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ORIGINAL ARTICLE Psychiatric manifestations in cerebrotendinous xanthomatosis

MJ Fraidakis^{1,2}

Cerebrotendinous xanthomatosis (CTX) is a rare and severe, but treatable, inborn disorder of bile acid biosynthesis and sterol storage with autosomal recessive inheritance and variable clinical presentation. CTX treatment consists of chenodeoxycholic acid and must be started as early as possible to prevent permanent disability. Psychiatric manifestations are rare and non-specific, and often lead to significant diagnostic and treatment delay. Therefore, better recognition of the gamut of psychiatric manifestations in CTX can diminish the risk of misdiagnosis and irreversible neurological deterioration. We hereby describe the psychiatric features in CTX. A complete review of all published cases of CTX in the medical literature was undertaken and the case reports with psychiatric presentation were collected and analyzed. We also describe the psychiatric features in relation to the neurological semeiology in six patients with CTX diagnosed at the La Salpêtrière Hospital. We conclude that psychiatric manifestations in CTX follow a bimodal/ bitemporal pattern, appearing early in the disease course in the form of a behavioral/personality disorder associated with learning difficulties or mental retardation, or manifesting in advanced disease in the setting of dementia as rich neuropsychiatric syndromes, such as frontal, orbitofrontal or frontotemporal syndromes of cortico-subcortical dementia encompassing behavioral/personality disturbance, affective/mood disorders or psychotic disorders. Behavioral/personality disturbance in childhood or adolescence, especially when accompanied by learning difficulties, should therefore lead to further investigation to exclude CTX, as early diagnosis and treatment is critical for prognosis.

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Keywords: behavior/personality disorder; cerebrotendinous xanthomatosis; chenodeoxycholic acid; cholestanol; psychiatric manifestations; young-onset dementia

INTRODUCTION

Cerebrotendinous xanthomatosis (CTX) (OMIM 213700) is a rare and severe inborn disorder of bile acid biosynthesis and sterol storage with autosomal recessive inheritance and variable clinical presentation.^{1–4} The enzymatic defect of sterol 27-hydroxylase impairs the synthesis of the bile acids, cholic and chenodeoxycholic, with a resultant overproduction and accumulation of several bile acid synthetic pathway byproducts in large deposits in many organs, including the central nervous system (CNS).^{4–6} Diagnosis is strongly suggested by specific clinical features, such as juvenile cataracts, tendon xanthomas or chronic diarrhea, and is confirmed by the lab finding of elevated cholestanol in the plasma or cerebrospinal fluid, and by molecular genetic analysis.^{3,4,6} Early diagnosis is of paramount importance, as treatment can halt disease progression.^{7,8} The mainstay treatment consists of chenodeoxycholic acid (CDCA) per os administration, 250 mg three times daily.^{3,4,7,8} Cholestanol levels can be further reduced by the addition of a statin, usually simvastatin or pravastatin, but with uncertain evidence-based clinical benefit, apart from the prophylactic effect against atherosclerosis.9 Timely therapy is effective in alleviating some of the neurological symptoms, but unfortunately, as a rule, diagnosis is made with a delay of up to many years if ever.¹⁰

More than 400 cases have been described previously worldwide in the medical literature.¹⁰ Given the autosomal recessive inheritance mode, higher prevalence has been reported in some closed communities, such as the Sephardic Jews of Morrocan origin.¹¹ The pathogenic mechanisms in CTX begin to operate early

in life and disease manifestations are usually evident by the second decade. Onset occurs insidiously, with systemic or neurological symptoms and signs.^{3,4,6} Early typical signs are chronic diarrhea, juvenile bilateral cataracts, Achilles (or other) tendon xanthomas, psychomotor retardation, cognitive impairment with learning difficulties or mental retardation, cerebellar ataxia and epilepsy, which can affect up to 50% of patients.^{3,4,6,12-16} Other late-stage neurological features are pyramidal tract signs with extensor plantar responses, progressive spastic paraplegia, progressive cerebellar ataxia and dysarthria, nystagmus, peripheral polyneuropathy with distal muscle wasting and, more rarely, movement disorders, such as parkinsonism and palatal myoclonus, and in advanced-disease pseudobulbar and bulbar syndromes.^{4,6,13,14} Intellectual deterioration may progress to veritable cortical/subcortical dementia, vet mental function may remain apparently normal in some patients.¹³ Systemic manifestations are premature atherosclerosis with cardiovascular morbidity, pulmonary dysfunction and osteoporosis predisposing to bone fractures.^{17–19} Psychiatric manifestations have been reported only sporadically.^{14,20–43}

Magnetic resonance imaging (MRI) is helpful in demonstrating early lesions in the cerebellum, in the form of hyperintensities in the dentate nuclei and hemispheres, and in folia atrophy, on T2-weighted images.^{44,45} Other MRI lesions can be seen in the brain stem, often symmetrically in the pyramidal tracts, the medial lemnisci and the inferior olives, whereas supratentorially, slight, non-specific signal alterations are seen periventricularly, always on T2 images.⁴⁴⁻⁴⁶ In advanced untreated cases, symmetrical dentate

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hypointensities have been reported with T1-weighted imaging, suggesting calcium and hemosiderin deposition, also detectable by transcranial ultrasonography.^{44–46} The cerebral cortex and centrum semiovale appear normal on MRI images, similar to that usually seen histologically.^{13,28,44–46} Macroscopically, large granulomatous lipid deposits (1–2 cm) with extensive demyelination can be found in the cerebellar hemispheres, the cerebellum being most conspicuously affected by lipid deposition, which can otherwise also involve the brain stem and spinal cord.^{13,28} Microscopically, white matter is replaced by neutral fat, needle-like clefts and cystic spaces; foamy vacuolated macrophages and multinucleated giant cells can be found in the affected CNS areas.²⁸

Psychiatric and behavioral manifestations in CTX are rare, especially in the younger patients, and are often misattributed to psychiatric disease when grave or are overlooked when subtle.¹⁰ Personality changes with irritability and aggressivity, depression, delusional syndrome, even catatonia, frank psychosis and suicidal ideation have all been reported in scant reports previously. Dementia often develops in advanced CTX, and in older patients it may be heralded or may coexist with florid psychiatric manifestations.^{20,24,34}

We hereby describe the psychiatric manifestations from a cohort of 13 patients with CTX evaluated at the Pitié-Salpêtrière University Hospital. Psychiatric manifestations in the setting of early or advanced dementia in our cohort are described to serve as a counterpoint to the psychiatric picture of younger CTX patients without dementia. A complete review of all published cases of CTX in the medical literature from 1936 to date was also undertaken and the case reports with psychiatric presentation were collected and analyzed. The results from our series are compared with the findings from the review of the medical literature. Conclusions are drawn on the spectrum of psychiatric manifestations in CTX in relation to patient age and disease stage.

MATERIALS AND METHODS

We reviewed 13 cases of genetically confirmed CTX, diagnosed and followed up at the Neurology Department of the La Salpêtrière University Hospital for the presence of psychiatric manifestations at any stage of the disease history. Patients and their parents were interviewed to obtain the best estimates for lifetime diagnoses. Furthermore, all psychiatric and medical charts, as well as school notes, were collected to confirm the clinical information, both in terms of symptoms and of time course.

A Medline search was done (1968-2012) on CTX. From the reference list of the papers thus collected, we found all articles on the subject, which were published before 1968, as far back as the original publications of Schneider (1936) and van Bogaert *et al.*^{47,48} monography. These papers were collected from medical libraries and were reviewed thoroughly on the basis of the documentation of CTX case reports. Original and review articles in English, French, Italian, Spanish, German and Dutch were read. Articles written in Japanese were not reviewed, but their abstracts were taken into account. Although the occasional review articles in Hebrew and Russian were not read, they did not contain any case reports. From the review of the above mentioned medical literature, we managed to collect data on 425 cases, which is most probably an underestimate of the worldwide disease prevalence. Although many case descriptions were poor in information, we extracted the case reports that contained even the slightest hint of a psychiatric semeiology. In our analysis of the psychiatric manifestations of CTX, however, we did not include the cases that were simply described as having developed learning difficulties, mental retardation or dementia. Nevertheless, we included all case reports wherein psychiatric manifestations were mentioned in the setting of CTX-related dementia.

Case reports (La Salpêtrière Hospital)

Early psychiatric manifestations (not associated with dementia)

Patient 1 (P1). The 24-year-old male patient was born to nonconsanguineous parents. Psychomotor development was normal. The ability to read was acquired normally. At the age of 6 years, frequent falls, clumsiness and learning difficulties in writing, spelling and arithmetic were

noticed. He began attending the local Child Development Center at the end of first grade and attended regular sessions with a reading specialist. Interpersonal relational problems and behavioural disturbance were noticed at elementary school. He was reported to be hyperactive, impulsive and sometimes violent. At the age of 13 years, he was hospitalized after an episode of aggressive behavior towards his mother during a family quarrel when he threatened her with a knife. Psychiatric diagnoses at adolescence were borderline intelligence, attention-deficit hyperactivity disorder, predominant hyperactive-impulsive subtype and oppositional-defiant disorder, and no pharmacologic treatment was administered. His high-school education was disorganized and underachieving because of his psychiatric problems and further intellectual decline. His behavioral disturbance with irritability, aggressivity and oppositional behavior persisted throughout adolescence. He showed indifference and aggressivity towards his mother, non-adherence to rehabilitation orders and negligence of basic hygiene. At 21 years of age, neuropsychological testing revealed diminished global cognitive efficiency, difficulties in abstract reasoning, deficit in attention and dysexecutive syndrome affecting the instrumental capacities (Mini Mental Scale (MMS): 21/29; Frontal Assessment Battery (FAB)^{49,50}: 16/18). At physical and neurological examination, elongated facies, pes cavus and cerebellar ataxia were found, and electroneuromyography evidenced mild peripheral neuropathy. MRI scanning showed discrete cerebellar and parieto-occipital white matter hyperintensities. Cholestanol levels were abnormally high and genetic analysis found mutation of the CYP27 alleles in a compound heterozygous state (E6:R395C/E8:R479C). In retrospect, his mother remarked that he had suffered from chronic diarrheas since early childhood. Standard treatment resulted in marked improvement of his psychiatric condition.⁸

Patient 2 (P2). She is the 19-year-old sister of P1. Psychomotor development was normal. Her medical history was non-specific, except for pes cavus with hammer toes discovered at the age of 7 years, as well as chronic diarrhea since childhood. During school years, the patient was active in sports but was underperformed academically, and exhibited oppositional behaviors and deficits in attention. She received methylphenidate (10-20 mg per day) for attention-deficit hyperactivity disorder from ages 9 to 11 years. The treatment was discontinued because of worsening of impulsivity, aggressiveness and cognitive decline, forcing her to leave school. Neurological examination and electroneuromyography found mild signs of sensorimotor axonal peripheral neuropathy. Results of brain MRI scanning and ophthalmological examinations were normal. Cholestanol levels were abnormally high and genetic analysis found mutation of the CYP27 alleles in a compound heterozygous state (E6:R395C/E8:R479C). The diagnosis of CTX was made after her brother's diagnosis, when she was 17 years old. Standard treatment resulted in substantial improvement of her psychiatric condition.8

Patient 3 (P3). The patient is a sister of P4. She was hospitalized in her twenties for a psychiatric illness that was characterized as psychotic episode. Bilateral Achilles tendon xanthomas were noticed at clinical examination. Retrospective psychiatric evaluation revealed only learning difficulties in childhood and adolescence with poor scholastic performance. The patient was lost to follow-up and did not receive specific treatment for CTX. Cholestanol levels were abnormally high and, similar to her brother, she was a carrier of the the pathogenic IVS4-1G > A mutation in the *CYP27A1* gene in a homozygous state.

Late psychiatric manifestations (associated with dementia)

Patient 4 (P4). The patient is a 50-year-old male, born to nonconsanguineous parents. Psychomotor development was normal. He has several siblings, among whom one sister was with a psychiatric illness and Achilles tendon xanthomas diagnosed with CTX. His overt disease history began at 13 years of age with a grand mal seizure. At 44 years, irritability with general hyperhidrosis was interpreted as anxiety/depression syndrome. In the following months, his condition progressed with apparition of gait instability, dysarthria, swallowing problems and apathy. Physical examination at that time confirmed gait ataxia, dysarthria, hyperreflexia of deep tendon reflexes and psychomotor slowing. MRI imaging revealed midbrain and dentate nuclei hyperintensities at T2 and FLAIR. Cholestanol levels were elevated and the diagnosis was confirmed by the identification of the pathogenic IVS4-1G>A mutation in the CYP27A1 gene in a homozygous state. Treatment with CDCA was initiated at age 44 years. Despite treatment, neurologic signs did not improve and he was markedly apathic when last seen.

	Gender	Origin	D (years)	X (years)	N (years)	Ψ	Ψ^*	$\Psi_{\sf S}$	Ψ_{D} (axis)	Cog
P1	М	Fr–Sp	22	25	ado	6	Yes*	Aggr., irritability, ADHD, ODD	Behavioral/personality	LD/MR
22	F	Fr–Sp	17	20	16	16	Yes*	Aggr., irritability, ADHD, ODD	Behavioral/personality	LD/MR
2	F	French	20	35	ch.	ado	No	Aggressivity, psychosis	Psychotic	LD
P4	М	French	48	50	13	44	No	Irritability, anxiety/depr., apathy	Mood/affective	Dementia
² 5	F	French	45	49	7 m	43	No	Anxiety/depression	Mood/affective	Dementia
DG	F	French	58	58	ch.	50	No	Irritability, emotional lability	Mood/affective	Dementia

Patient 5 (P5). The patient is a 49-year-old female. Psychomotor development was normal. Epilepsy with petit mal seizures appeared at the age of 7 months. She was schooled until the sixth class and later followed special education because of learning difficulties. She recalls that her gait problems began at the age of 30 years and progressed with time. At 42 years, a pyramidal syndrome with spasticity of all members was found at clinical examination and her gait deteriorated more rapidly thereafter. At 43 years, she was unemployed, on invalidity pension, living with her parents and suffering from depression attributed to her sorrow over her husband's death. Clinical examination revealed spastic paraparesis with gait ataxia, hyperreflexia of deep tendon reflexes with bilateral Babinski and Hoffmann signs, and mental retardation (IQ: 50; MATTIS: 118/ 144). At the age of 44 years, neuropsychologic testing scores were 21/30 and 10/18 for MMS and FAB, respectively. Ultimately, metabolic screening revealed elevated cholestanol and the diagnosis of CTX was confirmed with genetic analysis. The patient did not have visible tendon xanthomas. MRI imaging demonstrated bilateral cerebellar white matter hyperintensities, periventricular white matter hyperintensities, and diffuse cortical and subcortical atrophy subtentorially and supratentorially. A combinated regimen of pravastatin (20 mg per day) and CDCA (250 mg three times daily) was initiated at 45 years, and the patient was followed up. The patient experienced a substantial and quantifiable benefit from treatment. On a follow-up, 6 months later, MMS was 25/30 and FAB was 12/18, the remaining neurologic status being stable. On a second follow-up 1 year after the last, FAB was 13/18. On a third follow-up 9 months later, the patient reported sustained improved gait and cognitive benefits (better concentration capacity and less apragmatism). MMS was 28/30 and FAB was 15/18.

Patient 6 (P6). A 56-year-old female patient of French descent born to non-consanguineous parents. She was operated several times during childhood because of bilateral cataract. Gait difficulties and dysequilibrium appeared in her thirties and worsened steadily thereafter during the course of 20 years, resulting in severe spastic paraparesis and ataxia that rendered the patient wheel-chair bound. Along the same timeline, signs of dementia appeared and progressed (anterograde and retrograde amnesia, frontal syndrome signs, emotional lability and irritability). At the age of 56 years, the patient was hospitalized. Neuropsychiatric testing confirmed global dementia with MMS at 21/30 and dysexecutive syndrome with FAB at 12/18. MRI imaging revealed diffuse cerebral cortical and subcortical atrophy, as well as cerebellar cortical atrophy, diffuse moderate leukoencephalopathy, hyperintensities of the corticospinal tract at the capsular and cerebral peduncle levels, and T1-T2-FLAIR hypointensities of the dentate nuclei, corresponding to calcifications. Biochemical analysis revealed elevated cholestanol and sterol precursors of cholesterol, and the diagnosis of CTX was confirmed with genetic analysis. Treatment with CDCA was initiated at 56 years and resulted in alleviation of psychiatric symptoms, but neurological signs did not improve.

RESULTS

In the cohort of 13 patients at the La Salpêtrière Hospital, 6 patients exhibited psychiatric symptoms at some point in the disease course, and in half of them psychiatric symptoms occurred in the setting of organic dementia (Table 1). Out of the three younger patients without dementia at the time of psychiatric symptoms (P1, P2 and P3), only two siblings had a purely psychiatric presentation of CTX, occurring in their early-to-middle adolescence and associated with mild cognitive and intellectual decline without mental retardation (P1 and P2). Notably, they both presented with behavioral and personality disturbance, initially diagnosed as attention-deficit hyperactivity disorder and oppositional-defiant disorder. Agressivity, irritability and insubordination was a major component of their psychiatric tableau as has been seen in other CTX cases we reviewed.^{21,22} Both siblings received standard treatment (per os administration of Chenofalk, 250 mg three times daily), and follow-up psychiatric examination confirmed a clear and substantial improvement of their psychiatric condition.⁸ Standard treatment of the three older patients with CTX-related organic dementia also led to the alleviation of psychiatric symptomatology and/or mild improvement of cognitive status (P4, P5 and P6). In two families testing of the members at risk proved the diagnosis in a second sibling, and in one family this led to timely treatment with beneficial results for the psychiatric condition of the patients (P1 and P2).

Psychiatric manifestations in CTX have only been described in sporadic case reports and in only one case series.²⁰ In total, we found that 54 patients of the 425 CTX cases published in the medical literature in the period 1937-2012 have had psychiatric manifestations ranging from personality changes and behavioral disorder to frank affective and psychotic disorders²⁰⁻⁴³ (Tables 2 and 3). This amounts to an incidence of 12.7% for psychiatric manifestations in the CTX population. This figure is informative but only approximative. The true figure could be higher because of underreporting of psychiatric signs in the poorly described case reports.¹⁰ It could even be lower because of underdiagnosis of CTX in general.¹⁰

There was significant variation of psychiatric phenotype in the above mentioned 54 cases (Tables 2 and 3). The most common manifestation was behavioral/personality disorder (n = 24, 5.6%) followed by mood/affective disorder (n = 10, 2.3%)and psychosis (n = 5, 1.2%), whereas there was only one case report of catatonia in the setting of CTX (n = 1, 0.2%) (Figure 1). Patients with mood/affective disorders reportedly suffered from depression, except for one patient who suffered from dysthymia (Tables 2 and 3).

It was possible to determine the approximate age of onset and mode of disease onset for 34 out of the total of 54 patients found in the bibliography, with CTX and psychiatric manifestations at some point in the disease course (Tables 2 and 3). As expected, in most cases (26 out of these 34) CTX-related symptoms or signs led to hospital referral before the age of 26 years, as usually CTX is an inborn metabolic disease of childhood onset. Pure psychiatric presentation of CTX occured most often during childhood and never occurred after the age of 26 years (Tables 2 and 3, Figures 2

4

Psychiatric	manifestation	۱S	in	CT)
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Reference	Р	G	Origin	D	Х	Ν	Ψ	Ψ^*	Ψ_{D} (Axis)	Cog	Р
Philippart and van Bogaert ²³	P1	F	Flemish	42	42	10	10	Yes*	Behavioral/personality	MR	P1
Shapiro ²⁴	P2	Μ	US (Caucasian)	45	50	34	49	No	Mood/affective	Dementia	P2
Burnstein <i>et al.</i> ²⁵	P3	М	US (Caucasian)	42	42	32	32	No	NS	Dementia	P3
Laurent et al. ²⁶	P4	F	French	43	43	40	43	No	Behavioral/personality	Dementia	P4
Laurent <i>et al.</i> ²⁶	P5	М	French	39	39	29	39	No	Behavioral/personality	Dementia	P5
Laurent et al. ²⁶	P6	М	French	28	28	23	9	Yes*	Behavioral/personality	MR	P6
Laurent et al. ²⁶	P7	F	French	49	49	8	<49	No	Mood/affective	Dementia	P7
Berginer et al. ²⁰	P8	М	Israel	42	43	22	< 32	Yes	Behavioral/personality	MR	P8
Berginer et al. ²⁰	P9	м	Israel	25	18	18	37	No	Psychosis	Dementia	P9
Berginer <i>et al.</i> ²⁰	P10	M	Israel	22	12	12	27	No	Catatonia	MR	P10
Berginer <i>et al.</i> ²⁰	P11	F	Israel	32	32	12	22	Yes	Behavioral/personality	MR	P11
Wevers et al ²⁷	P12	F	Dutch	37	37	25	25	Yes*	NS	MR	P12
Soffer et al 28	P13	F	Morrocan Jewish	30	37	< 5	24	Yes	Rehavioral/personality	MR	P13
Verrins et al 29	P14	F	Dutch	33	35	17	17	No	NS	MR	P14
Sperbake <i>et al</i> ³⁰	P15	M	Germany	52	52	< 6	< 6	Yes*	Rehavioral/personality	ID	P15
Sugama et al ³¹	P16	F	lananese	11	11	38	28	No	Behavioral/personality	Dementia	D16
Dotti et al 32	D17	M	Italian	30	2	2	2	2	Behavioral/personality	2	D17
Dotti et al. ³²	D10	E	Italian	10	2	2	2	2	Beychoois	: 2	D10
Dotti et al. ³²	D10	1	Italian	52	: 7	: 7	:	:	Mood/affective	: 2	D10
Dotti et al. ³²	F 19		Italian	22	:	:	:	:	Rehavioral/personality	:	F 19
Dotti et al. ³²	P20 D21		Italian	20	? 2	? 2	? 2	? 2	Bendvioral/personality	: 2	P20 D21
Dotti et al.	PZI		Italian	39	? 2	? 2	? 2	? 2	Psychosis Babayiaral/marsanality	? 2	PZI
Dotti et al.	PZZ DDD	F F	Italian	39	? 2	? 2	? 2	? 2	Behavioral/personality	? 2	P22
Dotti et al.	P23	г г	Italian	42	? 2	? 2	? 2	? 2	benavioral/personality	? 2	P23
Dotti et al.	P24	F	Italian	4/	?	? 2	? 2	?	benavioral/personality	<i>?</i>	P24
Dotti et al.	P25	IVI	Italian	54	?	? 2	? 2	?	Niood/affective/depr.	<i>?</i>	P25
Dotti et al.	P26	IVI	Italian	51	<i>?</i>	?	<i>?</i>	<i>?</i>	Psychosis	?	P26
Dotti et al.	P27	F	Italian	3/	{	!	?	<i>. !</i>	Psychosis	<i>?</i>	P27
Lee et al.	P28	F	Taiwan	31	41	6	29	Yes	Mood/affective	MR	P28
Lee et al.	P29	M	Taiwan	32	43	<5	Ado	Yes*	Behavioral/personality	MR	P29
Lee et al.	P30	F	laiwan	3/	48	<5	Ado	Yes*	Behavioral/personality	MR	P30
Grandas et al. ¹⁴	P31	F	Spain	39	51	26	26	Yes*	Mood/affective	MR	P31
Bartholdi et al.	P32	F	Swiss	51	51	6	49	No	Mood/affective	Dementia	P32
Guyant-Maréchal et al. ³⁴	P33	М	French	53	53	15	44	Yes	Behavioral/personality	Dementia	P33
Siman-Tov et al. ³³	P34	М	Israel (Ashkenazi)	29	29	4	29	No	Behavioral/personality	MR	P34
Price Evans et al. ²²	P35	F	Saudi	22	17	10	10	Yes*	Behavioral/personality	MR	P35
Price Evans et al. ²²	P36	М	Saudi	30	18	10	18	No	Behavioral/personality	MR	P36
Gonzalez-Cuyar et al.36	P37	F	USA	49	49	?	40 s	No	NS	MR	P37
de la Fuente <i>et al.</i> ³⁷	P38	М	Spain	52	40	18	18	Yes*	Behavioral/personality	Dementia	P38
Szlago et al. ³⁸	P39	М	Argentina	17	17	0	Ado	Yes	Behavioral/personality	MR	P39
Zacherl et al. ³⁹	P40	F	Germany	44	44	20	?	No	Mood/affective	Dementia	P40
Filippi <i>et al.</i> ⁴⁰	P41	Μ	Chile	39	20	< 20	Ado	No	Mood/affective	Dementia	P41
Bonnot <i>et al.</i> ⁸	P42	Μ	French-Spanish	23	24	6	<10	Yes	Behavioral/personality	LD	P42
Bonnot <i>et al.</i> ⁸	P43	F	French-Spanish	17	19	7	<10	Yes	Behavioral/personality	LD	P43
Chang <i>et al.</i> ⁴¹	P44	М	Chinese	24	29	?	?	No	Anxiety	MR	P44
Chang et al.41	P45	F	Chinese	31	48	?	?	No	Mood/affective	Dementia	P45
de la Fuente <i>et al</i> . ⁴²	P46	?	Spain	?	?	?	?	?	NS	?	P46
de la Fuente <i>et al.</i> ⁴²	P47	?	Spain	?	?	?	?	?	NS	?	P47
de la Fuente <i>et al</i> . ⁴²	P48	?	Spain	?	?	?	?	?	NS	?	P48
de la Fuente et al 42	P49	, 7	Spain	, 7	, ,	, ,	, ,	, ,	NS	2	P49
de la Fuente <i>et al</i> ⁴²	P50	7	Spain	7	7	7	7	7	NS	7	P50
de la Fuente <i>et al</i> ⁴²	P51	2	Spain	2	;	2	2	2	NS	7	P51
de la Fuente et al 42	D50	: 7	Spain	: 7	: 7	: 7	: 7	: 7	NS	: 7	P51
do la Euonto <i>et al</i> ⁴²	F 52 D52	: 2	Spain	: 2	: 2	: 2	: 2	: 2	NC	: 2	P 52
de la Eucete et al^{42}	P 3 3	: 2	Spain	: 2	: 2	: 2	: 2	: 2		: 2	P 3 3
ue la ruente et al.	r54	1	Spain	1	1	:	1	1	CVI	1	r54

Abbreviations: Ado, adolescence not specified; Cog, psychiatric manifestations associated with LD, MR or dementia; CTX, cerebrotendinous xanthomatosis; D, age at diagnosis; depr., depression; F, female; LD, learning difficulties; M, male; MR, mental retardation; N, age at neurological debut; NS, not specified; X, age at last examination; Ψ , age at psychiatric debut; Ψ^* , psychiatric initial presentation, psychiatric first referral of CTX or none of these; Ψ_s , psychiatric semeiology; Ψ_D (Axis), type of psychiatric disorder.

and 3). From the 34 out of the 54 patients for whom records on the age of onset and mode of onset were found, 17 patients had been brought to clinical attention because of psychiatric manifestations irrespective of patient age or coexisting neurological, ocular or dermatological CTX manifestations (Table 2, color-coded red). However, in only 9 out of these 17 CTX cases were psychiatric manifestations the first sign of disease that led to clinical evaluation, and this occurred always at a young age (6–26 years; Figure 2, Table 2, marked with an * in column Ψ^*). It must be pointed out that cognitive decline or established mental retardation coexisted in seven out of these nine cases at the time of psychiatric presentation, whereas in the remaining two cases psychiatric symptomatology occurred in the absence of overt

cognitive decline. In the remaining 8 out of the above mentioned 17 cases, psychiatric manifestations had appeared at an older age, and a more advanced disease stage and mental retardation or dementia were already established (Table 2, color-coded red, not marked with * in column Ψ^*). In the other 17 out of the above mentioned 34 CTX cases, psychiatric manifestations were a late feature of the disease in the setting of established CTX with either mental retardation or dementia, and in most cases the diagnosis had been made many years previously; hence, the psychiatric symptomatology did not aid the clinical diagnosis (Table 2, not color-coded in column Ψ^*). The gamut of psychiatric pathology in these patients was more broad, encompassing affective/mood disorders, psychotic disorders and psychiatric



Table 3. Detailed description of psychiatric history and manifestations of CTX patients drawn from the medical literature Psychiatric symptoms and signs (age) P1: (BEHAVIORAL/PERSONALITY disorder) Behavioral/personality disorder, (<10) P2: (MOOD/AFFECTIVE disorder) Depressive episode, suicidal ideation and threats, aggressivity, assault threats, disinhibition, poor impulse control, fecal incontinence (49) P3; NS (32) P4: (BEHAVIORAL/PERSONALITY disorder) Apathy, spasmodic laughter/cry (43) P5: (BEHAVIORAL/PERSONALITY disorder) Behavioral alteration, logorrhea, psychomotor excitation, distractibility, fleeting ideas, emotional lability, no spasmodic laughter/cry (39) P6: (BEHAVIORAL/PERSONALITY disorder) Behavioral/personality disorder (childhood), psychiatric institution (23) P7: (MOOD/AFFECTIVE disorder) Mood instability, affective disorder, predominantly depressive periods, phases of agitation/excitation even aggressivity (49) P8: (BEHAVIORAL/PERSONALITY disorder) Behavioral/personality disorder, irritability, agitation (6 psychiatric hospitalizations before 32) personality disturbance, hypersexuality?(32) paranoid id., suicidal (41) P9: (PSYCHOSIS) Psychosis atypical, paranoid delusion (37) P10: (CATATONIA) Catatonia reactive to family stress?, akinetic, mute, fixed postures, urinary-fecal incontinence (27) P11: (BEHAVIORAL/PERSONALITY disorder) Aggressivity, agitated, required many hospitalizations (22) P12: NS (25) P13: (BEHAVIORAL/PERSONALITY disorder) Behavioral/personality disorder, aggressivity, destructive behavior (24) P14: NS (between 17-33) P15: (BEHAVIORAL/PERSONALITY disorder) Behavioral/personality disorder (early childhood) P16: (BEHAVIORAL/PERSONALITY disorder) Behavioral/personality disorder, lack of affection and neglect towards family, restlessness, disinhibition, apathy, abulia, PREDEMENTIA (38-40) P17: (BEHAVIORAL/PERSONALITY disorder) Behavioral/personality disorder, behavioral changes P18: (PSYCHOSIS) Psychosis P19: (MOOD/AFFECTIVE disorder) Depression P20: (BEHAVIORAL/PERSONALITY disorder) Behavioral/personality disorder, Behavioral changes P21: (PSYCHOSIS) Psychosis P22: (BEHAVIORAL/PERSONALITY disorder) Behavioral/personality disorder, Behavioral changes P23: (BEHAVIORAL/PERSONALITY disorder) Behavioral/personality disorder, Behavioral changes P24: (BEHAVIORAL/PERSONALITY disorder) Behavioral/personality disorder, Behavioral changes P25: (MOOD/AFFECTIVE disorder) Depression P26: (PSYCHOSIS) Psychosis P27: (PSYCHOSIS) Psychosis P28: (MOOD/AFFECTIVE disorder) Depression, irritability, insomnia, crying spells, poor appetite, agitated; bullied constantly by her siblings with CTX since early teens (29) P29: (BEHAVIORAL/PERSONALITY disorder) Behavioral/personality disorder, aggressivity towards sister (adolescence) P30: (BEHAVIORAL/PERSONALITY disorder) Behavioral/personality disorder, Aggressivity towards sister (adolescence) P31: (MOOD/AFFECTIVE disorder) Depression (>26, 26-37) P32: (MOOD/AFFECTIVE disorder) Depression severe (49) P33: (BEHAVIORAL/PERSONALITY disorder) Behavioral/personality disorder, poor judgment, self-neglect, altered sexual behavior, dysexec, irritability, anxiety (44) PERSONALITY disturbance profound, breakdown in social behavior (53) P34: (BEHAVIORAL/PERSONALITY disorder) Behavioral/personality disorder, spasmodic laughter P35: (BEHAVIORAL/PERSONALITY disorder) Behavioral/personality disorder, BEHAVIORAl disorder, aggressivity, irritability, awake at nights, anxiety, fight and hitting and hitting people (10) P36: (BEHAVIORAL/PERSONALITY disorder) Behavioral/personality disorder, odd behavior could lose his job, but no aggressivity (>18) P37: NS (40s) P38: (BEHAVIORAL/PERSONALITY disorder) Behavioral/personality disturbance at military, introverted (adolescence), paranoid delusions, undifferentiated schizophrenia and schizoid personality disturbance (20-40) P39: (BEHAVIORAL/PERSONALITY disorder) Behavioral/personality disorder, ADHD (late childhood, adolescence) P40: (MOOD/AFFECTIVE disorder) Depression, benzodiazepine abuse, suicidal ideation P41: (MOOD/AFFECTIVE disorder) Mood/affective, bipolar disorder, schizophrenia P42: (BEHAVIORAL/PERSONALITY disorder) Behavioral/personality disorder, aggressivity, irritability P43: (BEHAVIORAL/PERSONALITY disorder) Behavioral/personality disorder, aggressivity, irritability P44: (MOOD/AFFECTIVE disorder) Anxiety P45: (MOOD/AFFECTIVE disorder) Dysthymia P46: NS P47: NS P48: NS P49: NS P50: NS P51: NS P52: NS P53: NS P54: NS

Abbreviations: ADHD, attention-deficit hyperactivity disorder; CTX, cerebrotendinous xanthomatosis; NS: not specified. In parentheses, age at onset of psychiatric symptomatology/semeiology.

manifestations typical of subcortical dementia, which, in some cases, were compatible to a clinical diagnosis of frontotemporal dementia (Figure 4).

Documentation of the type of psychiatric diagnosis existed for 41 out of the total of 54 patients found in the bibliography, with CTX and psychiatric manifestations at some point in the disease course (Tables 2 and 3). Behavioral/personality disorder was the most frequent psychiatric manifestation in the small group of young CTX patients with pure psychiatric presentation (n = 9) of

CTX, whereas in the larger group of CTX patients with nonpsychiatric presentation (n = 32) of CTX it was as frequent as all the other psychiatric manifestations together (Figure 3). For 31 out of the 41 CTX cases with documentation of the type of psychiatric diagnosis, there was also available information on its age of onset (Figure 4). Behavioral/personality disorder was the most frequent psychiatric manifestation in all age groups, but especially so in patients younger than 26 years, followed by mood/affective disorder. In patients older than 26 years, psychiatric pathology was more complex, with richer semeiology often in the context of predementia/dementia (Figure 4).

In conclusion, based on the 425 CTX case reports we reviewed, the overall incidence of psychiatric manifestations was 12.7%, whereas only 9 CTX cases, for which documentation was adequate to allow further analysis, presented with essentially psychiatric symptoms early in the disease course and in the absence of overt neurological disease (overall incidence 2.5%). In seven out of nine of these cases, the manifestation of a pure psychiatric presentation was behavioral/personality disturbance (overall incidence 1.7%). Interestingly, all seven CTX patients, with premonitory psychiatric manifestations in the form of behavioral/personality disorder, were younger than 18 years (range 6–18 years) and all suffered from learning difficulties or mental retardation (Tables 2 and 3). It is concluded therefore that psychiatric manifestations in CTX follow a bimodal/bitemporal pattern. They appear (a) either early in the disease course, that is, during childhood or adolescence, in the form of a behavioral/personality disorder associated with learning difficulties or mental retardation, or (b) they manifest in advanced disease in older patients, often in the context of dementia as rich neuropsychiatric syndromes, such as frontal, orbitofrontal or frontotemporal syndromes of cortico-subcortical dementia encompassing behavioral/personality disturbance, affective/mood disorders and/or psychotic disorders.

DISCUSSION

CTX (also known as sterol 27-hydroxylase deficiency, cholestanolosis, van Bogært-Scherer-Epstein disease or simply van Bogært's disease) is a rare, inborn neurometabolic disorder of bile acid biosynthesis and sterol storage with autosomal recessive inheritance and variable clinical presentation.^{1-5,23,26,49-53} The mutated gene, the sterol 27-hydroxylase gene (CYP27), on the 2q33ter locus, codes for a mitochondrial cytochrome P450, which associates with two protein cofactors (adrenodoxin and adrenodoxin reductase) and catalyzes the hydroxylation at position C27 of the side chain of sterol intermediates, which constitutes the first step in their oxidative cleavage.^{5,54–57} More than 50 different mutations of the gene have been described, most of them within the adrenodoxin-binding and the heme-binding sites of the enzyme.^{43,58–64} As a consequence, CDCA synthesis is deficient and so is the negative feedback that CDCA exerts on cholesterol synthesis, with resultant increase in cholesterol synthesis and shunting of cholesterol towards cholestanol (5a-dihydrocholesterol) and other bile alcohol synthesis byproducts.^{5,51–58,65} Accumulation of cholestanol, as well as cholesterol and various bile alcohols, in large deposits in many organs, particularly in the CNS, tendons, vascular system and lungs, is the pathophysiological mechanism behind CTX.^{5,51,52,54,65-69} In the brain, cholestanol deposition far exceeds that of other sterols, ultimately comprising up to 30% of total sterols in the cerebellum. 36,54,70 Interestingly, despite increased cholesterol synthesis, plasma total cholesterol remains below or within the normal-range byproducts.^{5,49,51–58,70} Laboratory diagnosis rests on the findings of elevated cholestanol in the plasma and cerebrospinal fluid, and the presence of bile alcohols (such as lathosterol, 7α -hydroxylated bile acids and so on) and bile alcohol glycuronides in the plasma and/or urine.^{51,52,63,64,70} In bile, cholestanol is increased but CDCA is virtually absent.^{51,52,63,64,70} Diagnosis is confirmed by mutation analysis, and genetic counseling is necessary for at-risk relatives, as appropriate treatment can alleviate symptoms and slow disease progression in symptomatic patients, and, most importantly, prevent or mini-mize disability in asymptomatic homozygotes.^{43,59} The mainstay treatment consists of CDCA (Chenofalk) oral supplementation, 250 mg three times daily, which is the lacking bile acid that reinstitutes the negative feedback and balance in the cholesterol and bile acid synthetic pathways, respectively, and normalizes cholestanol levels in the blood.^{3,4,6,70–72} Cholestanol levels can be further reduced by the addition of a statin, usually simvastatin or pravastatin, but without an evidence-based clinical benefit.^{9,70–73} Treatment is effective in that it can alleviate some constitutional, neurologic and even psychiatric symptoms.^{4,6–8,70–80} Electrophysiological, neuropsychological and neuroradiological evaluation results support the notion that timely and adequate specific treatment may partially reverse neural tissue damage in the CNS and peripheral nervous system.^{41,72–80}

More than 400 cases have been described previously worldwide in the medical literature, and psychiatric manifestations of CTX have been reported only rarely and sporadically.²⁰⁻⁴³ To date. no comprehensive review of the psychiatric manifestations of CTX has been published. Psychiatric manifestations in CTX present a variability that reflects the variability of the rest of CTX semeiology. These encompassed behavioral/personality disorders, affective/ mood disorders, psychotic disorders and dementia-associated psychopathology (Tables 2 and 3). We found that psychiatric manifestations in CTX follow a bimodal/bitemporal pattern, that is, they would either appear early in the disease course in the form of behavioral/personality disorder associated with learning а difficulties or mental retardation, or they would manifest in advanced disease in the context of dementia as rich neuropsychiatric syndromes, such as frontal, orbitofrontal or frontotemporal syndromes of cortico-subcortical dementia encompassing affective/mood disorders, psychotic disorders or behavioral/ personality disturbance (Tables 2 and 3).

Unfortunately, many of the sporadic cases in the medical literature were poorly documented. The Dotti et al.³² and the de la Fuente et al.⁴² series comprised, respectively, 11 and 9 patients with psychiatric symptoms, yet these were presented monolectically and the evaluation of their importance and temporal correlation with non-psychiatric manifestations was not possible. Among 13 CTX patients with ocular signs scrutinized by Dotti et al.³², 11 had a psychiatric disturbance (Table 2), of which 5 had behavioral changes, 4 had psychosis and 2 had depression. Thus, clearly, in this series the psychiatric manifestations were overrepresented, which is in contrast to the rarity of psychiatric signs in CTX. In the only published case series specifically focusing on the psychiatric spectrum of CTX, Berginer et al.²⁰ reported four patients with disparate psychiatric syndromes. In this series, the first patient required numerous hospitalizations before and after the age of 32 years for irritability and personality changes with occasional hypersexuality and ended up with paranoid delusion and suicidal ideation at the age of 41 years. The second patient presented at the age of 37 years with atypical psychosis and paranoid delusion. The third patient suffered a severe catatonic episode at the age of 27 years and also had urinary-fecal incontinence. The last patient had been referred to a psychiatrist at the age of 22 years for aggressivity and agitation, and ultimately required many hospitalizations. In all patients of this series, however, the diagnosis of CTX had been easily made 10-15 years before the appearance of the psychiatric manifestations, as all four patients had had pes cavus and Achilles tendon xathomas since early or late adolescence, two out of four patients had learning difficulties and cognitive deterioration since childhood, and two out of four patients had bilateral cataracts. Importantly, by the time of the psychiatric evaluation, three out of four patients had already developed spastic paraparesis and cerebellar ataxia, that is, the disease had advanced enough to extensively involve the CNS.

Psychiatric manifestations and dementia in CTX are primarily the result of diffuse white matter pathology affecting multiple distributed intrahemispheric and interhemispheric neural networks. White matter pathology is caused by alterations of myelin lipid composition with markedly disproportional cholestanol incorporation into the glial cellular membranes, substitution of other sterols and extensive demyelination of white matter



Figure 1. Different psychiatric manifestations related to cerebrotendinous xanthomatosis (CTX) and their frequency distribution. Patients with mood/affective disorders all suffered from depression, except for one patient who suffered from dysthymia. NS: not specified. Behavioral/personality and mood/affective disorders are clearly the predominant psychiatric manifestations in CTX (patients, n = 54).



Figure 2. Incidence of disease presentation according to age group. Pure psyciatric and non-psychiatric presentation are plotted as bars of different color for the same age group. Pure psychiatric presentation of cerebrotendinous xanthomatosis (CTX) occurs most often in childhood (childhood: ≤ 12 years; adolescence: 13 years $\leq age \leq 18$ years; early adulthood: 19 years $\leq age \leq 26$ years; late adulthood: age > 26 years (patients, n = 34).

tracts.^{53,54,56,57} However, alterations of the composition of the lipid bilayer of neuronal plasma membranes may also have repercussions in neuronal integrity and/or function at the level of soma, synapse or circuitry.^{41,44,45,53,79} Moreover, intracerebral lipid deposition in the form of xanthomas and local inflammatory reaction take their toll on myelinated axonal tracts, central grey matter formations and neuronal cell bodies.^{66–69} Thus, both demyelination and inevitable neuronal loss ultimately lead to the neurologic and complex psychiatric manifestations of CTX. This



Figure 3. Frequency of behavioral/personality disorder in cerebrotendinous xanthomatosis (CTX) patients with pure psychiatric presentation as compared with patients with non-psychiatric presentation of CTX. Behavioral/personality disorder is the most frequent psychiatric manifestation in the group of CTX patients with pure psychiatric presentation (n = 9) of CTX, whereas it is as frequent as all the other psychiatric manifestations together in the group of CTX patients with non-psychiatric presentation (n = 32) of CTX (patients, n = 41).



Figure 4. Distribution of cerebrotendinous xanthomatosis (CTX) patients with psychiatric manifestations according to age group and type of clinical psychopathology (childhood: ≤ 12 years; adolescence: 13 years $\leq age \leq 18$ years; early adulthood: 19 years $\leq age \leq 26$ years; late adulthood: age > 26 years (patients, n = 31).

hypothesis is supported by anatomopathological findings, and both conventional and advanced neuroimaging.^{3,4,13,28,41,} ^{44–46,53,79–81} Despite the definite organic etiopathogenesis of the neurobehavioral manifestations of CTX, depressive and anxiety symptoms, which constitute the second most common psychiatric manifestation in CTX, may sometimes be attributable in part to the physical and psychosocial impact of a chronic neurologic disease



and cognitive decline on role functioning and quality of life, especially in younger patients without dementia.⁸²

Multiple sclerosis (MS) is another CNS disease, where primarily white matter and secondarily grey matter are damaged in a random fashion, albeit by a different pathogenetic mechanism. MS may cause cognitive disturbance or dementia and, less often, organic psychopathology encompassing behavioural/per-sonality disturbance, mood/affective disorders or psychosis.^{83–104} Psychiatric syndromes, such as bipolar disorders and atypical psychosis, often unassociated with cognitive disturbance or dementia, have been described, and euphoria sclerotica is a characteristic psychosyndrome in MS.^{83–85,89,91,94–96,99,104–108} Pure psychiatric presentation in MS is rare, as is the case for CTX.^{83,89,91,106,107} Major depressive disorder, which often is prodromal leading to diagnostic delay, is overrepresented in MS patients, as are suicidal ideation, anxiety and sleep disorders, and they are mainly believed to be secondary to physical illness and/or disease-modifying treatment.^{84–87,90,93,95,96,98,99,102,104,107,108} Athough in MS a detrimental autoimmune process is the culprit,

and not an enzymatic metabolic defect as in CTX, the relative infrequence, variability and complexity of psychiatric signs in both these diseases can be explained by the stochasticity and widespread nature of neural tissue damage, and the prerequisite of a cumulative effect and critical burden of white/grey matter loss or dysfunction.^{85,86,90,103,104,106–113} Moreover, one can further classify the organic neuropsychiatric syndromes associated with CTX and MS, as primarily related to the integrity of subcortical myelinated systems, as is also reflected in similarities in their neuropsychological profiles with cognitive slowing, dysexecutive syndrome and attentional disorder.^{107–112} Indeed, the characteristics of dementia in the setting of CTX and MS merit their classification as white-matter dementias.^{109,112} Systematic advanced neuroimaging studies with positron emission tomography, single photon emission computerized tomography, magnetic resonance spectroscopy and diffusion tensor imaging are needed to clarify the importance of white matter functional, biochemical and microstructural abnormalities in the cognitive and neurobehavioral syndromes associated with CTX at various stages in the disease course, correlate these findings to neuropsychological assessment parameters and scores, as well as monitor the effects of disease-modifying treatment.^{41,79,80,113,114}

Several other hereditary lipid storage disorders that cause leukoencepalopathy or leukodystrophy may result in or rarely present with complex organic neuropsychiatric phenotypes by affecting the cerebral white matter integrity and/or the neuronal elements and circuitry.^{108–144} In most lipid storage diseases of the nervous system (neurolipidoses) or leukodystrophies, which can present at some point with acute, progressive or relapsing psychosyndromes (Tay–Sachs and Sandhoff disease, Fabry disease, metachromatic leukodystrophy, adrenoleukodystrophy, Niemann-Pick type C, ceroid neuronal lipofuscinoses), pharmacotherapy of psychiatric manifestations has either poor effect or even worsens the condition, for example, as in adult-onset GM₂ gangliosidoses, where phenothiazines and tricyclic antidepressants may have a detrimental effect by inhibiting the activity of lysosomal enzymes, thereby increasing lipid storage in neuronal cells.¹¹⁵⁻¹⁴³ This is a little known domain of neuropsychiatry, because neurodegenerative/neurometabolic diseases are rare and seldom end up in the principal care of the psy-chiatrist.^{8,115,117,134,142} The latter happens when these conditions present either as complex neuropsychiatric syndromes or, more often, as cognitive/behavioral syndromes or learning difficulties in young age.^{8,115,117,119,120,122,125,127,131,133,134,142}

Although psychiatric manifestations in CTX are not uncommon, psychiatric presentation in CTX is rare and, therefore, is the initial pharmacotherapy of CTX with psychoactive drugs.⁸ Our own clinical experience from the CTX patients evaluated at the La Salpêtrière University Hospital is that pharmacotherapy (anxiolytic,

antidepressive or antipsychotic treatments) and psychotherapy (cognitive-behavioral treatments) have little or only temporary effects in CTX-related organic psychosyndromes without improving prognosis, both in patients with advanced disease and dementia, and also in younger patients with mild cognitive disturbance. However, if psychiatric symptomatology is compounded by secondary depression or anxiety disorder in CTX, it may be partially amenable to symptomatic pharmacotherapy with antidepressants or anxiolytics, such as selective serotonin reuptake inhibitors, and a treatment trial in a CTX patient with mild or major depressive or anxiety disorder is anavoidable, as it is warranted.⁸² However, specific disease-modifying treatment with CDCA has the potential to both alleviate symptomatology and improve prognosis by slowing down neurodegeneration and disease progression, thus preventing or even reversing psychiatric manifestations.⁸ This is all the more important for the young undiagnosed patient with CTX that presents to the child and department, adolescent psychiatry with an elusive psychosyndrome that is slowly but steadily progressive and is often associated with cognitive disturbance.^{8,144} Therefore, timely diagnosis and specific treatment is the ultimate goal in the management of neuropsychiatric manifestations of CTX.

In conclusion, psychiatric manifestations in CTX are variable, encompassing personality, affective and psychotic disorders, and either appear late in the course of the disease in the context of an organic dementia complicating a chronic neurologic tableau of spastic paraparesis and the cerebellar or early in the course of the illness preceded by cognitive disturbance. Chronic diarrheas, tendon xanthomas and cataracts are salient early features of this systemic illness that herald the neurological or psychiatric signs and should raise the diagnostician's index of suspicion.145-The earlier the onset of psychiatric symptoms, the milder is their severity, with personality/behavioral disorder, irritability and aggressivity most prominent in the 'younger than 25' age group. Irritability, aggressivity, attention-deficit hyperactivity disorder and oppositional-defiant disorder in childhood or adolescence, which easily can be disregarded or misdiagnosed as 'travails of youth' or 'peer-group influence', should therefore prompt a thorough neurologic evaluation with a complete physical and ophthalmologic check-up, especially in a background of consanguinity that increases the probability for autosomal recessive inherited disorders.¹⁴⁴ The attending psychiatrist should be aware of CTX, its psychiatric manifestations and its characteristic physical signs, and strive to exclude it, especially as anecdotal reports have shown a beneficial effect of timely CDCA treatment on psychiatric symptoms and overall prognosis.8,145-148 We hope that this review will raise awareness of this devastating but treatable neuropsychiatric inherited disorder among psychiatrists.

CONFLICT OF INTEREST

The author declares no conflict of interest.

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