Update on benign convulsions with mild gastroenteritis

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Benign convulsions with mild gastroenteritis (CwG) are characterized by afebrile convulsions associated with viral gastroenteritis in previously healthy infants and children. The main causative pathogens are rotavirus and norovirus. CwG occurs frequently in both East Asian and Western countries. The prevalence of CwG was reportedly not decreased by the introduction of rotavirus vaccines, and the prevalence of norovirus-associated CwG has been increasing annually. Convulsions in CwG are usually clustered, do not last longer than 5 minutes, and are mostly generalized. Laboratory diagnostics, electroencephalography (EEG), and imaging findings are usually normal. There is a probability of mild, transient abnormal findings on EEG or imaging limited to the acute disease phase. Although several reports have suggested that pathogens that affect the central nervous system through direct or indirect mechanisms could be related to the pathophysiology of CwG, its mechanism is not fully understood. Several antiepileptic drugs are effective during convulsions; however, long-term antiepileptic treatment is not required as CwG usually has a good prognosis.

Key words: Seizures, Gastroenteritis, Rotavirus, Norovirus

Epidemiology

The ratio of RaCwG to rotavirus gastroenteritis (RVGE) is approximately 1.29% to 5.5% (Table 1).12,13,18,23,24 The ratio of RaCwG to CwG was reportedly 48.4% to 55.8%, but the data in these reports were compiled before rotavirus vaccines were widely used.17,25 One study showed that the ratio of NaCwG to NVGE was higher than that of RaCwG to RVGE.24 However, other studies reported a lower ratio.13,29 Meanwhile, the ratio of NaCwG to CwG was reportedly 14.5% to 5.5% (Table 1).3,7,25 The ratio of NaCwG to CwG was reportedly 14.5% to 5.5% (Table 1).3,7,25

Introduction

Acute gastroenteritis (AGE) is a common disease in pediatric patients, affecting more than 100 million children annually and sometimes requiring hospitalization.1,2 Rotavirus and norovirus are the main causes of AGE, and several reports have shown that these viruses can cause central nervous system (CNS) complications such as meningitis, encephalitis, and encephalopathy.1-7 Benign convulsions with mild gastroenteritis (CwG) was first reported by Morooka9 in 1982 as mild gastroenteritis that causes convulsions without fever, severe dehydration, electrolyte imbalance, and hypoglycemia. In 1995, Komori et al.9 reported the following characteristics: (1) afebrile convulsions generated within 5 days of acute viral gastroenteritis in previously healthy infants and children between the ages of 6 months and 3 years; (2) absence of moderate or severe dehydration; (3) possible recurrence of seizures over several days; (4) normal serum electrolyte levels, cerebrospinal fluid (CSF) analysis results, and blood glucose level; (5) a good prognosis; and (6) usually caused by rotavirus. Since then, several reports have broadened the range of definitions for patients aged <6 years with CwG.10-16 CwG is frequently reported in East Asian countries as well as in the United States and Europe.17,18 After the introduction of rotavirus vaccines, some changes in epidemiology and clinical characteristics occurred in rotavirus-associated CwG (RaCwG) as well as norovirus-associated CwG (NaCwG).12,13,18,19 There have also been several recent reports on the diagnosis and treatment of CwG.20-22 Therefore, this review includes recent reports of CwG and covers the epidemiology, clinical manifestations, pathophysiology, biomarkers, and treatment with the aim of promoting the recognition and management of CwG.
Our recent study of nationwide data from health insurance reviews and associated services in South Korea showed that the prevalence of RaCwG was 0.013%–0.024% and that of NaCwG was 0.00001%.

We also found that the annual prevalence of RVGE decreased significantly after rotavirus vaccines were introduced in Korea; however, the prevalence of RaCwG did not change. In contrast, the annual prevalence of NaCwG is increasing (Fig. 1).

Geographically, the prevalence of RVGE decreased irrespective of the region after the introduction of rotavirus vaccines, but the ratio of RaCwG to RVGE increased in urban areas (Fig. 2).

However, these studies have limitations because they did not compile data directly from patient charts; rather, they defined patients using only diagnostic codes. Therefore, further multicenter trials are necessary to determine the actual prevalence of these conditions.

CwG commonly occurs in children aged <6 years and peaks at age 12–24 months. The ratio of male to female patients with CwG was reportedly 1:0.7–2.6. In addition, it mainly occurs between October and May. After rotavirus vaccination, the age at RaCwG onset repor-
Comparisons between RaCwG and NaCwG showed that the ratio of males to females with RaCwG was 1:0.9–1.8,10,12,13,19,23 while that of those with NaCwG was 1:1.2–2.2.13,15. RaCwG commonly occurred between December and April, especially in winter,10,14,19 while NaCwG commonly occurred between November and December. NaCwG was less prevalent than RaCwG in spring.14,15

Clinical manifestations

In CwG, seizures occur within 6 days of intestinal symptoms.11,14,15,17,23,27,28,30,31,33,36 The interval between enteric symptom onset and seizure onset was reportedly longer in RaCwG than in NaCwG14 and in RaCwG than in febrile seizures associated with rotavirus infection.23

One of the clinical characteristics of CwG is that between 2–4 and up to 20 episodes of clustered seizures occur.11,14,15,17,23,26,30,31,34,36 and 35%–70.9% of CwG patients experienced clustered seizures within 24 hours of viral infection.27,33-35 The number of clustered seizures and total seizure frequency in CwG was greater than the number of febrile seizures with gastroenteritis.31

Each seizure episode usually lasts less than 5 minutes.11,15,17,23,27,31,34,36 Seizure semiology in CwG is mostly generalized, although focal seizures are occasionally reported.11,15,17,23,27,31,34,36 In particular, focal seizures were more frequently observed in patients with CwG than febrile seizures in patients with gastroenteritis.31 Seizure relapse was also reported in CwG at a rate of approximately 4.7% (~7.7%); however, there were no cases of the subsequent development of epilepsy following CwG.11,32,33

Although one of the characteristics of CwG is no abnormal findings in laboratory tests, some authors have conducted research with broader characteristic criteria of CwG.14,17,30 Several studies demonstrated that CwG may be associated with hyperuricemia or hyponatremia.29,34,37 A recent study found that severe ketosis occurring in patients with CwG is associated with lower blood glucose and sodium levels.35 Serum aspartate aminotransferase levels were elevated in patients with RaCwG compared with patients with RVGE without seizures or patients with febrile seizure associated with RVGE.23 Lower serum Ca²⁺ levels were also found in RaCwG patients (9.22±0.50 mg) than in patients with rotavirus gastroenteritis without seizure (9.66±0.46 mg); therefore, the author suggested that the disruption of Ca²⁺ homeostasis plays a critical role in triggering seizures.38

Although electroencephalography (EEG) findings are usually normal in CwG,11,30 some reports have shown abnormal EEG findings such as a focal onset of spikes and slow waves.14,15,17,23,27,28,31,33,34 Abnormal EEG findings, mostly spikes in the asymmetric central region, are more frequent in CwG than in febrile seizures.31 In addition, posterior slowing was observed more frequently in NaCwG than in RaCwG.14 However, these abnormal findings normalized during follow-up EEG taken 2–6 months later.11

Brain magnetic resonance imaging (MRI) findings are usually normal;11,17,27,28,30,31 however, some reports stated that focal abnormal lesions in the splenium of the corpus callosum appeared transiently.23,31,36,39-42 In particular, rotavirus infection was a risk factor for splenial lesions in patients with CwG.36 A reversible splenial lesion on MRI resembles clinically mild encephalitis/encephalopathy with a reversible splenial lesion including the following diagnostic criteria: (1) onset of neuropsychiatric symptoms within one week after the onset of fever; (2) complete recovery without sequelae, mostly within 10 days after the onset of neuropsychiatric symptoms; (3) high-signal-intensity lesion in the splenium of the corpus callosum in the acute disease stage; (4) lesion that may involve the entire corpus callosum and the cerebral white matter in a symmetric fashion; and (5) lesion disappears within a week, with neither
residual signal changes nor atrophy. Therefore, caution is needed when diagnosing CwG with reversible splenial lesions.\(^{43,45}\) Abnormal findings consistent with vasogenic edema were observed in NaCwG,\(^ {46}\) which normalized during the follow-up period (Table 2).\(^ {36,39,42,46}\)

### Pathophysiology

Despite many reports on the pathophysiological mechanism of CwG, it remains unclear. Pathogens detected in CwG patients include rotavirus, norovirus, adenovirus, astrovirus, and sapovirus; of them, adenovirus, astrovirus, and sapovirus are rarely detected.\(^ {14,15,47}\) Several studies have attempted to identify the mechanisms of specific pathogens of rotavirus and norovirus.\(^ {3,47-57}\) Rotavirus has 2 outer capsid proteins, VP4 and VP7, which are used to classify rotavirus as G and P genotypes.\(^ {58}\) Two studies investigated the association between the CwG and rotavirus genotypes, but they are controversial. Yang et al.\(^ {47}\) analyzed the genotypes of 13 patients with RaCwG and found that none were associated with the disease. However, Choi et al.\(^ {40}\) investigated rotavirus genotypes in 82 patients with rotavirus infection and found that G2P was significantly associated with neurologic complications. Thus, further large-scale studies are required to clarify this relationship.

Several studies have sought to identify evidence regarding the direct invasion of rotavirus into the CNS. In the animal model and the sera of children, rotavirus antigenemia was confirmed.\(^ {49,51}\) This result suggests that rotavirus can penetrate the gastrointestinal tract and cause viremia.\(^ {49,51}\) Rotavirus has been shown to infect neuronal cells in vivo and in vitro,\(^ {52,59}\) and rotavirus RNA was detected in the CSF of patients with RVGE and convulsions.\(^ {50,51}\) Another hypothesis is that mediators may indirectly induce CwG. One study showed that NO metabolite levels in the serum and CSF from rotavirus gastroenteritis patients with convulsions were much higher than those in patients with purulent meningitis, encephalitis, febrile convolution, or in the healthy control group.\(^ {60}\) These findings indicate that NO may have a pathophysiological role in convulsions associated with rotavirus infection. Moreover, cytokines can indirectly affect the brain. Levels of interleukin (IL)-17 in particular are increased in convulsions following gastroenteritis versus rotavirus encephalopathy, indicating that IL-17 is a convulsion-related factor.\(^ {61}\) The endoplasmic reticulum-associated rotavirus protein, VP7, is used to indicate sorting signals to target vascular proteins on axons through nonclassical Golgi-independent mechanisms. These signals can also be detected in viral targets in different areas of the brain.\(^ {52}\)

Nonstructural protein 4 (NSP4), a glycosylated protein of rotavirus, plays a leading role in the pathogenic mechanism of rotavirus by increasing cell membrane permeability to Ca\(^ {2+}\).\(^ {53}\) Yeom et al.\(^ {54}\) investigated serum anti-NSP4[A] immunoglobulin G (IgG) and anti-NSP4[B] IgG in patients with RVGE and found that levels of both antibodies increased in the nonseizure versus seizure group. Therefore, the authors suggested that serum anti-NSP4 IgG might protect against rotavirus-associated seizures.

On the other hand, some studies revealed the pathophysiology of norovirus-associated neurologic complications and found evidence of norovirus penetration into the gastrointestinal tract; in fact, its RNA was found in the serum and CSF.\(^ {3,55-57}\) However, research on the pathophysiology of NaCwG is still lacking.

Molecular genetic studies aimed to determine the factors contributing to the occurrence of CwG. One study examined the association between CwG and channelopathy, as benign infantile convulsions were considered channelopathies by other molecular genetic studies.\(^ {62-65}\) However, no mutations in the neuronal sodium channel alpha 1 subunit (SCN1A) gene were found in patients with CwG.\(^ {66}\) Mutations in the proline-rich transmembrane protein 2 (PRRT2) gene are known to cause benign familial infantile epilepsy, paroxysmal kinesigenic dyskinesia, and infantile convulsions with paroxysmal choreoathetosis.\(^ {67,70}\) However, no mutations in PRRT2 were found in CwG patients.\(^ {71}\) A case report recently suggested that mutations in the alpha-II-spectrin (SPTAN1) gene, which is involved in early-onset epileptic encephalopathy with progressive brain atrophy, severe intellectual disability with cerebellar malformations, and relatively milder phenotypes with or without epilepsy, were found in patients with CwG.\(^ {72}\) Thus, more molecular genetic studies are required, including those on SPTAN1 gene mutations.

### Table 2. Clinical characteristics of benign convulsions with mild gastroenteritis

| Characteristic                                      | Value                                      |
|----------------------------------------------------|--------------------------------------------|
| Family history of epilepsy/febrile seizure         | 9.1%–17.1%                                 |
| Interval between enteric symptoms and first seizure| 0–6 Days after enteric symptoms             |
| Seizure type                                        | Usually generalized seizures or focal seizures |
| Seizure frequency                                   | Usually 2–4 episodes                        |
| Seizure duration                                    | Usually within 5 minutes                    |
| Interval between the first and last seizure         | Within 24 hours                             |
| Laboratory findings                                | Usually normal, hyperuricemia and relatively low levels of sodium or Ca\(^ {2+}\) may be observed |
| Cerebrospinal fluid                                | Normal                                     |
| Electroencephalogram                               | Usually normal                             |
| Radiological examinations                          | Usually normal                             |
| Prognosis                                           | Good                                       |

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Biomarker for differential diagnosis

An attempt was made to use S100B protein and neuron-specific enolase (NSE) as biomarkers. S100B is an intracellular calcium-binding protein that is dominant in astrocytes.79 Several studies have suggested that serum S100B levels are associated with cerebral parenchymal damage.74-78 NSE is a -y- subunit of enolase, a major component of the neuronal cytosol.79 Chen et al.20 compared serum S100B and NSE protein levels in 46 patients with CwG versus 35 patients with febrile convulsions. They found that serum NSE levels were significantly higher in the CwG group than in the febrile seizure group; however, S100B levels were not significantly different. This feature was not associated with the number of convulsions or total convolution duration in the CwG or febrile seizure groups. Therefore, the authors suggested that NSE could play a role in the differential diagnosis of CwG and febrile convulsions associated with mild gastroenteritis.

Treatment and prognosis

Several studies aimed to identify well-tolerated antiepileptic drugs in the acute phase of CwG. One randomized, single-blind, placebo-controlled trial suggested the efficacy of intravenous phenobarbital 10 mg/kg.20 However, another study compared phenobarbital to lidocaine and showed a significantly higher efficacy for the latter.80 Another study reported that 1-day treatment with carbamazepine or lidocaine effectively treated clustering seizures in CwG.81 Similarly, another study showed that 1-day treatment with low-dose carbamazepine was effective for CwG.82 Some studies demonstrated that fosphenytoin and chloral hydrate are also effective for CwG.83,84 In addition, a recent study showed that a single dose of lacosamide was well tolerated in patients with CwG.83 Although clustering seizures frequently occur in the acute phase of CwG, the prognosis is favorable, and antiepileptic therapy is not required after seizure cessation.

Conclusions

CwG is an afebrile seizure associated with mild gastroenteritis that occurs during healthy infancy and early childhood. Rotavirus and norovirus are its main causative pathogens. After the introduction of the rotavirus vaccine, the prevalence of CwG did not decrease, whereas that of NaCwG has increased annually. CwG is characterized as a generalized cluster seizure for which EEG and MRI show transiently abnormal findings that normalize. Some differential features were found in the clinical manifestation, EEG, and MRI findings between RaCwG and NaCwG. Its pathophysiology remains unclear; however, there is evidence of direct or indirect effects of pathogens on the CNS. Recent efforts have been made to design biomarkers for the differential diagnosis of CwG and other convulsions. Some antiepileptic drugs are effective in the acute phase of CwG, and long-term treatment is usually not required. Further studies are necessary to determine the pathogenesis of CwG, and continuous monitoring of epidemiological changes is required.

Footnotes

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