Evaluating a digital tool for supporting breast cancer patients: a randomised controlled trial protocol (ADAPT)

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Abstract

Background: There is a growing number of mHealth tools for breast cancer patients but a lack of scientific evidence for their effects. Recent studies have shown a mix of positive and negative impacts on users. Here we will assess the impact of OWise Breast Cancer, a mobile application for self-monitoring symptoms and managing care, on the process of self-management.

Methods: This randomised controlled trial with early-stage breast cancer patients will assess the effect of OWise use on patient activation at 3 months from diagnosis measured by the PAM-13 questionnaire. We will also assess differences in changes in health-related quality of life, psychological distress, health status and NHS health resource utilisation over the first year from diagnosis. Participants will be randomly allocated (1:1) to standard care or standard care plus OWise. Participants will complete questionnaires before starting anti-cancer treatment and at 3 months, 6 months and 12 months from diagnosis. Clinical and patient-reported outcome data will be linked to health resource utilisation data from Discover, an integrated care record of primary, secondary and social care in North West London. We will measure contamination in the control group and adjust the sample size to mitigate the dilution of effect estimates. A per protocol analysis will be conducted as a sensitivity analysis to assess robustness of the primary results.

Discussion: This study aims to generate evidence for the effectiveness of OWise at improving patient activation for women with early-stage breast cancer. The results will show the impact of using the tool at the patient level and the NHS health system level. The outcomes of the study will have implications for the application of OWise across the NHS for breast cancer patients and expansion into other tumour types. Assessing publicly available mHealth tools poses a challenge to trialists due to the risk of contamination. Here we apply various methods to measure, mitigate and assess the effects of
contamination.

Trial registration: The study was registered at clinicaltrials.gov (NCT03866655) on 7 March 2019.

Background

Breast cancer is the most common form of cancer diagnosed in the United Kingdom (UK), with around 55,200 new cases each year [1]. In 2010, the National Health Service (NHS) in England spent around £675 million for the care of patients with breast cancer, with current NHS and broader societal costs likely exceeding this value [2]. With the incidence of breast cancer expected to increase in the UK over the next 15 years [1], the NHS is promoting increased self-management of care [3].

Patients and providers are looking to mHealth applications for potential self-management benefit in cancer populations [4]. mHealth is the application of mobile technology by patients or health care providers to monitor health and improve outcomes [5]. A recent review has found 12 studies assessing mHealth tools to support self-management in breast cancer patients [6]. Many of the studies assessing mHealth tools found promising results with a wound monitoring application reducing health resource utilisation [7], an electronic daily journal stabilizing daily functional activity [8] and an application providing information and support improving self-efficacy and quality of life and reducing symptom interference [9]. However, one application providing tailored information before surgery increased levels of anxiety and depression in patients [10]. The inconsistency in effects highlights the need to rigorously assess the impact of mHealth tools before encouraging use.

OWise Breast Cancer is a new mhealth technology for the self-management of care in breast cancer patients. OWise provides tailored medical information, a tracker to self-monitor symptoms and functions to manage care including an appointment calendar,
modifiable question list and consultation recording device [11]. O Wise, listed in the NHS Apps Library, is freely available for download [12]. O Wise was developed outside an academic setting but followed the mHealth development and evaluation process defined by Whittaker et al [13]. Programmers designed the tool in an iterative process with patients and conducted thorough user testing.

A qualitative study evaluating O Wise in the Netherlands showed patients and providers found the tool usable and felt it had the potential to help patients take in more information from consultations, manage appointments and feel more in control during treatment [14]. To understand the impact of O Wise on health behaviours, health-related quality of life (HRQoL) and NHS resource utilisation, comparative data is needed.

**Conceptual model**

The conceptual model for O Wise is based on the Individual and Family Self-Management Theory. This theory posits that self-management is the process by which individual and family health knowledge and behaviours are used to reach certain health outcomes (Figure 1) [15]. The theory takes into account individual, medical, social and environmental factors that influence the process of self-management.

O Wise aims to improve HRQoL and reduce health resource utilisation by intervening on the self-management process. The digital tool aims to increase knowledge and beliefs by providing tailored medical information and recommended questions in the modifiable question list. O Wise aims to improve self-regulation skills and abilities with the symptom tracker and appointment calendar. The proximal outcome we will measure is patient activation and distal outcomes are HRQoL, psychological distress, health status and health care costs. Studies have previously linked patients activation to better HRQoL, improved care experiences and lower use of NHS resources [16–18].

**Study Design**
Aim
This study aims to understand the impact of OWise on health behaviours, HRQoL and health care utilisation in early-stage breast cancer patients compared to standard care alone.

Study design
We will evaluate the effectiveness of OWise using a multi-centre, individually randomized, parallel controlled trial recruiting 122 patients. The intervention group will receive OWise plus standard care, while the control group will receive standard care alone to assess superiority. Due to the nature of the digital tool, it is not possible to blind participants or providers. Patients in both groups will complete patient-reported outcome measures (PROMs) to assess outcomes at baseline, 3 months, 6 months and 12 months from diagnosis. See Figure 2 for the SPIRIT diagram showing the schedule of enrolment and assessment and Additional File 1 for the SPIRIT-PRO checklist.

Randomisation
Patients will be randomly assigned (1:1) to the intervention or control group [19,20]. Randomisation will be stratified by age group and centre. Age is grouped by (1) under 60 years old and (2) 60 years old and over as internet access drops between age groups 45–55 and 55–65 [21] and the incidence of breast cancer in the UK is evenly distributed around age 60 [1]. The Institute of Cancer Research Clinical Trials and Statistics Unit Randomisation Service will generate the randomisation sequence and allocate the group by phone.

Participants
Females (aged 18 years or over) newly diagnosed with early-stage breast cancer as a first primary diagnosis will be eligible to take part. Eligibility was restricted to early-stage and first primary diagnoses as metastatic patients may have confounding care and
psychosocial experiences and patient activation naturally increases with time after a breast cancer diagnosis [22]. Patients must complete the baseline measure before starting anti-cancer treatment. All participating sites are located in the UK, a list of which can be found on the registration website. Exclusion criteria include private care, difficulty reading in English, significant cognitive impairments or poor mental health and no internet access.

**Intervention Group**

OWise is an mHealth tool accessible online or by mobile application [11]. The tool offers tailored medical information, a modifiable question list with tailored recommended questions, a medical terms glossary, useful links to local resources, a tracking tool for symptoms, an appointment calendar and a consultation recording device.

In the study, only patients randomised into the intervention group will receive information about OWise. The individual enrolling the patient will provide instructions for creating an account and navigating the tool. Participants will be free to use the tool as much as they wish to mimic real-world use of the application. The tool is free to download and accessible to the participant beyond the study period.

**Control Group**

Participants in the control group will receive all standard information including leaflets and links to resources that patients usually receive at the time of a new breast cancer diagnosis. Participants in the control group will not be given information about the tool but will also not be explicitly prohibited from using the tool.

**Primary objective**

The primary objective of this study is to test whether use of OWise increases patient activation scores at three months follow-up by at least four points more than standard care.

**Secondary objectives**
(1) To test whether any difference in the change in patient activation between the two groups still exists after controlling for potential covariates.

(2) To test whether the use of OWise leads to a smaller decrease in health status at three months follow-up than standard care after controlling for potential covariates.

(3) To test whether the use of OWise leads to a smaller decrease in HRQoL at three months follow-up than standard care alone after controlling for potential covariates.

(4) To test whether the use of OWise leads to a lower increase in psychological distress at three months follow-up compared to standard care alone after controlling for potential covariates.

(5) To test whether the use of OWise reduces the rate of resource utilisation in the first year following diagnosis compared to standard care among patients registered in Discover, an integrated health and social care record in North West London.

(6) To test whether the use of OWise reduces the average cost per patient in the first year following diagnosis compared to standard care among patients registered in Discover.

(7) To describe the change in patient activation in the intervention group compared to the control group in the first year following diagnosis.

(8) To describe the level of OWise uptake in the intervention group in the first year following diagnosis.

(9) To describe the change in the pattern of patient activation, HRQoL, psychological distress and health status in the intervention group compared to the control group in the first year following diagnosis.

**Procedure**

**Recruitment**

We will continuously sample all patients meeting eligibility criteria diagnosed within the recruitment period. A member of the clinical team will identify eligible patients in multi-
disciplinary team meetings or clinic lists and invite potential participants at diagnosis. The name of the digital tool will not be disclosed when inviting patients. If a patient shows interest, a researcher will provide further information either in-person or over the phone. Patients can decide to take part any time before starting anti-cancer treatment. After meeting eligibility criteria and providing written informed consent, participants will be randomised. The researcher will inform the participant of their allocation and provide instructions for accessing the online PROM collection tool. Participants in the intervention arm will be required to complete the baseline measure before using O Wise.

Measures
Primary outcome measure
Patient Activation Measure (PAM–13)

Patient activation describes the knowledge, skills and confidence a person has in managing their health and care [23,24]. The PAM–13 is a 13-item questionnaire that measures patient activation [25]. Each item has four response options from (1) ‘strongly disagree’ to (4) ‘strongly agree,’ and ‘not applicable.’ PAM–13 scores will be calculated according to the guidelines [25]. Scores range on a scale of 1–100 corresponding to four activation levels: 1 (≤47.0) not believing activation important, 2 (47.1–55.1) a lack of knowledge and confidence to take action, 3 (55.2–67.0) beginning to take action and 4 (≥67.1) taking action [25]. This measure has been used widely among cancer patients and across the UK [26–28] and has robust evidence of reliability and validity [23,29,30]. Patient activation, as measured by the PAM–13, can be targeted by interventions and change over time [24]. Previous work has shown that higher patient activation is associated with HRQoL and lower health care utilisation [16–18,31].

Secondary outcome measures
European Organisation for Research and Treatment
of Cancer Quality of Life Questionnaire - Core30 (EORTC-QLQ-C30 version 3) and Updated Quality of Life Breast Cancer Module (QLQ-BR45)

This 30-item instrument measuring HRQoL has five functional scales (physical, role, cognitive, emotional and social), a global quality of life scale, eight symptom scales or items (fatigue, pain, nausea and vomiting, dyspnoea, loss of appetite, sleep disturbance, constipation and diarrhoea) and a single item assessing perceived financial impact [32]. The QLQ-BR45 contains five functional scales or items (body image, future perspective, sexual functioning, sexual enjoyment and breast satisfaction) and 7 symptom scales or items (systemic therapy side effects, upset by hair loss, arm symptoms, breast symptoms, endocrine therapy symptoms, skin mucosis symptoms and endocrine sexual symptoms) [33]. Scores will be calculated according to EORTC guidelines [34]. All scores range from 0-100. Higher scores on functional scales and global quality of life indicate better function and HRQoL, respectively. Higher scores on symptom scales and items indicate higher symptom burden [35]. The measures have strong evidence of validity and reliability in early-breast cancer patients and have been used in a number of clinical trials allowing for comparisons [33,36].

Hospital Anxiety and Depression Scale (HADS)

This 14-item questionnaire measures psychological distress, with seven items assessing anxiety and seven items assessing depression with the summed total score reflecting the level of psychological distress [37]. Three continuous scales will be calculated (anxiety, depression and overall psychological distress) according to HADS guidelines [37,38]. Higher scores indicate more psychological distress [37,38]. The HADS has evidence of reliability and validity in early breast cancer patients [36,39].

EuroQol 5-Dimension 5-Level (EQ-5D-5L)
This instrument assessing health status consists of five items and a visual analogue scale [40]. The items cover five dimensions (mobility, self-care, usual activities, pain/discomfort and anxiety/depression). Each dimension has five response levels from ‘no problems’ to ‘extreme problems.’ The visual analogue scale records the patient’s self-rated health from 0-100, with the highest score indicating ‘The best health you can imagine’ and the lowest score indicating ‘The worst health you can imagine.’ Responses to each item combine to form a five-digit number that describes the patient’s health state. A corresponding index value will be assigned according to a recent valuation study conducted in England [41]. The EQ-5D-5L has a large base of evidence and for validity and reliability in breast cancer patients and can also be used to conduct economic analysis [42].

Health Care Utilisation

Health care utilisation will be assessed using data routinely collected in Discover. The Discover linked dataset includes coded information on health and social care resource utilisation of individuals registered with a GP practice in the North West London region. Information is collected about the number and type of appointments from primary, secondary and social care for each patient between diagnosis and one year follow-up.

Health Care Costs

The Discover linked dataset also provides the current costs of health and social care to Clinical Commissioners in North West London based upon Commissioner local pricing. Information on the cost of each type of appointment is calculated routinely and collated together across health care settings to provide a measure of health and social care utilisation of each patient.

OWise Uptake

With patient informed consent, we will evaluate OWise uptake by reviewing timestamps that indicate logging in or modification of a specific function. This information will allow us
to evaluate whether participants use the tool, which function patients use and how long the tool is used for.

**Contamination**

Patient responses to items at each measurement time point will indicate contamination. A set of 19 items will ask participants to identify the use of supportive care services including self-management mobile phone applications or websites. If a participant says yes, we will ask them to name the source in a free-text box to determine whether or not OWise has been used. Prior to recruitment close, the statistician will assess the level of contamination and increase the sample size commensurately [43].

**Data management**

The study steering committee determined a data monitoring committee was unnecessary for this study as it poses a minimal risk to patient safety and uses only routinely collected or patient-reported data.

**Patient-Reported Outcome Measure Data**

Participants will complete PROMS using PROFILES (Patient Reported Outcomes Following Initial treatment and Long term Evaluation of Survivorship), an online PROM collection and data management system developed in the Netherlands and implemented at the Royal Marsden [44]. Follow-up time points will be managed by the Royal Marsden as the coordinating centre. PROM responses cannot be viewed by the researcher or clinical team until extracted at the end of the study when it will be linked to other study data by the study identification number.

**Clinical Data**

With patient informed consent, researchers will collect relevant clinical data from the local electronic patient record and store it digitally at the Royal Marsden. Clinical data will be extracted at the end of the study and linked to other study data by the study identification
Health care utilisation

Study participants will be identified in Discover by NHS number and flagged with the study identification number. The Imperial College Health Partners Discover team, will access a de-identified version of the data to analyse health and social care resource utilisation on behalf of the Royal Marsden. Health and Social Care resource utilisation data will be extracted at the end of the study and linked to other study data by the study identification number.

OWise Engagement

Participants will be provided with a unique invitation code linked to their study identification number to input when creating an OWise account. Timestamp data will be identified by the invitation code and extracted at the end of the study. The data will be linked to other study data by the unique invitation code.

Sample size

Sample size calculations are based on the change in PAM–13 score from baseline to three months. A difference of four points is considered clinically relevant [29,45]. In a similar study, the mean PAM–13 score at baseline for the intervention group was 61.3 (SD 16.61) and 67.9 (SD 16.85) at three months [46]. For the control group, the study reported a mean PAM–13 score at baseline of 62.1 (SD 17.30) and 62.8 (SD 14.94) at three months. Based on these findings, this study is planned to detect a mean change difference of 5.90 assuming a common standard deviation of 10.0. Using an 80% power the study will recruit 47 patients per group. This was calculated using a 2-sided test with alpha = 0.05. We will increase the sample size taking 23% attrition at three months into account (47/(1–0.23)) to 61 patients per group [47] and, as mentioned above, increase the sample size accordingly if contamination is found near the end of recruitment.
Analysis

The CONSORT-EHEALTH recommendations for reporting randomised trials for developing and evaluating eHealth interventions will guide trial reports [48]. Primary and secondary outcomes will be assessed using intention-to-treat analysis where all participants are analysed according to the arm to which they were randomised. We will conduct a sensitivity analysis separately as described below. The data will be analysed after all patients have completed the one year follow-up. Data will be reported descriptively at each time point. Mean and standard deviation or median and range will be reported for continuous outcomes. Frequency and percentage will be reported for categorical outcomes.

The association between potential covariates and the primary and secondary endpoints will be explored using univariate analysis. Any variables with a p-value of <0.1 will be included in the multivariable model. Multivariable analysis controlling for potential covariates associated with the particular outcome will be conducted using logistic regression for binary outcomes and multiple linear regression for continuous outcomes.

Two-sided p-values of <0.05 will be considered statistically significant.

Primary endpoint

We will compare the PAM–13 score change between the intervention arm and the control arm using independent t-test. Data will be log transformed to achieve normality as appropriate. We will also compare the mean change in PAM–13 score in the intervention and control arm in a multiple linear regression model including potential covariates.

Secondary endpoints

We will compare the mean change in EQ–5D–5L index score and visual analogue score, EORTC-QLQ-C30 and BR45 scale scores in the intervention and control arm in simple and multiple linear regression models including potential covariates.
We will compare the mean change in the three HADS scale scores in the intervention and control arm in simple and multiple linear regression models including potential covariates. Based on the continuous overall psychological distress score, patients are classified as ‘distressed’ when they have a score of ≥8, and ‘not distressed’ when they have a score <8. Frequency, percentages and any appropriate 95% confidence intervals of this dichotomization at baseline and 3 months will be presented. Chi-square or Fisher’s exact test will be used to compare the level of distress between the intervention and control arms.

We will present the mean rate of resource utilisation and cost per patient in the two groups by type of resource (primary, secondary and social care) and for total NHS resources used. Simple and multiple linear regression models including potential covariates will compare the mean rate of total resource utilisation and the mean cost per patient in the two groups.

We will describe the average scale scores of the four validated measures in the two groups at the four time points and show graphically the trend in scale scores in each group. We will also compare the mean change in scale scores of the measures across the four time points between the intervention and control arm using a mixed models approach.

To describe O Wise uptake, the average number of times logging in at daily intervals throughout the follow up period will be described and the trend of mean logging in over time will be graphed. We will also show the average frequency of use for each function of the tool over time.

**Sensitivity analysis**

Per protocol analysis will be performed as a sensitivity analysis to assess the impact of contamination on the primary analysis. In the sensitivity analysis, all participants in the
control arm that report using OWise will be excluded. If the sensitivity analysis produces results dissimilar from the primary analysis, we will determine the primary results are not robust and further research is required.

**Missing data**

Missing data of multi-item scales will be handled according to questionnaire guidelines. Where guidelines are unavailable, items will be mean-imputed if at least half of the items from the scale are answered. Descriptive statistics are based on complete case analysis. We will analyse available data before imputation for the groups comparison and use the complete case data as a form of sensitivity analysis.

**Dissemination**

Any protocol modifications will be submitted for approval to the research ethics committee, reflected in the online registration and disseminated by email to site principal investigators and trial coordinators. To mitigate attrition, the coordinating centre will engage participants with newsletters via email or post. These will also discuss any changes to study procedures relevant to participants and results of the study. Each party involved will continue to own the data they collected, i.e. The Royal Marsden will own the clinical and PROM data, Discover on behalf of the NWL data custodians will own the health and social care resource utilisation data, and Px Healthcare will own the OWise uptake data. The statistician and health economists will have access to the final linked trial dataset. There are no plans to provide public access to the full protocol, participant-level data or statistical code. The researchers aim to publish results in a peer-reviewed journal and share via social media and conferences. Authorship will be determined by the owner of the data included in the publication.

**Discussion**

This study aims to evaluate the impact of OWise on patient activation, psychosocial
outcomes and health and social care utilisation. In the face of an expanding mHealth field, robust, comparative studies are vital to understand the impact of such tools on patients. Evaluating mHealth technology available to the public poses a challenge to study design due to the high risk of contamination. To mitigate contamination, we will not disclose the specific tool to participants unless randomised to the intervention group. We felt individual randomisation was appropriate over cluster randomisation as health care providers are not directly involved in the administration of the intervention and O Wise use in the UK is low [43]. Specific items in the questionnaires will measure contamination at each time point. The items will ask patients to identify any supportive care tools or information sources used, including websites and mobile phone applications, with free-text boxes.

Previous work suggests adjusting the sample size for expected contamination [43]. However, no previous literature has reported contamination levels in similar studies. We decided instead to assess the level of contamination before the end of recruitment and increase the sample size if necessary. To assess the impact of contamination on effect estimates, we will also conduct a sensitivity analysis using per-protocol analysis [49]. This will test the robustness of our primary intention-to-treat analysis, the gold standard method for randomised controlled trials [50].

This study will use a new web-based PROM collection tool implemented by the Royal Marsden called PROFILES [44]. Electronic capture of PROMS provides a number of benefits including flexibility for participants, more accurate and timely data collection and reduced time and costs to conduct research [51]. There is also growing evidence for equivalence between paper and electronic PROM collection [52].

This study design may be limited by relying on patient-reports of O Wise use to measure contamination, However, this is unavoidable due to data protection arrangements of the
tool. Participants may also use similar mHealth tools which could confound the results.
The open-ended items assessing contamination will measure the use of other tools and enable us to control for these effects in analysis as much as possible. This study will also be limited by updates of the application. Post-hoc analysis of differences before and after the updates will allow us to assess whether and changes in the application reduce or enlarge any effects or change participant uptake.
This study will allow us to assess whether OWise, a patient-focused mHealth technology, can have an impact on self-management processes, HRQoL and NHS health resource utilisation. With the comparative nature of the study and conduct in the NHS system, this will have broad implications for the adoption of this tool by the NHS in future. If successful, this application can be modified to meet the needs of other tumour groups. This study is also applying new methods in a growing field of mHealth evaluation and can serve as an example for researchers in future.

Trial Status
Protocol version 2.0 31/05/2019 was approved on 11/07/2019 with recruitment pending.
End of recruitment is planned for 30 June 2020.

List Of Abbreviations

UK United Kingdom
NHS National Health Service
PROM Patient-reported Outcome Measure
HRQoL Health-Related Quality of Life
EORTC-QLQ-C30 European Organisation for the Research and Treatment of Cancer—Quality of Life Questionnaire—Core 30
BR–45 Breast Cancer 45
EQ-5D-5L EuroQol 5-Dimension 5-Level

HADS Hospital Anxiety and Depression Scale

Declarations

Ethics approval and consent to participate

The Royal Marsden NHS Foundation Trust is the study sponsor responsible for initiating and managing the study and the coordinating centre. The sponsor will monitor the trial at one year and the end of the study. It was reviewed by the Royal Marsden NHS Foundation Trust and Institute of Cancer Research Combined Clinical Research Committee (CCR4965) and the London-Brent Research Ethics Committee (IRAS250002) and Health Research Authority (19/LO/0725). The Discover Research Access Group reviewed and approved the study on 18/07/2019. The study was registered at clinicaltrials.gov (NCT03866655) on 7 March 2019 (https://clinicaltrials.gov/ct2/show/NCT03866655?id=NCT03866655&rank=1). This research will be carried out in accordance with the Declaration of Helsinki (1996). The study will be conducted in accordance with the conditions of ethical approval. Before participation, all participants must give written informed consent. The planned first enrolment is 1 August 2019.

Consent for publication

Not applicable

Availability of data and material

Not applicable

Competing interests

EL: Nothing to declare
SM: Nothing to declare
JN: Nothing to declare
SS: Nothing to declare
AL: Nothing to declare
KM: Nothing to declare
WvdG: nothing to declare
OH: Nothing to declare

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**Authors’ contributions**

All authors were part of the study steering committee. OH is the principal investigator responsible for scientific leadership and final decision on PROMs selected. EL led study design, writing the protocol and study set up. SM, JN and SS provided clinical expertise for study design and recruiting patients. AL provided expertise for the economic analysis design and provision of analysis. KM led the statistical design of the trial. WvdG provided technical supervision and review of the study design. All authors read and approved the final manuscript.

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Figures
Figure 1

The Individual and Family Self-Management Theory
| TIMEPOINT | Enrolment | Allocation | Post-allocation | Close-out |
|-----------|-----------|------------|----------------|-----------|
| ENROLMENT: |           |            |                |           |
| Eligibility screen | X |            |                |           |
| Informed consent | X |            |                |           |
| Allocation |            | X          |                |           |
| INTERVENTIONS: |           |            |                |           |
| Provision of O Wise |    | X          |                |           |
| ASSESSMENTS: |           |            |                |           |
| Demographics | X | X          |                |           |
| PAM-13 | X | X          | X              | X         |
| EORTC-QLQ-C30 | X | X          | X              | X         |
| EORTC-QLQ-BR45 | X | X          | X              | X         |
| HADS | X | X          | X              | X         |
| EQ-5D-5L | X | X          | X              | X         |
| Clinical diagnosis |            |            |                |           |
| Treatment and hospitalisation data |            |            |                | X         |
| Discover data |            |            |                | X         |
| O Wise use data |            |            |                | X         |

Figure 2
Schedule of enrolment and assessments

Supplementary Files
This is a list of supplementary files associated with the primary manuscript. Click to
download.

SPIRIT_Checklist_Attachment 1.doc
PIS_v2.0_31.05.2019_Attachment 2.pdf
Consent_v2.0_31.05.19_Attachment 3.pdf