Autism and propionic acid

P Goof*

**Editorial**

The author has referenced some of his own studies in this editorial. These referenced studies have been conducted in accordance with the Declaration of Helsinki (1964) and the protocols of these studies have been approved by the relevant ethics committees related to the institution in which they were performed. All human subjects, in these referenced studies, gave informed consent to participate in these studies.

Derrick MacFabe’s hypothesis that propionic acid generated by gut bacteria induces autism, while compelling, leaves one question largely unanswered: Why do children with inborn propionic acidemia rarely show autistic behaviour? Although exacerbations of propionic acidemia (PA) bear “some resemblance” to autism spectrum disorders (ASD), MacFabe noted, only one case of autism associated with PA has been reported in the literature; these authors stated: “In the consensus conference about diagnosis and management of PA hosted in Washington, D.C. in January 2011, there was no reported association among the neurological sequelae of the disease between PA and autism.”

Other physicians who treat children with PA also report they rarely show autistic disorders: Sabine Scholl-Bürgi: “In our PA patient group none has an ASD.” Professor of Paediatrics: “I see lots of kids with PA and UCD [urea cycle disorders], but few (perhaps none) have ASD.” As the professor noted, children with inborn urea cycle disorders (UCD) also rarely show autistic behaviour.

Krivitzky and colleagues: “[C]hildren in this cohort [UCD] show other behavioural/emotional strengths, including a minimal percentage with previous diagnoses of Autism spectrum disorders, mood disorders, and other psychiatric disorders.”

Gropman and colleagues, however, concluded that patients with partial deficiencies of urea cycle enzymes and late-onset presentations may show signs of autism.

What do children with propionic acidemia or urea cycle disorders have in common that might protect them against ASD? Both are inborn errors of metabolism which induce high concentrations of blood ammonia, among other hazards. Children with urea cycle disorders lack one or more of the enzymes that convert ammonia to urea in the liver. Scholl-Bürgi and colleagues cited evidence implicating secondary inhibition of these enzymes in PA. Filipowicz and colleagues, however, concluded: “The results presented suggest that defective formation of glutamate/glutamine, rather than a block in the urea cycle, is the likely mechanism associated with hyperammonemia in patients with propionic acidemia.”

In tissues that produce large amounts of highly toxic ammonia (e.g. skeletal muscles) the enzyme glutamine synthetase converts ammonia to the nontoxic amino acid glutamine for safe transport in plasma to the intestines and conversion to urea by the liver.

Ammonia generated by brain neurons is first detoxified by α-ketoglutarate, forming the excitatory transmitter glutamate; astrocyte glutamine synthetase then combines glutamate with ammonia from neurons and from blood to form glutamine, which astrocytes release back to neurons to reform glutamate (and GABA).

When plasma ammonia is high in urea cycle disorders or hepatic encephalopathy, plasma glutamine is also high. When plasma ammonia is high in PA, however, plasma glutamine is normal or low (glutamine paradox).

Al-Hasnan and colleagues: “The mechanism that disrupts this correlation in propionic acidemia is unclear. Metabolic acidosis, which is not generally seen in UCD, may be a factor. It has been demonstrated that acute changes in pH have an effect on glutamine/glutamate metabolism. Acidosis enhances glutaminase in the kidneys, whereas in the liver it inhibits both glutaminase and glutamate dehydrogenase while stimulating glutamine synthesis.”

Tuchman and Yudkoff: “... in patients with severe PA, plasma glutamine levels correlate poorly with ammonia levels. The cause for this phenomenon is unknown, but may be that glutamine synthetase is inhibited by a putative toxin (e.g., propionyl-CoA), which accumulates in PA, or a low concentration of ATP, which is expected in PA, may affect this ATP-dependent enzyme.”

Plasma glutamine is also low in children and adults with ASD citations in 26 and brain glutamate/glutamine (glx) also usually low despite their frequent high plasma ammonia.

Ghanizadeh: “The low level of plasma glutamine... is suggested as a screening test for detecting autism in children especially those with normal IQ. The decreased level has been reported before in all children with autism.”

Wakefield and colleagues suspected bacteria in their diseased intestines generated more ammonia than their

*Corresponding author
Email: autismstudies1@gmail.com

1 Autism Studies, LA Pine, OR, USA

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impaired liver could clear, which reached the brain. Finding serum glutamine low in these children, and knowing liver dysfunction impairs synthesis of astrocyte glutamate transporters, they proposed their brain glutamine was also low.22

In children with propionic acidemia7,12,23 or urea cycle disorders11, by contrast, CSF/brain glutamine is often high. Scholl-Bürgi and colleagues: “In contrast to plasma, cerebral glutamine concentrations in PA are often elevated even in metabolic stable situations. Accumulation of intracellular glutamine measured as Glx peak has been shown by magnetic resonance spectroscopy. Elevated CSF concentrations have been reported in PA patients during a stroke-like episode or in hyperammonaemia.”23

Davison and colleagues, on the other hand, found that when PA was stable, brain glutamine was greater than normal in white matter, but much less than normal in the basal ganglia – subcortical gray matter structures deep within each hemisphere: “MRS studies undertaken during metabolic stability before any severe acute episodes beyond the neonatal period demonstrated decreased Glx in basal ganglia compared to the normal MRI comparator group but a trend to increase in white matter. Glutamine alone was significantly decreased in basal ganglia during metabolic stability.”

During exacerbations of PA, basal ganglia Glx fell further: “The alterations seen in glutamine and glutamate in basal ganglia are of particular note. Glx was significantly decreased during severe acute episodes, with a smaller (non-significant) decrease noted in basal ganglia in studies acquired during metabolic stability.” They concluded: “The metabolite alterations seen in propionic academia in the basal ganglia during acute encephalopathy reflect loss of viable neurons, and a switch to anaerobic respiration. The decrease in glutamine + glutamate supports the hypothesis that they are consumed to replenish a compromised Krebs cycle and that this is a marker of compromised aerobic respiration within brain tissue.”24

Does the fall in brain glutamine during exacerbations of PA explain why they bear “some resemblance” to ASD (MacFabe)?

Horder and colleagues studied brain metabolites in adults with ASD by MRS at 1.5 Tesla20. Their thoughtful report speaks for itself: “In summary, we found preliminary evidence that adults with ASD (both narrowly and broadly defined) have significant differences in brain glutamate and/or glutamine metabolism. This may be a final ‘common pathway’ in the disorder, and underpin some clinical symptoms.”

“Taken together, these results demonstrate that, rather than being a ‘global’ neurobiological abnormality, Glx changes seen in ASD are highly regionally specific, suggesting that the underlying neurobiological cause(s) are also localized. . . We are only able to report a correlation between ASD in adults . . . and reduced basal ganglia Glx levels. Hence, we cannot be certain whether the differences in Glx are the cause of the ASD symptoms. At 1.5 T, it is not possible to distinguish between the compounds that contribute to the ‘Glx’ signal, that is, glutamate and glutamine. Future studies at 3 T or higher are needed to distinguish these compounds, but previous studies have cautiously attributed reductions in Glx to glutamate, as glutamate constitutes the most abundant central neurotransmitter.”18

An MRS study of fever’s benefit in ASD might also speak for itself. The frequent ability of infectious fever to relieve autistic behaviour dramatically has long tantalized parents, practitioners, and researchers. Is the decisive factor in this phenomenon the great amounts of glutamine skeletal muscles release into blood as provisional fuel to compensate the loss of appetite (anorexia) of fever?25,26

The risky (but effective) antipsychotic drug risperidone (Risperdal), which calmed 54% of ASD children and adults (but aggravated 20%)27 also suggests brain glutamine may be low.

Risperidone may suppress serotonin and dopamine activity at synapses, but also stimulates glutamate uptake by astrocytes and activity of glutamine synthetase28.

Propionic acid interacting with ammonia may also play a role in ASD. Burrus: “A reaction between ammonia and propionic acid should result in the production of beta-alanine, a chemical similar in composition to gamma-aminobutyric acid (GABA), an inhibitory neurotransmitter which has been shown to be present in higher quantities in autistic patients.”29

In 2012 I sent my paper on fever, ammonia, and glutamine in ASD to all practitioners formerly listed on the Autism Research Institute (ARI) website. Ten practitioners replied they regularly give ASD children 250 mg to 8 g/day of oral glutamine to heal their intestines. One said 5 g/day was “fantastic” to heal intestines. Several warned of hyperactivity in some children; others said it was rare. Only two, however, reported improved behaviour from glutamine. One was Franco Verzella, an Italian MD who gives ASD children 5-7 g/day of oral glutamine, after first cleansing their intestines. He stated that oral glutamine has induced “multifactorial and multisystemic is the condition, so that the improvement has different aspects in different children. Most common: sedation, less stereotypes, better sleep, more concentration.” At the ARI ‘think tank’ in Baltimore April 2013, I mistakenly reported that Verzella gave ASD children 20-30 g/day of glutamine! He meant that amount was sometimes given to heal adult intestines.

A neurologist at the think tank warned that oral glutamine has induced
In the small intestine, oral glutamine nourishes enterocytes and other rapidly replicating cells, and also breaks down to glutamate and ammonia shunted to the liver, and citrulline, which the kidneys convert to arginine — required to detoxify ammonia to urea in the liver. Jon Pangborn, senior ARI biochemist, warned that intestinal bacteria and yeast can degrade oral glutamine and other amino acids to toxic metabolites. He recommends cleansing the gut before giving ASD children any amino acid except taurine, which helps detoxify ammonia.

MacFabe’s plausible hypothesis of ASD speaks for itself — but its implications are far-reaching. If propionic acid generated by gut bacteria induces autism, we might expect children with inborn propionic acidemia to show autistic behaviour. Yet they rarely do. In light of their usual high brain glutamine — and frequent low brain glutamine in ASD — does high brain glutamine protect them?

Postscript
A preliminary version of this communication was sent to all attendees of the 2013 ARI think tank. These replies appear most useful:

John Green (MD): “I have found that children with dihydroxyphenylpropionic acid elevations in the urine very often have substantial elevation in autistic behaviours, responsive to antibiotics such as vancomycin or flagyl. The problem with this observation is that antibiotics may help these kids for other reasons, so it certainly doesn’t close the logical loop on MacFabe’s hypothesis. I think glutamine bears close the logical loop on MacFabe’s hypothesis; and William Ellis of St. John’s Cathedral, Spokane, for encouragement and support for these studies.

References
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