An Association of Fragile X Associated Tremor/Ataxia Syndrome (FXTAS) with Three Common Neurological Problems: A Review

Shuja Yaqub, Robert Soo Hoo, Devyn Cottor, Hassaan Tohid*

Center for Mind and Brain, University of California, Davis, 95618, United States

*Corresponding author: Hassaan Tohid, Center for Mind and Brain, University of California, Davis, 95618, United States, E-mail: hassaantohid@hotmail.com

Abstract

Aim: To study the association of Fragile X associated tremor/ataxia syndrome (FXTAS) with Alzheimer’s disease and Lewy Body Dementia.

Method: A systematic search was performed in various data bases and journals of Neurology. In total over 600 articles were reviewed and out of those, 39 were selected based on selection criteria.

Results: FXTAS is associated with Alzheimer’s disease and Lewy Body dementia because so many symptomatic and pathophysiological similarities are found between FXTAS and these two neurodegenerative diseases.

Conclusion: The association of FXTAS with neurodegenerative illnesses like Alzheimer’s disease and Lewy Body dementia is present because of many similarities between them. However, some dissimilarity does exist and some questions remain in addressing the pathophysiology of this association. Therefore we recommend more research in the near future to understand this association to develop and utilize wide-reaching therapies.

Keywords: FXTAS; FXTAS Alzheimer’s disease; FXTAS Alzheimer’s Disease; FXTAS Alzheimer; FXTAS dementia; FXTAS memory; FXTAS symptoms; FXTAS Lewy Body dementia; Lewy body FXTAS.

Introduction

When the word “dementia” is discussed. Most people probably think of Alzheimer’s disease (AD). However, there many other rare causes of dementia. One of these rare causes is Fragile X–associated tremor/ataxia syndrome (FXTAS). FXTAS is a rare neurodegenerative disease seen mostly in men aged 50 and above[1]. One in 3000 men is affected by the syndrome. FXTAS arises in male and female carriers of a premutation expansion (55 to 200 CGG repeats) of the fragile X mental retardation 1 (FMR1) gene. Common symptoms include memory problems, behavioral instability and cognitive decline especially in executive function, memory and visuospatial function. These neurodegenerative symptoms are synchronized with cortical-subcortical dementia[2]. Other symptoms seen in FXTAS patients include tremors and ataxia, which ultimately affect tasks like handwriting and lead to balance problems, limb weakness, muscle rigidity, speech problems, and lack of response to stimuli (e.g. pin prick)[3]. Furthermore, bladder and bowel control could also be affected in FXTAS. The degeneration involved in FXTAS is confined mostly to white matter[4].

Since the establishment of neuropsychiatry as a recognized sub-specialty of medicine, interest in diseases like FXTAS has increased, not least due to the association of neuropsychological symptoms like apathy, delusions, hallucinations, sleep problems and cognitive impairments with neurodegenerative diseases. Neuropsychiatric symptoms such as these are not exclusive to FXTAS and occur in other common disorders such as Alzheimer’s, Parkinson’s and Huntington’s. These symptoms also present in 50% to 80% patients with frontotemporal and Lewy body dementias.

In this review, we aim to relate fxtas in particular to AD and LBD. Because of the common symptoms observed in these conditions, it is conceivable that there is an association between the three conditions. Thus, this review will broaden our understanding of the possible association of these conditions with each other.

Received Date: Feb 26, 2016
Accepted Date: July 21, 2016
Published Date: July 25, 2016


DOI: 10.15436/2377-1348.16.795
Patients with FXTAS dementia have been compared to Alzheimer’s disease and FXTAS.

**Discussion**

**Alzheimer’s disease and FXTAS**

Patients with FXTAS dementia have been compared with patients with Alzheimer’s disease (AD)\(^3\). Some similarities and dissimilarities have been found between the two medical conditions. For example, language performance is higher in FXTAS than in AD patients due to the involvement of both parietal lobes in AD, which ultimately causes impairment in language. Language and speech are not affected for the majority of FXTAS patients, however a cerebellar type of dysarthria is found in some patients with advanced disease\(^8\). Also evidence suggests that FXTAS patients score poorly compared to normal controls on Block Design\(^7,8\).

Seritan et al 2008 confirmed that the measures of executive functions like verbal fluency, are expected to show reduced values in FXTAS, with more frontal involvement. They also found that FXTAS patients performed far better on a measure of simple attention. They also found that there was no significant difference between the working memory scores of FXTAS dementia patients and AD patients. This demonstrated that the working memory impairment in FXTAS dementia patients is not significantly different than the typical shortfalls seen in AD patients\(^9\). While according to Grigsby et al\(^9\) attentional control impairment is found in some FXTAS patients when compared with AD patients. They also studied that FXTAS patient with cognitive impairment demonstrated moderate to severe shortfalls in working memory\(^6\).

Some gender differences have been found in FXTAS patients, according to two studies on female carriers of the FMR1 premutation. Decreased cerebellar volume and higher severity of FXTAS symptoms and increased length of the CGG repeat expansion were found in male premutation carriers only and not in females. However in women with FXTAS, the cognitive deficit is moderate in intensity and milder brain changes on MRI are observed as compared to their male counterparts\(^10\). This could be due to the protective effect of the second X chromosome or estrogen in females\(^10\).

Dementia with FXTAS has a cortical-subcortical pattern, because of the involvement of cortical (hippocampal, frontal) and subcortical (middle cerebellar peduncles, white matter) regions (95).

**Neurobiology of Alzheimer’s disease and FXTAS**

Alzheimer’s disease is another very prevalent neurodegenerative disease associated with cognitive dysfunction. Caused by protein plaques and neurofibrillary tangles that destroy the synaptic pathways and affect the neuronal connections. The main proteins involved in this disease are amyloid peptide protein (APP) and the tau protein. This process is caused by secretase enzymes (β and γ) which are located on cell membranes and form β-amyloids that are released into the plasma and cerebrospinal fluid where they aggregate and form plaques\(^12\). The formation of these plaques results in a breakdown of the synaptic microtubules and hyper phosphorylation of tau proteins. In FXTAS, cognitive degeneration is primarily caused by the mutation of the fragile X mental retardation gene (FMR1) which results in mediated RNA toxicity and defected mRNA. This can affect the brain’s motor and cognitive functions. FMR1 in relation to Alzheimer’s disease was studied by Abigail J. Renoux et al 2014\(^13\) where FMRP (fragile X mental retardation protein FMRP, a protein constructed under the instruction of FMR1 gene) was found to have little expression in brains of AD patients. This suggests that there is little involvement of FMRP in AD neuropathology. However, similarities between AD and FXTAS have been discovered in that both neurodegenerative diseases involve APP and tau protein aggregates. Neuropathological evidence of both FXTAS cosinophilic inclusions and neurofibrillar tangles and neuritic plaques, consistent with AD are found\(^14\).

Due to the significant involvement of proteins in both neurodegenerative disorders Sokol et al 2011\(^11\) carried out a study that examined the presence and link of APP FMRP between the diseases. This was conducted by using FMR1 knockout mice which exhibited high quantities of APP translation.
This suggests that the FMR1 mutations can result in increased APP levels and may explain future cognitive deficits. Although the study focused on the Fragile X syndrome specifically, the analysis of FMR1 mice could be linked with FXTAS as well, as FXTAS is also associated with FMR1 and FMRP. The study suggested that amyloid plaques did not form due to the lack of FMR1 gene in the knock-out mice. However, in FXTAS as the disease is caused by elevation of FMR1-mRNA as stated by Kathryn Lovell et al 2008[16], thus still enabling amyloidogenic processes to occur which could be a partial cause of cognitive impairment that befalls FXTAS patients.

Samples of the entorhinal cortex layer II of the medial temporal lobe from human brain showed decreased levels of 5-hydroxymethylcytidine (5-hmC) and DNA (Cytosine-5-)-Methyltransferase 1 (DNMT1) in neurons of AD patients[17], 5-methylcytidine (5-mC) and 5-hydroxymethylcytidine (5-hmc) were also studied by Choulia et al. 2013. They found a similar reduction in 5mC and 5-hmC in the hippocampus of AD patients[18]. On the other hand a study also revealed opposite results, with augmented levels of 5mC and 5hmC in AD brains. These high levels of 5mC were seen in frontal cortex of AD patients[19].

Another study revealed augmented levels of Ten-eleven translocation methylcytosine dioxygenase 1 (TET1), 5mC, and 5-carboxylcytosine (5caC), in the hippocampus of AD patients reported by Coppieters et al. (2014) found a similar reduction in 5mC and 5-hmC in the hippocampal region of AD brains. Furthermore, levels of 5-mC as well as the levels of 5-hmC showed a significant negative correlation with amyloid plaque load in the hippocampus[19]. On the other hand a study also revealed opposite results, with augmented levels of 5mC and 5hmC in AD brains. These high levels of 5mC were seen in frontal cortex of AD patients[19].

Diagnosis

A study conducted by Sachdev et al. 2013[32] suggested that Alzheimer’s is not exclusive to the gray matter of the brain but in fact affects the white matter of the brain as well. As a result, the experiment carried out by Filley CM et al. 2015[33] examined white matter disease and cognitive impairment within carriers of the FMR1 premutation. White matter mainly consists of myelin, which is a layer of fat used to augment connectivity between nerve fibers. Thus, the degeneration of white matter in Alzheimer’s can suggest that processing speed is slower. The investigation included 13 premutation carriers compared with 7 healthy controls that were examined using magnetic resonance spectroscopy (MRS) and diffusion tensor imaging (DTI). Areas of interest included the middle cerebellar peduncles and corpus callosum, both areas of white matter and cognitive function. The study used performances of the Controlled Oral Word Association Test (COWAT), Symbol Digit Modalities Test and Behavioral Dyscontrol Scale and correlated the results with fractional anisotropy (FA) acquired from DTI. FA examines the connectivity (processing speed) of the brain. The results revealed that FXTAS patients had the lowest mean FA, suggesting that FXTAS results in cognitive deficits and reduced processing speed within the brain similar to AD, and also correspond to how white matter disease results in reduced connectivity of the brain. This can perhaps enable easier diagnosis of both FXTAS and Alzheimer’s as DTI and FA would be able to reveal signs of white matter disease.

Symptoms

The summarized report of FXTAS produced by Hagerman RJ et al. 2015[1] also included in depth analysis of the cognitive deficits and impairments associated with the disease. Various features of the FXTAS phenotype include mild cognitive impairment, behavioral instability, and cortical-subcortical dementia, both of which relate to the features of Alzheimer’s disease. These common features sometimes affected the true diagnosis of the disorder. Features of cognitive impairment include loss of short term memory and reduced competence in memory and executive functions as well as various behavioral deficits such as attention and aggression.

Dementia within FXTAS patients can occur both atypically and gradually. It also occurs in succession with other disorders such as tremors and ataxia. An investigation by Philip K. Mothersead et al 2005[34] identified a 61 year old man with gradual neurodegeneration beginning with ataxia and tremors and progressing to cognitive deficits such as loss of short term memory as well as an inability to carry on a conversation. The patient also suffered severe injuries related to his gait ataxia such as a broken arm or leg. The man was examined using a number of tests such as the Wechsler Adult Intelligence test – III and Wechsler Memory Scale – III in which scores were significantly below normal levels. After detailed cognitive examination the patient was also listed as a research subject at the University of California-Davis within the MIND institute. The patient had significant ataxic tremor and cognitive impairments such as confusion of his left and right as well as behavioral deficits such as anxiety and obsessive compulsive behavior. This phenotype was significantly similar to the FXTAS phenotype in that the patient exhibited both tremors and ataxia as well as significant cognitive degeneration. The patient’s brain MRI revealed signal intensities in the cerebellar peduncles and cerebral atrophy previously noted as an important trait in a study conducted by Chaussenot et al. 2008[31]. The patient’s CGG repeat was also within the premutation range at 93 and upon discovering that he was a grandparent of children with FXS the man was diagnosed with FXTAS. Overall, the subject presented a neuropsychological and physical decline from an intelligent individual in a rare case where the patient had both Alzheimer’s disease and FXTAS thus resulting in rapid decline by dementia.
when Adams JS et al 2007[10] carried out their experiment concerning females with FXTAS and the volumetric brain changes associated with them they discovered that the females demonstrated a less brain parts involvement in comparison with males who were also affected. The patients also followed consistent patterns of brain atrophy and white matter disease which can be linked to features of Alzheimer’s as previously represented by Filley CM et al 2015[33].

Early diagnoses of Alzheimer’s and FXTAS aren’t exclusively determined by volumetric expansions and examinations of the white matter. Another way to distinguish cognitive deficits and features is through messenger-ribonucleic acid (mRNA), which are single stranded RNA molecules 22 nucleotides in length. They work by suppressing protein genesis and are involved in a wide range of processes such as cell cycle control, stem cell differentiation and neurogenesis. Despite the numerous functions mRNA is also expected to be a future potential biomarker for Alzheimer’s disease[34]. However, future studies possibly in the next quarter of the 21st century will provide more details whether it could be used as a potential biomarker or not. The study detailed the many functions of mRNA and specifically examined the impacts on Alzheimer’s as well as other neurological disorders. Areas affected by AD pathology are cortex, hippocampus, and the cerebellum. Moreover, the investigation also regarded the fragile X mental retardation diseases as effective demonstration of mRNA dysfunction in relation to the disorder. The mRNA is required for regulating synaptic plasticity as well as maintaining dendrite development. However, an alteration or absence of mRNA results in the deactivation of the Dicer enzyme (essential for RNA synthesis) in the Drosophila fruit fly, hence leads to tau-induced degeneration which also causes in Alzheimer’s disease via neurofibrillary tangles leading to subsequent cognitive impairments.

Lewy body dementia and FXTAS

Lewy body dementias are specifically caused by lewy bodies which are irregular accumulations of protein that form within the nerves. The buildup of these proteins results in neurodegeneration resulting in Parkinsonism and cognitive impairment[36].

FXTAS could be associated with Lewy body according to some studies[37,38]. In spite of the possible similarities of the Lewy Body Dementia phenotype, some evidence from the study by Loesch et al 2015[39] suggests that unlike similarities found within pathogeneses of some diseases for example, tau proteins in Alzheimer’s; the pathogenesis for some FXTAS cases is different from cytoplasmic inclusions which are mainly associated with Lewy Body Dementia. The study also focuses on particular cases of FXTAS where a patient has contracted atypical forms of Parkinsonism as well as dementia but not the extreme forms such as mild cognitive impairment and ataxia. This irregular occurrence can be described as the “FXTAS spectrum” where some features of the syndrome overlap with other disorders such as Parkinson’s with Lewy Body Dementia.

In the report summarizing the emerging topics in FXTAS Deborah A. Hall et al 2014[7] many cognitive disorders were identified specifically within older people. These disorders were able to co-exist within the patient, as post-mortem examinations of these subjects revealed that they not only had multiple sclerosis & AD but also Lewy Body Dementia. This can suggest how disorders can synchronize within patients that have FXTAS. It is possible, that these issues were only discovered as a result of the autopsy of the patients. The features of parkinsonism and Alzheimer’s disease as described by Hagerman et al 2015, Filley CM et al 2015, Andreea L. Seritin et al 2010[1,2,5] are potentially difficult to distinguish as they bare close similarity to other diseases.

Conclusion

Fragile X–associated tremor/ataxia syndrome (FXTAS) is a rare neurodegenerative disease associated with memory problems, behavioral instability and cognitive decline, especially in executive and visuospatial abilities. Research exists which shows that patients with FXTAS may exhibit symptoms commonly seen in Alzheimer’s disease and Lewy Body dementia. This mini review highlights the fact that these three neurological disorders (AD, LBD and FXTAS) can overlap in symptoms and somewhat in neurobiology: especially in that all of these disorders present with dementia. We believe articles like these will help future clinicians and scientists explore the unidentified aspects of any possible association. Further research is needed as there is more about dementia in these disorders that is unknown, and as new research is conducted it will broaden our understanding of dementia not only in FXTAS, LBD & AD, but about all possible causes of dementia.

Acknowledgements

The authors are grateful for the help, support and suggestions by Dr. Jin Chen Yang (UC Davis).
References