection. The WP3 survey work is essential to capture information on recent developments of this type that have yet to feature in published academic articles. We will also produce an accessible guide to the current content of international MDS, and relate each data item to the component of resident care or experience being measured.

We will address the issue of commercial sensitivity, with carefully worded questions on willingness to contribute to an MDS and barriers to data sharing. The survey will be discussed, piloted and launched with the cooperation of the Care Providers Alliance (who represent 11 national associations of independent and voluntary adult social care providers) and study steering group members from LaingBuisson, Four Seasons, Quantum Care and

the Registered Nursing Home Association. Follow up by email (x2) and telephone (x2) will be used to boost response rate.

Outcomes: An overview of the range of data generated and collated by care home providers in England to support care delivery and to fulfil regulatory and governance requirements.

Matrix development. The findings of the scoping review and care home survey will be synthesised into a matrix that presents potential MDS content items, alongside characteristics such as ease of access, availability, cost of collection, and perspective (resident, clinical provider, resource manager, etc.). This visual display of potential MDS content, with pros and cons of individual items and scales clearly articulated, will be a resource for use in a priority setting exercise with the expert national collaboration (see below) and in discussion with stakeholders and the research team. Used in this way, it will inform the content and direction of WP5

Consultation and priority setting: Working with the national expert consultation group (see text box below) we will run a consultation exercise across the 4 regions. This consultation will use the evidence and expert opinion to build on the earlier stakeholder input to rank the evidence for resource requirements, ease of access, utility and relevance of items and scales identified.

Work Package 4: Data linkage of existing administrative data sources to inform the prototype MDS.

The NHS collects a huge amount of data, including the electronic medical record in general practice, administrative hospital data, and operational data sets from the emergency services, urgent care and community health. Local authorities (LA) also collect data on packages of social care funded publicly, and needs assessments. This work package will identify, document (create metadata) and link these existing data sources together to establish a basis for the MDS. This will demonstrate what a MDS could look like without the need of additional data collection by care homes.

Led by The Health Foundation (THF) who have experience bringing data together for research and evaluation within its secure data environment (34, 80-84) this WP focuses on two Integrated Care Systems (ICS) sites who have agreed to participate (Surrey Heartlands & Nottinghamshire) to:

- i. Identify and document sources of data on **all** care home residents in the two ICS held by health and social care.
- ii. Create two linked administrative datasets on care home residents, one for each ICS.
- iii. Demonstrate the benefit of routinely linking these data together by providing descriptive analysis of the linked datasets, and share the results with the ICS and care home managers involved.

(i) Identify and document sources of data on care home residents held by health and social care: To test the feasibility of using data routinely held by NHS and Social Care to populate a MDS for improving quality of care as well as for research. The team will:

• Map out the data sets available in two Integrated Care Service sites, including existing information about the contents and scope of the data (Meta data), existing information governance agreements, and privacy impact assessments.

• Explore using address information stored in GP records. Address information can be used to identify care home residents that are self-funded, and not included in LA datasets.

Informed by the national findings from the expert consultation group, by input from the care home based PPIE groups and the PPIE panel and WP 1-3 the data sets will be prioritised based on which are likely to produce most value to the minimum data set. It will include hospital administrative data (from the Secondary Uses Service), GP records, local authority social care data, and data from urgent and emergency care and community health.

(ii) Create two linked administrative datasets on care home residents, one for each ICS. Based on the priorities identified above, relevant data sets will be transferred, linked and processed in the secure data environment at THF. This is a complex process that involves agreeing with each of the relevant stakeholders the purposes of the data collection, the way in which patients and the public will be kept informed of the data processing, and the safeguards that will be put in place. All data will be processed in the ISO 27001 accredited secure data environment at the Health Foundation. It will also be 'pseudonymised' before the data are transferred. However, three linkage keys will be created: two of these will relate to the individual (one formed by encrypting the NHS number, and another by encrypting a combination of the person's initials, gender and date of birth, to be used to validate the linkage by NHS number or when the NHS number is not present) (81). The other key will relate to the address of the person involved, formed by encrypting the unique property reference number (84). We will support the data providers to assign the unique property reference number to their data sets where they are not already present.

The data linkage process will operate in two stages, the first will identify the information required to create the linkage keys required to link the data. Once created, we will extract the relevant information on care home residents from the various administrative data sources. Information on clinical needs will be collected from NHS data (for example GP prescriptions and hospital data), and other information such as mobility assessments from social care datasets.

(iii) Analysis of the linked datasets, and presentation of results with the ICS and care home managers. Using the administrative datasets, we will demonstrate the benefit of routinely linking these data by providing descriptive analysis of these data to stakeholder in the ICS sites and care home managers involved in WP 5. Previous engagement with stake holders (34) demonstrated that descriptive statistics (e.g. a breakdown of the reason of admission to hospital for potentially avoidable admissions at care home level) are useful for care home managers. The specification of the descriptive analysis will be agreed with local stakeholders to ensure relevance and make explicit the data from routine data that could be linked with a care home generated MDS.

Once the linked datasets are established, we will liaise with analytical teams to gauge interest in using these data for further analysis and/or evaluation to further maximise the benefit of linking these datasets (e.g. the Improvement Analytics Unit, a partnership between NHS England and the Health Foundation, works closely with ICSs to evaluate their impact and might benefit from an existing linked dataset). This additional analysis is outside the scope of the current grant application.

National Expert Consultation Group Meeting 2: Priority setting

"Meeting two with the regional expert groups will review the proposed MDS that is developed from the combined evidence from WP 1-4. Using nominal group techniques to engage with experts, and additional meetings with staff, resident and family groups organised in 4 care homes from the integrated care system sites of Work Package 5. This will build on the earlier stakeholder input to rank the evidence for utility and relevance of items and scales identified. This consultation will use the evidence and expert opinion to help resolve questions around: What to measure (and how), at what level to capture it (individual level so it can be linked to other sources or aggregate care home level to compare services) and How to share it (securely and appropriately) and with whom. Findings will feed into WP5

WP5 Longitudinal pilot of the MDS This WP has four aims:

- Assess the feasibility of collecting data directly from care homes and matching this to native data to populate a complete MDS;
- Assess the quality of the MDS data, comprising care home-collected data alongside native data, to create a MDS with the minimum number of scales/attributes required;
- Demonstrate the utility of the matched MDS data to stakeholders (ICSs, CCGs, local authorities, providers and residents and their families)
- Systematically assess potential barriers and facilitators to wider implementation of the MDS

Design: A longitudinal pilot of the MDS (including an electronic prototype collecting the care home components of the MDS and a protocol for integrating these with native data from NHS and social care databases to form the full dataset) in 20 care homes in each of our two partner ICS sites (N=40 homes).

WPs 4 and 5 will work with the two ICSs to identify four Clinical Commissioning Groups (CCGs), two per ICS site, committed to supporting a pilot study in care homes in their areas and working with us to identify priority research and service development questions that the MDS might help them address at the local level.

Work packages (1-4) will have informed the content of the MDS, taking into account its utility to key stakeholders in our two ICS sites (WP4). The final MDS, in addition to routinely collected data and care home-generated indicators is likely to include validated measures of assessment of resident priorities, function, symptoms, treatment and measures specific to social care. We will also capture data from care homes at baseline on the proportion of self-funders, number of NHS continuing care packages and funding to support rehabilitation and respite care. Although the final content of the MDS will not be finalised until year three (after WPs1-4), preliminary ethical and governance approvals will be sought in years one and two (including approval from the Association of Directors of Adult Social Services ADASS). Clinical Research Networks will support resident recruitment within participating care homes.

Represenation Representation is based on identifying the proportion of the population of care home residents in the two Integrated Care Systems (ICS) with a certain characteristic or outcome (e.g. proportion conveyed to hospital) using a confidence interval-based sample size. Using Surrey Heartlands as the example, assuming 50% of those in the population of 7,560 care home residents (90% occupancy of 8,399 beds) have the factor of interest, a sample of 240 residents would be required to estimate the expected proportion with 5% precision and 90% confidence. As such, if 50% of the 240 residents had the factor of

interest, we would be 90% confident that 45%-55% of the care home population have the factor of interest.

Sampling: The study teams at Kent and Nottingham will each recruit and work with 20 care homes for older adults (total N=40 homes). Data held by the Care Quality Commission indicates that Surrey Heartlands has 202 care homes (8,399 beds) for older adults across three CCGs and Nottinghamshire has 236 homes (8,541 beds) across six CCGs. Care homes for learning disabilities will not be included in this study.

Assuming an occupancy rate of 90% (4), the sample size required to give a true representation of the finite older care home population, with 90% confidence and 5% margin of error, is 237-260 residents, from the smallest (in Nottinghamshire) to largest (in Surrey Heartlands) two CCGs by care home beds (85). To achieve this sample size, we will first stratify care home selection by size, maintaining an even split of nursing and residential homes in each ICS, where possible.

Small care homes (fewer than ten beds) will be excluded. Although small care homes (less than 20 beds) still account for a large percentage of the market. However, this situation is rapidly changing, mainly because of the economic pressures the sector is experiencing. The average size of care homes ceasing to operate in 2016- 17 was 29. There are fewer, on average larger, care homes, but it is the smaller homes that are closing. https://ipc.brookes.ac.uk//publications/Market Shaping in Adult Social Care.pd

Smaller care homes are excluded for three reasons,

- To reflect current changes in the sector
- Residents in smaller homes are likely to differ from those in larger homes which would potentially create a small level of bias. For example, catering to a younger population with learning or long term physical and mental health problems requiring 24/7 care. The focus of this work is older residents
- To ensure an adequate sample size (i.e. to avoid under-recruitment).

Larger care homes in each ICS will be over-selected – we anticipate that approximately two out of every three care homes selected will be above average size (40 beds). For recruitment, we will then utilise a census approach where all eligible care home residents (long-stay/permanent beds) in participating homes will be invited to take part. We assume that there will be a 33% response rate using this approach (86). With 90% occupancy, we anticipate a sample of 320 in Surrey Heartlands ICS and 280 in Nottinghamshire ICS. This sample size allows for sample attrition of approximately 1 in 6 residents across each ICS. Median length of stay for residents in care homes has been estimated at 15-18 months in England (87, 88), but this will vary by location and provider. We will liaise with care home managers to assess any potential issue with attrition and look to identify ways to resolve this, e.g. target higher response rates for specific homes, re-sampling of residents.

All permanent/long-stay residents will be eligible for inclusion in the study. We will support recruitment of adults without mental capacity by using established consultee process and an appointed independent consultee for residents with nobody to act on their behalf. Exclusion criteria would include residents receiving respite or on temporary/short-stay care, or people identified by the care home staff as being in the last week of life at the point of study inception. Temporary or short stay residents (e.g. intermediate care) represent the minority of residents in care homes. For the development of the MDS, we want to focus on identifying data that reflects how health and social care needs are documented over time. If however,

the consultation and data collection demonstrate that this group should be included (for example because the NHS often pays for continuing and intermediate care) then that will be addressed and this group will be included.

Training To provide appropriate support and training participant homes in each ICS will be broadly grouped into three cohorts, according to their readiness to participate using questions based on the domains of the Alberta Context Assessment tool (89) and developed by members of the team for NHS England Vanguard sites (<u>http://vuh-la-uhra.herts.ac.uk/handle/2299/18200</u>). We expect care homes to identify 2-3 key members of staff to have responsibility for completing the MDS and propose to run regional training in each ICS, one per cohort, for staff from participating homes to attend (N=40-60 members of staff trained in each ICS). We will begin with the care homes most ready to participate (cohort one), so that learning from these sessions can improve the training and support for cohorts of homes identified as needing more support. We anticipate training will require 2 days. It will include: familiarisation with the study, the scales included in the electronic prototype and practising completing scales, entering the data online and information about data security and storage.

Method The frequency of data input for the prototype MDS will have been determined as part of the co-design process in WP4. If this emulates MDS systems used in other countries, it is likely that data will be inputted monthly. To understand how this is operating, we propose a longitudinal design with three data collection points (baseline, 4 and 12 months), consisting of an audit of the data from all consenting eligible care home residents in participating homes.

A longitudinal design will enable us to assess the feasibility of the tool by looking at completion rates and implementation over time and will also provide insights into seasonal variation. This will enable seasonal factors to be controlled for in the analysis when responding to the priority questions identified by the CCGs (e.g. hospital admissions from care homes and what affects this). It will also generate data into how seasonal pressures on the care home sector affect data completion rates.

Linking the primary data collected with the resident specific administrative linked data developed in WP4 about the residents we will further compare what information can be accurately captured in existing administrative data, and what data has to be collected at care home level to create a valuable MDS. Linking the two datasets together provides an opportunity to validate the identification of care home residents in the administrative data, especially where self-funders are concerned.

The precise means of collecting data directly from care homes to complete the minimum dataset will be considered as part of the co-design process and informed by earlier work streams.Our current proposal, (subject to further exploration as part of the study) though, is to use REDCAP cloud, for which the University of Nottingham holds a license, for homes to upload pseudonymised data. The pseudokey and identifiers would be shared separately with staff at the Health Foundation using a password protected encrypted file (AES 256). They would then use this to link the pseudonymised data from REDCAP with native variables from NHS and social care datasets.

After each wave of data collection, we will give participating homes from each cohort aggregated summaries of their MDS data, benchmarked against other resident data from the whole sample in their ICS. This will include administrative comparisons, such as MDS completion rates, as well as benchmarking indicators of residents' health, use of services

and outcomes (e.g. ASCOT, if collected). Previous research (Towers et al, 2016, submitted) suggests homes find this information valuable and that it can increase participation in research.

| Cohort in each ICS | Training | Time 1(baseline) | Focus Groups | Time 2 (5-6 months) | Focus Group | Time 3 (11- 12 months) | Focus Group |
|--------------------------|----------|---------------------|-----------------|---------------------------|----------------|---------------------------|----------------|
| 1 (most ready) | Month 26 | Month 27 | Month 28 | Month 32 | Month 33 | Month 38 | Month 39 |
| 2 | Month 28 | Month 29 | Month 30 | Month 34 | Month 35 | Month 40 | Month 41 |
| 3 (least ready) | Month 29 | Month 30 | Month 31 | Month 35 | Month 36 | Month 41 | Month 42 |

The trained care home staff in each ICS cohort will be brought back together after each wave of data collection to participate in focus group interviews (up to 2 hours long) exploring:

- Implementation issues
- Applicability and utility for the sector
- Modifications to the MDS.

Focus groups will be facilitated by two researchers and audio-recorded. We will keep focus group interviews to a manageable size (up to 10 per group) running additional groups to accommodate the views of staff from all participating homes, if required. At the end of data collection period, we will conduct telephone or face-to-face interviews with key representatives in each ICS or CCG (n=6, 3 interviews per ICS) to explore their views of the implementation process and the relevance of the data to their local priorities. To systematically assess potential barriers and facilitators to the wider implementation of the MDS, the implementation aspects of these focus groups will be structured using the Consolidated Framework for Implementation Research (CFIR) (90).

A final stakeholder event will be held in each participating ICS area to present summary data on the MDS, constituent variables and linked health and social care outcomes. Group discussions will explore the utility and feasibility of the final MDS.

Data synthesis and analysis WP5 will generate care home level data from the MDS and pilot the process of matching this to native data sources, collated by The Health Foundation. In addition, there will be field notes (from training and support given by the research team), focus group data and interview data. As such, there will be both quantitative and qualitative data collected by the study at different levels (resident, care home, CCG and ICS). Plans for analysis of this data are described according to the aims they address below:

Assess the feasibility of collecting data directly from care homes and matching this to native data to populate a MDS Analysis of the implementation process will be structured to test the findings of WP4 of what enables uptake and use of MDS in care home settings and what is perceived by different stakeholders as key to successful uptake (shared aims, utility, ease of use)., Data from each ICS will be analysed separately, to identify local area themes and then together to identify common themes for national learning. We propose to use Nvivo 11 for windows (QSR International Pty Ltd) to store and analyse the qualitative data.

Assess the quality of the MDS data and explore the psychometric properties of the measure as whole, with a view to item reduction and improving factor structure

Working with Kent's psychometrics researcher Stephen Allan, basic descriptive statistics will describe measurement attributes of variables within the MDS, including completion rates, median/mean and measures of distribution, and floor and ceiling effects (proportion of candidates returning lowest or highest score respectively). Regression analysis will be used to consider the influence of baseline care home contextual factors from the CFIR tool on completion rates.

Exploratory factor analysis will be used to identify the underlying factor structure of the data held in the MDS and to identify any redundant items. Suitability for exploratory factor analysis will be assessed using Bartlett's test of sphericity and the Kaiser-Meyer-Olkin (KMO) measure of sampling adequacy. It is of interest to further explore the opportunity for variable reduction, with a view to making the final MDS as minimal as possible whilst describing all necessary attributes. The factor structure of variables held in the MDS will be assessed and subsequently reduced using Rasch analysis (partial credit model). Rasch analysis assesses whether items or variables in a measure capture the same latent trait, and this approach is routinely used to reduce the number of items in a scale (Rasch, 1960). Item goodness-of-fit will be assessed using infit mean square <1.2. Variables in the MDS exceeding the cut-off criteria will be subsequently removed. For dimensionality, the variance for the Rasch model is expected to explain atleast 50% variance, with the remaining components explaining < 5% of the variance (eigenvalue < 2.0). We will also examine the internal consistency of participants' responses (i.e. whether items tap into a single underlying construct) using Cronbach's alpha statistic (Cronbach, 1951). If the internal reliability improves by removing one or any set of variables from the MDS, these will be removed.

Demonstrate the utility of the matched MDS data to the local CCGs. To demonstrate the utility of MDS data, we will work with The Health Foundation team and the four CCGs that have supported the pilot studies in care homes to assess priority areas. There are a number of potential analyses and the choice of which to use will depend on the data held in the MDS. These include substitution analysis, drivers of average total cost, and evaluating specific interventions. For example, one area that could be explored is the impact of social care related quality of life on health care use, the latter measured by number of GP visits, hospitalisations, total costs to the NHS, etc. Another would be exploring the sensitivity of outcomes measures in the MDS to changes in residents' health and functional abilities over time, controlling for baseline characteristics. Quantitative analysis using appropriate econometric techniques will be used to analyse the data, exploiting the longitudinal nature of the data.

National Expert Consultation Group Meeting 3

The final meeting will be held in London and will bring together the participants from the four regional groups and the Study Steering Committee. Participants will receive a briefing prior to attending. The findings will be presented and followed by facilitated group discussions of their implications and of recommendations that will feed into the final report. Previous consultation work has demonstrated the value of discussion being organised according to background (e.g. expert by experience, NHS and care home staff, commissioners, IT and software developers, Regulator) and then meeting to discuss and review their recommendations and priorities for implementation.

DISSEMINATION, OUTPUTS AND ANTICIPATED IMPACT

The goal is to produce resources for those researching, living in, and working in and with care homes and to ensure these are clinically relevant and nationally informative. The national collaborations and organisational partners are the basis of our dissemination strategy.

We will strengthen the dissemination to care home residents and their families through a number of reports throughout the lifecycle of the project, tailored to be accessible to residents and their families, to be shared through national and local networks including Age UK, Alzheimer's Society, My Home Life, Residents and Relatives Association, older peoples' fora (such as for example Norfolk Older People's Strategic Partnership, National Pensioners' Convention). Reports will also be made available to appropriate magazines such as Saga, and newspapers with an appropriate readership. Information will also be made available via social media such as Facebook and YouTube and via local and national radio, e.g. BBC Radio 4 Inside Health. For researchers, and drawing on current practice from related studies, blogs from PPIE members will be included as part of the study's updates and online presence and be featured on the ENRICH website.

We will ensure that the 2 care home based Patient and Public Involvement and Engagement (PPIE) groups, and the PPIE panel meet at appropriate intervals to be able to contribute to the outputs from Work Packages 1, 2, 3, 4 and 5. We will also extend PPIE to include the perspectives of Providers (Patient, Public, and Provider Involvement and Engagement) and explore with the PPIE group how best to engage staff in discussion (either separately or together).

Where this is agreed with PPIE/PPPIE members to be appropriate, we will disseminate our evidence-based messages using real-life illustrative stories told from the perspective of residents, relatives and staff, in their own words. These stories will be incorporated into the deliverables from the study (please also see response to question about useful outputs).

The study will have a website, and maintain a social media presence (monthly blog, twitter, Facebook). Alongside the national consultation and review process that is threaded through the project each WP will work with the CLAHRC (ARCS) in the care home collaboration, to share briefings about findings and resources relevant to their partners and to ensure that local implementation issues are identified early. Conference presentations and papers in professional and academic journals will be produced from each WP. We will organise 3 half-day dissemination events targeting different audiences and will work with the Kings Fund and My Home Life, the Professional Records Standardisation Board, Health Education England and the Care Provider Alliance.

Outputs

WP1 – Care home resource and guidance (available online and as briefings) for researchers and those supporting care home research and innovation

WP2 - Trial repository (ongoing development post-funding)

WP3 – Evidence of what is known about the implementation and content of Minimum data sets relevant for English systems of care and a prototype care home generated resident MDS

- WP4 Resident dataset developed from routinely collected health and social care data
- WP5 Prototype MDS that aligns with clinical and care records standards

Whole project outputs

- Guidance for the public on how to access information on resident outcomes and approaches to their measurement
- An implementation strategy for MDS that links NHS, Local Authority, and other care home generated data

- A resource on agreed core outcomes for care home research, together with recommended assessment tools
- Evidence on what needs to be in place to support MDS uptake by researchers, commissioners and providers to inform resident care planning and evaluate outcomes

The linked dataset would be available for analytical teams outside the Health Foundation as well. Setting this up would be part of the project. Assuming this does not incur additional costs for access, this should be free of charge. It is technically feasible to provide remote access to the HF secure environment, once IG is addressed.

Anticipated Impact for stakeholders

For residents and family: US research using MDS has demonstrated how information about resource use and resident care can inform care home selection and quality assessment. A MDS supports identification of residents' care priorities, implementation of evidence based approaches, ensures secure transfer of personal information and reduces the burden of research on the individual.

Care home staff & providers: A MDS prioritises collection of resident focused information, increases visibility of residents' needs and areas for building capacity in the workforce to support them. Through integration of data it makes explicit care homes' roles and impact in intermediate care, palliative care and service innovation. It reduces research burden on care home staff.

MDS as a resource for integrated working: Shared care records and the use of linked health and social care datasets reduce duplication of effort, identify care gaps and improve productivity. Previous research and the Vanguard care homes demonstrated the importance of recognising care homes as part of the health care economy. A MDS could support working methods that facilitate assessment, review, and continuity of care across organisations.

Information Governance

The review work, the regional consultations, the prior experience of the Health Foundation team and involvement of key stakeholders in the participating ICS study sites will contribute to the development of these protocols with particular attention to the specific needs of care home and other third sector providers

Supporting equity of access to health and social care: A 20 year history of research describes the ad hoc nature and inequity of health care provision to care home residents. This research provides the resource to address this and ensure equity of access to health care.

Research and planning: Despite a proliferation of care evaluation measurement tools many are of questionable quality and relevance to the UK systems of care. The development of a RCT repository for secondary data analysis on key issues (e.g. medication optimisation), resources to build research capacity in care homes plus standardized, organised, accessible MDS offers the resources necessary to measure the impact of different initiatives and models of service delivery. As noted above, a MDS enables implementation of evidence-based practices and thus will offer researchers a powerful tool to support the co-production of research and implementation of research findings.

For care home providers: The study addresses the heterogeneity of the care home market and ever increasing range of IT solutions for data capture. A MDS can inform how future systems standardise data for individual care management and service improvement **National government and wider society:** Understanding the role of care homes in the continuum of care for complex adults. An improved understanding of the care home population and evidence of the positive role care homes play in UK society would contribute to improving attitudes towards long-term care provision and resource

Further development: if we are able to demonstrate the feasibility and utility of a <u>Minimum</u> Data Set. We anticipate that it will be added to and adapted as care in this sector and digital capability expands. Further work will be needed to standardise its use for system wide uptake across the different providers and developers of IT systems for care homes

PROJECT / RESEARCH TIMETABLE

| Plan of investigation and timetable | |
|---|-----------------------------------|
| Research team set up meetings. Meetings with staff in Integrated Care Systems sites to put in place permissions and governance for accessing data (Nottinghamshire and Surrey Heartlands) Recruitment national expert consultation groups in four regions. Confirmation of study steering committee | Completed prior to start of study |
| Set up 2 care home based Patient and Public Involvement and Engagement (PPIE) groups to meet regularly throughout the project. Set up a PPIE panel to meet regularly throughout the project (AK,SF JM,IL,SB) | 1-3 months then ongoing |
| Work package 1 and 2 Reviews and trial repository Design development and testing of care home research resources (GP,SK,LI,JB with input from CG,AK, AG) | 1-18m |
| WP 3 Realist review, scoping review, survey and consultation with expert consultation group Development of prototype MDS (CG,BH, KS,SB with input from ALL) | 7-21m |
| WP 4 Integration of resident health and social care resident data sets in Nottinghamshire and Surrey Heartlands(AS, with input from JB,AMT,AG,CG) | 12-36m |
| National expert consultation meetings X 3 Meeting1:10m Meeting2: 24m Meeting 3 38m (CG,KS,IL,I | BH,AG,JM,SB,LI) |
| WP 5: Ethics submission, recruitment and MDS intervention and analysis(AMT,AG with input from ALL) | 20 -44m |
| Dissemination of preliminary findings and expert consultation Final Report Workshops and dissemination of findings (ALL) | 38-48m |

PROJECT MANAGEMENT

Claire Goodman will lead and oversee the project supported by an administrator. The research team will meet every 8 weeks alternating face to face meetings in London with ZOOM meetings. Work Package leads and their research staff will meet monthly and circulate updates to the rest of the team.

The study steering committee chaired by Des Kelly (Residents and Relative Association trustee) will meet every six months. Agreed members include **PPIE**; John Wilmott, John Thurman (carers with family in care homes). **Data analysis, synthesis and datasets** William Laing (LaingBuisson), Professors Mike Clarke (individual participant data (IPD methods) Ruth Hancock (LA and care home datasets), Esme Moniz-Cook (care home outcome measurement) **Care home sector**: Maria Ball (Quantum Care), Roberta Roccella (Four Seasons Health Care), **Health**: Peter O'Brien GP commissioner, William Roberts (former NHS England National Lead, Enhanced Health in Care Homes now Head of Health and Social Care Innovation Unit, **Data governance**: Lorraine Foley (Professional Record Standards Body), **IT** Iain Turner National Registered Nursing Home Association (expertise in digital inclusion)

Jonathan Papworth CEO Person Centred Software, Tom Ward, (Head of Health and Care Data Exchange CQC and NHS tbc). **International advisor**: Dr George Heckman (InterRAI Fellow University of Waterloo Research Institute for Aging Canada).

ETHICS AND REGULATORY APPROVALS

The study will adhere to the UK Framework for Health and Social Care research and require HRA and Association of Directors of Adult Social Service (ADASS) approvals. Working with The Health Foundation team we will establish the individual permissions, information governance and approvals required to support the resident data sharing and commercial sensitivities. Where additional data has to be collected prospectively, and then linked, we will work to a consent acceptable to NHS digital.

PATIENT AND PUBLIC INVOLVEMENT AND ENGAGEMENT

Sue Fortescue, a former carer and retired IT specialist, is a team member. Building on the preparation of the bid and links with representative organisations the aim is to have an infrastructure that fosters ongoing conversations, both formal and informal, both within and across work packages. To keep the person being cared for at the centre of our thinking in ways that inform delivery of care/patient benefit we will convene two care home based resident PPI groups that meet over the life of the study.

Previous work by members of the team (91) have identified the value of developing job specifications with PPIE members to ensure that expectations around involvement are shared. To achieve this, and maintain continuity of involvement for the life of the study a UH researcher will be the link person with administrative support for PPIE representatives providing feedback on study progress and the impact of their involvement (92, 93).

In addition to the PPIE involvement in the national expert consultation group and the study steering groups. Involvement from staff in direct care roles and management roles, family carers and representatives of people living with dementia will be achieved by setting up a PPIE panel that meets throughout the life of the research project and offers a hub-and-spoke model of PPIE where each work package is working with up to three PPIE representatives.

PROJECT RESEARCH / EXPERTISE

The research team has extensive experience of working in and with care homes and have in the last 10 years together and independently completed over 35 externally funded care home studies. Particular expertise includes social care and building capacity in care home research (JM,AMT,SB,KS,AK,AG) social care and primary care workforce (KS, BH CG) integrated working between health and social care (CG BH AG), Geriatric medicine, nursing and quality improvement(ALG,JB,SB,CG,JM), Health economics(LI), implementation science (IL,CG) data analytics, tracking residents' care across health systems, large health and social care datasets (AS,JB,AG), PPIE (SF,AK JM,IL) Evidence review, realist review and data synthesis(CG,KS GP,SK), method expertise in assessment tool and MDS development (AG,AMT).

SUCCESS CRITERIA AND BARRIERS TO PROPOSED WORK

The success of the study will be judged by

• The level of engagement and consensus achieved at each stage of the study from care home representatives, residents and families, health and social care commissioners, providers and the regulator

- The delivery of online and published research resources evaluated as useful by researchers and care home staff and a RCT repository that can improve the effectiveness and efficiency of care home research
- The delivery of a MDS that links routine resident data with care home generated data that supports commissioning, planning, delivery and evaluation of resident care

| RISKS | Mitigation |
|--------------------|---|
| The project is too | The study has addressed the whole commissioning brief linking findings about the range of |
| ambitious and not | measures and outcomes used in care home research, implementation challenges with work on |
| achievable within | the development and use of a MDS. The allocation of senior staff to lead each WP and build a |
| the timescale of | cumulative programme of work ensures that it will deliver to time and target. |
| the study | |
| | Success relies on the active engagement of the different stakeholders. The proposal is |
| | grounded in a national collaboration of care home researchers working with local NIHR |
| | research organisations, social care providers (National Care Forum, Care England, Care |
| | Providers Alliance) and relatives of residents. Complemented by links with Your Care Rating |
| | (www.yourcarerating.org) CQC, Cabinet Office, international care home networks and MDS |
| | initiatives. The need for a MDS to standardise how information is shared and improve the public |
| | understanding of care homes is a priority. The study is ambitious but there is the cross sector |
| | commitment, expertise and a national infrastructure of support to sustain it to completion. |
| The research | The NIHR Dissemination Centre review (11) emphasised the need to move beyond descriptive |
| resources and | work, resources and a repository for secondary data analysis builds research capacity and the |
| RCT repository | scope for analysis of resident and staff needs. |
| are not used | |
| Heterogeneity of | There is a cross sector consensus that a workable MDS could support residents 'access to |
| care home market | health care, experience of care and quality review and reduce duplication of effort. This project |
| &multiple systems | provides the structure for standardisation and cross sector working. Work packages 1-4 |
| of data collection | systematically address the range of resident data and related outcomes, how care homes use |
| leading to partial | and collect data and the potential to link this to existing datasets to create a MDS. The |
| uptake and | involvement of the Professional Records and Standardisation Board in the consultation and |
| resistance to | dissemination events ensures that outputs from the project could be incorporated into national |
| change | records. |
| Existing MDS and | This study addresses the content, utility and feasibility of using existing MDS. We have argued |
| linked commercial | they provide a partial account of the resident experience, have not gained traction in England, |
| systems are | superimpose a layer of administration and do not link with routinely collected care data. |
| established. This | Dr George Heckman is an InterRAI Fellow at Waterloo University with experience of the |
| duplicates prior | Canada Health Infoway (CHI), a single point of contact for health information standards. He has |
| work | agreed to be an international expert to maximise opportunities for shared learning. |
| Resident consent | I his is addressed throughout the study and the findings will demonstrate about what needs to |
| and data | be in place to develop a resident MDS that merges multiple data sources. See also statement |
| governance | about information governance and development of protocols for data sharing. |
| Care nome and | By using established approaches from design and implementation science potential obstacles |
| NHS STATT WIII NOT | to implementation are identified early. Ongoing engagement with stakeholders will allow us to |
| co-operate. | reduce and streamline data entry demands on staff. The Involvement of the Professional |
| | Record Standards Body, the experience of The Health Foundation with INHS England plus the |
| Doto protoction | Pric involvement will help to intuite proof the MDS. |
| Data protection | Standard operating procedures will ensure data is protected. All data processing will be on the |
| and resident | meanin Foundation's purpose-dulit secure data environment that is accredited to hold and |
| privacy | process patient information. |