1 Abstract

2 **Context:** Current research suggests that dark cocoa may reduce fatigue; however, the effect 3 on fatigue in people with MS (pwMS) has never been established. The objective of this 4 feasibility study was to explore the acute effect of high flavonoid cocoa on measures of fatigue and glycaemic response. Methods: This was a randomised crossover participant blind 5 6 exploratory study in 12 participants (2 male and 10 female) with MS-related fatigue (>4 on the Fatigue Severity Scale; FSS). After fasting overnight, participants consumed the high 7 flavonoid cocoa drink (350 mg gallic acid equivalents {GAE}/g) or a low flavonoid cocoa 8 9 control (120 mg GAE/g), consuming the alternative drink on the next visit. Fatigue was selfreported on a 100mm visual analogue scale at 30-minute time intervals for 2 hours post cocoa 10 11 consumption and every 2 hours for the rest of the day. Fatigability was monitored using a 6 12 minute walk test (6MWT) at the end of the visit (2 hrs), and activity monitors worn for 24 hours commencing at 12noon on the day of testing. The feasibility of performing the trial 13 including outcome measures was documented. Results: A moderate effect was found in self-14 15 reported fatigue throughout the day in favour of the high flavonoid group (Cohen's d 0.32, 95% non-central t CI -0.57-1.20). Fatigability measures did not change. Participants 16 17 consumed and enjoyed the cocoa, all participants completed the study and outcome measures were accepted. Conclusion: The results of this study support further trials to investigate the 18 feasibility and efficacy of pure cocoa as a dietary supplement for fatigue in pwMS. 19 20

21 Abbreviations

BG, blood glucose; GR, glycaemic response; VAS, visual analogue scale; GAE, gallic acid
equivalents; pwMS, people with Multiple Sclerosis; FFQ, food frequency questionnaire; BI,
Barthels Index; FSS, fatigue severity scale

25 Introduction

Fatigue is one of the most debilitating symptoms in people with Multiple Sclerosis (pwMS),
greatly affecting quality of life (Tabrizi and Radfar, 2015). The exact cause of fatigue in MS
is unknown, however various mechanisms may influence fatigue severity.

Foods rich in flavonoids may show potential for reducing fatigue, through several 29 proposed mechanisms. There is currently available evidence that suggests oxidative stress 30 may contribute to the pathology in MS, which in turn may be improved or inhibited by the 31 antioxidant properties in flavonoids (van Horssen et al., 2008). In addition, it has been 32 33 suggested that the functional properties of flavonoids allow for penetration through the bloodbrain barrier, potentially leading to improved neurosignaling, as well as rehabilitation of 34 neuronal function (Solanki et al., 2015). A pathological inflammatory response may be 35 36 responsible for the fatigue experienced in MS, for example TNF-alpha levels have been found to be elevated in fatigued pwMS compared to those who were non fatigued (Braley and 37 Chervin, 2010). Luteolin, a naturally occurring flavonoid, has been found to benefit the 38 disease course of pwMS, for example by inhibiting activated peripheral blood leukocytes and 39 mast cells and mast cell dependent T cell activation (Theoharides, 2009). Katz, Doughty and 40 Ali (2011) suggest that cocoa may be beneficial towards MS remission as its flavonoid 41 content may promote blood flow to the brain, and may therefore lead to additional nerve 42 repair, better metabolic clearance from the brain and greater oxygen availability. Therefore 43 44 foods containing flavonoids may be used in conjunction with other disease modifying treatments (DMTs) in pwMS to reduce relapses and improve the severity of the symptoms 45 experienced. 46

47 Cocoa is rich in flavonoids and is a popular and easily accessible product. A recent
48 systematic review and meta-analysis investigated 42 randomized control trials, and found
49 cocoa to be significantly beneficial for vascular endothelial function and inflammation

(Hooper et al., 2012). Cocoa has been shown to improve fatigue in people with Chronic
Fatigue Syndrome (CFS). Sathyapalan et al. (2010) conducted a double blinded, randomised,
pilot crossover study, daily providing participants with 45g of high flavonoid chocolate. After
eight weeks participants reported significant reductions in fatigue and disability. Poor sleep
quality has previously been shown to be significantly correlated with fatigue in MS (Attarian
et al., 2004). Flavonoids have also been shown to improve sleep quality and therefore may
reduce daytime fatigue in those with MS (Ngan & Cunduit, 2011).

Flavonoid rich foods have also been shown to influence postprandial blood glucose levels (Coe et al. 2013). Glucose tolerance may be altered in pwMS (Mahler et al., 2012; Wens et al. 2013; White et al., 2006) and an association has been found between the availability of glucose to the brain and perceived fatigue (Roelcke et al., 1997). This has therefore raised the question as to whether improved glucose tolerance may reduce the fatigue experienced in those with the disease. However to date there has been no exploration of the response in this group.

The current randomised crossover exposure response participant blind exploratory study will assess the effect of high flavonoid cocoa versus low flavonoid cocoa on fatigue and fatigability as measured by mobility in pwMS. Glycaemic response (GR) after the consumption of the drink, was also measured.

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70 Method

This was a randomised crossover participant blind exploratory study in 12 participants (aged 71 54 ± 10.56 years, 2 male and 10 female) with MS-related fatigue. Participants were expected 72 73 to attend two test visits at Oxford Brookes University and in a randomised order (determined electronically by a random number table) consumed either the low flavonoid control or the 74 high flavonoid cocoa on different days, with at least three days between test visits (Figure 1). 75 The present study was approved by the Oxford Brookes University Research Ethics 76 Committee: UREC Registration No: 150938. All procedures were carried out accordingly to 77 78 Declaration of Helsinki guidelines and policies, and retained data was managed accordingly to the Oxford Brookes University's policy on Academic Integrity. 79 80

81 **Procedure**

PwMS were recruited from local support groups throughout the Thames Valley and via advertisements posted at Oxford Brookes University. After expressing interest in the study, participants where provided with the study information sheet and were given a minimum of 24 hours to review the information and ask the researchers any questions regarding the trial. Once potential participants agreed to take part in the trial and after initial eligibility was checked over the phone, a combined screening and first test visit was arranged where signed consent was taken.

Participants were asked to keep a 24 hour food diary the day before each visit, and to repeat this diet before the next test day. In addition to reduce variability in testing, participants were also asked to avoid vigorous exercise, and to limit their alcohol and caffeine intake on the day prior to testing (≤ 2 units and ≤ 3 cups respectively). Additionally, participants were asked to fast overnight for 10 - 12 hours prior to visits, which began between 7-10 am, and on the first assessment visit a health questionnaire was administered 95 (asking about smoking habits, current or previous diseases, current medication or supplement96 intake, dietary habits).

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98 Screening and descriptive data

At the screening/ first test visit demographics were recorded including MS subtype, compliance with the fasting protocol, blood pressure (mmHg) was recorded and mean fasting blood glucose (BG) was measured. Participant independence in daily living was assessed using the Barthels Index (BI; Nicholl et al., 2004). Each test occasion lasted no longer than three hours, and participants were required to leave a minimum of 24 hours between test days.

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106 **Participants**

Participants were excluded if they reported any sudden changes in MS symptoms 107 within the last three months, had a change in their DMTs and/ or medications that could 108 109 influence fatigue in the past three weeks, had a metabolic disease or were presently on medication interfering with insulin or glucose metabolism, had been diagnosed with a 110 condition other than MS affecting the CNS, had an allergy or intolerance to ingredients used 111 during testing, experienced fatigue from any condition other than MS, were pregnant or 112 lactating, were clinically depressed, had a BMI outside 18.5 - 30 kg/m² (body composition 113 was confirmed using Tanita BC-418MA), had impaired glucose tolerance (7.8-11.1 mmol/L), 114 or had a fatigue severity score less than 4 on the Fatigue Severity Scale (FSS; Krupp et al. 115 1989). This scale asks nine questions about various aspects of perceived fatigue, 1 = not116 fatigued at all and 7 = very fatigued. 117

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119 Acute response to flavonoid drink/ intervention

120 Participants were randomly administrated a high polyphenol test or low polyphenol control cocoa drink (Table 1), to consume within a maximum time of 15 minutes. Drinks were 121 matched as closely as possible for available carbohydrate (avCHO) and energy content. Due 122 123 to the idea that pwMS are following the Overcoming Multiple Sclerosis (OMS) diet, which excludes diary from the diet, the drink was made with Alpro rice milk (Tesco, UK). The total 124 polyphenol content of the drinks had previously been established (Santos and Coe, 2016), 125 with the high flavonoid cocoa powder containing 350mg gallic acid equivalents (GAE)/g, 126 whilst the low flavonoid control powder had instead been established to contain 120mg 127 128 GAE/g.

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130 Fatigue VAS

Fatigue was recorded on a horizontal 100mm VAS every 30 minutes following drink consumption and throughout testing, categorizing 0mm as 'not at all fatigued' and 100mm as 'extremely fatigued' (Kos et al., 2006). Participants continued to record fatigue every two hours after testing was completed, until six hours after leaving the lab.

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136 Fatigability

Fatigability was monitored using a 6 minute walk test (6MWT) performed at the end of thevisit (2 hrs), and through activity monitoring.

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140 Activity monitoring

141 A GENEActiv (Geneactive, UK) was used to record physical activity for a 24 hour period.

142 Data was sampled at 100Hz at a +/-8g range at 3.9mg resolution and recorded from the non-

dominant wrist. Post measurement, data was epoched to 1 seconds samples and analysed in a

bespoke spreadsheet (Excell, Microsoft Office 2011, US) and expressed as percentages of

physical activity level (sedentary, light, moderate, vigorous) per hour, according to sample
frequency adjusted Single Vector Magnitude cut-offs described by Esliger et al. (2011).

148 6 Minute Walk Test

The 6MWT has previously been proved to be an accurate tool to establish physical fatigue in MS (Goldman, Marrie and Cohen, 2008). On each test occasion, the 6MWT was performed approximately 120 minutes after the test meal was consumed. Participants were instructed to walk back and forth along a 14m long corridor at a pace they deemed comfortable, rounding a cone at the end of each lap. During the walk, participants were at all times accompanied by appropriately trained personnel, and were informed they may stop and rest if needed. The walk was timed using a hand held stopwatch and distance walked was measured in metres.

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157 **GR**

GR was measured using the previously validated method of Wolever (2004) and redesigned 158 by Coe et al. (2013), and is in line with procedures recommended by the Food and 159 Agriculture Organization/World Health Organization. A total of eight blood measures were 160 taken using an automatic blood glucose analyser (Glucose 201+, Hemocue AB, Sweden). 161 Mean fasting BG was calculated using two 5µl finger-prick samples at -5 and 0 min, and 162 measurements were collected at 15, 30, 45, 60, 90 and 120 minutes after the cocoa 163 164 consumption. BG levels were compared to WHO guidelines to determine if any of the participants had fasting BG or postprandial BG values outside of the healthy range. 165

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167 Statistical Analysis

This trial was not designed to determine efficacy and therefore no formal sample size
calculation was under taken. Descriptive statistics were expressed, including demographic
characteristics.

Fatigue raw data, and GR-AUC data was calculated using a fixed effects model with 171 two treatments (control versus test) adjusted using the Tukey-Kramer and comparing time 172 and treatment interaction. GR VAS AUC was calculated geometrically using the trapezoidal 173 rule at each time point relative to baseline values (Wolever, 2004). For outcome data the 174 Linear Mixed Models (LMM) procedure of SAS 9.4 was used to determine the mean changes 175 176 in measures, as response variables, according to exposures (test and control) and three repeated measurements, using baseline as a covariate. Further and based on the differences of 177 LS (Marginal) means between two groups (test versus control; pairwise comparisons), 178 179 provided by LMM analysis, powers, effect sizes (Cohen's d) and their 95% non-central confidence limits were calculated. 180

For physical activity, a model with time as a repeated factor, a treatment factor with two levels (placebo and test) and 3rd factor with 3 levels (sedentary, light, moderate and vigorous) was considered. Day and night required introduction of one more factor with two levels (day and night). Activity data was aggregated over 6 hours (4 levels) for sedentary, light and moderate activity. Activity awake versus sleep data was aggregated for day and night (2 levels). A random effect model aggregated for total activities with sequence incorporated.

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190 **Results**

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A summary of descriptive data may be viewed in Table 2. Participants who had had 192 193 MS for a number of years presented with relapsing remitting, primary and secondary progressive subtypes. All were high functioning and independent in daily activities. As can 194 be seen in Table 2 four out of the 12 participants were on special diets, three of who were on 195 the OMS diet, and one on a gluten free diet. Seven people were taking Vitamin D 196 supplements. Two participants reported daily taking the antidepressant medication 197 198 Amitriptyline (10mg and 50mg) and two additional participants reported currently taking drugs for fatigue including Modafinil and Amandatine (200mg and 100mg). 199 200 No adverse events or side-effects were reported or observed during this study and all 201 participants finished the drink on all occasions. Participants were overall content with the taste and sensory properties of the drink, none reported a dislike of the drink. Participants 202 seemed to find it feasible to comply to consuming a relatively similar diet the day before each 203 204 test day. 205

206 Fatigue Visual Analogue Scales

A moderate effect was found in self-reported fatigue throughout the day in favour of the high flavonoid group post consumption (Cohen's d 0.32, 95% non-central t CI -0.57-1.20). Figure 2 indicates a trend for fatigue to be reduced in the hours after leaving the lab through until evening.

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212 Activity monitoring

There was more activity during the sleeping hours (10pm-9am) after the consumption of thelow flavonoid compared to the high flavonoid drink (Figure 3). Moderate physical activity

215	tended to be higher five hours post high flavonoid cocoa consumption compared to moderate
216	physical activity after the control (Cohen's d 0.47, 95% CI: -0.36-1.27).

218 6 Minute Walk Test

- All participants completed the 6MWT on both visits. There was a wide range in distance
- walked with an average of 273 ± 115 m, ranging from 120-475m (Figure 4). This was
- expected due to the varied levels of disability among the participants, of which three needed
- the assistance of an aid (cane or frame) to complete the walk.
- 223

224 Glycaemic Response

- All participants had normal fasting BG levels and 2 hour postprandial BG levels as defined
- by the WHO (diabetes if fasting BG of >7.0 mmol/l or 2 hour postprandial BG >11.1 mmol/l
- and impaired glucose tolerance if fasting BG >7.0 mmol/l and 2 hour postprandial BG >7.8
- and <11.1 mmol/l); however two participants were exclude from the analysis due to
- abnormally high BG levels at time points between 0-120 mins. There was a trend for the high
- flavonoid cocoa to slightly decrease GR in the early stages postprandial (Cohen's d 0.07,
- 231 95% t CI -94-0.81; Figure 5).

232 Discussion

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The results of this trial ultimately show that a single drink of flavonoid rich cocoa produced a modest effect size reduction in perceived fatigue ratings compared to a low flavonoid control drink. This is the first study in pwMS to explore flavonoid use as a treatment for fatigue. The positive effect on fatigue alongside good tolerance and no adverse effects support the need for a powered trial of the longer-term effects. The results of the study may help direct future studies to identify specific questions and employ more efficient designs.

240 A single dose of flavonoids produced a modest effect on fatigability as measured the 24 hour activity levels yet showed no impact on the 6MWT. Specifically, later in the day five 241 242 hours after cocoa consumption an effect was observed in activity levels. However, 243 performance on these mobility measures will have been affected by a number of other biopsychosocial factors and it may have been that the 6MWT was administered too early to 244 see an effect on activity levels. In the current study there was a trend for activity during the 245 246 day to be greater after the high flavonoid cocoa consumption, and activity during the sleeping hours was less. This observation is a positive observation as both poor sleep quality and 247 reduced physical activity are common in MS (Attarian et al., 2004) and a number of 248 participant's commented on poor sleep quality the night before testing. 249

Moreover, the fatigue VAS does not differentiate between primary fatigue which is a direct result of the disease, and secondary fatigue which is a result of other symptoms. High flavonoid cocoa has previously been shown to improve factors which may lead to secondary fatigue such as sleep quality, and it may be speculated a further frequent intake of high flavonoid cocoa may allow for the beneficial mechanisms to take effect, possibly reducing secondary fatigue while allowing for improved fatigue VAS accuracy (Bisson et al., 2008). Also from assessing the activity data, participants overall seemed to perform low levels of physical activity and spent a majority of their time in sedentary/ light activity. Therefore
although further studies would need to be performed, it appears that high flavonoid cocoa
may contribute to improving physical activity during the day and better sleep patterns at night
in those with MS.

This was a one-day study which may not have been long enough to produce a larger 261 effect. Sathyapalan et al. (2010) administered high flavonoid cocoa daily over a period of 262 eight weeks to find significant differences. A meta-analysis of Hooper et al. (2012) similarly 263 suggests positive changes to health may depend on the quantity of cocoa consumed. The dose 264 of polyphenols present in the high flavonoid drink in the current study may have been a 265 limitation as the drinks contained 350 mg of gallic acid equivalents (GAE)/g and 120 mg 266 GAE/g for the high flavonoid and low flavonoid beverages, respectively. In Field et al. 267 268 (2011), on results obtained for spatial memory and performance on aspects of choice reaction time task, high flavonoid cocoa containing 773 mg flavonoids was used, with significant 269 results found compared to the control (low flavonoid cocoa). 270

271 The underlying mechanisms were not fully explored in this acute response to a single dose trial; however, foods rich in flavonoids have been shown to stabilise postprandial BG 272 levels (Davies et al., 2012) and consequently reduce fatigue in healthy individuals (Micha et 273 al., 2011). In this study there was no effect of dark cocoa on BG, however there was a trend 274 for the high flavonoid cocoa to reduce the BG in the early stages after consumption and 275 276 stabilise levels later after consumption. Although all participants showed normal BG levels as defined by WHO, there were some abnormally high patterns in GR during the two hours after 277 the cocoa drink consumption. Therefore the GR in people with MS should be considered in 278 future trials when assessing nutrition and fatigue, and the effect of pure cocoa on fasting BG 279 would need further exploration. 280

Diet has recently been shown to be an important factor in influencing clinical outcomes in MS (Grossman & Wahls, 2016). Research has found that pwMS are not only willing to consider dietary approaches to manage their symptoms (Brenton & Goldman, 2016) yet many are already on various diets in order to do so (Schwarz et al., 2008). It was therefore hypothesised that this population would be more open to a dietary intervention. Indeed results show that four out of 12 participants were on special diets for their condition, and seven were taking some form of Vitamin D supplements.

Although not for managing depression, two participants reported daily intake of the 288 289 antidepressant drug Amitriptyline (50 and 10mg), while two additional participants reported currently taking drugs for fatigue including Modafinil and Amandatine (200mg and 100mg). 290 291 Amitriptyline has previously been shown to cause sleepiness, as well as drowsiness in 292 patients (Frost et al., 2011). Contrastively, Modafinil and Amandatine have previously been reported to decrease fatigue (Ashtari et al., 2009, Brown et al., 2010). Therefore these 293 medications may have increased variability and decreased ability to see a difference in 294 results. However, pwMS are on these medications and therefore the study is more reflective 295 of real world practice. 296

No participants in the trial vocalised issues with the outcome measures or with the 297 fasting protocol. From the accelerometer results, it seemed that participants wore the watch 298 for the entire testing period. Overall we propose that the flavonoid cocoa drink was well 299 300 tolerated and enjoyed by all participants and is both suitable for a trial of longer term use and feasible that this drink would be consumed as part of a daily diet. There were no adverse 301 events, and all participants completed the study. Participants were compliant with the study 302 protocol and approved of outcome measures. This, in combination with the moderate effect 303 size on reducing fatigue, shows promise for further studies into the role of flavonoid rich 304 305 cocoa on fatigue in MS.

307 Limitations

308 Due to the novel and exploratory nature of the study a pragmatic approach was used 309 determine sample size and no formal sample size calculation was under taken. However, post 310 hoc analysis found moderate effect sizes at measurement points where differences in fatigue 311 were greatest in the high flavonoid group.

In the current study, whilst the participants were, assessors were not blinded to the 312 intervention. However, the protocol was delivered in randomised format and according to 313 314 fixed standardised operating instruction sets and procedures. Also a number of measures including the accelerometer data and the fatigue ratings were recorded at home following the 315 test day. Furthermore all data processing was performed blinded and a statistician blinded to 316 317 the groupings performed analysis. The inclusion/ exclusion criteria in this study was very broad including those with all types of MS and on any medication for their condition and/ or 318 other health conditions (excluding glycaemic control medication). Therefore the 319 320 heterogeneity of the participant characteristics in addition to the small sample size may have reduced the observed effect. However the study enabled the procedures to be tested across 321 322 subtypes and establish a follow on study as feasible.

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