Clinical Reasoning: A 14-year-old girl with headache, seizures, and confusion

Lijun Xiao, MD, Wenping Gu, MD, PhD, Bin Jiao, MD, PhD, Yunhai Liu, MD, PhD, and Xiaosu Yang, MD, PhD


Correspondence
Dr. Gu
Guwenping393@sohu.com

Section 1

A 14-year-old girl without any relevant medical history was transferred to our institution due to worsening headache along with nausea, vomiting, and generalized tonic-clonic (GTC) seizures for 7 days. She was also noted to have multiple psychological and behavioral abnormalities for 1 day. Her headache was described as severe holocephalic pain aggravated when lying down and alleviated after vomiting. No throbbing or phonophobia/photophobia was observed. Her seizures occurred once or twice a day, lasting about 1 minute each and resolving spontaneously. Symptoms were refractory to rotundine (dopamine D1 receptor antagonist), azasetron (antiemetic), mannitol, oxcarbazepine, and phenobarbital.

Psychological and behavioral abnormalities were noted including singing, raving, crying, and laughing intermittently. On physical examination, exotropia, hyperpigmentation of the skin, and long fingers were observed. On neurologic examination, the patient was unable to follow commands but responded to noxious stimuli. Funduscopic examination revealed papilledema. Plantar responses were extensor bilaterally.

D-dimer was high (2.72 mg/L, normal <0.5 mg/L), which suggested hypercoagulation and thrombogenesis. CT of head and intracranial arteries showed cerebral infarction and subarachnoid hemorrhage (figure 1, A) without arterial aneurysm or vascular malformation. Lumbar puncture revealed an opening pressure that was elevated at 300 mm of water, elevated red blood cells (>1,000 cells/mm³), and elevated white blood cells (about 20 cells/mm³). CSF glucose, chloride, and protein were normal.

A continuous EEG showed diffuse background slowing with epileptiform discharges including frequent spikes and sharp waves present over the bilateral frontal and temporal regions. CT venography (CTV) revealed cerebral venous sinus thrombosis (CVST) with filling defects of the sigmoid and right lateral sinuses (figure 1, B). There was no family history of headache, epilepsy, or CVST.

Question for consideration:
1. What is the cause of the patient’s CVST?
Figure 1 Images of the reported case

(A) Brain CT shows cerebral infarction (triangle), subarachnoid hemorrhage (bold arrow), and thrombus of superior sagittal sinus (thin arrow). (B) CT venography (CTV) reveals cerebral venous sinus thrombosis (CVST) with filling defect of the right sigmoid sinus and right lateral sinuses (white arrow). (C) CTV after a month of treatment; the right sigmoid sinus and right lateral sinuses unobstructed but narrowed (white arrow). (D) X-ray suggests that joint space of bilateral wrist was narrowed. (E) The patient's long fingers. The circles show hyperpigmentation of skin, especially in the interphalangeal joints. (F) Histopathologic study of the patient's slightly darker skin reveals hyperkeratosis (yellow arrowhead) and verrucous hyperplasia in the basal layer of the epidermis (white arrowhead) (magnified 10 × 10 times under the microscope). (G) Pedigree of the family with homocystinuria. (H) Gene direct sequencing identified 2 novel compound heterozygous mutations: c.551T>C (p.L184P) and c.949A>G (p.R317G) (indicated by the red arrow). I-1: Heterozygous mutations of c.949A>G, the c.551 TT is normal; I-2: heterozygous mutations of c.551T>C, the c.949AA is normal; II-1, II-2: compound heterozygous mutations of c.551T>C and c.949A>G. (I) Orthologous protein sequence alignment of cystathionine β-synthase from different species. The mutated residue showing conservation is shaded in red. Red shaded amino acids proteins show that the 2 novel missense mutations occurred at highly conserved positions in these species. DD = Dictyostelium discoideum; HS = Homo sapiens; MM = Mus musculus; OC = Oryctolagus cuniculus; RN = Rattus norvegicus; splice isoform III; SC = Saccharomyces cerevisiae; TC = Trypanosoma cruzi.
Section 2

The primary cause for CVST is usually hypercoagulability due to acquired and genetic risks (table 1). The patient had no evidence of common risks in table 1 such as infection, vitamin deficiency, abnormal protein C or protein S, estrogen-related factors, or others. However, she did have high level of plasma total homocysteine (Hcy) (149.40 μmol/L, normal <15 μmol/L).

Hcy causes approximately 4.5% of cases of CVST. It is a nonessential but indispensable sulfur-containing amino acid in humans. Hcy levels in the bloodstream can rise in 3 different ways: (1) protein structure modifications; (2) oxidative stress induction; and (3) excitotoxicity. Hyperhomocysteinemia (HHcy) can lead to CVST and thromboembolism by causing prothrombotic conditions, endothelial dysfunction, and impaired thrombolysis. HHcy can also result in multisystem damage (table 2) by the 3 main pathways mentioned above.

### Table 1 Causes of cerebral venous sinus thrombosis

<table>
<thead>
<tr>
<th>Acquired risks</th>
<th>Genetic risks</th>
</tr>
</thead>
<tbody>
<tr>
<td>Connective tissue disease</td>
<td>The prothrombin G20210A mutation</td>
</tr>
<tr>
<td>Hematologic system diseases</td>
<td>Factor V Leiden gene mutations</td>
</tr>
<tr>
<td>Infections</td>
<td>Protein C, protein S, and antithrombin III deficiencies</td>
</tr>
<tr>
<td>Cancer</td>
<td>Genes associated with homocysteine metabolism</td>
</tr>
<tr>
<td>Pregnancy and puerperium</td>
<td>Others</td>
</tr>
<tr>
<td>Surgery, trauma, and other mechanical precipitants</td>
<td>Unidentified</td>
</tr>
<tr>
<td>Exogenous hormones</td>
<td>Unidentified</td>
</tr>
<tr>
<td>Drug-associated conditions</td>
<td>Unidentified</td>
</tr>
</tbody>
</table>

**Table 2** Multiple target organs or systems damage caused by hyperhomocysteinemia reported in the literature and the 2 patients

<table>
<thead>
<tr>
<th>Target organs/systems</th>
<th>Clinical characteristics reported in the literature</th>
<th>II-1 (older sister)</th>
<th>II-2 (proband)</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Eye</strong></td>
<td>Ectopia lentis, exotropia, or severe myopia</td>
<td>Congenital binocular ectopia lentis, impaired vision</td>
<td>Congenital binocular ectopia lentis, exotropia, impaired vision</td>
</tr>
<tr>
<td><strong>Skeleton</strong></td>
<td>Dolichostenomelia and arachnodactyly, osteoporosis and higher risk of fractures, bone deformities</td>
<td>Long fingers, kyphoscoliosis</td>
<td>Long fingers, the physiologic curvature of thoracic vertebra straightened, narrowed joint space of bilateral wrist, osteoporosis, decreased BMD (values −3.3)</td>
</tr>
<tr>
<td><strong>Skin (not common)</strong></td>
<td>Light skin and brittle red to blond hair, malar flush</td>
<td>Not found</td>
<td>Hyperpigmentation, especially in the interphalangeal joints, histopathologic study revealed hyperpigmentation, hyperkeratosis, verrucous hyperplasia in the basal layer of the epidermis</td>
</tr>
<tr>
<td><strong>CNS</strong></td>
<td>Intellectual disability, psychiatric and behavioral problems, seizures, and extrapyramidal sign</td>
<td>Psychological and behavioral abnormalities, irascibility, intellectual disability, ataxia, unstable gait</td>
<td>Intellectual disability, scored 70 on WISC-IV, 25 of 30 on MMSE, 17 of 30 on MoCA, psychological and behavioral abnormalities, low self-learning ability</td>
</tr>
<tr>
<td><strong>Vascular system</strong></td>
<td>Venous or arterial thromboembolism</td>
<td>Not found</td>
<td>Cerebral venous sinus thrombosis</td>
</tr>
<tr>
<td><strong>Others</strong></td>
<td>Congenital defects including congenital heart defect and neural tube defect</td>
<td>Not found</td>
<td>EMG, ECG, cardiac color ultrasound, peripheral artery and peripheral venous examinations all normal</td>
</tr>
</tbody>
</table>

Abbreviations: BMD = bone mineral density (value below −2.5 means higher risk of fractures); MMSE = Mini-Mental State Examination; MoCA = Montreal Cognitive Assessment; WISC-IV = Wechsler Intelligence Scale for Children, 4th edition.
In this case, the patient had CVST and multisystem dysfunction due to HHcy (table 2), including ocular anomalies, skeletal system deformities (figure 1, D), skin diseases (figure 1, E and F), and CNS abnormalities.

**Questions for consideration:**
1. What are the common causes of HHcy?
2. How can we confirm the root cause of HHcy in our patient?
Section 3

The common causes for HHcy can be divided into exogenous factors such as overintake of cysteine (Cys) or methionine (Met) or dietary deficiencies of vitamins B6, B12, or folate (Figure 2). Exogenous factors of HHcy were considered unlikely—there was no history of dietary restrictions or overuse of vitamins. However, the patient’s older sister (II-1 in the pedigree of Figure 1G) also had multiple medical problems (Table 2). Thus, the patient’s HHcy may have been due to inherited factors, which tend to be a considerable cause of HHcy.

For our patient, we first screened for common genetic defects and biochemical abnormalities. Investigations revealed elevated plasma Met (437.77 μmol/L) without a normal level of methylmalonic acid on plasma aminoacid chromatography. These results were consistent with cystathionine β-synthase (CBS) and methylenetetrahydrofolate reductase (MTHFR) deficiency (Table 3).

To confirm the diagnosis, an experimental study was conducted using all exons of CBS and MTHFR genes in the pedigree (Figure 1, G). We found novel compound heterozygous mutations of the CBS gene, c.551T>C (p.leucine 184 proline) and c.949A>G (parginine 317 glycine) (Figure 1, H), which were coseparated and predicted as damaged and possibly damaged according to SIFT (sift.jcvi.org/) and PolyPhen (genetics.bwh.harvard.edu/pph2/). Interestingly, the 2 novel mutations occurred at highly conserved positions across different species (Figure 1, I).

To our knowledge, this is the first time this has been reported in the literature, including from the National Heart Blood and Lung Institute Exome Sequencing Project, 1000 Genomes Project, or the Single Nucleotide Polymorphism Database, and 50 healthy controls from

![Figure 2 The biochemical metabolism of homocysteine (HCy)](A) Hcy and serine can transfer to cystathionine, and the reactions are catalyzed by vitamin B6 and cystathionine β-synthase (CBS) resynthesis. (B) Remethylation to methionine of Hcy pathways. Vitamin B12, betaine, and betaine homocysteine methyltransferase (BHMT) are independent. (C) In the cycle of methylenetetrahydrofolate (THF), vitamin B2 and methylenetetrahydrofolate reductase (MTHFR) are indispensable. Cys = cysteine; Met = methionine; MS = methionine synthase; MSR = methionine synthase reductase; SAM = S-adenosyl methionine; SAH = S-adenosyl homocysteine.

### Table 3 Different genes/proteins deficiency causing different variations of organic acids

<table>
<thead>
<tr>
<th>Genes deficiency</th>
<th>Protein</th>
<th>Hcy (normal 15 μmol/L)</th>
<th>Met (normal 50 μmol/L)</th>
<th>MMA (normal &lt; 4 μmol/L)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Methionine synthase</td>
<td>MS</td>
<td>↑</td>
<td>↓</td>
<td>—</td>
</tr>
<tr>
<td>Methionine synthase reductase</td>
<td>MSR</td>
<td>↑</td>
<td>↓</td>
<td>—</td>
</tr>
<tr>
<td>Cystathionine β-synthase</td>
<td>CBS</td>
<td>↑</td>
<td>↓</td>
<td>—</td>
</tr>
<tr>
<td>Methylenetetrahydrofolate reductase</td>
<td>MTHFR</td>
<td>↑</td>
<td>↑</td>
<td>—</td>
</tr>
<tr>
<td>Genes deficiency associated with cobalamin (MMACHC, MMADHC, ABCD4, HCFC1)</td>
<td>Protein associated with cobalamin</td>
<td>↑</td>
<td>—</td>
<td>↑</td>
</tr>
</tbody>
</table>

Abbreviations: ABCD4 = ATP-binding cassette-D 4; HCFC1 = host cell factor c-1; MMACHC = methylmalonic acid homocystinuria type C protein; MMADHC = methylmalonic acid homocystinuria type D protein.
Whole-exome sequencing bioinformatics was performed and confirmed that there were no other potential associated genes mutation in this pedigree. Therefore, these results indicate that the compound heterozygous mutations of CBS gene, c.551T>C and c.949A>G, which have not been previously reported, had a strong relationship with homocystinuria in this pedigree.

**Question for consideration:**
1. What is the next step in management? How should one treat the CVST? How should one treat the HHcy?
**Section 4**

For CVST, low-molecular-weight heparin was given to the patient followed by oral anticoagulants. For HHcy, the current treatment strategies primarily focus on correcting the biochemical abnormalities with supplementation of vitamin B6 (100–200 mg/d), folic acid (5 mg/d), and vitamin B12 (5 mg/d), along with a Met and Cys restricted diet. Betaine is another possible treatment for these patients as it provides an alternate remethylation pathway to convert excess Hcy to Met (figure 2), especially in patients who could not achieve target levels of Hcy by other means.

One month later, the CTV image showed that the previously obstructed sinuses were unobstructed but narrowed (figure 1, C). Three months later, the patient attended school, as her learning ability and behavior had improved. After a year, the patient had no further thrombotic events and a normal Hcy. In addition, the patient was counseled about birth control and avoiding estrogen-containing contraceptives, which may increase risk of thrombosis including CVST. If the patient were to have a future pregnancy, prophylactic anticoagulation with low-molecular-weight heparin would be recommended postpartum and during the third trimester to reduce the risk of thromboembolism.

**Discussion**

We report a case of CVST due to homocystinuria. Homocystinuria due to CBS deficiency (OMIM #236200) is a rare disorder of sulfur amino acid metabolism, with elevated plasma concentrations of tHcy and Met and increased excretion of Hcy in urine. So far, at least 164 mutations in CBS have been reported (cbs.lf1.cuni.cz/index.php). In our case, novel compound heterozygous pathogenic mutations of CBS, c.551T>C and c.949A>G, led to HHcy causing CVST and multiple system dysfunction, skin hyperpigmentation, and straightened vertebra.

The case expands the phenotypes and mutation spectrum of CBS resulting in homocystinuria. This discovery is helpful in presymptomatic molecular diagnosis, prenatal diagnosis, management of patients with homocystinuria, and genetic counseling of families. The treatment management and corresponding prognosis indicate that early diagnosis, early prevention, and early treatment are beneficial to prognosis. Gene detection technique is of great value to diagnose disease and to find the new gene mutation in clinical genopathy and it should be widely used in clinical practice.

This case also highlights that (1) for CVST in young patients without common risk factors, HHcy is a possible cause; and (2) for patients with HHcy, once exogenous factors of HHcy are ruled out, genetic testing should be done, especially if there is evidence of multiorgan dysfunction or a strong family history.

**Author contributions**

Lijun Xiao: study design, acquisition of clinical data, analysis of data, drafting the manuscript. Wenping Gu: study concept and design, analysis of data, revising the manuscript. Bin Jiao: statistical analysis, drafting the manuscript. Yunhai Liu: acquisition of clinical data, study supervision. Xiaosu Yang: study concept and design, analysis of data.

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**References**

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