Since the start of the AIDS epidemic more than three decades ago, doctors, family and friend caregivers, and patients have observed that some people with the disease experience decline in brain function and movement skills as well as slight or sudden shifts in behavior and mood. This disorder is called HIV-Associated Neurocognitive Disorder or "HAND." Although advances in antiretroviral therapy from the past two decades have decreased the severity of HAND, symptoms still persist in 30-50% of people living with HIV. For many people, these symptoms continue to affect activities of daily living.

The most severe form of HAND called HIV--Associated Dementia (HAD) now occurs in less than 5% of people who have access to antiretroviral therapy. The more common form of HAND is Mild Neurocognitive Disorder (MND). Symptoms of MND include behavioral changes; difficulty in making decisions; and difficulty with learning, attention, concentration, and memory. Some patients develop tremors or loss of coordination. A third form of HAND has been described in research studies as Asymptomatic Neurocognitive Impairment (ANI). Study participants with ANI have impaired performance on neuropsychological tests yet don't recognize having any symptoms. Early studies suggest that these participants do poorly on tests that evaluate their daily function suggesting they have real impairment that goes unrecognized. Others note that participants with ANI are more likely to develop symptomatic impairment (e.g., MND) than are participants who test normally on cognitive tests. The following figure summarizes the breakdown of HAND and its various forms.

Risk factors for developing HAND are less clear for people who are adhering to antiretroviral therapy and who have undetectable amounts of HIV in their body fluid (aka 'viral load'). For those who are not taking these medications, treatment is recommended. It is especially important that patients with detectable viral loads seek care and adhere to medications to achieve an undetectable viral load. Both age and other illnesses appear to increase risk. In untreated patients, a low CD4 count
remains a risk factor. CD4 cells or T-helper cells are a type of white blood cell that fights infection.

There is no “typical” course of the ailment. Several groups have described daily fluctuation in abilities. Most of the time, HAND remains relatively mild in patients with suppressed viral load. HAND may be severe or progress rapidly. Some people experience only cognitive disturbances or mood shifts while others struggle with a combination of mental, motor, and behavior changes. How much these changes disrupt a person’s day-to-day life differs from one individual to the next and from one stage of the disease to another.

Facts

Managing HAND is sometimes difficult because side effects from medications, other HIV infections, nutritional imbalances, depression, and anxiety as well as the effects of comorbid diseases (e.g., vascular disease and liver impairment) can all contribute to cognitive, behavioral, and mood disturbances.

Because neurological changes often develop gradually, it can be a challenge to determine exactly when isolated incidents or symptoms should be given a diagnosis of HAND. In the era of HIV where most people are on treatment, it is becoming clearer that cognitive impairment is often due to more than one cause.

Before the arrival of combination anti-retroviral therapy (a.k.a. cART and sometimes referred to as HAART) in the second half of the 1990s, dementia (the most severe form of HAND) was a frequent finding in late-stage disease. More recently, experts have estimated that less than 5 percent of people on cART develop HAD although many more suffer from mild changes that can affect daily living.

Symptoms

Unlike many of the infections associated with HIV and AIDS, HAND has a wide range of possible symptoms, which can vary greatly from one person to the next. The symptoms of HAND fall into three broad categories:

- **Cognitive**: problems in concentration (difficulty following the thread of a conversation, short attention span, inability to complete routine tasks, or trouble finishing a sentence), memory loss (trouble recalling phone numbers, appointments, medication schedules, agreements, or previous conversations), and a generalized slowdown in mental functions (difficulty understanding and responding to questions, or loss of sense of humor or wit).

- **Motor**: poor coordination, weakness in legs, difficulty maintaining balance, tendency to drop things, decline in clarity of handwriting, loss of bladder or bowel control.

- **Behavioral**: personality changes (increased irritability, apathy toward loved ones or life in general, loss of initiative, withdrawal from social contact); mood swings (depression, excitability, emotional outbursts); impaired judgment (impulsive decision-making, loss of inhibitions); and, on occasion, symptoms of psychosis (hallucinations, paranoia, disorientation, sudden rages).

Cognitive problems are often—even if not always—the first to become
apparent to a person with HAND and their family members, friends, caregivers, and health care providers. Motor and behavioral changes, if they occur, frequently become evident in later stages of the syndrome.

**Diagnosis**

Many factors can contribute to the same types of symptoms as HAND making diagnosis a complex and challenging task. Depression, other psychiatric disturbances, reactions to medication, and nutritional deficiencies can all lead to similar symptoms. Infections, common among people with advanced HIV, can also lead to these symptoms although typically only among those not on cART (e.g., toxoplasmosis, lymphoma, progressive multifocal leukoencephalopathy (PML), and cryptococcal meningitis.

An accurate diagnosis of HAND, therefore, requires a comprehensive examination that generally includes a mental status test, a brain scan, and sometimes lab tests on the cerebrospinal fluid (the fluid that bathes the brain and spinal cord) which are obtained through a procedure known as a spinal tap or lumbar puncture. A mental status exam can help identify whether a person is suffering memory loss, difficulties with concentration and other thinking processes, mood swings, and other symptoms. The best diagnosis requires that a third party (e.g. friend, partner, or other family member) corroborate the behavior/memory changes.

Because no single test definitively answers the question of whether someone has HAND, the final diagnosis is made by weighing all the evidence together. Time and repeated measures are helpful in confirming a diagnosis.

**Treatment**

Although there is no cure for HAND, the single most important treatment is adherence to antiretroviral therapy to maintain a suppressed viral load in the blood. The specific medicines that make up the cART regimen appear to be less important that just being on a regimen. In rare cases, doctors may have to consider how well these medicines get into the central nervous system, and there are studies underway to see if some medications may help symptoms better than others. These new findings, however, should not cloud our understanding that the suppression of plasma virus to undetectable or unquantifiable levels in blood appears to be most critical.

Knowing the diagnosis is also sometimes therapeutic as there is stigma and stress related to these symptoms and how they affect daily activities. These can be addressed with proper diagnosis. Knowing that the illness exists may also prompt compensatory approaches and, ultimately, decrease stress and anxiety.

In addition to treating HAND itself, it is important to find ways to treat the secondary symptoms when possible. Anti-depressants, anti-psychotics, and anti-anxiety drugs can help relieve some of the mental distress people with HAND may experience. However, some of these medications may cause complications when taken along with antiretroviral therapy or other drugs; caution is needed in choosing the best approach. Consultation with an HIV-knowledgeable doctor is recommended.

It is also important that patients engage in their own care to prevent factors that can contribute to cognitive symptoms. This includes aggressive treatment of depression, good health care
maintenance to address common comorbidities (e.g., hypertension, lipid abnormalities, liver impairment), avoiding non-prescription drugs and excessive alcohol, working with care providers to assure that the medications taken are all required, and exercise. There is growing evidence that physical exercise is important. Optimally, this can be done while engaging in other activities since social integration and activities may also be helpful. Keeping engaged with enjoyable activities likely translates into benefits.

**Advice for Caregivers**

Caring for someone with HAND can be stressful. Part of the difficulty is that caregivers must cope with not just the practical implications of the situation—for example, making sure the person remembers to take medications on time—but also with the loved one’s own fears and despair about the deterioration of his or her abilities. Exhaustion, fear, isolation, and grief of the loss of the relationship and partnership that once was are common for caregivers. The fact that HAND can impact behavior (e.g., apathy, depression) and that many people suffer from HAND but don't feel they have symptoms makes caregiving even more challenging.

Encouraging someone to obtain a proper diagnosis is an important step. Because the loss of control can be terrifying, some individuals with HAND may be unwilling to admit that they are suffering from cognitive loss. Caregivers should be prepared for possible resistance to seeking help. It is important to remind the patient as gently as possible that there may be ways of addressing and improving the situation once the diagnosis is clarified.

This is also the time to make sure that all important legal documents (visit [www.caregiver.org/where-find-my-important-papers](http://www.caregiver.org/where-find-my-important-papers)) such as advance directives, financial durable powers of attorney, POLST (physician orders for life sustaining treatment) forms, and estate plans have been prepared with the help of a knowledgeable attorney for both the patient and their care partner.

Just as in the early stages of Alzheimer’s or other cognitive disorders, there are steps a caregiver can take to make the situation as manageable as possible. In addition to getting informed about HAND, available treatments, and working with a competent care team, there are steps one can take in their daily life to maximize function and decrease stress.

One of the primary coping strategies is to keep the home environment familiar. Memory prompts such as calendars, lists, beepers, and pillboxes can be a tremendous help in alleviating frustration for both the caregiver and the person living with the condition. Matching the ability of the person to the prompt tools used is key to successful use of these aids. For some patients with more severe impairment or dementia, consider simplifying communication by gaining eye contact before you speak, repeating the names of visitors, and talking slowly and clearly while retaining an adult tone of voice and level of respect. It is often helpful to limit the number of tasks that need to be done at the same time and to break down directions into one or two-step commands.

For caregivers themselves, the demands of coping with a person suffering from cognitive loss—whether related to AIDS or another disorder—can be challenging. Feelings of isolation, depression, guilt, rage,
despair, and fear are common. It can be crucial for caregivers to seek help themselves whether in the form of individual counseling or support groups with others in similar circumstances.

Caregivers must build time in the week to take breaks and recharge. Hiring outside assistance such as a home health aide to help care for an individual with HAND (particularly individuals with the advanced form, dementia) can be essential to achieving a degree of balance in life. In addition, many social service agencies provide caregiver consultation, respite for caregivers (a break from care), daycare programs, and other practical support services. Some caregivers feel ashamed that they cannot do everything and are reluctant to access such support especially if the one who is ill resists. But getting the support you need in order to maintain your ability to carry on with caregiving responsibilities and to stay healthy while doing so—even over possible objections—can be a genuine expression of love and concern.

Resources

**Southern Caregiver Resource Center**
891 Kuhn Drive Ste. 200
Chula Vista, CA 91914
(858) 268-4432 | (800) 827-1008 (in CA)
Fax: (858) 268-7816
E-mail: scrc@caregivercenter.org
Website: www.caregivercenter.org

The Southern Caregiver Resource Center offers services to family caregivers of adults with chronic or disabling conditions in San Diego and Imperial counties. Services include information and referral, counseling, family consultation and case management, legal and financial consultation, respite care, education and training, and support groups.

**Family Caregiver Alliance**
(415) 434-3388 | (800) 445-8106
Website: www.caregiver.org
E-mail: info@caregiver.org

Family Caregiver Alliance (FCA) seeks to improve the quality of life for caregivers through education, services, research and advocacy. FCA’s National Center on Caregiving offers information on current social, public policy and caregiving issues and provides assistance in the development of public and private programs for caregivers, as well as a toll-free call center for family caregivers and professionals nationwide.

**Medical Treatment and Research**

**UCSD Memory, Aging and Resilience Center**
https://health.ucsd.edu/specialties/psych/senior/general/memory-aging-resilience/Pages/default.aspx

**UCSD Owen Clinic**
http://owenclinic.ucsd.edu/Pages/default.aspx

**Neurology Caregiver Corner**
https://wiki.library.ucsf.edu/display/NeurologyCaregiverCorner/Home

An international leader in the field of memory disorders and dementia. The center offers comprehensive evaluations for the diagnosis and treatment of patients, conducts research on new therapies, and offers support groups for patients, family members and friends.

**Project Inform**
www.projectinform.org

**Information, Advocacy, and Support**

**Gay Men’s Health Crisis**
www.gmhc.org

**The Body**
www.thebody.com
This website publishes an online magazine and other materials on all aspects of HIV/AIDS, including dementia, and also offers a bulletin board, chat rooms and additional opportunities to interact with others.

**Recommended Reading**

**Fact Sheets**

*Caregiving at Home: A Guide to Community Resources*

*Special Concerns for LGBT Caregivers*

*Taking Care of YOU: Self-Care for Family Caregivers*

This fact sheet was prepared by Family Caregiver Alliance and updated and Reviewed by P. Esmaeili, BS, N. Boissier, LCSW and V. Valcour, MD, PhD., UCSF Memory and Aging Center. © 2015 Family Caregiver Alliance.

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