Prevalence of serum antibodies to caudate nucleus in autistic children

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Abstract

Autism may involve autoimmunity to brain. We studied regional distribution of antibodies to rat caudate nucleus, cerebral cortex, cerebellum, brain stem and hippocampus. The study included 30 normal and 68 autistic children. Antibodies were assayed by immunoblotting. Autistic children, but not normal children, had antibodies to caudate nucleus (49% positive sera), cerebral cortex (18% positive sera) and cerebellum (9% positive sera). Brain stem and hippocampus were negative. Antibodies to caudate nucleus were directed towards three proteins having 160, 115 and 49 kD molecular weights. Since a significant number of autistic children had antibodies to caudate nucleus, we propose that an autoimmune reaction to this brain region may cause neurological impairments in autistic children. Thus, the caudate nucleus might be involved in the neurobiology of autism.

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Autism is an early-onset disorder of the developing central nervous system (CNS). The disorder causes severe deficits of higher mental functions such as social interaction, language, communication, imagination and cognition. Today, autism is the fastest-growing developmental disability in children. Autism affects more than one-half of a million Americans and millions more worldwide but very little is known about the etiology and pathogenesis of the disorder. Current theories include genetic factors, immune factors, environmental factors and yet other unidentified factors. We focused on immune factors such as autoimmunity and provided supporting evidence for ‘autoimmune hypothesis’ in autism [9–14]. Autoimmunity in autistic children is shown by several autoimmune factors: brain-specific antibodies [10], impaired lymphocyte functions [3,6,15–17], abnormal cytokine regulation [11], viral associations [10,12,13] and indirect association of certain immunogenetic factors [16]. In the present study, our goal was to determine if autistic children harbor brain-specific antibodies, particularly with regards to different brain regions. We selected five brain regions for our study: caudate nucleus (Cn), cerebral cortex (Cx), cerebellum (Cb), brain stem (Bs) and hippocampus (Hpc). Laboratory results described here provide initial evidence for an autoimmune reaction to caudate nucleus in children with autism.

The study included 68 autistic children aged 4–12 years and 30 normal children aged 5–12 years. Of the 68 autistic children, eight had a diagnosis of pervasive developmental disorder (PDD) and the remainder 60 cases had a diagnosis of autism. The clinical diagnosis of autism and/or PDD relied essentially on the standard DSM-IV criteria of the American Association of Psychiatrists, Washington, DC. The Institutional Review Board or the ethical committee of Utah State University reviewed and approved our research protocol that involved the use of blood samples from children. All children were at the baseline without any treatment with prescription medications at the time of blood collection or at least 2 weeks prior to it.

Brain antibodies were detected by our standard immunoblotting technique [10,13]. Sprague–Dawley rats were sacrificed to remove brain, which was dissected to collect Cn, Cx, Cb, Bs and Hpc regions. Each brain region was separately homogenized in Tris-buffered saline containing 0.05% Tween-20 (TBST) and stored frozen at −20 °C. Neural proteins were separated in 12% Ready Gels (Bio-Rad, Richmond, CA) by sodium dodecyl sulfate-polyacrylamide gel electrophoresis, transferred to nitrocellulose membranes by the double sandwich technique, followed by blocking with 1% bovine serum albumin in TBS. For immunoassay, 3–4
mm wide blots were incubated with a screening dilution of 1:25 of patient or control sera for 1 h. They were washed four times with TBST, followed by 1 h incubation with 1:2500 diluted alkaline phosphatase conjugated-goat anti-human polyvalent immunoglobulins (Sigma, St. Louis, MO). After four washings with TBST, the blots were developed in substrate solution according to instructions from the manufacturer of the AP substrate kit (Bio-Rad). A reaction was scored positive only if a purplish-blue band appeared. To prevent potential bias, all samples were analyzed in a double-blind fashion and the code was broken after the completion of the laboratory analysis. For molecular weight determination, we used kaleidoscope pre-stained protein standards (Bio-Rad) that included myosin (237 kD), β-galactosidase (134 kD), bovine serum albumin (85 kD), carbonic anhydrase (43.1 kD), trypsin soyabean inhibitor (32 kD), lysozyme (17.4 kD) and aprotinin (7.2 kD).

Immunoblotting screen showed antibody positive reactions to neural proteins of the Cn, Cx and Cb. Autistic serum showed a positive reaction but normal serum did not (Fig. 1). We found that the highest number of antibody-positive autistic sera was with the Cn region (49%) as compared to Cx region (18%) or Cb region (9%) (Fig. 2). All sera were negative for antibodies to Bs and Hpc regions either because there was no antibody reaction (as with Bs region) or it was non-specific and indistinguishable between patients and controls (as with Hpc region). This finding suggests that the neurons in the caudate nucleus might be a target of autoimmune pathology in autism.

Because Cn was the most immunoreactive brain region, an attempt was made to extract proteins from this region. The protein gel pictures are not included here but the protein patterns of Cn homogenate and Cn supernatant (CnS) resembled quite closely. As shown in Fig. 3, antibodies in autistic sera reacted with two major proteins of Cn homogenate having molecular weight of approximately 160,000 (band A) and 115,000 (band B). These two proteins also showed positive reactions with CnS but an additional protein band (band C) was positive at a molecular weight of
approximately 49,000 (Fig. 3). Considering the fact that Cn and CnS contain a large number of proteins, it is interesting to find only two to three proteins positive for antibodies. This must imply a significant level of specificity with respect to autoantigens. To that end, it is worth noting that the caudate nucleus is enriched with serotonergic receptors [7], and antibodies to serotonergic receptors have been found in autistic children [14]. Naturally, while other possibilities exist, we think it is instructive to consider the possibility of an autoimmune reaction to serotonergic nerve-endings of the caudate nucleus. Once they have been fully characterized, these proteins could potentially serve as markers of neuropathology in autism. This topic is currently under investigation in our laboratory.

Fig. 3. Representative blots showing antibody positive reactions of caudate nucleus with autistic sera. As shown by black arrows, there were two antibody-positive proteins (bands A and B) in the caudate nucleus (left hand blots) and caudate nucleus supernatant (right hand blots), but the antibody reaction also occurred with an additional protein (band C) of the caudate nucleus supernatant.

Studies of autistic brain are extremely rare despite pathognomonic manifestations in people affected with the disorder. Brain autopsies, albeit of only a few cases, have shown size reduction in brainstem [5] and loss of cellular density of Purkinje Cells in the cerebellum [8]. The other abnormalities include limbic structures and forebrain [1]. Magnetic resonance imaging studies have shown hypoplasia of cerebellar vermal lobules VI and VII [2]. Delayed maturation of the frontal cortex [18] and developmental abnormalities of the myelin sheath [10] have been described in the brain of autistic children. As reported here, we did not find antibodies to brainstem or hippocampus and only a few cases had antibodies to cerebellum or cerebral cortex. This finding may indicate that autoimmune process is probably not the reason for abnormalities in these regions of the autistic brain. Besides these regions, we found antibodies to caudate nucleus in approximately one-half of the autistic children, which is a significantly high number of cases that we studied. These antibodies may impair neuronal functions in caudate nucleus thereby manifest neurological and behavioral symptoms in children with autism. Some credence to this line of thinking may come from a recent study of Tourette syndrome children who also harbor anti-striatal antibodies; upon infusion into the rat striatum, these antibodies caused neuronal dysfunction that was similar to the one found in Tourette syndrome children [4]. While the further characterization of Cn-derived autoantigens is in progress, we think that this new immunological evidence may suggest the involvement of caudate nucleus in the neuropathology of childhood autism.

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References


