Inability to Reduce Morbidity of Diarrhea by ORS: Can We Design a Better Therapy?

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Abstract

Diarrheal disease is a worldwide problem that still causes significant morbidity and mortality among children. Currently, oral rehydration solution (ORS) is the standard of care for acute diarrhea in pediatric patients. While effective in reducing mortality, ORS does not alleviate diarrheal symptoms, thus reducing caregiver compliance and therapeutic efficacy. This manuscript will briefly review the current problem of pediatric diarrhea and the shortcomings of current therapies; however, the focus of this review is to examine the intestinal calcium-sensing receptor (CaSR). The author summarizes the evidence suggesting that targeting the CaSR will enable clinicians to address all four major pathophysiological mechanisms of diarrheal disease, and substantiates the need for future research regarding this therapy.

History of Oral Rehydration Solution Use in Diarrhea

Despite a decrease in worldwide diarrhea-related mortality over the past 30 years, diarrheal disease remains the fourth leading cause of death in children under the age of five with 499,000 dying in 2015. While mortality has decreased 39.2% from 2005-2015, morbidity has not declined to a comparable degree. In 2015, there were 2.39 billion estimated cases of diarrhea. Of these cases, approximately 957.5 million occurred in children under the age of 5 (1). In the United States from 2001-2006, the cost of hospitalizations, emergency room visits and outpatient visits for diarrhea in pediatric patients was approximately $10 billion per year, not including indirect costs for the family. Notably, 77% of children visit an outpatient clinic for diarrhea by the age of five (2). The most common cause of acute diarrhea is infectious gastroenteritis, however, the majority of diarrheal-related deaths are the result of dehydration and electrolyte imbalance, not infection (3).

The development of the current oral rehydration therapy dates back to over fifty-years ago. In 1958, Robert K. Crane and his colleagues discovered that intestinal Na+ and glucose...
absorption are coupled. The researchers also learned that Na\(^+\)/glucose cotransport plays a significant role in the reabsorption of water (4). Understanding this mechanism proved vital in developing a treatment for Cholera. While Cholera toxin effectively inhibits many intestinal electrolyte transporters, Na\(^+\)/glucose cotransport remains undisturbed. In 1968, researchers working in Bangladesh and India exploited this principle and formulated a mixture of glucose, sodium, potassium, and chloride, now called oral rehydration solution (ORS) (5).

Cholera causes secretory diarrhea; the toxin increases fluid secretion and decreases fluid absorption in the intestine. The alpha-subunit of the Cholera toxin constitutively activates Adenyl Cyclase, increasing the amount of cyclic adenosine monophosphate (cAMP) in the cell, which in turn stimulates Protein Kinase A (PKA). When activated, PKA phosphorylates the Cystic Fibrosis Transmembrane Conductance Regulator (CFTR) channel, causing secretion of Cl\(^-\) and water. In addition, high levels of cAMP inhibit NaCl absorption, Na\(^+\)/H\(^+\) exchange, and Cl\(^-\)/HCO\(_3\) exchange, resulting in decreased electrolyte and water absorption (6, 7). Na\(^+\)/glucose cotransport is cAMP independent; therefore, solute and water can be reabsorbed despite the presence of high cAMP levels. Certain strains of Escherichia coli and other bacteria produce enterotoxins that cause secretory diarrhea in a similar fashion. Consequently, clinicians have expanded the use of ORS to treat diarrhea-related dehydration caused by pathogens other than Cholera, including viruses.

The pathophysiology of rotavirus gastroenteritis can be partially attributed to a viral-encoded enterotoxin NSP4, which also results in excessive Cl\(^-\) secretion. However, this secretion is distinct from increased cAMP dependent CFTR activity and the identity of the Cl\(^-\) channel remains unknown. Importantly, NSP4 also impairs intestinal cells through the inhibition of Na\(^+\)/glucose cotransport, the main mechanism by which ORS enhances absorption (8). As a result, ORS therapy may not be as effective in patients infected by rotavirus due to this inhibition.

Currently, ORS is the only oral therapy available for acute diarrhea in children. The use of ORS has significantly decreased mortality from diarrheal disease, however, its usage has not increased but decreased since 2000, with fewer than 33% of children with diarrhea under the age of 5 using the therapy (9). Often, caregivers are reluctant to use ORS because it takes a relatively long time to produce positive results. Additionally, despite attempts to modify it, including lowering osmolality to reduce the risk of increased diarrhea volume, ORS does nothing to prevent excessive secretion of fluid, the major mechanism by which Cholera and other infectious organisms cause diarrhea. In fact, the glucose contained in ORS stimulates calcium-activated chloride secretion in the small intestine, transiently increasing diarrhea volume (10). Again, this translates to decreased levels of caregiver compliance and to the misconception that ORS is not an effective therapy.

**Current Anti-diarrheal Therapies**

There are four categories of anti-diarrheal treatment: pro-absorptive, anti-secretory, anti-motility, and anti-inflammatory (Figure 1). Clinical management of acute diarrhea in pediatric patients is purely pro-absorptive and involves replacement of fluid using ORS or
parenteral rehydrating solution (3). Currently, there are no approved therapies in the United States that stop fluid loss or decrease gut motility which are efficacious and safe to use in children. Several experimental therapies have been considered. Loperamide, an anti-motility drug that has been used with success in adults, can cause serious side effects such as ileus, lethargy, and death in children (11). A promising CFTR and calcium-activated chloride channel (CaCC) inhibitor, Crofelemer, proved to be a weak and partial antagonist of CFTR (12), limiting its potential positive effects against diarrhea caused by excessive CFTR activity.

Although ORS decreases mortality from dehydration and electrolyte imbalance, it does not alleviate symptoms. ORS does not target cAMP/cGMP accumulation or the influence of the enterotoxin NSP4, the major mechanisms causing secretory diarrhea by numerous bacteria and rotavirus respectively. Patients continue to experience diarrhea in spite of treatment. As a result, many clinicians and caregivers turn to anti-microbial agents (7). While antibiotics are useful in some cases of acute bacterial diarrhea, they must be used cautiously in order to prevent pathogen resistance and opportunistic infection. Certain enteric organisms such as salmonella should not be treated with antibiotics, as they can prolong the duration of illness (13). Most importantly, antibiotics do not immediately prevent fluid and electrolyte loss, the most common cause of diarrhea-related mortality.

While ORS is an effective lifeline for those experiencing acute diarrhea, it is not curative. Even while consuming ORS, patients experience severe symptoms and can suffer recurrent episodes of diarrhea. Children with frequent diarrheal disease are more likely to be malnourished and have impaired growth and cognitive development (14). Malnutrition is known to increase the incidence and duration of diarrhea, thus creating a viscous cycle (15). Unfortunately, there are no current therapies that address this issue in a cost effective manner. A pediatric therapy that both lessens diarrheal excretion and rehydrates is urgently needed.

A Promising New Target for Therapy

A promising target for such a therapy is the extracellular Calcium-Sensing Receptor (CaSR). CaSR is a G protein coupled receptor that is expressed in various human tissues including the small and large intestine. This receptor primarily functions to maintain calcium homeostasis and to detect luminal nutrients including: calcium, polyamines, L-amino acids, and oligopeptides (16). Stimulation by such a variety of agonists can result in multiple distinct active conformations of the receptor, allowing for different biological effects to take place in the cell (17).

CaSR has been studied in mammals, amphibians, birds, and marine species. In the intestine of marine organisms, CaSR has been found to prevent water loss to hyperosmotic seawater surroundings (18). This function is conserved in human intestine and is the basis for the development of CaSR targeted anti-diarrheal therapy (7). Inside the intestine, CaSR has been located in secreting crypt epithelium, absorbing surface epithelium, motility-modulating enteric nervous tissue, and inflammatory cells (19, 20, 21, 22, 23). Researchers investigating the role of CaSR in these cells have discovered activation of the receptor decreases cellular
cyclic nucleotides and reverses all four pathological changes occurring in diarrhea: excessive secretion, malabsorption, increased gut motility, and a heightened inflammatory response.

**Understanding the Calcium-Sensing Receptor**

The two major mechanisms of secretory diarrhea, increased secretion and decreased absorption of electrolytes and fluid, are both altered by CaSR activation (16, 22, 23, 24, 25, 26). Studies using rat colonic epithelium show CaSR activation counteracts the effects of Cholera toxin, *Escherichia coli* heat stable enterotoxin STa and other secretagogues by inhibiting cyclic nucleotide accumulation and anion secretion (24). By stimulating the enzyme phosphodiesterase (PDE), CaSR leads to the degradation of cyclic nucleotides. Decreased cAMP/cGMP concentrations reduce CFTR translocation to the apical membrane and enhance both Na\(^+\)/H\(^+\) and Cl\(^-\)/HCO\(_3\)^\(^-\) exchange. Thus, CaSR activation simultaneously normalizes fluid excretion and reabsorption.

Cyclic nucleotides also play a critical role in regulating the enteric nervous system (ENS), which controls about 50% of fluid secretion in infectious diarrhea (27). When stimulated by cAMP, endocrine intestinal cells secrete neurotransmitters, such as serotonin and neurotensin. These peptides are activators of the ENS and initiate an enteric neural response through the stimulation of submucosal plexus neurons, which ultimately augment secretion and inhibit absorption (6). Similarly, the ENS is activated in rotavirus-induced diarrhea. It has been postulated that NSP4 may bind to enterochromaffin cells, prompting the release of serotonin, leading to stimulation of the ENS and fluid loss (8).

When mammalian small intestine is treated with Cholera toxin, fluid secretion is reduced by two neurotoxins: tetrodotoxin (TTX) and hexamethonium (6). TTX blocks voltage gated Na\(^+\) channels, while hexamethonium prevents acetylcholine from binding to nicotinic receptors; both effectively suppress the ENS. A study by Cheng et al. examined the effect of CaSR on ENS-mediated fluid secretion by analyzing short circuit current (I\(_{sc}\)) through rat colons. The group applied secretagogues to the epithelium with and without TTX. I\(_{sc}\) increased to a greater extent in the absence of TTX, indicating fluid secretion is greater without ENS inhibition. Later, CaSR was activated using R568, a calcimimetic. Interestingly, CaSR activation decreased I\(_{sc}\) in tissue that was not treated with TTX. When TTX was administered, R568 did not significantly reduce I\(_{sc}\), suggesting that the stimulated CaSR reduces fluid secretion by manipulating ENS activity (22).

Finally, a study examining CaSR knockout mice showed that CaSR activation promotes an anti-inflammatory response in the intestine that could prove to be beneficial in diarrhea caused by enterocolitis and inflammatory bowel disease. CaSR knockout mice have decreased expression of tight junction molecules, such as claudin 2, 8 and 14, which help form a strong intestinal barrier. Additionally, their intestinal barrier function is compromised, as seen by significantly reduced transepithelial electrical resistance and increased flux to fluorescein isothiocyanate-conjugated dextran (28). This leaky barrier allows microbes to easily traverse from the intestinal lumen through the paracellular space into the lamina propria, activating an inflammatory response. Mice without CaSR intestinal expression also have reduced secretion of anti-microbial peptides that protect against
bacterial translocation from the lumen into the blood. A gene array analysis of CaSR−/− mouse colonic epithelium showed an increase in expression of several proinflammatory cytokine-encoding genes, including IL-1β, TNF-α, and IL-6, further supporting the role of CaSR in intestinal immune homeostasis (28).

An Improved Oral Rehydration Solution

CaSR is a very encouraging target for diarrheal disease, as activation can reduce symptoms caused by all four pathophysiological mechanisms, while other drugs target only one. Fortunately, CaSR agonists are easily accessible, inexpensive, and safe to use in the pediatric population. The primary ligand for CaSR is calcium, an essential nutrient that is commonly found and fortified in foods. Allosteric activators of CaSR include the polyamine spermine that is found in breast milk and the amino acid tryptophan. CaSR adopts multiple active conformations, as it is constantly exposed to these agonists in the intestine. Adding nutrients to ORS would be simple and could possibly increase the efficacy of the current therapy. Small additions of spermine can dramatically enhance activation of CaSR, even when calcium is at a physiologically normal level (25). Secondarily, polyamines and amino acids would provide essential nutrition to malnourished children suffering from diarrhea. Different agonist combinations should be investigated to identify which combinations and doses are the most therapeutic with the fewest adverse effects.

As expected, studies have shown that these nutrients are safe to use. Intake of oral calcium slightly above the recommended daily allowances (RDA) has shown positive effects, reducing diarrhea in preliminary clinical trials (29, 30). It is probable that CaSR activation played a role in reducing diarrhea volume and accelerated the recovery period in these patients. A transient increase in calcium intake did not alter blood-calcium homeostasis. The patients urinary and blood calcium levels remained normal, and the only noticeable side effect was mild constipation that subsided with completion of treatment. Calcium absorption from the intestine is a process that is heavily influenced by the presence of vitamin D. Without additional vitamin D, the majority of extra calcium remains in the intestine, locally stimulating CaSR without systemic effects.

While these results are promising, the clinical trials were completed in very specific patient populations and cannot be generalized. Some of the patients included were hypocalcemic to begin with, therefore, increasing their calcium intake slightly above the RDA is unlikely to have a negative effect. It is probable that a calcium concentration above the RDA would be necessary to produce a sufficiently potent therapy even with the addition of amino acids and polyamines, creating a potential safety concern. Randomized controlled trials are needed to establish a concentration of calcium that is both effective and safe. Calcimimetic drugs, such as the chemical compound R568, are more potent and could have therapeutic effects at smaller concentrations (24). Nevertheless, calcium is a more attractive remedy because it is widely available and inexpensive. A new formulation of ORS with calcium and other nutrient CaSR agonists could more easily be implemented as standard treatment around the world.
Conclusion

The CaSR is a promising target for pediatric anti-diarrheal therapy. In contrast to conventional ORS, CaSR based treatment is a four-pronged approach, addressing all major pathophysiological mechanisms of diarrhea. Initial clinical studies have produced positive results, demonstrating that a CaSR-ORS would be safe, affordable, and possibly more effective than the current gold standard. By relieving the symptoms associated with diarrheal disease, CaSR-ORS could inspire more confidence in caregivers, increasing overall compliance and reducing mortality worldwide.

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Figure 1. Four pathways leading to the formation of diarrhea
A: Impaired absorption, B: Excessive secretion, C: Intestinal barrier breakdown and enhanced inflammation, D: Overly active enteric nervous system. CaSR activators reduce changes in all the four diarrhea-causing pathways. CaCC: Calcium-activated chloride channel; CFTR: Cystic fibrosis transmembrane conductance regulator; CTX: Cholera toxin; ENS: Enteric nervous system; IEC: Intestinal epithelial cell; NTS: Neurotransmitters