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http://sure.sunderland.ac.uk/policies.html or alternatively contact sure@sunderland.ac.uk. **TITLE:** Interrupting prolonged sitting with frequent short bouts of light-intensity activity in people with type 1 diabetes improves glycaemic control without increasing hypoglycaemia: The SIT-LESS randomized controlled trial

AUTHORS: *Matthew D. Campbell PhD^{1,2,3}, Anwar M. Alobaid MSc^{4,5}, Mark Hopkins PhD⁴, Paddy C. Dempsey PhD^{3,6,7,8}, Sam M. Pearson MD², Noppadol Kietsiriroje MD^{2,9}, Rachel Churm PhD¹⁰, Ramzi A Ajjan MD²

AFFILIATIONS: ¹ John Dawson Drug Discovery and Development Institute, University of Sunderland, Sunderland UK; ² Leeds Institute of Cardiovascular and Metabolic Medicine, Faculty of Medicine, University of Leeds, Leeds, UK; ³ Institute of Metabolic Science, University of Cambridge, Cambridge Biomedical Campus, Cambridge, UK; ⁴ School of Food Science and Nutrition, Faculty of Environment, University of Leeds, Leeds, UK; ⁵ Ministry of Health, Farwaniya Hospital, Kuwait city, Kuwait; ⁶ Diabetes Research Centre, University of Leicester, Leicester General Hospital, Leicester, UK; ⁷ Baker Heart and Diabetes Institute, Melbourne, Australia; ⁸ Institute for Physical Activity and Nutrition (IPAN), School of Exercise and Nutrition Sciences, Deakin University, Geelong, Victoria, Australia; ⁹ Endocrinology and Metabolism Unit, Faculty of Medicine, Prince of Songkla University, Thailand, UK; ¹⁰ Applied Sports Technology, Exercise and Medicine (A-STEM) Research Centre, Faculty of Science and Engineering, Swansea University, Swansea, UK

CORRESPONDING AUTHOR:

Dr Matthew D. Campbell John Dawson Drug Discovery and Development Institute University of Sunderland, Sunderland UK Email: matthew.campbell@sunderland.ac.uk

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ABSTRACT

Aims: To examine the impact of interrupting prolonged sitting with frequent short bouts of light-intensity activity on glycaemic control in people with type 1 diabetes (T1D).

Materials and methods: 32 inactive adults with T1D (aged 27.9 \pm 4.7 years, 15 men, diabetes duration 16.0 \pm 6.9 years and HbA1c 8.4 \pm 1.4% [68 \pm 2.3 mmol/mol]) underwent two 7-hour experimental conditions in a randomised crossover fashion with >7-day washout consisting of: uninterrupted sitting (**SIT**), or, interrupted sitting with 3-minute bouts of self-paced walking at 30-minute intervals (**SIT-LESS**). Standardised mixed-macronutrient meals were administered 3.5-hours apart during each condition. Blinded continuous glucose monitoring (CGM) captured interstitial glucose responses during the 7-hour experimental period and for a further 48-hours under free-living conditions.

Results: SIT-LESS reduced total mean glucose (**SIT** 8.2 ± 2.6 vs. **SIT-LESS** 6.9 ± 1.7 mmol/L, P=0.001) and increased Time in Range (TIR; 3.9-10.0 mmol/L) by 13.7% (**SIT** 71.5±9.5 vs. **SIT-LESS** 85.1±7.1 %, P=0.002). Hyperglycaemia (>10.0 mmol/L) was reduced by 15.0% under **SIT-LESS** (**SIT** 24.2±10.8 vs. **SIT-LESS** 9.2±6.4 %, P=0.002), whereas hypoglycaemia exposure (<3.9 mmol/L) (**SIT** 4.6±3.0 vs. **SIT-LESS** 6.0±6.0 %, P=0.583) was comparable across conditions. **SIT-LESS** reduced glycaemic variability (CV%) by 7.8% across the observation window (P=0.021). These findings were consistent when assessing discrete time periods, with **SIT-LESS** improving experimental and free-living postprandial, whole-day, and night-time glycaemic outcomes (P<0.05).

Conclusions: Interrupting prolonged sitting with frequent short bouts of light-intensity activity improves acute postprandial and 48-hour glycaemia in adults with T1D. This pragmatic strategy is an efficacious approach to reducing sedentariness and increasing physical activity levels without increasing risk of hypoglycaemia in T1D.

Clinical Trial Registry Number: ISRCTN13641847

1 INTRODUCTION

2 Physical activity is a critical element of diabetes care and is universally recommended to all 3 individuals with diabetes¹. Recently, guidelines have evolved to stipulate that in addition to 4 traditional structured moderate-vigorous intensity physical activity, individuals should limit 5 prolonged periods of sitting by incorporating frequent episodes of low-intensity physical activity into the day². This recommendation is based upon data demonstrating a dose-dependent 6 7 relationship between sedentary behaviour and cardiometabolic morbidity, worsening glycaemic 8 management, and increased weight gain, irrespective of physical activity status^{3,4}. Further, emerging evidence demonstrates that interruption of prolonged sitting with frequent short bouts 9 of activity improves acute postprandial and whole-day glucose levels, with glycaemic 10 improvement continuing until the next morning⁵⁻⁹, resulting from enhanced contraction-induced 11 and/or energy deficit-induced insulin sensitivity³¹, and/or a greater reliance on insulin-12 independent contraction-mediated glucose disposal³². However, these data remain preliminary 13 14 and limited to individuals with, or at risk of developing, type 2 diabetes. 15 Within the context of type 1 diabetes (T1D), most individuals struggle to meet physical activity 16 guidelines¹⁰ and spend a greater proportion of time sedentary than people without T1D¹¹. For 17 example, a recent large cross-sectional survey of 18,028 adults with T1D, reported that ~60% 18 did not achieve recommended physical activity levels¹⁰ a finding which supports some¹²⁻¹⁴, but 19 not all previous studies¹⁵. 20

21

22 Many people with T1D report fear of hypoglycaemia and an inability to manage their diabetes as major barriers to becoming active and engaging in regular moderate-to-vigorous physical 23 activity participation¹⁴, yet, few mention this fear when asked about lower-intensity activities 24 such as walking¹⁶. although many individuals with T1D do little-to-no exercise, they are often 25 26 willing to increase participation in lower-intensity physical activity and are keen to learn how to reduce sedentary behaviours^{14,16,17}. However, little information is available for individuals 27 28 with T1D or for the healthcare professionals who support them with regards to strategies for reducing sedentariness and their potential impact on hypoglycaemia risk^{14,16,17}. 29

30

Should findings from recent research in individuals with type 2 diabetes translate to those with T1D, interrupting sitting with frequent, short, light-intensity activity breaks, may serve as a pragmatic strategy for enabling inactive T1D individuals to incorporate more physical activity into their everyday lives and improve glucose management. This may be particularly beneficial for those who are unable or unwilling to engage in structured moderate-vigorous physical activity and an important stepping-stone towards achieving physical activity recommendations. However, no research has investigated the impact of such a strategy on glucose control in people with T1D. Therefore, the aim of this study was to examine the acute postprandial and subsequent 48-hour free-living glucose responses to interrupting prolonged sitting with frequent, short bouts of light-intensity activity in inactive people with T1D.

41

42 METHODS

43 Study Design

44 This randomised crossover trial was undertaken at the University of Sunderland between May 45 2021 and December 2022. The study received ethical approval from the Health Research Authority (HRA; London - Surrey Research Ethics Committee; Ref 20/LO/0650) and was 46 prospectively registered (ISRCTN13641847). All patients who participated provided written 47 informed consent with study procedures complying with the Declaration of Helsinki. 48 49 Participants completed an initial medical screening visit and two laboratory-based experimental visits each of which were separated by a minimum of 7 days (Supplementary Figure 1). 50 51 Experimental conditions were randomly assigned using a computerized random number 52 generator (www.randomization.com) with study personnel and participants blinded to 53 experimental condition order up until commencement of the first experimental visit.

54

55 *Participants*

56 Patients with autoantibody confirmed T1D treated on a stable (>6 months) insulin regimen 57 consisting of continuous subcutaneous insulin infusion (CSII) or multiple daily injections 58 (MDIs) were recruited in-clinic and via university recruitment streams from the North-East 59 region of the United Kingdom. Patients were eligible for inclusion if aged between 18 and 60 60 years with a duration of diabetes >2 years on enrolment and classified as inactive as per international physical activity guidelines^{1,2}; specifically, this consisted of failing to achieve a 61 minimum of 150 minute of moderate-vigorous intensity physical activity per week. Pregnancy, 62 the presence of significant functional limitations, dietary intolerances, overt diabetes 63 complications, or hypoglycaemia unawareness – as determined by the Clarke method¹⁸ – were 64 65 exclusion criteria.

66

67 Pre-experimental procedures

68 After initial telephone screening, potentially eligible participants underwent medical screening 69 at our laboratory for assessment of pre-treatment clinical characteristics including medical 70 history, anthropometry, blood pressure, and self-reported physical activity status using a validated assessment tool¹⁹. During this visit eligible participants then underwent initial study 71 72 orientation and were fitted with a blinded continuous glucose monitoring (CGM) device 73 (FreeStyle Libre Pro iQ, Abbott, UK). Participants were provided with a food diary to record 74 diet and insulin regimen and were provided with a pedometer to which recorded total step count 75 during each 24-hour period of the 48-hours before and after the first experimental laboratory 76 visit; this information was then used to replicate diet, insulin administration and physical 77 activity levels during the second experimental period. During this time, participants were required to abstain from exercise, caffeine, and alcohol in the 48-hours prior to each 78 experimental condition. Prearranged, standardized text messaging and/or email prompts were 79 80 used to maximize participant compliance.

81

82 For standardisation of glycaemic control prior to each laboratory visit, a standardised mixed-83 macronutrient meal (Supplementary Table 1) was provided to participants to consume on the 84 evening before each experimental visit; following consumption of this meal, participants were 85 instructed to avoid further food intake including calorific beverages, except for extremes of glucose readings managed as appropriate with corrective insulin boluses for hyperglycaemia 86 87 and glucose supplementation for hypoglycaemia. The aim was to ensure fasting status upon arrival to each experimental visit as detailed below. On the morning of each experimental visit, 88 89 study personnel contacted participants to ensure fasting status and confirm glucose levels were 90 within the range 4-12 mmol/L. Experimental visits were re-arranged if participants experienced 91 one or more sustained (>90-minutes) hyperglycaemic or sustained (>30 minute) hypoglycaemic 92 episodes. To limit the potential impact of menses on glycaemic measures for menstruating 93 female participants, procedures were arranged to occur within two-consecutive weeks during 94 their follicular phase (self-reported).

95

96 *Experimental procedures*

97 A schematic of the experimental procedures is presented in Supplementary Figure 1.
98 Participants attended our temperature-controlled (21-23°C) laboratory on a morning (~08:00)
99 following an overnight fast. On both occasions participants consumed standardized mixed100 macronutrient breakfast and lunch meals at 3.5-hours apart with start time equivalent on both

101 experimental arms. Each meal sought to replicate a typical Western diet with an energy density 102 of ~855 kcal, and a macronutrient profile of ~42% energy from carbohydrate, ~16% energy 103 from protein, and ~42% energy from fat (Supplementary Table 1). The carbohydrate content of 104 each meal was individualized equating to 1g carbohydrate per kilogram body mass. Participants 105 were instructed to administer their usual prandial insulin bolus immediately prior to each meal, 106 the dose of which was calculated using an individuals' established insulin-to-carbohydrate ratio, 107 with dose, timing and site of injection replicated across visits. Water was consumed ad libitum 108 during the first visit with the volume recorded and replicated during visit two; standardized 109 (within subject) lavatory visits were incorporated into the protocol to minimize unscheduled physical activity; however additional lavatory visits were permitted if needed. On one arm 110 111 (SIT), participants remained at rest and seated in a reclining chair for the duration of the visit. On a second arm (SIT-LESS) study procedures were replicated but sitting was interrupted by 112 performing 3-minute bouts of self-paced light-intensity walking at 30-minute intervals, 113 114 commencing 60-minutes after each meal; this equated to a total of 36-minutes of physical 115 activity across the 7-hour period. During each laboratory visit, participants had access to 116 television, books, and internet, and were supervised consistently by study personnel to ensure 117 resting periods were maintained. At 3.5-hours post-lunch, participants were discharged from 118 the laboratory with further free-living glycaemic assessment captured remotely via CGM for a further 48-hours. To minimize potential confounding of food intake, participants were provided 119 120 with an evening and breakfast meal to consume in sequence, replicating eating times within each study arm (Supplementary Figure 1). Any additional nutritional intake during the 121 122 subsequent 48-hour observation window was recorded on visit one, and subsequently replicated 123 on visit two. All meals provided to the participants consisted of commercially available foods 124 with standardized heating and preparation instructions. During the >7-day washout between 125 experimental conditions, participants resumed their habitual diet and physical activity patterns, 126 excluding the 48-hours pre-experimental period prior to the next experimental visit.

127

128 Continuous Glucose Monitoring (CGM)

Blinded CGM was used to capture interstitial glucose concentrations with sensor insertion a minimum of 72-hours prior to each data capture window to minimize artifacts during initialization. Sensors were inserted into the subcutaneous tissue on the back of the upper arm with insertion site marked with indelible ink to replicate senor insertion site during sensor replacement; existing CGM users continued to use their CGM as normal but were provided with a study-prescribed CGM to ensure consistency in CGM data capture. Data were 135 retrospectively downloaded and analyzed using manufacturer software (FreeStyle Libre software version 3.12; https://www.libreview.com) with the criterion of >80% data capture 136 137 within each 24-hour period across each experimental observation window (~5 days on each study arm) with no more than two consecutive hours of missing data during each 24-hour period 138 139 to be considered valid²⁰. From downloaded data, mean glucose, percentage of time in range (TIR: 3.9-10.0mmol/L), time above range (TAR:>10.0 mmol/L and >13.9 mmol/L), and time 140 141 below range (TBR: <3.9 mmol/L and <3.0 mmol/L), and glycaemic variability (CV%) were calculated as per international guidelines for the use of CGM in clinical trials ^{20,21}. 142

143

144 Data analysis

The primary outcome was 48-hour glycaemic control as assessed by mean glucose. We 145 estimated that 32 paired observations would be required to achieve 95% power to detect a 146 1.6mmol/L between group difference in mean glucose with an SD of 1.5mmol/L (moderate 147 effect size; Cohen d = 0.64) in the primary outcome variable. Our post-hoc power assessment 148 149 confirmed that our sample size was sufficient to achieve a minimum statistical power of 80% across our secondary outcomes (TIR, TAB, TBR, glycaemic variability). Across both 150 151 conditions, a total of 26,368 individual CGM derived glucose readings over a combined total 152 of 10 days were analysed, with missing data accounting for <1% (211 of 26,368). CGM data were summarised into three periods: (1) 48-hour pre-experimental phase, (2) experimental 153 154 phase, and (3) 48-hour post-experimental free-living phase. The 48-hour post-experimental free-living phase was further summarised into free-living day time periods (awake time: 08:00-155 156 23:00), and night-time periods (sleep time: 23:00-08:00).

157

158 We employed a series of generalized linear mixed models with random intercepts and 159 Bonferroni-corrected *post-hoc* pairwise comparisons to evaluate the differential effects of SIT 160 versus SITLESS on acute postprandial and 48-hour mean glucose, TIR, TBR, and TAR, as well as glycaemic variability (CV%). Linear regression analyses were utilised to examine 161 potential relationships between pre-treatment clinical characteristics (age, sex, BMI, HbA1c, 162 residual C-peptide, diabetes duration, and treatment regimen (CSII vs. MDI) and the magnitude 163 164 of treatment response across CGM metrics. Dietary intake, insulin administration, and physical activity (total step count) were summarised for each 24-hour period within the 48-hour post-165 intervention period and assessed for conditional differences over time using repeated measures 166 167 ANOVA. To assess mealtime glucose exposure, we calculated net incremental area under the curve (net iAUC) as previously reported²². Statistical analyses were performed using SPSS 168

- 169 software (IBM Statistics, version 28), with statistical significance accepted at a threshold of
- 170 $P \le 0.05$ and residuals examined for serial correlation, heteroscedascity, and normality. Data are
- 171 presented as mean±SD unless stated otherwise.
- 172

173 *Role of the funding source*

- 174 This study was funded by Diabetes UK (project grant: 20/0006154). The funder was not
- involved in the design of the study, data collection, analysis, or interpretation.
- 176

177 **RESULTS**

Thirty-two participants with T1D (age 27.9±4.7 years, 15/17 males/females, body mass index 178 179 (BMI) 26.5 ± 3.5 kg/m², diabetes duration 16.0 ± 6.9 years, HbA1c $8.4\pm1.4\%$ [68 ± 2 mmol/mol], CSII:MDI n=15:17) were randomised and completed both experimental conditions 180 181 (Supplementary Figure 2). Patients displayed similar glycaemic control across the 48-hours preceding each laboratory visit (Table 1), with similar mean glucose (SIT 7.7±1.1 vs. 182 SITLESS 7.5±2.1 mmol/L; P=0.631), and TIR (3.9-10.0 mmol/L; SIT 79.1±12.5 vs. 183 SITLESS 81.1±19.9 %; P=0.561). Exposure to hyperglycaemia and hypoglycaemia were also 184 comparable across conditions (P>0.01; Supplementary Table 1). Two patients re-arranged their 185 186 visits due to hypoglycaemia. Dietary intake, insulin regimen, and physical activity levels were 187 also similar across conditions (P>0.05).

188

Glucose concentrations at experimental start time were comparable between conditions (SIT 189 190 7.3±1.5 vs. SIT-LESS 7.2±1.8 mmol/L, P=0.774; Figure 1). During the laboratory phase, SIT-191 LESS attenuated postprandial glucose responses following administration of the breakfast (net 192 iAUC: SIT 1690±597 vs. SIT-LESS 1329±420 mmol/L.min P<0.001) and lunch (net iAUC: 193 SIT 1754±735 vs. SIT-LESS 1557±558 mmol/L.min P=0.001) test meals, resulting in lower 194 mean glucose (SIT 8.5±2.0 vs. SIT-LESS 7.1±1.8 mmol/L, P=0.008; Figure 1 and Table 1) and increased TIR by 17% (3.9-10.0 mmol/L; SIT 71.6±19.3 vs. SIT-LESS 84.6±14.8 %, 195 P=0.004; Table 1) as a consequence of reduced hyperglycaemia (TAR <10.1mmol/L: SIT 196 26.5±27.5 vs. SIT-LESS 8.6±18.3 %, P=0.005; Table 1); exposure to hypoglycaemia remained 197 198 comparable across conditions, irrespective of pre-treatment HbA1c, with similar TBR (<3.9mmol/L: SIT 2.7±8.4 vs. SIT-LESS 3.34±10.2 %, P=0.795; Table 1) and total number of 199 200 hypoglycaemic episodes at a threshold of 3.9 mmol/L (SIT 4 vs. SIT-LESS 5). 201

202

*** INSERT FIGURE 1 ***

203

204 The glycaemic lowering impact of SIT-LESS continued into the free-living period (Figure 1 205 and Table 1), with lower subsequent 48-hour mean glucose under SIT-LESS (SIT 8.1±1.3 vs. 206 SIT-LESS 6.9±1.5 mmol/L, P=0.001) and increased TIR by 13.0% (SIT 71.6±19.3 vs. SIT-207 LESS 84.6±14.8 %, P=0.004). TAR (>10.0 mmol/L) was reduced by 14.4% under SIT-LESS (SIT 23.8±18.6 vs. SIT-LESS 9.4±11.6 %, P=0.001), with TBR (<3.9 mmol/L) comparable 208 209 across conditions (SIT 4.6±5.0 vs. SIT-LESS 6.0±9.9 %, P=0.529). SIT-LESS reduced 48hour glycaemic variability (CV%) by 7.2% (P=0.035). These findings were consistent when 210 211 assessing discrete time periods with SIT-LESS improving postprandial, whole-day, and nighttime TIR (P<0.05; Figure 1 and Table 1). Dietary intake, insulin administration, and objectively 212 213 assessed physical activity levels were similar across conditions during the subsequent 48-hour 214 free-living period (P<0.05; Supplementary Table 2).

215

A significant HbA1c-by-condition interaction effect (P=0.007, F=8.635, η^2 =0.249, β =-0.801), and BMI-by-condition interaction effect (P=0.030, F=5.293, η^2 =0.169, β =-0.773) were observed for the magnitude of change between **SIT** and **SIT-LESS** in mean glucose. Higher pre-treatment HbA1c and BMI were associated with greater improvements across mean glucose, TIR, TAR, and TBR, but not glycaemic variability (Figure 2; Table 2). Age, sex, diabetes duration, residual C-peptide, and treatment regimen (CSII vs. MDI), did not significantly mediate any of the responses observed (P>0.05).

- 223
- 224

*** INSERT FIGURE 2 ***

225

226 **DISCUSSION**

227 This study is the first to evaluate the impact of interrupting prolonged sitting with frequent 228 short bouts of light-intensity activity on glucose control in people with T1D. This intervention 229 improved acute postprandial glucose control, reducing mean glucose concentrations, 230 improving TIR whilst reducing glycaemic variability without increasing exposure to 231 hypoglycaemia. Glycaemic improvement was sustained for at least 48-hours under free-living 232 conditions. Overall, these findings build on previous experimental work in people with or at risk of type 2 diabetes, and support the extension of current physical activity guidelines² to 233 234 individuals with T1D, specifically regarding the interruption of prolonged sitting with frequent, 235 short-duration, light-intensity activity breaks.

236

In people with diabetes, prolonged uninterrupted sitting is associated with worsening glucose 237 control and increased weight^{4,23} which collectively and independently predict both macro- and 238 microvascular complications^{24,25}. In the present study, we show that simply interrupting 239 240 prolonged sitting with regular light-intensity activity breaks results in a net glucose lowering effect of ~1.3mmol/L, with the greatest level of improvements in those with higher pre-241 242 treatment HbA1c and BMI. This clinically relevant margin, which if maintained over the longterm, has previously been shown to result in a reduction of HbA1c of $\sim 2\%^{26}$, translating to a 243 38% reduced risk of a macrovascular event, a 40% reduced risk of a microvascular event, and 244 a 38% reduced risk of premature mortality at a HbA1c threshold of 7% or higher²⁷; this is 245 substantial given recent data indicating that fewer than 30% of people with T1D achieve the 246 247 HbA1c treatment target of $< 7.5\%^{28}$.

248

Importantly, glucose lowering was achieved without increasing the risk of hypoglycaemia. We, 249 250 and others, have previously shown that moderate-vigorous physical activity predisposes to an 251 increased risk of hypoglycaemia during, immediately following, and late after moderatevigorous intensity exercise¹, and, that fear of exercise-induced hypoglycaemia is a major barrier 252 to regular participation in physical activity¹⁴. Whereas exercise is often viewed as daunting and 253 254 unachievable by many patients, translation of our data into clinical practice and patient 255 education may help to reduce fear of hypoglycaemia surrounding physical activity and enable 256 better glycaemic control when adopting lower-intensity activities. In addition, it is likely that 257 the adoption of our strategy to target sedentary time with short-duration light-intensity activity 258 breaks may serve as a logical starting point for inactive individuals with T1D to develop and 259 build upon achievable and positive behavioural routines that increase overall physical activity 260 levels.

261

The assessment of acute postprandial glucose control provides novel insightful data. We 262 observed a ~17% improvement in TIR under SIT-LESS, resulting almost exclusively from a 263 264 reduction in hyperglycaemia. Moreover, 75% of patients under SIT-LESS achieved TIR >80% and 56% achieved TIR =100% during their laboratory stay, compared to 38% and 6% under 265 266 SIT, respectively. During this time, glycaemic variability was reduced by 6% with all patients achieving the target CV% of $<36\%^{29}$ whilst concurrently avoiding increased exposure to 267 hypoglycaemia. Further, this effect persisted over the course of the subsequent 48-hour free-268 living observation window with an improvement in daytime TIR of ~12%, with 66% of patients 269

270 under SIT-LESS achieving TIR >80% - double that achieved under SIT. Given that no 271 differences were observed in dietary intake, insulin administration, or objectively assessed 272 physical activity levels during this period, it is likely that persistence in glycaemic improvement under SIT is due to the residual effect from the interrupted sitting intervention rather than 273 274 secondary to a change in behaviour. As such, our data demonstrate that the majority of patients adopting our strategy are able to achieve and exceed current mealtime glycaemic targets²⁹. This 275 276 a major finding given the inherent complexity and difficulty associated with optimising 277 postprandial glucose management in T1D and that controlling postprandial glucose excursions 278 is a key component of achieving recommended HbA1c levels and minimising disease burden. 279 In reality, many patients are exposed to increased glycaemic variability and hypoglycaemia 280 during mealtimes, both of which are significant sources of frustration for patients and factors that increase the risk of cardiovascular events and premature mortality independent of HbA1c³⁰. 281 282

A remarkable finding of the present study was that the magnitude of glycaemic improvement 283 284 across our chosen CGM metrics (mean glucose, TIR, glycaemic variability) persisted beyond 285 our controlled experimental observation window for up to a further 48-hours under free-living 286 conditions. Importantly, time spent in nocturnal hyperglycaemia was on average 16% lower 287 under SIT-LESS with minimal exposure to hypoglycaemia. Whereas our data highlight the 288 detrimental and persistent effects of high levels of prolonged sitting in T1D, they also clearly 289 demonstrate the glycaemic benefits of interrupting prolonged sitting and offer a strategy for 290 incorporating more physical activity throughout the day whilst avoiding increased exposure to 291 potentially dangerous hypoglycaemia. It remains unknown however, whether adopting a SIT-292 LESS protocol on consecutive days, or on multiple days per week results in further glucose 293 lowering. Future work should assess the impact and safety of sustained adoption of SIT-LESS 294 to establish whether combining activity days has continued or increased glucose lowering 295 power.

296

Within our study, we also examined the potential impact of pre-treatment clinical characteristics on the magnitude of treatment response. Our data show that baseline HbA1c and BMI status are important clinical characteristics that strongly associate with the magnitude of glucose lowering, with patients presenting with poorer glucose control and increased BMI demonstrating, on average, the largest degree of glucose lowering. The measures employed within this study do not enable an exploration into the putative mechanisms underpinning the improvements in glycaemic control observed under **SIT-LESS**, nor the interaction between 304 HbA1c and BMI with treatment response. However, the standardisation of insulin 305 administration and dietary intake across conditions, is suggestive of enhanced contractioninduced and/or energy deficit-induced insulin sensitivity³¹, and/or a greater reliance on insulin-306 independent contraction-mediated glucose disposal³². As such, interrupting sitting may present 307 308 an opportunity not only to tackle suboptimal glucose control, but also increase insulin 309 sensitivity in those presenting with insulin resistance. Overweight, obesity, and insulin resistance have recently been shown to be highly prevalent within the T1D population and to 310 be strongly associated with the risk of micro- and macrovascular complications independent of 311 HbA1c²⁴. Therefore, future studies are warranted that explore the longer-term impacts of 312 interrupted sitting on insulin resistance in T1D. Furthermore, it would be beneficial to explore 313 314 whether the additive effects of exercise and diet-induced energy deficit on glycaemic improvement extend to physical activities at the lowest-end of the physical activity continuum, 315 316 in order to optimise lifestyle change prescription.

317

318 Strengths of this study include: the rigorous well-controlled randomised crossover study design 319 allowing for within and between participant comparisons, increasing internal validity and 320 reliability of the data collected, and permitting a smaller sample size whilst ensuring adequate 321 statistical power. We standardised condition run-in periods with strict but pragmatic assessment 322 and replication of confounding variables including diet, physical activity, fasting metabolic and 323 glycaemic status, and experimental start time; comprehensive and blinded glucose profiling 324 under controlled and extended free-living conditions with negligible data loss (<0.1%); full 325 retention of study participants that reflect a relatively broad and representative demographic; 326 and, the simple and practical nature of the intervention which enables widespread promotion 327 and adoption. Key study limitations are that this is a single centre study with a conservative 328 sample size which prevented subgroup analyses. Further, we assessed physical activity volume 329 using total step count and were unable to assess other dimensions of physical activity and therefore cannot rule out the possibility that undetected changes in physical activity could have 330 331 impacted glucose outcomes during the free-living period. Future research is needed to 332 determine whether such an intervention can be optimised (frequency, intensity, and duration of 333 walking breaks), and tailored specifically to accommodate patients with mobility issues, functional limitations, the presence of overt diabetes complications and other comorbidities, as 334 335 well as those with insulin resistance. In addition, future studies should establish if such a 336 strategy can be maintained by patients in free-living environments over the long-term and 337 whether this translates to reduced risk of long-term complications and improved quality of life.

338

339 CONCLUSION

340 We show, for the first time, that interrupting prolonged sitting with frequent short bouts of light-341 intensity activity improves acute postprandial glucose control resulting in glucose lowering, 342 improved TIR and reduced glycaemic variability without increased risk of hypoglycaemia, with sustained improvement for up to a further 48-hours. Although longer-term efficacy needs to be 343 344 established, our findings provide the first experimental evidence for the value of frequent low intensity physical activity for improving glycaemia in individuals with T1D. This simple and 345 346 acceptable approach may help to enable inactive individuals to incorporate more physical 347 activity into the day and improve diabetes management. Interruption of sitting with light 348 activities could be particularly useful for those who are unable or unwilling to engage in structured exercise, and this approach can be seen as an important "stepping-stone" toward 349 350 regular participation in structured moderate-vigorous physical activity or exercise. It should be 351 emphasised that, unlike moderate-vigorous exercise, the improvement in glycaemia with our 352 simple intervention did not result in increased hypoglycaemia and therefore we propose that healthcare professionals consider advising patients to regularly interrupt prolonged sitting. 353 354 Large scale studies are warranted to fully evaluate both the short- and long-term impact of this 355 simple intervention in the management of individuals with T1D.

AUTHORSHIP

MDC: conceptualization, funding acquisition, investigation, methodology, resources, supervision, visualization, writing - original draft, writing – review and editing; AMA: investigation, visualization, writing - review and editing; MH: investigation, visualization, writing - review and editing; PCD: conceptualization, funding acquisition, investigation, methodology, resources, supervision, visualization, writing – review and editing; SMP: investigation, visualization, writing - review and editing; RC: investigation, visualization, writing - review and editing; RAA: conceptualization, funding acquisition, investigation, methodology, resources, supervision, visualization, investigation, writing - review and editing; RAA: conceptualization, funding acquisition, investigation, methodology, resources, supervision, visualization, writing – review and editing; RAA: conceptualization, funding acquisition, investigation, methodology, resources, supervision, visualization, and accept responsibility to submit for publication. This study was funded by Diabetes UK (20/0006154). The funder was not involved in the design of the study, data collection, analysis, or interpretation.

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CONFLICT OF INTEREST STATEMENT

No interests to declare.

DATA SHARING

Deidentified participant data collected during the trial alongside the study protocol and statistical analysis plan will be made available beginning 3 months and ending 36 months following article publication for investigators whose proposed use of the data has been approved by an independent review committee identified for this purpose. Proposals may be submitted up to 36 months following article publication and should be directed to matthew.campbell@sunderland.ac.uk to gain access, data requestors will need to sign a data access agreement. After 36 months the data will be available in our university's data warehouse but without investigator support other than deposited metadata.

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	SIT	SIT-LESS	P Value
Pre-experimental phase (48-hour run-in period)			
Mean glucose (mmol/L)	7.7±1.1	7.5±2.1	0.631
Percent TIR 3.9-10.0 mmol/L (70-180 mg/dL)	79.1±12.5	81.1±19.9	0.561
Percent TAR >10.0 mmol/L (>180 mg/dL)	16.5±12.5	14.7±19.8	0.638
Percent TAR >13.9 mmol/L (>250 mg/dL)	2.7±4.1	4.9±13.1	0.293
Percent TBR <3.9 mmol/L (<70 mg/dL)	4.5±5.2	4.3±7.4	0.903
Percent TBR <3.0 mmol/L (<54 mg/dL)	0.6±1.4	4.0±7.0	0.326
Glycaemic variability (CV%)	31.4±10.6	28.7±9.7	0.104
Experimental phase response	<u>.</u>		
Mean glucose (mmol/L)	8.5±2.0	7.1±1.8	0.008**
Percent TIR 3.9-10.0 mmol/L (70-180 mg/dL)	70.9±27.4	88.0±19.9	0.007**
Percent TAR >10.0 mmol/L (>180 mg/dL)	26.5±27.5	8.6±18.3	0.004**
Percent TAR >13.9 mmol/L (>250 mg/dL)	6.9±14.3	1.7±6.5	0.072
Percent TBR <3.9 mmol/L (<70 mg/dL)	2.7±8.4	3.3±10.2	0.795
Percent TBR <3.0 mmol/L (<54 mg/dL)	0.7±3.7	0.2±1.2	0.536
Glycaemic variability (CV%)	24.4±13.0	18.1±9.2	0.013*
Free-living phase response		· · ·	·
Mean glucose (mmol/L)	8.1±1.3	6.9±1.5	<0.001***
Percent TIR 3.9-10.0 mmol/L (70-180 mg/dL)	71.6±19.3	84.6±14.8	0.004**
Percent TAR >10.0 mmol/L (>180 mg/dL)	23.8±18.6	9.6±11.6	<0.001***
Percent TAR >13.9 mmol/L (>250 mg/dL)	4.5±5.8	1.5±3.67	0.007**
Percent TBR <3.9 mmol/L (<70 mg/dL)	$4.6{\pm}5.0$	6.0±9.85	0.568
Percent TBR <3.0 mmol/L (<54 mg/dL)	1.3±2.3	1.8±4.5	0.529
Glycaemic variability (CV%)	31.7±12.4	24.5±11.9	0.035*

Mean glucose (mmol/L)	8.2±1.4	7.1±1.7	0.002**		
Percent TIR 3.9-10.0 mmol/L (70-180 mg/dL)	71.0±18.6	82.5±19.0	0.023*		
Percent TAR >10.0 mmol/L (>180 mg/dL)	24.3±18.8	11.0±15.7	0.003**		
Percent TAR >13.9 mmol/L (>250 mg/dL)	19.6±16.1	9.0±12.1	0.017*		
Percent TBR <3.9 mmol/L (<70 mg/dL)	4.7±6.1	11.0±10.9	0.478		
Percent TBR <3.0 mmol/L (<54 mg/dL)	1.5±3.6	2.2±4.9	0.536		
Glycaemic variability (CV%)	24.0±7.6	19.2±8.7	0.044*		
Combined free-living night-time periods					
Mean glucose (mmol/L)	8.0±1.5	6.7±1.4	0.003**		
Percent TIR 3.9-10.0 mmol/L (70-180 mg/dL)	71.6±23.3	86.6±14.3	0.003**		
Percent TAR >10.0 mmol/L (>180 mg/dL)	22.9±22.2	7.2±11.1	0.001**		
Percent TAR >13.9 mmol/L (>250 mg/dL)	3.5±6.3	0.5±1.6	0.007**		
Percent TBR <3.9 mmol/L (<70 mg/dL)	5.4±7.2	6.8±11.0	0.606		
Percent TBR <3.0 mmol/L (<54 mg/dL)	0.9±1.7	3.5±5.9	0.159		
Glycaemic variability (CV%)	44.5±18.3	39.4±22.3	0.374		

Note: Day time and nighttime periods calculated as the combined mean for each respective period. TIR = time in range; TAB = time above range; TBR = time below range; CV = coefficient of variation. *Indicates a conditional difference at P<0.05; **Indicates a statistically significant conditional difference at P<0.01; ***Indicates a statistically

Table 2. Association between pre-treatment clinical characteristics and treatment response

	HbA1c	BMI			
Experimental and free-living phase response					
Mean change in mean glucose (mmol/L)	$\beta = -0.801$ (-1.39 to -0.78); P < 0.001***	$\beta = -0.773$ (-0.53 to -0.283); P =<0.001***			
Mean change in percent TIR 3.9-10.0 mmol/L (70-180 mg/dL)	$\beta = 0.462$ (2.18 to 13.14); P = 0.008**	$\beta = 0.481$ (0.97 to 5.22); P = 0.005**			
Mean change in percent TAB >10.0 mmol/L (>180 mg/dL)	$\beta = -0.686$ (-14.72 to -6.37); P < 0.001***	β = -0.740 (-5.94 to -2.93); P < 0.001***			
Mean change in percent TBR <3.9 mmol/L (<70 mg/dL)	$\beta = 0.343$ (-0.064 to -5.84); P = 0.049*	$\beta = -0.404$ (-0.208 to -2.45); P = 0.022*			
Mean change in glycaemic variability (CV%)	$\beta = 0.052$ (-4.241 to 5.624); P = 0.777	$\beta = 0.108$ (-1.36 to 2.47); P =0.558			

Note: Data presented as unstandardized β -coefficients (95% confidence interval); BMI = body mass index; *Indicates a statistically significant association at P<0.05; **Indicates a statistically significant association at P<0.01; ***Indicates a statistically significant association at P<0.01.

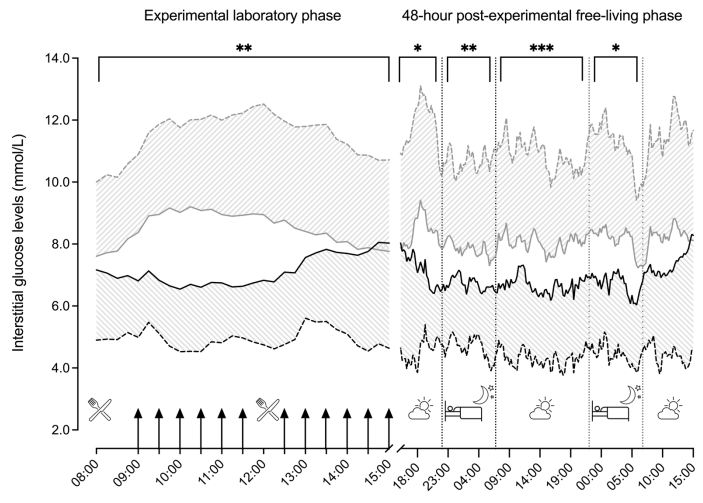
FIGURE LEGENDS

Figure 1. Glycaemic responses to interrupting sitting with frequent short bouts of light-intensity activity. Grey trace = **SIT** (uninterrupted sitting); Black trace = **SIT-LESS** (interrupted sitting with 3-minute bouts of self-paced light-intensity walking at 30-minute intervals as indicated by black vertical arrows). *Indicates a statistically significant conditional difference during each respective time-period at P<0.01; ***Indicates a statistically significant conditional difference during each respective time-period at P<0.01; ***Indicates a statistically significant conditional difference during each respective time-period at P<0.01; ***Indicates a statistically significant conditional difference during each respective time-period at P<0.001; Vertical dashed line breaks indicate nocturnal periods. Data presented as mean (solid trace) with SD (dashed trace); To improve clarity, +SD is presented for SIT, and -SD is presented for SIT-LESS.

Figure 2. Individualized magnitude of change in treatment response between **SIT** and **SIT-LESS** across: A: mean glucose, B: mean TIR, C: mean TAR, D: mean TBR, E: mean GV. Circles = pre-treatment HbA1c \geq 7.5 mmol/mol; Triangles = pre-treatment HbA1c <7.5 mmol/mol. White data points = normal weight (<25 kg/m²); Grey data points = overweight (25-29.9 kg/m²); Black data points = obese (>29.9 kg/m²). Numbers represent individually annotated participant data points. Treatment response calculated by subtracting mean **SIT-LESS** responses from mean **SIT** responses. **SIT** = uninterrupted sitting; **SIT-LESS** = interrupted sitting with 3-minute bouts of self-paced light-intensity walking at 30-minute intervals. TIR = time in range (3.9 – 10.0 mmol/L), TAR = time above range (>10 mmol/L), TBR = time below range (<3.9 mmol/L), GV = glycaemic variability (CV%). *Indicates a statistically significant association with magnitude of treatment response at P<0.05. **Indicates a statistically significant association with magnitude of treatment response at P<0.01; ***Indicates a statistically significant association with magnitude of treatment response at P<0.01;

FIGRURES

Figure 1.



Time (24-hour clock)

Figure 2.

