Effect of a smart temperature logger on correctly storing biological disease-modifying antirheumatic drugs at home: a pre-post study

Lex I Haegens,1 Victor J B Huiskes,2,3 Charlotte L Bekker,3 Bart J F van den Bemt2,3

ABSTRACT

Objectives Biological disease-modifying antirheumatic drugs (bDMARDs) require specific storage temperatures, but are frequently stored outside the recommended range of 2–8°C. As incorrect storage may affect therapy effectiveness and consequently lead to higher disease activity, compliance with recommended storage temperatures should be improved. eHealth interventions can provide insight into storage temperatures and alerts in case of deviations from recommended temperatures. Therefore, this study aims to assess the effect of a smart temperature logger on correctly storing bDMARDs at home by patients with rheumatic diseases.

Methods A pre-post study was performed in a hospital in the Netherlands. The baseline period consisted of 12 weeks of storage temperature measurement with a passive temperature logger, and the intervention period consisted of 12 weeks of storage temperature measurement with a smart temperature logger. This smart logger included a smartphone application which provided insight into storage temperatures and real-time alerts when exceeding recommended temperatures. The main outcome measure was the difference in the number of patients who stored their bDMARDs correctly between baseline and intervention. Secondary outcomes were the difference in the proportion of measurement time within 2–8°C between baseline and intervention, the distribution of measurement time among temperature categories, and the patient’s acceptance measured using a questionnaire based on the Technology Acceptance Model.

Results In total, 48 participants (median age 55 years [IQR 47–64], 53% male) were analysed. The proportion of participants correctly storing bDMARDs increased from 18.8% (n=9) during baseline to 39.6% (n=19) during intervention (p=0.004). The median proportion of measurement time between 2–8°C improved by 6% (IQR 0–34%) (p<0.0001). Technology acceptance was scored as moderate.

Conclusions Temperature monitoring and real-time feedback with a smart temperature logger shows potential to improve at-home storage of bDMARDs, provided that continuous connection is realised to ensure real-time alerts and data collection.

INTRODUCTION

Biological disease-modifying antirheumatic drugs (bDMARDs) are safe and effective drugs for the treatment of inflammatory rheumatic diseases if the treatment target is not achieved with conventional synthetic DMARDs alone.3 bDMARDs require storage between 2–8°C, as described in the Summary of Product Characteristics (SmPC) recommendations.6–8 Via a strictly monitored cold chain, bDMARDs are produced, stored and transported within this temperature range. But from the moment drugs are dispensed to the patient, storage conditions are uncontrolled and out of sight of healthcare professionals. Despite instruction to the patient regarding storage conditions by the pharmacy staff on dispensing, research has shown that over 90% of patients using bDMARDs do not store their drugs according to the SmPC instructions.9–11 Moreover, one study found that 26% of patients with rheumatoid arthritis stored their bDMARDs below 0°C or above 25°C for more than 2 hours consecutively, and 6% below 0°C for at least 24 hours consecutively.9 Storing bDMARDs outside the recommended temperature range can lead to formation of protein aggregates, which in turn may lead to formation of anti-drug antibodies.14 Anti-drug antibodies are immunogenic and result in reduced drug responses for bDMARDs.15,16 In short, incorrect storage of
bDMARDs may affect therapy effectiveness and lead to higher disease activity.\(^{17}\)

Compliance with recommended storage temperatures of bDMARDs should thus be improved. eHealth technologies have previously been applied for at-home monitoring of body temperature as a vital sign\(^ {18,19}\) or foot temperature in diabetes, for example,\(^ {20,21}\) and options for at-home temperature monitoring of medication storage have been outlined previously.\(^ {22}\)

eHealth can therefore be a valuable instrument to support patients in correctly storing drugs at home by giving patients insight into the actual storage conditions within their refrigerator. MedAngel One is a wireless temperature logger connected with a smartphone application that continuously monitors storage temperature and alerts users in real-time when actual storage temperatures exceed recommended storage temperatures. Temperature monitoring with real-time alerts might help patients to store bDMARDs adequately within the recommended temperature range. It is currently unknown if a temperature logger with a smartphone application is able to positively affect storage conditions of bDMARDs at home, and whether patients find such a device acceptable and easy to use.

Therefore, the primary objective of this study was to assess the effect of using a smart temperature logger on storing of bDMARDs at home according to the recommended temperature range by patients with inflammatory rheumatic diseases. Additionally, user experience was assessed using a questionnaire based on the Technology Acceptance Model.

**METHODS**

**Design and population**

This one-group pre-post-test study was performed between February 2021 and July 2022 in the outpatient pharmacy of the Sint Maartenskliniek, which is the largest rheumatology clinic in the Netherlands with 18 850 patients treated in 2021. Patients were eligible for participation if all the following criteria were met: (1) receive treatment by a rheumatologist for rheumatoid arthritis, psoriatic arthritis or spondyloarthritis (based on International Classification of Diseases, 10th Revision (ICD-10) codes in the patient’s medical record); (2) receive a bDMARD from the outpatient pharmacy prescribed by their rheumatologist; (3) possess a smartphone or tablet capable of running the software application accompanying the smart temperature logger; and (4) sufficiently write and understand the Dutch language.

**Sample size**

Prior research has shown that only 7% of patients with rheumatoid arthritis stored their drugs within the recommended 2–8°C.\(^ {9}\) We hypothesise that our intervention will be able to improve this to 20%. With a power of 0.8 and a two-sided \(\alpha\) of 0.05, a sample size of 124 was calculated. Including 20% loss to follow-up, we aimed for a sample size of 150 participants.

**Procedure and data collection**

Eligible patients received study information by email and were asked to provide digital informed consent. Participants were enrolled at their next bDMARDs delivery by the pharmacy, following the study procedure depicted in figure 1.

After inclusion, participants received bDMARDs for exactly 12 weeks from the outpatient pharmacy, delivered at home or dispensed in the pharmacy. A sealed passive temperature logger (TempyTag\(^ {23}\)) was placed on top of one drug package to measure storage temperature continuously, with instructions to use the package with the logger last, to ensure 12 weeks of temperature registration. Participants were instructed to store their bDMARDs as usual.

After 12 weeks, participants received their subsequent delivery of bDMARDs. During this intervention period, participants received a ‘smart’ intervention temperature logger. A passive temperature logger was also provided to ensure temperature registrations in the baseline and intervention periods were measured by the same device. During the intervention period, participants were instructed to instal the smartphone application accompanying the intervention temperature logger, register their temperature logger, and actively monitor their storage condition using the smartphone application. To ensure both data were gathered and past storage conditions were viewed, participants received weekly reminders via email to connect to and synchronise with their temperature logger. No specific instructions on how to handle alerts were provided other than participants should alter the way they store their bDMARDs in order to comply with the recommended temperature range.

After the intervention period, participants received a return envelope to return the passive temperature logger. Participants were allowed to keep the smart temperature logger for future use free of charge. Additionally, participants received an email with an invite to complete a digital questionnaire regarding user experience.

**Intervention**

The intervention used in this study, the MedAngel One, is a Bluetooth temperature logger that allows users to (retrospectively) monitor storage temperature of temperature-sensitive drugs 24/7 using the accompanying application for mobile phones. Additionally, the application provides the user with real-time alerts on their smartphone when temperature exceeds pre-defined temperature limits (set at 2–8°C for all participants), assuming an active Bluetooth connection. Although several temperature loggers exist, the MedAngel One was chosen as this temperature

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**Figure 1** Overview of study procedure. TAM, Technology Acceptance Model.
logger features both a smartphone and a tablet application giving insight into current and past recorded temperatures, and additionally can give a real-time alert in case of excursions outside 2–8°C.

Primary outcome
The primary outcome was the difference in the proportion of patients who correctly stored their bDMARDs within 2–8°C between baseline and the intervention period. Correct storage was defined as storing bDMARDs within 2–8°C for the total duration of measurements (1) without excursions outside this temperature range for more than 48 hours cumulatively, and (2) without excursions below 0°C or above 25°C for a duration of at least 2 hours consecutively, in accordance with a previous study performed by Vlieland et al.9

Secondary outcomes
Secondary outcomes were (1) the difference in proportion of total measurement time between 2–8°C between baseline and the intervention period, (2) the difference in distribution of measurement time across multiple temperature groups (<0°C, 0–2°C, 2–8°C, 8–15°C, 15–25°C and >25°C) between baseline and the intervention period, and (3) user experience with the software application.

Questionnaire
The questionnaire to assess individual participants’ acceptance of the intervention was based on the Technology Acceptance Model.24 This model states that technology use is dependent on the intention to use a technology. This intention is in turn dependent on the attitude towards using a technology, which is a product of the perceived usefulness and perceived ease of use of a technology.

Questions were derived from Davis24 (Perceived usefulness), Brooke’s System Usability Scale (SUS)25 (perceived ease of use), and the user version of the smartphone application Rating Scale (uMARS) by Stoyanov et al.26 Items from the uMARS questionnaire were screened and selected for applicability to this study by LH and CB. Selected questions were initially applied to the smartphone application and translated by LH, checked by CB, and adjusted by LH and CB until all discrepancies were resolved (English translation in online supplemental material A).

Data analysis
The start and end of measurement periods were determined for individual temperature loggers. Within participants, the shortest measurement time from either baseline or intervention temperature logger was applied to the other temperature logger measured from the start of measurements to ensure corresponding lengths of measurement times within participants (online supplemental material B). For the primary outcome measure, both baseline and intervention periods from individual participants were assessed binarily as correct or incorrect, according to the criteria outlined in the methods section. The difference in the proportion of participants who correctly stored bDMARDs between baseline and intervention was calculated by McNemar’s test for paired categorical data.

Proportions of measurement time were compared between baseline and intervention period by either paired t-test or Wilcoxon Signed Rank test, depending on normality of the data. Technology acceptance was assessed using the scoring methods of the incorporated questionnaires, if applicable, and reported using mean and SD or median and IQR, depending on the normality of the data.

RESULTS
Participants
A total of 640 eligible patients were invited to participate, of whom 151 agreed (24% response rate) and 140 (93%) started the baseline period. Of these participants, 89 (59%) resulted in retrievable baseline temperature measurements, after which 48 (32%) participants resulted in both retrievable baseline and intervention temperature measurements (figure 2). The questionnaire was completed by 90 participants (64%), as this was sent to all participants completing intervention measurements, regardless of retrievability of temperature data.

The participants’ median age was 55 (IQR 47–64) years with just over half the population being male (53%). The participants’ median time since diagnosis was 7 (IQR 2–17) years, and 47% had rheumatoid arthritis (table 1). Participants excluded from per-protocol analysis did not differ significantly from included participants (data not shown). After matching measurement times of baseline and intervention periods within participants, the mean (SD) total measurement time per patient per study phase was 71.8 (19.4) days.

Primary outcome
The proportion of participants who correctly stored bDMARDs increased significantly from 18.8% (n=9) during baseline to 39.6% (n=19) during the intervention period (p=0.004). The main improvement was seen in the number of participants who complied with the criterion of a maximum of 48 hours outside 2–8°C in total, with nine participants (18.8%) complying in the baseline period and 20 (41.7%) in the intervention period. Data on numbers of patients who stored bDMARDs in accordance with individual requirements for correct storage can be found in the online supplemental material C.

Secondary outcomes
Proportion within 2–8°C
The median proportion of measurement time per patient between 2–8°C improved statistically significantly with 6% (IQR 0–34%) (p<0.0001) (figure 3) from median 69% (IQR 19–96%) during baseline to 96% (IQR 70–99%) during intervention. Thirty-seven participants (77%) improved in the proportion of total measurement time within 2–8°C with a median of 24% (IQR 3–47%) between baseline and intervention. Nine participants (19%) worsened in proportion to the total measurement time within 2–8°C in total with a median of 4% (IQR 1–9%) between baseline and intervention. Two participants (4%) did not improve or worsen, of whom one participant had 100% and the other had 0% of total measurement time within 2–8°C during both baseline and intervention. A figure with individual baseline and intervention measurements can be found in the online supplemental material D.

Temperature distribution
The proportion of total measurement time within 2–8°C increased significantly from 59% to 79% (p<0.0001) and the proportion of total measurement time within 8–15°C decreased significantly from 36% to 19% (p<0.0001) (figure 4).

Technology acceptance questionnaire
Of the 90 participants who completed the intervention, 71% (n=64) completed at least one section and 49% (n=44)
fully completed the questionnaire. Results per questionnaire section below are based on participants who completed that particular section.

**Perceived ease of use**
All respondents completed the perceived ease of use section of the questionnaire. The mean (SD) SUS score was 61 (17), with 40% rating ease of use as good or excellent (SUS > 71.4). The mean (SD) score on the selected items from uMARS was 2.9 (1.0) out of 5.

**Perceived usefulness**
Sixty-two respondents (97%) completed this section of the questionnaire. Overall, participants’ perceived usefulness of the smartphone application was neutral to slightly positive (figure 5). Only usefulness of the application for disease management (‘By using this application, I think my rheumatic disease is doing better’) and engagement with their rheumatic disease (‘The application makes me more engaged with my rheumatic disease’) were perceived lower, with both statements scoring a median of ‘slightly disagree’.

**Attitude towards using**
Sixty respondents (94%) completed this section of the questionnaire. The application received a median of 3 (IQR 2–3) out of 5 stars: 27 of these respondents (45%) would use the application at least monthly; 33 respondents (53%) indicated they would use the application once per year or not at all; 47 (78%) respondents would recommend the application to other people who could benefit from it; and eight (13%) participants were willing to pay for the app.

**DISCUSSION**
This study showed that a smart temperature logger was able to increase significantly the proportion of participants who stored their bDMARDs correctly. Furthermore, participants stored bDMARDs significantly longer within the recommended

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**Table 1** Baseline characteristics of participants

<table>
<thead>
<tr>
<th>Characteristic</th>
<th>Value</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Sex</strong></td>
<td>Male (%)</td>
</tr>
<tr>
<td><strong>Age</strong></td>
<td>Median (IQR), years</td>
</tr>
<tr>
<td><strong>Diagnosis</strong></td>
<td>Rheumatoid arthritis, Psoriatic arthritis, Ankylosing spondylitis</td>
</tr>
<tr>
<td><strong>Disease years</strong></td>
<td>Median (IQR), years</td>
</tr>
<tr>
<td><strong>Prescribed bDMARD</strong></td>
<td>Adalimumab, Etanercept, Abatacept, Tocilizumab, Other</td>
</tr>
</tbody>
</table>

bDMARD, biological disease-modifying antirheumatic drug; IQR, Interquartile Range.
temperature range of 2–8°C using the intervention when compared with baseline. Most of this improvement was found in a shift from storage between 8–15°C to between 2–8°C. Technology acceptance was scored as moderate. However, technical defects with both baseline and intervention loggers impacted the number of participants included in the analysis and intended intervention exposure. Defects included baseline temperature loggers with (partly) irretrievable data and intervention temperature loggers with insufficient connection stability, possibly resulting in absence of real-time alerts and retroactively viewable temperature data.

This study investigated the effect of a smart temperature logger on the proportion of participants who stored bDMARDs correctly. We found that 18.8% of participants correctly stored bDMARDs during the baseline period. Although this number is higher than the 6.7% found by Vlieland et al, at-home storage of bDMARDs is still suboptimal without intervention.9 The proportion of measurement time within 2–8°C of 59% found in this study is comparable with the proportion of correct bDMARD storage time of 54.8% by Vlieland et al9 and other studies reporting on correct storage of bDMARDs.10 28 No studies have investigated an intervention to improve this suboptimal at-home storage of bDMARDs.

The intervention used in this study significantly improved the proportion of participants who correctly stored bDMARDs and the proportion of storage time within 2–8°C. However, the criteria for correct storage in this study were stricter than storage requirements stated in the SmPCs which state that bDMARDs can be stored at temperatures of up to 30°C for prolonged periods of time, ranging from 5 to 28 days.2–8 Therefore, the question remains whether this improvement has a clinically relevant effect (eg, on therapy effectiveness and safety, or disease activity) as improvement could have taken place in participants who already fell within the SmPC stated storage requirements.

In spite of the significant increase of storage time at the right conditions between baseline and intervention, in 19% of the participants the proportion of measurement time between 2–8°C decreased from baseline to intervention. This might be explained by the fact that patients inadvertently may have adjusted the way of storage wrongly but were insufficiently alerted by the smart temperature logger due to technical defects.

This study showed the potential of a smart temperature logger in improving at-home storage conditions of bDMARDs. However, technical problems encountered during the study should be resolved before the intervention is applicable in a real-world setting. Therefore, based on the experiences in this study, a smart temperature logger should possess several (technical) features. Connection stability between the smart temperature logger and the user’s smartphone should be guaranteed to

Figure 3 Individual difference in proportion of total measurement time within 2–8°C between baseline and intervention period. Dots represent individual participants.

Figure 4 Bar chart showing distribution of measurement time over temperature categories. *Statistically significant difference (p<0.05).
ensure real-time feedback on storage temperature. Due to the limited range of Bluetooth, the connection between smartphone and temperature logger could be improved by using an internet connection. Additionally, data collection should be ensured to enable users to monitor storage temperature retrospectively. Presuming Bluetooth signals might have difficulty penetrating certain refrigerators and is limited in connectivity range, other connections such as WiFi could be explored.

Although a smart temperature logger seems promising for improving at-home storage of refrigerated drugs under the right conditions, one could argue whether continuous monitoring is needed or whether this can be applied for specific therapy phases. For example, continuous monitoring and real-time feedback when temperature limits are exceeded may be useful during the first dispensing of refrigerated drugs, as patients get to understand how to store their drugs properly in combination with their refrigerator at therapy initiation. This may sufficiently improve at-home storage of bDMARDs at the right conditions in the long term and obviate the need for long-term continuous intervention use. On the other hand, as bDMARDs are expensive drugs, with prices ranging from €8000 to €12000 per patient per year depending on the bDMARD and dosing interval, disposal of medication due to incorrect storage can lead to economic losses. Furthermore, intervention exposure (ie, number of alerts generated) was not measurable due to technical defects, and participants possibly did not receive real-time feedback on storage temperature and might therefore be underexposed to the intervention. Therefore, it remains uncertain whether improvements in storage conditions are related to real-time feedback or to (retrospective) insight in storage temperature. Second, generalisability might be limited as participants were recruited digitally. Furthermore, participants were obliged to possess a smartphone. This potentially attracted a more digitally-inclined population when compared with the general rheumatic diseases population. Additionally, part of the effect of the intervention may be due to the written instructions patients received at the beginning of the study on how to adjust their storage behaviour when they received an alert of temperatures exceeding 2–8°C. Third, due to the pre-post design of this study with a single group, participants may have been aware of being observed and as a result altered their bDMARDs storage. This risk was minimised by not providing feedback on storage temperature during the baseline period. Moreover, participants received no instructions regarding bDMARDs storage at the start of the baseline period, only at the start of the intervention period. Lastly, seasonal weather changes may have influenced storage conditions within refrigerators, but since data collection for both the baseline and intervention period took place across all seasons, we assume this risk was minimal.

Figure 5  Median and IQR scores for statements regarding usefulness of the application accompanying the intervention temperature logger.
CONCLUSION

Temperature monitoring and real-time feedback with a smart temperature logger shows potential to improve at-home storage of refrigerated medication, provided that the smart temperature logger has adequate connection stability and ensured data collection.

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Contributors LH, VB, CB and BB drafted the manuscript; BB is guarantor of the study.

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Competing interests None declared.

Patient consent for publication Not applicable.

Ethics approval This study was performed in line with the principles of the Declaration of Helsinki (version 2013). The Medical Ethics Research Committee of Arnhem-Nijmegen waived official ethical approval (case-number 2020–7127) as this study was deemed not subject to the Medical Research Involving Humans Act. Informed consent was obtained from all individual participants before inclusion in the study.

Provenance and peer review Not commissioned; externally peer reviewed.

Data availability statement Data are available upon reasonable request.

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ORCID iD
Lex L Haegens http://orcid.org/0000-0003-1850-4365

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**Supplementary material**

**A. English translation of technology acceptance questionnaire.**

Below is the English translation of the originally Dutch questionnaire sent to participants to measure technology acceptance. The questionnaire is divided into 3 parts.

**Part 1: Perceived ease of use**

*Question 1 to 10 originate from the System Usability Scale, and are answered on a 5-point Likert scale (Strongly disagree, Somewhat disagree, Neutral, Somewhat agree, Strongly agree).*

1. I think that I would like to use the app frequently.
2. I found the app unnecessarily complex.
3. I thought the app was easy to use.
4. I think that I would need the support of a technical person to be able to use the app.
5. I found the various functions in the app were well integrated.
6. I thought there was too much inconsistency in the app.
7. I would imagine that most people would learn to use the app very quickly.
8. I found the app very cumbersome to use.
9. I felt very confident using the app.
10. I needed to learn a lot of things before I could get going with the system.

*Question 11 to 17 originate from user version of the Mobile Application Rating Scale (uMARS) Sections A through D, and are answered on 5-point Likert scales related to each individual question.*

11. Is the app interesting to use? Does it present its information in an interesting way?
12. Does the app allow you to customize the settings and preferences that you would like to (e.g. sound, content and notifications)?
13. Is the app content (visuals, language, design) appropriate for the target audience?
14. How accurately/fast do the app features and components work?
15. Is arrangement and size of buttons, icons, menus and content on the screen appropriate?
16. Is the app content correct, well written, and relevant to the goal/topic of the app?
17. Is the information within the app comprehensive but concise?

**Part 2: Perceived usefulness**

*Questions 18 to 23 originate from uMARS section F, and are answered on a 5-point Likert scale (1 – strongly disagree to 5 – Strongly agree).*

18. This app has increased my awareness of the importance of addressing the health behaviour.
19. This app has increased my knowledge/understanding of the health behaviour.
20. The app changed my attitudes toward improving this health behaviour.
21. The app has increased my intentions/motivation to address this health behaviour.
22. This app would encourage me to seek further help to address the health behaviour (if I needed it).
23. Use of this app will increase/decrease the health behaviour.
Part 3: Attitude towards using

Questions 24 to 27 originate from uMARS section E, and are answered on 5-point Likert scales adjusted to each question.

24. Would you recommend this app to people who might benefit from it?
25. How many times do you think you would use this app in the next 12 months if it was relevant to you?
26. Would you pay for this app?
27. What is your overall (star) rating of the app?

B. Start- and end-times of measurement period.

Start of measurement period was defined as either 1) 24 hours after delivery or dispensing by pharmacy, or 2) 24 hours after the temperature logger first measured a temperature below 8 degrees Celsius in the case of delivery via mail. For participants where temperature loggers did not measure temperatures below 8 degrees Celsius, start of measurement period was defined as 24 hours after the temperature logger first measured a temperature below 15 degrees Celsius.

End of measurement period was defined as either 1) 12 weeks after the start of the measurement period, minus 24 hours or 2) the moment storage temperature exceeded 15 degrees Celsius for at least 12 hours consecutively without subsequent cooling below 15 degrees Celsius for at least 48 hours consecutively, minus 24 hours.

A forgiveness of 24 hours was incorporated to account for adjustment periods of temperature loggers for both start and end of measurement periods.
C. Numbers of patients that stored bDMARDs in accordance with individual requirements for correct storage.

Correct storage baseline vs intervention

<table>
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<th>Incorrect</th>
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<td><strong>39</strong></td>
<td><strong>48</strong></td>
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Correct storage, criterium ‘no more than 2 hours consecutively below 0 degrees’ baseline vs intervention

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<tbody>
<tr>
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<tr>
<td>Incorrect</td>
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<tr>
<td><strong>Total</strong></td>
<td><strong>43</strong></td>
<td><strong>5</strong></td>
<td><strong>48</strong></td>
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Correct storage, criterium ‘no more than 2 hours consecutively above 25 degrees’ baseline vs intervention

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<tr>
<td>Incorrect</td>
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<td><strong>Total</strong></td>
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<td><strong>0</strong></td>
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Correct storage, criterium ‘no more than 48 hours in total outside 2-8 degrees’ baseline vs intervention

<table>
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<tr>
<td>Incorrect</td>
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<tr>
<td><strong>Total</strong></td>
<td><strong>9</strong></td>
<td><strong>29</strong></td>
<td><strong>48</strong></td>
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</tbody>
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D. Individual baseline and intervention measurements on proportion of measurement time within 2 – 8 degrees Celsius.

Graph depicting the individual difference in proportion of measurement time between 2 – 8 degrees Celsius between baseline and intervention measurements. Red lines depict participants that improved from baseline to intervention, blue lines depict participants that declined from baseline to intervention period.