Patient work from a context and time use perspective: a mixed-methods study protocol

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ABSTRACT
Introduction Self-management is widely promoted but less attention is focused on the work required from patients. To date, many individuals struggle to practise self-management. ‘Patient work’, a concept that examines the ‘work’ involved in self-management, is an approach to understanding the tasks, effort, time and context from patient perspective. The purpose of our study is to use a novel approach combining non-obstructive observations via digital devices with in-depth qualitative data about health behaviours and motivations, to capture the full range of patient work experienced by people with type 2 diabetes and chronic comorbidities. It aims to yield comprehensive insights about ‘what works’ in self-management, potentially extending to populations with other chronic health conditions.

Methods and analysis This mixed-methods observational study involves a (1) prestudy interview and questionnaires, (2) a 24-hour period during which participants wear a camera and complete a time-use diary, and a (3) poststudy interview and study feedback. Adult participants living with type 2 diabetes and chronic comorbidities will be recruited using purposive sampling to obtain a balanced gender ratio and of participants using insulin and those using only oral medication. Interviews will be analysed using thematic analysis. Data captured by digital devices, diaries and questionnaires will be used to analyse the duration, time, context and patterns of health-related behaviours.

Ethics and dissemination The study was approved by the Macquarie University Human Research Ethics Committee for Medical Sciences (reference number 201700718). Participants will carry a wallet-sized card that explains the purpose of the study to third parties, and can remove the camera at any stage. Before the poststudy interview begins, participants will view the camera images in private and can delete any images. Should any images be used in future publications or presentations, identifying features such as human faces and names will be obscured.

INTRODUCTION
Self-management is widely promoted but less attention is focused on the work required from patients. It is a constant commitment of learning new skills, breaking old habits and managing different emotions over the length of the illness. Patients and their families often need to adjust their daily routines to accommodate for their health condition, such as making time to administer medications, visit medical professionals or prepare special meals. Yet, many individuals struggle to practise self-management effectively, especially for those with long-term chronic conditions.

Past studies reported on the barriers and challenges to self-management, focusing on patients with multiple chronic conditions, the difficulty involved in self-management, time invested, competing priorities, support sources required and how self-management differs between the patients’ and healthcare professionals’ perspectives. Yet, few studies reported on the actual ‘work’, or the tasks involved, from the patient, as well as whether self-management fits within the patient’s context.

When self-management is poorly integrated into people’s lives, the work required has the potential to generate additional stress,
contributing to non-compliance and negatively impacting on patient health outcomes. Unfortunately, many health interventions are currently implemented without considering the work involved from patient perspective or the context of a patient’s life. Treatment and self-management recommendations, while given with goodwill, often fail to acknowledge the specifics of a patient’s daily routine, physical environment, social obligations, as well as internal beliefs and attitudes. In parallel, digital health interventions hold the promise to improve self-management. However, numerous studies evaluating digital self-management interventions showed a high participant dropout rate, with some studies having retention rates of 1% by the end of the study period. Specifically, participants reported the devices’ lack of ‘fit’ into their daily lives as reason for discontinuation.

Understanding whether there is a dissonance between patients’ existing routines and the work involved in self-management is important to overcome barriers and challenges. The concept of ‘patient work’, derived from health ergonomics, describes the tasks (physical and cognitive) conducted by patients to manage their health, as well as the holistic sum of contexts (physical, social, mental and organisational) that influence the work conducted. It is an approach to understanding the actions people take (or not take) in self-management. While self-management focuses on the strategies people employ, patient work breaks down these strategies into day-to-day tasks and examines how the effort and time involved, as well as the contextual and ergonomic factors, affect the way self-management is practised and why some tasks are carried out while others are neglected.

The purpose of this study is to examine the patient work involved in managing with multiple chronic conditions, focusing on the daily lives of people with type 2 diabetes and chronic comorbidities.

**Patient work research**

Previous patient work research had focused on patients living with cardiovascular diseases such as heart failure and stroke. However, no studies have focused specifically on people living with more than one chronic condition.

In addition, previous patient work approaches relied heavily on self-report methods (interviews, direct observation (researchers being present continuously in participants’ homes) or theoretical modelling (analysing macroergonomic elements). These data collection methods are limited by the level of reliability in self-report data (eg, social desirability effect) and the degree of discomfort participants experience during direct observation.

The need for direct observation by researchers also meant the duration of visits had to be limited, ranging from 30 min to 1 hour at a time, making these short periods unlikely to capture the ‘natural’ routines of participants as there is high likelihood they could be affected by the observer effect. To our knowledge, no studies have used digital devices that automatically and non-obstructively collect data on patient work over a longer time frame.

**Type 2 diabetes and other comorbidities**

The nature of type 2 diabetes, which impacts the entire metabolic process, means that patients need to carry out a variety of different activities. These may include diet modification, weight management, exercise, complication prevention, blood glucose tests and insulin injections.

Metabolically related comorbidities such as hyperlipidaemia, cardiovascular diseases and renal disorders are frequently present in people living with type 2 diabetes, warranting the inclusion of comorbidities to reflect the true extent of patient work. Comorbidity is also frequently a criterion for exclusion in research studies, suggesting this study could give rare insights into the lives of people living with type 2 diabetes, as well as a greater understanding of self-management in patients with chronic comorbid conditions.

**AIMS**

This study will investigate the patient work conducted by people living with type 2 diabetes and at least one chronic comorbidity. It will address three gaps identified in the literature: the few studies reporting the ‘work’ involved in self-management from patient perspective (patient work), the methodological gap in using digital devices to collect data non-obstructively on self-management activities and contexts over a longer time frame (24 hours period) and focusing on people with complex challenges to self-management (those with multiple chronic conditions).

**METHODS**

This is a mixed-methods observational study that involves three stages: (1) prestudy interview and questionnaires, (2) a 24-hour period during which participants wear a camera and complete a time-use diary, and a (3) post-study session with interview and study feedback. Our study adapted the protocols used and validated by Kelly et al. which were employed in research projects conducted by the Centre for Time Use Research and the British Heart Foundation Centre on Population Approaches for Non-Communicable Disease Prevention, both at the University of Oxford.

**Sample and recruitment**

Eligible participants are: (1) aged 18 years and older; (2) consenting adults who can speak, write and read in English; (3) have been diagnosed with type 2 diabetes and at least one chronic comorbidity; (4) willing to complete a time-use diary for 1 day; and (5) agree to wear the body-worn camera for 1 day. Exclusion criteria include those who are currently experiencing: (1) pregnancy; (2) critical illness; (3) end-stage disease; (4)
dementia; (5) severe mental disorders that prevent one from giving informed consent; and (6) dependent on others for basic activities of daily living such as dressing and eating. Participants will receive $A60 of shopping vouchers as remuneration.

To capture patients at various stages of diabetes, 40 participants will be recruited, with balanced gender ratio, and balanced numbers of participants using only oral medication and those using insulin. The number of participants chosen was informed by previous qualitative studies in patient work. This sample size offers the power to achieve rich and nuanced data, while acknowledging the different time intervals between data points for each data capture tool. Previous patient work studies analysed 60–80 hours of interviews or direct observations per study. This study will capture approximately 2 hours of interview data and approximately 5000 images per participant. Therefore, a sample size of 40 participants is comparable to the depth of data captured by previous studies.

Recruitment will be carried out via Macquarie University general practice and general practitioners (GP) and endocrinologists in the surrounding suburbs. These medical practitioners manage a large and diverse patient base across northern Sydney and cover a wide range of culturally and economically diverse population groups.

Promotional flyers will be distributed to the participating medical practices and includes general study information and a link to the study website, where potential participants can check the eligibility criteria (www.tinyurl.com/patientswork). The study will also be promoted on the Macquarie University Facebook page and via an emailed newsletter associated with the University.

Healthcare practitioners will not be notified if their patients agreed to participate in the study to reduce the possibility of coercion.

Databases of medical practices involved in recruitment will be searched to identify eligible patients for the study. Only personnel employed by the medical practices will conduct these searches. Participating healthcare practitioners will also send invitation letters to eligible patients, with a prepaid reply slip included. Patients interested in participating will be asked to return the reply slip with their contact details, so a member of the research team can contact them.

Telephone calls will also be made to eligible patients to inform them about the study. If during the initial phone call the potential participant agrees to consider participating, the researcher will send a detailed email to the participant with a digital version of the information pamphlet as well as a link to the study website. The potential participant then either agrees to participate or declines. If the potential participant agrees to participate, the researcher will arrange a suitable time for interviews to occur. All participants will provide written informed consent before data collection commences.

Initial contact for all participants will be a telephone call conducted by the research team, to check whether the participant fits the eligibility criteria, assure confidentiality and anonymity during data collection and address any questions. This approach will minimise participant contact with the research team and clinicians. While self-nominated participants may be subject to participation bias, high completion rates and compliance are likely.

Figure 1 Flow diagram illustrating the tasks involved at each stage of the study.
The following instruments and equipment will be used for data collection:

- Time-use diary. The self-report time-use diary is a validated tool\(^39\) and the version used in this study is the Harmonised European Time-Use Diary\(^42\) (UK version). For details of the time-use diary, please see online supplementary appendix 2. This method of using free-text time-use diaries to self-report activities has high statistical validity and good acceptance rates with volunteer participants (ranging from 56% to 100%).\(^39\)\(^46\)
- Photos. With permission from the participant, photos of their medications, medical devices and home surroundings will be taken to inform about the participant’s physical environment. Should the participants reside in an environment that can affect their healthcare (eg, a house with stairs and a risk for falls), these issues will be taken into consideration. Participants will be alerted should an immediate and significant risk be identified.
- Body-worn camera. The body-worn camera (Edesix VB-300, Edinburgh) will automatically record silent continuous video footage. It can be attached to clothing and/or worn on a lanyard. Previous studies with body-worn cameras showed high degrees of acceptability, with acceptance rates ranging from 70% to 100%.\(^39\)\(^46\)\(^47\) especially when participants are reassured they can remove the camera if they feel the recording may intrude on their privacy.\(^39\)\(^46\)
- Demographic questionnaire. The demographic questionnaire (online supplementary appendix 3) will be administered during the prestudy session and takes about 10 min to complete.
- The Multimorbidity Illness Perceptions Scale. The Multimorbidity Illness Perceptions Scale\(^41\) (online supplementary appendix 4) is validated statistically and was developed to capture the perception of patients with comorbidities about their health.\(^41\) Illness perception is known to affect and predict health-related behaviour, to the extent that patients who perceive their diseases as more controllable are more likely to engage in adaptive strategies,\(^48\) thus directly affecting the patient work they engage in. The Scale will be administered during the prestudy session and takes approximately 5 min to complete.
- Participant experience questionnaire. A participant experience questionnaire (online supplementary appendix 5) will be administered during the poststudy session. This is estimated to take approximately 10 min.
- Researcher notes. A summary of each participant will be written up within a week after the poststudy session, summarising the researcher’s observations of the participant, home context and any feedback from the participant at different stages of the study.

**Patient and public involvement**

The concept study was conceived during a 1-day Consumer Symposium conducted at Macquarie University, where 10 health consumers gave feedback and critique regarding health research.

Health Consumers New South Wales, a patient advocacy group, was then consulted regarding consumer engagement and advice was given to the researchers regarding consumer involvement during study conception, recruitment, analysis to dissemination.

A pilot trial of the protocols with the equipment was conducted with two patients. Summaries of the findings were presented to the participants.

During study conception, the researchers also engaged with Healthdirect, a national organisation providing digital health information for health consumers, and the Northern Sydney Primary Health network and local GP network, which has a consumer council.

Results will be disseminated to these bodies, which will have broader implications to policies directed to consumers. Publications arising from the study will also be forwarded to each participant via emails.

**Data collection instruments and equipment**

The study will be carried out in three stages as described in figure 1 and table 1. Using multiple types of data, the study generates a holistic report of patient work conducted by people living with type 2 diabetes and chronic comorbidities in terms of observed physical work, organisational efforts, burden of disease and attitudes and beliefs underlying health behaviours.

The 24-hour data collection is more likely to capture a more representative day compared with previous patient work studies, which only captures patient activities for a few hours at most. While it will be hard to find a day that is truly representative of one’s life, the researchers ask the participants to nominate a day for the study that the participants themselves consider to be ‘normal’.

**Stage 1: prestudy session**

Prior to the prestudy session, participants will be provided with the Participant Information Booklet (online supplementary appendix 1) and will sign a written informed consent form (online supplementary appendix 6) before data are collected. Participants have the right to decline at this stage and to retract consent at any time throughout the study.

Ideally, the prestudy interview will occur in the participant’s home for researchers to note ergonomic factors. To ensure researcher safety, two researchers will make the home visits. Participants may come to Macquarie University for the interview if they prefer.

Following consent and before the prestudy interview, participants will also be asked to complete the demographic questionnaire (online supplementary appendix 5) and the Multimorbidity Illness Perceptions Scale\(^41\) (online supplementary appendix 4).

**Prestudy interview**

After the completion of the questionnaires, a semistructured interview will be conducted to gain information about the participant’s life history, medical history and
existing factors that impact patient work. Participants will be invited to talk about their health and lifestyle, with areas of focus separated into life history, social life and mental health, physical health and medical history, physical environment, and organisational effort and attitude (online supplementary appendix 7). With the participant’s permission, the interview will be audio recorded to assist with data coding and thematic analysis (eg, daily activities, rationales behind behaviours and factors that influence why participants behave in certain manners).

Lastly, participants will be provided with a body-worn camera and the Harmonised European Time Use Surveys UK Time-Use Diary (online supplementary appendix 2), together with relevant instructions. Photographs of the participant’s medications, medical devices and home environment will also be taken at this stage.

**Stage 2: participant self-recording data collection**

During the 24-hour study period, participants will be asked to wear the camera around their neck on a lanyard during waking hours (12–16 hours). The time-use diary will be completed by the participant over the course of the same day for 24 hours. Participants will be given instructions (figure 2) to recharge the camera overnight if the battery becomes low.

**Stage 3: poststudy session**

A poststudy session will occur immediately after the 24-hour study period has concluded. The camera footage will be downloaded and converted into images using software developed in-house, with one screenshot taken for every 10 seconds of footage. The participants will be able to view all the images alone before the research team views them, allowing the participant to delete any or all unwanted images.

**Poststudy ‘reconstruction’ interview**

The participant will be interviewed for approximately 30–40 minutes after data collection (online supplementary appendix 8). Participants will be asked to describe the study day, using the screenshots and time-use diary as prompts, with a focus on health-related tasks. The interview will be audio recorded with the participant’s permission to allow for thematic analysis of transcripts.

During the second interview, the researchers will also ask the participant whether the study day was ‘normal’ for them, and whether anything special occurred.

After the interview, the participants will be asked to complete the participant experience questionnaire (online supplementary appendix 5) and provide feedback on the methodology.

**DATA ANALYSIS**

Both qualitative and quantitative data will be gathered and analysed. The varied data sources are complementary and focus on different aspects of patient work. For example, self-reported data regarding daily tasks via the time-use diary can be enriched through an examination of the screenshots from the camera.

**Quantitative data analysis**

Video recordings from the body-worn camera will be automatically converted into series of 10-second screenshots, which are then processed for analysis of daily activities (eg, eating, sleeping). These screenshots will be tagged using the TensorFlow and/or Clarifai (New York, USA) Application Programming Interface, which automatically generates tags that identify objects and describe content within the image (eg, a woman and a car). A subset of these tags will be manually verified for accuracy.

Time-use diary data will be analysed in conjunction with camera screenshots. Participants’ activities reported in the time-use diary and observed via screenshots will be coded using an adapted version of the Harmonised European Time Use Surveys activity frame, and assessed for the duration, frequency, timing and patterns.

Questionnaire data (participant demographics, study feedback and the Multimorbidity Illness Perceptions Scale) will be reported using descriptive and inferential statistics. All statistical analyses will be conducted using SPSS v.25 (IBM).

**Qualitative data analysis**

Both prestudy and poststudy interviews will be transcribed verbatim. Analysis of the interview transcripts will
be conducted using the NVivo thematic analysis software package (NVivo, Melbourne, Australia).\textsuperscript{45} Two members of the research team will independently code the transcripts in a systematic manner. Any coding discrepancies between the two will be resolved through reaching a consensus.

Photographs of the participant’s medications, medical devices and home environment and camera screenshots will provide context to the interview. The photographs will also be used to give contextual feedback regarding the participant’s daily activities, as revealed in the video recordings and time-use diaries. Notes taken by researchers after the poststudy interview for each participant will be used to help with photo and qualitative data analysis.

ETHICS AND INFORMATION DISSEMINATION

Ethical considerations arising from this study are addressed below.

Consent and freedom to withdraw

Participants’ questions or concerns will be addressed individually and with all due diligence. Participants will be free to withdraw from the study at any point and can remove the camera at any time during the study day without giving a reason. The decision to join the study will have no consequence on their relationship with either Macquarie University or their health professionals.

Third-party and bystander consent

The research team will provide participants with a wallet-sized card that includes the statement below with contact information of the research team, which can be shown to others (to reduce the psychological burden of the participant) should anyone approach the participants to ask about the camera.

I am volunteering for a research project. The device is a wearable camera and the images will be used to record my daily activities. I am happy to remove it if you would like me to.

Should anyone request video recordings to be deleted, we ask participants to take note of the time that requests were made, and to assure the enquirer that the related recordings will be removed at the poststudy session. Participants are also asked to inform their household members, friends and acquaintances about the study, and to seek permission from these parties prior to recording.

Personal privacy

Participants will be asked to keep the body-worn camera off during situations when anyone (including themselves) is dressing, undressing, using the bathroom, showering or during any situations where privacy is of concern. Participants will be advised that they should always prioritise their own personal safety and comfort above the study’s requirements.

Furthermore, participants will be able to view and delete any (or all) of the extracted images before researchers view the images or videos at the start of the poststudy session. The videos will be deleted immediately after screenshots are extracted. Only members of the research team will have access to the screenshots. None of the video or screenshots will be released to the public, academic peers or participants. Should participants provide written informed consent for the research team to use their images in publications or presentations, the images will be blurred such that no one is identifiable.

To ensure the participant’s privacy, names and other identifying information will be removed in the photos of medications.

Participant feedback

Participants will be asked about their experiences in the study during the poststudy interview. A questionnaire to assess the participant’s experience (online supplementary appendix 5) will also be administered. Participants will be provided with contact details of the research team (email and phone number) for contact during the study.

Psychological support

In the unlikely case that participants become psychologically distressed during either of the interview sessions, a staff member with counselling experience will be present in both interviews to deal with any anxiety the participant may experience. Should participants experience high levels of distress during the study day, they are advised that they should immediately attend their local hospital emergency department and GP, and/or call Lifeline or emergency services.

Participant identification

Participant identities will not be masked during data collection. Participants will be individually identifiable due to the small sample size. However, their identities will only be known to the research team, and only members of the research team will have access to identifiable data. Participants will be individually identifiable due to the small sample size. However, their identities will only be known to the research team, and only members of the research team will have access to identifiable data. Data will be separated from contact details and stored in separate locations. No identifying information about the participant will be published or presented.

Information dissemination

Great caution will be taken in the handling and storage of the data by the research team so that risks to privacy are minimised, in accordance with the Australian Code for the Responsible Conduct of Research. Only members of the research team will have access to the data. The videos will be downloaded into a secure browser in a password-protected computer or Macquarie University servers. All physical copies of surveys and questionnaires will be locked in secure cabinets at Macquarie University. The data will be held for a minimum of 5 years. Anonymity will be upheld throughout reporting, and participants will only be referred to by their participant number.
DISCUSSION

At the time of writing, the study had engaged 40 potential participants and 23 participants completed the study, more than half of the number of participants the study aims for.

The reasons some potential participants did not take part in the study included: declined consent due to personal reasons (which the participant did not disclose), unable to communicate in English to a sufficient standard (as judged by the researcher during the initial phone call), too busy with work, too busy due to full-time care of family members, going on a holiday, was sick on the scheduled visit and withdrew from the study, and was not home when the researchers visited and withdrew from the study. No participant who completed the study had withdrawn their consent to date.

Completion rates for the time-use diary is very high, with 21 out of 23 participants successfully filling the diary throughout the study day. The two participants who did not fill in the diary did so during the poststudy interview, with the assistance of recorded footage.

All participants complied with continuous camera recording. There was a variation of camera recording time between participants, due to some participants turning the camera off for some periods of time out of privacy concerns.

One challenge the study had faced was communicating camera operations effectively to the participants. Some participants had difficulty turning off the camera and changing the camera battery due to reasons such as forgetfulness and confusion regarding the orientation of the camera buttons. To address the issue, the research team had devoted more time during the prestudy interview to physically demonstrate how to use the camera, which had successfully improved participant understanding of camera operations.

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Contributors  Study conceptualisation: AYSL, EC. Study design: AYSL, KY, TH, JG. Data collection preparation and analysis plan: KY, TH, AYSL, LL, FR. Recruitment and liaison: KH, SV. First draft: KY, AYSL. Draft revision: All authors provided feedback and approved the manuscript.

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Competing interests  Some participants may be recruited via the Macquarie Health General Practitioner clinic and Macquarie Hospital, where SV is practising as a general practitioner and KH is practising as an endocrinologist, respectively. Therefore, the two team members may be responsible for the treatment of some participants. However, SV and KH will not know whether a patient is a participant unless the participant informs them.

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