Signs, meanings and practices of people living with human t-cell lymphotropic virus type 1 or tropical spastic myelopathy

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Abstract

Background: Human T-cell lymphotropic virus type 1 (HTLV-1) spreads silently in the world’s population and causes several syndromes. Among these, HTLV-1 associated myelopathy, also called tropical spastic paraparesis (HAM/TSP), affects the nervous system. It causes sensorimotor losses, spasticity, muscle weakness, voiding and sexual dysfunction, pain, and balance disorders. There is limited knowledge of the feelings, experiences, and coping mechanisms associated with this neglected disease. The objective of the present qualitative study was to investigate the signs, meanings, and practices of people with HAM/TSP, through narratives obtained from focus groups and individual semi-structured face-to-face interviews.

Results: Thirty-eight individuals diagnosed with HAM/TSP participated in the study. The following categories and subcategories emerged from the participants: Signs—physical signs, symptoms, and discovery of the disease; Meanings—reaction to diagnosis and knowledge of disease, fears, and expectations; Practices—daily life, leisure, religious, and treatment activities.

Conclusions: People with HAM/TSP suffer from symptoms that limit their social participation, and they are affected by complex and multidimensional feelings. This awareness can contribute to the implementation of public policies—focused on the real perspective of these patients—that provide more directed, empathic, and harmonious care for these individuals.

Keywords: Tropical spastic paraparesis, Human T-lymphotropic virus 1, Qualitative research, Psychosocial approach

Introduction

Human T-cell lymphotropic virus type 1 (HTLV-1) was the first retrovirus identified in humans [1]. Its infection occurs either vertically (breastfeeding) or horizontally (through sexual intercourse, blood transfusion, or contamination with piercing objects) [2]. It is estimated that there are 5 to 10 million infected people worldwide [3], of whom more than 2 million are in Brazil [4]. Salvador is the city with the highest seroprevalence in the country, with 1.76% of the population affected. The majority of infected people are female with black skin color and low socioeconomic status [5].

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feelings, perspectives, and ways of coping, can help to uncover questions that quantitative research does not achieve. Indeed, although much is known about the biomedical aspects of HAM/TSP, little is known about the effect of the disease and therapeutic protocols on psychosocial parameters [13]. Assessing the perception of affected people can reveal subjective aspects that affect the results of therapeutic interventions and clinical evolution. It can also help establish grounds for health actions recommended by the World Health Organization (WHO) for neglected diseases [14].

Qualitative research aims to gather systematic knowledge on the ways of thinking and acting for specific populations. The analysis of signal systems, meanings and practices is grounded in the study of patients’ narratives about their experience with a disease. Based on interpretative anthropology, this analytical model investigates the subjective experience of illness and treatment in different sociocultural contexts [15]. Thus, the main objective of the present study was to investigate the signs of the disease process, meanings attributed to the experience and treatments of the disease, and reactive and self-care practices, expressed in the cultural networks of people living with HAM/TSP.

Methods
Study design and participants
This study was a descriptive and exploratory examination with a qualitative approach. The study sought to systematize narratives from participants via focus groups (FGs) and face-to-face semi-structured interviews. Information analysis was based on the system of culturally constructed signs, meanings, and practices. It was conducted at the Advanced Physiotherapy Clinics (CAFIS) of the Teaching Assistance Ambulatory, Bahiana School of Medicine and Public Health, located in Salvador, Bahia, Brazil. The population was comprised of patients diagnosed with HAM/TSP who were enrolled in the Integrative and Multidisciplinary Centre for Assistance and Research to Family and HTLV Carriers (CHTLV) of the Bahiana School of Medicine and Public Health in 2014.

Recruitment and sampling
Enrollment in the study was intentional and voluntary; it occurred via an invitation that was presented to CHTLV participants who were participating in another research project [16]. Fifty-six people who were part of the clinical trial were invited to participate in the present study. Thus, 38 individuals were included in the final sample. They were distributed in the FGs, represented by the letter P and numbered from 1 to 38 (which represented each participant).

Participants were included if they had a confirmed diagnosis of HTLV-1 infection by enzyme-linked immunosorbent assay (ELISA) and western blot exam, clinical diagnosis for HAM/TSP according to Castro-Costa et al. criteria [17], were enrolled in CHTLV, and were above 18 years of age. Patients were excluded if they had difficulties understanding questions or communicating.

Procedures
Data collection was performed by a researcher with experience in the FG and semi-structured interview techniques. The main researcher (the facilitator) is a physiotherapist, a master in Human Development and Social Responsibility for 25 years with 32 years of clinical practice. The facilitator was accompanied by two undergraduate students from the Physical Therapy course at the Bahiana School of Medicine and Public Health. The students were trained by the researcher. One served as an observer, responsible for making notes of aspects observed in the dynamics of the FG, and the other undergraduate student was responsible for audio recordings. The researcher spoke in person to the participants to invite them to a FG. On the eve of each FG, the researcher confirmed the time and place of the meeting via telephone.

A total of 11 FGs were created, out of which 6 were categorized as pre-training, meaning they took place after the first evaluation for the RTC. These 6 FGs were constituted as follows: FG1 (1 man and 4 women); FG2 (2 men and 3 woman); FG3 (2 men and 1 woman); FG4 (4 men and 4 women); FG5 (2 men and 4 women); FG6 (2 men and 3 women). Some participants reports that they had no time, transportation, company, or financial availability or they had other commitments to medical consultations and/or treatments at the same time as the survey. After training and the last evaluation at the ECR, 13 participants of the pre-training group attended the 5 post-training FG in addition to 3 new participants. These groups were formed as follows: FG7 (1 man and 3 women); FG8 (3 women); FG9 (2 men and 1 woman); FG10 (2 men and 1 woman) and FG 11 (3 women). FGs were performed for approximately 1.5 h.

FGs are group discussions, organized to explore a specific set of issues, where participants can interact, explore each other’s arguments, and express topics they consider important [18]. It also provides the interpretation of beliefs, values, concepts, conflicts, confrontations, and points of view [19]. These features are why it was one of the techniques selected for this research.

The FG environment should be pleasant, comfortable, and warm. Thus, we opted for the use of incense, which lightly perfumed the room, relaxing music with sounds of water and nature, and snacks throughout the meeting [20]. Each FG was conducted in a private setting in the
physiotherapy clinic (CAFIS). The location of the room allowed the meetings to proceed without external interference. This design facilitated the debate and ensured privacy, comfort, easy access, and a neutral environment [21]. The availability of the chairs in a circle allowed face-to-face interaction, good eye contact, and the same field of view for everyone [21]. After clarifying the objectives of the study and providing written informed consent, the participants were invited to relax and the procedures recommended by Iervolino and Pelicione [22] were adopted. Relaxation and integration techniques were used prior to the application of the method. Semi-structured interviews were conducted in a private, air-conditioned room at the CAFIS. The guide for the questions used in the FG and interviews are described below.

**Guiding questions of the pre-treatment FG**
- How did you discover the disease?
- How did you get the diagnosis?
- What were the symptoms?
- What is the relationship of exercises to your health problem?
- What do you expect from participation in this project?
- What reasons could prevent your participation in this project?

**Guiding questions for post-treatment FG**
- Were your expectations reached?
- How was your participation?
- Did you comply with the protocol?
- Did you notice changes in your condition?

Semi-structured interviews were also performed with 12 participants, out of which 8 were FG participants and 4 new members that participated in the RCT control group. 7 men and 5 women were interviewed. Twelve interviews were sufficient to reach data saturation because the information generated in the last interviews was repeated and no new code emerged. The average duration of the interviews was approximately 1 h. Semi-structured interviews were conducted in a private, air-conditioned room at the CAFIS. Semi-structured interview questions: How is your day-to-day activity? How do you do your exercise as part of your day-to-day routine?

**Analytical approach**

The analysis of the systems of signs, meanings and practices, investigates experiences of the illness and healing process in different socio-cultural contexts [15]. After the focus group transcripts, interviews, and field notes were carried out by the main researcher, the transcribed information was read and reread by 2 researchers separately, thus increasing reliability. Content analysis techniques were performed manually to detect units of meaning and nexus categories. After identifying the codes, the authors met to discuss the findings of the themes related to the signs (category refers to symptoms that indicate the disease or suffering described in the patients’ narratives), meanings (category refers to how the person perceives the problem and how s/he constructs, in their imagination, the received diagnosis), and practices (category refers to the creation of strategies to solve a specific problem of myelopathy). After the data was revised multiple times and a consensus was reached, the findings were compiled, and a matrix describing the most representative themes was created in the results section.

**Results**

The sample consisted of 38 people with HAM/TSP; the mean age was 54.2 ± 10.28 years. The majority of participants were women (57.89%), of African descent (47.37%), married (42.10%), had completed primary schooling (36.84%), and of the average socioeconomic class (55. 26%). The mean disease duration was 11.45 ± 8.31 years and mean body mass index (BMI) was 24.67 ± 3.21 kg/m2. Twenty (52.64%) participants did not practice physiotherapy regularly and 23 (60.53%) used a walking or gait devices (Table 1).

From the analyses of the information collected in the FGs and interviews, the following categories emerged: signs—physical signs, symptoms, and discovery of the disease (Fig. 1); meanings—reaction to the diagnosis, knowledge about the disease, fears and expectations (Fig. 2); practices—activities of daily living, leisure activities, religious activities, and treatment (Fig. 3).

**Signs**

The Signs category refers to symptoms that indicate the disease or suffering described in the patients’ narratives (Fig. 1).

Most participants shared the same symptoms of HAM/TSP, especially pain, pelvic floor dysfunction, weakness in the legs that led to loss of balance and falls. In the present study, dermatitis and uveitis were mentioned by only 1 participant each. Regarding discovery of the disease, most participants discovered the infection through blood donation, followed by prenatal examination and sexual partner examination. Loss of functionality also led patients to seek medical services.

**Meanings**

The category Meanings refers to how the person perceives the problem and how s/he constructs, in their imagination, the received diagnosis.

Fourteen respondents reported feelings of sadness and depression upon receiving the HAM/TSP diagnosis. Three patients felt the desire to die, feelings that demonstrate suffering and anguish permeate the imagination of these people. In contrast, 6 participants adopted a resilient...
stance as a way to adapt to change and overcome adversity.

Among the fears and expectations, most respondents cited marital problems, financial loss, loss of productive capacity, need for family support, fear of people’s reaction, and fear of falling. They also welcomed the potential benefits of health care and expressed hope about research and treatment. We observed the presence of family support, mainly from mothers and wives, in most of the narratives. However, in some cases the support came from strangers.

Practices
The Practices category refers to the creation of strategies to solve a specific problem of myelopathy.

Regarding the practices of the exercise programme, all respondents reported greater autonomy and balance. Obstacles such as lack of self-discipline, financial problems, low levels of education, walking difficulties, and cultural factors make it tough to adhere to exercise protocols. Ten participants indicated a preference for performing exercises in the clinic, even though they knew how to do the exercises alone. According to them, in addition to socialization, working in the clinic increases their confidence in safely, correctly executing the exercises. There is a need for multidisciplinary rehabilitation centers for outpatient treatment of individuals with HAM/TSP that are comprised of professionals prepared to promote care for this population.

Nine interviewees indicated that going to church, regardless of religion, is an act of exercise of faith and socialization. Preaching was also cited by 5 patients. Praying was another way of contacting the Divine. Only 1 interviewee, of the African religion Candonblé, reported talking to his orixa as a way of exercising faith and asking for problem-solving advice.

| Variables                                      | N = 38 | N (%)        |
|-----------------------------------------------|--------|--------------|
| Age, years (M ± SD)                           | 54.20 ± 10.28 |
| Sex, Female                                   | 22 (57.89)% |
|      Male                                     | 16 (42.11)% |
| Skin color, Black                            | 18 (47.37)% |
|      Mixed race                              | 17 (44.73)% |
|      White                                   | 3 (7.9)%   |
|      Yellow                                  | 0 (0)%     |
| Socioeconomic class, A                       | 1 (2.63)%  |
|      B                                       | 3 (7.89)%  |
|      C                                       | 21 (55.26)%|
|      D                                       | 12 (31.59)%|
|      E                                       | 1 (2.63)%  |
| Marital status, Single                       | 10 (26.32)%|
|      Married                                  | 16 (42.10)%|
|      Separated                               | 3 (7.89)%  |
|      Widowed                                 | 9 (23.68)% |
| Education Level, Illiterate                  | 1 (2.63)%  |
|      Elementary School Incomplete            | 10 (26.32)%|
|      Elementary School complete              | 14 (36.84)%|
|      Secondary Education                     | 7 (18.42)% |
|      Higher Education                        | 6 (15.79)% |
| Walking or gait devices, Does not use         | 15 (39.47)%|
|      Uses                                    | 23 (60.53)%|
| Physiotherapy, Yes                           | 18 (47.36)%|
|      No                                      | 20 (52.64)%|
| Disease duration, years (M ± SD)             | 11.45 ± 8.31|
| BMI, kg/m² (M ± SD)                          | 24.67 ± 3.21|

*Mean, SD Standard deviation; Social Class A and B = high, C = average; D and E = low (parameters IBGE, Brazil); BMI Body mass index*
Discussion
General aspects of the sample
The sociodemographic characteristics of the participants are similar to prior reports, in which most of the samples were composed of women [2, 3]. Our data further confirmed that the population most affected by HAM/TSP has lower education and socioeconomic levels, is of African descent, and is married [4]. Most of the participants were middle-aged adults, which suggests a period of latency of the virus for several decades [6].

The median time of clinical manifestations varied from months to decades, which is a finding consistent with previous studies [4]. The use a walking or gait devices was present in more than half of the participants. This data highlights the pronounced functional disability, as reported in previous studies [12, 23]. Despite this incapacity, the majority of the interviewees did not perform physiotherapy, a finding similar to a previous study [10]. The benefits of rehabilitation programs for this population are evident [9–11]. In the arsenal of techniques employed by physiotherapists, exercises are the main modality used to improve functional capacity and quality of life. The positive effects of home exercise programs underscore the prospect of assisting this population in today’s contexts [24].

Signs
The highest reported symptom was daily pain, especially in the lower back and lower limbs [7, 8], which is data consistent with previous studies [24–26]. Pain is the principal complaint of those affected by the HAM/TSP and causes significant loss of quality of life [27]. Pelvic floor dysfunction was also reported by participants, especially women [28, 29]. This situation interferes with social life because loss of urine, unpleasant odor, and need for absorbents indicates the fragility of these people. Additionally, the urgent need to urinate several times at night interferes with sleep quality and impairs sexual performance, self-care, and the willingness to perform various activities, all of which reduce self-esteem and self-confidence [30].
Weakness in the legs, loss of balance, and falls were also cited as frequent symptoms according to related literature [7]. The risk of falling is a public health problem because the consequences bring great suffering and increased expenses resulting from surgeries and hospitalization [31, 32]. These issues cause more withdrawal social behavior.

| Reaction to Diagnosis                     | Participant’s voice                                                                 |
|------------------------------------------|-------------------------------------------------------------------------------------|
| 1) Indifference (2)                      | “Knowing I was calm, because I did not know what it was, there they explained [to] me and I was nervous.” (G6P31); “It did not shake me.” (G6P32) |
| 2) Sadness (14)                          | “I was very sad […] I was really sad, (pause), horrible.” (G2P6); “At first it was very difficult, I went into depression, I did not want to see anyone, I did not go anywhere.” (G1P3) |
| 3) Desire for death (3)                  | “[…] I came to SARAH one day, I do not even like to remember, (choking voice) I was looking like I was throwing myself under a car.” (G2P6) |
| 4) Resilience (6)                        | “[…] My illness I have already accepted, the good thing for us is to accept because otherwise it is worse, if we do not accept it is not helpful.” (G8P73) |

| Knowledge about Disease                  |                                                                                   |
|------------------------------------------|-------------------------------------------------------------------------------------|
| 1) Unawareness of HTLV (8)               | “[…] I did not even know what that was, I had never even heard of it, when I heard about HTLV, I thought I had AIDS, it’s confusing, they’re related.” (G1P2) |
| 2) Searching information on the Internet (4) | “[…] curious as I am, I searched the Internet to really know what this was.” (G10P35) |
| 3) Information in reference Centres (12) | “When I was in SARAH I learned to extract urine every 6 hours (catheterisation).” (G1P3) |

| Fears and Expectations                   |                                                                                   |
|------------------------------------------|-------------------------------------------------------------------------------------|
| 1) Conjugal problems (7)                 | “I discovered in 2001, it was precisely at that time that I got divorced […] but the greatest impact was not to know about the disease, because from that point on came the separation (pause, cry and silence).” (G6P14) |
| 2) Financial losses (1)                  | “I think work was the joy I had, without work I was swept off the floor, retirement is little in comparison to what I earned, I earned a lot more.” (G7P33) |
| 3) Loss of productive capacity (1)       | “[…] it was a thump when the INSS expert said that he was going to send me to a reorganisation in the administrative sector, which I would no longer teach.” (G6P14) |
| 4) Need for family support (8)           | “[…] today my quality of life has improved a lot and since I have a family that gives me support, I have daughters, which helps a lot.” (G2P2); “[…] my wife is wonderful, takes good care of me, I have nothing to complain about.” (G3P1) |
| 5) Fear of people’s reaction (10)        | “People in general have a prejudice of everything: social, racial, cultural and because of this, I do not want to go through this, because I know that if people are prejudiced with me I will suffer, and that is where my immunity will go.” (G10P29) |
| 6) Fear of falls (5)                     | “Because the pavements all are broken, we can slip.” (G1P5); “I had to leave work because I was not taking any more because of the falls, I took a lot of falls.” (G1P3) |
| 7) Reception of the health team (20)     | “The distance does not matter; the important thing is the affection that you give to us.” (G9P34); “I cannot speak all the good that the staff did for me, so that was helping me […]” (G10P29) |
| 8) Hope with research and treatment (10)  | “To maintain balance, […] I am sure that according to the teachings and guidelines the tendency is to even improve.” (G1P2); “Improve in pain, in weakness.” (G2P3) |
| Categories and subcategories (number of participants) | Participant’s voice |
|-----------------------------------------------------|---------------------|
| **Daily Life Activities**                           |                     |
| 1) Domestic practices (15)                          | *"I clean the yard, I make food, I do everything, I have a 4-year-old son, I play basketball, I ride a bicycle, I have a dog, I have a large acrylic for I go to the market, I shop, I do everything." (EP36)* |
| 2) Going to the street (2)                          | *"[...] I eat lunch on the street, I watch television, I stay on the computer, I live alone, I do not do very of this. I do not like it very much and because I am not able anymore (domestic activities)." (EP35): "I am retired. I do not do anything. I live alone. I have a person who makes food, cleans the house and everything else." (EP36)* |
| 3) Hiring employees (2)                             | *"Sometimes I go out, I do something to complement my salary, I like to stay in the street. I do not like to stay in the house anymore, if I stay at home, I just think of foolish things, in this agony. I never got married." (EP35): "The thing I most do is go to the market. Every time I stretch I go to the market, because it’s physiotherapy for me." (EP36)* |
| 4) Difficulties with collective transportation (6)   | *"For me it will be transportation, because transportation is difficult." (G1P10): "I myself when it’s raining can’t imagine coming here, because it’s a difficult time for us to come." (G1P5)* |
| **Leisure Activities**                              |                     |
| 1) Sports (2)                                       | *"I go out with my children, ride a bicycle and take my son and everything by bicycle." (EP17): My fun is more the beach, I swim, I take my son to swim, sometimes surfing." (EC7)* |
| 2) Culture (6)                                      | *"I go to the beach, I go to carnival, I go out to dance, I try to have fun, I travel during Saint John festivities, I dance forró and samba." (EP28): "Ah, I like to cross-stitch." (EP7): "Sometimes I go in the machine, sometimes I go to the sewing machine. I make crafts with scraps of cloth." (EP37): "I travel inland." (EP12)* |
| 3) Entertainment (11)                               | *"One time or another there is something to read, I can have a movie to watch, who has a phone and a computer does not have solitude." (EP7)* |
| **Religious Activities**                            |                     |
| 1) Go to church (9)                                 | *"I am evangelical, I go to church, I am part of the CWA, which is Christian Women in Action, every Sunday morning, I go to church." (EP2): "I really like going to church." (EP6)* |
| 2) Prayers (2)                                      | *"Agreed, I make my prayers." (EP37): "Read the word of God." (EP5)* |
| 3) Preaching (5)                                    | *"[...] as I am a witness of Jehovah, I go forth, I preach." (G3P1): "I am evangelical, I even preach the gospel." (G3P11)* |
| 4) Consultation with spiritual guide: orixá (1)      | *"I have a candomblé yard [...] My Orixá sent me to see a doctor with his head, he cannot speak. For 8 years he’s been trying to tell me about it, put it in my head." (G5P2)* |
| **Treatment**                                       |                     |
| 1) Improvement in autonomy (5)                      | *"[...] I feel more willing, I go out myself, I have been to Faria de Santana alone, I thank God." (G9P12): "[...] when I do physiotherapy I feel stronger, more excited, right, because it is a hope even though physiotherapy will not cure the disease, it will make me much better." (G1P2)* |
| 2) Improved functionality (5)                       | *"You do not need anyone, I do it myself." (G10P34): "You do not need a physiotherapist [...], I do every other day." (G10P17): "To me it does not matter." (G10P1)* |
| 3) Improving self-esteem (14)                        | *"When I do, I feel like there’s an improvement, it’s important to me, it’s everything, that’s all. My greatest improvement is due to exercise." (G4P19)* |
| 4) Lack of discipline (4)                           | *"It is laziness, it is embarrassment, it is emotional problems, it is family problems, it is religious problems, it is economic problems, in fact, everything influences." (EP7): "There is a little bit of laziness." (G1P5)* |
| 5) Preference for outpatient treatment (10)          | *"[...] I do but I think that in the presence of the physiotherapist is much better." (G7P5): [...] physical therapy I think is good, but really doing it here we feel better, because it has people to correct, encourage us [...] would do it alone, but I want to come here too, because it encourages more." (G6P12)* |
and decrease the ability to perform day-to-day work and activities that were previously executed. Furthermore, dermatosis has been documented with a high frequency in symptomatic [33] and asymptomatic individuals [8, 34]. Genital candidiasis, associated in most cases with urinary incontinence [35], was also reported. The association between ocular manifestations and people infected with HTLV-1 has also been reported [36]. Although the literature evidences the presence of dermatosis and eye problems in patients with HAM/TSP, these symptoms were rarely reported in the present study.

The discovery of the disease was casual in general. These findings indicate the need to include ELISAs in routine basic care for the early detection of HTLV-1 in endemic areas, such as Salvador [5]. Blood banks have been the main detection site, thanks to Ordinance No. 1376 of the Ministry of Health [37]. The prohibitive cost of testing is a limiting factor for screening in economically disadvantaged countries [38]. A study conducted in Japan showed that through public policy actions, with the insertion of serological testing in pregnant women and interruption of breastfeeding of seropositive women, vertical transmission was reduced from 20 to 3% [39].

Meaning

Most respondents shared the presence of mental disorders, in the face of disease progression, as well as loss of the ability to work and enjoy leisure time. The reaction to the diagnosis corresponds to the stages of mourning described by Elizabeth Kubler Ross [40]. Acceptance of the disease does not occur immediately after diagnosis. Untroubled acceptance will usually only occur after decades, when the patients become more adherent to the therapeutic programs [41].

Health professionals and society in general have a profound lack of knowledge about HTLV infection, and this knowledge gap has implications for both diagnosis and care practice. The fact that only 1–5% of the infected develop symptoms makes it difficult to implement public policies aimed at this population and perpetuates the spread of the virus [8]. Indeed, HTLV is an “invisible” virus; its spread generally occurs silently in the population [41]. Coping with the disease should be approached in the biopsychosocial model recommended by the WHO [42], namely by a team aware of the psychological stages expected after the diagnosis and who can guide the minimization of adverse reactions. The rise of the Internet has favored health literacy [43]. However, the number of websites that do not guarantee quality information is still significant. Only limited content is available on the HTLV-1.

Health centers are sought in the hope of finding other practices that the one day a cure will be discovered [44]. HTLV-1 is a sexually transmitted disease, thus often affecting marital relationships. Suffering often stems from a sense of betrayal due to a partner’s extramarital affairs and/or a lack of encouragement in seeking new partners due to fears of infecting others. Issues include loss of libido, perineal hypersensitivity, vaginismus, pelvic floor hypertonia caused by neuro-motor changes, and impaired sexual activities and pleasure [30]. In addition, premature menopause is common in infected women [28]. This complex scenario reduces the quality of life. Socio-educational actions are fundamental for the practice of protected sex and reduction of the number of sexual partners that increase the risks of contamination.

Another factor involved in chronic diseases is the commitment of financial resources of patients and their families. The transition from productive worker to retired due to disability markedly reduces wages and consequently lowers the socioeconomic status [45]. The most affected individuals are frequently the family providers; changes in social roles become necessary. In addition to a reduction in the ability to provide for her/his family, symptomatic people often become dependent and vulnerable [12]. Spending on diapers, transportation, and exams increases. These purchases, coupled with reduced income, require substantial family rearrangements and decrease the purchasing power of all involved.

Loss of days at work, medical leave, and early retirement generate high costs for the entire society [45]. The subjects often feel unable to contribute socially and feel shame and fear of bothering others, which generates great suffering and social exclusion. The families undergo major changes in their composition [46]. They are the primary source of support for HTLV-1-seropositive people. The family should seek support, security, affection, and respect, all of which are fundamental elements for better coping.

Practices

Individual or group exercise programs, at the clinic or at home, are recognized as useful for improving function in this population [9–11]. However, difficulties with self-discipline limit the implementation of home programs in Brazil [10].

The chronic degenerative evolution of HAM/TSP leads to the need for the use of walking aids, such as canes, walkers, and wheelchairs and total dependence on care from others in the final phase of the disease [23, 27]. This condition reduces recreational and sports activities, which lead to physical inactivity and increase the risk of comorbidities [31]. Cell phones and computers were reported as important forms of leisure.

It is also clear that spiritual practice affects the health of people with HTLV. Although religious diversity in Brazil demonstrates that the country has become secular, the search for a religious meaning for practical
experience is fundamental to addressing health problems. In the present study, most women sought to exercise their faith in various ways while men were in the minority. Instead, they chose to pray. Spirituality is part of the constitution of all men, regardless of any spiritual experience [47].

Implications for clinical practice and future research
This study is the first qualitative research that involves people with HAM/TSP, assisted at CTHTLV. The knowledge gained from this study by authorities and health professionals may promote greater understanding of the needs that permeate this population. This awareness can contribute to the implementation of public policies, focused on the real perspective of these patients, through more targeted care that is empathic and harmonious for these individuals.

Future research should aim to solve the problems presented in the current study, including accessibility, public policies, knowledge about the virus, physiotherapy, and leisure, among others.

Conclusions
It can be concluded that people with HAM/TSP suffer from symptoms that are limiting their social participation. They are affected by complex and multidimensional feelings. Activities of daily living, including leisure, spiritual, and therapeutic assistance, help in coping with this neglected health condition. Public policies should be implemented to allow an integrated multidisciplinary team to minimize the effects of this infection on the quality of life of these people.

Abbreviations
ADAB: Assisting teaching ambulatory; CTHTLV: Integrative and Multidisciplinary Center for Assistance and Research to Family and HTLV Carriers of the Bahiana School, Salvador; CATIS: Advanced physiotherapy clinic; EBMSP: Bahia School of Medicine and Public Health; ELISA: Enzyme-linked immunosorbent assay; HAM/TSP: Human T-cell lymphotropic virus type 1 associated myelopathy/tropical spastic paraparesis; HEMOBA: Bahia State Hematology and Hemotherapy Foundation; HTLV-I: Human T-cell lymphotropic virus type 1; RCT: Randomized clinical trial; SARAH: Network of rehabilitation hospitals; WHO: World Health Organization

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Authors’ contributions
Study concept and design contributors were GOS, MP and KNS. Data analyses and interpretation were performed by GOS, AML and AVG. The manuscript was written by GOS and KNS and revised by all authors. The authors read and approved the final manuscript.

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Competing interests
The authors declare that they have no competing interests.

Ethics approval and consent to participate
The project was approved by the Research Ethics Committee of the Bahiana School of Medicine and Public Health, under the CAAE 13568213.8.0000.5544, and obeyed the Helsinki Declaration of 1964 and subsequent amendments. All participants who agreed to participate in the study signed the informed consent form.

Consent for publication
All individual participants provided consent for the publication of this study’s results as a part of the informed consent process.

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