Exploring attitudes towards use of technology to support stroke survivors living at home: A quantitative and qualitative content analysis study in Spain

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Abstract
Purpose: The aim of this study was to better understand the attitudes towards the use of technology to support chronic stroke survivors in a home-based setting.
Methods: A quantitative study was used on the data obtained from a face to face survey with the sample group, incorporating quantitative statistical analysis.
Results: Participants reported positive attitudes towards using technology for their own independent health management. The purpose of the home-based technology was different: source of information, supporting self-management, pharmacological treatment reminders... and differed according to age, educational level and survivor disability. Installing devices and sharing information remains a challenge.
Conclusions: 100% of stroke survivors living in the community with any type of disability, reported that they would like to use technology as a tool to help improve their health status.

Keywords
Telehealth, stroke rehabilitation, decision making system, self-management, self-care

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Background
Stroke is the leading cause of long-term disability in western countries1,2 resulting in life altering changes for both the stroke survivor and their closest family, sometimes, resulting in profound difficulties and needs. Different locations of the stroke cause different results. A stroke on the right side of the brain can lead to paralysis on the left side of the body, vision problems (and spatial unawareness – “hemispatial neglect”), quick, inquisitive behavioural style (including denial/ unawareness), memory loss, a left sided stroke, on the other hand, may lead to paralysis on the right side of the body, speech/language problems, slow, cautious behavioural style (aware of the problems), memory loss as well. Fatigue is also a common problem after all brain injuries. Remaining symptoms after the stroke might cause difficulties in handling things with two or one hand, difficulty or inability to walk, complications such as falls, fractures, dysphagia, balance and dexterity issues, problems for understanding, speaking, reading, calculating and visual recognition, difficulty in perceiving and/or processing visual or auditory

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information, difficulty to start, follow and/or remem-
ber instructions and sequences. . . . 3

Most strokes are due to ischemic assault which kills
one third of the patients leaving another third with
severe disabilities. Fortunately, as a result of the
improvement of the acute medical treatment, there’s
an increasing number of stroke survivors. 4

Surviving a stroke and living with its effects involves
a long and challenging process for patients and their
families including rehabilitation process, changes in
the physical, social, emotional aspects, and furthermore
the majority of these patients must follow a tight con-

trol of cardiovascular risk factors and life-style changes
to prevent the risk of suffering a recurrent stroke. 5

Limited data exists on the long-term needs of
community-dwelling stroke survivors but for the tech-
nologies, scarcity of data is the norm. Prior qualitative
research (via interviews and focus groups) on end user’s
requirements for e-rehabilitation are published in the
literature however, they do not consider a holistic point
of view of living after stroke neither they focus the
attention on the needs of new technologies in rehabil-
itation (robots, video games, telemedicine . . .) which
should be adapted to the end user’s requirements. 6,7

Nevertheless, there is an opportunity to explore pre-
viously unreported factors, particularly those related to
technology (e-Health/m-Health) such as smartphones
applications for improving healthy lifestyles, enhancing
adherence to pharmacological and non-
pharmacological treatment, empowering the patient
and caregivers with self-management tools, tracking
wearables to promote outdoor exercise or to detect
complications (falls, arrhythmia, patient getting
lost . . .), home-based telehealth, detecting the risk of a
recurrent stroke in real time among other ideas.

As health care providers concerned with quality of
life of stroke survivors, our aim was to better under-

stand the attitudes of chronic, home-based, stroke sur-
vivors regarding technology (e-Health/m-Health). As a
secondary objective, differences between age, disability
and educational level were evaluated.

Material and methods

Research design

For this study, questionnaires were used to capture the
required data to investigate and gather patients’ expe-
riences and interest in potential technology which could
be beneficial for chronic stroke survivors living at home
with a disability.

Ethics committee approval was obtained on behalf
of the STARR project registered in clinicaltrial.gov
(NCT03580642).

Participants and context

Post stroke chronic phase adults (both male and
female) with any type of impairment, living within
the community and able to voluntary participate and
understand the instructions were identified and invited
to participate in the outpatient neurorehabilitation
consultation of a Cruces University Tertiary Hospital,
Spain, from September to December 2017.

The objective and the purpose of the study were
explained and the patient was free to read and sign
the Informed Consent Form before the collection of
any data, they were allowed to voluntarily withdraw
their consent at any moment. Caregivers were welcome
to attend the interview and also sign the consent to
participate. Individuals were excluded if any of the fol-
lowing applied: unable to cope with the interview or
with severe language and/or cognitive impairments
assessed by Mississippi Aphasia Screening Test (<45)
and Montreal Cognitive Assessment (MoCA <26)
respectively. Participants were enrolled by a conve-
nience sampling method. Participants did not receive
any honorarium.

Given that age, educational and disability level have
been reported as confounders in the literature, they
were analysed separately. The sample was divided
and compared the differences between age groups
(group 1 = ≤45 years old, group 2 = 45–65 years old,
group 3 = ≥65 years old), educational (primary, sec-
tary and university education) and disability level mea-
sured by Barthel Index.

The researchers, 2 trained physical medicine rehabil-
itation (PMR) MDs, male and female, with a wide
background in neurorehabilitation. The researchers
were impartial and only had professional familiarity
with the patients. The reasons and interests in the
research topic were explained to the participants.

Questionnaire administration and data collection
procedure

The interview guide and the interview questions, were
designed by researchers at Lund University, Sweden,
CEA, France and Osakidetza, Spain, in collaboration
with patient and care organization partners in the
STARR project. The themes explored by the interviews
were consequences of the stroke, daily activities before
and after the stroke, adherence to treatment and posi-
tion towards technology. The interview technique con-
sisted of a questionnaire (online Appendix 1), including
both close-ended and open-ended questions in one ses-
sion. These questionnaires were translated and admin-
istered in Spanish which is the mother language of the
participants.
The interview was held in a comfortable atmosphere in a consultation of outpatient setting of the Tertiary University Hospital with a face-to-face interview. The questionnaire was completed in 20–30 minute discussions moderated by the researcher. Participants were verbally asked pre-determined questions, to which they wrote down their responses, where necessary additional explanations were given. The results of the questionnaires were not returned to participants for further comment.

Data was extracted from medical records based on demographics, stroke features, vascular risk factors, medical care and planned rehabilitation, including previous functional situation, education level and impairment details.

Identifying information was removed from the documents and written notes were taken when necessary.

Data analysis and statistical analyses

Researchers independently coded the data, developing a formal coding framework and categories which were completed after a second peer debriefing session. During the analysis process, when differences were observed, the researchers carefully examined any potential source of bias ensuring consistency. Researchers duplicated and verified the data in other following peer sessions.

Descriptive statistics were used to present socio-demographic data and medical information. Barthel index was used to categorize stroke severity as independent, mild, moderate and severe. Quantitative statistical analysis was utilised.

The qualitative variables were described in percentages and quantitative variables with median and range.

We analyzed the data with SPSS (version 23.0) statistical software package, testing subgroup differences using chi-square analysis or Fisher exact test. Reported P values are two-sided. Significance level was specified at 0.05.

Results

Participants characteristics

A total of 56 participants, including 22 with mild impairment in communication (14) and/or in cognition (8) fulfilled the inclusion criteria for this study. The majority of the participants were male with a mean age of 67.50 years. No refusal to participate nor dropouts were reported. The main socio demographic factor characteristics and medical information are shown in Table 1.

Consequences of stroke

In terms of disability, all participants suffered a type of impairment, with a final Barthel index of 77.03.

Stroke has a large number of negative consequences on survivors’ everyday life. The participants in study talked about many physical difficulties, for example difficulties with mobility of their upper and/or lower limb as well as a general reduction of the physical activity, balance issues and fatigue.
93% were ambulatory to some extent, although some needed assistance such as a cane, crutch and/or foot-up orthoses.

Study participants also evoked communication problems, cognitive difficulties, anxiety, depression and emotionalism. They expressed a negative feeling about the new situation (Table 2).

**Rehabilitation**

All the survivors needed rehabilitation (RHB) during their stay in the hospital with being 30% discharged to a RHB nursing home.

The most frequent combination of RHB were physiotherapy and occupational therapy. Mean average time of treatment was 9.8 months excluding the botulinum toxin A treatment periods (range 1 to 14 months) (Table 3).

**Current situation**

When describing life **before** stroke, participants often evoked working activities (90% of people in working age), domestic activities such as cooking, sport and hobbies, as well as social activities such as visiting family, friends or taking care of grandchildren.

When talking about life **after** stroke, patients mentioned activities such as dressing one’s self, grooming, eating, cleaning and preparing simple meals.

### Table 2. Consequences of stroke.

| Employment situation previous to stroke (working/retired) | 21 were of working age, (37.5%): 19 were employed. 35 were retired (62.5%) |
| --- | --- |
| Employment situation after stroke | From the 21 people who were in working age, none returned back to work |
| Basal Barthel index: mean | (25–100) |
| – Independent | 54 (96.42%) |
| – Mild dependent | 0 |
| – Moderate dependent | 1 (1.78%) |
| – Severe/totally dependent. | 1 (1.78%) |
| Familiar situation before stroke | |
| – Living on their own | 6 (10.71%) |
| – Living with relatives | 50 (89.29%) |
| o Needed help of a third person | 1 |
| Familiar situation after stroke | |
| – Living on their own | 4 (7.14%) |
| – Living with relatives | 52 (92.86%) |
| o Needed help of a third person | 16 |
| Final Barthel index: mean | 77.03 (10–100) |
| – Independent | 14 (25%) |
| – Mild dependent | 8 (14.28%) |
| – Moderate dependent | 22 (39.28%) |
| – Severe/totally dependent. | 12 (21.42%) |
| Consequences of stroke | |
| – Upper limb mobility difficulty | 41 (73.21%) |
| – Walking difficulties | 39 (69.64%) |
| – Balance problem | 12 (21.42%) |
| – Communication problem | 14 (25%) |
| – Swallowing problem | 3 (5.35%) |
| – Perception problem (ie neglected limb) or attention problem | 5 (8.92%) |
| – Memory loss | 8 (14.28%) |
| – Emotional problem | 6 (10.71%) |
| – Vision problem | 4 (7.14%) |
| – Fatigue | 17 (30.35%) |
| Able to walk independently | 31 (55.35%) |
| Need assistance: | |
| – Wheelchair | 4 (7.14%) |
| – Cane | 14 (25%) |
| – Crutch | 5 (8.92%) |
| – Foot up | 6 (10.71%) |
| Their feeling about the new situation is negative | 17 (30.35%) |
These data show that they need a considerable time to do basic activities of daily life.

Walking was considered a very frequent and important activity after stroke as it gives autonomy and is part of the non-pharmacological treatment to keep the physical activity level and control cardiovascular risk factors (high blood pressure, dyslipidaemia, diabetes mellitus, and obesity). Social activities such as visiting friends and family or taking care of relatives was less frequent depicting life after stroke. On the contrary, they mentioned watching TV, reading activities that were not present in their discourses on life before stroke (Table 4).

**Technologies in stroke survivors**

A vast majority of the participants (91.07%) were familiar with technologies especially with television (78.57%) and cell phone (71.42%). Tablet (17.85%) and tele-assistance (8.92%) were not very popular.

There was strong consensus (92.85%) about the eagerness of using technologies in a health care setting, however their perception of use was different. More than 70% of the participants would like to employ the technology for self-management and for gathering information (causes of stroke, how to identify a new stroke, what to do, different types of stroke, recovery time...). The therapeutic function of the technologies related to health was divided in non-pharmacological (rehab exercises) and pharmacological treatment. More than half would be grateful to use it for rehab treatment exercises (i.e. exercises for upper limb such as prevention of painful shoulder, dexterity of the hand, coordination, safe balance exercises, aerobically exercise program, strength exercise program, stretching spastic muscles...) while only a quarter would like for treatment info such as medication reminders (i.e. “have you taken your medication?” “If response is negative, do not forget to take it before 8:00 pm”). All the survivors wanted to use the technology to improve their condition, increase their autonomy, self-efficacy and consequently their self-management (Figure 1).

Despite having a positive disposal for eHealth, the vast majority (89.5%) would accept it only for certain activities or issues.
Fortunately, a significant majority of stroke survivors affirmed following the medical recommendations and instructions for the pharmacological treatment only 5.35% did not.

With regard to data sharing, the majority of patients of the responders would allow sharing information with the relatives and/or health staff, as shown on Table 5.

Whilst 19.64% would want to keep their data private.

Almost half of the participants would accept installation of devices at home (Table 6).

Subgroup analysis by age. For technology as a disease information source, a dominant positive feeling emerged, as 100% of the younger patients, 86.7% of medium age and 59.4% of older age identified this function as useful. The interest in using technology to support self-management (100%; 87.50%; 67.60% respectively), for rehabilitation exercises (100%, 66.7% and 63.6% respectively) followed similar distribution. However, the distribution of percentages were different for pharmacological treatment reminders (0%, 25%, 27.3% respectively), but most of these differences are not statistically significant.

Similar point of views apply for the readiness to install devices at home 100% of the younger patients, 85% of medium age and 52% of older age ($p = 0.025$) being these differences statistically significant.

Subgroup analysis by educational level and disability level. There was a directly proportional relation between the educational level and the intention of installing devices at home being this tendency similar with the disposal for sharing information with third parties, in other words, those participants with higher level of education, such as university studies, showed greater tendency to install devices and to share information compared to those with no education (Table 7).

There were no statistically significant differences with disability level and technology use, eagerness of using it, functionality and data sharing.

Discussion

This is the first type of study in Spain to obtain detailed information on the experience and potential uses referring to new technologies (e-Health/m-Health) in the chronic stroke survivors in the home, focused on the experience of technologies can offer. Identifying the users’ attitude towards technology can help to define interventions to support best stroke survivors living in the community, reduce dissatisfaction, improve adherence to physical activity exercises, to medication intake and to improve quality of life of patients and caregivers indirectly. There are other domains for instance, everyday living, work, leisure, social support, driving and finances... Although they can also be fulfilled with serious games, tele-rehabilitation, robotic devices, virtual reality, wearables/sensors, tablets, health devices and others.

Our results showed physical, cognitive and physiological problems, which might have a huge negative impact of their daily life. These findings are not new and correspond the reviewed literature.3

Previous studies on long-term unmet needs were concentrated on needs resulting from functional deficits after stroke, such as management of body function, participation in basic and instrumental activities of

| Table 5. Data sharing. |
|------------------------|
| Yes | No | NA |
| Medical doctor | 44 (78.57%) | 11 (19.64%) | 1 (1.78%) |
| Family | 38 (67.85%) | 11 (19.64%) | 7 (12.5%) |
| Friends | 18 (32.14%) | 29 (51.78%) | 9 (16.07%) |
| None | 11 (19.64%) | | |
| All | 3 (5.35%) | | |

| Table 6. Interview answers. |
|----------------------------|
| Already use technological devices | 51 (91.07%) |
| Want to use technological devices in a health care setting | 52 (92.85%) |
| Would like to use technology |
| | yes | no | don't know/depends |
| For disease information |
| | yes | 36 (64.29%) | 15 (26.79%) | 5 (8.92%) |
| For auto-control |
| | yes | 42 (75%) | 13 (23.22%) | 1 (1.78%) |
| Technology devices for treatment |
| | For RHB + exercises | 25 (44.64%) |
| | Pharmacologic treatment control | 9 (16.07%) |
| | Everything | 4 (7.14 %) |
| | Don’t know | 18 (32.14%) |
| Ready to install gadgets at home |
| | yes | 31 (55.36%) |
| | No | 16 (28.57%) |
| | Don’t know | 9 (16.07%) |

| Table 7. Educational level. |
|-----------------------------|
| Info sharing | Yes | No |
| Primary school | 86% | 14% |
| Secondary school | 95% | 5% |
| University studies | 100% | 0% |
daily living, or secondary needs of new socio-familial or environmental factors. It is important to understand the specific needs perceived by stroke survivors for a patient-centered health and social care. Furthermore, the unsatisfied needs perceived by patients may differ from those perceived by healthcare professionals and caregivers. Furthermore, younger survivors potentially could have a higher functional needs to be fulfilled, not only home-rehabilitation exercises but also intellectual fulfillment, work, holiday and family support as they need a higher degree of recovery to participate in the society. Following this line of reasoning, clinicians may utmost focus on the emotional support and their skills to adapt to the new living situation rather than, clarifying repeatedly stroke’s functional recovery process. Definitely, rehabilitation efforts and reintegration on the society remains a challenge in this age group.

In the study, a significant proportion of participants reported be familiar with technology, having a positive attitude towards its use in health-related basis. Notably, all of them wanted to improve their condition. This is not surprising as other studies have shown that patients, especially chronically ill patients, are positive towards being involved in their care and rehabilitation processes.

The results of this study showed differences in the purpose of the technology. Attitude towards using technologies were most prominent in home-rehabilitation exercise program instead of cardiovascular risk factor control, maybe due to sampling recruitment in rehabilitation consultation basis. In addition, installing devices at home is convenient in younger patients (p < 0.05). Undeniable, patients are no longer passive receivers of care. The findings suggest higher involvement in the decision making process as they preferred to choose the aim of using the technology. For example, 76% would like to employ the technology for self-management, 71% for learning information about disease, medium age group in particular. However, when interviewed about treatment, they felt confident in pharmacological management, especially younger patients who showed dissatisfaction of pharmacological treatment reminders.

Nonetheless, it has been generally acknowledged for years that non-adherence rates for chronic illness regimens and for lifestyle changes are around 50%, admitting the success of a medical treatment is largely determined by compliance. Given that non-compliance of drug treatment may lead to complications is crucial to control the cardiovascular risk factors and to prevent a recurrent stroke. To foster adherence, patient centered approach is essential, listening to survivors and care-givers to discover the unmet needs and what is important for them.

For data sharing, results showed a notable variation in opinion which need to be evaluated in the context of an application, taking into consideration the aim of the information sharing.

It is not completely clear whether and how technologies can be implemented in stroke rehabilitation in different settings but, it is worth considering flexibility in use of the technology so that it could allow personalization to varying abilities, interests and situations. These factors associated with unmet needs could help guide policy decisions, particularly for tailoring care and support services provided after discharge in home basis.

STARR (The Decision Support and self-management system for stroke survivors) project and the system developed in it are targeting the self-management of stroke risk factors. Existing predictive models of stroke risk will be used, a modular, affordable, and easy-to-use system to inform stroke survivors will be developed aiming to inform about the relation between their daily activities and the risk of having a secondary stroke leading to better prevention and reduction of secondary strokes and to a more efficient participation of survivors in medical decision-making process.

STARR project could be a solution to manage the attitude towards technology unsatisfied need considering age, disability level, educational level, home environment. The system should be flexible and able to engage patient’s participation increasing internal and external motivation.

Limitations
Some limitations exist in the study, including modest sample size, aphasic population is not evaluated and needs to be considered. Those without rehabilitation consultation review such as the institutionalised or transient ischemic attack were not studied and may have different unmet needs. Subgroups are not homogeneous, it was not feasible to obtain young stroke survivors as the prevalence of this disease is not common within this age group. Time delay between data collection, analysis and publication exits, however, based on our experiences, and those gathered from colleagues and patients, the attitudes towards technology and responses to technology change at a slower than the technology itself and vary more on a generation basis.

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Supplemental material
Supplemental material for this article is available online.

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