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Ubiquitous Assessment of the Recovery of Cancer Patients Using Consumer-Level Activity Trackers †

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Abstract: Performance Status (PS) variability is a powerful tool to evaluate overall condition, treatment needs and survival chances of cancer patients. Traditionally, its assessment has relied on the experience of oncologists when interpreting results of clinical tests and when interviewing the patients. Meanwhile, consumer-level activity trackers have obtained good results in behavior-change oriented intervention trials and Fitbit devices have demonstrated enough reliability to provide objective data related to physical activity, but the clinical possibilities of the data collected has been neglected. This work presents a system design for ubiquitous assessment of PS by means of objective and quantifiable data from different sources: medical history, self-reported quality-of-life questionnaires and a commercial activity tracker Fitbit Alta HR. The system proposed aims to contextualize and model the recovery process of breast cancer patients during chemotherapy treatment.

Keywords: ubiquitous; e-Health; chemotherapy; cancer; performance status; physical activity; Fitbit

1. Introduction

Performance Status (PS) variability is a powerful tool to evaluate overall condition, treatment needs and survival chances of cancer patients. It measures the ability of patients to perform the activities of daily living by comprehending several dimensions of the disease, such as physical activity performance, quality of life and the toxicity levels of the treatment [1]. The Eastern Cooperative Oncology Group (ECOG) [2] and Karnofsky Performance Status (KPS) [3] scales have served this purpose for decades and they are still widely used among oncologists.

However, the ambiguity and broad dimensions rooted in the definition of PS scales make them prone to subjectivity and interpretation, forcing oncologists to evaluate PS mainly from personal interviews with patients that may be under severe conditions of pressure, anxiety and uncertainty, along with the secondary effects of the treatment [4,5]. This subjectivity and the tight relationship of physical activity with PS [6] are the main reasons for introducing activitytrackers when monitoring the recovery of cancer patients.

Several clinical trials have already used wearable activity trackers in oncology trials, specially with breast cancer patients and survivors [7]. Research-grade monitors like Actigraph GT3X+ or the ActivPAL were the most used, but there is a growing trend in including consumer-level activity trackers like Fitbit devices due to their popularity, reliability and cost-effectiveness. Fitbit devices have been successfully used within the context of behavior change interventions such as healthy lifestyle promotion, physical activity raise and weight control in population with different conditions such

as breast cancer survivors, pregnant moms and patients with type 2 diabetes mellitus [8–10], but the possible clinical relevance of the data collected is still being neglected. Despite the subtle differences interpreting results across the literature, most of them point out that Fitbit devices may provide reliable objective data referred to physical activity, especially within free living environments [11–14].

All this lays down the basis for the system proposed, where commercial activity trackers, namely Fitbit, provide ubiquitous, objective and quantifiable measures of the physical activity of breast cancer patients, thus aiming to reduce biases and subjectivity in PS evaluation.

2. State of the Art

In previous work we addressed automatic monitoring of PS from a broad perspective of cancer patients [15]. In this work we focus in breast cancer patients since they have been widely studied with research-oriented and commercial activity trackers. According to Gresham et al., they represented the 65% of the total amount of oncology trials related to activity trackers between 2005 and 2016 [7]. This over-representation of breast cancer (both patients and survivors) across oncology trials matches their high involvement and implication in their treatment and recovery. Moreover, the overall good acceptance reported (82% median adherence) is similar to the those reported in more recent studies promoting healthy lifestyle behavior in breast cancer patients (92.13% adherence) [16] and survivors (88.13% adherence) [8]. For these reasons, breast cancer patients reveal themselves as the most appropriate group to study the ubiquitous assessment of PS.

Regarding consumer-level activity trackers, Fitbit devices reported better reliability than the one we used in previous work [15,17], however, despite their promising results and growing acceptance in clinical trials, “discretion should be used when considering the use of Fitbit devices as an outcome measurement tool in research or to inform health care decisions” [17]. In this section we discuss some of the most relevant contributions related to the reliability provided by Fitbit devices: steps count, energy expenditure, physical activity levels, heart rate and sleep tracking.

2.1. Fitbit's Steps Count

Steps count is one of the main variables used for measuring physical activity levels, even in oncology trials [7,18]. The good accuracy reported by wrist-worn Fitbit devices in controlled and free-living settings makes this activity trackers ideal for the proposed system [17]. Some trials included in the systematic review of Feehan et al. [17] represent the conditions to be encountered when assessing PS in breast cancer patients, for example, a trial with 22 prostate cancer survivors monitored with Fitbit One during 7 days under free-living settings found no significant differences in steps count compared to GT3X+ ($r = 0.94$) [12]; and 22 healthy women monitored with Fitbit Flex during 7 days found no statistical difference in daily steps count compared GT3X+ [19].

On the other hand, the work of Ummels et al. rejected the validity of 9 commercial activity trackers (Fitbit One and Fitbit Flex among them) for 130 patients with a chronic disease [20]. However, the activities described in the protocol were most of them homework tasks, and very short in duration (19 activities in 28 to 33 min). The short-lasting activities and the implicit low-consistent patterns of movement within them (e.g., 1 min of simulated cleaning of windows, 7 m of walking weaving around cones) may have over-hindered the steps count for the tested devices; besides, not including research-oriented activity trackers impedes any comparison to trials relying on validated monitors like GT3X+. Other works show that the validation problem of consumer-level devices does not seem to be rooted in the chronic condition of patient, but in their overall mobility [17], for example, Alharbi et al. obtained a very high correlation of Fitbit Flex steps count compared to GT3X+ ($r = 0.95$) in a community-based phase III cardiac rehabilitation population [21].

2.2. Fitbit's Energy Expenditure & Physical Activity Levels

There is agreement on Fitbit overestimating energy expenditure and misclassifying physical activity levels (sedentary, light, moderate, vigorous) when compared to GT3X+, especially for moderate

and vigorous activity [12,13,17,19,22]. Winfree et al. have successfully tackled this issue with a Naïve-Bayes classification model that leverages Fitbit Flex data as its input that removes statistical differences between GT3X+ MVPA (moderate-to-vigorous physical activity) and Fitbit Flex MVPA [23]. This approach makes shows that it is possible to extract meaningful knowledge from the original data provided by Fitbit, so this deserves to be studied along with reliable clinical data.

2.3. Fitbit’s Heart Rate & Sleep Tracking

Regarding other measures of Fitbit such as heart rate, there is clear agreement in its low reliability and high sensitivity to artifacts and sensor displacement, so this data should be interpreted carefully [11,17,24]. Sleep tracking is also reported to be systematically overestimated [17], thus it needs to be addressed with caution too.

3. System Design

This section presents a system design based in previous work [15], adapting it to Fitbit devices and breast cancer patients. Figure 1 presents the system design for the experiment proposed. Patients will participate voluntarily through informed consent and always under the protection of General Data Protection Regulation (GDPR) guidelines; data will be anonymized and stored securely within a remote sever located in our faculty laboratory.

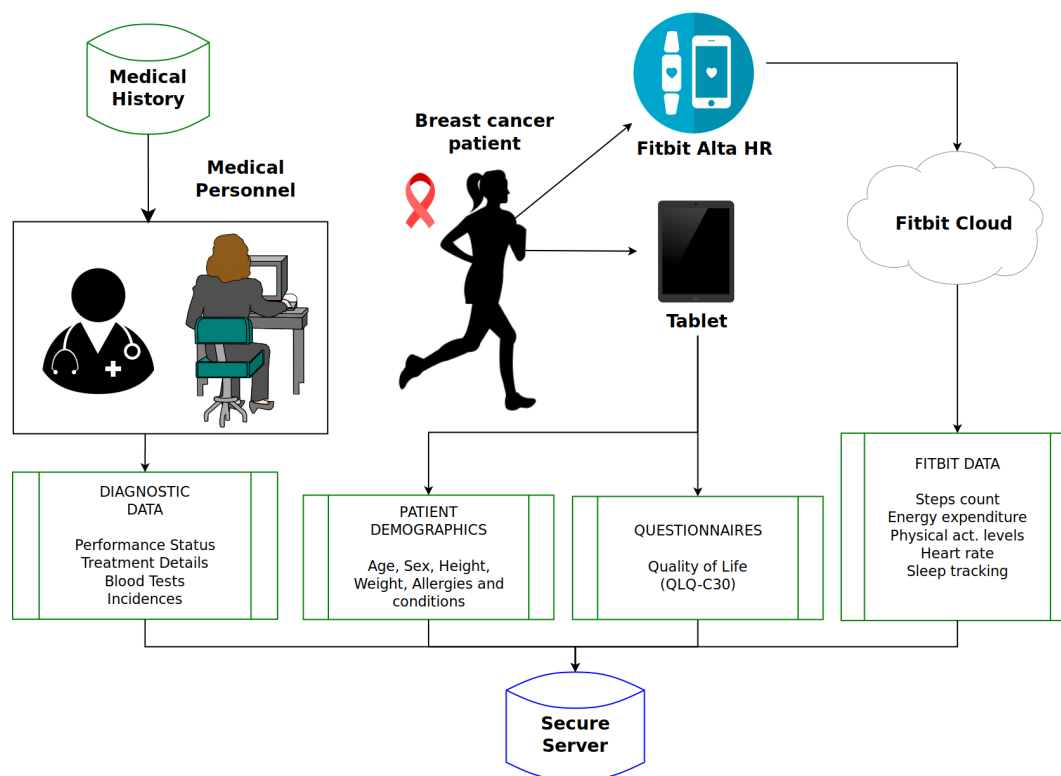


Figure 1. System design.

3.1. Data Sources

There are three main data sources: medical history, self-reported quality-of-life questionnaires and the Fitbit Cloud, which hosts the data collected by a Fitbit Alta HR activity tracker. On the left of Figure 1, there are data from medical records, the most important part of our system: the experts’ knowledge. Clinical PS evaluations of patients will be the main outcome to measure; treatment details, blood tests and incidences will be gathered too during the recovery process to remove factors of confusion like anemia—which is related to sensations of fatigue and tiredness—. The frequency of

these data will depend on the drugs and treatment administered, varying among 7, 14 or 21 days. This data needs introduced into our database manually through web forms connected to a secure web service.

Patient will report its demographics in the start of the trial when registering. They will report their quality of life with the QLQ-C30 questionnaire at the beginning, middle and end of their treatment to contrast self-reported functional and symptom scales. This data will be retrieved by using web forms connected to our database by a web service. The frequency of this data is even lower than medical records, but this methodology mimics previous intervention trials to avoid overwhelming patients with constant questionnaires.

Fitbit Alta HR will monitor the activity of each patient during the entire chemotherapy treatment (minimum of 12 weeks). This device measures every variable discussed in Section 2, so that will contextualize the data from medical history and questionnaires in an objective and quantifiable way. The sampling frequency of the device is very high (compared to the rest of the data) and it uploads all the data directly to the Fitbit Cloud. This cloud provides a REST API that allows data to be collected in our database with granularity from seconds (heart-rate) to minutes (steps count, energy expenditure, physical activity levels, sleep tracking).

3.2. Data Security

Security of patients is the most important issue while handling their sensitive data, therefore, this subsection explains the conditions in which data is stored and collected to meet security standards under GDPR legislation. To ensure physical security the server will be placed at the CITIC-UGR Research Center in the University of Granada (Granada, Spain). Access to the center will be restricted to university personnel, and besides, the server will be located in a room with limited access to the researchers involved in the project. The server will not be used by third persons in a direct nor indirect way. The information stored in the server will be encrypted so that there is no possible way to access the data, not even by stealing the hard drives or by other means. The database collecting the data will have limited access, only available to the research team. Any possible communication between a secure web environment (HTTPS) and the database will be made locally so that information is not visible nor accessible from the outside. Moreover, to ensure maximum security of external access, a firewall will limit the number of available ports for connections, only enabling ports 22 (SSH) and 443 (HTTPS). Both ports will be secured by OpenSSL libraries that ensure the confidentiality and integrity of the information by encrypting data exchanges with the outside.

4. Discussion

This work proposes a system to overcome the limitations of subjective PS evaluation by combining ubiquitous and objective physical activity data, clinically validated self-reported questionnaires (QLQ-C30) and the clinical data within medical records (e.g., PS assessed by oncologists, blood tests, symptoms, etc.). The consistency of Fitbit devices when measuring steps count, and the possibilities offered by energy expenditure and sleep tracking, support a more objective assessment of PS. Moreover, breast cancer patients reveal themselves as a trustworthy group to test the system proposed due to the high adherence and acceptance outcomes along activity-tracker related trials [7].

The inclusion of continuous and ubiquitous data of each patient will provide the ability to study it from a more individual perspective, focusing on tendencies and on the evolution of recovery across time. Diagnostic data will remain always as source of truth despite the critiques of PS scales, since the expert's knowledge is what we are aiming to support. The vision of this work is to ultimately provide oncologists with decision support systems for better assistance of cancer patients with affordable tools like commercial activity trackers.

5. Conclusions

Performance status variability is a powerful tool to evaluate treatment needs and survival chances of cancer patients. Traditionally, its assessment has relied on personal interviews with patients and the experience of oncologists. Consumer-level activity trackers have managed good results among behavior-change oriented intervention trials, but the clinical possibilities of the data collected has been neglected. Fitbit devices have demonstrated enough reliability to supply objective data during the entire recovery process of breast cancer patients. This work presents a system design to evaluate performance status by means of objective and quantifiable data from different sources: medical history, self-reported quality-of-life questionnaires and continuous monitoring of a Fitbit Alta HR activity tracker. Its future implementation will allow us to extract the relevance of each variable in performance status assessment, and to model the recovery process of breast cancer patients during chemotherapy. A trial design with breast cancer patients and the system proposed has been approved by the Ethics Committee of the Virgen de las Nieves Hospital.

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