ABSTRACT

Objective: To identify HIV coping strategies among HIV-positive older adults. Method: An exploratory-descriptive study with a qualitative approach conducted with HIV-positive older adults in two HIV Specialized Care Services and using the Theory of Social Representations as a theoretical framework. A semi-structured interview technique was used, which was analyzed by the lexical analysis method using the IRAMUTEQ software program. Results: Forty-eight (48) older adults participated in the study. The HIV coping strategies adopted by HIV-positive older adults are to cling to religiosity and spirituality, to adhere to treatment, to have institutional support from health professionals and support from social networks, especially family and friends, and to choose to maintain confidentiality of the diagnosis. Conclusion: HIV coping strategies, among other things, are configured for seropositive older adults as sources of empowerment, hope, possibility of life and acceptance.

DESCRIPTORS

Aged; HIV; Adaptation, Psychological; Social Support; Geriatric Nursing.
INTRODUCTION

There has been a change in the course of the HIV/AIDS epidemic disease with its evolution in recent years, since the epidemiological profile has shown a gradual increase in cases in the age group of 50 years or older in both genders, resulting in an increase in older people living with HIV.

Such a reality may be related to social and individual aspects. In other words, it can be the result of both unfavorable socio-demographic and economic conditions, as well as the lack of information, creating an uncomfortable situation where older adults exercise their sexuality with complete freedom in the sense that they do not protect themselves and feel immune, and on the other hand the society and health professionals who believe they are asexual.

Thus, health professionals sometimes do not investigate HIV among older people, and due to age-related immune system weakness coupled with the lack of suspicion regarding a diagnosis, they may therefore present AIDS symptoms which are confused with a common manifestation of aging or with another chronic disease. Thus, the ideal situation is that HIV investigation in older adults becomes part of the routine of professionals, since the earlier the diagnosis, the better the patient’s prognosis.

In this context, the discovery of HIV infection among older adults requires a new look at life as a way to adapt and positively manage the challenges arising from the diagnosis, thus aiming toward developing coping strategies in order to face living with HIV. These strategies serve as mechanisms to alleviate suffering, conflict, and weakness, thus empowering people, giving hope, and increasing their resistance to stress.

In view of the above, the aim of this study was to identify strategies for coping with HIV among HIV-positive older adults.

METHOD

STUDY DESIGN

An exploratory-descriptive study implementing a qualitative approach.

SCENARIO

The study was developed in two HIV Specialized Care Services (SAE) located in public hospitals in the metropolitan region of Recife, Pernambuco, Brazil. It is noteworthy that Recife has eight SAE, and the choice for these two was because they serve a larger number of people living with the virus, as according to data provided by the institutions themselves; one had 6,806 registered patients, and the other, 2,630 patients.

Both institutions stated that they did not have control over the number of patients treated according to age group in their system; a fact which made it impossible to reach an exact number of the older adult population in the referenced SAE.

The Theory of Social Representations (TSR) was used in its procedural approach as a theoretical framework. Social representations demonstrate the way in which individuals understand everyday events, indicating common sense theories based on their knowledge of their position regarding conflicting events.

POPULATION

Forty-eight (48) older adults who met the following inclusion criteria were selected by convenience to participate in this study: age 50 years or older, diagnosed with HIV for at least 6 months, and outpatient follow-up in one of the SAE included in the study. Any people without the necessary cognitive conditions to participate were excluded. The choice of the number of participants to compose the sample was guided by a consensus which exists among theorists studying TSR, which considers 30 as the minimum number to define a representation. Thus, the data saturation criterion was considered from this minimum number.

Participants were collected at the outpatient clinic during the day shift by oral invitation to participate in the study by order of arrival of patients for medical consultation, followed by the signing of the Informed Consent Form (ICF) and consequent application of data collection instruments. There was no dropout during data collection.

DATA COLLECTION

Data collection was carried out from April to May 2017 using a sociodemographic and clinical questionnaire extracted from a validated instrument, and a semi-structured interview script prepared by the authors, containing questions which explored the representational contents about HIV, aspects inherent to the diagnosis, and coping strategies for living with HIV.

DATA ANALYSIS AND PROCESSING

The interviews were audio recorded with the consent of the participants, with subsequent transcription and preparation of the corpus for analysis. The average duration of each interview was 24 minutes. The participants are identified with the letter “I” derived from the word Interviewee in order to preserve their identity, along with the sequence number of the interview (example: I1).

After the database was organized, it was inserted and processed in the software Interface de R pour les Analyse Multidimensionnelles de Textes et de Questionnaires (IRAMUTEQ), in which a brief description of its contents was obtained. One opted for subsequent analysis using the Descending Hierarchical Classification (DHC) method.

Each text was evaluated and divided into text segments by DHC, which were classified according to their vocabularies, constituting lexical classes. The contents of two classes which addressed the coping strategies of HIV-positive older adults about HIV were analyzed and discussed in this study, since it was in these that the excerpted object for discussion was found.

ETHICAL ASPECTS

The study development complied with national and international standards of ethics in research involving human
The church helps me to face it, it gives me a little more hope (I11).

Religiosity and spirituality

The first category presents religion and faith in God as a strategy for coping with living with HIV. I would say that I ask God very much for the strength to keep fighting for my exams, my consultations, my treatment (I9).

Religion helps me cope a lot. God is who sustains me to face this, and he has been very faithful in my life (I10).

The church helps me to face it, it gives me a little more hope (I11).

Religion helps me to cope (I17).

The Spiritism religion also helps me to cope (I17).

I think God helps me to confront it and especially my family (I30).

There are also reports of hope placed in God’s action towards HIV, in which the confidence of the HIV-positive older adults can be perceived in either a miracle or in scientific success in coping the disease.

God created the laboratory and gave intelligence to man. My faith in God helps me a lot to cope (I19).

God enables humans to discover these medications and through them we succeed (I35).

But nothing is impossible to God. God has already healed me of cancer, so He can cure me of HIV (I42).

Religiosity and spirituality strengthen older people with HIV in facing their sickness, helping them to have their hopes renewed and contributing to a subjective well-being.

Adhesion to treatment

The second category reveals adherence to treatment as another strategy for coping with HIV and is of great value when considering treatment, especially antiretroviral drugs as a way of survival and for improving the quality of life.

This treatment is a great thing and is very important for me to live a little longer (I3).

The treatment is very important, it’s my life. If it wasn’t for it, I don’t know if I would be alive (I7).

For me, the treatment is there for me to get well. I mean, well, I won’t get better, but I keep taking the medicine, which is a way to help me cope until it’s time to die. This treatment means that I am alive (I18).

My real thought is that I have to take this medicine because it is God to me (...) I don’t think I will die without God allowing it, and the antiretroviral doesn’t allow it, and I lead a normal life (I10).

This treatment is great and this antiretroviral is very important because if I run out of medication the tendency is to get worse (I17).

Treatment means I can have a better life expectancy, because by taking the antiretroviral drug properly, in the future I may even die from something that has nothing to do with HIV (I26).

I cope by taking the antiretroviral drug (I29).

Adherence to treatment is a form of self-care, contributing to adapting to the new life after diagnosis.

Institutional support

The third category complements the previous category, since information and attention of health professionals to older adults in the context of treatment adherence were listed as important social support mechanisms for coping with HIV, as a relationship of trust between patient, doctor and other team members is one of the factors which positively influences adherence.

The treatment I receive from service professionals, my doctor and others is very important to me. Thank God for this treatment (I7).

The psychologist was talking to me and that’s when I reacted, I got better and started thinking the way I think now. The moment I started talking to the psychologist everything got better in my head, I confronted it, standing firm and I didn’t listen to what they thought of me and kept talking (I10).

Treatment is a way to fight HIV, I know I have to continue with it until the end of my life. The treatment by the professionals I receive here I also find excellent, from the attendants to the doctors (I13).

Here at SAE there is care and concern that health professionals have with us, and out there, there isn’t. They [health professionals] helped me a lot and gave me a lot of strength to face it. This treatment is my life because a person with HIV gets very weak and then recovers. It’s like a plant root dying in the sun, then the person puts it in their house and waters it and it returns. So the treatment is that. I’m very grateful for the medicine and the people who work here (I20).

The staff is very good. In addition to fighting the disease, the professionals gave me very important mental and spiritual strength (I22).

It can be seen that the dialogue and welcoming fostered by the health team contribute to the older adults with HIV...
Living with HIV: coping strategies of seropositive older adults

feeling welcome and valued, corroborating their physical and mental well-being, the process of acceptance of their health condition and adherence to the antiretroviral therapy.

SOCIAL SUPPORT NETWORK

The fourth category covers social support networks with an emphasis on family and friends as key elements in this coping process. As these are people with whom older people often live with and trust, they soon know about their diagnosis when they feel safe to tell them. Thus, the participants mentioned that family support greatly contributes to living with the virus. I faced it, including I had the support of my family, because all my family knows (I4).

I faced it with the help of my brothers (I8).

My family gave me a lot of support. Thank God this happened, because contempt drives man to despair (I19).

The support of my sisters helps me a lot (I26).

I face it normally. I believe my son helps me, and it strengthens me a lot (I39).

Disclosure of the diagnosis is a situation which causes distress and uncertainty in the minds of people living with HIV, however it is known that older adults will feel more empowered to deal with the virus when receiving support and assistance from friends and family. In this sense, it can be seen that the older adults feel supported after disclosing their diagnosis when there is a supportive relationship without discrimination.

My family supported me. Thank God nothing has changed (I6).

My current wife and sister are the only ones who know. I told them from the beginning, and their relationship with me hasn’t changed. They gave me a lot of support (I13).

My whole family knows, just the family. They did not isolate themselves from me, they have always treated me well and gave support (I17).

Much of my family knows and some friends. Our relationship remained the same, and they support me a lot (I34).

Today my family is more careful with me than they were before I had HIV. They support me a lot (I42).

Thus, it is clear that family members and friends can be considered two major drivers for the well-being and coping of HIV-positive older adults living with HIV. After all, the support of loved ones can help them to accept themselves and thus create mechanisms to move forward.

DIAGNOSIS CONFIDENTIALITY

The fifth category addresses the diagnosis confidentiality as a way to address HIV seropositivity because of fear of the reaction from some family members or close relatives, choosing not to tell anyone about their condition or to disclose to only one or a few reliable people.

God forbid to tell someone I have HIV. They will want to put me aside again, they will want to despise me. I already told my sister to tell the people of the street and all the rest of my family only when I die and God has taken me (I13).

I try not to tell anyone and not be talking nonsense. There is a lot of prejudice. If they knew they would discriminate me, they would even stone me if it was possible (I6).

My son doesn’t know so far. He also doesn’t know that his father died of it, I didn’t tell him and I don’t want him to know, especially a neighbor, God forbid. If one person knows, the whole city knows. I never wanted other people to know because of prejudice, because I know that prejudice would mainly come from neighbors, because such a disease, whether you like it or not, there is always someone who has prejudice (I26).

I was embarrassed and didn’t have the courage to tell the children, so much so that they don’t know until today, only my wife. Besides my wife, only health professionals know (I31).

Only my daughters know in my family, because I know that if the rest of the family finds out that I have it, they will reject me (I36).

So no friend knows, no neighbor, and even worse my family, because in my family there is only snakes. Not even my daughter knows. She’s a psychologist, but she’s very explosive and I don’t trust her (I46).

It is noted that the fear of prejudice and consequent discrimination and rejection is the main factor which leads to the option of maintaining diagnosis confidentiality from the family and society, a representation rooted in the imagination of people in general.

DISCUSSION

As shown in the results, religiosity and spirituality, adherence to treatment, institutional support, family support, and diagnostic confidentiality were defined as coping strategies for HIV.

Regarding religiosity and spirituality, it should be noted that their concepts overlap in the context of common sense and are used as synonyms; however, each has its particularities. Religiosity is an organized system of practices, beliefs or rituals in which people engage and approach the transcendental or sacred, serving as a way to express spirituality. On the other hand, spirituality involves individual beliefs and represents a way for a person to connect with their faith and maintain a commitment to the divine, which may or may not culminate in religious rituals, influencing or being influenced by social, cultural, and psychological factors, among others. This means it is characterized by a personal relationship with the transcendent, of an extraphysical character, referring to something broader, which may also include religion.

A study with older adults showed that spirituality and personal beliefs helped in coping with chronic disease, giving them a feeling of relief and serenity. Other studies with people living with HIV reinforce this data by conceiving prayer, personal faith, and faith in some God as indispensable sources of comfort and strength when facing adversities and health issues.
Still from this perspective, research has shown that people living with HIV who had faith in God or followed a religion were able to more easily adhere to treatment and to follow it correctly compared to those who did not belong to any religious group or had faith in something sacred\(^{(16)}\).

Thus, it is noted that spirituality and/or religiosity is presented as a strategy for coping with HIV. It is even considered that religiosity and/or spirituality can be a strategy of self-care not only on physical issues, but mental, since believing in the transcendental helps to ease the pain and face the adversities of being HIV positive.

In this context, it was possible to infer that the hope for curing HIV among the older adults is also based on faith, but in different ways, being linked to a direct relationship with the transcendental, and on the other hand mediated by God, in which man will make scientific discoveries and find the cure through the wisdom given by the divine.

In this case, it is considered that there is an association between ideas and conceptions which engender the consensual and reified universe, and in the latter, science and faith complement each other to promote the well-being and quality of life of people living with HIV.

Moreover, by identifying the social representations of individuals and the way in which they are constructed, it is possible to understand their reality and the factors which may interfere with their daily life\(^{(27)}\). From this perspective, in this study it was observed that the social representations of older adults regarding HIV are composed of a set of images, concepts and positive feelings related to antiretroviral therapy (ART), which results in removing the perception of the proximity of death, creates expectations for the future and promotes closeness with family members to help coping with living with the virus.

It is noteworthy that treatment success cannot be restricted to medication alone, as adherence also involves other factors such as dietary and behavioral changes, meaning that it corresponds to the adequacy to new habits, regular appointments, periodic examinations and care with prevention, food and hygiene\(^{(18-19)}\). In a study with people living with HIV aged 18 years and over, including older adults, it was found that the group gave greater importance to healthy lifestyle habits including proper diet, exercise, leisure activities, more attention to sleep and rest, among other care actions for health promotion, in addition to the therapeutic program. Medications represent healthcare needed to continue living normally and to improve quality of life\(^{(20)}\).

In the social thinking of the group studied in this research, treatment is considered important for coping with the disease, as it provides health, reduces morbidity and increases life expectancy. Thus, it is valid to recognize that treatment drives the desire to live and overcome the obstacles imposed; however, it is still a challenge, especially for older adults\(^{(21)}\).

Thus, the work of health professionals to promote a multidisciplinary approach is emphasized in this context, and they should foster a dialogue which enables patients to be prepared for starting therapy and the possible side effects which may occur, providing calmness and safety\(^{(22)}\). Providing sufficient health services and a welcoming attitude of professionals are key elements in the motivation to maintain treatment continuity, especially at the beginning when the psychological impact is greater and people living with HIV need to feel cared for by someone\(^{(23)}\).

In a study aimed at assessing the quality of life of older people, the authors found that the best scores were related to the care and respect from the health team with this clientele during care, which favored trust and bond with the older adults, in addition to contributing to adherence to ART\(^{(24)}\).

The health services which attend this type of clientele are care enhancers, with a reflection on the quality of life of this group, providing more humanized care with care actions that go beyond drug treatment, consultations and examinations, including psychological support, which greatly helps people in coping with living with HIV\(^{(20)}\).

It is noteworthy that a committed and interested multi-professional team is reflected in the motivation of self-care among people living with HIV, as evidenced in this study by recognizing the appreciation they receive from health professionals as a foundation in coping, not only due to their skills, but also for the ethics and commitment to their care process. In addition, health education or guidance provided by health professionals is of great importance in contributing to lifestyle change in pursuit of a healthier life.

In addition to institutional support, it is worth highlighting family and friends as the main social support networks in this process. It is common to use social support networks as a way to confront the stressful situation when it comes to any kind of difficulty in treatment. This means that the person living with HIV often defines their family or relatives as an indispensable source of solidarity and encouragement, alleviating daily challenges and assisting in maintaining a positive outlook\(^{(23,25)}\).

From this perspective, a study conducted with older adults revealed that 80% of them considered their family their reason for living and the main source of strength to face a chronic disease, as it is responsible for meeting both their affective needs and the care needs; in other words, they provide love and care in facing the limitations of old age or illness, protecting older people from negative feelings and making them feel important and belonging to a group\(^{(31)}\).

At the same time, other studies with people living with HIV have revealed that the family is the main incentive to deal with HIV, playing a decisive role in changing habits and routines and encouraging adherence and maintaining treatment\(^{(26-27)}\).

However, there are older adults who choose to keep the diagnosis confidential even from their social network. In this regard, it is believed that such an attitude is due to the fear of contempt, as social representations of HIV linked to danger and prejudice still exist in society and in the imaginations of the older adults themselves. Thus, they think that if they reveal seropositivity, their social networks are likely to suffer from the shock of diagnosis or to be afraid of becoming infected as well, expressing prejudiced attitudes and discriminatory behaviors, which may be reflected in the lack of support offered and consequently in the coping of seropositive older adults.

The fear of not being accepted in the social group or feeling abandoned and lonely are the main reasons why a person living with HIV is ashamed and decides to keep their condition confidential. Two studies of seropositive...
older adults revealed that they were sometimes reluctant to break the secrecy of their serological status, and when they revealed it was only to a single confidant, who is almost always a family member.\(^{26,28}\)

In a study of older adults with HIV, the authors considered that the concern with concealing the seropositivity condition would be related to the fear of non-acceptance and a feeling of guilt, showing that such conditions may impact an increase in anxiety, depression and desire to die.\(^{24}\)

As a result of this fear of being marginalized, criticized and discriminated against, older adults with HIV are isolated from contact with friends, co-workers and neighbors as a way of protecting themselves.\(^{29-30}\)

The social representations of HIV linked to family interaction and support are almost always focused on overcoming the diagnosis and the possibility of reorganizing life and facing HIV seropositivity.

This study was limited by the fact that data collection was performed without the interview script having undergone a pilot test for validation.

CONCLUSION

Religiosity and spirituality are sources of psychological strengthening and hope in relation to HIV. In turn, adherence to treatment, especially with regard to ART, presents itself as a concrete possibility of living, constituting something positive for the seropositive older adult.

The support coming from health professionals (institutional) and from family and friends (social network) also contributes to coping with HIV, since older adults tend to be more confident and more easily adhere to treatment by feeling respected and welcomed by these people. However, because they are aware of discriminatory and prejudiced actions by society towards an individual with HIV, seropositive older adults sometimes tend to use secrecy as another form of coping, which is reflected in their desire to be accepted and to protect themselves.

It is considered that the objective of this study was achieved, since it was possible to extract the main daily coping strategies for living with HIV from the participants’ statements. The study is expected to contribute to improving the knowledge of the entire scientific community, especially health professionals who directly deal with and treat HIV-positive older people in order to promote care focused on their individual needs and encourage the development of strategies which help older adults coping with their serological condition positively.

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