Frontotemporal Lobar Degeneration – Agrammatic Primary Progressive Aphasia: Case Report

Degeneración lobar frontotemporal-afasia progresiva primaria tipo agramatical: presentación de caso

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ABSTRACT
We present a case of a 57-year-old male architect with a master’s degree in economics who ran his own business and gave lectures until 2016, with a clinical presentation of 16 years of evolution which began with anxiety symptoms, excessive concern about trivial events, as well as difficulty in planning and decision making. In the last 3 years, there is evidence of social isolation, difficulty in finding words and decreased verbal fluency; he omits connectors in the grammatical composition in association with ritualistic, stereotyped and perseverative behavior (makes several paintings from the same photograph).

Keywords language; aphasia; anomia; primary progressive nonfluent aphasia; frontotemporal lobar degeneration.

RESUMEN
El artículo presenta el caso de un arquitecto de 57 años de edad con Maestría en Economía, que dirigió su propio negocio y dio conferencias hasta 2016, con un cuadro clínico de 16 años de evolución que comenzó con síntomas de ansiedad, preocupación excesiva por hechos triviales, así como dificultad en planificación y toma de decisiones. En los últimos 3 años hay evidencia de aislamiento social, dificultad para encontrar palabras y disminución de la fluidez verbal; omite conectores en la composición gramatical en asociación con comportamientos ritualistas, estereotipados y perseverantes (hace varias pinturas de la misma fotografía).

How to cite: Ariza-Galindo CJ, Santacruz-Escudero JM, Lozano-Rengifo MJ, Segura-Valencia AI. Frontotemporal lobar degeneration – agrammatic primary progressive aphasia: case report. Univ. Med. 2021;62(1). https://doi.org/10.11144/Javeriana.umed62-1.apha
**Introduction**

Clinically, pathologically and genetically, Frontotemporal Lobar Degeneration (FTLD) is a group of heterogeneous disorders that mainly affect the frontal and temporal lobes, usually in people younger than 65 years at the time of diagnosis (1). It represents the second most common form of dementia in people younger than 65 years, and diagnosis tends to be more difficult in older people (2). It has a prevalence of 10 to 30 cases per 100,000 people between 45 and 65 years of age (3). The most common clinical syndrome, accounting for more than half of cases, is the behavioral variant (4). However, the diagnosis of variants involving speech and language is becoming increasingly important.

One of the less frequent language variants in terms of prevalence is agrammatic or non-fluent PPA, where a clinical presentation characterized by agrammatism, labored speech, paraphasias and difficulty in motor speech planning has been described (5).

**Case A**

The case corresponds to a 57 years-old patient, born and living in Bogota. He studied architecture and did a master’s degree in Economics, had his own company, lectured to entrepreneurs until 2016, and currently works as an independent architect. The patient says that for 16 years he has had anxiety symptoms, which have exacerbated, with excessive worries about trivial events, difficulty in planning and decision making.

In the last three years, affective symptoms have become more frequent, he worries about financial difficulties, has become more expressive and communicates his feelings and thoughts to his relatives more often, but participates less in social gatherings. His wife describes that the patient tells some facts out of context, with circumlocutions, and that he has difficulty in drawing conclusions and finding words; his verbal fluency has diminished, in association with ritualistic, stereotyped and perseverative behavior. In addition, he has difficulty in remembering some proper names, he must write down appointments and commitments more frequently, because it is difficult for him to organize his time. Currently, he persistently expresses subjective complaints regarding his memory (“he forgets where he leaves the keys,” “he forgets commitments and appointments”). A family member reports that they create strategies for planning activities (several weeks in advance they make a plan of the day-to-day activities), and with this they have noted an improvement in relation to the subjective complaints of forgetting appointments, commitments or objects. In problem solving, he remains typecast in one or two solutions, without allowing other possible solutions. In relation to language, they have noticed less verbal fluency, it is difficult for him to find words, especially in periods of anxiety, but also in quiet moments, when he narrates events centered on himself, excluding the other participants, slow processing, without paraphasia, omitting connectors in the grammatical composition (e.g. “home we go soon”) that sometimes it is possible to understand. No episodes of disorientation. Empathy and affection for other people has decreased. Associated with these changes in language, he has become much more interested in drawing. However, he has changed the pattern of his drawings and has been typecast into continuing to create paintings from a single photograph.

**Discussion**

Frontotemporal lobar degeneration is a clinically, genetically, and pathologically heterogeneous neurodegenerative disorder that causes selective neuronal loss and gliosis of the frontal and temporal lobes of the brain. Frontotemporal lobar degeneration is characterized by neuronal loss, gliosis, and microvacuolar changes of frontal lobes, anterior temporal lobes, anterior...
The cingulate cortex, and insular cortex. Subtypes are associated with characteristic patterns of abnormal protein deposition (6). Initial changes occur in the anterior cingulate cortex, fronto-insular cortex, orbitofrontal cortex, and cingulate-frontal transitional zones. These regions have von Economo neurons and fork cells in layer 5 of the cortex, which are thought to play a central part in the integration of cortical and subcortical networks, and degenerate very early in behavioural-variant frontotemporal dementia (7). Either the microtubule-associated protein tau, the TAR DNA-binding protein with molecular weight 43 kDa (TDP-43), or the fused-in-sarcoma (FUS) protein account for nearly all cases of frontotemporal lobar degeneration. The corresponding pathological subtypes of frontotemporal lobar degeneration are frontotemporal lobar degeneration-tau, frontotemporal lobar degeneration-TDP, and frontotemporal lobar degeneration-FUS (6).

As mentioned previously, the diagnosis of PPA is based on objective deterioration of language and speech, while other cognitive domains such as episodic and topographic memory, and constructive praxis are relatively well preserved (8); therefore, in the initial stages the affection of daily life activities is totally attributable to language problems (5). It should be noted that, although many patients do not have memory impairment, those who do may be misdiagnosed with Alzheimer’s disease with left hemisphere involvement, where language symptoms are more prominent (3).

Globally, a highly variable incidence of non-fluent PPA has been described, ranging from 2.2 to 3.5 cases per 100,000 people. 45% of FTLD cases are due to PPA, and less than half of the cases are due to non-fluent PPAg (9). Some studies have described a much higher prevalence of non-fluent PPA in females, while others report no difference between the two genders (4, 9); some have even suggested that the age of onset may be much later than in other forms of FTLD (4). One study showed an average age of onset of 63 years for non-fluent PPA vs. 57.5 years for FTLD patients and 59.3 years for the semantic PPA variant (4), with an average survival of approximately 7 years for non-fluent PPA (5).

Clinically, patients with non-fluent PPA present with low quality labored speech. Some studies have shown that the average speech rate produced by nfvPPA patients is about 45 words per minute compared to 140 words per minute in healthy controls. This group of patients shows slow speech, with effort and grammatical errors and/or apraxia of speech, that is, in the planning and motor sequencing of the lips, tongue and breathing necessary to articulate speech (4). They usually have difficulty finding words, speak in short phrases, omit articles (a, the) and show pronunciation difficulties, similar to what is observed in Broca’s aphasia (9). Patients may have aphemia, but in the early stages they can communicate correctly in writing (9). Anatomically, the correlation with the clinical picture is explained by the involvement of Brodmann’s area 44 and 45 (Broca’s area) in the lower left frontal convolution and the anterior insula (5).

To diagnose these disorders, it is essential to evaluate language along with other domains. Different tests can be used such as the verbal fluency test, the Boston Naming Test, and the Boston Diagnostic Aphasia Battery, among others. Generally, in early stages it is common for the patient to have difficulty finding words, but frank anoma is rare. Comprehension usually remains intact, except for longer complex sentences. The executive function may show subtle dysfunction, but memory and visuospatial skills remain relatively free early in the course of the disease (3).

For a patient to meet the clinical criteria for non-fluent variant primary progressive aphasia, they must have a speech with apraxia or motor speech agrammatism as well as supporting features (Table 1).
Table 1. Features that distinguish dementia of the frontal type from dementia of the Alzheimer’s type in the early stage of the disease

| Feature                        | Frontotemporal Dementia | DAT (dementia of the Alzheimer’s type) |
|-------------------------------|-------------------------|----------------------------------------|
| Memory                        | Initially relatively spared. Verbal memory is preserved. | Impaired encoding and storage of recent information. Poor memory for recent events. |
| Language                      | Poor | Poor |
| Verbal fluency                | Better for categories than letters. | Poor |
| Spelling                      | Poor, better for letters than categories. | Poor |
| Comprehension                 | Poor | Poor, worse comprehension in speech comprehension. |
| Repetition                    | Poor | Poor |
| Attention/executive function  | Severe impairment, loss of planning. | Severe impairment, deficits of executive attention. |
| Visuospatial abilities        | Preserved | Preserved |
| Behavioral                    | Early personality changes; more behavioral than psychiatric features. Frequent catatonic-like behaviors. | Late personality changes; more psychiatric than behavioral features. Suspicion and paranoia. |
| Neuroimaging findings         | Frontal lobe atrophy | Posterior lobe atrophy. Specific atrophy associated by rapid loss of hippocampal cells. |

Source: adapted from Harciarek M, Jodzio K. Neuropsychological differences between frontotemporal dementia and Alzheimer’s disease: a review. Neuropsychol Rev. 2005 Sep;15(3):131–45

Neuroimaging studies for non-fluent variant primary progressive aphasia may be normal at the time of presentation. However, the changes found are consistent with atrophy in the brain MRI or hypometabolism in the PET or SPECT scan in the lower left frontal convolution near the perisylvian area and involves the Broca’s area (10).

Treatment is symptomatic. The FDA has not approved drugs for frontotemporal degeneration. The use of acetylcholinesterase inhibitors has been shown to worsen symptoms in this group of patients (10, 11). In a double-blind, placebo-controlled trial, the use of memantine in patients with frontotemporal degeneration showed no benefit on behavior or cognition (11, 12).

Behavioral symptoms in frontotemporal degeneration variants can be managed with selective serotonin inhibitors. Atypical antipsychotic drugs can be used in frontotemporal degeneration, but caution should be exercised, given the vulnerability to extrapyramidal side effects (12). The use of non-pharmacological therapies that include behavioral, environmental and physical techniques, in addition to caregiver education, may help minimize or redirect unwanted behaviors, while language deficits may be slightly improved with speech therapy in patients with variant primary progressive aphasia (10, 13).

Conclusion

Although the agrammatic or non-fluent variant of PPA is a rare entity, with limitations in its early diagnosis and treatment, it should be considered within the differential diagnosis of young people with affective and language symptoms, since its early diagnosis can improve the quality of life of the patient and their family.

Conflicts of interests

We declare that we are aware of the journal’s conflict of interest policy. The author declared no potential conflicts of interest with respect to the research, authorship, and/or publication of this article.

Ethical approval

Our case report has the ethical approval of the Institutional Committee to report individual cases or series of cases.

Financial disclosure

We declare that this research work is original and is not being evaluated by any other scientific journal. We declare that the corresponding ethical authorities have been taken into account to carry out this research work, all authors agree with the present version of the manuscript and there are no conflicts of interest. We declare that no financial support or sponsors were required to carry out the research and / or the preparation of the article.
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