Utilizing Consumer Technology (Apple's ResearchKit) for Medical Studies by Patients and Researchers: Proof of Concept of the Novel Platform REach

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Abstract

Medical research suffers from declining response rates, hampering the quest for answers to clinically relevant research questions. Furthermore, objective data on a number of important study variables, such as physical activity, sleep, and nutrition, are difficult to collect with the traditional methods of data collection. Reassuringly, current technological developments could overcome these limitations. In addition, they may enable research being established by patients themselves provided that they have access to a user-friendly platform. Using the features of Apple's ResearchKit, an informed consent procedure, questionnaire, linkage with HealthKit data, and “active tasks” may be administered through a publicly available app. However, ResearchKit requires programming skills, which many patients and researchers lack. Therefore, we developed a platform (REach) with drag and drop functionalities producing a ready-to-use code that can be embedded in existing or new apps. Participants in the pilot study were very satisfied with data collection through REach and measurement error was minimal. In the era of declining participation rates in observational studies and patient involvement, new methods of data collection, such as REach, are essential to ensure that clinically relevant research questions are validly answered. Due to linkage with HealthKit and active tasks, objective health data that are impossible to collect with the traditional methods of data collection can easily be collected.

(J Participat Med 2018;10(2):e6) doi: 10.2196/jopm.9335

KEYWORDS

data collection; HealthKit; platform; PRIDE Study; smartphone; wearables

Introduction

The growing number of smartphones in both the developed and developing world (2014: 2.6 billion; 2020: 5.9 billion) and the rapidly expanding coverage of Long-Term Evolution (4G) networks [1], combined with numerous wearable devices such as activity trackers and smart watches, provide unique possibilities to reach potential study participants worldwide on a device they use multiple hours per day [2]. In addition, current technological developments may enable research being established by patients themselves provided that they have access to a user-friendly platform. For example, in the Health app that is available on iPhones since 2014, health and fitness data collected in other apps and wearable devices are put together in one place through HealthKit. Consequently, Apple introduced ResearchKit in March 2015, an open source framework that enables researchers to create iOS apps for medical research [3]. The first studies using ResearchKit for data collection have recently been published [4-7].

The core ResearchKit framework consists of 3 modules handling informed consent, surveys, and “active tasks”. For the latter, data are collected through the iPhone sensors including the...
accelerometer, gyroscope, Multi-Touch display, and microphone. However, although ResearchKit claims to be an easy-to-use platform for researchers to create research apps, involvement of a developer may in fact be necessary to incorporate all needs. For example, iOS and Swift programming skills are necessary to develop an app for a medical study using ResearchKit. Therefore, we developed REach, a platform that enables both patients and researchers to collect data through an app using the main features of ResearchKit. Its reliability and usability were assessed in a pilot study among postpartum women.

**How the Innovation Works**

REach was developed by the Radboud REshape Innovation Center in cooperation with patients and researchers from various medical disciplines, including epidemiology, health technology assessment, pediatrics, and medical informatics. It consists of two sections: a Web app in which the investigator (patient or researcher) can set up the study, and an app available in the App Store with which data are collected from participants. Using drag and drop, an informed consent procedure, questions, and active tasks may be easily added to a study in the Web app (Figure 1).

A full informed consent procedure, consisting of displaying the consent documents, participant name entry, and the participant’s signature, is available in REach. The core of ResearchKit has already received numerous endorsements as a secure platform because the data are stored highly encrypted, only on the smartphone itself [8]. It is considered one of the most, if not the most, secure platforms available at present. Once the participant signs the informed consent form, the document is available for the investigator as a PDF file for archiving purposes.

Within the Web app, the investigator may build a regular questionnaire with instructions, multiple choice questions, open-ended questions (literal or numerical), rating scales, and date/time questions. Comparable to Web-based questionnaires, validity checks may be included to improve data quality. In addition, HealthKit questions may be added, in which the participant is asked for consent to share already collected data on for example body weight, heart rate, and steps. This enables investigators to easily collect unique and objective data on the health of study participants.

In addition to completing the questionnaire, data may be collected from the participants by having them performing active tasks. A number of active tasks are predefined in ResearchKit, which enable inviting participants to perform activities under partially controlled conditions using iPhone sensors for data collection. These active tasks fall into 6 categories: motor activities, fitness, cognition, voice, audio, and hole peg [9]. Currently, two active tasks are available within REach. With tone audiometry, the minimum amplitude for the participant to recognize the sound is determined. Reaction time uses the smartphone’s accelerometer and gyroscope to collect data on device motion.

Once the study has been set up in the Web app, the investigator may start data collection. Studies in the app can be open to the public or by invitation only. For the latter, the email address of the potential participant should be available. The app may be free or paid, enabling possibilities for crowdfunding. There are no restrictions on the geographical location or number of study participants, but an individual can only participate once in a certain study. Furthermore, the app is available in multiple languages, depending on the language settings of the iPhone (default: English). In the dashboard, the investigator may monitor the status of the study (number of views, number of completed consent forms, and number of completed participations). The resulting data file can be downloaded at any time during or after completion of data collection and imported in statistical software packages.
Pilot Testing

Methods

Data collection through REach was pilot tested in a subsample of participants in the PRegnancy and Infant DEvelopment (PRIDE) Study [10]. All women who delivered a live-born singleton and completed the PRIDE Study’s first postpartum Web-based questionnaire (2 months after the estimated date of delivery) between October 2015 and March 2016 were invited to test REach (n=463). The app was developed by a researcher without needing to consult the developer. Testing of the app involved providing informed consent, completing 10 questions...
comparable to those in the Web-based questionnaire to get more
insight into its reliability, and possible sharing of HealthKit
data. The active tasks were not used in this population.

Using REach, we collected data on the occurrence of pregnancy
complications (“check all that apply” format), estimated date
of delivery and actual date of delivery (to calculate gestational
age), birth weight, birth length, head circumference, birth
defects, and closed-ended questions on mode of delivery,
presentation, and infant sex. The questions in the Web-based
questionnaire on perinatal outcome, including birth weight
(intraclass correlation coefficient [ICC] 0.90) were previously validated [11].

Furthermore, participants were asked to complete a short
evaluation questionnaire, which included the System Usability
Scale [12]. The app and evaluation questionnaire were
administered at least 2 weeks after the postpartum Web-based
questionnaire. To assess reliability, we calculated kappa statistics
and ICCs with 95% confidence intervals for categorical and
continuous variables, respectively. IBM SPSS Statistics for
Windows, Version 22 (IBM Corp, Armonk, NY) was used for
all statistical analyses.

**Results**

A total of 31 women tested the app. The results of the reliability
analyses for the categorical variables are shown in Table 1. In
general, there were very few discrepancies between the data
collected through the app and through the Web-based
questionnaire for pregnancy complications, mode of delivery,
presentation at birth, and infant sex. The relatively high numbers
of false negatives for nausea and vomiting of pregnancy and
extreme fatigue mainly included women who reported these
complication in the baseline questionnaire, which is
administered at the end of the first trimester. Therefore, the
effect of time may play a bigger role than underreporting of
these two complications in the app itself.

One woman reported diagnosis of a birth defect in both the app
and the questionnaire. Agreement between the questionnaire
and app was excellent for birth weight (ICC 1.00, 95% CI
1.00-1.00; Figure 2), but substantially lower for birth length
(ICC 0.73, 95% CI 0.50-0.86; Figure 3). However, this outlier
seemed to be caused by the respondent making a typo in the
questionnaire. Omitting this subject from the analysis increased
the ICC to 0.97 (95% CI 0.93-0.98).

HealthKit data were shared by 11 of the 31 participants (35%);
only the number of steps per day was shared. For the remaining
20 participants, we cannot distinguish between those who
granted sharing of data but had no data available, and
participants who denied permission to share data. Although
insight into this matter would be interesting from a research
perspective, HealthKit does not allow sharing of these data to
avoid information leaks and to protect user privacy.

We received 25 evaluation questionnaires. The participants did
not report problems using the app. The mean score on the
System Usability Scale was 83.9 (SD 10.7), indicating a nearly
excellent level of satisfaction [13]. On a scale from 1 (worst)
to 10 (best), the mean rating was 7.8 (0.7). Only 2 participants
(8%) preferred a Web-based questionnaire to completing the
questions through the app; the majority either preferred the app
(56%) or had no preference (36%).

**Table 1.** Comparison of app and questionnaire data for categorical variables. N/A: not applicable.

<table>
<thead>
<tr>
<th>Variable</th>
<th>App positive</th>
<th>Questionnaire positive</th>
<th>Questionnaire negative</th>
<th>App negative</th>
<th>Questionnaire positive</th>
<th>Questionnaire negative</th>
<th>Kappa statistic</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Pregnancy complications</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Nausea and vomiting of pregnancy</td>
<td>6</td>
<td>5</td>
<td>20</td>
<td></td>
<td></td>
<td></td>
<td>0.68</td>
</tr>
<tr>
<td>Extreme fatigue</td>
<td>3</td>
<td>16</td>
<td>12</td>
<td></td>
<td></td>
<td></td>
<td>0.13</td>
</tr>
<tr>
<td>Gestational hypertension</td>
<td>1</td>
<td>0</td>
<td>29</td>
<td></td>
<td></td>
<td></td>
<td>0.65</td>
</tr>
<tr>
<td>Preeclampsia</td>
<td>0</td>
<td>0</td>
<td>31</td>
<td></td>
<td></td>
<td></td>
<td>N/A</td>
</tr>
<tr>
<td>Gestational diabetes</td>
<td>0</td>
<td>0</td>
<td>31</td>
<td></td>
<td></td>
<td></td>
<td>N/A</td>
</tr>
<tr>
<td>Thyroid disorders</td>
<td>0</td>
<td>0</td>
<td>31</td>
<td></td>
<td></td>
<td></td>
<td>N/A</td>
</tr>
<tr>
<td>Pelvic girdle pain</td>
<td>5</td>
<td>1</td>
<td>24</td>
<td></td>
<td></td>
<td></td>
<td>0.79</td>
</tr>
<tr>
<td>Anemia</td>
<td>1</td>
<td>0</td>
<td>29</td>
<td></td>
<td></td>
<td></td>
<td>0.65</td>
</tr>
<tr>
<td><strong>Mode of delivery</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Unassisted vaginal delivery</td>
<td>24</td>
<td>0</td>
<td>7</td>
<td></td>
<td></td>
<td></td>
<td>1.00</td>
</tr>
<tr>
<td>Cesarean section</td>
<td>7</td>
<td>0</td>
<td>24</td>
<td></td>
<td></td>
<td></td>
<td>1.00</td>
</tr>
<tr>
<td>Breech presentation</td>
<td>2</td>
<td>0</td>
<td>31</td>
<td></td>
<td></td>
<td></td>
<td>1.00</td>
</tr>
<tr>
<td>Male infant</td>
<td>12</td>
<td>0</td>
<td>18</td>
<td></td>
<td></td>
<td></td>
<td>1.00</td>
</tr>
</tbody>
</table>
Figure 2. Comparison of birth weight (in grams) reported by mothers in the REach application and in the Web-based questionnaire (N=31).

Figure 3. Comparison of birth length (in cm) reported by mothers in the REach application and in the Web-based questionnaire (N=29).
Conclusion

In the era of declining participation rates in observational studies and patient involvement [14,15], adding new methods of data collection to the toolbox of medical researchers, such as REach, is essential to ensure that clinically relevant research questions are validly answered. Despite some methodological limitations of the pilot study, including the relatively small sample size and selective participation as only iPhone users could be included, study participants were very satisfied with data collection through smartphones and measurement error seemed minimal. Although no formal validation analyses were conducted in the few other studies in which ResearchKit was used for data collection, data quality was also reported to be high and consistent [5-7].

Due to the linkage with HealthKit and the incorporation of active tasks, objective health data that are impossible to collect with the traditional methods of data collection can easily be collected. However, HealthKit data will probably not be available for the complete study population due to declining to share this information or not using the HealthKit on the iPhone at all, yielding the possibility for selection bias.

REach is currently available through the website of the Radboud REshape Innovation Center [16]. More extensive tests of the platform, including patient-initiated studies, are ongoing and possibilities for platforms for other mobile operating systems, such as Android, are now being explored.

Conflicts of Interest
None declared.

References


9. ResearchKit. Active tasks. URL: http://researchkit.org/docs/docs/ActiveTasks/ActiveTasks.html [accessed 2017-11-02] [WebCite Cache ID 6ugHLYzo]


Abbreviations

**PRIDE Study**: Pregnancy and Infant Development Study

**ICC**: intraclass correlation coefficient