Neurodevelopment, Intestinal Function, and Autism

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Abstract

In brain development, functional and cognitive development matures after the sensory and sensorimotor regions are developed. Stimulation with information processing, whether from external environment or from visceral sensations, is important for developing functional networks and refining synaptic plasticity. The autistic brain is a neurodevelopmental disorder in which time windows shaped by the dynamic enteric microbial assembly and cortical neuroconnectivity may deviate developmental trajectories for brain function and behaviors. Association between gut microbiota and cognition in human infants has been recognized. Memory related to hippocampus neurogenesis in the acquisition of episodic memory as well as spatial working memory and path integration abilities can be related to vagal input from the gut as well as the enteric microenvironment. Alterations in microbiota could also be associated with postnatal neuro-inflammation with brain-gut dysregulation and the aberrant developmental processes could lead to diminished social motivation and increased activation to stimuli associated with their restricted interest. The resultant aberrant development trajectory may account for the memory representations which, though processed and activated by the hippocampus during successful retrieval, are not searched for, transferred, or monitored efficiently in autism spectrum disorders (ASD). These could manifest with their characteristics of stereotypical behavior, communication, and social interaction deficits, and related mixed picture of neural connectivity.

Effective treatment to restore normal school work and social interactions has been found possible by early management of the enteric system, relieving inflammation and improving sleep. Management of the enteric system could normalize the microbial elements influencing neuro-inflammation, and also sustaining signals to the hippocampus and brain for neurogenesis. Together with restoring the internal environment and management of aberrant processes, these have a high potential for ASD returning to a normal developmental trajectory for useful activities and life.

Keywords: Autism spectrum disorders; The enteric system; Enteric microbial assembly; Hippocampus memory; Aberrant development trajectory

Sensory Development before Functional and Cognitive Development

Studies of brain development noted sensory and motor functions maturing earlier as compared to higher-order integrative functions maturing later [1-4]. This can be correlated with respective cortical gray matter maturation in young children compared to older children and adolescents [5].

Rapid cortical gray matter growth in overall volume with regional differences is seen in the first 2 years of life. The sensory and sensorimotor regions are the first to grow and mature starting before birth. These enable motor, somatosensory, and auditory systems to function prior to birth, and vision for the newborn [6-8].

Brain regions develop around information processing sensory modalities. Regions expanding faster in the first year include the inferior frontal gyrus and angular gyrus, cortical regions involved with language [9], the fusiform gyrus, involved with face recognition and color processing [10,11] and the inferior temporal gyrus, involved with higher-order visual processing, including shape and faces [12,13]. The insula is also one of the most rapidly growing regions.

These are followed by frontal and parietal cortex expansion in the second year of life, probably associated with development of functional networks, since the default network which involves parietal and prefrontal cortex does not fully develop until 2 years of age [14]. Functional network hubs are noted earlier in sensorimotor regions [15,16] and in motor, sensory, auditory, and visual primary cortex in infants [17].

Maturational through Stimulation

The maturational trajectories are subjected to a variety of genetic and environmental factors. While heredity has a part to play, activity and experience would shape formation and elimination of synapses in the developing brain [18]. This probably accounts for the heterogeneity of developmental trajectories of subcortical structures in older children [19,20]. Nevertheless, the relative contribution of genetic and environmental factors to human gray matter development in this period of rapid growth and development is not clear.

Memory plays a significant part. There is a rapid increase in hippocampal volume in the first two years of life [5] while it continues to grow to 14 years of age [21,22]. It is less in the first year compared with the other subcortical structures, but becomes one of the faster growing structures in the second year of life. It supports, with more mobility of the child, the acquisition of episodic memory [23] as well as spatial working memory and path integration abilities [24,25].
Gut Influence on Brain Development

Visceral sensations develop as the enteric nervous system (ENS), some called the 'little brain' [26], develops during interactions of the neural crest-derived precursors (mostly vagal neural crest cells [27]) with the enteric microenvironment. Luminal stimuli activate mucosal enteroendocrine cells, which secrete signaling molecules, such as serotonin, that stimulate intrinsic primary afferent neurons (IPANS) to initiate peristaltic and secretory reflexes [28]. The insula, the most rapidly expanding regions in the first year, is involved with a variety of functions: awareness of interoceptive or visceral sensations, pain, body movement, emotions, vocalizations, and perhaps even consciousness [29-31].

Assembly of the developing nervous system for a final functional neural circuitry is dependent of a series of temporally regulated developmental processes. Development is not just following senses. Studies recently have provided much evidence that, besides the mainframe brain, microorganisms in the gut play a role in neurodevelopment [32]. Germ free (GF) mice displayed an exaggerated stress response, which could be restored by oral replacement of a single strain of bacterium, Bifidobacterium infantis [33].

Replacement for GF mice, using the microbiome from specific pathogen free mice raised with a normal gut flora, restored the stress response as well as reduced anxiety-like behavior and increased locomotor activity of GF mice to basal levels when given at an early developmental stage but not at a later age [33]. Serotonin levels in the hippocampus of GF mice were raised and correlated with reduced anxiety, but re-colonization of the mice after weaning did not restore the neurochemical differences (though it could revert the altered anxiety-like behavior) [34]. Time windows of microbial community might be critical in shaping the brain function and have long-lasting effects on behaviors. It needs be further revealed about the action of the gut microbiome on specific neuronal populations which could affect neurodevelopment of brain circuits. But perturbations in the delicate synergetic host-microbiota relationship may have serious consequences and lead to brain, digestive, and metabolic disorders [35-37].

Normal gut microbiota plays a critical role in shaping brain functions [38-41]. Microbe colonization after birth and gut microbiota assembly in the first 3 years of life, can shape development of the gut [42,43], endocrine system [36], and brain [38,44,45]. Gut microbiota and cognition in human infants are associated, as fecal microbial community diversity in infants affects later Mullen score (scale of early learning), visual reception scale, and expressive language scale at two years of age [46].

Developmental Outcome

Neurodevelopmental events including neurogenesis, axonal and dendritic growth, synaptogenesis, and refinement of these synaptic connections need be orchestrated to shape the functional neural circuitry matching needs that are critical for normal cognitive, motor, and emotional development. While plasticity allows a wide spectrum of outcome derivatives, systems matching [47,48] requires that the generated numbers of neurons and the appropriate synaptic density be matched exactly to the requirements of the pertinent neural circuit.

In the mainframe brain, development begins with an intricate process of neural cell generation, differentiation and migration, which is regulated intrinsically by expression of transcription factors as well as extrinsically by extracellular signals or morphogens [49,50]. This is followed by neuronal axonal growth as being guided by various attractive and repulsive cues and regulated by local mRNA translation [51,52]. Synapses are formed when the target synaptic partners are linked by the axons and then finally pruned to organize the final neural circuits [49,53]. The functions of these connections are further refined by synaptic plasticity.

Developmental outcome is certainly multifactorial. The brain is a plastic organ where both the intrinsic CNS milieu and extrinsic cues play important roles in shaping and wiring neural connections. Early life experiences can have profound and persistent effects on behaviors and traits with consequences for later life behavior and disease risk [54-56]. From psychosocial adjustment and self-regulation and from wiring of neural networks, the balance of plasticity and stability, being critical for information processing and storage, precedes the associative learning.

Early life stress during this critical period can induces alterations in many body systems. As simple as maternal separation can produce alterations of the intestinal barrier function, altered balance in enteric microflora, exaggerated stress response and visceral hypersensitivity [57]. There is bidirectional communication pathway between gut bacteria and the central nervous system [45]. The microbiota–gut–brain axis exerts a profound influence on key brain processes, such as neuroinflammation, activation of the stress axes, neurotransmission, and neurogenesis, in addition to modulating complex behaviors, such as sociability and anxiety [35-37,58-61].

The gut microbial community is dynamic during the first 3 years of life before stabilizing to an adult-like state [62]. Microbiome composition and specificity differ across host species and lifestyles [63]. The complex and diverse alterations in gene expression in multiple neurotransmission pathways, including glutamatergic, GABAergic, serotonin and dopamine pathways as well as neurotransmitter transporters and ion channels can have diverse impacts on the brain [64]. Gut bacteria influence these central processes through their ability to synthesize neurotransmitters including gamma-aminobutyric acid (GABA), noradrenaline, and dopamine, modulate activation of the immune system, and produce metabolites, such as short-chain fatty acids (SCFAs), that possess neuroactive properties [65]. Moreover, additional pathways link the gut microbiota and the brain, through the vagus nerve and through the modulation of key dietary amino acids such as tryptophan [66-69].

Gastrointestinal Improvement Precede Recovery in Treated Autistic children

A series of autistic children (age 2.5-6.5 yrs) demonstrated the significance of gastrointestinal (GI) function. From 2008-13, 8 autistic children diagnosed by psychometric assessment by psychologists came to InteMed Specialist Centre. 6 were treated with a host of medicine and herbs for 1.5-2 years while 2 did not receive treatment. They were evaluated once every two weeks for GI health in terms of bowel motion, appetite, smell stools and GI discomfort, sleep quality in terms of entry and restlessness, temper, concentration, and school acceptance and results. All other educational and social support were carried on unchanged as given outside the clinic. 5 patients improved greatly in final outcome, temper got significantly improved (5/6), and scoring high in schools (2/6). During recovery, appetite, speed of finishing feeds, sleep restlessness, concentration, temper, and school...
acceptance improved mostly in that sequence (Figure 1). 5 had treatment stopped within 1.5-2 years, while one with Asperger syndrome continued with stubbornness recovered after 2.5 years though negative and grudging thoughts changed positive only after 3 years. One child mainly stubborn was not compliant, feeding improved but worsened with frequent running nose and cough, and overall recovery variable with pattern not distinct. Follow-up in 2018 could not reach this child, being in foreign lands. For the remainder 5 patients, follow-up showed, for school: results good or upper grade in 3 and middle in 2, none need for special classes; for emotion: generally acceptable with peers in 5/5 and temper good in 1, easy to cry in 2, wrangling over issues in 2, none stubborn, all taken off special classes, nervous for work in 2, and none having mannerism; thus on the whole exceptionally effective.

The batch of medicine included chlorpheniramine 2mg and senna 3 mg daily. This is special low dose for longer-term use, even though they may not be obviously constipated, had no dependency problem displayed in all patients even afterwards. During treatment, it was particularly impressive that GI function in terms of appetite, speed of finishing feeds and smelly stools recovered within 6-24 weeks and tended to precede the whole sequence of recovery. Though cases were too few to be confident, there is some suggestion that the earlier GI function recovered, the earlier the rest of other symptoms recovered.

Gut motility in general depends on the gut luminal environment (including the gastrointestinal microbiota and fermentation), as well as factors related to the immune system, the ENS, and the central nervous system. Transit time is a key determinant and stool consistency strongly associated with gut microbiota richness and composition [70]. The increase of transit time is positively correlated with the increase of methanogens, of breath methane concentrations and pH, while inversely correlated to the proportion of sulfate reducing bacteria and SCFAs, generated by enteric bacterial fermentation, may induce neurodegenerative diseases [71]. Besides, gut microbial cell wall components continually interact with the innate immune system to induce the secretion of cytokines. Neuroinflammation-related brain injuries are associated with [73] and cytokine imbalance is involved in autism spectrum disorders and schizophrenia [74].

There can be a lot more processes about the brain-gut. The ENS is a division of the autonomic system put in close apposition to effector systems that it controls; enterohormones also co-working. Brain development depends on nutrition and immune development from the gut. Long-chain polysaturated fatty acids, monounsaturated fatty acids, insulin-like growth factor 1 are among a long list of constituents and regulators of brain cell proliferation, apoptosis, myelination, neurogenesis, maturation and differentiation [75].

Aberrant Development Trajectory in Autistic Children

Autism spectrum disorders (ASDs) are neurodevelopmental disorders characterized by the presence of stereotypical behavior, communication, and social interaction deficiencies. ASD etiology remains unknown and pathogenesis is sought for genetic and environmental factors [76]. Cognitive models include domain-specific models elaborating primary deficit in social cognition, and domain general models elaborating primary deficit in nonsocial or domain-general processing. The disrupted cerebral connectivity hypothesis postulates that its clinical symptoms originate from deficiencies in the way the brain coordinates and synchronizes activity amongst different brain regions [77]. Physiological and behavioral indicators can be found in high-risk groups less than a year old that predict the disease [78,79]. In a recent paper, atypicalities of structural and functional development of the brain related to prenatal and perinatal causes (hypoplasia of the pons just after neural tube closure; and a deficient GABA developmental switch in the perinatal period) have been postulated to explain the diverse phenotypes of ASD [80]. Nevertheless, these specific prenatal and perinatal causes are expected to be infrequent, yet ASD is common, affecting approximately 2.25% of children [81,82].

Considering aberrant developmental processes, the Social Motivation Hypothesis views that an early neurobiological difference in response to rewarding social input could in turn lead to diminished social motivation, further increasing motivation for restricted interests [83]. Without appropriate input signals to develop the neural organization over a long period, decreased gray matter volume in the region, and synaptic strength is modified in activity- and experience-dependent ways, causing change in connectivity. Thus an array with high variability in patterns of widespread local overconnectivity, a mixed picture of local over- and underconnectivity as well as mixed patterns of long range under- and overconnectivity have been observed in ASD [77]. Meta-analysis of 13 functional magnetic resonance imaging studies found aberrant reward circuitry activation to both social and nonsocial rewards and increased activation to stimuli associated with their restricted interest [84]. Whether due to a circuit dysfunction [85] in reward and motivation, autistic brains might fail to register social experiences as rewarding, further reducing social interactions and social abilities, ultimately leading to heterogeneous social deficits with this "aberrant development trajectory".

Dopamine dysfunctions have been reported in ASD, and autist-like behavior could arise from dopamine dysfunctions in midbrain dopaminergic modulatory systems affecting social motivation and goal-directed motor behavior [86]. Dopamine affects plasticity, synaptic transmission and the network activity in the hippocampal circuitry for memory [87]. Findings suggest that while memory
representations are processed and activated by the hippocampus in both ASD and controls during successful retrieval, these are not searched for, transferred, or monitored in an efficient way during episodic memory retrieval as a result of widespread disrupted connectivity. In brief, memory deficits in the ASD could be driven by retrieval-related impairments that reduce the probability of recollection success [88]. Aberrant connectivity may lead to structural demonstrable differences in many brain areas, especially for developmental process involved in response to rewarding social input, which in turn may lead to the diminished social motivation.

Molding with Gut-related Brain Development

Neurogenesis with new neurons continues in the hippocampus to play an important role in learning and memory and responses to stress, even till adulthood [89,90]. Germ-free mice with absence of a gut microbiota have impaired abilities in spatial and object, but not olfactory memory related to less c-Fos-positive CA1 hippocampal cells [91].

ASD is associated with other GI abnormalities, particularly alterations in microbiota composition and function [39,92-96]. Studies demonstrated that autism-like behavioral and GI phenotypes are associated with altered microbiota in two separate mouse models of ASDs [39,97]. Gut microbiota dysbiosis [96] and altered fecal flora have been observed [97,98]. Inflammation and neuro-immune system dysregulation can be a prominent clinical features of ASD [99-101]. Pro-inflammatory cytokines were found elevated [75] and dysfunctional immune responses could affect core behaviors in ASD [102]. Tumor necrosis factor alpha (TNF-α) were positively correlated with the ASD severity [103]. ASD blood monocytes showed altered immune responses [104] while blood monocyte-derived macrophages show strong impairments in the endocannabinoid system [105]. These are endogenous agonists of cannabinoid receptors, involved in the suppression of synaptic transmission and mediate signaling in the brain and the ENS. Besides monocytes are precursors of macrophages and dendritic cells.

Neuro-inflammation can affect developmental trajectories. Inflammatory cells in the brain such as microglia and astrocytes regulate synaptic structure and function. Neuroglia responses are activated toward pro-inflammatory processes involving astroglia and microglia, and demonstratable by brain immunohistochemical analysis in ASD brain [106]. Microglia cells like the monocytes are more committed to molecular pro-inflammatory changes. Astrocytes are also involved in ASD development possibly related to the G-protein coupled receptors [107]. Activation of the immune system probably dynamically affects synaptic organization and function in the developing brain and microglia-mediated elimination of synapses [108].

Possibilities for Treatment

Our series open up a new treatment perspective for ASD through GI management and concurrent repatterning brain development. Synaptic remodeling and developmental reorganization could be important in development of cognitive abilities and the necessary behavioral transition when growing up [109]. The experience of our cases including others outside the series suggests that treatment before 6 years of age would have a high chance of achieving normal school and social outcomes, while late treatment only restore emotion stability and strengthen communication and learning ability. The effectiveness of therapy probably are made by normalization of GI function and concurrent actions to remedy the 'aberrant developmental trajectory'.

There is a critical window in early life during which microbial colonization influences adult neurogenesis, including that in the hippocampus. Microbe colonization beginning around birth, produces a dynamic assembly during the first 3 years of life [62], and then gradually changes in composition onwards [110]. Increased intestinal permeability (leaky gut) has been associated with ASD, and is found in 37% of autistic patients and in 21% of their relatives [111]. With GI dysfunction present even in relatives and being subjectable to significant lowering with gluten-casein-free diet [112], a treatable postnatal element is present and may contribute to its associated neuro-inflammation in ASD. Gliogenesis essentially developed perinatally and postnatally [113,114]. Glial cells include astrocytes modulating the chemical environment by altering ion gradients and neurotransmitter transduction, oligodendrocytes producing myelin, and microglia to remove cellular and foreign debris within the central nervous system. Mal-assembly of gut microbiota during early childhood could enhance the individual's susceptibility to environmental insults and poor diet whence GI dysfunction is resulted with a negative impact on mental health. Although the pathogenesis of ASD remains largely unknown [115], neuro-inflammation contributes to a significant subset of ASD [116]. The use of herbs including Potentilla chinensis Ser. with inflammatory effects in our series may have contributed. Other herbs used in the early stage of treatment including Poria cocos (Sow.)Wolf, Trichitum astvium L., Ziziphus zizyphus, Taxillus chinensis, Lili bulbus, Fructus oryzae and Morus alba L. were also used to strengthen the GI system and the body. At a later stage for older children, Citrus aurantium L., Citrus reticulata Blanco, Bambusa tuludoides Munro and Pinella ternata (Thunb.) Breit. were used to improve temperament. The use of probiotics has been proposed as a therapeutic intervention for ASD [117,118]. For ASD children with immune dysfunction, intravenous immunoglobulin infusion was demonstrated useful in behavioral issues, eye contact, and social interactions [119] and even with standardized cognitive and behavioral tests [120]. Along the same line, manipulation of the endocannabinoid system for the gut-brain axis, which can block and reduce the development of colitis [121], has yet to be explored in autism.

With the ongoing neuoinflammatory insults alleviated, concurrent restoration of the mental abilities from the deterring 'aberrant developmental processes' that block and lead to a diminished social motivation, would restore the necessary early rewarding social input. The hippocampus is activated by GI signals through the vagus nerve between the GI tract and the brain [122]. The hippocampus is linked with learning and memory control and with feeding behavior [123]. Vagus nerve stimulation (albeit non-physiological electrical stimulation) enhances memory [124,125], facilitates hippocampal neurogenesis, and increases hippocampal expression of brain-derived neurotrophic factor [126] and induce neuronal plasticity [127]. Certain herbs may restore brain synaptic remodeling and reorganization.

Normalization for ASD then depends on actions to re-direct and pattern the aberrant developmental trajectory. With developmental plasticity, early life experiences can have profound and persistent effects on traits expressed throughout the life course, with consequences for later life behavior. The effectiveness of treatment in the above cases could be also be related to sleep improvement, as enabled with antihistamine and restored by sleep-promoting herbs including spine date seed (Ziziahi Spinosae Semoen). ASD tends to be
associated with difficulty in falling asleep, wake up in the night frequently and a low frequency of saccadic eye movement during rapid eye movement (REM) sleep [128]. Slow wave sleep is also shortened in ASD, and sleeping time, particularly the proportion of REM sleep, is reduced [129]. As cerebral plasticity has a very important relationship with sleep, our series showed improvement of sleep precedes improvement of other symptoms.

There are other postnatal remedies demonstrated as effective. Oxytocin as a nasal spray can modulate human social behavior and improvement [130]. Postnatally, a lot can be done as shown by our series. Postnatal synaptic plasticity during the synaptic pruning begins at birth. Late steps of neurodevelopment including axon and dendrite growth and arborization, and experience-dependent synapse modification, are primarily involved in the development of neurocircuitry, and open to management to normalize developmental trajectory.

Conclusion

ASD as a neurodevelopmental disorder may have a lot of remedial postnatal elements, including the deleterious GI function and gut microbiota assembly, as well as aberrant developmental processes that deter useful interactions and lead to the diminished social motivation. Management of the GI system could target not only the microbial elements influencing neuro-inflammation and glial cells, but also the ENS sustaining signals to the hippocampus and brain for neurogenesis. Restoring the internal environment, management of the aberrant developmental processes may be facilitated by better sleep and stayed by behavioral modification or by herbs. The concurrent treatment of postnatal process has a high potential for ASD returning to normal trajectory for useful activities and life.

References


