Piloting the objective measurement of eating weight and speed at a population scale: a nested study within the Avon Longitudinal Study of Parents and Children [version 2; peer review: 1 approved, 1 approved with reservations]

Previously titled: 'Piloting the objective measurement of eating behaviour at a population scale: a nested study within the Avon Longitudinal Study of Parents and Children'

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Abstract

Background: Effective measurement and adaption of eating behaviours, such as eating speed, may improve weight loss and weight over time. We assessed whether the Mandometer, a portable weighing scale connected to a computer that generates a graph of food removal rate from the plate to which it is connected, together with photo-imaging of food, might prove an effective approach to measuring eating behaviours at large scale.

Methods: We deployed the Mandometer in the home environment to measure main meals over three days of 95 21-year-old participants of the Avon Longitudinal Study of Parents and Children. We used multi-level models to describe food weight and eating speed and, as exemplar analyses, examined the relationship of eating behaviours with body mass index (BMI), dietary composition (fat content) and genotypic variation (the FTO rs9939609 variant). Using this pilot data, we calculated the sample size required to detect differences in food weight and eating speed between groups of an exposure variable.
**Results:** All participants were able to use the Mandometer effectively after brief training. In exemplar analyses, evidence suggested that obese participants consumed more food than those of "normal" weight (i.e., BMI 19 to <25 kg/m^2^) and that A/A *FTO* homozygotes (an indicator of higher weight) ate at a faster rate compared to T/T homozygotes. There was also some evidence that those with a high-fat diet consumed less food than those with a low-fat diet, but no strong evidence that individuals with medium- or high-fat diets ate at a faster rate.

**Conclusions:** We demonstrated the potential for assessing eating weight and speed in a short-term home setting and combining this with information in a research setting. This study may offer the opportunity to design interventions tailored for at-risk eating behaviours, offering advantages over the "one size fits all" approach of current failing obesity interventions.

**Keywords**
Eating behaviour, obesity, Mandometer, ALSPAC

This article is included in the Avon Longitudinal Study of Parents and Children (ALSPAC) gateway.
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Amendments from Version 1

In response to the recent comments from the first reviewer, we have changed the manuscript in the following ways (full details in the official response to the reviewer):
- Updated the title to “Piloting the objective measurement of eating weight and speed at a population scale: a nested study within the Avon Longitudinal Study of Parents and Children”
- Discussed the results in the context of possible treatments and interventions, rather than therapy.
- Mentioned reactivity as a mechanism by which the Mandometer could be used in interventions/treatments.
- Clarified the study design as a pilot study and how the participants were told to use and how we assessed whether participants were able to use the Mandometer effectively.

Any further responses from the reviewers can be found at the end of the article

Introduction

In many countries, the prevalence of obesity and related complications such as type 2 diabetes continue to increase1–3. Interventions aimed solely at improving diet and physical activity ignore other potentially modifiable and important factors. For instance, there is evidence that specifically addressing eating behaviours, such as reducing eating speed, may improve weight loss and maintain weight improvement over time4,5. In 1974, Schachter and Rodin suggested that obese individuals eat at an increased rate, which can dissociate satiety from the amount of food ingested, potentially leading to overeating6. Indeed, an experimental increase in the speed of eating in “normal” weight volunteers (i.e., a body mass index (BMI) ≥19 to <25 kg/m²) caused overeating and delayed the development of satiety, potentially mirroring the pattern of eating in a group of obese patients7. The hypothesis that eating rate is causally related to food intake (and therefore body weight) has been translated into the successful management of adolescent obesity in a randomised controlled trial (RCT) using a device able to record food weight throughout the course of a meal8.

Developed as an intervention to aid food consumption, the Mandometer is a portable weight scale connected to a small computer that can create a graph representing the rate of food ingested, potentially leading to overeating. Indeed, an experimental increase in the speed of eating in “normal” weight volunteers (i.e., a body mass index (BMI) ≥19 to <25 kg/m²) caused overeating and delayed the development of satiety, potentially mirroring the pattern of eating in a group of obese patients. The hypothesis that eating rate is causally related to food intake (and therefore body weight) has been translated into the successful management of adolescent obesity in a randomised controlled trial (RCT) using a device able to record food weight throughout the course of a meal.

The association between obesity and eating behaviour has been widely discussed in the literature, with variation in weight playing a role in food consumption and differences in eating behaviour forming a component of obesity predisposition9–12. Whilst a greater amount of energy (and therefore food) is required to maintain a greater body mass, variation in eating behaviour could, in part, be a cause of the obesity epidemic, with higher food consumption, lower responsiveness to satiety cues and greater responsiveness to external food cues leading to increased weight gain13–12.

Furthermore, over the last ten years, it has been well documented that various common genome-wide, single nucleotide polymorphisms (SNPs) are associated with both adult and childhood obesity13. As an example, in 2007 it was established that variation in and around the FTO gene locus on chromosome 16 was associated with obesity in children and adults14,15, with studies pointing to the causal variant playing a role in regulation of nearby genes (i.e., ARID5B, IRX3 and IRX5) that have downstream effects on thermogenesis to lipid storage, adipocyte size, increased fat stores and body weight gain16,17. Once this gene had been identified as being associated with obesity, researchers started examining how polymorphic variation impacts on obesity risk18. The A allele of rs9939609 has been linked to a tendency to increased energy dense food consumption19 and increased food intake but decreased satiety responsiveness20,21. Functional magnetic resonance imaging studies have also demonstrated variation in food cues by FTO genotype and hormonal studies have suggested that the same genetic variation may be involved in the regulation of Ghrelin, a key orexigenic hormone, activity22,23.

Alongside adiposity and related genetic variation, differential dietary composition, macronutrient distribution and patterns are associated with eating behaviours, including consumption quantity and speed24,25. For example, intake of foods with a high fat content is associated with a higher consumption and eating speed26,27, possibly explained by the impact of an excess amount of fat on satiation and the impact of increased energy density on energy intake28. Given the complex interplay between adiposity, nutritional content and both internal and external environmental cues to food intake, it remains difficult to disentangle the relationships between eating behaviours and adiposity-related traits.

Together, these studies suggest that unravelling some of the potential mechanisms underlying adiposity, genetic propensity and dietary composition as correlates of eating behaviour may be identifiable and may provide new avenues for future interventions and treatment strategies. Despite this, methods used to assess these behaviours tend to be labour- and technology-intensive and likely too costly for more extensive population studies. Therefore, this study aimed to assess whether the Mandometer, together with imaging of food to estimate macronutrient content and total calories on the plate, might prove a less intensive and more economical approach to examining eating behaviours.
Methods
The Avon Longitudinal Study of Parents and Children (ALSPAC) is a large geographically homogeneous prospective birth cohort from the southwest of England established to investigate environmental and genetic characteristics that influence health, development and growth of children and their parents. Full details of the cohort and study design have been described previously and are available at http://www.alspac.bris.ac.uk. Please note that the study website contains details of all the data that is available through a fully searchable data dictionary and variable search tool (http://www.bristol.ac.uk/alspac/researchers/our-data/).

Briefly, 14,541 pregnant women residing in the former county of Avon with an estimated delivery date of between the 1st of April 1991 and the 31st of December 1992 (inclusive) were enrolled to the study. Out of those initially enrolled, 13,998 children who were still alive at 1 year have been followed up to date with measures obtained through regular questionnaires and clinical visits, providing information on a range of behavioural, lifestyle and biological data. Ethical approval for the study was obtained from the ALSPAC Ethics and Law Committee and the Local Research Ethics Committees. Consent for biological samples has been collected in accordance with the Human Tissue Act (2004) and informed consent for the use of data collected via questionnaires and clinics was obtained from participants following the recommendations of the ALSPAC Ethics and Law Committee at the time. Written informed consent was obtained from mothers at recruitment, from the main carers (usually the mothers) for assessments on the children from ages 7 to 16 years and, from age 16 years onwards, the children gave written informed consent at all assessments.

Eating behaviour measurement
The Mandometer (Microdiktat, Sweden) was developed at the Section of Applied Neuroendocrinology and Mandometer Clinic, Karolinska Institutet, Stockholm, Sweden. The device is a portable weighing scale connected to a small computer that generates a graph representing the rate of food removal from the plate to which it is connected (with weight of food (grams) on the y-axis and time (minutes) on the x-axis). In the therapeutic setting, the user puts a self-determined portion of food on the scale (plate) and the connected computer records and displays to the user, in real-time graphics, the weight loss from the plate as the user eats from time zero (total portion size) to the time they have finished eating.

Study recruitment and methods
In this pilot study, we aimed to collect data on eating behaviours (specifically, food weight and eating speed) recorded using the Mandometer in ‘home-based, monitored eating sessions’ within a random sample of the ALSPAC cohort. Recruitment was through a positive response to a randomly selected, mailed-out invitation with information sheet about the study to individuals in the ALSPAC cohort when their mean age was 21 years. Those wishing to take part came to the central clinic centre for ALSPAC and went through the study requirements with a research nurse. If happy to take part, they signed a consent form and were then instructed how to use the Mandometer in a baseline data collection clinic at Oakfield House, Bristol until the participants felt happy to repeat the process at home on their own. Of the 1,117 invites sent out, 214 accepted and 95 individuals participated in the study.

Over three consecutive days (usually covering one week day and the weekend), participants were asked to go home and eat three separate meals of their choosing (usually dinner) at home recording total weight of meal consumed (grams) and speed of eating in grams/second without the device providing any feedback (so called “blind-meals”). They also took a photograph of each meal using a digital camera with a short description to assess total calorie content and major food types eaten, alongside a questionnaire to provide written detail of the foods consumed.

When three meals had been recorded, the individual under study contacted the research nurse who arranged for a courier to pick up all the equipment. Mandometer data was downloaded on to MandoBase, the central repository for Mandometer data in Sweden. Digital photographs were uploaded on to a central computer prior to analysis. Using the photos and information provided by the participant for each meal, an experienced dietician identified the foods present and visually estimated the weight of each food type before entering data in Dietplan 6 (Forestfield Software Ltd, United Kingdom).

Upon recruitment (i.e., when participants were 21 years of age), participants also attended a clinic where they were weighed on Seca scales (nearest 0.1 kg) and their height was measured using a Harpenden Stadiometer (nearest 0.1 cm) whilst wearing light clothing and socks. The participants’ BMI was calculated as weight (kg) divided by height (m) squared.

Exposure data
As exemplar analyses typically of that likely to be undertaken with data on eating behaviours, we assessed whether eating behaviour was associated with genetic variation, a contemporaneous measure of BMI and average dietary composition of the meals consumed.

Genetic variation. As part of the ALSPAC study sample, participants were genotyped using the Illumina HumanHap550 quad genome-wide SNP genotyping platform. Participants were excluded due to having at least one of: incorrectly recorded sex, minimal or excessive heterozygosity, disproportionate levels of individual missingness, evidence of cryptic relatedness or non-European ancestry. SNPs with a minor allele frequency (MAF) of <1% and call rate of <95% were removed and only SNPs that passed an exact test of Hardy-Weinberg equilibrium (P<5x10^-7) were included. For this project and at the time of recall, imputation of genotypes was conducted with MACH 1.0.16 Markov Chain Haplotyping software, using CEPH individuals from phase 2 of the HapMap project as a reference (release #22), where imputation quality was high (>0.9). Variation in the FTO gene is known to be associated with BMI and explains the most variance in BMI of any known genetic variant.
experiment, we observed “spikes” in the data (Figure 1) and hypothesised that these could be caused by potential user interactions with the Mandometer (e.g., users placing cutlery on the plate after taking a bite). Attempting to alleviate these potential artefacts, we performed basic smoothing of the data. Here, we only allow weight to decrease. If an increase in weight was detected at a particular snapshot, then the weight was set as the previous snapshot (i.e., we forced the weight to remain constant during these events) (Figure 1).

Multi-level models were used to account for the hierarchical structure of the data (i.e., repeated measurements of food weight taken within meals and several meals measured within participants). This three-level arrangement was controlled for by allowing one random intercept term for meals and another for participants. These random effects partitioned the variance into (i) between participant, (ii) between meal and (iii) within meal estimates, which account for the non-independence of food weight within meals and within participants. For a given exposure (for example, the FTO rs9939609 genotype), the model for food weight is represented by the following equation.

\[
\text{mass}_i = \beta_0 + u_{0i} + v_{0i} + (\beta_1 + u_{i1} + v_{i1}) \times \text{time}_i + \beta_2 \times 1_{A/T} + \beta_3 \times 1_{A/A} + \beta_4 \times 1_{T/T} + \beta_5 \times 1_{A/T} \times \text{time}_i + \beta_6 \times 1_{A/A} \times \text{time}_i + \epsilon_{0i} + \epsilon_{2i} + \epsilon_{4i}
\]

where \( i = 1, \ldots, n \) indexes the participant, \( j = 1, \ldots, m \) indexes the meal, \( k = 1, \ldots, n_j \) are the repeated measurements of food weight within each meal. \( u_{0i} \) is the random intercept for each participant \( i \), which allows participants to have different average starting meal weights from the overall sample intercept (\( \beta_0 \)). \( v_{0i} \) is a random intercept for meals within participants, which allows meals to have different weights. The random slope terms \( u_{j1} \) and \( v_{ij} \) allow different eating speeds for participant \( i \) and meal \( j \) within participant \( i \), respectively.

\( 1_{A/T} \) is an indicator variable, which takes the value 1 if participant \( i \) has an A/T genotype and 0 otherwise and, similarly, \( 1_{A/A} \) is an indicator for participant \( i \) having a A/A genotype (here, \( T/T \) is the reference group). \( \beta_1 \) gives the average change in meal weight for each unit of time (seconds) or the rate at which food is consumed; \( \beta_2 \) gives the mean difference in average meal weight between T/T and A/T participants; \( \beta_3 \) gives the mean difference in average meal weight between T/T and A/A participants; \( \beta_4 \) is the mean difference in average food intake rate between T/T and A/T participants; \( \beta_5 \) is the mean difference in average eating rate between T/T and A/A participants; \( \epsilon_{0i} \) captures the within meal variability, which has a constant variance \( \sigma^2_{\epsilon} \). A value of zero shows that each meal is eaten in a perfectly linear fashion and that there is no error about this line.

**Sensitivity analysis**
We carried out a simple multi-level model comprising only random intercepts for individuals and meals to try and realise gross effects. This assumes that participants in each category eat at the same speed, which allows all of the variation in eating speed to be attributed to the individual categories; thereby, making it easier to detect differences between them that are otherwise difficult to determine with low sample sizes.
Using this pilot data, we were able to carry out a sample size calculation to find the number of individuals required to adequately power (80%) an analysis of eating speed and food weight across BMI, the FTO genotype or nutritional categories. Given the lack of previous results in this area, the effect size to detect was set by convention to 0.5 standard deviations (SDs) of food weight and eating speed. The SD of food weight was straightforward to calculate from the observed data. Eating speed was calculated using the best linear unbiased predictor (BLUP) for each meal, and the SD was calculated across these eating speeds.

Using multiple meals per individual leads to clustered data (i.e., each new meal for the same individual is not entirely independent). We present sample sizes required if each individual were to eat two, three or four meals each. The total number of meals needed can then be calculated by multiplying the original sample size by the design effect. This is calculated as $1+(k-1)\times ICC$, where $k$ is the cluster size (i.e., two, three or four) and the ICC (intra-class correlation) is the proportion of variation in the outcome that is accounted for by the between-meal variation. The ICC is calculated from the multi-level models as the ratio of the between meal variation (i.e., the random intercept and slope variance for $v_{0j}$ and $v_{1j}$) to the total variation.

**Results**

Of the 95 people who participated in the Mandometer exercise, there were 89, 60 and 81 participants who had measured BMI, genotype and nutritional information available, respectively, with 54 having all three alongside descriptive characteristics (Table 1). Approximately one-fifth of these participants were male (25.93%), with a mean age of 21 years (SD = 0.66) at the time of this study. On average, individuals had a BMI of 24.10 kg/m$^2$. 

![Figure 1. Raw (top panel) and smoothed (bottom panel) Mandometer data (N=50).](image-url)
Table 1. Sample characteristics of individuals with measures of body mass index, the FTO genotype and nutritional information available (N=54).

<table>
<thead>
<tr>
<th>Variable</th>
<th>N</th>
<th>Mean (SD) or percentage</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sex (% male)</td>
<td>54</td>
<td>25.93</td>
</tr>
<tr>
<td>Age (years) of clinic</td>
<td>54</td>
<td>21.02 (0.66)</td>
</tr>
<tr>
<td>Average time taken (seconds) to eat meal</td>
<td>54</td>
<td>544.76 (448.37)</td>
</tr>
<tr>
<td>Average starting food weight (g) for meal</td>
<td>54</td>
<td>355.74 (155.92)</td>
</tr>
<tr>
<td>BMI (kg/m²)</td>
<td>54</td>
<td>24.10 (4.27)</td>
</tr>
<tr>
<td>BMI categories</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Normal</td>
<td>36</td>
<td>66.67</td>
</tr>
<tr>
<td>Overweight</td>
<td>14</td>
<td>25.93</td>
</tr>
<tr>
<td>Obese</td>
<td>&lt;5</td>
<td>7.41</td>
</tr>
<tr>
<td>FTO rs9939609 genotype</td>
<td>54</td>
<td></td>
</tr>
<tr>
<td>T/T</td>
<td>14</td>
<td>25.93</td>
</tr>
<tr>
<td>A/T</td>
<td>32</td>
<td>59.26</td>
</tr>
<tr>
<td>A/A</td>
<td>8</td>
<td>14.81</td>
</tr>
<tr>
<td>MAF of FTO rs9939609</td>
<td>54</td>
<td>0.44</td>
</tr>
<tr>
<td>Fat Content</td>
<td>54</td>
<td></td>
</tr>
<tr>
<td>Low (&lt;30%)</td>
<td>14</td>
<td>25.93</td>
</tr>
<tr>
<td>Medium (≥30% &lt;35%)</td>
<td>15</td>
<td>27.78</td>
</tr>
<tr>
<td>High (&gt;35%)</td>
<td>25</td>
<td>46.30</td>
</tr>
</tbody>
</table>

BMI = body mass index; MAF = minor allele frequency; SD = standard deviation.

Table 2. Results from the multi-level model of Mandometer eating behaviour across three meals stratified by body mass index (BMI) (N=89).

<table>
<thead>
<tr>
<th>Category</th>
<th>Estimate (95% CI)</th>
<th>P-value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Average meal weight (normal BMI; g)</td>
<td>358.09 (335.03, 381.14)</td>
<td>1.49x10⁻²⁰³</td>
</tr>
<tr>
<td>Average difference in meal weight (overweight vs. normal BMI; g)</td>
<td>9.02 (-36.41, 54.46)</td>
<td>0.70</td>
</tr>
<tr>
<td>Average difference in meal weight (obese vs. normal BMI; g)</td>
<td>136.17 (52.19, 220.15)</td>
<td>0.001</td>
</tr>
<tr>
<td>Average eating speed (normal BMI; g/s)</td>
<td>1.46 (1.22, 1.70)</td>
<td>6.64x10⁻³³</td>
</tr>
<tr>
<td>Average difference in eating speed (overweight vs. normal BMI; g/s)</td>
<td>-0.18 (-0.65, 0.29)</td>
<td>0.45</td>
</tr>
<tr>
<td>Average difference in eating speed (obese vs. normal BMI; g/s)</td>
<td>0.18 (-0.69, 1.06)</td>
<td>0.68</td>
</tr>
</tbody>
</table>

Estimates represent the average difference in meal weight (g) or eating speed (g/s) in participants with a “normal” BMI, in overweight participants vs. participants with a “normal” BMI or in obese participants vs. participants with a “normal” BMI (where indicated in the “Category” column).
(A/A, $P=0.17$) consumed more food. There was some evidence that homozygote individuals ate at a faster rate (0.77g/second faster; 95% CI: -0.01, 1.55; $P=0.05$); however, there was no strong evidence that heterozygote individuals ($P=0.63$) consumed food at a faster rate (Table 3 and Figure 3). Due to the relative sample size of genotype groups, estimates are imprecise.

**Fat content**

On average, individuals who consumed a low-fat diet using the Mandometer (i.e., <30% fat across 3 meals) started off (intercept of our multi-level models) with 400.25g of food (95% CI: 356.53, 443.96) and ate at a rate (baseline slope of our multi-level models) of 1.23g/second (95% CI: 0.91, 1.54). There was some evidence that those with a high-fat diet consumed -48.28g less food (95% CI: -101.76, 5.19; $P=0.08$), while there was no strong evidence that those with a medium-fat diet ($P=0.49$) ate more. Furthermore, there was no strong evidence that individuals with a medium fat diet ($P=0.70$) or a high-fat diet ($P=0.66$) consumed food at a faster rate (Table 4 and Figure 4). As with the other groups, the estimates are imprecise due to the low sample size.

**Sensitivity analyses**

Using the random intercept model, the average meal weight for “normal” weight category was estimated to be 266.23g (95% CI: 246.04, 286.43). On average, participants with a “normal” BMI consumed food at a rate of 0.29g/second (95% CI: 0.29, 0.29). In this model, there was evidence that obese individuals had larger meals than individuals with a “normal” BMI (170.67g; 95% CI: 97.11, 244.23; $P<0.0001$). In addition, there was evidence that overweight and obese participants ate at a faster rate than participants with a “normal” BMI ($P<0.0001$). Overweight participants ate 0.13g/second quicker (95% CI: 0.12, 0.13) than participants with a “normal” BMI, while obese participants ate 0.41g/second faster on average (95% CI: 0.39, 0.42) (Table 5, Figure 5).
Table 3. Results from the multi-level model of Mandometer eating behaviour across three meals stratified by FTO genotype (N=60).

<table>
<thead>
<tr>
<th>Category</th>
<th>Estimate (95% CI)</th>
<th>P-value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Average meal weight (T/T genotype; g)</td>
<td>339.83 (290.36, 389.31)</td>
<td>2.58x10^{-41}</td>
</tr>
<tr>
<td>Average difference in meal weight (A/T vs. T/T genotype; g)</td>
<td>23.36 (-35.62, 82.34)</td>
<td>0.44</td>
</tr>
<tr>
<td>Average difference in meal weight (A/A vs. T/T genotype; g)</td>
<td>53.69 (-22.99, 130.36)</td>
<td>0.17</td>
</tr>
<tr>
<td>Average eating speed (T/T genotype; g/s)</td>
<td>1.45 (0.94, 1.96)</td>
<td>2.24x10^{-8}</td>
</tr>
<tr>
<td>Average difference in eating speed (A/T vs. T/T genotype; g/s)</td>
<td>-0.15 (-0.75, 0.45)</td>
<td>0.63</td>
</tr>
<tr>
<td>Average difference in eating speed (A/A vs. T/T genotype; g/s)</td>
<td>0.77 (-0.01, 1.55)</td>
<td>0.05</td>
</tr>
</tbody>
</table>

Estimates represent the average difference in meal weight (g) or eating speed (g/s) in participants with a homozygous T/T genotype, in participants with a heterozygous A/T genotype vs. participants with a homozygous T/T genotype, or in participants with the homozygous A/A genotype vs. participants with a homozygous T/T genotype (where indicated by the “Category” column).

Figure 3. Smoothed output from the Mandometer device stratified by FTO genotype (top panel), and relationships between meal weight and eating behaviour stratified by FTO genotype (bottom panel) (n=60).
Table 4. Results from the multi-level model of Mandometer eating behaviour across three meals stratified by total fat content (N=81).

<table>
<thead>
<tr>
<th>Category</th>
<th>Estimate (95% CI)</th>
<th>P-value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Average meal weight (low fat; g)</td>
<td>400.25 (356.53, 443.96)</td>
<td>5.19x10^{-72}</td>
</tr>
<tr>
<td>Average difference in meal weight (medium fat vs. low fat; g)</td>
<td>-22.47 (-86.82, 41.88)</td>
<td>0.49</td>
</tr>
<tr>
<td>Average difference in meal weight (high fat vs. low fat; g)</td>
<td>-48.28 (-101.76, 5.19)</td>
<td>0.08</td>
</tr>
<tr>
<td>Average eating speed (low fat; g/s)</td>
<td>1.23 (0.91, 1.54)</td>
<td>1.54x10^{-14}</td>
</tr>
<tr>
<td>Average difference in eating speed (medium fat vs. low fat; g/s)</td>
<td>0.09 (-0.37, 0.55)</td>
<td>0.70</td>
</tr>
<tr>
<td>Average difference in eating speed (high fat vs. low fat; g/s)</td>
<td>0.08 (-0.30, 0.47)</td>
<td>0.66</td>
</tr>
</tbody>
</table>

*Estimates represent the average difference in meal weight (g) or eating speed (g/s) in participants with a low fat meal content, in participants with a medium fat vs. low fat meal content, or in participants with a high fat vs. low fat meal content (where indicated by the “Category” column).*

Figure 4. Smoothed output from the Mandometer device stratified by total fat content (top panel), and relationships between meal weight and eating behaviour stratified by total fat content (bottom panel) (N=81).
Table 5. Results from the multi-level model (using random intercepts for individuals and meals) of Mandometer eating behaviour across three meals stratified by body mass index (BMI) (N=89).

<table>
<thead>
<tr>
<th>Category</th>
<th>Estimate (95% CI)</th>
<th>P-value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Average meal weight (normal BMI; g)</td>
<td>266.23 (246.04, 286.43)</td>
<td>3.26x10^{-147}</td>
</tr>
<tr>
<td>Average difference in meal weight (overweight vs. normal BMI; g)</td>
<td>23.47 (-16.35, 63.29)</td>
<td>0.25</td>
</tr>
<tr>
<td>Average difference in meal weight (obese vs. normal BMI; g)</td>
<td>170.67 (97.11, 244.23)</td>
<td>5.43x10^{-06}</td>
</tr>
<tr>
<td>Average eating speed (normal BMI; g/s)</td>
<td>0.29 (0.29, 0.29)</td>
<td>&lt;1x10^{-299}</td>
</tr>
<tr>
<td>Average difference in eating speed (overweight vs. normal BMI; g/s)</td>
<td>0.13 (0.12, 0.13)</td>
<td>&lt;1x10^{-299}</td>
</tr>
<tr>
<td>Average difference in eating speed (obese vs. normal BMI; g/s)</td>
<td>0.41 (0.39, 0.42)</td>
<td>&lt;1x10^{-299}</td>
</tr>
</tbody>
</table>

1 Estimates represent the average difference in meal weight (g) or eating speed (g/s) in participants with a “normal” BMI, in overweight participants vs. participants with a “normal” BMI or in obese participants vs. participants with a “normal” BMI (where indicated in the “Category” column).

Figure 5. Smoothed output from the Mandometer device stratified by body mass index (BMI, top panel), and relationships between meal weight and eating behaviour (bottom panel) stratified by BMI (using random intercepts only for individuals and meals) (N=89).
The average meal weight for the T/T genotype was estimated to be 214.14g (95% CI: 176.32, 251.97) and individuals with this genotype consumed food at a rate of 0.15g/second (95% CI: 0.15, 0.15). There was evidence that individuals with the A/T and A/A genotype had larger meals (P=0.0003). Those with the A/T genotype had 83.68g more food (95% CI: 38.57, 128.79) and those with the A/A genotype had 106.39g more food (95% CI: 48.30, 164.48). Furthermore, there was evidence that the A/T and A/A genotypes consumed food at a faster rate (P<0.0001). Those with the A/T genotype ate 0.29g/second faster than those with the T/T genotype (95% CI: 0.29, 0.30), and those with the A/A genotype ate 0.26g/second faster than those with the T/T genotype (95% CI: 0.26, 0.27) (Table 6, Figure 6).

Finally, we also observed a similar trend when stratifying by average fat content. The average meal weight for individuals with a low-fat diet was 325.94g (95% CI: 290.30, 361.58). Individuals in this group consumed food at a rate of 0.40g/second (95% CI: 0.40, 0.41; P<0.0001). In this model, individuals with a medium fat diet had a similar amount of food (P=0.98) but ate at a rate of 0.17g/second faster (95% CI: 0.16, 0.18; P<0.0001) than those with a low-fat diet. Those with a high-fat diet ate 70.98g less food (95% CI: 27.41, 114.55; P=0.001) and 0.15g/second slower (95% CI: 0.14, 0.15; P<0.0001) than those with a low-fat diet (Table 7, Figure 7).

### Power

Using the pilot data, the SDs of food weight and eating speed were estimated to be 123g and 0.3g/second, respectively. To achieve 80% power to detect a true difference between groups of 67g in food weight (alpha 0.05), 67 individuals in each group were required. If this difference was additive across BMI categories, then a total sample size of 201 individuals eating one meal each would be required for a three-group exposure variable of interest (i.e., 201 total meals). To adequately power the same analysis for eating speed, 73 individuals were required in each group, or 219 individuals (i.e., 219 total meals) across three, say, BMI groups.

The ICC of meals within individuals was calculated to be 0.97, suggesting that meals were very similar within individuals, and thus multiple meals would not give much extra power and may actually reduce sample size. The design effects for two, three and four meals per person were 1.97, 2.94 and 3.91, respectively. If two meals were recorded for each individual, then 432 total meals (1.97 times the 219 meals estimated for the eating speed analysis above) would be needed, or 216 individuals (432 divided by 2 meals per person - a saving of three people). Similarly, for both three and four meals per person, we would require a sample size of 215 people for both.

### Discussion

In this study, we have shown that it is possible to measure in-depth eating behaviours (i.e., food weight and eating speed) within a healthy population. For future analyses, our calculations suggest that, in order to reliably detect differences in self-selected portion size, we would need 67 individuals in each group (i.e., 201 individuals eating one meal); whereas, to detect differences in eating speed, we would need 73 individuals in each group (i.e., 219 individuals across the three groups). Importantly, these sample sizes reflect the phenotypic precision of the Mandometer and relatively low sample sizes required

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**Table 6. Results from the multi-level model (using random intercepts for just individuals and meals) of Mandometer eating behaviour across three meals stratified by genotype (N=60).**

<table>
<thead>
<tr>
<th>Category</th>
<th>Estimate (95% CI)</th>
<th>P-value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Average meal weight (T/T genotype; g)</td>
<td>214.14 (176.32, 251.97)</td>
<td>1.31x10^-28</td>
</tr>
<tr>
<td>Average difference in meal weight (A/T vs. T/T genotype; g)</td>
<td>83.68 (38.57, 128.79)</td>
<td>0.0003</td>
</tr>
<tr>
<td>Average difference in meal weight (A/A vs. T/T genotype; g)</td>
<td>106.39 (48.30, 164.48)</td>
<td>0.0003</td>
</tr>
<tr>
<td>Average eating speed (T/T genotype; g/s)</td>
<td>0.15 (0.15, 0.15)</td>
<td>&lt;1x10^-29</td>
</tr>
<tr>
<td>Average difference in eating speed (A/T vs. T/T genotype g/s)</td>
<td>0.29 (0.29, 0.30)</td>
<td>&lt;1x10^-29</td>
</tr>
<tr>
<td>Average difference in eating speed (A/A vs. T/T genotype; g/s)</td>
<td>0.26 (0.26, 0.27)</td>
<td>&lt;1x10^-29</td>
</tr>
</tbody>
</table>

1Estimates represent the average difference in meal weight (g) or eating speed (g/s) in participants with a homozygous T/T genotype, in participants with a heterozygous A/T genotype vs. participants with a homozygous T/T genotype, or in participants with the homozygous A/A genotype vs. participants with a homozygous T/T genotype (where indicated by the “Category” column).
Figure 6. Smoothed output from the Mandometer device stratified by genotype (top panel), and relationships between meal weight and eating behaviour (bottom panel) stratified by genotype (using random intercepts only for individuals and meals) (N=60).

Table 7. Results from the multi-level model (using random intercepts for just individuals and meals) of Mandometer eating behaviour across three meals stratified by total fat content (N=81).

<table>
<thead>
<tr>
<th>Category</th>
<th>Estimate (95% CI)(^1)</th>
<th>P-value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Average meal weight (low fat; g)</td>
<td>325.94 (290.30, 361.58)</td>
<td>7.47x10^{-22}</td>
</tr>
<tr>
<td>Average difference in meal weight (medium fat vs. low fat; g)</td>
<td>-0.83 (-53.29, 51.62)</td>
<td>0.98</td>
</tr>
<tr>
<td>Average difference in meal weight (high fat vs. low fat; g)</td>
<td>-70.98 (-114.55, -27.41)</td>
<td>0.001</td>
</tr>
<tr>
<td>Average eating speed (low fat; g/s)</td>
<td>0.40 (0.40, 0.41)</td>
<td>&lt;1x10^{-299}</td>
</tr>
<tr>
<td>Average difference in eating speed (medium fat vs. low fat; g/s)</td>
<td>0.17 (0.16, 0.18)</td>
<td>&lt;1x10^{-299}</td>
</tr>
<tr>
<td>Average difference in eating speed (high fat vs. low fat; g/s)</td>
<td>-0.15 (-0.15, -0.14)</td>
<td>&lt;1x10^{-299}</td>
</tr>
</tbody>
</table>

\(^1\)Estimates represent the average difference in meal weight (g) or eating speed (g/s) in participants with a low fat meal content, in participants with a medium fat vs. low fat meal content, or in participants with a high fat vs. low fat meal content (where indicated by the “Category” column).
to achieve resolution in tests by pertinent exposures in this study (i.e., genotype, BMI and meal composition). Other useful details relate to variation in speed of eating across a meal, potentially reflecting satiety responsiveness. It has been demonstrated that those individuals who slow their speed of eating as the meal progresses (so called ‘decelerators’) tend to rate their feeling of fullness higher than linear eaters at the end of a standard meal\(^4^3\). In addition, individuals whose eating speed is naturally linear tend to respond to Mandometer training to eat slower by decreasing food intake.

We used a subsample of the ALSPAC birth cohort to assess the utility of the Mandometer as an instrument to assess eating behaviour at the population level, and whether these eating patterns were associated with BMI, genetic variation associated with BMI and overall nutritional content as exemplar analyses. Using multi-level models, we observed some evidence to suggest that obese participants consumed more food than “normal” weight participants (i.e., BMI $\geq$19 to $<25$ kg/m\(^2\)) and that those with a homozygote \(F\!)O\ genetic variant (i.e., an indicator of higher weight) ate at a faster rate.

We also tested a simpler multi-level model (i.e., one that included random intercepts only for the individual and the meal) as a sensitivity analysis. This simpler model allowed for all the variation in eating speed to be attributed to the individual categories and made differences between the phenotypic groups easier to detect, despite the low sample size. Here, we observed evidence for differences between our phenotypic groups. For example, we observed that overweight and obese individuals consumed more food than those with a “normal” BMI. Furthermore, we also observed that overweight and obese

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**Figure 7.** Smoothed output from the Mandometer device stratified by total fat content (top panel), and relationships between meal weight and eating behaviour (bottom panel) stratified by total fat content (using random intercepts only for individuals and meals) (N=81).
individuals ate at a faster rate when compared to individuals with a “normal” BMI. These trends also replicated across genotypic and nutritional content strata.

In this study, participants each ate several meals meaning that there is a level hierarchy of repeated measures of food weight (i.e., within meals and within participants). To deal with this hierarchy appropriately, we used multi-level models to control for the correlation in meal weight and eating speed within individuals over time and across meals. Findings from sensitivity analyses using only random intercepts in our multi-level models provided some evidence that differences in measured weight categories (i.e., overweight/obese vs. “normal” weight), genotypes (i.e., A/A, A/T vs. T/T) and dietary fat content varied in terms of total food consumption and eating speed. Despite this, not including random slopes in these models are likely not realistic, given they remove inter-individual eating speeds. Therefore, these results should be taken with caution. A further limitation is the assessment of food type and weight on the Mandometer plate, which was assessed by only one experienced dietician, and the limited variability and generalisability in the assessment of only one meal per day, which may not capture full dietary information of the participants. Finally, the use of categorical BMI and fat content groups instead of continuously measured BMI and fat intake reduced power to detect associations between eating speed and these traits. However, this was done to mirror the three-level categories of the FTO genetic variant and was a pilot exercise to demonstrate the potential for the future use of the Mandometer as an assessment for eating behaviour.

Conclusions

This study, together with others (e.g., those that have examined eating behaviours in school children in Sweden\(^4\)), demonstrated that the Mandometer device has potential in assessing eating behaviour when used in a research setting. It may become an additional tool in nutritional epidemiology for examining phenotype-genotype relationships between pro-adiposity eating behaviours and at-risk polymorphisms. Those studies using fMRI, or other laboratory-based testing such as video analysis of eating behaviour\(^4\), offer granular detail but are very costly and time consuming. We have shown that with one clinic visit, which realistically and easily could be transformed into a teaching video, we could examine eating behaviours in the home environment over a period of time. Furthermore, given the potential of the Mandometer to change eating behaviour via reactivity between the participant and device, it may offer the opportunity to design effective and efficient interventions that are tailored for genetically determined at-risk eating behaviours. Therefore, this offers advantages over the “one size fits all” approach of current obesity interventions that in general achieve only modest improvement in medium to long term adiposity\(^46,47\).

Data availability

Underlying data

ALSPAC data are available through a system of managed open access. Data for this project was accessed under the project number B1038. The application steps for ALSPAC data access are highlighted below.

1. Please read the ALSPAC access policy which describes the process of accessing the data in detail, and outlines the costs associated with doing so.

2. You may also find it useful to browse the fully searchable research proposals database, which lists all research projects that have been approved since April 2011.

3. Please submit your research proposal for consideration by the ALSPAC Executive Committee.

You will receive a response within 10 working days to advise you whether your proposal has been approved. If you have any questions about accessing data, please email alspac-data@bristol.ac.uk. The study website also contains details of all the data that is available through a fully searchable data dictionary: http://www.bristol.ac.uk/alspac/researchers/data-access/data-dictionary/.

Acknowledgements

We are extremely grateful to all the families who took part in this study, the midwives for their help in recruiting them, and the whole ALSPAC team, which includes interviewers, computer and laboratory technicians, clerical workers, research scientists, volunteers, managers, receptionists and nurses. We also want to acknowledge Hashem Shihab, who conducted the initial analyses for this manuscript.

References


Open Peer Review

Current Peer Review Status: ✓ ?

Version 2

Reviewer Report 05 May 2021

https://doi.org/10.21956/wellcomeopenres.18248.r43588

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Phil Morys
Montreal Neurological Institute, McGill University, Montreal, Quebec, Canada

Wade and colleagues present a pilot study using a relatively large sample size to test the usefulness of a Mandometer in describing eating patterns and relating them to weight status, genotype and dietary fat intake. This is a very carefully conducted and well-designed study which is an excellent introduction to using the Mandometer in future research. The Authors find that meal weight and eating speed differ in groups of individuals with different BMI, genotype and dietary fat intake. More importantly, however, they conduct power analyses to calculate the minimum number of participants to detect between group differences in meal size and eating speed. The Authors are very thorough throughout their analyses and describe the limitations of the study very well. However, some analytical choices were not clearly motivated in the manuscript, some of statistical assumptions were not checked prior to conducting the analyses, and I believe that the Authors could do a slightly better job in describing their results both in terms of being explicit about their sample sizes in each tested group, but also in terms of checking whether their results are accurate, or perhaps driven by outliers or smaller group sizes. I have a small number of mostly minor comments that I hope will help improve this great manuscript.

1. Why were the participants divided into groups only based on fat content in their meals? Why not carbohydrates or proteins? This decision needs to be motivated.

2. In the first paragraph of the Results section - 25% is one-fourth, not one-fifth

3. Why were the participants divided into groups only based on fat content in their meals? Why not carbohydrates or proteins? This decision needs to be motivated.

4. I would like the Authors to discuss how the fact that individuals might have eaten different meals (e.g. breakfast vs. dinner) could affect their results. If, for example, obese participants only ate breakfast using the Mandometer, then their meal weight would presumably be lower. Do the Authors have the information about which meal was eaten? Can this be taken
into account in their analyses?

5. It is unclear from the text (but clear from the Tables and Figures) whether the results are presented for the sample of 54 participants or for the full sample for each respective measure.

6. There is only information about distribution of participants in different groups for the sample of 54 participants, but not for other samples. This should be available at least in the supplement.

7. Seeing huge differences in the sample size between obese and lean individuals (but not only) I wonder whether the Authors tested for the assumption of homoscedasticity between the two groups. The variance between groups should be the same, otherwise confidence intervals, parameter estimates and significance cannot be properly calculated in the multilevel models. If the variance is different, this should at least be mentioned as a limitation and I think the Authors should be explicit about their group sizes in the Results section. It is a pilot study after all, so one cannot expect it to be perfect and the outcomes should be judged by the reader, but the reader needs to know what exactly is happening in the described research.

8. I would like to see a scatter plot of starting meal weight and weight status/BMI. From Fig. 2a it seems like the group differences here might be driven by an outlier in the obese group. A similar problem might occur in the genetic analysis. Did the Authors check this?

9. How exactly is the eating speed calculated? Is this measure adjusted for the initial weight of the meal? Would this not make sense?

10. The Authors state that ‘due to the relative sample size of genotype groups, estimates are imprecise’ - why is this different than in the BMI groups? Shouldn't this statement also be valid for normal weight, overweight and obese groups?

Is the work clearly and accurately presented and does it cite the current literature?
Yes

Is the study design appropriate and is the work technically sound?
Yes

Are sufficient details of methods and analysis provided to allow replication by others?
Yes

If applicable, is the statistical analysis and its interpretation appropriate?
Partly

Are all the source data underlying the results available to ensure full reproducibility?
Yes

Are the conclusions drawn adequately supported by the results?
Competing Interests: No competing interests were disclosed.

Reviewer Expertise: Neurobiological mechanisms of obesity and metabolic syndrome, neuroimaging, eating disorders, eating behaviour

I confirm that I have read this submission and believe that I have an appropriate level of expertise to confirm that it is of an acceptable scientific standard, however I have significant reservations, as outlined above.

Author Response 07 Jun 2021
Kaitlin Wade, University of Bristol, Bristol, UK

We thank the Reviewer for reading over our manuscript and providing such useful comments. We have responded to each comment below and amended the manuscript accordingly in a manner that we hope is satisfactory.

Comment 1: I am a bit puzzled by the general aims of this manuscript. In the abstract the Authors say that the goals are to define interventions tailored for at-risk eating behaviours, yet in the Introduction, they state that the study aims to assess whether the Mandometer is a less intensive and more economical approach to examine eating behaviours. I think a potential for designing interventions, if any, should also be mentioned in the Introduction.
Response 1: The reviewer is correct, and we apologise for any confusion and inconsistency between the Abstract and Introduction of the manuscript. We have amended by editing the abstract to mirror what is stated in the introduction. As such, the abstract aims now read as follows: “We assessed whether the Mandometer, a portable weighing scale connected to a computer that generates a graph of food removal rate from the plate to which it is connected, together with photo-imaging of food, might prove a less intensive and more economical approach to measuring eating behaviours at large scale”. We have added a small statement about how this device may offer an opportunity to design interventions tailored for at-risk individuals to the last sentence of the introduction.

Comment 2: Why were the participants divided into groups only based on fat content in their meals? Why not carbohydrates or proteins? This decision needs to be motivated.
Response 2: As this was a pilot study, we explored fat content as an example correlate of eating speed and weight to see whether this could be done for protein and carbohydrates using similar methodology in a larger study. We have now made this clearer in the manuscript.

Comment 3: In the first paragraph of the Results section - 25% is one-fourth, not one-fifth.
Response 3: Thank you for pointing this out – this has been amended.

Comment 4: I would like the Authors to discuss how the fact that individuals might have eaten different meals (e.g., breakfast vs. dinner) could affect their results. If, for example, obese participants only ate breakfast using the Mandometer, then their meal weight would presumably
be lower. Do the Authors have the information about which meal was eaten? Can this be taken into account in their analyses?

Response 4: The random intercept for each participant included within the multi-level models do allow for differential meal weight within participants; therefore, this is, in part, accounted for in the way that we have modelled the data. However, the reviewer highlights an important point in that it is plausible that individuals in different weight categories may have chosen to systematically record different meals with the Mandometer device. Therefore, the difference in initial meal weight (and potentially eating speed) may be due to the choice of meal recorded by those individuals. It is worth noting, however, that participants were asked to record a “cooked” meal of their choosing (information of which has now been clarified in the manuscript) and this resulted in mainly evening meals being recorded (~90% of meals were eaten after 4pm). Of those who recorded meals that were likely to reflect breakfast or lunchtime meals, only two individuals consistently recorded meals at lunch, both of whom were in the “normal” weight category. Otherwise, all other individuals recorded meals at varying times (again, usually dinner). Although this is unlikely to have impacted our results, it is something that could be of interest in further evaluations in larger samples. This has been added to the discussion as a future consideration.

Comment 5: It is unclear from the text (but clear from the Tables and Figures) whether the results are presented for the sample of 54 participants or for the full sample for each respective measure.

Response 5: We have added sample sizes where relevant in the results.

Comment 6: There is only information about distribution of participants in different groups for the sample of 54 participants, but not for other samples. This should be available at least in the supplement.

Response 6: We have now added a table showing the sample characteristics of the pool of 95 participants with non-missing data to the main text and referenced accordingly in the manuscript.

Comment 7: Seeing huge differences in the sample size between obese and lean individuals (but not only) I wonder whether the Authors tested for the assumption of homoscedasticity between the two groups. The variance between groups should be the same, otherwise confidence intervals, parameter estimates and significance cannot be properly calculated in the multilevel models. If the variance is different, this should at least be mentioned as a limitation and I think the Authors should be explicit about their group sizes in the Results section. It is a pilot study after all, so one cannot expect it to be perfect and the outcomes should be judged by the reader, but the reader needs to know what exactly is happening in the described research.

Response 7: The reviewer has highlighted an important point and having checked this in the current sample, there does not seem to be a substantial difference in the variances between BMI, genotype and fat content groups (box plots provided below). Therefore, whilst unlikely to have a substantial impact in the current study, this is certainly something that needs to be considered in future, larger studies if using similar categorical exposure variables.
BMI

Genotype
Fat content
Comment 8: I would like to see a scatter plot of starting meal weight and weight status/BMI. From Fig. 2a it seems like the group differences here might be driven by an outlier in the obese group. A similar problem might occur in the genetic analysis. Did the Authors check this?

Response 8: Given that this was a pilot study demonstrating the utility of the Mandometer device, exploring the impact of a small number of outliers on the results presented was not within the remit of the current analysis. However, this is definitely something that would need to be considered in future evaluations of this relationship in larger studies.

Comment 9: How exactly is the eating speed calculated? Is this measure adjusted for the initial weight of the meal? Would this not make sense?

Response 9: For each participant, eating speed was calculated as the number of grams consumed by the participant (as measured by the Mandometer device) divided by the number of seconds it took to consume the food (i.e., the slope of the line showing the relationship between seconds on the x-axis and food weight on the y-axis). The equations in the manuscript describe the specific models, where our multi-level models, specifically the random intercepts model, allowed for different mean meal
weights and eating speeds between each participant and meal that a participant consumed. The intercept term represents the average initial weight of all meals, with the random intercept allowing this to vary between individuals and between different meals within each individual.

Comment 10: The Authors state that ‘due to the relative sample size of genotype groups, estimates are imprecise’ - why is this different than in the BMI groups? Shouldn’t this statement also be valid for normal weight, overweight and obese groups?
Response 10: Given that this is a pilot study, it is true that the sample sizes of all groups are small and therefore estimates are more imprecise than what would be achieved with larger sample sizes. We have removed this statement specifically from the section presenting the results of the genotype groups and added this to the discussion as an overall limitation.

Competing Interests: N/A

Reviewer Report 25 January 2021

https://doi.org/10.21956/wellcomeopenres.18248.r42304

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Falko F. Sniehotta
NIHR Policy Research Unit Behavioural Science, Newcastle University, Newcastle, UK

Thanks, nice paper. I don't have any further comments.

Is the work clearly and accurately presented and does it cite the current literature? Yes

Is the study design appropriate and is the work technically sound? Yes

Are sufficient details of methods and analysis provided to allow replication by others? Yes

If applicable, is the statistical analysis and its interpretation appropriate? Yes

Are all the source data underlying the results available to ensure full reproducibility? Yes

Are the conclusions drawn adequately supported by the results?
Yes

**Competing Interests:** No competing interests were disclosed.

**Reviewer Expertise:** I am a behavioural scientist and have an interest in applied obesity research. I am also interested in measurement reactivity.

I confirm that I have read this submission and believe that I have an appropriate level of expertise to confirm that it is of an acceptable scientific standard.

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**Author Response 05 May 2021**

Kaitlin Wade, University of Bristol, Bristol, UK

Thank you!

**Competing Interests:** No competing interests were disclosed.

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**Version 1**

Reviewer Report 21 December 2020

https://doi.org/10.21956/wellcomeopenres.17657.r41015

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Falko F. Sniehotta
NIHR Policy Research Unit Behavioural Science, Newcastle University, Newcastle, UK

Congratulations, this is an interesting and well written report. There are just a few odd points I would like to highlight.

1. The title refer to an "objective measurement of eating behaviour" when the Mendometer is mainly an objective measure of consumption quantity and speed. It would be more helpful to make this clear in the title of the paper, not only for future evidence synthesis.

2. In the context of overweight and obesity the manuscript refers to the term therapy. This term makes all sorts of assumptions about the nature of obesity and the evidence for treatment options which are probably not defensible. Why not referring to intervention?

3. Surprising that the paper does not refer to reactivity. Monitoring of intake with or without feedback is a common behaviour change technique and the Mandometer has the potential of changing the behaviour it measures. This deserves a mention, here is the latest relevant systematic review https://www.sciencedirect.com/science/article/pii/S08954356193114611.
4. There is a confusion about the nature of the study. It is twice referred to as feasibility study and once as a pilot. What is it? Assuming it is a feasibility study, it would be helpful to clarify what aspect of feasibility was tested and how. In the results, the manuscript reads “All participants were able to use the Mandometer effectively at home after brief training.” How exactly was this measured? Also, if it is a feasibility study, this should be reflected in the title.

Aside from these points, I thoroughly enjoyed reading the paper.

References

**Is the work clearly and accurately presented and does it cite the current literature?**
Yes

**Is the study design appropriate and is the work technically sound?**
Yes

**Are sufficient details of methods and analysis provided to allow replication by others?**
Yes

**If applicable, is the statistical analysis and its interpretation appropriate?**
Yes

**Are all the source data underlying the results available to ensure full reproducibility?**
Yes

**Are the conclusions drawn adequately supported by the results?**
Yes

**Competing Interests:** No competing interests were disclosed.

**Reviewer Expertise:** I am a behavioural scientist and have an interest in applied obesity research. I am also interested in measurement reactivity.

I confirm that I have read this submission and believe that I have an appropriate level of expertise to confirm that it is of an acceptable scientific standard.
satisfactory.

Comment 1: The title refer to an “objective measurement of eating behaviour” when the Mendometer is mainly an objective measure of consumption quantity and speed. It would be more helpful to make this clear in the title of the paper, not only for future evidence synthesis.
Response 1: We totally agree with the reviewer, as we tried to make this clear throughout the text so have mirrored this by changing the title, as suggested by the Reviewer, to “Piloting the objective measurement of eating weight and speed at a population scale: a nested study within the Avon Longitudinal Study of Parents and Children”.

Comment 2: In the context of overweight and obesity the manuscript refers to the term therapy. This term makes all sorts of assumptions about the nature of obesity and the evidence for treatment options which are probably not defensible. Why not referring to intervention?
Response 2: We agree with the Reviewer and have changed the manuscript such that we discuss the results in the context of possible treatments and interventions, rather than therapy. Specifically, in the introduction, we have changed the word “therapy” in the following sentence such that it now reads as follows: “Together, these studies suggest that unravelling some of the potential mechanisms underlying adiposity, genetic propensity and dietary composition as correlates of eating behaviour may be identifiable and may provide new avenues for future interventions and treatment strategies.”

Comment 3: Surprising that the paper does not refer to reactivity. Monitoring of intake with or without feedback is a common behaviour change technique and the Mandometer has the potential of changing the behaviour it measures. This deserves a mention, here is the latest relevant systematic review https://www.sciencedirect.com/science/article/pii/S08954356193114611.
Response 3: The point is very well taken. The interactivity of the Mandometer device does indeed have the potential of changing the behaviour it measures and one of the most likely mechanisms by which the Mandometer could be used in interventions/treatment of obesity. We have mentioned this in the conclusive statement of the discussion, specifically: “Furthermore, given the potential of the Mandometer to change eating behaviour via reactivity between the participant and device, it may offer the opportunity to design effective and efficient interventions that are tailored for genetically determined at-risk eating behaviours.”

Comment 4: There is a confusion about the nature of the study. It is twice referred to as feasibility study and once as a pilot. What is it? Assuming it is a feasibility study, it would be helpful to clarify what aspect of feasibility was tested and how. In the results, the manuscript reads “All participants were able to use the Mandometer effectively at home after brief training.” How exactly was this measured? Also, if it is a feasibility study, this should be reflected in the title.
Response 4: We apologise for this uncertainty in the study design – the work we present is indeed more of a pilot study rather than a feasibility study, as the main aim was to undertake a study that could be repeated at larger scale with a similar design. We have removed the “feasibility study” wording and replaced with “pilot study” for clarity. With regards to the inference about participants being able to use the
Mandometer, participants were shown how to use the device at a baseline collection clinic by a trained research nurse until the participants felt happy to repeat the process at home on their own, but this was not tested formally – future and larger studies could indeed expand on this and formally test the usage accuracy. We have made this clearer in the “Study recruitment and methods” of the Methods section of the manuscript.

**Competing Interests:** N/A