A Proposed Physiopathological Pathway to Hyperammonemic Encephalopathy in a Non-Cirrhotic Patient with Fibrolamellar Hepatocellular Carcinoma without Ornithine Transcarbamylase (OTC) Mutation

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Patient: Male, 31
Final Diagnosis: Fibrolamellar hepatocellular carcinoma
Symptoms: Encephalopathy
Medication: —
Clinical Procedure: —
Specialty: Gastroenterology and Hepatology

Objective: Rare disease
Background: Hyperammonemic encephalopathy is a potentially fatal condition that may progress to irreversible neuronal damage and is usually associated with liver failure or portosystemic shunting. However, other less common conditions can lead to hyperammonemia in adults, such as fibrolamellar hepatocellular carcinoma. Clinical awareness of hyperammonemic encephalopathy in patients with normal liver function is paramount to timely diagnosis, but understanding the underlying physiopathology is decisive to initiate adequate treatment for complete recovery.

Case Report: A 31-year-old male with fibrolamellar carcinoma and peritoneal carcinomatosis presented with rapid onset hyperammonemic encephalopathy. Despite usual treatment for hepatic encephalopathy, his hyperammonemia was aggravated. A physiopathological pathway to encephalopathy resulting from hepatocellular dysfunction or portosystemic shunting was suspected and proper treatment was initiated, which resulted in complete remission of encephalopathy. Thus, we propose there is a physiopathological path to hyperammonemic encephalopathy in non-cirrhotic patients with fibrolamellar carcinoma independent of ornithine transcarbamylase (OTC) mutation. An ornithine metabolism imbalance resulting from overexpression of Aurora Kinase A as a result of a single, recurrent heterozygous deletion on chromosome 19, common to all fibrolamellar carcinomas, can lead to a c-Myc and ornithine decarboxylase overexpression that results in ornithine transcarboxylase dysfunction with urea cycle disorder and subsequent hyperammonemia.

Conclusions: The identification of a physiopathological pathway allowed adequate medical treatment and full patient recovery from severe hyperammonemic encephalopathy.

MeSH Keywords: Ammonia • Brain Diseases, Metabolic • Liver Neoplasms • Ornithine Carbamoyltransferase Deficiency Disease

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Background

Protein digestion results in the production of amino acids, which are metabolized by the liver by oxidative deamination or transamination into ammonia, which are then converted to urea and excreted by the kidneys. Any disturbance of this cycle may lead to hyperammonemia and as a result, hyperammonemic encephalopathy (HE).

There are three well-established physiopathological pathways that lead to disruption of the nitrogen excretion cycle. An excessive nitrogen load, for instance in the case of gastrointestinal bleeding or urinary diversion, may oversaturate the hepatic metabolism capacity. Another pathway disruption is an inability of the urea cycle to metabolize a normal nitrogen load, as in hepatic failure or in specific enzymes deficiencies. Additionally, the nitrogen load coming from intestinal protein digestion may bypass the liver, which occurs with porto-systemic shunting [1].

The most common cause of hyperammonemic encephalopathy is liver failure. Hepatocellular dysfunction and porto-systemic shunting are the most important causes of inability of the urea cycle to properly excrete nitrogen metabolites resulting in hyperammonemia in patients with cirrhosis [2]. Nevertheless, there are many non-hepatic causes of HE related to inborn errors of metabolism, such as ornithine transcarbamylase (OTC) mutation.

We present the case of a non-cirrhotic patient with fibrolamellar hepatocellular carcinoma (FHC) who developed hyperammonemia and encephalopathy despite having normal hepatocellular function. A physiopathological path involving an overexpression of Aurora Kinase A (AURKA) due to a heterozygous deletion on chromosome 19 (which is present in every FHC) is proposed.

Case Report

A 31-year-old male with no comorbidities and no risk factor for chronic hepatic disease presented with a history of three weeks of abdominal circumference enlargement. A computed tomography (CT) disclosed a large 18 cm hepatic mass occupying the right liver, as well as ascites and signs of carcinomatosis (Figure 1). Serum bilirubin level was 0.9 mg/dL (reference interval: 0.2–1.0 mg/dL), INR was 1.34 (reference interval: 0.8–1.2). He had normal hepatic enzymes at presentation. There were no stigmata of chronic hepatic disease at physical examination, other than ascites, that could be explained by peritoneal carcinomatosis.

The patient underwent diagnostic laparoscopy and biopsy of the hepatic tumor, and peritoneal implants were obtained (Figure 2). A large hepatic mass was found on the right liver with signs of rupture. Ascites and carcinomatosis were also present. The left lobe was enlarged probably due to compensatory hypertrophy as a result of the right portal vein tumor thrombosis. The non-tumoral liver was normal, with no signs suggesting chronic hepatic disease.

The patient’s postoperative period was uneventful and the patient was dismissed from the hospital two days after the procedure. Pathology identified hepatic fibrolamellar hepatocellular carcinoma (FHC) with metastatic peritoneal seeding. Immunohistochemistry examination determined a profile consistent with FHC (Table 1).

One week after surgery, he was admitted to the hospital due to vague symptoms such as fatigue and mild abdominal pain. Moderate ascites had returned. He presented no signs of infection or dehydration. Laboratory tests disclosed normal liver function, with serum bilirubin 0.8 mg/dL (reference interval: 0.2–1.0 mg/dL), INR 1.16 (reference interval: 0.8–1.2), serum venous lactate 11 mg/dL (reference interval: 6.3–18.9 mg/dL). One day after admission, he presented with lethargy and confusion and was admitted to the ICU.

A head MRI disclosed no secondary malignancies, no signs of intracranial hypertension, and no other significant findings. Blood ammonia level was 204 mcg/mL (reference interval: 9–30 mcg/mL). Usual treatment for hepatic encephalopathy was initiated with neomycin, lactulose, and L-ornithine-L-aspartate. A continuous 18-hour electroencephalography disclosed lowered basic rhythm, frequent diffuse triphasic waves, and sporadic outbreaks of moderate amplitude theta waves, consistent with HE.

Despite initial therapy, blood ammonia level increased to 280 mcg/mL and the patient presented with neurological deterioration and coma. He required endotracheal intubation and mechanical ventilation. Hemodiafiltration and proper parenteral nutrition were initiated. Despite all procedures to treat HE and continuous hemodiafiltration, his serum ammonia level reached 312 mcg/mL.

At this time, as treatment was not achieving the therapeutic results expected, another physiopathological cause other than the usual excessive nitrogen load, hepatic dysfunction, and porto-systemic shunting to HE was suspected. In fact, the reticulocyte production index was 0.5% (reference interval: 0.5–2.5%), lactic dehydrogenase was 280 IU/L (reference interval: 105–333 IU/L) and the hematocrit was stable, suggesting there was no hemolysis secondary to porto-systemic shunting. Moreover, hepatic function laboratory tests were still normal.
between normal range and the compensatory hypertrophy of the left lobe as a result of tumor thrombosis of the right portal vein suggested healthy non-tumoral hepatic parenchyma.

This led to the investigation of less common causes of hyperammonemia, such as a urea cycle disorder. Since ornithine transcarbamylase (OTC) deficiency is the most common urea cycle enzymatic defect associated with late onset HE in adults without hepatic dysfunction, this diagnosis was suspected. As urinalysis disclosed very elevated orotic acid (10 mmol/mol of creatinine, reference interval: 0.4–1.2 mmol/mol of creatinine) and plasma amino acid chromatography revealed reduced citruline (3.0 mcmol/L, reference interval 16–51 mcmol/L), very low arginine (15.5 mcmol/L, reference interval 43–407 mcmol/L) and reduced ornithine (19.0 mcmol/L, reference interval 15.0–80.0 mcmol/L), confirming OTC deficiency, treatment with sodium benzoate (3 g) and arginine (3 g) administered every four hours via nasogastric tube was promptly initiated. A multi-gene panel genetic testing for inborn errors of metabolism was performed and no mutations were observed (Figure 3). OTC gene was tested and was not mutated.

Within 18 hours, blood ammonia level had reduced to 107 mcmol/L. As the conscious level improved, the endotracheal...
tube was removed. The patient was dismissed from the intensive care unit one day after, with no clinical signs of HE. After treatment was initiated, there were no more clinical episodes of encephalopathy and the patient’s neurological examinations were normal. Before hospital discharge, the patient had chemotherapeutic treatment with a multikinase inhibitor (sorafenib) and GEMOX (gemcitabine-oxaliplatin). After two months on chemotherapeutic treatment, an abdominal CT disclosed stable disease.

Three months after the introduction of sodium benzoate and arginine, and adequate diet, without any other medical treatment for HE (i.e., neomycin, lactulose, and L-ornithine-L-aspartate were discontinued), even with ongoing chemotherapy treatment, there were no clinical signs of encephalopathy and the patient’s ammonia blood levels ranged between 40 and 60 mcM/L.

**Discussion**

As hyperammonemic encephalopathy may be present even in patients with normal hepatic function, a high level of suspicion of encephalopathy is paramount in order to reach a timely diagnosis and to not miss the possibility of reversion and cure. Clinical presentation can be very variable and symptoms are usually episodic. Initial signs of HE may be inversion of sleep...
pattern, mild confusion, lethargy, and personality changes. There may be asterixis. However, HE can develop into somnolence, disorientation, marked confusion, and even coma. If left untreated, it may lead to intracranial hypertension, seizures, and death [2].

Early diagnosis relies on measuring plasmatic ammonia level in suspected patients. Although plasmatic ammonia levels do not always correlate directly with the degree of encephalopathy, subsequent dosages may be used to monitor treatment response, and clinical improvement usually accompanies decreasing levels of blood ammonemia [2, 3].

Image studies such as CT and magnetic resonance imaging (method of choice) may help establishing diagnosis. Typical findings of hyperammonemia are hyperintense lesions in the globus pallidus [4, 5]. Other image study modalities, such as magnetic resonance spectroscopy, single-photon emission CT, and positron emission tomography may add some information on the degree of impairment of cerebral water homeostasis, metabolic changes, brain edema, and astrocyte dysfunction and contribute to assessment of the severity and prognosis of neurological impairment [5–7].

Another useful diagnostic modality is the electroencephalography. Common findings are abnormal power of theta activity in mild or latent encephalopathy and low mean dominant frequency identified by biphasic power spectrum (delta and theta peaks) or high power delta activity in more severe cases [8–11].

As long as the diagnosis HE persists, proper treatment must be initiated, and it depends of identifying and understanding the myriad of conditions and physiopathological pathways that may lead to HE in patients with normal liver function.

Fibrolamellar hepatocellular carcinoma is a rare tumor of unknown etiology that almost always arises in non-cirrhotic livers of young adults without chronic viral hepatitis. It was first described in 1956 by Edmonson as a subtype of hepatocellular carcinoma [12]. Since 2009, this specific type of hepatic tumor has been associated with hyperammonemia in the literature [13–17].

There is no consensus in the literature about the exact causative physiopathological mechanism of hyperammonemia in patients with FHC. Some authors propose that a portosystemic shunt resulting from large tumors occupying significant portions of the liver may impair the hepatic capacity of nitrogen waste clearance [13, 14].

However, despite the fact that our patient had an 18 cm tumor occupying almost the entire right hepatic lobe with tumor thrombosis of the right portal hepatic veins, the left hepatic lobe was substantially enlarged probably as a result of compensatory hypertrophy. This compensatory hypertrophy of the left liver along with normal reticulocyte production index, serum lactic dehydrogenase, unconjugated bilirubin, and stable hematocrit suggested no significant portosystemic shunt bypassing the liver. It also demonstrated regenerative capacity of the non-tumoral liver that requires healthy parenchyma. Pathological examination of the liver from biopsies that were performed during the exploratory laparoscopy showed no signs of chronic hepatic disease and there were no laboratory or physical signs of hepatic failure. All these factors raise questions regarding why the liver lost only the capacity to perform nitrogen waste clearance and no other energy consuming metabolic pathways were affected.

Another explanation to hyperammonemia in patients with hepatic cancer is chemotherapy-related toxicity [15–17, 18]. Several chemotherapy agents have been associated with HE, such as oxaliplatin, vincristine, 5-fluorouracil, cyclophosphamide, methotrexate, etoposide, and gemicitabine [15]. Nevertheless, our patient had never received chemotherapy before developing HE.

As previously reported in the literature, a more convincing explanation to the development of HE in patients with FHC is a disorder in the urea cycle [15–17]. The urea cycle (also known as ornithine cycle) was described by Krebs and Henseleit in 1932, being the first metabolic cycle discovered [19]. It is responsible for converting waste nitrogen from protein digestion and protein catabolism into urea, which is then excreted by the kidneys. The urea cycle consists of five consecutive reactions, two mitochondrial and three cytosolic, and converts one amino group from ammonia, two molecules of nitrogen from ornithine, and one molecule from aspartate to urea. Two transporters (citrin and ornithine transporter-1) and a cofactor enzyme (N-acetyl glutamate synthetase) are also involved in the urea cycle [20]. Therefore, a total of eight enzymes are involved in this metabolic cycle:

- N-acetylglutamate synthetase (NAGS);
- Carbamoyl phosphate synthetase (CPS1);
- Citrin (aspartate glutamate translocase);
- Argininosuccinate synthetase (ASS);
- Argininosuccinate lyase (ASL);
- Arginase (ARG);
- Ornithine translocase (ORNT1);
- Ornithine transcarboxylase (OTC).

Disorders in the urea cycle are deficiencies of any of these factors and result in the accumulation of ammonia and other precursor metabolites. In fact, OTC deficiency has been previously associated with FHC [15–17, 21]. However, the specific reason for the suppression of OTC enzyme activity or for a deficiency of ornithine availability and consequent decreased ornithine cycle functioning has never been determined [15, 22].
The answer to that question probably began to be solved in 1972 by Weber and colleagues [23]. They demonstrated in rats a decrease in OTC activity reaching 1% of that observed in normal livers of control animals parallel to the increase in the hepatomas growth rate. This decrease had a close link to the increase in hepatomas growth rates, thus providing support to the Molecular Correlation Concept described by Weber and Lea [24]. A decrease in the urea cycle functioning determined by the reduction on OTC activity results in a decline in the utilization of aspartate and carbamoyl phosphate that could be spared for biosynthesis of DNA and RNA and thus become a biological advantage to the hepatic tumors.

It was also demonstrated that, despite the fact that OTC is not a rate-limiting enzyme of the urea cycle, its metabolic location in the cycle puts it in competition with other enzymes that use ornithine as a substrate. This has been shown to be the case with ornithine decarboxylase (ODC) [23].

Ornithine decarboxylase catalyzes the decarboxylation of ornithine, which is the first rate-limiting and committed step in polyamines synthesis, particularly for putrescine, spermine, and spermidine molecules that are indispensable for mammalian cell growth. They are key substrates for DNA stabilization and repair and are important antioxidants. So, ornithine decarboxylase activity is essential for cell growth and the reduction of this enzyme may induce apoptosis in DNA damaged cells [25]. Weber et al. also demonstrated an imbalance in the activities and ratios of OTC and ODC. While in normal rat livers the OTC activity is very high compared to ODC, in hepatomas there is a proportional reduction on OTC parallel to increase in ODC activity [23]. Thus, the ratio of activity of ODC/OTC rise along with the increase in hepatomas growth rates, resulting in an augmented utilization of ornithine for polyamine synthesis and the resulting nucleic acid biosynthesis by hepatoma cells. This imbalance becomes progressively increased parallel to the tumor growth rate, as showed by the Molecular Correlation Concept of an imbalance in enzymatic activities in opposing and competing metabolic pathways involved in tumor growth [26]. Therefore, in contrast to recent publications on the association of FHC and HE that focused on a probable paraneoplastic effect of the tumor on the direct reduction on the activity of OTC, the hypothesis is that an augmented activation of ODC results in the consumption of ornithine, and resulting urea cycle disturbance (Figure 4). However, the reason for this ODC activation is still missing in this clinical scenario.

This takes us back to the characteristics of the FHC itself. The etiology of this tumor is unknown and there is limited data on its pathophysiology. Elevations on blood alpha fetoprotein in patients with FHC are less likely to occur than in patients with traditional hepatocellular carcinoma and the genome sequencing of such tumors (mutations, pathways and structural variants) have demonstrated that it is in fact a distinct disease [27,28].

The analysis of fresh frozen specimens of FHC has demonstrated the presence of a single, recurrent heterozygous deletion of chromosome 19 that resulted in a functional chimeric protein of the heat shock protein DNAJB1 and a catalytic subunit of protein kinase A, PRKACA [27–29]. This finding was very sensitive and specific (100% of the FHC presented this chimeric transcript) for this tumor [28–30].

The lack of a high background of mutations throughout the genome of this tumor and the absence of an identifiable second hit mutation necessary to carcinogenesis suggests that the chimeric DNAJB1-PRKACA kinase is necessary and probably sufficient for the tumorigenesis of FHC [26]. The most probable molecular pathway involved is that of the DNAJB1-PRKACA chimera, which results in changes of AURKA expression within the tumor [30,31]. Transcriptome sequencing has demonstrated increased expression of AURKA (a known oncogene) in FHC samples [31].

It has been previously demonstrated that AURKA accumulations in the nucleus of hepatocellular carcinomas upregulates c-Myc transcription by binding to its promoter, which contains a high conserved CCCTCCCCA in the NHE region of CpG islands [32]. C-Myc is a regulator gene that codes a multifunctional nuclear phosphoprotein that is a transcription factor of paramount importance to cell cycle progression, apoptosis, and cellular transformation. It was first described in patients with Burkitt lymphoma [33]. A persistent expression of c-Myc leads to overexpression of many genes involved in cellular proliferation, resulting in carcinogenic effects [32]. AURKA and c-Myc mediate each other’s expression at the transcriptional level, affecting cellular proliferation, growth, and ATP production, and playing an important role in the carcinogenesis of hepatic tumors [32].

So, increased expression of AURKA correlates with that of c-Myc.

Finally, one of the c-Myc oncogene targets is ODC [34]. Overexpression of ODC secondary to c-Myc signaling resulting in increased polyamines has been observed in many tumors samples and is one of the first and most important carcinogenic steps in a variety of cancers [35].
To summarize, a recurrent heterozygous deletion of chromosome 19 common to all FHCs results in an increased expression of AURKA. This leads to an overexpression of c-Myc that upregulates ODC function that consumes ornithine in polyamines synthesis. The consequence is a reduction in intracellular ornithine bioavailability and consequent decreased urea cycle functioning resulting in HE.

In our particular patient, a multi-gene panel genetic testing for inborn errors of metabolism was performed, including the OTC gene, and no pathological variants were observed (Figure 3). He also presented very elevated plasmatic histidine level (451.0 mcMol/L, reference interval 6.0–250.0 mcMol/L) and low serine plasmatic level (19.0 mcMol/L, reference interval 20.0–120.0 mcMol/L) that could be indicative of ODC overexpression.

As the fibrolamellar hepatocellular carcinoma presents increased expression of AURKA, the chemotherapeutic regimen option for this patient was sorafenib (a multikinase inhibitor) and GEMOX. Two months after beginning chemotherapy, a CT of the abdomen and thorax disclosed stable disease. Four months after the acute encephalopathy episode and treatment with sodium benzoate and arginine was introduced, ammonia blood levels remained low and there were no other signs of HE.

**Conclusions**

Hyperammonemic encephalopathy in patients with normal hepatocellular function is a potentially fatal complication of rapidly growing liver tumors. A high level of suspicion is necessary to prompt diagnosis, and adequate treatment relies on the proper understanding of the physiopathological mechanism involved in order to reach complete remission of encephalopathy. We advise that all patients with large hepatic tumors should have ammonia blood level dosed at initial evaluation and repeatedly during treatment. We propose that an overexpression of AURKA, which occurs in all FHCs, is the initial key event that results in ornithine metabolism imbalance that leads to a urea cycle disorder resulting in hyperammonemia and encephalopathy.
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References:


Statement

All authors declare that they have no conflicts of interest.