Video Reflexive Ethnography Methodology to Explore the Use of Variable Rate Intravenous Insulin Infusions

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Research Article

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Abstract

Background: The use of variable rate intravenous insulin infusion (VRIII) is a complex process that has consistently been implicated in reports of error and consequent harm. Investment in patient safety has focused mainly on learning from errors, though this has yet to be proved to reduce error rates. The Resilient Health Care approach advocates learning from everyday practices. Video reflexive ethnography (VRE) is an innovative methodology used to capture, reflect on and thereby improve these. This study set out to explore the use of VRIII with VRE, a secondary aim being to describe VRE’s feasibility and acceptability.

Methods: This study was conducted in a Vascular Surgery Unit. Quantitative data (e.g. blood glucose measurements) were collected from electronic patient records. Qualitative data were collected using VRE methodology. The latter involved videoing healthcare practitioners caring for patients treated with VRIII and discussing the resulting video-clips with participants in reflexive meetings. Transcripts of these were subjected to thematic analysis and the quantitative data used to judge the outcomes of the video-observed tasks. Feasibility in relation to recruitment and data collection, as well as the acceptability of using VRE, were assessed based on participant responses during the study.

Results: The use of VRE in conjunction with quantitative data revealed that context-dependent adaptations (seeking verbal orders to treat hypoglycaemia) and standardised practices (using VRIII guidelines) were strategies used in everyday work. Reflexive meetings highlighted the challenges faced while using VRIII (lack of knowledge of the appropriate medications to be prescribed with VRIII) and encouraged participants to suggest solutions (face-to-face, VRII-focused training). The use of VRE was judged acceptable, based on the researcher’s interpretation of participants’ willingness to participate, and feasible, since all patients and 83% of the healthcare practitioners who were approached to participate agreed to do so.

Conclusions: VRE deepened understanding of VRIII by shedding light on its essential tasks and the challenges and adaptations entailed by its use. The use of VRE to explore VRIII in a single unit was judged feasible and acceptable. However, future research might focus on collecting data across various units and hospitals to develop a full picture of the use of VRIIIs.

Background

Diabetes mellitus is a common, serious, and chronic disorder that affects more than 422 million adults globally [1] and is characterised by hyperglycaemia where fasting blood glucose (BG) levels are greater than 7.0 mmol/L (126 mg/dl) [2]. In the UK, the National Diabetes Inpatient Audit (NaDIA) found that the prevalence of hospitalised adult patients with diabetes has steadily risen from 15% in 2011 to 18% in 2017 [3]. Hyperglycaemia is associated with an increased risk of complications and mortality, a longer hospital stay, and a higher admission rate to the intensive care unit [1, 4].

Variable rate intravenous insulin infusions (VRIII) are considered the treatment of choice to achieve optimal BG levels in patients who are unable to eat, who will miss more than one meal, who have severe illness, e.g. sepsis, and where there are special circumstances, e.g. acute coronary syndrome [2]. VRIII is extremely effective at reducing BG levels quickly for hospitalised patients because of the rapid onset of action of the intravenous route of administration, compared to other administration routes. This characteristic, however, carries the risk of causing patient harm because it is not possible to reverse its action if used inappropriately [5]. Therefore, patients treated with VRIIIs need special care in areas such as regular BG measurements, an adjustment of the VRIII rate according to BG readings, and awareness of the appropriate time to transition from using VRIIIs to subcutaneous insulin if required. The mismanagement of any of these areas is dangerous and can lead to complications such as hypoglycaemia (BG < 4 mmol/L) and ketoacidosis [3].

Patient safety is a vital concern in hospitals and numerous initiatives have been introduced to enhance patient safety when using VRIIIs. Such initiatives included producing practical guidelines for the use of VRIIIs [6], using prefilled, ready-to-administer, injectable medication to mitigate against patient harm due to preparation errors [7], and recommending independent verification for dose calculations, prescription, dispensing, and administration of certain high-alert medications (e.g. VRIII) in order to identify potential errors before medications are administered to patients [8]. The current initiatives are predominantly based on traditional safety approaches that focus on top-down solutions (e.g. standardised policies) introduced into healthcare systems and/or by external experts by learning from specific errors and producing solutions by eliminating errors and filling in any knowledge gaps (Safety-I) [9]. Healthcare systems are considered complex adaptive systems (CASs). By definition, CASs are systems that include individual agents – doctors, nurses, pharmacists, and patients – who act and interact with each other and with the surrounding environment (e.g. equipment and technology) in dynamic and unpredictable ways that change over time through learning, feedback, and adaptation [10]. Although great efforts have already been made to enhance patient safety while using VRIIIs, the dependence on traditional safety approaches has proved problematic, failing to demonstrate convincing reductions in risk, error or death [11].

A different way of thinking about safety has been introduced, switching the focus from approaches that concentrate on identifying and eliminating errors to more comprehensive approaches that advocate learning from how things go right as well as how they might go wrong (Safety-II) [12]. A relatively new area of theory called Resilient Health Care (RHC) adopts the Safety-II way of thinking, focusing on understanding how everyday work is actually done rather than how it is assumed to be done [13]. RHC’s proponents argue that performance variability is an
integral part of how people and systems deal with expected and unexpected situations, and that their capacity to successfully adapt is what makes a successful system work, enabling good outcomes in spite of problems and challenges [14]. It suggests a shift in focus from learning only from errors to a complementary perspective of learning from how things go right as well as how things go wrong [15]. One of the important constructs in RHC is the distinction between how everyday work is actually done (‘Work as Done’, WAD) and how work is assumed to be done (‘Work as Imagined’, WAI) [16].

In order to understand how healthcare practitioners work in real life situations (WAD), ledema et al. used the term ‘exnovation’ (‘innovation from within’) to focus our attention on how healthcare practitioners enact and understand the internal complexity of their clinical work. In contrast to innovation introduced from elsewhere, exnovation is the process of strengthening insight into in situ everyday work, and enabling practitioners to enact on that insight [17]. It is clear that the complexity of care is often taken-for-granted. For that reason, understanding WAD requires an in-depth exploration of everyday work and of the hidden, taken-for-granted clinical work performed by frontline practitioners in situ. Video reflexive ethnography (VRE) is a qualitative methodology involving collaboration between researcher and participants and aims at rendering the complexity of ordinary everyday tasks visible through the videoing of everyday work. VRE has three phases: (a) familiarisation with the work through informal observations; (b) video-recording of everyday work; and (c) reflexive meetings in which participants and researcher review edited video footage [17]. Video feedback with those involved in the everyday work enables activation of insights articulated by the practitioners, patients, and researchers collaborating to learn about the work, and to make sense of context (reflexivity). Research shows this process results in participants recommending and initiating designing realistic and realisable solutions to enhance safety and care [17].

The present study is part of a project that has a published protocol [18] with the overarching aim of exploring RHC in the use of VRIII. As far as we know, no previous study has used VRE methodology to explore WAD in the use of VRIII. Therefore, the aim of this study was to understand, by using VRE and quantitative data, how VRIII is used in a Vascular Surgery Unit. The secondary aim was to describe the feasibility of using VRE in relation to (1) the recruitment of participants, (2) the data collection procedure, and (3) the acceptability of VRE within the Vascular Surgery Unit.

### Methods

#### Study design

The current study was informed by the RHC approach [19, 20] and the four theoretical principles of VRE: exnovation, reflexivity, collaboration, and care [17]. RHC and VRE principles shaped our ontological stance, which suggested a need for an in-depth understanding of the taken-for-granted clinical tasks associated with the use of VRIIs. This stance led us to think about the types of knowledge that would be of most value in exploring WAD. We reasoned that it would be important to focus on how clinical work was delivered, and on how adaptations were made in expected and unexpected situations. Therefore, a qualitative dominant (QUAL+ quan) mixed-methods approach was used. Our approach depended mainly on an exploratory sequential design [21] in which qualitative data were first collected and then used to inform what quantitative data were needed to make judgments on the outcomes of the observed tasks [22]. Findings were integrated by merging the qualitative VRE data and the quantitative data (e.g. BG measurements) to complement each other and better understand WAD while using VRIIs. This study follows the Good Reporting of A Mixed Methods Study (GRAMMS) guidelines [23] and was conducted between September 2019 and July 2020.

#### Study setting

The study was conducted in a Vascular Surgery Unit in a tertiary hospital in England that provided emergency and elective treatment for patients with vascular diseases, e.g. peripheral vascular disease and diabetic foot disease. The Unit had 10 patient bays (24 beds), mostly occupied for the duration of the observations. There was a treatment room where the nurses prepared medications and assembled equipment. The Unit used a syringe pump for insulin infusion and a volumetric pump for IV glucose containing fluids. Insulin and glucose containing infusions were administered via a single cannula via a Y-connector with dual anti-reflux valves. The Unit used an electronic prescribing, monitoring, and administration (ePMA) system within the electronic patient record system (EPR). Two nurses conducted independent verification of prescription, patient, infusion pump programming, capillary blood glucose (CBG), VRIII initial rate and for each rate change. There was generally one nurse per six patients and foundation year one/two (FY1/2) doctors were regularly present.

#### Recruitment

Participant recruitment was conducted over five months. Based on the published study protocol [18], the aim was to include two patients and all healthcare practitioners responsible for the use of VRIII in their care. Box 1 shows the eligibility criteria and the recruitment process for both patients and healthcare practitioners.
### Eligibility criteria and recruitment process.

| **Eligibility criteria** |

**Inclusion criteria**
- Healthcare practitioners who are:
  1. Willing to be observed by video recording.
  2. Working in the Vascular Surgery Unit.
  3. Managing/dealing with patients on VRIII.
- Patients who are:
  1. Aged ≥ 18 years old.
  2. Receiving VRIII for at least 24 hours to treat elevated BG.
  3. Under the care of a healthcare practitioners who have consented to participate in this study.
  4. Able to provide informed consent.

**Exclusion criteria**
- Healthcare practitioners who are:
  1. Not willing to be observed by video recording.
  2. Not working in the Vascular Surgery Unit.
  3. Not involved in the use of VRIII.
- Patients who are:
  1. Not willing to be observed by video recording.
  2. Not prescribed VRIII.
  3. On IV insulin and glucose infusion for hyperkalaemia (potassium levels >5.5mmol/L).
  4. Unable to provide informed consent.
  5. Non-English speakers.

**Recruitment**

**Healthcare practitioners**
To recruit potential healthcare practitioners, an invitation letter and participant information sheet outlining the purpose of the study, the methodology, and the design was first sent by the Unit clinical and managerial lead to all potential healthcare practitioners. Then the researcher (MI) met the healthcare practitioners working in the Unit in two ward meetings and explained the study aims and process. Informed consent was then taken from interested healthcare practitioners. A poster with details about the study was also placed in the staff room until the completion of data collection.

**Patients**
The study site collaborator with a pharmacist team identified potential patient participants. The researcher confirmed with the senior nurse the appropriateness of the patient before recruiting them. This is in addition to whether the patient had the capacity to consent. Once agreed, the researcher provided an invitation letter and participant information sheet to the patient and explained the purpose and objectives of the study and that as part of filming the work of healthcare practitioners, some parts of their body might appear in the video recordings (arm, leg, etc.). The researcher asked the patient if they had any questions about the study before taking written informed consent.

### Data collection

**Qualitative VRE**
The VRE involved three stages:

**Stage 1: Familiarisation and informal observation**
There were three aims to this stage: 1) To familiarise the researcher (MI) with the environment and build a level of trust with healthcare practitioners, thus facilitating the video observations. 2) To informally observe the staff working in the Unit, in order to understand how VRIII was being used and in turn determine the tasks that needed to be filmed. 3) To familiarise MI with the ePMA system in order to understand how prescribing, monitoring, documenting and other tasks were being reported when using VRIIIIs. This stage involved taking field notes while conducting 15 hours of general observations in three to five-hour sessions. It also involved shadowing three healthcare practitioners while they were using VRIII for short periods of 30–60 minutes each.

**Stage 2: Video observation**
The focus of this stage was to capture how VRIII was used in situ. Two digital cameras, one static next to the patient's bed and the other handheld by the researcher, were used. The static camera focused on the tasks accomplished around the insulin pump. For the tasks that occurred
elsewhere, e.g. handover, the assembly of the VR III and other equipment, preparation and documentation, the handheld camera was used. The video footage was transferred to the project’s shared drive as soon as possible and then deleted from the camera.

Stage 3: Reflexive meeting discussion

The aim of this stage was to engage the healthcare practitioners collaboratively in analysing their work.

1- Choosing the clips of interest for the reflexive meetings

The research team (MI, RL, CC, FG, RI) and the study site lead nurse (PW) chose the clips for use in the discussion based on the main criteria presented in Box 2.

Box 2. Criteria used to choose the video clips for use in the reflexive meetings.

1. The video clips were chosen on the basis that they could be used to address the research question with the healthcare practitioners, enabling them to describe and scrutinise how VR III is used in everyday clinical work. The guiding principles of VRE are:

- Does the footage reveal a range of tasks or a range of different people doing similar things in previously un(der) appreciated ways (reflexivity)?
- Can the clips show different perspectives on a task and create the possibility for more appropriate understandings, perspectives or solutions (exnovation)?
- Do the chosen clips and the feedback discussion maintain the psychological safety of the participants (care)?
- Do the chosen clips could help engaging the healthcare practitioners in collectively identifying safety interventions in the in situ, taken-for-granted everyday work (collaboration)?

2. The time available for the reflexive session

- The video clips were chosen in such a way that they could be discussed in 30–45 minutes.

2- Conducting the reflexive meetings

MI conducted reflexive meetings through the Microsoft Teams platform and verbally confirmed continuing consent with healthcare practitioners before the start of the meetings. The strategy for facilitating and running the reflexive meetings was guided by the four principles of VRE [17]. MI was an ‘outsider’ facilitator who used specific clips to facilitate the reflexive meetings in order to achieve specific aims [17]. MI described to the participants the footage they would be watching and facilitated the discussion by asking pre-prepared questions throughout the session (see reflexive meeting discussion guide in see Additional file 1) in order to explore what worked well, what might be learned from everyday clinical work, what problems and challenges were faced, and what adaptations and adjustments were made while using VR III. Each of the reflexive meetings was audio-recorded and transcribed verbatim for further analysis.

Quantitative approach

The EPRs of the patient participants were accessed after completion of videoing to identify CBG measurements, the VR III regimen selected, the monitoring frequency of CBG and blood ketones, the duration of persistent hyperglycaemia and the number of hypoglycaemic episodes. The quantitative data covered the whole 24 hours during which video observations were captured.

Data analysis

Qualitative VRE

All the video recordings, including verbal utterances and actions, were interpreted and transcribed by MI. Quantitative data from patient medical records were added to the transcripts to provide a comprehensive written account of the tasks captured in the video recordings and to facilitate the choice of clips of interest to be used in the reflexive meeting discussions. All the verbatim transcripts of the reflexive meeting discussions were analysed using inductive thematic analysis [24]. MI coded transcripts then constructed sub-themes and themes within and across these codes. In an iterative process, the research team (MI, RL, CC, FG and RI) then met to discuss the overarching themes in order to identify significant broader patterns of meaning that represented how the healthcare practitioners used VR III.

Quantitative approach

The quantitative data were collected and compiled by MI and discussed with the study site collaborator (CC) in order to ensure the data collection and reporting processes were robust. The time period during which CBG was at target range was identified. The number of CBG readings of <4.0 mmol/L (hypoglycaemia), and CBG readings of >12.0 mmol/L (hyperglycaemia), were recorded. The duration of persistent hyperglycaemia, and the incidence of hypoglycaemia episodes (any CBG reading of <4.0 mmol/L in a four-hour period), were identified. The number of times the patient required an IV glucose infusion to treat hypoglycaemia was also recorded.
Qualitative and quantitative data were interpreted and reported using a narrative approach in which the researcher presented the two sets of findings in various sections, weaving both together and writing up the results on a theme-by-theme basis [21].

Results

Qualitative VRE data

More than 100 tasks (e.g. administer VRIII and IV fluids) were observed while using VRIII to treat elevated CBG (see Box 3 for examples).

Box 3. Summary of the main tasks observed to treat elevated CBG using VRIIIs.

The treatment of elevated CBG using VRIIIs started with confirming the potential need for VRIII in the ward round/board meeting. Each morning between 8am and 10am there was a ward round during which consultants, registrars, FY1/2 doctors, and a senior nurse checked each patient’s status, examined each patient and reviewed their progress, laboratory results, and medications. After the ward round finished at around 10.30am there was a board meeting attended by various healthcare practitioners including nurses, consultants, registrars, FY1 doctors, medical students, receptionists, pharmacists, and occupational therapists. The staff discussed patients’ cases and approved their treatment plans. Nurses came to check the confirmed plans for their patients, with a senior nurse documenting the plans on a yellow paper.

Ensuring the right medications were prescribed, based on CBG readings, was conducted by doctors who were required to prescribe VRIII, IV fluids, and IV glucose, stop all diabetes medicines, except long-acting insulin analogues which were to be continued.

Before assembling the components of VRIII, nurses checked that the medications matched the prescription on the EPR. To assemble the components, several steps were performed including, but not limited to, wiping a blue preparation tray with alcohol wipes, assembling equipment (syringe, extendable administration line, IV administration set, drawing up needles, syringe for the flush, sodium chloride 0.9% ampoule for the flush, chlorhexidine alcohol wipes), and drawing up the solution from the sodium chloride 0.9% ampoule using the blunt fill needle.

Nurses then recorded on the ePMA the medications to be administered (VRIII and IV fluids). Independent verification of the VRIII/IV fluids was conducted before administering insulin/IV fluids by checking the label on the insulin/IV fluid syringe (name, dose, expiry date), confirming this by signing the details on the EPR.

Ensuring CBG and blood ketones were within the normal range was conducted by bedside monitoring of CBG and blood ketones every 1-7 hours. Nurses were required to regularly monitor cannula, patient complaints, and site of injection, based on the patient’s clinical status.

Nursing handovers were conducted three times a day for each patient, sometimes during lunch breaks.

It was clear that electronic documentation of CBG/ketone readings, VRIII rate, insulin/IV fluids administration, and VIP score was a crucial step by which healthcare practitioners made sure each task they accomplished using the EPR system was documented.

Quantitative approach

Table 1 shows the demographics and the main data concerning CBG monitoring obtained from the two patients’ medical records.

| Table 1: Demographics and CBG monitoring data from two patients’ medical records |
|---------------------------------|------------------|------------------|
| **Patient 1**                  | **Patient 2**    |
| Age (years)                    | 87               | 84               |
| Gender                         | Male             | Male             |
| Time intervals between each CBG reading (average; range) | 4.5 hours; 2−7 hours | 2 hours; 1−3 hours |
| Percentage of time during which CBG was at target range i.e. 4.0 to 12 mmol/L | Day 1: 4% of 24 hours | Day 2: 25% of 24 hours |
| Percentage of CBG readings of <4.0 mmol/L or >12 mmol/L | Day 1: 90% of CBG readings >12 mmol/L | 0% |
| Duration of persistent hyperglycaemia | Day 1: persistent hyperglycaemia over 24 hours | Note: there was no DKA associated with the persistent hyperglycaemia |
| Incidence of hypoglycaemia episodes | None reported | 3 episodes |
| Number of times IV glucose infusion was administered for managing hypoglycaemia | 0 | 2 |

Two male patients were recruited and took part in the study. Most of the time, the BG readings of both patients were out of the target range. One patient had persistent hyperglycaemia over 24 hours and the other experienced three episodes of hypoglycaemia for which he was treated with two 20% IV glucose infusions.
Qualitative and Quantitative

As a result of analysing the reflexive meeting transcripts and the quantitative data, three broad themes were identified. (1) Safety strategies: standardise, adapt, and learn to ensure delivery of patient care. (2) Lack of knowledge and insufficient organisational infrastructure as the main challenges in the use of VRIII. (3) Suggestions for enhancing the effectiveness of current safety strategies. (See Additional file 2 for the themes, sub-themes, codes, and quotes of the reflexive meetings’ analysis.) These themes are described in turn below.

1. Safety strategies: standardise, adapt, and learn to ensure delivery of patient care

During the reflexive meetings, healthcare practitioners reported that the use of standardised improvements provided by the hospital facilitated the delivery of patient care using VRIII. Specifying a time to start the VRIII in the VRIII guidelines, the availability of VRIII guidelines, the use of ready-to-administer insulin infusion syringes, and the availability of the treatment algorithm provided as a laminated printed sheet inside the 'hypo box', have between them improved the practice of using VRIII. Most participants said that the use of EPR produced more accurate and clearer prescriptions compared to handwritten ones. The importance of using checking and verification was emphasised as a strategy for ensuring patient safety while using VRIII, e.g. independent verification and countersigning before administering the VRIII and fluids, to ensure that nurses get the right information (right dose, expiry rate); checking the CBG before starting the VRIII; and checking the patient's blood ketones if the CBG level was >12 mmol/L.

I think... because insulin is such... can be such a... if you get it wrong it can cause a serious problem, so I still do think that it should be double-checked. Any infusion like that we would get double-checked anyway. (Nurse 1)

The use of VRE revealed that context-dependent adaptation was the most frequently described and observed strategy used to ensure the delivery of patient care. Most of the time, these context-dependent adaptations had positive outcomes in terms of patient care delivery, leading to the tasks being accomplished. For example, nurses seeking verbal orders for IV glucose when the doctors were busy, and nurses administering IV glucose then checking if it was prescribed to prevent delays in treating hypoglycaemia and consequent patient harm. There were, however, other outcomes that led to harm or inadequate care. One of the inadequate adaptations occurred when, as described by the NAs, if the nurse was busy and the patient had hypoglycaemia, they would give the patient a sugary drink. The NAs were not aware that the patient was on a VRIII and nil by mouth. By being unaware of the VRIII and the patient’s status, giving oral rather than IV glucose might delay the patient going into theatre.

If it’s not prescribed then I would just speak to the doctor, tell them it’s urgent and if... ask if I can either have a verbal order if they’re not able to prescribe it there and then because it could lead to an emergency if it’s not corrected. (Nurse 1)

2. Lack of knowledge and insufficient organisational infrastructure as the main challenges in the use of VRIII

The two sub-themes identified were ‘lack of familiarity leads to fear’ and ‘deficient organisational infrastructure risks effective work’.

The sub-theme ‘lack of familiarity leads to fear’ was expressed in the context of prescribing. The most prominent prescribing issues were related to a lack of clinical knowledge about the appropriate IV fluids to use; the need for prescribing long-acting insulin analogues alongside the VRIII; the need to discontinue other diabetes medicines; and the importance of prescribing IV glucose (for hypoglycaemia). Nurses reported that these problems can occur because of a lack of experience (especially seen with new doctors); inadequate training for VRIII indications, side effects, and monitoring; and poor training of senior doctors in how to use electronic systems to prescribe VRIII and IV fluids. This view was echoed by an SpR, who described the EPR as a relatively new system and acknowledged that the training to use it was mainly given to junior doctors. SpRs were not used to using it, which could lead to delay or errors in the electronic prescription.

Box 4 shows a vignette that describes the problem with prescribing IV fluids and how it was approached.
Box 4. Prescribing IV fluids.

A: The nurse checked the prescription for VR III and IV fluids on the EPR before gathering equipment in preparation for VR III and IV fluids infusion. The nurse could not find the IV fluid prescription on the EPR and decided to check with the SpR. There were three other nurses in the room at the same time, who were preparing medications for other patients.

B: The SpR (on the left) was trying to prescribe the required IV fluid, but was unsure of the types of fluids available. He struggled to prescribe IV fluids using the EPR. The nurse (on the right) assisted the SpR, showing him how to choose the IV fluid on the EPR. The nurse shared some critical information about the patient's status, including the nil by mouth status and the potassium level, details important in deciding which IV fluid was best suited to the patient's situation. The SpR disregarded the potassium level and chose an IV fluid that was inappropriate for the patient's condition based on the hospital VR III guidelines. The nurse confirmed the IV fluid rate with the SpR and went back to the treatment room to check the EPR.

C: The nurse (on the right) put on disposable gloves and assembled the equipment for using insulin infusion and IV fluids. The nurse prepared a blue tray (a tray in which prepared equipment is placed) and wiped it with alcohol wipes whilst waiting for the prescription to appear on the EPR. The nurse approached the senior nurse (on the left) for help as she could not find the IV fluid prescription on the EPR, although the nurse had twice spoken about it with the SpR. The senior nurse decided to talk to the SpR to resolve the issue of prescribing the IV fluid.

Other challenges, such as fear of hypoglycaemia and a lack of confidence treating it, were mainly related to lack of experience, and hesitant new staff. (See Additional file 3 for a video clip on confirming the treatment of hypoglycaemia with a senior nurse).

Challenges associated with deficient organisational infrastructure were exemplified by the requirement for frequent CBG monitoring with an insufficient workforce; insufficient knowledge gained from e-learning courses; and the near-patient wireless CBG meter results sometimes not being updated on the EPR.

Yeah we do struggle sometimes with the equipment because sometimes if after the procedure we put the meter in the [docking] station and it doesn't really update, I mean the system, it's [not] updated so sometimes a nurse will ask if the blood sugar testing was done so yes, I tell them that it's been done but it's not showing on the computer. (NA3)

3. Suggestions for enhancing the effectiveness of current safety strategies

The reflexive meetings allowed healthcare practitioners to suggest solutions that they regarded as being essential to enhancing quality and safety while using VR IIIIs. Their suggestions were categorised into staff-focused and system-focused enhancement strategies. Staff-focused strategies included improving staff knowledge about prescribing and monitoring VR IIII; predicting the need for a VR IIII even if one is not prescribed; using proactive measures to prevent further hypoglycaemia in patients with ongoing risk, e.g. educating healthcare practitioners about the most common symptoms of hypoglycaemia to anticipate the risk; being mindful of the patient's need for VR IIII; and individualising the treatment plan based on the patient's clinical status, rather than simply following protocols.

So I guess identifying symptoms of a patient who's having hypo or hyper will be more helpful in performing this task. What to expect so you know what's happening. (NA3)

System-focused strategies were mainly focused on improving the current training approaches and investing in resources to improve the prescribing of VR IIIIs. A common suggestion for improving training sessions was directed towards conducting more specific training on VR IIII prescribing, recognising the side effects of VR IIII, and how to deal with its side effects. When asked about training to enhance knowledge, the participants were unanimous in the view that face-to-face or in-house training would be much better than e-learning.

If we have physical training, like... physical is better than just e-learning with just reading, reading, and answering the questions. (NA1)

Feasibility of VRE use

The recruitment strategy

The recruitment strategy was feasible and suitable. Twenty healthcare practitioners (nurses, senior nurses, pharmacist, registrars, consultants, FY1/2 doctors and NAs) working in the Unit provided informed consent prior to identifying and recruiting patients. Of these, 12 healthcare practitioners were involved in the care of the two patients who was subsequently identified and recruited for the study. Ten of the 12 healthcare practitioners agreed to take part in the study; three nursing assistants (NAs), two nurses, three senior nurses, one specialist registrar (SpR) and one FY1 doctor. Two declined to participate as they were not trained to use VR IIIIs, the focus of this study.
The role of the study collaborator and the pharmacy team was vital in identifying potential patients for recruitment. However, the workload of the team may have affected the frequency of their checks for potential patients and also their ability to contact the researcher before a patient started VRIII.

Data collection stages

The familiarisation stage had met the three aims previously mentioned, as well as allowing the researcher to determine the optimum camera placement (including the researcher’s own position) for capturing both the VRIII and the healthcare practitioners’ work. During the video-observation stage, patient 1 was observed for two days (four hours on day 1 and three on day 2). The researcher stopped observing patient 1 on the second day, based on the healthcare practitioners’ request, as the patient had developed sepsis i.e. a life threatening clinical syndrome results from extreme body response to infection [25]. Patient 2 was observed for one day for a total of six hours. Although it was planned to video both patients’ cases over 24 hours (day and night) [18], the researcher was unable to stay in the hospital overnight and healthcare practitioners were too busy to help with the video recordings when the researcher was not available. As a result, the 13 hours of video recording were all captured during the day.

Previous VRE studies had conducted reflexive meetings attended by a group of participants who had been involved in the video clip [26-28]. In the case of the present study, the hospital site paused all research activity because of the coronavirus (COVID-19) pandemic and the research team responded by conducting virtual reflexive meetings with the healthcare practitioners. The 10 healthcare practitioners involved at the video-recording stage were approached via email and telephone. Three did not reply, two were not able to participate because of work and family issues, leaving five who agreed to take part. When it came to organising the reflexive meetings, the aim was to conduct a meeting with all healthcare practitioners who appeared in the video footage. However, it was not feasible to meet all healthcare practitioners, or even a group of two, at the same time. One-to-one reflexive meetings, lasting between 30 and 45 minutes, were conducted instead. Figure 1 summarises the recruitment and analysis stages of the study.

The acceptability of using VRE within the Vascular Surgery Unit

Acceptability of VRE use was evaluated based on the participants’ responses during the data collection period. Both patients were willing, indeed keen, to be video-recorded. Despite meetings and discussions about the study and the value of VRE, healthcare practitioners, especially the nurses, were concerned about patient confidentiality. This was understood to be because healthcare practitioners were not used to being video-recorded for any purpose. The research team engaged healthcare practitioners in designing appropriate ways to video both themselves and patients. These included where to locate the cameras, how to identify potential patient participants, and when to start and stop videoing. This allayed concerns from healthcare practitioners.

In the first video recording, the researcher noticed that participants were initially trying to modify their work in order to deliver the best care (Hawthorne effect) [29]. However, there was no observable sustained change in participants’ behaviour when they were being filmed. For example, while the nurse initially cleaned her hands as per the Aseptic Non-Touch Technique guidelines, after five minutes of recording she no longer used alcohol rub or soap and water but instead applied non-sterile gloves before checking the EPR.

Discussion

This study was the first to explore the use of VRIII in a Vascular Surgery Unit using VRE and quantitative data. Feasibility studies enable researchers to assess whether or not the process of developing and implementing methods or interventions can be relevant and sustainable [30, 31]. The results of this study demonstrated that VRE methodology was feasible in exploring the use of VRIII in a Vascular Surgery Unit and potentially finding ways to improve patient care.

The VRE methodology uncovered the actual and potential hidden complexity in everyday work that encompasses various types of tasks and clarified how it was handled in situ. Engendering healthcare practitioners’ reflexivity helped the researcher to explore how and why some tasks were accomplished in particular ways and to align more closely with the reality of everyday work, the result being a better understanding of WAD while using VRIII. Various methods had previously been used to explore WAD and resilience in healthcare, e.g. field observation, interview, and focus groups [32–34]. These methods can partially capture an approach to understanding WAD. However, no studies had been conducted to explore WAD in the use of VRIIIIs using a VRE methodology in which healthcare practitioners are involved in reviewing, analysing, and reflecting on their work.

Exploring the use of VRIIIIs using VRE and quantitative data

Although the study findings were obtained on the basis of video-recordings of two patients and 10 healthcare practitioners, the use of VRE along with quantitative data, provided data that enabled analysis of VRIII use, and in turn recommendations to improve its use. Healthcare practitioners and/or the system used a range of strategies to ensure the delivery of patient care. These include using guidelines for VRIII and context-dependent
adaptations such as delegating the CBG monitoring task to other colleagues when the nurse was busy. The use of VRE allowed healthcare practitioners to recognise their personal roles and to appreciate the importance of context-dependent adaptations as a strategy for dealing with unexpected situations in ways ensuring patient safety. This finding broadly supports the work of other studies demonstrating how adaptations are required in everyday work to provide safety improvements and resilience in the system [35–37].

Previous studies highlighted the impact of using VRE in various clinical settings by demonstrating achievements such as enabling participants to become explicit about their own practices and problems in healthcare-associated infections [38], developing meaningful solutions for problems in an intensive care unit [39], and enhancing team capacity to enact person-centred care to improve dementia care [40]. In this study, VRE helped healthcare practitioners to be more reflexive and explorative of challenges, acknowledging that the work may need changing and suggesting practical solutions tailored to their work.

The use of VRE highlighted some of the challenges experienced by healthcare practitioners when using VRIII. These related to a lack of clinical knowledge and experience when prescribing the appropriate IV fluids, failure to appreciate the necessity of continuing to administer long-acting insulin, or lack of awareness of the type of fluids available. Rickard et al. identified that 64% of the IV fluids prescribed with VRIII did not have the recommended potassium concentration [41]. This is consistent with what was observed in this study where there was a discrepancy between the fluid prescription and patients’ clinical status. Another challenge identified in this study was the use of the ePMA to prescribe VRIII and IV fluids. This finding is likely to be related to the fact that, prescribers rotate from other organisations, so the ePMA system might be new to them. This explanation matches the conclusion reached in other NHS Trusts, where the initial implementation of the ePMA system was more time consuming than the paper method, although as staff became more familiar with it the process of prescribing, monitoring, and administering became more efficient [42].

The study hospital used different ways to improve staff knowledge and enhance patient safety, however, some participants said there was no specific training on VRIII’s use or its complications. The healthcare practitioners stated that completing the diabetes e-learning module with a pass did not necessarily mean they had gained the practical benefits expected by the module. With this in mind, some suggested that face-to-face in-house diabetes and VRIII-focused training sessions, tailored to the practical aspects of their work, would be more effective than e-learning.

**Challenges encountered while using VRE and suggestions for change**

Some challenges were encountered during the VRE stages. At the patient recruitment stage, the researcher relied on the acute patient cases admitted to the Unit to be recruited. Patient recruitment might have been enhanced by having access to the surgery schedule for elective cases, enabling alignment of the patients’ schedule with the times when the researcher was available in the Unit.

A key concern with using video approaches is their effect on practice and on the communication between the participants and the patients, and the participants and their colleagues. Existing literature confirmed that there is no evidence video-recording causes significant alteration to the way participants usually behave [29, 43]. Although at the beginning of this study the healthcare practitioners tried to change their work because of the mere presence of the researcher’s handheld camera, the prolonged presence of the researcher in the field unit helped the healthcare practitioners get accustomed to the camera. Using a chest-mounted camera would likely have made the recording more efficient by enabling hands-free recording and filming from a different angle without making healthcare practitioners worried when the researcher followed them while holding a camera.

In terms of acceptability, the most prominent concern healthcare practitioners raised was related to the privacy and confidentiality of the patients. Confidentiality can be ensured by masking participants’ identifiers, by discussing anticipated threats to confidentiality and anonymity with participants, and by giving them explicit information, on the consent form, about the status and use of video-recordings in the research [44, 45]. However, as the healthcare practitioners have not used to have cameras in the hospital, the researcher had a sense that healthcare practitioners were using concerns about patient confidentiality to disguise their own reluctance to be filmed and the potential consequences of being revealed to be part of an error and patient safety incident.

**Strengths and limitations of the study**

Quality and safety improvement initiatives cannot be understood outside their context, and initiatives can only influence work when healthcare practitioners agree that strategies proposed for implementing these initiatives will improve their work [46]. The novelty of the present study’s use of VRE methodology lies in the fact that its innovations arise from within established work (exnovation), and from within practitioners’ collective sense-making of their work.

This study identified several strategies that might enhance safety in the use of VRIII in the study hospital, including improving electronic prescriptions by providing preparatory training sessions for senior and junior doctors; face-to-face training; and teaching sessions on VRIII that are focused on the practical needs of the healthcare practitioners. CBG monitoring and independent verification of prescribing and administering VRIII are important strategies for enhancing safety, meaning that attention should be directed towards investing resources in ensuring the consistency and continuity of experienced staff over time. The study’s findings could potentially be discussed with healthcare practitioners, NHS
safety professionals and diabetes guideline developers, with a view to exploring the applicability of the suggested solutions and assessing how they might influence future VRIII guidelines and policies.

Patient and public involvement is recommended as best practice. Although there was no active patient involvement in this study, healthcare practitioners were proactively involved in the study conduct and analysis, engaging in designing ways to video themselves and patients as well as analysing their work in the reflexive meetings, a process revealing opportunities for improvement by learning lessons from the everyday work that would otherwise remain undetected.

The small number of participants engaged in this study and the low number of healthcare practitioners returning for the reflexive meetings could be considered a limitation of this study. However, using a mixed method approach to explore a specific phenomenon, i.e. the use of VRE in a Vascular Surgery Unit, along with engaging participants in analysis of their own work, enhanced the credibility and transferability of the study findings. Earlier studies using VRE methodology in healthcare, conducted reflexive meetings as group discussions intended to investigate participants’ knowledge and experience of their work [27, 38, 47]. A hallmark of an effective feasibility study is that its method can be adapted when necessary to achieve the most promising results [30, 48]. In this study, adaptation needed to be made in relation to conducting the video reflexive meetings because of the COVID-19 pandemic. Although there were initial concerns that having only one healthcare practitioner in each reflexive meeting might affect the depth of the discussion, it was found that healthcare practitioners were very keen to discuss the video clips, openly analyse their work and suggest solutions for improving the delivery of patient care.

We are aware that the current findings of this study provided snapshots of how VRIII was used to treat elevated CBG. Future work should focus on collecting more data to develop a richer, deeper view of the use of VRIII. Additional studies using VRE, need to tap into a wider range of patients and healthcare practitioners across many units in the same hospital used for this study and across different Trusts, to build a more comprehensive understanding of WAD in the use of VRIII and thus better inform policy makers and clinical work.

**Conclusions**

To the best of our knowledge, this study is the first to have explored the use VRE and quantitative data to understand how everyday work is done in reference to the use of VRIII. The results of this study demonstrated the feasibility of the recruitment and data collection procedures used. VRE methodology was generally accepted by patients and healthcare practitioners, as the majority of those recruited consented and agreed to take part in the study. The VRE had the utility to explore everyday work and to add depth to understanding how everyday tasks were accomplished. It also generated increased reflexivity in healthcare practitioners, which was a precursor to enhancing their knowledge and insights into how to act amidst complexity by proposing practical solutions tailored to their needs.

**Abbreviations**

BG: Blood glucose

CAS: Complex adaptive system

COVID-19: Coronavirus

DISN: Diabetes inpatient specialist nurse

EPR: Electronic Patient Record

IV: Intravenous

NA: Nurse assistant

NaDIA: The National Diabetes Inpatient Audit

NHS: The National Health Service

RHC: Resilient Health Care

VRE: Video Reflexive Ethnography

VRIII: Variable rate intravenous insulin infusion

WAD: Work as Done

WAI: Work as Imagined
Declarations

Ethics approval and consent to participate

The study was approved by the University of Reading Research Ethics Committee (UREC) (ref: 18/03), NHS Research Ethics Committee (REC) (ref: 18/SC/0456), Health Research Authority (HRA) (ref: 18/SC/0456), and the Oxford University Hospitals NHS Foundation Trust through the Research and Development Approval Processes (ref: 13827). The participants provided written informed consent before the video recordings and verbally confirmed their continuing consent before the commencement of the reflexive meetings.

Consent for publication

Not applicable.

Availability of data and materials

The unpublished data used in the current study are available from the corresponding author on request.

Competing interests

Dr Rosemary Lim is an Associate Editor of the journal BMC Health Services Research. All other authors declare that they have no competing interests.

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Authors’ contributions

MI, RL, CC and RI made a substantial contribution to the design of the work. MI collected the data, transcribed the video footage and analysed the video and the reflexive meeting transcripts. MI, RL, CC, RI and FG extensively discussed how to analyse and interpret the data. RL supervised the analysis of the reflexive meeting transcripts and all authors participated in discussing the themes. MI drafted the initial manuscript. All authors revised the manuscript critically for intellectual content, agreed and approved the final version to be published.

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Figures
Figure 1
Study flow diagram

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