Niemann-Pick disease type C (NPC) is an autosomal recessive neurometabolic disorder that rarely presents in adulthood, and is associated with cognitive decline, various movement disorders (ataxia, chorea, dystonia, and myoclonus), a vertical supranuclear gaze palsy (VSGP), and seizures. A recent case report demonstrated a delay in diagnosis of eight years when a patient with NPC presented with psychosis. This article reviewed all cases seen at the Mayo Clinic with a possible diagnosis of NPC between 1976 and 2000. Of the 52 possible cases, five had an established diagnosis of adult onset NPC. Of these, two presented with psychosis and were not diagnosed with NPC for 5 and 15 years, respectively. NPC may initially present in adulthood with psychosis, and when psychosis is associated with VSGP, various dyskinesias, and seizures, NPC should be suspected.

RESULTS
Of these five adult onset NPC cases two presented with psychosis.

Case 1
A 61 year old woman was transferred to the Mayo Clinic with a 15 year history of an undiagnosed, progressive neuropsychiatric disorder, rendering her bed bound and unable to eat.

The patient initially developed depression and hypersomnia at age 46. Her symptoms improved with an antidepressant. At 49, she was hospitalised for mood lability, loquacity, delusions, and hypervigilance. She was treated with thiothixine, imipramine, and lithium then discharged. At age 50, she developed a gait disorder with postural instability, was noted to “lean backwards” and have retropulsion. This was attributed to tardive Parkinsonism. She was treated with levodopa and bromocriptine, and her gait improved. However, she again became hypersomnolent, as well as dysphagic, which required gastrostomy tube placement. One year later, auditory hallucinations began, and she became paranoid, hyperreligious, and obsessive, for example, making an excessive number of long distance telephone calls. Her differential diagnoses included: schizoaffective, bipolar, and organic affective disorders. She was then treated with thiroidazine and amitriptyline. At age 55, she was noted to have saccadic pursuits, a hypokinetic and ataxic dysarthria, bradykinesia, and her first documented seizure. Between ages 55 and 61, she became demented, rigid, and virtually mute. She was transferred to the Mayo Clinic for evaluation at age 61.

On neurological examination, the patient was essentially mute. She had cranio cervical dystonia, axial rigidity, and stimulus sensitive myoclonus. Eye movement examination was hampered by severe blepharospasm. MRI of the head showed subtle small foci of increased T2 signal changes within pons and cerebral white matter (see fig 1A). Bone marrow biopsy revealed a normocellular marrow with increased foamy macrophages. Skin fibroblast testing showed 5.5% cholesterol esterification and a positive filipin staining consistent with a diagnosis of NPC. Genetic testing was not performed.

She died at age 62.

Case 2
A 32 year old woman presented to Mayo Clinic with a five year history of an undiagnosed neuropsychiatric disorder.

Her birth and development had been unremarkable. At the age of 27, she developed a “nervous breakdown,” associated with paranoid delusions and was treated with haloperidol. Later that year, her psychosis resolved, but she developed facial dystonia, a shuffling gait and dysarthria, attributed to a tardive syndrome. Subsequently she experienced an annual recurrence of her psychiatric symptoms, particularly the paranoid delusions. At 32, she became dysarthric, ataxic and
neurological examination revealed a vertical supranuclear gaze palsy (VSGP). She was give intravenous thiamine for possible Wernicke's encephalopathy. A small bowel biopsy was performed revealing lymphangiectasias, “suggestive of Whipple’s” and she was treated with intravenous antibiotics. However, her decline continued, and she developed urinary incontinence and insomnia.

Neurological examination at Mayo Clinic revealed oromandibular dystonia, an ataxic and hypokinetic dysthria, mild hyperreflexia, and gait ataxia. MRI of the head showed increased T2 and fluid attenuated inversion recovery signal in bilateral basal ganglia and periventricular areas (see fig 1B). Duodenal biopsy was negative. Brain biopsy revealed mild gliosis. One month later, neurological examination revealed virtual mutism, definite corticospinal tract signs, and a VSGP. Duodenal biopsy demonstrated scattered sea blue histiocytes. Skin fibroblast testing showed 26% cholesterol esterification and indeterminate filipin staining. She was diagnosed with variant type NPC. A year later, a repeat cranial MRI showed extensive white matter disease. Genetic testing was not performed.

**DISCUSSION**

Of our five confirmed cases with adult onset NPC two (40%) presented with psychosis. This is higher than the 25% (4 of 16) noted in the review by Shulman et al and may reflect a referral bias of complicated cases to the Mayo Clinic. Alternatively, the higher prevalence of psychosis at onset noted in this small series of adult onset NPC cases, may, more accurately reflect true prevalence, as our cases were limited to those with cytochemical confirmation.

Patients with inborn errors of metabolism are usually diagnosed in childhood. To date, there are less than 25 reported cases of adult onset NPC. Compounding the diagnostic confusion surrounding our two patients was their initial symptom: psychosis. In the single extant report of a patient with adult onset NPC presenting with psychosis, several renowned movement disorder authorities underscore that such a case is extraordinary. Our two patients demonstrate that adult onset NPC presenting with psychosis is not unique, though it may be rare. Our case histories reveal clues that may aid clinicians in recognising similar patients. For example, the presence of a VSGP in a progressive disorder narrows a patient’s diagnostic possibilities considerably to: progressive supranuclear palsy, corticobasal degeneration, multiple systems atrophy, dementia with Lewy bodies, Whipple’s disease, Wilson’s disease, Huntington’s disease, neuroacanthocytosis, and NPC.

Another potentially illuminating feature of our two cases is that the patients developed probable tardive parkinsonism and other dyskinesias readily after receiving neuroleptics. This is reminiscent of what occurs in patients with dementia patients with Lewy bodies who receive neuroleptics. Additionally, it may be difficult to ascertain whether patients with neuroleptic exposures, who later develop dyskinesias, have a tardive phenomenon or not, as it was in our patients. In either case, one should consider the possibility of an undiagnosed disease affecting the basal ganglia in psychotic patients who abruptly develop dyskinesias after ingesting neuroleptics.

Organomegaly is common in patients with inborn errors of metabolism, including NPC; however, hepatosplenomegaly may be present in only 50% of patients with adult onset NPC, so its absence should not dissuade one necessarily from this diagnosis. The presence of seizures also is typical in adult onset NPC.

In summary, adult onset NPC may present in adulthood with psychosis. When a patient less than 60 years of age presents with the tetrad of psychosis, a VSGP movement disorders, and seizures, NPC is a possible aetiology.

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Adult onset Niemann-Pick disease type C presenting with psychosis

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