New developments in the treatment of hyperammonemia: emerging use of carglumic acid

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Abstract: Hyperammonemia is a true neonatal emergency with high toxicity for the central nervous system and developmental delay. The causes of neonatal hyperammonemia are genetic defects of urea cycle enzymes, organic acidemias, lysinuric protein intolerance, hyperammonemia–hyperornithinemia–homocitrullinemia syndrome, transient hyperammonemia of the newborn, and congenital hyperinsulinism with hyperammonemia. In some of these conditions the high blood ammonia levels are due to the reduction of N-acetylglutamate, an essential cofactor necessary for the function of the urea cycle, or to the reduction of carbamoyl-phosphate synthase-I activity. In these cases, N-carbamylglutamate (carglumic acid) can be administered together with the conventional therapy. Carglumic acid is an analog of N-acetylglutamate that has a direct action on carbamoyl-phosphate synthase-I. Its effects are reactivation of the urea cycle and reduction of plasma ammonia levels. As a consequence it improves the traditional treatment, avoiding the need of hemodialysis and peritoneal dialysis. In this review we evaluate the possible field of application of carglumic acid and its effectiveness and safety.

Keywords: hyperammonemia, N-carbamylglutamate, carglumic acid, urea cycle disorder, metabolic disorders

Introduction
In humans catabolism of proteins and other nitrogen-containing molecules results in the production of nitrogen. Waste nitrogen is excreted largely as urea with smaller amounts as creatinine, ammonium ion, amino acids, and other nitrogenous compounds. In the brain, ammonia is an important product of the metabolism of the neurotransmitters glutamate and aspartate and of the neurotransmitter monoamines. It is also the product of the synthesis of glutamate from glutamine in nerve endings. At least some of the glutamate released from the nerve endings is neutralized via uptake by the astrocytes where it is combined with ammonia to form the electrophysiologically inactive molecule glutamine. Despite its central importance in metabolism, high concentrations of ammonia are generally toxic, particularly to the central nervous system, and the compound must be maintained at relatively low levels in the tissues.

Many inborn errors of metabolism cause increases in blood ammonia levels. Hyperammonemia is a condition that can cause neurological complications, coma, and even death. Indeed, high concentrations of ammonia may partially saturate the “enzymatic detoxifier” of the astrocytes, impeding the brain’s capacity for self-protection, which contributes to the neurological dysfunction. Therefore, high levels of ammonia are a real emergency and should be treated promptly. First-line treatment...
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does not contain any new information. It is a continuation of the previous text, discussing the treatment of hyperammonemia with N-carbamylglutamate (NCG). The review evaluates the field of application and the possibility for the treatment of hyperammonemia with NCG.

N-carbamylglutamate in urea cycle disorders

The urea cycle represents the principal mechanism of ammonia detoxification. Five enzymes are required for ureagenesis: CPS-I, ornithine transcarbamylase, argininosuccinate synthase, argininosuccinate lyase, and arginase. The urea cycle is also regulated by NAG, an essential cofactor necessary for the function of CPS-I. NAG is produced in the mitochondrial matrix from glutamate and acetyl coenzyme A by N-acetylglutamate synthase (NAGS). Deficiency in any one of these enzymes

Figure 1 Schematic representation of the urea cycle and related pathways.
Abbreviations: ASS, argininosuccinate synthase; ASL, argininosuccinate lyase; ARG, arginase; CPS-I, carbamoyl-phosphate synthase I; MMA, methylmalonic acidemia; NAGS, N-acetylglutamate synthase; OTC, ornithine transcarbamylase; PA, propionic acidemia.
causes a urea cycle disorder which results in ammonia accumulation.6

N-acetylglutamate synthase deficiency
A NAGS deficiency is a rare, autosomal recessive urea cycle disorder, and was described for the first time by Bachmann and colleagues in 1981.7 In this congenital error there is a wide range of severity of symptoms, as well as in the age of onset. Most frequently, the onset occurs in the early neonatal period with the signs and symptoms of hyperammonemia: poorly fed, vomiting, and neurological alteration such as hypotonia, abnormal movements of limbs, irritability, generalized seizures, afterward lethargy, and coma. Late forms may present with acute attacks of hyperammonemia in a previously healthy individual or with chronic neurological, gastrointestinal, and psychiatric clinical signs. The symptoms could be chronic nausea, vomiting, ataxia, confusion, and irritability, and in acute cases could lead to coma. The laboratory findings show an increase in blood levels of ammonia and markedly elevated concentrations of glutamine and alanine. In contrast to most of the other urea cycle disorders, plasma citrulline and urinary orotic acid are normal as well as urine organic acids and blood acylcarnitines. These clinical and biochemical features are present both in NAGS and CPS-I deficiencies. The diagnosis could be confirmed either with determination of NAGS and CPS-I activity in liver tissue in presence of arginine or by molecular diagnostic analysis.8 The majority of early forms have <5% NAGS residual activity and often have frameshift or nonsense mutations. Indeed, the late form presentation is associated with partial NAGS deficiency and hypomorphic alleles with single amino acid substitutions.8–11 The treatment of acute hyperammonemia consists of an intravenous administration of adequate calories, fluids, and electrolytes and in a restriction of protein intake but providing a mixture of essential amino acids. NCG is a specific treatment of hyperammonemia due to NAGS deficiency and should be administered as early possible when it is suspected.12 Also sodium benzoate (200–500 mg/kg/24 h), sodium phenylacetate (250–500 mg/kg/24 h), and arginine (200–600 mg/kg/24 h) should be considered. In case of failure of the above treatment, peritoneal dialysis must be initiated. In 1995, Guffon et al described a case of a newborn with NAGS deficiency treated by NCG; this treatment was initiated on the 25th day of life with a total dose of 80–100 mg/kg/day, with optimal results. A 1-year follow-up showed an infant with normal somatic and neurological development and a good metabolic balance.13 In the following years, 3 other cases, 1 neonatal form and 2 cases of partial NAGS deficiency, were described.14–16 NCG was administered (60–100 mg/kg/day in 4 single doses) and a decrease of blood ammonia was observed: this enabled an increase in protein intake. In 2004, Caldovic et al studied how the incorporation of an isotopic label from ammonium chloride into urea increased after the treatment with NAG. This study provided strong evidence that NCG therapy restores ureagenesis in NAGS deficiency. Later, in 2008, Tuchman et al presented a stable isotope 13C tracer method to measure ureagenesis and evaluate the effect of NCG in humans. The results of these investigations showed the reduction of plasma ammonia and glutamine levels and confirmed that NCG could be an effective treatment for inherited and secondary NAGS deficiency.18 Nordenström et al and Gessler et al reported other clinical cases of NAGS deficiency and concluded that the immediate treatment of hyperammonemia with NCG entails a favorable long-term outcome.

Currently NCG is approved in the European Union and the United States for the treatment of hyperammonemia due to NAGS deficiency. The initial daily dose should be 100 mg/kg, up to 250 mg/kg if necessary. It should then be adjusted individually in order to maintain normal ammonia plasma levels.

Carbamoyl-phosphate synthase I deficiency
The therapy with NAG in the kinetic variant or partial CPS-I deficiency has been discussed by O’Conner et al in 1985.2 In 1996 Kuchler et al administered NCG in an infant on suspicion of a NAGS deficiency. After the treatment, the levels of ammonia and glutamine decreased to the normal range. The results of liver biopsy showed a normal activity of NAGS and a decrement of activity of CPS-I. The authors concluded that NCG stimulates the remaining CPS-I and increases ureagenesis, for this reason NCG might be used in patients with partial CPS-I deficiency.

N-carbamylglutamate in organic acidemias
Neonatal hyperammonemia can be caused by some organic acidemias. Since the 1970s, hyperammonemia has been known to be a complication of propionic acidemia (PA) and methylmalonic acidemia (MMA), due to an inhibition of carbamoyl-phosphate synthesis, the first step of the urea
cycle. In fact, in such disorders, propionyl-CoA accumulates and inhibits NAGS. The reduced synthesis of NAG, the natural stimulator of CPS-I, hinders the conversion of ammonia to urea in the liver.21

Propionic acidemia
Propionic acid is an intermediate metabolite of branched-chain amino acids (valine, theonine, methionine), odd-chain fatty acids, and cholesterol degradation. Propionic acid is carboxylated to methylmalonic acid by a mitochondrial biotin-dependent enzyme: propionyl CoA carboxylase. It is composed of 2 subunits, α and β; the subunit α binds biotin. A failure of this enzyme determines the PA (MIM 606054). This inborn disease is inherited as an autosomal recessive trait. The genes involved are located in the following chromosomal regions 13q32 and 3q21-q22, and many mutations have been identified in different patients. Two forms are described based on the clinical manifestations: a severe form in which symptoms develop during the first days of life, and a milder form in which a patient may have an episode of severe ketoacidosis after a healthy period. The clinical findings are poor feeding, vomiting, dehydration, hypotonia, lethargy, severe ketoacidosis with rapid progress to coma, and death. Laboratory findings during the acute attack are a severe metabolic acidosis, hypoglycemia, and hyperammonemia. Propionic acid and methylcitric acid are markedly elevated in the blood and urine of infants with PA. To reach a definite diagnosis, a measurement of the enzyme activity in leukocytes or cultured fibroblasts is required. The treatment consists of hydration, correction of acidosis, with intravenous glucose, insulin, carnitine, sodium benzoate, and vitamin therapy, associated with a hypercaloric protein-free diet. Gebhardt et al described 2 patients with PA and hyperammonemia who were treated with NCG. Their blood ammonia levels were decreased even by the first dose of NCG and normalized in a 6-hour period. In both patients dialysis was avoided.22 Also Jones and collaborators published 2 cases of patients with PA and high ammonia levels. The first newborn responded to administration of NCG optimally with a fall in plasma ammonia levels. The second patient did not respond to NCG therapy, but in this case the dose of NCG was lower (25 mg/kg rather than 250 mg/kg).23 Other cases of PA treated with NCG have been described since these cases were described. In these studies the administration of NCG improved detoxification of ammonia.24-27 The effectiveness of NCG in decreasing ammonia levels in patients with PA was demonstrated by Tuchman et al. They investigated the in vivo synthesis of [13C]urea following administration of [1-13C]acetate, a stable isotope tracer, and NCG: in the patient with PA they observed an increase of 13CO2 in breath, of [13C]urea in plasma and production of absolute [13C]urea, although this was lower than in healthy subjects.24 Similar results were published by Ah Mew et al:28 they administered [13C]sodium acetate to 7 patients with PA and neonatal hyperammonemia, and measured the concentration of 13CO2 in blood and [13C]urea, before and after the start of NCG therapy. They demonstrated that NCG increases ureagenesis and decreases the blood ammonia and glutamine levels in PA patients.

Methylmalonic acidemia
Methylmalonic acid is normally derived from propionic acid as part of catabolism of branched-chain amino acids, cholesterol, and odd-chain fatty acids. The enzyme methylmalonyl CoA mutase converts the L-methylmalonic acid to succinic acid. This enzyme requires as a cofactor adenosylcobalamin, a metabolite of vitamin B12.29 Deficiency of enzyme mutase or, less frequently, of adenosylcobalamin determines the accumulation of methylmalonic acid in plasma. Two forms of mutase apoenzyme deficiencies are identified, the mutation mut0 responsible for complete enzyme deficiency and mut responsible for partial enzyme deficiency.30 These patients are not responsive to B12 vitamin therapy, unlike the patients with a defect in the formation of adenosylcobalamin. The clinical findings of the patients with MMA due to mut0, mut, and defect in metabolism of vitamin B12, are similar. Severe forms develop in the first days of life with feeding problems, vomiting, hypotonia, metabolic acidemia, and consensual tachypnea. Also this condition when not treated promptly may progress to coma and death. The milder forms evolve later with hypotonia, feeding failure, and developmental delay.31 Laboratory findings are hyperammonemia and metabolic acidosis with increase of methylmalonic acid in blood. In addition propionic acid and its metabolites are found in the urine. Diagnosis can be confirmed by measuring mutase activity in cultured fibroblasts, or by identifying the mutant gene. Intravenous treatment must be immediately started with carnitine, glucose, insulin, arginine, sodium benzoate, and hypercaloric protein-free diet. It has been demonstrated that the administration of NCG decreases blood ammonia levels. Gebhardt et al described a case of a 6-day-old newborn affected by MMA and hyperammonemia (805 µmol/L). After the initial conventional treatment and peritoneal dialysis his plasma ammonia levels were 610 µmol/L. NCG therapy was started and blood ammonia levels dropped to 100 µmol/L 3 hours after the first NCG dose administration.32 Also Levrat et al reported a case of MMA in which the hyperammonemia was resolved with insertion of NCG therapy.33
Maple syrup urine disease
Branched-chain amino acids (leucine, isoleucine, and valine) are metabolized by a mitochondrial thiamine pyrophosphate (vitamin B1)-dependent complex enzyme system: branched-chain α-ketoacid dehydrogenase. This enzyme is composed of 3 catalytic components, E1 (decarboxylase component, formed by 2 subunits E1α and E1β), E2 (acyltransferase component), and E3 (dehydrogenase component). Deficiency of this enzyme determines maple syrup urine disease (MSUD, MIM 251000), so called for the typical sweet odor of maple syrup found in body fluids, especially urine. Based on clinical manifestation and response to vitamin B1, administration, 5 types of MSUD have been classified. All these clinical phenotypes are inherited as an autosomal recessive trait. The gene for E1α, E1β, E2, and E3 subunits resides on chromosomes 19, 6, 1, and 7, respectively. There is no mutual relationship between clinical phenotypes and genotypes. The classic MSUD is the most severe form with major clinical findings. At birth the newborns are healthy; the clinical manifestations develop in the first week of life. These are poor feeding, vomiting, and in the following days lethargy and coma may appear. These patients are hypertonic, with muscular rigidity and severe opisthotonos. The neurological manifestations are convulsions, hypoglycemia, and cerebral edema may also be present. In acute state, the neuroimaging may show cerebral edema most frequently in cerebellum, dorsal brainstem, cerebral peduncle, and internal capsule. In later ages of life hypomyelination and cerebral atrophy may be observed. Laboratory findings show increased levels of leucine, isoleucine, and valine in plasma; these amino acids and their respective ketoacids are elevated in urine and in cerebrospinal fluid. The definitive diagnosis can be done by measuring the enzyme activity in leukocytes or cultured fibroblasts. The treatment of acute state is hydration with glucose infusion and a high caloric protein-free diet. In some patients peritoneal dialysis or hemodialysis are necessary. In 2009 for the first time, Kalkan Ucar et al used NCG in a patient with decompensated MSUD. These authors demonstrated that the administration NCG together with conventional therapy decreases the blood levels of ammonia and can be added to improve the acute metabolic compensation in patients with MSUD.

Isovaleric acidemia
Isovaleric acidemia (MIM 243500) is a rare autosomal recessive inborn disease caused by deficiency of isovaleryl coenzyme A dehydrogenase (IVD). IVD is a mitochondrial flavoenzyme which catalyzes the conversion of isovaleryl-CoA to 3-methylcrotonyl-CoA in the leucine degradation. Two forms of this disease exist: an acute one in which the symptoms develop in early hours of life with vomiting, severe metabolic acidosis, lethargy, convulsion, and coma, and a chronic intermittent form defined by episodes of vomiting, dehydration, metabolic acidosis, and alteration of mental status; these events develop in the first months or years of life. Characteristic “sweaty feet” may be present in acute attacks. If untreated, these patients may die or may develop severe central nervous system dysfunctions. Laboratory findings during the acute attacks show metabolic acidosis, neutropenia, and thrombocytopenia. Hypocalcemia, hyperglycemia, and moderate to severe hyperammonemia may be present in some patients. The diagnosis is confirmed by the increase of blood levels of the isovaleric acid and its metabolites and by the elevations of isovalerylglucine and 3-hydroxyisovaleric acid as well as other metabolites in urine. Mass screening in newborn is possible by tandem mass spectrometry (MS/MS) for the analysis of the acylcarnitine profile in blood spot. The gene of IVD has been mapped to chromosome 15q14-15q15. Also it is possible to measure the enzyme activity in leukocytes or cultured fibroblasts. Today many mutations of this gene have been identified. In acute attacks the treatment consists of hydration, correction of metabolic acidosis by infusing sodium bicarbonate, promotion of anabolism by increased caloric intake (infusing glucose), and reduced leucine intake. Treatment must be initiated promptly. The long-term treatment is a diet with lower protein intake (1.5 g/kg/day), and administration of L-carnitine (100 mg/kg/day) and glycine (150–250 mg/kg/day) to prevent the accumulation of isovaleric acid and its toxic metabolites. Coude et al demonstrated in rat hepatocytes that high levels of isovaleric acid inhibited ureagenesis due to a decrease in NAG levels, causing hyperammonemia. For this reason, in patients with isovaleric acidemia associated with hyperammonemia the administration of NCG may be added to conventional therapy. A clinical case report of IVA successfully treated with carbamylc acid was recently communicated by Kasapkara et al.

N-carbamylglutamate in hyperinsulinism hyperammonemia syndrome
The hyperinsulinism hyperammonemia syndrome (HI/HA) is an inborn error of metabolism. It was described for the first time by Zammarchi et al in 1996. HI/HA syndrome is caused by a mutation of the enzyme glutamate
dehydrogenase (GDH) encoded by GLUD1. These mutations result in higher activity of GDH. About 80% of cases are due to de novo mutations; however a dominant pattern has been demonstrated. GDH is a mitochondrial matrix enzyme, expressed at high levels in liver, kidney, brain, and pancreatic β-cells. It is composed of 6 subunits and oxidizes the glutamate to α-ketoglutarate and ammonia using NAD and/or NADP as a cofactor. The clinical manifestations of affected infants are variable episodes of hypoglycemia associated with hyperammonemia. Hypoglycemia may occur with fasting and in response to protein feeding. Patients with HI/HA syndrome show a large acute insulin response to leucine administration and the blood amino acid concentrations are normal. In these children blood ammonia levels are persistently elevated between 2 and 5 times over normal limit, but the hyperammonemia is not associated with lethargy, irritability, or coma. Stanley has supposed that overactivity of GDH in the brain decreases the levels of glutamate and glutamine, protecting the central nervous system from the neurotoxicity of the accumulation of these elements. In this metabolic disorder the physiological and pathological basis of hyperammonemia is that the gain in these elements. The reduction of hepatic NAG, the obligatory metabolite. The reduction of hepatic NAG, the obligatory activator of the first enzyme of the urea cycle, leads to the inhibition of hepatic mitochondrial CPS-I.

N-carbamylglutamate in valproate-induced hyperammonemia?

Valproate-induced hyperammonemic encephalopathy is an unusual, but serious, adverse effect of valproic acid treatment that can lead to death, and is characterized by a decreasing level of consciousness, focal neurological deficits, cognitive slowing, vomiting, drowsiness, and lethargy. The mechanism by which hyperammonemia develops is still unclear. Hyperammonemia may result from the stimulation of kidney glutaminase in the renal cortex, from the depletion of mitochondrial acetyl CoA, and decreased production of NAG, but also from the accumulation of propion, a valproate metabolite. The reduction of hepatic NAG, the obligatory activator of the first enzyme of the urea cycle, leads to the inhibition of hepatic mitochondrial CPS-I.

Two patients with valproate-induced hyperammonemia have been treated with carglumic acid. NCG was effective in decreasing blood ammonia levels and its administration may be added to conventional therapy, as suggested by recent guidelines.

Side effects

The most common adverse reactions (occurring in more than 13% of patients) observed in patients receiving NCG, regardless of causality, are vomiting, abdominal pain, pyrexia, tonsillitis, anemia, ear infection, diarrhea, nasopharyngitis, and headache. In newborns, NCG is usually well tolerated without significant adverse effects. Tachycardia, profuse sweating, increased bronchial secretion, hyperthermia, and restlessness have been reported only at a dosage of 650 mg/kg/day.

Conclusion

We have described the beneficial effects of NCG to decrease blood ammonia levels and to improve ureagenesis in different conditions presenting with hyperammonemia. NCG is safe, fast acting and easy to administer. Dosing varies depending on the clinical condition and indication. Restarting the urea cycle, NCG increases ureagenesis and diminishes blood ammonia levels. It also improves the acute therapy and thus reduces the need for or the duration of peritoneal dialysis and hemodialysis. Favorable long-term outcomes in promptly treated patients with hyperammonemia suggest the need to start the NCG therapy as soon as possible, even before the exact diagnosis is set up.

Abbreviations

ASL, arginosuccinate lyase; ASS, arginosuccinate synthase; CPS-I, carbamoyl-phosphate synthase I; HI/HA, hyperinsulinism hyperammonemia syndrome; MMA, methylmalonic acidemia; MSUD, maple syrup urine disease; NAG, N-acetylglutamate; NAGSI, N-acetylglutamate synthase; NCG, N-carbamylglutamate; OTC, ornithine transcarbamylase; PA, propionic acidemia.

Disclosure

The authors report no conflicts of interest in this work.

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