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Effectiveness of at-home skin temperature monitoring in reducing the incidence of foot ulcer recurrence in people with diabetes: a multicenter randomized controlled trial (DIATEMP)

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ABSTRACT

Introduction The skin of people with diabetic foot disease is thought to heat up from ambulatory activity before it breaks down into ulceration. This allows for early recognition of imminent ulcers. We assessed whether at-home monitoring of plantar foot skin temperature can help prevent ulcer recurrence in diabetes.

Research design and methods In this parallel-group outcome-assessor-blinded multicenter randomized controlled trial (7 hospitals, 4 podiatry practices), we randomly assigned people with diabetes, neuropathy, foot ulcer history (<4 years, n=295), or Charcot's neuro-arthropathy (n=9) to usual care (ie, podiatric treatment, education, and therapeutic footwear) or usual care plus measuring skin temperatures at 6-8 plantar sites per foot each day (enhanced therapy). If ΔT >2.2°C between corresponding sites on the left and right foot for two consecutive days, participants were instructed to reduce ambulatory activity until this hotspot disappeared and contact their podiatrist. Primary outcome was ulcer recurrence in 18 months on the plantar foot, interdigital, or medial/lateral/anterior forefoot surfaces; secondary outcome was ulcer recurrence at any foot site.

Results On the basis of intention-to-treat, 44 of 151 (29.1%) participants in enhanced therapy and 57 of 153 (37.3%) in usual care had ulcer recurrence at a primary outcome site (RR: 0.782 (95%CI 0.566 to 1.080), p=0.133). Of the 83 participants in enhanced therapy who measured a hotspot, the 24 subsequently reducing their ambulatory activity had significantly fewer ulcer recurrences (n=3) than those in usual care (RR: 0.336 (95% CI 0.114 to 0.986), p=0.017). Enhanced therapy was effective over usual care for ulcer recurrence at any foot site (RR: 0.760 (95% CI 0.579 to 0.997), p=0.046). **Conclusions** At-home foot temperature monitoring does not significantly reduce incidence of diabetic foot ulcer recurrence at or adjacent to measurement sites over usual care, unless participants reduce ambulatory activity when hotspots are found, or when aiming to prevent ulcers at any foot site. Trial registration number NTR5403.

INTRODUCTION

With a life-time incidence up to 34%, foot ulceration is a common complication in people with diabetes that poses a large burden

Significance of this study

What is already known about this subject?

- A meta-analysis of four randomized controlled trials (RCTs) published in 2020 reported a pooled OR of 0.51 (95% CI 0.31 to 0.84, p=0.009) favoring athome skin temperature monitoring over usual care in preventing foot ulcers in people with diabetes.
- All RCTs showed large Cls around the effect and have a high risk of bias, with the meta-analysis concluding that this intervention deserves further evaluation in larger studies.

What are the new findings?

- At-home monitoring of plantar foot skin temperature non-significantly reduced the incidence of foot ulcer recurrence at or adjacent to a measurement site by 22% relative to usual care.
- Adherence to reducing ambulatory activity when a "hotspot" is found has a large positive effect on clinical outcome.
- The intervention significantly reduced the incidence of ulcer recurrence at any foot site with 24% relative to usual care.

How might these results change the focus of research or clinical practice?

Foot care providers should selectively recommend at-home foot temperature monitoring to people with diabetes and high ulcer risk who are likely to reduce ambulatory activity when a hotspot is found or as method to increase foot awareness and help prevent an ulcer at any foot site.

on the patient and healthcare system.^{1–3} The risk for ulceration is particularly high in people with a foot ulcer history: 40% within 1 year after healing.¹ Therefore, prevention of foot ulcers and their recurrence is important, but it is underexposed in research and clinical practice.⁴ The need for more randomized controlled trials (RCTs on this topic has

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been emphasized, together with widespread adoption of evidence-based ulcer prevention programs.⁵⁶

The most common mechanism of ulceration, particularly on the plantar foot, involves the cumulative effect of repetitive stress during ambulation that goes unrecognized because of peripheral neuropathy.¹ International guidelines therefore recommend pressure-relieving footwear for people at high risk of plantar ulceration.⁶ But the alarmingly high ulcer recurrence rates necessitate adjunctive modalities for prevention.¹ It has been suggested that foot ulceration is preceded by increased local skin temperature due to inflammation (enzymatic autolysis) of the tissue caused by accumulating mechanical stress from being ambulatory.^{7 8} These increased local skin temperatures can be assessed by patients in their own homes using thermometry, thus providing a method for early recognition of this sign of impending ulceration.⁹ This method allows people at-risk and care providers to act timely by reducing ambulatory activity or providing (further) pressure relief to reduce the local inflammation. Three RCTs investigated such at-home foot temperature monitoring and showed large effects in reducing incidence of foot ulcer recurrence in people with diabetes.^{10–12} Therefore, this approach has been recommended in international guidelines.⁶

Notwithstanding these findings and recommendations, skin temperature monitoring is currently rarely used in preventative foot care. This may be because of generalizability issues with the three RCTs being conducted by the same research group in one geographical location in the USA in the early 2000s. A more recent pilot RCT from Norway using the same intervention found no beneficial effect on ulcer recurrence, but was underpowered.¹³ An important scientific argument relates to the fact that all four previous RCTs had as their primary outcome an ulcer at any foot site, while temperatures were only measured at six predefined local sites on the plantar foot. If measuring local skin temperature is the key to this intervention, one would expect a reduction in ulcer incidence at or adjacent to measurement sites, but not necessarily elsewhere on the foot. Ulcer sites are not reported in these four trials, which limits interpretation of results. More practical reasons for a lack of implementation may be the minimal attention for this intervention in ulcer prevention guidelines until 2015, the scarcity of proper measurement equipment, and the burden of long-term daily measurement and false-positive readings on a population already monitoring many aspects of their disease.⁶¹⁴¹⁵

In addition to these arguments, recent meta-analyses have reported uncertainty over the effect of this intervention, deserving further evaluation in larger studies.^{16–18} For these reasons, we assessed the effectiveness of at-home infrared foot temperature monitoring on the incidence of foot ulcer recurrence in people with diabetes and hypothesized to find significantly less ulcers at or adjacent to measurement sites when compared with usual care.

RESEARCH DESIGN AND METHODS Study design

This study was an investigator-initiated, multicenter, outcome-assessor blinded, two study-arm parallel-group RCT with a 1:1 allocation ratio. The medical ethics committee of Amsterdam University Medical Center (UMC) as coordinating center approved the protocol for this RCT (ID 2015_105), which has been published in detail elsewhere,¹⁹ and is summarized below. All participants gave informed consent before taking part.

Setting and recruitment

We enrolled participants from the multidisciplinary diabetic foot clinics of three university hospitals and four community hospitals, and from four affiliated podiatry practices, spread across the Netherlands (ie, in 6 of the 12 provinces).

Participants

The inclusion criteria were: diagnosis of diabetes mellitus type 1 or 2; age ≥ 18 years; loss of protective sensation (LOPS) as a result of peripheral neuropathy; history of a foot ulcer or an amputation in the 4years preceding randomization, or diagnosis of Charcot neuro-osteoarthropathy in chronic state (ie, International Working Group on the Diabetic Foot risk grade 3);⁶ ambulatory status (ie, not wheel-chair bound); regular foot care provided by a podiatrist or willingness to undergo such care; and ability to follow study instructions. Exclusion criteria were: foot ulcer or open amputation site; active Charcot neuro-osteoarthropathy; foot infection; chronic limb-threatening ischemia, as defined by previously published criteria;²⁰ bilateral amputation proximal to the tarsometatarsal joint; severe illness that would make 18-month survival unlikely; or current use of at-home foot temperature monitoring.

Interventions

Usual care, as provided in the Netherlands according to evidence-based guidelines,⁶ consisted of: (a) professional foot care and foot screening once every 1–3 months by a podiatrist; (b) therapeutic (custom-made) footwear, if indicated based on ulcer risk and foot condition; and (c) education about self-care practices, ulcer risk factors, and ulcer etiology. Education was provided via verbal and written information by a clinician or an investigator at baseline and ad libitum by clinicians during follow-up clinic visits. All participants were advised to contact their podiatrist if they identified any area of concern on their foot.

Enhanced therapy consisted of usual care plus at-home measurement of plantar foot skin temperature, per instruction once per day in the morning directly after waking up. Participants used an infrared thermometer (TempTouch, Diabetica Solutions, San Antonio, Texas, USA)¹¹ to measure skin temperature at six predefined sites on the plantar surface of both feet where foot ulcers most commonly occur:²¹ hallux, second and third

toe, first, third and fifth metatarsal heads. A maximum of two additional plantar foot sites were measured if a previous ulcer or preulcerative lesion (ie, abundant callus, hemorrhage, or blister) had been or was present at another than a predefined site. This selection of measurement sites was different than in previous trials, where only six predefined sites (four at the forefoot, one midfoot, and one heel) were used.¹⁰⁻¹³ Participants with a minor or unilateral major amputation that prevented measurement at a predefined site, measured at an adjacent site, or used average temperatures during run-in for comparison, as described in our protocol.¹⁹ Participants recorded each temperature value and the difference between corresponding sites on both feet in a customized form. These forms were returned to the investigator after 2weeks at the start and on a 4weekly basis thereafter.

If the temperature difference at corresponding sites was >2.2°C for two consecutive days, it was defined as a "hotspot". Participants were instructed verbally and in writing on their form to then substantially reduce their ambulatory activity, that is, by at least 50% as judged subjectively, until the temperature difference normalized to <2.2°C, and to contact their podiatrist for further instruction and, if needed, treatment.¹¹ Participants recorded these actions in their forms. When foot temperature difference exceeded 4°C or did not normalize in 2 days, participants were instructed to immediately contact and see their podiatrist. Participants received mobile-phone text reminders two times a week for the first 6weeks and once every 2 weeks for the remainder of follow-up, to encourage them in measuring their foot temperatures and in reducing ambulatory activity and contacting their podiatrist if a hotspot was found.

Procedures

After providing informed consent, participants underwent assessment at their study center by an investigator. Demographic and disease-related characteristics were obtained. LOPS was assessed and confirmed to be present when the pressure of a 10g (5.07) Semmes-Weinstein monofilament was not felt at ≥ 2 sites of plantar hallux, first and fifth metatarsal heads on both feet or when the vibration of a 128Hz Tuning fork placed on the apex of the hallux was not felt.²² Peripheral artery disease was assessed and classified according to previously published procedures,²⁰ first by palpation of the dorsalis pedis and posterior tibial pulses of both feet. If non-palpable, the participant's medical file was checked for vascular status and, if status was not clear, toe pressures were assessed. Foot deformity was assessed clinically for presence of hammer/claw toes, prominent metatarsal heads, hallux valgus, pes planus, pes cavus, and Charcot deformity and classified into one of four categories according to the severity of deformity present.²³

If after assessment study eligibility was confirmed, participants were randomly assigned to usual care or enhanced therapy using an independent onlineaccessible computer-generated allocation sequence that used the nondeterministic minimization method.¹⁹ Randomization was stratified according to participating center and sex.

Participants allocated to enhanced therapy watched a video with instructions for at-home measuring and recording of foot temperatures, and for activity reduction and podiatrist contact when a hotspot was found. These participants did a first complete measurement of foot temperature to demonstrate ability in doing so, and to identify if any further instruction was needed. Participants were then handed all necessary equipment and materials to take home.

All participants were followed for 18 months for study outcomes. An investigator contacted participants every 3 months by phone, or more frequently if needed, to ask about study execution, foot ulcer development, and any contact with a foot care provider. If participants identified or suspected an ulcer in-between regular podiatry visits, they were instructed to immediately see their podiatrist or multidisciplinary foot team for diagnosis and foot care. The foot care provider debrided the specific area if required to assess outcome, and in case a (suspected) ulcer was classified according to the University of Texas system, took photographs of the lesion and completed an outcome case report form.^{19 24} The investigator received these materials and then sent them anonymized to a panel of three to five foot care specialists for blinded assessment of ulcer outcome, as described in our protocol.¹⁹

Outcomes

The primary outcome for this study was the proportion of participants with a recurrent foot ulcer in 18 months at a primary site (ie, the plantar foot, interdigital space or medial, lateral, or anterior forefoot). These are primary sites because they are at or adjacent to a temperature measurement site where any increased temperature is most likely picked up, increasing validity for group comparisons on effectiveness of this intervention. A foot ulcer was defined as a full-thickness lesion through the dermis without reference to time present.²⁵ Study endpoints were ulceration at a primary site, death, or 18-month follow-up. Secondary outcomes analyzed using intention-to-treat (ITT) were for ulcer recurrence at a primary site (i) complicated by ischemia or infection; (ii) per-center category (ie, UMC, community hospital, or podiatry practice); and (iii) per center; and analyzed perprotocol in participants who: (iv) measured foot temperature >70% of days until endpoint and (v) reported to have reduced ambulatory activity when a hotspot was found. Other secondary outcomes analyzed using ITT were for: ulcer recurrence at (i) the previous ulcer site; (ii) an exact measurement site; and (iii) any foot site (which was the primary outcome in previous trials.^{10–13} Any serious adverse event (SAE) was promptly reported to the accredited medical research ethics committee for assessment.

Pathophysiology/complications

Sample size calculation

Based on data from a previous trial including the same study centers,²³ we anticipated a 44% ulcer recurrence incidence in 18 months for usual care. Using a more conservative estimate for relative reduction of incidence of 35% compared with the 61%–85% found in three previous trials,^{10–12} we anticipated a 28.6% ulcer recurrence incidence for enhanced therapy. With α 0.05 (two-sided), power 80%, χ^2 analysis, and ITT analysis for which primary outcome data from all participants could be obtained, 304 eligible participants were required.

Statistical analysis

Identification of group allocation and statistical analysis were performed after the last follow-up visit of the last participant and was conducted using SPSS statistical software (V.26.0, SPSS, Chicago, Illinois, USA). All tests assessed group effects were two-sided and used p<0.05 as significance level. Baseline participant characteristics were assessed with independent sample t-tests when data were normally distributed or Mann-Whitney U tests when data were not normally distributed. Effectiveness of the intervention was assessed using Pearson's χ^2 analysis. Ulcer outcome data from patients who died during study follow-up were based on outcome at moment of death (last observation carried forward). From participants who discontinued their study participation, information on outcome at 18 months for the purpose of the ITT analysis was obtained from their medical file after informed consent from the participant was obtained. Outcome of ulcer recurrence over time was assessed using log-rank testing and presented as Kaplan-Meier plots that were censored for death.

RESULTS

Baseline characteristics

A study flow diagram is shown in figure 1. Participants were recruited between November 5, 2015 and June 12, 2018, and the last participant follow-up was on December 12, 2019. A total of 295 participants were included based on a foot ulcer history and nine on having a Charcot foot. Baseline participant characteristics are shown in table 1.

Ulcer recurrence at a primary site

Ulcer outcome data is provided in table 2. A total of 101 participants (33.2% of the total group) had a recurrent foot ulcer at a primary site in 18 months. All ulcers were in participants with a foot ulcer history. Of participants included in the UMCs, significantly more ulcerated (45.5%) compared with those included in the community hospitals (32.1%) or podiatry practices (22.0%, p=0.005). There was no significant effect of sex or ethnicity on the primary outcome.

ITT analysis

In the enhanced therapy group, 44 of 151 (29.1%) participants had a recurrent ulcer at a primary site, which was not significantly different from the 57 of 153 (37.3%)

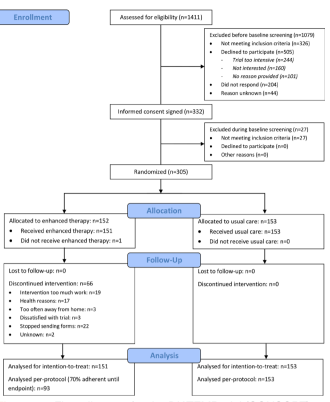


Figure 1 Flow diagram for the DIATEMP trial (CONSORT).

participants in the usual care group (RR 0.782 (95%CI: 0.566 to 1.080), p=0.133). Kaplan-Meier curves were also not significantly different between groups (Log Rank 1.907, p=0.167) (figure 2A).

Of the 134 participants (44.0%) enrolled in a community hospital, 13 (of 68, 19.1%) in enhanced therapy had a recurrent foot ulcer at a primary site, which is significantly lower than the 30 (of 66, 45.5%) in usual care (RR 0.421 (95% CI 0.241 to 0.733), p=0.001) (table 2).

Adherence and hotspots

Ninety-four participants in enhanced therapy (62.3% of total) measured foot temperature at least 70% of days until a study endpoint. Seventeen participants (11.2%) never measured foot temperature and 51 (33.8%) did not have a hotspot during follow-up, as analyzed from returned weekly logs. A total of 83 participants (55.0%) had at least one hotspot during follow-up. Of these 83, 24 (28.9%) reported reducing their ambulatory activity level with at least 50% and 14 (16.9%) reported contacting their podiatrist with at least one hotspot, of which 12 did both. With 32.5% of the hotspots found, the participant either reduced ambulatory activity or contacted the podiatrist. In 506 of the total 5862 weeks (ie, 112.7 person-years) of registered temperature measurements, a hotspot was found; this equates to 4.5 hotspots/person-year.

Per-protocol analysis

Thirty-two of the 94 (34.0%) participants who were adherent to temperature monitoring had a recurrent

Table 1 Baseline characteristics of the int	ention-to-treat popu	lation		
Characteristic	All	Enhanced therapy	Usual care	Missing values
Number of participants	304	151	153	
Age (years)	64.6±10.5	65.0±10.6	64.2±10.5	
Male sex	220 (72.4%)	109 (72.2%)	111 (72.5%)	
Ethnic origin: Caucasian	283 (93.1%)	140 (92.7%)	143 (93.5%)	
Type of diabetes				4 (1.3%)
Type 1	66 (21.7%)	30 (19.9%)	36 (23.5%)	
Type 2	234 (77.0%)	119 (79.9%)	115 (75.3%)	
Years diagnosed with diabetes	20±14	20±14	21±15	3 (1.0%)
HbA1c (mmol/mol)	60.7±16.0	60.5±16.5	60.9±15.5	65 (21.4%)
Body mass index (kg/m²)	29.8±5.3	29.6±5.4	30±5.3	1 (0.3%)
Retinopathy	151 (49.7%)	83 (55.0%)	68 (44.4%)	2 (0.7%)
Nephropathy	60 (19.7%)	31 (20.5%)	29 (19.0%)	1 (0.3%)
Dialysis	4 (1.3%)	0 (0.0%)	4 (2.6%)	
Smoking or history of smoking	169 (55.6%)	88 (58.3%)	81 (52.9%)	
≥1 unit alcohol intake per week	199 (65.5%)	102 (67.5%)	97 (63.4%)	
Living alone	105 (34.5%)	52 (34.4%)	53 (34.6%)	
Using a walking aid	89 (29.3%)	49 (32.5%)	40 (26.1%)	
Education			, , , , , , , , , , , , , , , , , , ,	2 (0.7%)
Low	117 (38.5%)	61 (40.4%)	56 (36.6%)	
Medium	96 (31.6%)	45 (29.8%)	51 (33.3%)	
High	89 (29.3%)	44 (29.1%)	45 (29.4%)	
Employed	75 (24.7%)	38 (25.2%)	37 (24.2%)	
Footwear				2 (0.7%)
Conventional	97 (31.9%)	52 (34.4%)	45 (29.4%)	
Semi custom-made	37 (12.2%)	19 (12.6%)	19 (12.4%)	
Full custom-made	168 (55.3%)	80 (53.0%)	89 (58.2%)	
Walking barefoot at home	113 (37.2%)	54 (35.8%)	59 (38.6%)	
Participating center category				
University Medical Center	88 (28.9%)	42 (27.8%)	46 (30.1%)	
Community hospital	134 (44.1%)	68 (45.0%)	66 (43.1%)	
Podiatry practice	82 (27.0%)	41 (27.2%)	41 (26.8%)	
Participating center	02 (21.070)			
Amsterdam UMC, location AMC	95 (31.3%)	47 (31.1%)	48 (31.4%)	
Amsterdam UMC, location VUmc	18 (5.9%)	9 (6.0%)	9 (5.9%)	
Maastricht UMC+	25 (8.2%)	12 (7.9%)	13 (8.5%)	
Ziekenhuisgroep Twente	49 (16.1%)	23 (15.2%)	26 (17.0%)	
Reinier de Graaf Gasthuis	41 (13.5%)	21 (13.9%)	20 (17.0%)	
Maxima Medisch Centrum	64 (21.1%)	32 (21.2%)	32 (20.9%)	
Medisch Spectrum Twente	12 (3.9%)	7 (4.6%)	5 (3.3%)	
Frequency of professional foot care, every:	12 (0.370)	7 (4.070)	0 (0.070)	
1–4 weeks	82 (27.0%)	41 (27.2%)	41 (26.8%)	
5–8 weeks	199 (65.5%)	102 (67.5%)	97 (63.4%)	
>8 weeks				
LOPS based on inability to sense	23 (7.6%)	8 (5.3%)	15 (9.8%)	
10g monofilament	276 (90.8%)	138 (91.4%)	138 (90.2%)	
rog monomament	210 (30.070)	130 (91.470)	130 (90.2%)	

Table 1 Continued				
Characteristic	All	Enhanced therapy	Usual care	Missing values
128Hz tuning fork only	28 (9.2%)	13 (8.6%)	15 (9.8%)	
Peripheral artery disease				
Grade 1	230 (75.7%)	114 (75.5%)	116 (75.8%)	
Grade 2	74 (24.3%)	37 (24.5%)	37 (24.2%)	
Foot deformity				
Absent	17 (5.6%)	10 (6.6%)	7 (4.6%)	
Mild	58 (19.1%)	32 (21.2%)	26 (17.0%)	
Moderate	202 (66.4%)	99 (65.6%)	103 (67.3%)	
Severe	27 (8.9%)	10 (6.6%)	17 (11.1%)	
Minor lesions at entry*	121 (39.8%)	54 (35.8%)	67 (43.8%)	31 (10.2%)
Amputation†				
No amputation	223 (73.4%)	110 (72.8%)	113 (73.9%)	
Lesser toe(s)	29 (9.5%)	14 (9.3%)	15 (9.8%)	
Hallux or ray	39 (12.8%)	19 (12.6%)	20 (13.1%)	
Forefoot	6 (2.0%)	5 (3.3%)	1 (0.7%)	
Major	7 (2.3%)	3 (2.0%)	4 (2.6%)	
Previous ulcer site				1 (0.3%)
Plantar forefoot	95 (31.3%)	50 (33.1%)	45 (29.4%)	
Medial/lateral/interdigital/apex forefoot	104 (34.2%)	52 (34.4%)	52 (34.0%)	
Plantar midfoot/hindfoot	23 (7.6%)	11 (7.3%)	12 (7.8%)	
Dorsal side of the foot	72 (23.7%)	34 (22.5%)	38 (24.8%)	
No previous ulcer (ie, Charcot foot)‡	9 (3.0%)	4 (2.6%)	5 (3.3%)	
Months between healing of most recent ulcer and study entry	7 (2–14)	8 (2–15)	6 (2–12)	10 (3.3%)‡
Months duration of last two previous ulcers	4 (2–9)	4 (2–8)	4 (2–9)	1 (0.3%)

Data are n (%), mean±SD, or median (IQR).

*Minor lesion defined as a hemorrhage, blister, abundant callus, or erythema, identified at entry and confirmed present from photographic assessment.

†In case of bilateral amputation, the highest level was chosen.

‡Including nine participants that were included based on having a history of Charcot neuro-osteoarthropathy and having no history of ulceration.

AMC, Academic Medical Center; LOPS, loss of protective sensation; UMC, University Medical Center; VUmc, Vrije Universiteit Medical Center.

foot ulcer at a primary site (table 2). This was not significantly different from the 57 of 153 in the usual care group (RR 0.914 (95% CI 0.645 to 1.295), p=0.610). Of the 24 of 83 participants who reported to have reduced ambulatory activity when finding a hotspot, three (12.5%) had a recurrent foot ulcer at a primary site. This was significantly lower than the 21 of 59 participants (35.6%) who found a hotspot but did not reduce ambulatory activity (RR 0.351 (95% CI 0.115 to 1.069), p=0.035), and significantly lower than the 37.3% in usual care (RR 0.336 (95% CI 0.114 to 0.986), p=0.017).

Secondary analyses of ulcer recurrence at alternative sites

Thirty-two ulcers (31.7% of all ulcers) recurred at a previous ulcer site and 59 (58.4% of all ulcers) at a

measurement site, with a significant group effect found for the former (table 2). A total of 126 participants (41.4%) had a recurrent ulcer at any foot site, of which 54 were in enhanced therapy and 72 were in usual care (RR 0.760 (95% CI 0.579 to 0.997), p=0.046). Kaplan-Meier curves were not significantly different between groups (Log Rank 3.514, p=0.061) (figure 2B).

Adverse events

Ninety-five SAE occurred in 70 participants during follow-up, of which 8 deaths and 87 hospital admissions occurred (32 because of diabetic foot disease). Incidence of SAE was not significantly different between study groups (p=0.154) and none were reported to be related to the intervention.

Table 2 Study outcomes Outcome parameter Image: Control of the study outcomes	Enhanced the second		Polotivo rick (05% Ol)	Dyelus
Outcome parameter	Enhanced therapy	Usual care	Relative risk (95% CI)	P value
Number of participants	151	153		
Ulcer recurrence at primary site			0.700 (0.500 to 4.000)	0.400
Participants with ulcer	44 (29.1%)	57 (37.3%)	0.782 (0.566 to 1.080)	0.133
Ulcer site	0 (00 50/)			0.316
Hallux plantar/apex	9 (20.5%)	10 (17.5%)		
Toes plantar/apex	7 (15.9%)	4 (7.0%)		
Interdigital spaces	3 (6.8%)	3 (5.3%)		
Plantar metatarsal heads	15 (34.1%)	19 (33.3%)		
Medial border first ray	3 (6.8%)	9 (15.8%)		
Lateral border fifth ray	4 (9.1%)	6 (10.5%)		
Midfoot plantar	2 (4.5%)	4 (7.0%)		
Heel plantar	1 (2.3%)	2 (3.5%)		0.004
Reported cause of ulcer	04/77 00/)	45 (70.000)		0.994
Mechanical stress	34 (77.3%)	45 (78.9%)		
Direct trauma	3 (6.8%)	4 (7.0%)		
Ischemia	1 (2.3%)	1 (1.8%)		
Unknown	6 (13.6%)	7 (12.3%)		0.005
Ulcer per center category				0.005
University Medical Center	21 (50.0%)	19 (41.3%)	1.211 (0.765 to 1.195)	0.413
Community hospital	13 (19.1%)	30 (45.5%)	0.421 (0.241 to 0.733)	0.001
Podiatry practice	10 (24.4%)	8 (19.5%)	1.250 (0.549 to 2.846)	0.594
Ulcer per participating center				0 774
Amsterdam UMC, location AMC	15 (31.9%)	14 (29.2%)	1.094 (0.596 to 2.008)	0.771
Amsterdam UMC, location VUmc	5 (55.6%)	3 (33.3%)	1.667 (0.559 to 4.973)	0.343
Maastricht UMC+	8 (66.7%)	8 (61.5%)	1.083 (0.602 to 1.949)	0.79
Ziekenhuisgroep Twente	6 (26.1%)	11 (42.3%)	0.617 (0.271 to 1.402)	0.234
Reinier de Graaf Gasthuis	6 (28.6%)	8 (40.0%)	0.714 (0.301 to 1.694)	0.44
Maxima Medisch Centrum	2 (6.3%)	13 (40.6%)	0.154 (0.038 to 0.627)	0.001
Medisch Spectrum Twente	2 (28.6%)	0 (0.0%)	0.714 (0.447 to 1.141)	0.19
Complicated ulcer*	11 (25.0%)	19 (33.3%)	0.750 (0.400 to 1.408)	0.363
Ulcer in participants>70% adherent†	32 (34.0%)	57 (37.3%)	0.914 (0.645 to 1.295)	0.61
Ulcer in participants with reported activity reduction‡	3 (12.5%)	57 (37.3%)	0.336 (0.114 to 0.986)	0.017
Ulcer recurrence at alternative sites				
at previous ulcer site	10 (6.6%)	22 (14.4%)	0.461 (0.226 to 0.939)	0.028
at measurement site	23 (15.2%)	36 (23.5%)	0.647 (0.404 to 1.038)	0.067
at any foot site	54 (35.8%)	72 (47.1%)	0.760 (0.579 to 0.997)	0.046
Serious adverse events				0.154
Deaths	5 (3.3%)	3 (2.0%)		0.462
Hospital admissions	37 (24.5%)	28 (18.3%)		0.187

Data are n (%). Effects are shown as relative risk ratio with 95% CI for enhanced therapy relative to usual care.

*Complicated foot ulcer defined as a University of Texas depth 3 (ie, bone contact) or grade B, C, or D (ie, infection and/or ischemia present); analysis on proportion of complicated ulcers of all ulcers at a primary site.

†Per-protocol analysis in which for the enhanced therapy group 93 participants who measured their foot temperature on 70% or more of all follow-up days were included.

‡Per-protocol analysis in which for the enhanced therapy group 24 participants who reported to have reduced their ambulatory activity when finding a hotspot were included

AMC, Academic Medical Center; UMC, University Medical Center; VUmc, Vrije Universiteit Medical Center.

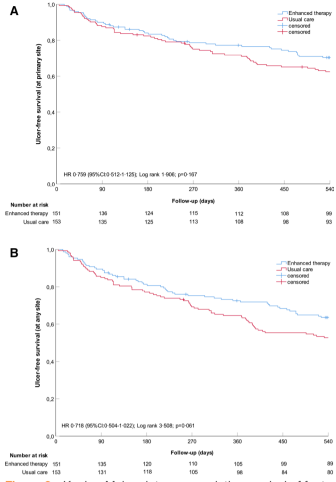


Figure 2 Kaplan-Meier plots on cumulative survival of foot ulcer recurrence over 18 months of follow-up with censored data for participants who died. Numbers at-risk are given per 3-month interval. (A) ITT on ulcer recurrence at primary site (primary); (B) ITT on ulcer recurrence at any site (secondary). ITT, intention-to-treat.

DISCUSSION

In the largest RCT on the topic to date, with enhanced therapy, we showed a non-significant relative 21.8% lower incidence of ulcer recurrence at a primary foot site compared with usual care alone. While not significant and potentially underpowered due to lower than expected ulcer incidence in the usual care group, the effect may still be attributable to the intervention and is a clinically important one given the burden of diabetic foot disease.² Our per-protocol analysis showed that adherence to monitoring foot temperatures had no effect on ulcer recurrence (relative 8.6% lower incidence); however, adherence to reducing ambulatory activity after identifying a hotspot did, with a relative 64.9% lower incidence found. Secondary analyses also showed that when ulcer recurrence at only the previous ulcer site (relative 53.9% lower incidence) or at any site on the foot (relative 24.0%lower incidence) was considered as outcome, enhanced therapy was effective over usual care. These results show a variable pattern of at-home foot temperature monitoring in prevention of foot ulcer recurrence.

We found a much smaller effect of enhanced therapy than the three American RCTs that used the same handheld thermometer and tested a similar population in a similar study design: relative 24% vs a 61%-85% lower incidence of ulcers an any foot site compared with usual care.¹⁰⁻¹² Our results are in line with the relative 22% lower incidence found in a more recent pilot RCT from Norway,¹³ and the relative 31% lower incidence per patient-year found in a very recent retrospective prepost temperature measurement cohort analysis.²⁶ Some study aspects were different, with the American trials being ~15 years older, performed by the same research group in one geographical region, having fewer participants with PAD, renal disease, or long-standing diabetes as risk factors and more with a diverse ethnic background than the European trials, and measuring at other predefined plantar foot sites. Furthermore, one American trial reported an absolute 35% higher adherence (64%) vs 29%) to reducing ambulatory activity when hotspots occurred compared with our study.¹¹ As reducing the cumulative stress on the foot is the primary suggested mechanism in ulcer risk reduction and temperature monitoring itself is only conditional to identify a hotspot and come into action, 78 a higher adherence is expected to result in better outcomes. Our per-protocol analysis supports this. It is unclear to what extent the variation in effect sizes between trials might be explained by above differences. Another RCT on this topic is ongoing,²⁷ and more are needed, to further clarify the preventive effect of at-home foot temperature monitoring.¹⁶¹⁷ More specifically, given the benefit of reducing ambulatory activity with a hotspot identified, studies should focus on the specific offloading actions required (including the continuous use of prescribed footwear) and on how to improve adherence to achieve a best possible effect from this intervention.^{28–30}

In secondary analyses, enhanced therapy was showed to be effective over usual care when the previous ulcer site was considered and showed a relative 35% reduction in ulcer incidence (although statistically not-significant) when the exact measurement sites were considered. This demonstrates that the smaller effect found for the primary outcome sites is mainly because of inclusion of adjacent sites. However, one should realize that only 32% of ulcers developed at a previous ulcer site, and other studies find even lower percentages.^{21 31} and only 58% of ulcers developed at a measurement site, limiting thermometry when only these sites are targeted. With a more liberal choice of ulcer at any foot site, the intervention was also effective over usual care. This may suggest a surrogate function of foot thermometry, increasing the participant's attention to the foot and the chance of picking up an early ulcer sign anywhere on the foot and acting on that. While previously a twice-per-day structured self-examination of the foot using a mirror to increase awareness did not show any benefit,¹¹ our effect found may be from being guided by quantitative measurements rather than just looking. Enhanced therapy was also effective over usual care for those participants enrolled in the community hospitals, but not for those enrolled in the UMCs or podiatry practices. Generally in the Netherlands, the most complex patients at highest risk are seen in UMCs and the least complex at lowest risk in the podiatry practices and we speculate that a single intervention may not differentiate adequately between study groups in these two settings; the optimum effect may be for those "medium" high-risk patients enrolled in community hospitals.⁴ Overall, these secondary analyses suggest that a benefit of the intervention may be dependent on outcome sites chosen, restricted to selected participants, and through a surrogate means of improving self-care. In support of this, a third of our study participants never identified a hotspot and would therefore not benefit from this intervention. Future studies should carefully consider participants and outcome sites and investigate above hypotheses so as to make targeted provision of this intervention possible⁴ and limit overtreatment and unnecessary patient burden. Furthermore, studies should assess the cost-effectiveness of this intervention (ongoing analysis of the current trial data) and the intervention as part of a multimodal treatment plan to move towards more personalized preventative care in diabetic foot disease (a new project we are starting).⁴

Adherence to monitoring foot temperatures was comparable to that found with other self-management strategies in diabetes,³² but disappointingly low for acting when hotspots occurred, while the working mechanism of the intervention is in this action. In explaining this, first, it may be that our instructions for reducing activity level were insufficiently clear, not clearly enough presented in the log or not memorized by the participant by the time a hotspot occurred, which could be months after study entry. Second, the complexity and burden of daily measuring and logging foot temperature in order for an event not to occur (ie, the "prevention paradox",⁴ may require too much effort of the participant to continue monitoring.⁴ ³³ Third, people with foot disease may develop the sense that they have little influence whatsoever on the outcome of foot ulceration, creating an otiose effort to control this.³⁴ Finally, participants may have judged that a hotspot found was not serious enough to require any action, which also relates to the problem of false-positive outcomes for this intervention.¹⁴ These aspects should be considered in the development of more user-friendly and effective technologies and methods for this purpose, which alarms users or their healthcare providers when a hotspot is found²⁶ and can provide specific instructions and encouragement for subsequent action to offload the foot, increasing patient engagement and benefit.

Several limitations apply. First, while ulcer recurrence incidence for enhanced therapy was as estimated, for usual care it was lower than estimated in the sample size calculation. This reduces the effect size and the statistical power in finding a potentially present effect of the intervention. Second, with handheld thermometry, participants may

not have measured exactly at the predefined sites. Third, many different options for outcome sites could be considered for analysis, which affected the interpretation of results, as our analyses showed. While we lacked evidence to support choosing sites adjacent to the measurement site as the primary outcome sites, we considered that choosing only the measurement site would limit validity, as many ulcers may occur elsewhere, as would, for the same reason, choosing any foot site. Handheld devices with automated/semiautomated measurement reporting increase efficiency and would allow assessing more locations.³⁵ Platform systems also increase usability and foot coverage and can automatically report measurement data, but are limited to measuring only the surface that is in contact with the platform (ie, mostly only part of the plantar foot surface).^{14 26} Temperature sensors in socks can overcome this limitation, but are also confined to a limited number of measurement sites.³⁶ Any choice made regarding outcome site and measurement method is to a certain extent flawed, and therefore reporting for different options for outcome sites is important. Fourth, we subjectively obtained adherence to activity reduction when a hotspot was found, based on self-report. Finally, given the effect on the study findings of this adherence, our instruction to participants at baseline and repetition thereof in text messages twice a week during follow-up may have been too complex, infrequent, or ineffective.³ Data from a recent trial confirm that text and voice reminders in using thermometry do not affect adherence and outcome.³⁷ This may be inherent to this treatment approach and addresses an important barrier to implementation.

In conclusion, at-home daily foot temperature monitoring in addition to usual care does not significantly reduce incidence of foot ulcer recurrence at or adjacent to measurement sites compared with usual care alone in people with diabetes, peripheral neuropathy, and a foot ulcer history or Charcot foot. Being adherent to monitoring foot temperature does not mitigate this effect, but when participants reduce their activity when a hotspot is identified, the intervention is effective over usual care. Also, when only ulcers at the previous ulcer site or ulcers at any foot site (including non-measurement sites) are considered, the intervention is effective over usual care. Thus, the effect of at-home foot temperature monitoring in preventing ulcer recurrence is not as straightforward as previously found and may be limited to those adherent to change in behavior when guided by temperature measurement, to specific foot sites that should be targeted, or as method to increase awareness for the foot. And although the intervention has potential, the findings highlight that it is not a solution on its own and a multimodal treatment approach is required to substantially and continuously reduce risk of ulcer recurrence in people with diabetes.

Correction notice This article has been corrected since it was published. The layout of Table 2 has been corrected for better understanding.

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Contributors SAB conceived and designed the DIATEMP trial, obtained ethical approval, and registered the trial. FN contributed to trial design. SAB, WBadS, and JGvB drafted or edited the trial protocols. SAB, WBadS, JJvN, and TEB-W collected data for the trial. TEB-W and JGvB provided clinical advice and had clinical responsibility for patient-related matters. WBadS performed the statistical analyses. SAB and JJvN verified the underlying data. SAB wrote the manuscript and all other authors read, commented, and edited the manuscript for intellectual content. All authors approved the final version of the manuscript. SAB is the guarantor of this work and accepts full responsibility for the work and/or the conduct of the study, had access to the data, and controlled the decision to publish.

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