METHOD ARTICLE

Updating the evidence on the effectiveness of the alcohol reduction app, Drink Less: using Bayes factors to analyse trial datasets supplemented with extended recruitment [version 1; referees: 1 approved with reservations]

Claire Garnett 1, Susan Michie 2, Robert West 1, Jamie Brown 1

1Department of Behavioural Science and Health, University College London, London, UK
2Department of Clinical, Education and Health Psychology, University College London, London, UK

Abstract

Background: A factorial experiment evaluating the Drink Less app found no clear evidence for main effects of enhanced versus minimal versions of five components but some evidence for an interaction effect. Bayes factors (BFs) showed the data to be insensitive. This study examined the use of BFs to update the evidence with further recruitment.

Methods: A between-subject factorial experiment evaluated the main and two-way interaction effects of enhanced versus minimal version of five components of Drink Less. Participants were excessive drinkers, aged 18+, and living in the UK. After the required sample size was reached (n=672), additional data were collected for five months. Outcome measures were change in past week alcohol consumption and Alcohol Use Disorders Identification Test (AUDIT) score at one-month follow-up, amongst responders only. BFs (with a half-normal distribution) were calculated for those for which we had outcome data (BF<0.33 indicate evidence for null hypothesis; 0.33<BF<3 indicate data are insensitive).

Results: Of the sample of 2586, 342 (13.2%) responded to follow-up. Data were mainly insensitive but tended to support there being no large main effects of the enhanced version of individual components on consumption (0.22<BF<0.83) or AUDIT score (0.14<BF<0.98). Data no longer supported there being two-way interaction effects. In an unplanned comparison, participants receiving the four most promising components averaged a numerically greater reduction in consumption than those not receiving any (21.6 versus 12.1 units), but the data were insensitive (BF=1.42).

Conclusions: Data from extended recruitment in a factorial experiment evaluating components of the Drink Less app remained insensitive but tended towards individual and pairs of components not having a large effect. There was weak evidence for a synergistic effect of four components. In the event of uncertain results, calculating BFs can be used to update the strength of evidence of a dataset supplemented with extended recruitment.

Keywords
Bayes Factors, digital interventions, alcohol reduction, smartphone apps
Corresponding author: Claire Garnett (c.garnett@ucl.ac.uk)

Author roles: Garnett C: Conceptualization, Data Curation, Formal Analysis, Investigation, Project Administration, Writing – Original Draft Preparation, Writing – Review & Editing; Michie S: Conceptualization, Funding Acquisition, Supervision, Writing – Review & Editing; West R: Conceptualization, Funding Acquisition, Supervision, Writing – Review & Editing; Brown J: Conceptualization, Funding Acquisition, Supervision, Writing – Review & Editing

Competing interests: No competing interests were disclosed.

Grant information: JB and RW are funded by Cancer Research UK (CRUK; C1417/A22962). CG and SM are funded by CRUK and the National Institute for Health Research (NIHR)'s School for Public Health Research (SPHR). Drink Less was funded by NIHR SPHR, the UK Centre for Tobacco and Alcohol Studies (UKCTAS), the Society for the Study of Addiction (SSA), and CRUK. The views expressed are those of the author(s) and not necessarily those of the NHS, the National Institute for Health Research or the Department of Health. The research team is part of the UKCTAS, a UKCRC Public Health Research Centre of Excellence. Funding from the Medical Research Council, British Heart Foundation, Cancer Research UK, Economic and Social Research Council and the National Institute for Health Research under the auspices of the UK Clinical Research Collaboration, is gratefully acknowledged.

Copyright: © 2019 Garnett C et al. This is an open access article distributed under the terms of the Creative Commons Attribution Licence, which permits unrestricted use, distribution, and reproduction in any medium, provided the original work is properly cited.

How to cite this article: Garnett C, Michie S, West R and Brown J. Updating the evidence on the effectiveness of the alcohol reduction app, Drink Less: using Bayes factors to analyse trial datasets supplemented with extended recruitment [version 1; referees: 1 approved with reservations] F1000Research 2019, 8:114 (https://doi.org/10.12688/f1000research.17952.1)

First published: 29 Jan 2019, 8:114 (https://doi.org/10.12688/f1000research.17952.1)
**Introduction**

A factorial experiment evaluating the effect of ‘enhanced’ versus ‘minimal’ versions of five components of the alcohol reduction app, Drink Less, found no clear evidence for simple effects but did find evidence that two-way combinations of certain ‘enhanced’ components together resulted in greater reductions than ‘minimal’ versions. This was a planned analysis but should be interpreted with caution as the two-way interactive effects were not specifically hypothesised a priori and were part of multiple interactions tested. Findings of this sort are not uncommon in experimental studies. One approach is to start another randomised trial specifically to test this hypothesis. A potentially more efficient alternative is to extend the trial with further recruitment and test this and other hypotheses using Bayes factors. We used this approach with the Drink Less app.

Bayes factors are a measure of strength of evidence and allow researchers to ‘top-up’ their results from one trial with additional data collected, regardless of the stopping rule, unlike frequentist statistics. The use of Bayes factors supports efficient, incremental model building, as evidence can be continuously accumulated until it is clear whether there is an association or not. The rapid accumulation of large amounts of data about digital behaviour change interventions (DBCIs) offers the opportunity to apply emerging methods to their evaluation. DBCIs often have the capacity to continue automatic data collection beyond the end of a trial with little or no additional resources. This paper will illustrate how Bayes Factors can be used to optimise a DBCI by updating evidence from an effectiveness trial using the example of Drink Less—an alcohol reduction app.

Bayes factors are the ratio of the average likelihood of two competing hypotheses being correct given a set of data and can overcome some of the issues associated with traditional frequentist statistics. They indicate the relevant strength of evidence for two hypotheses; when evaluating interventions, the two hypotheses are typically the alternative hypothesis (the intervention had the desired effect) and the null hypothesis (the intervention had no effect). Bayes factors, unlike frequentist statistics, can distinguish between two interpretations of a non-significant result: i) support for the null hypothesis of ‘no effect’ and ii) data are insensitive to detect an effect i.e. ‘unsure about the presence of an effect’. Calculating Bayes factors to supplement frequentist statistics is a quick and simple procedure with several software packages freely available (e.g. an online calculator developed by Zoltan Dienes’). Researchers are actively encouraged to supplement, or even replace, classical frequentist hypothesis testing with a Bayesian approach to provide greater interpretative value to any non-significant results. This is important as often non-significant results are misinterpreted as evidence for no effect; a review of trials conducted in addictions research found that the reporting of ‘no difference’ was only appropriate in a small number of papers reporting this.

The use of Bayes Factors also has another major advantage over the traditional frequentist approach that relates to the stopping rule. The traditional frequentist approach necessitates a strict stopping rule and a single analysis of data. Typically, this involves an a priori power calculation to specify the required sample size for data collection and the trial to end at that point. Subsequent ‘topping-up’ of existing data and re-analysing the new larger data set is ‘prohibited’. This is because any p-value between 0 and 1 is equally likely if the null hypothesis is true, regardless of how much data are collected. Therefore, given enough time and data collection, a significant p-value will always be obtained even if the null hypothesis is true. So if researchers find a non-significant result—which cannot distinguish between support for the null hypothesis and being insensitive to detect an effect—then a new study would be required to build on these findings. Restarting the process is a waste of research resources but necessary in the context of using a frequentist approach for analysis because additional data collected cannot be analysed. However, this is not the case when using Bayes factors, as they are driven towards zero when the null hypothesis is true and additional data are collected. Therefore, researchers may use Bayes factors to analyse additional data to complement an employed stopping rule.

In the evaluation of DBCIs, using Bayes factors is beginning to complement traditional frequentist statistics, and analysing additional data would be of particular benefit. Data collection for a DBCI effectiveness trial is typically automated and therefore does not require additional resources to continue after a pre-specified sample size is reached. Rapid evaluations of DBCIs and efficient accumulation of evidence can be used to inform future versions, keeping pace with advances in technology. Using Bayes factors to update findings about the relative plausibility of the two hypotheses allows researchers to assess the DBCI’s effectiveness in an ongoing manner. This remains useful when deciding about whether there is sufficient evidence to demonstrate effectiveness and, therefore, continued development. To the authors’ knowledge, no DBCIs have used additional data collected to supplement original effectiveness trial findings.

DBCIs require novel methods of evaluation that are quick and timely to inform the optimisation of the intervention. The multiphase optimisation strategy (MOST) is a method for building, optimising and evaluating multicomponent behavioural interventions. It involves a series of steps identifying the set of intervention components to be examined and evaluating the effects of these components. Factorial trial designs allow the simultaneous evaluation of the intervention components, which enables both the independent and interactive effects to be estimated. Using a factorial trial to evaluate a DBCI can overcome some of the challenges associated with using the traditional randomised controlled trial, such as prolonged duration from recruitment to publication and a high-cost trial implementation. The results from a factorial trial can be used to make decisions about which components to retain when optimising the intervention.

The Drink Less smartphone app is a DBCI aimed at supporting people who drink excessively to reduce their alcohol consumption. It was developed using evidence and theory, following MOST. The app was analysed in a full factorial trial to assess the effectiveness of its five intervention modules and their effects on app usage and subsequent usability ratings. The stopping rule for data collection, in line with the frequentist approach to analysis,
was pre-specified, although data collection continued under the same conditions as the original factorial trial. Analysis of the original trial data using Bayes factors indicated that the data were insensitive to detect main effects but that combinations of the modules appeared effective1.

**Aims**

The aims of this study are substantive and methodological:

1. To update the evidence on effectiveness of Drink Less app components singly and in combination. Specifically, what are the main and two-way interactive effects of the intervention modules on:
   i. Change in weekly alcohol consumption
   ii. Change in full Alcohol Use Disorders Identification Test (AUDIT) score

2. To demonstrate how Bayes Factors can be used to analyse additional outcome data collected in effectiveness trials and update beliefs about hypotheses.

**Methods**

**Design**

A between-subject full factorial (23) trial to evaluate the effectiveness of five intervention modules in the Drink Less app. The research questions were specified prior to the trial commencing and pre-registered on ISRCTN (registration number: ISRCTN40104069) and published in an open-access protocol paper15.

**Participants**

Participants were included in the study if they: were aged 18 or over; lived in the UK; had an AUDIT score of 8 or above (indicative of excessive drinking9); were interested in reducing their drinking; provided an email address and had downloaded a ‘trial version’ of the app (described below).

The sample size for the original factorial trial was 672 providing 80% power (with alpha at 5%, 1:1 allocation and a two-tailed test) to detect a mean change in alcohol consumption of 5 units between the ‘enhanced’ and ‘minimal’ versions for each intervention module50, comparable with a face-to-face brief intervention11. This assumed a mean of 27 weekly units at follow-up in the control group, a mean of 22 units in the intervention group and a SD of 23 units for both (d=0.22).

Recruitment was undertaken via promotion from organisations, such as Public Health England, Cancer Research UK, and listing the app in the iTunes Store according to best practices for app store optimisation.

**Measures**

Baseline measures included the AUDIT questionnaire and a socio-demographic assessment (age, gender, ethnic group, level of education, employment status and current smoking status). The primary outcome measure was self-reported change in past week alcohol consumption (the difference between one-month follow-up and baseline). The secondary outcome measure was self-reported change in full AUDIT score.

**Interventions**

The Drink Less app is a DBCI for people who drink excessively to help them reduce their alcohol consumption. It is freely available on the UK version of the Apple App Store for all smartphones and tablets running iOS8 or above. The content of the app did not change during the trial except for minor bug fixes.

The app is structured around goal setting: users can set their own goals based on units, cost, alcohol free days or calories with information on the UK drinking guidelines, units and alcohol-related harms. There are five intervention modules that aim to help them achieve their goal: Normative Feedback (providing normative feedback on the user’s level of drinking relative to others); Cognitive Bias Re-training (a game to retrain approach-avoidance bias for alcoholic drinks); Self-monitoring and Feedback (providing a facility for self-monitoring of drinking and receipt of feedback); Action Planning (helping users to undertake action planning to avoid drinking), and Identity Change (promoting a change in identity in relation to alcohol). In the trial version of the app, the five intervention modules existed in two versions: i) an ‘enhanced’ version containing the predicted active ingredients and ii) a ‘minimal’ version that acted as a control.

A detailed description of the content, development and factorial trial evaluation of the app is reported in two separate papers16,22.

**Procedures**

Data collection for the factorial trial began on 18th May 2016 and the required sample of eligible users was reached on 10th July 2016; follow-up data were collected until 28th August 2016. Trial data was collected continuously for a further four months until 19th December 2016 under the same conditions as the original factorial trial (i.e. a ‘trial version’).

Informed consent to participate in the trial was obtained from all participants on first opening the app. Users who consented to participate completed the AUDIT and a socio-demographic questionnaire, indicated their reason for using the app and provided their email address for follow-up (a prize of £100 was offered in an attempt to decrease the proportion of users leaving this field blank). Users were then provided with their AUDIT score and, those who met the inclusion criteria, were randomised to one of 32 experimental conditions using an automated algorithm within the app for block randomisation.

Follow-up was conducted 28 days after participants downloaded the app and the questionnaire consisted of the full AUDIT and usability measures. Follow-up was conducted in two ways: i) via email with a link to the questionnaire in an online survey tool (Qualtrics), which also sent up to four reminders, and ii) within the app. Participants included according to the original trial and stopping rule were due to complete the follow-up questionnaire up until 29th August 2016 and were contacted via email.
(through Qualtrics) and the app. Participants due to complete the follow-up questionnaire from 30th August onwards, were only contacted via the app.

**Ethical approval**

Ethical approval for Drink Less from the UCL Ethics Committee under the ‘optimisation and implementation of interventions to change health-related behaviours’ project (CEHP/2013/508).

**Analysis**

All analyses were conducted using R version 3.4.0. The analysis plan for this paper followed a similar analysis plan as for the original factorial trial (which was pre-registered on 13th February 2016; ISRCTN40104069). Participant characteristics were reported descriptively by intervention module. A factorial between-subjects design was used to assess the main and two-way interactive effects of the five intervention modules on the primary and secondary outcome measures. Analyses were conducted amongst responders only, those who completed the follow-up questionnaire. Bayes Factors were calculated for each analysis assessing the main and the two-way interaction effects of the five intervention modules on the outcome measures. The two-way interactions were defined as enhanced/enhanced versus minimal/minimal for each pair of intervention modules. The mean difference and standard error of the mean difference for each main and two-way interactive effect was calculated. A half normal distribution was used to specify the predicted effect. Peak at 0 (no effect) with a SD equal to the expected effect size. This is a conservative approach and represents a hypothesis that the intervention had a least some positive effect, with the effect being more likely to be smaller than larger. Bayes factors were calculated using an online calculator.

The expected effect size for the primary calculation of Bayes factors was a reduction of 5 units per week \( (d=0.22) \), reflecting a large effect and that of the power calculation for the original factorial trial. Bayes Factors were also calculated for a medium effect (reduction of 3 units per week), and a small effect (reduction of 0.5 units per week) to permit a relative judgment for screening purposes. The expected effect size for the secondary outcome measure was calculated by translating the estimated effect size for the primary outcome measure \( (d=0.22) \) into the equivalent mean difference score of 1.45 \( \text{mean}=19.1, \text{SD}=6.56 \) [based on original trial users, \( n=672 \)]. Bayes factors were interpreted in terms of categories of evidential strength (see Table 1).

**Results**

**Study sample**

The total sample size was 2586, of these 1914 (74.0%) were additional users to the original factorial trial (672, 26.0%). In total, 342 users (13.2%) completed the primary outcome measure in the follow-up questionnaire—the original users’ response rate was 26.6% and the additional users’ response rate was 8.5%. Figure 1 shows a flow chart of users throughout the study.

Socio-demographic and drinking characteristics of participants are reported in Table 2. Participants’ mean age was 37.2 years, 53.4% were women, 95.8% were white, 74.3% had post-16 qualifications, 87.0% were employed, and 30.0% were current smokers. Mean weekly alcohol consumption was 39.0 units, mean AUDIT-C score was 9.3, and mean AUDIT score was 19.1, indicating harmful drinking. Participants’ characteristics by intervention module are reported in Table 2. Generally, characteristics were similar for the enhanced and minimal version of each intervention module.

**Change in past week’s alcohol consumption**

The main effects of the intervention modules are reported in Table 3 for the change in past week’s alcohol consumption. Bayes factors showed that the data were insensitive to detect an effect for **Normative Feedback** for effect sizes of 5-, 3- and 0.5-unit reductions \( (0.47<\text{BF}<0.97) \). Data were insensitive to detect an effect for **Cognitive Bias Re-training** for effect sizes of 5-, 3- and 0.5-unit reductions \( (0.74<\text{BF}<1.06) \). Bayes factors showed that the data were insensitive to detect an effect for **Self-monitoring and Feedback** for effect sizes of 5-, 3- and 0.5-unit reductions \( (0.43<\text{BF}<0.95) \). Bayes factors showed that the data were insensitive to detect an effect for **Action Planning** for effect sizes of 5-, 3- and 0.5-unit reductions \( (0.83<\text{BF}<1.08) \). Bayes Factors for **Identity Change** showed support for the null hypothesis of no difference between the enhanced and minimal version of the module for a 5-unit reduction \( (\text{BF}=0.22) \), though data were insensitive to detect an effect for 3- and 0.5-unit reductions \( (0.34<\text{BF}<0.81) \). The data were insensitive to detect a two-way interactive effect between any pair of intervention modules for effect sizes of 5-, 3- or 0.5-unit reductions \( (0.35<\text{BF}<1.22) \), except for between **Self-monitoring and Feedback** and **Identity Change** for a 5-unit reduction which supported the null hypothesis \( (\text{BF}=0.31) \) (see Extended data, Supplementary Table 1).

**Change in AUDIT score**

The main effects of the intervention modules are reported in Table 4 for the change in AUDIT score. The data were insensitive to detect an effect on change in AUDIT score for: **Normative Feedback** \( (\text{BF}=0.60) \); **Cognitive Bias Re-training** \( (\text{BF}=0.98) \); and **Action Planning** \( (\text{BF}=0.95) \). The data supported evidence for the null hypothesis of no effect for **Self-monitoring and Feedback** and **Identity Change** for a 5-unit reduction which supported the null hypothesis \( (\text{BF}=0.31) \) (see Extended data, Supplementary Table 1).

**Table 1. Interpretation of Bayes factors.**

<table>
<thead>
<tr>
<th>Bayes factor</th>
<th>Interpretation</th>
</tr>
</thead>
<tbody>
<tr>
<td>&gt;30</td>
<td>Very strong evidence for ( H_1 )</td>
</tr>
<tr>
<td>10–30</td>
<td>Strong evidence for ( H_1 )</td>
</tr>
<tr>
<td>3–10</td>
<td>Moderate evidence for ( H_0 )</td>
</tr>
<tr>
<td>1–3</td>
<td>Anecdotal evidence for ( H_0 )</td>
</tr>
<tr>
<td>1</td>
<td>No evidence</td>
</tr>
<tr>
<td>0.33–1</td>
<td>Anecdotal evidence for ( H_0 )</td>
</tr>
<tr>
<td>0.10–0.33</td>
<td>Moderate evidence for ( H_0 )</td>
</tr>
<tr>
<td>0.03–0.10</td>
<td>Strong evidence for ( H_0 )</td>
</tr>
<tr>
<td>&lt;0.03</td>
<td>Very strong evidence for ( H_0 )</td>
</tr>
</tbody>
</table>

\( H_0 \), alternative hypothesis; \( H_1 \), null hypothesis.
Figure 1. Flow chart of users.

Total downloads
n= 7913

Completed AUDIT questionnaire
n=6390

Completed socio-demographic assessment
n=6074

Eligible users included in the trial
n=2586

Ineligible for trial, n= 3358
Opted out (n= 34)
Not from UK (n=224)
YOB >=1998 (n=76)
AUDIT <=7 (n=427)
Not interested in drinking less (n=403)
No e-mail address provided (n=2194)

*Hierarchical list of reason for ineligibility

Duplicate cases removed, n=130
Duplicate device ID (n=98)
Duplicate email address (n=32)

Completed follow-up:
Primary outcome measure – change in past week alcohol consumption, n=342
Intensive, n=190
Minimal, n=152

Cognitive bias re-training
Intensive, n=1289
Minimal, n=1297

Completed follow-up:
Primary outcome measure – change in past week alcohol consumption, n=342
Intensive, n=181
Minimal, n=168

Identity change
Intensive, n=1285
Minimal, n=1301

Completed follow-up:
Primary outcome measure – change in past week alcohol consumption, n=342
Intensive, n=181
Minimal, n=161

Action planning
Intensive, n=1286
Minimal, n=1300

Completed follow-up:
Primary outcome measure – change in past week alcohol consumption, n=342
Intensive, n=201
Minimal, n=141

Self-monitoring & feedback
Intensive, n=1292
Minimal, n=1294

Completed follow-up:
Primary outcome measure – change in past week alcohol consumption, n=342
Intensive, n=182
Minimal, n=160

*Hierarchical list of reason for ineligibility
Table 2. Participants’ characteristics at baseline. Data given as mean (SD), unless stated.

<table>
<thead>
<tr>
<th>Variable</th>
<th>All</th>
<th>Normative Feedback</th>
<th>Cognitive Bias Re-training</th>
<th>Self-monitoring &amp; Feedback</th>
<th>Action Planning</th>
<th>Identity Change</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Enh</td>
<td>Min</td>
<td>Enh</td>
<td>Min</td>
<td>Enh</td>
<td>Min</td>
</tr>
<tr>
<td>Age</td>
<td>37.2 (10.64)</td>
<td>36.8 (10.49)</td>
<td>37.5 (10.77)</td>
<td>37.0 (10.56)</td>
<td>37.3 (10.72)</td>
<td>37.3 (10.88)</td>
</tr>
<tr>
<td>% Women (%)</td>
<td>53.4% (1381)</td>
<td>52.7% (686)</td>
<td>54.1% (695)</td>
<td>53.0% (693)</td>
<td>53.8% (688)</td>
<td>53.0% (685)</td>
</tr>
<tr>
<td>% White (%)</td>
<td>95.8% (2477)</td>
<td>96.1% (1250)</td>
<td>95.5% (1227)</td>
<td>96.3% (1241)</td>
<td>95.3% (1236)</td>
<td>95.7% (1237)</td>
</tr>
<tr>
<td>% Post-16 qualifications</td>
<td>74.3% (1921)</td>
<td>74.4% (968)</td>
<td>74.2% (953)</td>
<td>74.3% (958)</td>
<td>74.2% (963)</td>
<td>74.1% (958)</td>
</tr>
<tr>
<td>% Employed (%)</td>
<td>87.0% (2250)</td>
<td>87.0% (1132)</td>
<td>87.0% (1118)</td>
<td>87.1% (1123)</td>
<td>86.9% (1127)</td>
<td>85.6% (1106)</td>
</tr>
<tr>
<td>% Current smokers (%)</td>
<td>30.0% (776)</td>
<td>29.7% (386)</td>
<td>30.4% (390)</td>
<td>29.8% (384)</td>
<td>30.2% (392)</td>
<td>28.9% (373)</td>
</tr>
<tr>
<td>Past week alcohol consumption (units)</td>
<td>39.0 (26.93)</td>
<td>39.0 (27.02)</td>
<td>39.0 (26.85)</td>
<td>39.7 (27.66)</td>
<td>38.3 (26.18)</td>
<td>39.2 (26.99)</td>
</tr>
<tr>
<td>AUDIT score</td>
<td>19.1 (6.66)</td>
<td>27.0 (19.18)</td>
<td>26.9 (18.98)</td>
<td>19.4 (6.78)</td>
<td>18.7 (6.52)</td>
<td>19.2 (6.63)</td>
</tr>
<tr>
<td>AUDIT-C score</td>
<td>9.3 (1.76)</td>
<td>9.3 (1.76)</td>
<td>9.3 (1.77)</td>
<td>9.3 (1.80)</td>
<td>9.3 (1.73)</td>
<td>9.3 (1.78)</td>
</tr>
</tbody>
</table>

Enh, enhanced; Min, minimal.

Table 3. Main effects of intervention modules on change in past week’s alcohol consumption. A negative number indicates a reduction over time.

<table>
<thead>
<tr>
<th>Variable</th>
<th>Mean change in alcohol consumption, Units (SD)</th>
<th>Bayes factor</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Enhanced</td>
<td>Minimal</td>
</tr>
<tr>
<td>Normative Feedback</td>
<td>-12.5 (25.70)</td>
<td>-12.7 (26.57)</td>
</tr>
<tr>
<td>Cognitive Bias Re-training</td>
<td>-13.4 (26.93)</td>
<td>-11.9 (25.22)</td>
</tr>
<tr>
<td>Self-monitoring and Feedback</td>
<td>-12.3 (24.97)</td>
<td>-13.0 (27.61)</td>
</tr>
<tr>
<td>Action Planning</td>
<td>-13.5 (24.70)</td>
<td>-11.6 (27.55)</td>
</tr>
<tr>
<td>Identity Change</td>
<td>-10.7 (27.76)</td>
<td>-14.8 (23.89)</td>
</tr>
</tbody>
</table>

Table 4. Main effects of intervention modules on change in AUDIT score.

<table>
<thead>
<tr>
<th>Variable</th>
<th>Mean change in AUDIT score (SD)</th>
<th>Bayes factor</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Enhanced</td>
<td>Minimal</td>
</tr>
<tr>
<td>Normative Feedback</td>
<td>-2.4 (5.55)</td>
<td>-2.04 (6.11)</td>
</tr>
<tr>
<td>Cognitive Bias Re-training</td>
<td>-2.5 (5.73)</td>
<td>-1.9 (5.88)</td>
</tr>
<tr>
<td>Self-monitoring and Feedback</td>
<td>-1.8 (5.62)</td>
<td>-2.8 (6.02)</td>
</tr>
<tr>
<td>Action Planning</td>
<td>-2.5 (6.06)</td>
<td>-1.9 (5.50)</td>
</tr>
<tr>
<td>Identity Change</td>
<td>-1.7 (5.89)</td>
<td>-2.8 (5.66)</td>
</tr>
</tbody>
</table>
The current data also remained insensitive to detect whether the four most promising components (Normative Feedback, Cognitive Bias Re-Training, Self-Monitoring and Feedback and Action Planning) may each have effects smaller than 5 units. An unplanned analysis provided weak anecdotal evidence of a synergistic effect of the ‘enhanced’ versions of these four intervention modules together. On both past week alcohol consumption and AUDIT score, and across several alternative effect sizes, there was support for no effect of the fifth intervention module, Identity Change. These findings, alongside results from analysing user feedback and usage data on the most frequently visited screens, guided the decision to remove the Identity Change module from the next major app update whilst retaining Normative Feedback and Cognitive Bias Re-Training, and Self-Monitoring and Feedback and Action Planning.

A major strength of this study is its illustration of how it is possible to evaluate data from trials of DBCs in an on-going manner. No additional resources were required to continue data collection within the original trial of Drink Less. Analysing the supplemented dataset has allowed us to update our findings and provided more confidence in our original decisions on which components to retain or remove. The stopping rule in frequentist statistics means that additional trial data collected as part of an effectiveness trial for a DBCI would go to waste. The use of Bayes factors in this situation prevents unnecessary waste of resources and enables researchers to continually update their evidence on a DBCI rather than collect and analyse individual data sets as part of separate trials.

A limitation of this study and the use of Bayes factors was that we were not able to use the intention-to-treat (ITT) approach in the analysis (as was done for the original trial), whereby those lost to follow-up (non-responders) were assumed to be drinking at baseline levels. Whilst Bayes factors can overcome a lot of the issues with the frequentist approach, they are not meaningful when assumptions are made that limit the variability in the data. Due to low overall follow-up rates (13.2%) in this larger sample, the ITT assumption that there was no change in the large majority of the sample drives the variability down, which in turn drives support for the null hypothesis. This highlights that Bayes factors were not useful in this study when using the ITT assumption, which limits the variability in the data.

The intervention modules of the Drink Less app do not have a large individual effect on reducing alcohol-related outcomes,

---

**Table 5.** Four modules in ‘enhanced’ vs four modules in ‘minimal’ versions for past week alcohol consumption.

<table>
<thead>
<tr>
<th>Variable</th>
<th>PWAC, Units (SD)</th>
<th>Bayes factor</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>All enhanced</td>
<td>All minimal</td>
</tr>
<tr>
<td>Responders only (change in PWAC)</td>
<td>-21.6 (20.36)</td>
<td>-12.1 (26.82)</td>
</tr>
<tr>
<td>Last observation carried forward (PWAC)</td>
<td>36.7 (28.48)</td>
<td>37.4 (26.59)</td>
</tr>
</tbody>
</table>

PWAC, past week alcohol consumption.
though they may have a small effect that the current data were unable to detect. There is weak evidence for a synergistic effect of the ‘enhanced’ versions of four intervention modules together: Normative Feedback and Cognitive Bias Re-Training, and Self-Monitoring and Feedback and Action Planning. This study has updated the existing evidence on the effectiveness of intervention modules in the Drink Less app. In the event of uncertain results following a primary analysis, Bayes factors can be used to ‘top-up’ results from DBCI trials with any additional data collected, therefore supporting efficient, incremental model building to inform decision-making.

Data availability

Underlying data
A dataset containing the extended trial outcomes is available on OSF. DOI: https://doi.org/10.17605/OSF.IO/KQMSB

Extended data
Extended data are available on OSF. DOI: https://doi.org/10.17605/OSF.IO/KQMSB

Supplementary Table 1. Two-way interactive effects of intervention modules on change in past week’s alcohol consumption.

Supplementary Table 2. Two-way interactive effects of intervention modules on change in AUDIT score.

Data are available under the terms of the Creative Commons Zero “No rights reserved” data waiver (CC0 1.0 Public domain dedication).

Grant information

JB and RW are funded by Cancer Research UK (CRUK; C1417/A22962). CC and SM are funded by CRUK and the National Institute for Health Research (NIHR)’s School for Public Health Research (SPHR). Drink Less was funded by NIHR SPHR, the UK Centre for Tobacco and Alcohol Studies (UKCTAS), the Society for the Study of Addiction (SSA), and CRUK. The views expressed are those of the author(s) and not necessarily those of the NHS, the National Institute for Health Research or the Department of Health. The research team is part of the UKCTAS, a UKCRC Public Health Research Centre of Excellence. Funding from the Medical Research Council, British Heart Foundation, Cancer Research UK, Economic and Social Research Council and the National Institute for Health Research under the auspices of the UK Clinical Research Collaboration, is gratefully acknowledged.

The funders played no role in the design, conduct or analysis of the study, nor in the interpretation or reporting of study findings.

References

7. Dienes Z: Making the most of your data with Bayes. [Internet]. Reference Source

Page 9 of 13


http://www.doi.org/10.17605/OSF.IO/KQMBB
This paper describes the effectiveness of DrinkLess, a mobile alcohol intervention, using Bayes factor to further compliment previously published findings. I have one major comment and several minor comments.

Major Comment:
1. The work extends our understanding of DrinkLess and its effectiveness in managing alcohol misuse; however, it would be helpful to make a clear statement on how the Bayes analysis has improved the value proposition of DrinkLess.

Minor Comments:
1. Abstract: “Amongst responders only” – is that the sample who took part in the follow-up questionnaire?
2. Abstract: “Unplanned comparison” appears to convey a negative connotation the authors could alter to “additional analyses”;
3. Abstract: “four most promising” could be misleading as you only had five components but we also have to be mindful that the data was insensitive;
4. Abstract: “reminded insensitive but tended” are you able to provide any BF for this statement?;
5. Introduction: It would be helpful to provide some discussion (specific examples) on how Bayes have been used in other domains to provide more insight by means of additional data;
6. Methods – Participants: “were interested in reducing their drinking” how was this measured? Were participants research aware, or were they targeted because they had previously stated an interest in reducing alcohol? Or could it be by downloading DrinkLess they were assumed to be interested in reducing their alcohol consumption?;
7. Methods – Participants: Was a geolocation restriction placed on participants? How can you be sure that users were from the UK?;
8. Methods – Intervention: What were the minor bug fixes, is a summary able to be provided as a supplement?;
9. Results: Results present AUDIT-C score, however this is not discussed previously.
10. Results: It would be helpful to have Table 2 represented as supplementary material for those who took part in follow-up;
11. Results: It would be of interest to discuss further the difference in AUDIT and AUDIT-C score and the role the final two questions (risk taking etc) play;
12. Discussion: “no additional resources were required” – is this the case, was the app provisioned for longer than anticipated;
13. Discussion: “our decision on which components to retain or remove” – a bit more discussion round this aspect would be helpful to the reader;
14. Discussion: A 13.2% follow-up rate appears to be very low, do the authors have any reasons for this?;

Is the rationale for developing the new method (or application) clearly explained?
Partly

Is the description of the method technically sound?
Yes

Are sufficient details provided to allow replication of the method development and its use by others?
Yes

If any results are presented, are all the source data underlying the results available to ensure full reproducibility?
Yes

Are the conclusions about the method and its performance adequately supported by the findings presented in the article?
Partly

**Competing Interests:** Researcher on the InDEx app project - an app designed to help armed forces personnel monitor their alcohol consumption

**Referee Expertise:** Mobile health with a focus on alcohol misuse

I have read this submission. I 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.
The benefits of publishing with F1000Research:

- Your article is published within days, with no editorial bias
- You can publish traditional articles, null/negative results, case reports, data notes and more
- The peer review process is transparent and collaborative
- Your article is indexed in PubMed after passing peer review
- Dedicated customer support at every stage

For pre-submission enquiries, contact research@f1000.com