Behavioral and cognitive core domains shared between autism spectrum disorder and schizophrenia

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ABSTRACT
Growing genetic evidence including results of genome-wide association studies and whole-genome sequencing is still perplexing scientists in the fields of human complex traits and compels them to reconsider the relationship between autism spectrum disorder (ASD) and schizophrenia. The developmental trajectory of schizophrenia may be characterized by difficulties in “theory of mind” tasks, poor insight or imagination, low empathy level, weak executive function, failure in social judgments, delayed language development, movement mannerisms, motoric rituals, strong preoccupation with unusual objects, bizarre habits, and high intelligence. Most of these characteristics are associated with the standard criteria for ASD and all of them are core domains or endophenotypes of ASD. The diagnosed ASD cases, which were found in individuals with schizophrenia, have a significant tendency to be high-functioning and to have atypical age of ASD manifestation. Episodic features including reality distortion, which can be seen in individual with ASD, may be associated with hyperarousal conditions with cognitive hypersensitivity, misattribution of negative experiences, and the perceptual thinking style. Together with the emerging new genetic concepts concerning human behavioral complex traits, the results and constructions of these recent studies may further warrant that reconsideration of the relationship between ASD and psychotic conditions is necessary. In addition, the possibility that psychotic conditions are the secondary derivative states in individuals with ASD or in individuals with subclinical ASD components should be constructively examined.

Keywords: Autism Spectrum Disorder (ASD); Schizophrenia; Reality Distortion; Psychotic Conditions

1. INTRODUCTION
Different perspectives cause distinct perceptions and interpretations of a condition. When the apparent difference among the perceptions or interpretations is recognized to be significant, a single condition could be misunderstood to be multiple separate categorical entities. In the conceptualization of the relation between symptoms and diagnostic categories, latent core structures underpinning the symptoms and causal networks consisting of the symptoms are recognized as important prerequisites for the categorization [1]. Therefore, probing the core domains or structures and considering the interrelationship between the domains may be quite worthwhile to evaluate the categorical identity of conditions.

The categorical identity of a lot of human behavioral and psychiatric conditions is now being compromised by recent results of genetic studies [2-9], twin studies [10-14], and endophenotypic reconsideration [15-18]. Such conditions include autism spectrum disorder (ASD), schizophrenia, attention-deficit hyperactivity disorder, learning disorders, obsessive-compulsive disorder, bipolar disorder, major depressive disorder, anxiety disorders, eating disorders, posttraumatic stress disorder, and personality disorders. In this genetic and endophenotypic complexity, ASD and schizophrenia are essential as the prototypic diagnostic categories whose genetic and phenotypic characteristics are being reexamined or compared among these conditions. The distinction between autism and the preceding clinical entity, schizophrenia, had been well attempted and should have already been completed [19]. However, the conviction that the ostensible lack of the first rank schizophrenic symptoms (Schneider) can distinguish autism from the early onset schizophrenia variant only implies that autistic children cannot be categorized as a subset of schizophrenia. Although the age of
onset had been recognized as an important firm ground for the distinction between autism and schizophrenia [19], the onset age of autism (ASD) in a standard diagnostic criteria (Diagnostic and Statistical Manual of Mental Disorders, DSM) is going to be changed from “prior to age 3 years” (DSM-IV-TR) to “in early childhood” (DSM-5). Significantly, it is additionally documented in DSM-5 that autistic symptoms may not become fully manifest until social demands exceed limited capacities. Therefore, even the taxonomic identity of schizophrenia should be reconsidered and a new insight whether schizophrenia is a subset of ASD should be examined [20-22]. Here we review recent results of studies in relation to behavioral and cognitive core domains shared between ASD and schizophrenia.

2. AUTISTIC DEVELOPMENTAL FEATURES OF SCHIZOPHRENIA

The developmental pathological model for schizophrenia was originally established on the basis of the speculated relationships between schizophrenia and some epiphenomena including nonspecific abnormal brain pathologies [23,24]. However, both the specific pathological marker and the molecular cause of schizophrenia remain, strictly speaking, unknown [25]. There is also no biological marker for ASD [26]. Recent findings are providing further support for developmental similarities in social cognitive deviations between ASD and schizophrenia, and the developmental trajectory of schizophrenia may be characterized by difficulties in “theory of mind” tasks, poor insight or imagination, low empathy level, weak executive function, failure in social judgments, and delayed language development [27-31]. These characteristics are all associated with the persistent deviations in social communication and social interaction in the standard criteria for ASD. The autistic pattern of social cognition is significant especially in individuals with the negative schizophrenia symptoms [27], and the executive function deviation is correlated with the severity of negative symptoms [30]. In addition, it is also well known that individuals with schizophrenia often display restricted and repetitive patterns of behavior, interests, or activities [32]. These repetitive behaviors in schizophrenia include movement manners, motoric rituals, strong preoccupation with unusual objects, and bizarre habits [32]. A longitudinal cohort study revealed that children with speech delays or rituals/habits were more likely to develop psychiatric experiences in early adolescence [22]. The diagnosed ASD cases, which were found in individuals with schizophrenia, have a significant tendency to be Asperger syndrome and to have atypical age of ASD manifestation [33].

The most unique and potentially meaningful developmental property of autistic cognition is savant skill. The estimated prevalence of the cognitive superiority in ASD varies from 10% to surprising numbers [34-36]. This discrepancy among estimations could be explained by the presence of high intelligence autistic individuals with low IQ [37]. Regardless of the IQ level, autistic individuals may possess high processing speed, prodigious memory capacities, and heightened primary sensory processing [37-39]. These cognitive superiorities are believed to have the same origin as the social difficulties in ASD [40], and the term, “autistic savant skills”, is used to describe one of the core cognitive features of ASD [37,41]. Importantly, creative people with high intelligence have an elevated risk for developing certain forms of psychopathology including schizophrenia [42,43]. To interpret the striking increase of genetic structural variations including copy number variations in the last common ancestor of humans and chimpanzees, an evolutionary trade-off theory is provided in association with ASD and schizophrenia [44,45]. The benefits in the trade-off theory may be something benefical resulted from expanding the range of genetic diversity [44] or reproductive something benefical associated with the creativity [42,46].

3. EPISODIC FEATURES IN INDIVIDUALS WITH AUTISM

Episodic characteristics or acute exacerbation of behavioral problems should be associated with the personal experiences and environmental changes and can be influenced by the individual background knowledge. Because what we can learn from experience and our responses to environmental changes are highly age-related, episodic contents of a behavioral condition should change by age. In children with ASD, temper tantrums and self-injurious behaviors correlate with autistic traits [47]. The clinical picture of ASD can change with increasing age and in different circumstances [48], and the behavioral plasticity or behavioral improvement is evident in supportive circumstances in individuals with ASD [26], suggesting possible emergence of age-related problematic episodes in unsupportive circumstances. Hyper- or hypo-reactivity to sensory input or unusual interest in sensory aspects of environment, which is introduced in DSM-5 as ASD criteria, may affect the episodic characteristics of ASD. Hypo-prior-based ubiquitous sensitivity in the perceptual Bayesian model [49] may be associated with hyper-arousal conditions with cognitive hypersensitivity in individuals with ASD. Fluctuation of the arousal conditions, which is induced by environmental changes, may also be associated with episodic features in ASD. Both the excessive preoccupation with unusual fantasy worlds in children with ASD and the “withdrawal to fantasy life” in schizophrenia [50] can be referred to as reality...
distortion. The reality distortion might clinically manifest itself or be latent according to the arousal level in perceptual thinkers [20,51]. In adolescents with ASD, not only negative symptoms but also positive schizotypal symptoms including catatonia correlate to ASD symptoms [52-54].

4. DISCUSSION

The reconsideration of the categorical identity of schizophrenia was launched in the discussion on the Kraepelinian dichotomy [55]. This traditional dichotomous view of the relationship between schizophrenia and bipolar disorder was challenged by family studies, twin studies, and recent molecular genetic findings [55]. The proposed spectrum from schizophrenia at one end to bipolar disorder at the other was historically suggested by Kraepelin by himself [56], and consequently involved ASD at the position adjacent to schizophrenia in the spectrum [57]. As described above, the developmental pathological model for schizophrenia was originally established on the basis of the speculated relationships between schizophrenia and some epiphenomena including nonspecific abnormal brain pathologies [23,24]. However, both the specific pathological marker and the molecular cause of schizophrenia remain, strictly speaking, unknown [25].

In the developmental hypotheses of schizophrenia, the clinical outcomes were supposed to be mediated by a range of mechanisms including genetic mediation, environmental influences, and individual cognitive experiences [55,57-59], and the childhood origins of schizophrenia were underscored [58,60]. It has been repeatedly claimed that ASD and schizophrenia are clinically distinct but have a clear degree of overlap that may reflect shared etiological mechanisms [2,61]. The distinction between ASD and schizophrenia is speculated to be explained by condition-specific genetic factors, molecular networks, brain functions, or neuroanatomical differences [61-63].

To reconsider the genetic and developmental relationships between ASD, schizophrenia, and bipolar disorder, four alternative hypotheses were recently introduced [64]. These are 1) ASD subsumed in schizophrenia, 2) independence, 3) diametric, and 4) partial overlap models. The diametric hypothesis is based on a proposal in which ASD and schizophrenia are distinct and at opposite ends of the same “social brain” spectrum [64]. Importantly, the reverse of the subsumed model is missing from the list. We independently proposed the missing model in which schizophrenia and bipolar disorder are subsumed in ASD [20]. In our hypothesis, we challenged the dogma that schizophrenia is a distinct clinical entity and schizophrenia is supposed to be mere one of the secondary derivative states in individuals with ASD [20]. The secondary derivative states may include schizophrenia, mood disorders, anxiety disorders, stress-related disorders, dissociative disorders, eating disorders, and abuse. The clinical outcomes may be determined by environmental influences and individual cognitive experiences. Because diagnosed ASD resides in the extreme of bell-shaped behavioral dimension that distribute quantitatively [26], the proposed developmental backgrounds for psychotic conditions include both diagnosed and undiagnosed ASD in the extreme. Behavioral or cognitive components of ASD characteristics in individuals with a normal developmental trajectory may also provide the underpinning for the development of psychotic conditions. In DSM criteria, the only distinctive symptom for the additional diagnosis of schizophrenia in individuals with ASD is apparent emergence of reality distortion (delusions or hallucinations). However, misattribution of stressful experiences (one of the inclinations of autistic adolescents) and the perceptual thinking style (one of the autistic characteristics) can constitute the underpinning of the reality distortion [20]. The possibility that schizophrenia is on the autism spectrum was also introduced by King and Lord [21], and Bevan Jones and colleagues explained the association between autistic traits and psychotic experiences by the possibility that ASD is an early precursor of psychotic experiences [22].

5. CONCLUSION

This concise review provides recent results of studies in relation to behavioral and cognitive core domains shared between ASD and schizophrenia. Together with the emerging new genetic concepts concerning human behavioral complex traits, the results and constructions may further compel scientists to reconsider the relationship between ASD and psychotic conditions. In addition, the possibility that psychotic conditions are the secondary derivative states in individuals with ASD or in individuals with subclinical ASD components should be constructively examined.

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Is autism spectrum disorder common in schizophrenia?


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