High Blood Pressure in a Urea Cycle Disorder: Case Report

Carolina Solé, MD1 María Arriaga, MD1 Elvira Cañedo, MD2

1 Department of Neonatology, Gregorio Marañón Hospital, Madrid, Spain
2 Department of Metabolism Diseases and Nutrition, Niño Jesús Hospital, Madrid, Spain

Address for correspondence Carolina Solé, MD, Neonatology Department, Gregorio Marañón Hospital, O’Donnell Street, 48, 28009 Madrid, Spain (e-mail: carolina.sole.delgado@gmail.com).


Abstract

Introduction Urea cycle disorders (UCDs) form a group of metabolic pathological conditions that might develop serious neurological consequences. Early diagnosis, before irreversible damage is established, is the most important prognostic and morbidity factor.

Case Report We present the case of a 5-day newborn with high blood pressure and respiratory distress. Diagnosis was type I citrullinemia. With appropriate citrullinemia guided-treatment blood pressure returned to normal.

Conclusion High blood pressure has been rarely described as a lead symptom for the debut of a UCD. We must take this into consideration as an early recognition and treatment of these disorders are of the utmost importance.

Case Report

Our patient is a 5-day male newborn, with no pathological clinical record and no parental consanguinity presenting in the emergency department for low volume urine output in the last 12 hours. Parents referred no other symptoms. On the physical examination it stands out respiratory distress and maintained high blood pressure (BP) in the four limbs (above 120/90 mm Hg). Heart rate, peripheral oxygen saturation (SatO₂), and temperature were within normal range. Full blood count, renal and liver function, C-reactive protein (CRP), venous blood gas, and urine sediment were examined and found normal except for a respiratory alkalosis. Also, urine and blood culture were taken, and empirical antibiotic treatment started. He was transferred to a tertiary care neonatal center to continue study and support.

On arrival to our center, he was ill with continuous respiratory moan and work of breathing. He also had an altered mental status of being lethargic. No other findings of dysmorphic features were observed. We corroborated the high BP both in arms and legs (respiratory rate of 50 rpm, and SatO₂ 98%).

Some additional tests were added, showing mild coagulopathy, an ammonia of 93 µmol/L, and slight left ventricular hypertrophy. Cranial and abdominal ultrasound were normal as the CRP for enterovirus, cytomegalovirus, herpes 1,2, zoster, and parechovirus.

After a few hours of admission, the metabolic screening center informs of an altered result for the patient (sample on dry paper at 48 hours of life) with elevated citrulline and normal arginosuccinate acid, suggesting urea cycle disorder (UCD), specifically type I citrullinemia as we can see in ►Table 1. At that time ammonia levels had risen to 243 µmol/L. Another plasma and urine sample were collected for amino acid analysis; (results in ►Table 2) then, treatment of type I citrullinemia-oriented case was initiated with sodium benzoate, sodium phenylbutyrate, L-arginine, and carnitine. Calory intake was maintained with glucose and lipids, with protein restriction. Despite the initiation of the treatment, ammonia quickly rose to a maximum of 875 µmol/L in the first 12 hours, requiring venous hemofiltration. After this, it was finally possible to control hyperammonemia.

The patient needed intensive care with mechanical ventilation, inotropic support, and hemofiltration for 3 days. Blood pressure normalized within 8 hours of the treatment.

From a neurological point of view: at admission, he was hypoactive with both limited range of movement and response.

Keywords ► hypertension ► citrullinemia ► arginine ► newborn

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to stimulus. Altered consciousness, rapidly evolved from stupor to drowsiness. Abnormal amplitude integrated electroencephalography (aEEG) with a burst suppression pattern. On the second day, he presented with clinical seizures of low amplitude clonisms of the head with acute increase work of breathing. It was necessary to provide phenobarbital and levetiracetam bolus to control seizures, maintaining levetiracetam as a prophylactic treatment. Significant clinical improvement was noticed on the third day of admission, with continuous pattern on the aEEG, normal awareness level, appropriate eye tracking, adequate peripheral and axial tone, present both osteotendinous and archaic reflexes, harmonious and symmetrical movement of the four limbs and normal strength. He had normal EEG. Brain magnetic resonance imaging showed altered signal on basal ganglia with hyperintensity in T1 of nucleus pallidus and hypointensity on white matter with punctiform hyperintensity lesions. Findings in line with UCD are shown in Figs. 1 to 5.

After control of HA, low protein intake (0.3 g/kg/d) was started with progressive increase of dose up to 1.6 g/kg/d. Well-tolerated and daily ammonia check was normal. Intravenous medication was changed to enteral L-arginine (350 mg/kg/d), carnitine (30 mg/kg/d), and glycerol phenylbutyrate. He was discharged home with multidisciplinary follow-up with pending results of the genetic study.

Discussion

The urea cycle is a metabolic pathway that transforms ammonia into urea. The result of the catabolism of amino acids is ammonia, a high toxic compound that cannot be excreted unless it is processed into a high solubility substance such as urea. Genetic defects in the enzymes involved in the cycle can occur, the result is a UCD. These can be:

- N-Acetyl glutamate synthase deficiency (NAGS; OMIM 237310).
- Carbamoyl phosphate synthetase deficiency (CPS-1; OMIM 237300).
- Ornithine transcarbamoylase deficiency (OTC; OMIM 311250).
- Citrullinemia type I (deficiency of argininosuccinate acid synthase [ASS]; OMIM 215700).
- Argininosuccinic aciduria (deficiency of argininosuccinate acid lyase [ASL]; OMIM 207900).
- Argininemia (deficiency of arginase [ARG]; OMIM 207800).

<table>
<thead>
<tr>
<th>Table 1</th>
<th>Results of metabolic screening 48 h of life</th>
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<tbody>
<tr>
<td>Amino acids</td>
<td>μmol/L</td>
</tr>
<tr>
<td>Alanine</td>
<td>327</td>
</tr>
<tr>
<td>Arginine</td>
<td>1760</td>
</tr>
<tr>
<td>Glutamine</td>
<td>306</td>
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<tr>
<td>Glutamic acid</td>
<td>409</td>
</tr>
<tr>
<td>Glicine</td>
<td>117</td>
</tr>
<tr>
<td>Leucine + isoleucine + hydroxyproline</td>
<td>46.4</td>
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<tr>
<td>Methionine</td>
<td>39.6</td>
</tr>
<tr>
<td>Ornithine</td>
<td>25.34–75.42</td>
</tr>
<tr>
<td>Phenylalanine</td>
<td>189</td>
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<tr>
<td>Proline</td>
<td>0.22</td>
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<tr>
<td>Tyrosine</td>
<td>115</td>
</tr>
<tr>
<td>Valine</td>
<td>107</td>
</tr>
<tr>
<td>Argininosuccinic acid</td>
<td>0.12</td>
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</tbody>
</table>

<table>
<thead>
<tr>
<th>Table 2</th>
<th>Amino acid profile at 5 d of age</th>
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<tr>
<td>Amino acids</td>
<td>μmol/L</td>
</tr>
<tr>
<td>Arginine</td>
<td>9</td>
</tr>
<tr>
<td>Citrulline</td>
<td>4,002</td>
</tr>
<tr>
<td>Glutamine</td>
<td>2,197</td>
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<tr>
<td>Lysine</td>
<td>438</td>
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<tr>
<td>Methionine</td>
<td>347</td>
</tr>
<tr>
<td>Ornithine</td>
<td>14</td>
</tr>
</tbody>
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| Fig. 1 | Axial T2-weighted image showing high intensity alteration on white matter with smaller low intensity signals. |
| Fig. 2 | T1-weighted low intensity signal on white matter and focal high intensity signal. |
We can include three deficiencies of membrane transporters:

- Hyperornithinemia, hyperammonemia, homocitrullinuria syndrome (deficiency of the mitochondrial ornithine transporter).
- Lysinuria with protein intolerance (due to altered transport of dibasic amino acids).
- Citrine deficiency or citrullinemia type 2 (aspartame-glutamate mitochondrial hepatic transporter deficit; Fig. 6).

In this case, given the increase in citrulline and the decrease in arginosuccinate, the diagnostic suspicion is an alteration in arginase synthetase and therefore a type I citrullinemia. Even so, regardless of the suspected diagnosis, the main acute manifestations of the UCD are those
Derived from hyperammonemia (HA). Ammonia is a highly toxic product, especially on the central nervous system. It is considered as a potentially lethal metabolic emergency. The clinical syndrome of the newborn (which is the form of debut in 60% of the cases) begins at 24 to 72 hours of life, producing decreased feeding, thermic dysregulation, vomiting, impaired level of consciousness (drowsiness that evolves into coma), and seizures. A common sign is central hyperventilation and respiratory alkalosis. This may evolve to respiratory and cardiac failure.

An ammonia level > 110 mmol/L in the neonatal period is considered pathological. In the case of suspicion, therapeutic measures should be initiated immediately. In our case, these measures were aimed at the diagnosis of suspected type I citrullinemia which does not respond to cofactors or cysteine acid (Carbaglu).

**Why High Blood Pressure?**
The patient debuted with arterial hypertension as a lead symptom. Hypertension screening was performed:

- Heart disease: there were no differences in BP or SatO2 between upper or lower limbs. Chest X-ray without cardiomegaly or acute lung edema was obtained. An emergency echocardiogram was performed showing only a concentric hypertrophy of the left ventricle.

- Acute renal failure, renal stenosis: Had normal renal function. Doppler ultrasound of the renal veins was performed without alterations.
- Cranial ultrasound was normal.

After the diagnosis of suspected type I citrullinemia, we reviewed the pathophysiology and previous literature. This disorder involves arginine deficiency. It cannot be synthesized and forms an essential part of amino acid, as can be seen in Fig. 7. L-arginine acts as a substrate for nitric oxide synthetase to form endothelial nitric oxide and citrulline.4

**Fig. 6** Urea cycle in the liver. Ammonia reacts with bicarbonate and with N-acetyl glutamate to produce carbamoyl phosphate (CPS). The CPS yields its carbamoyl group to ornithine to form citrulline. Citrulline is released to cytoplasm. The second amyl group from aspartate is condensed with citrulline to form arginosuccinate, which is hydrolyzed into free arginine and fumarate. The fumarate enters the Krebs cycle and arginine hydrolyzes to form urea (excreted in the urine) and ornithine, which re-enters the mitochondria to restart the cycle. Mitochondrial enzymes (upper part of the cycle): N-acetyl glutamate synthetase (NAGS) essential co-activator of CPS-1; carbamoyl phosphate synthetase (CPS-1); Ornithine transcarbamylase (OTC). Cytoplasmatic enzymes (or lower part of the cycle): arginosuccinate synthetase (ASS); arginosuccinate lyase (ASL); arginase (ARG). Ornithine transporter deficit (ORNT-a) produces HHH syndrome. HHH, hyperornithinemia, hyperammonemia, homocitrullinuria.

**Fig. 7** L-arginine acts as a substrate for nitric oxide synthetase (ONS) to form endothelial nitric oxide (NO) and citrulline.
A reduction in NO production plays a major role in vascular tone and hence increase in BP.\(^4,5\) As it can be seen in – Table 1, arginine levels at 48 hours of life were in the lower reference value. At that moment, our patient was asymptomatic. At 5 days of age, when he debuted with hypertension, arginine was below normal range. Once treatment was started, which included arginine supplementation, the patient’s BP normalized. The connection between hypertension and UCD had barely been described, with little representation in this specific literature. Recently, Teufel et al\(^6\) described a series of 17 patients in which 81% of them presented with BP >95th percentile at debut. They also, highlighted how this clinical parameter could be useful to discern a UCD debut from a neonatal sepsis. Although the study involves a small series of patients, it underlines and supports what we have seen in this particular case and the importance of having present UCD as a differential diagnosis in a newborn with high BP.

**Conclusion**

Hypertension and UCD maintain a clear pathophysiological relationship, barely described in the literature, which can prove to be very useful in daily clinical practice.

**Authors’ Contribution**

C.S.D. wrote the manuscript. M.A.R. and E.C. reviewed and corrected the manuscript. All authors approved the final version.

**Conflict of Interest**

C.S.D., M.A.R., and E.C. declare that they do not have a conflict of interest. Informed written consent was obtained from parents of the patient.

**References**