Mechanism of Action of Veverimer: A Novel, Orally Administered, Nonabsorbed, Counterion-Free, Hydrochloric Acid Binder under Development for the Treatment of Metabolic Acidosis in Chronic Kidney Disease

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ABSTRACT
Current management of metabolic acidosis in patients with chronic kidney disease (CKD) relies on dietary intervention to reduce daily endogenous acid production or neutralization of retained acid with oral alkali (sodium bicarbonate, sodium citrate). Veverimer is being developed as a novel oral treatment for metabolic acidosis through removal of intestinal acid, resulting in an increase in serum bicarbonate. Veverimer is a free-amine polymer that combines high capacity and selectivity to bind and remove hydrochloric acid (HCl) from the gastrointestinal (GI) tract. In vitro studies demonstrated that veverimer had a binding capacity of 10.7 ± 0.4 mmol HCl per gram of polymer with significant binding capacity (>5 mmol/g) across the range of pH values found in the human GI tract (1.5–7). Upon protonation, veverimer bound chloride with high specificity but showed little or no binding of phosphate, citrate, or taurocholate (<1.5 mmol/g), which are all anions commonly found in the human GI tract. Administration of veverimer to rats with adenine-induced CKD and metabolic acidosis resulted in a significant increase in fecal chloride excretion and a dose-dependent increase in serum bicarbonate to within the normal range compared with untreated controls. Absorption, distribution, metabolism, and excretion studies in rats and dogs dosed with 14C-labeled veverimer showed that the polymer was not absorbed from the GI tract and was quantitatively eliminated in the feces. Acid removal by veverimer, an orally administered, nonabsorbed polymer, may provide a potential new treatment for metabolic acidosis in patients with CKD.

SIGNIFICANCE STATEMENT
Metabolic acidosis is a complication of chronic kidney disease (CKD) as well as a cause of CKD progression. Veverimer is a high-capacity, selective, nonabsorbed, hydrochloric acid–binding polymer being developed as a treatment for metabolic acidosis. Veverimer binds and removes hydrochloric acid from the gastrointestinal tract, resulting in increased serum bicarbonate and the correction of metabolic acidosis. Veverimer is not an ion-exchange resin and does not deliver sodium or other counterions, and so it may be appropriate for patients with CKD with and without sodium-sensitive comorbidities.

Introduction
Metabolic acidosis is a common disorder in patients with non–dialysis-dependent, stage 3–5 chronic kidney disease (CKD) and is caused by the inability of the diseased kidney to quantitatively remove daily endogenous acid production (Alpern and Sakhaee, 1997; Hamm et al., 2015; Kraut and Madias, 2016). The result of this imbalance is that acid accumulates in the body and serum bicarbonate, the major extracellular acid buffer, and pH both fall from their normal ranges of 22–29 mEq/l and 7.36–7.44, respectively (Wesson et al., 2020). Chronic metabolic acidosis is recognized clinically as a persistent reduction of serum bicarbonate to less than the lower limit of normal, which is generally 22 mEq/l, in a patient with CKD and normal pulmonary function (Kraut and Madias, 2018; Raphael, 2018).

The human body continually generates acid through metabolism of the daily diet. Sources of acid include amino acids and nucleic acids, incomplete oxidation of carbohydrates, and...
demonstrated a high capacity and selectivity for HCl binding and removal in the GI tract, leading to an increase in serum bicarbonate. Veverimer binds and removes HCl without exchanging the proton for a cation, thus representing a novel means of increasing acid excretion in patients with CKD and metabolic acidosis.

Materials and Methods

Chemicals and Reagents. Veverimer, which was previously designated TRC101 (Bushinsky et al., 2018), was synthesized at Tricida, Inc. (South San Francisco, CA). Veverimer is a highly crosslinked, aliphatic amine polymer with the chemical structure presented in Fig. 1; the chemical name for veverimer is poly(allylamine-co-N,N′-diallyl-1,3-diaminopropane-co-1,2-diaminoethane). Veverimer was synthesized by first copolymerizing two monomers: allylamine hydrochloride [Chemical Abstract Service (CAS) Registry Number 10017-11-5] and N,N′-diallyl-1,3-diaminopropane dihydrochloride (CAS Registry Number 205041-15-2), followed by crosslinking of the polymer with 1,2-dichloroethane (CAS Registry Number 107-06-2). Veverimer has an intrinsic high binding capacity and selectivity for chloride. The hydrochloric acid binding capacity derives from the high amine content of the monomer components comprising the polymer, and the extensive crosslinking in veverimer provides size-exclusion properties and selectivity for binding chloride over larger anions.

Physicochemical Characterization of Veverimer. The particle size distribution of veverimer was determined via laser light diffraction with a Malvern Mastersizer 3000 particle size analyzer (Malvern Instruments, Malvern, UK). In each measurement, a veverimer sample (250 mg) was weighed into a closable 15-ml plastic tube, and 5 ml of methanol was added while vortexing at low speed. The tube was closed tightly, and vortexing continued at a higher speed for an additional 30 seconds. An aliquot of this suspension was then used to measure the particle size. The average for d(0.1), d(0.5), and d(0.9) of three sample preparations as a whole number (in micrometers) was recorded.

The solubility of veverimer was evaluated in a total of 12 test media covering a range of solvent polarities as well as in test media reflecting relevant physiologic conditions veverimer encounters during transit through the GI tract (Table 1). Water, methanol, and n-heptane were evaluated to represent aqueous and organic media, respectively.

In this report, we provide the initial in vitro and in vivo characterization of veverimer, a nonabsorbed HCl-binding polymer under development for the treatment of metabolic acidosis and slowing of kidney disease progression in patients with CKD (Bushinsky et al., 2018; Wesson et al., 2019a,b). Veverimer is composed of low-swelling, spherical polymeric beads that are approximately 100 μm in diameter. Size exclusion built into the three-dimensional structure of the polymer enables preferential binding of chloride versus larger inorganic and organic anions, such as phosphate, citrate, fatty acids, and bile acids. Using a rat model of CKD, veverimer
Anion concentration of in vitro test matrices designed to mimic GI tract compartments

<table>
<thead>
<tr>
<th>Matrix</th>
<th>Acetate (mM)</th>
<th>Chloride (mM)</th>
<th>Phosphate (mM)</th>
<th>Citrate (mM)</th>
<th>Taurocholate (mM)</th>
<th>Oleic Acid (mM)</th>
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<tbody>
<tr>
<td>SGF</td>
<td>0</td>
<td>98.0</td>
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<tr>
<td>SIB</td>
<td>0</td>
<td>36.0</td>
<td>20.0</td>
<td>0</td>
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<td>0</td>
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<tr>
<td>SOB</td>
<td>50.0</td>
<td>36.0</td>
<td>7.0</td>
<td>1.5</td>
<td>5.0</td>
<td>30.0</td>
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</tbody>
</table>

**TABLE 2**

Anion concentration of in vitro test matrices designed to mimic GI tract compartments

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**Mechanism of Action of Veverimer**

**In Vitro Hydrochloric Acid Binding Assays.** The in vitro chloride binding capacity and specificity of veverimer were assessed in three biologic matrices mimicking the environment of different GI compartments: 1) simulated gastric fluid (SGF) that mimics the acidic environment of the stomach (pH 1.2) and reflects an optimal condition for binding of HCl; 2) simulated intestine inorganic buffer (SIB) with a pH of 6.5; and 3) simulated intestine organic and inorganic buffer (SOB) with a pH of 6.2. After incubation, the polymer was isolated and washed with 10 ml of deionized water. The recovered polymer was next incubated in SOB buffer at 37°C for 2 hours; after incubation, the polymer was washed and with 10 ml of deionized water. Finally, the recovered polymer was incubated in a solution of 0.2 M sodium hydroxide solution at 37°C for 18 hours to elute anions, which were quantitated by IC analysis.

**pH-Dependent Equilibrium Chloride Binding.** To determine the pH-dependent equilibrium chloride binding, veverimer (4 mg/ml) was incubated for 18 hours in 100 mM sodium chloride solution at pH 3.0, 6.0, and 9.0. The pH values maintained in the individual test solutions ranged from 1.5 to 7.0. After incubation, the chloride content was determined by IC analysis.

**GI Compartment Transit Assay.** An in vitro GI compartment transit assay (GCTA) was used to measure the binding and retention of various anions to veverimer under conditions simulating the passage of the polymer through various compartments of the GI tract. Sequential incubation in three different matrices simulated the pH, ionic environment, and residence times associated with transit through different GI compartments (i.e., stomach, distal small intestine/colon, and colon). After incubation in each of the three matrices, all remaining bound ions were eluted from veverimer and quantitated by IC (ICS-2100; Thermo Scientific).

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**pH-Dependent Equilibrium Chloride Binding.** To determine the pH-dependent equilibrium chloride binding, veverimer (4 mg/ml) was incubated for 18 hours in 100 mM sodium chloride solution (i.e., test solution). The sample was continuously stirred during the incubation, and the pH of the solution was maintained at a predetermined value by slowly adding 0.1 N HCl using an automatic titrator (T50; Mettler Toledo). The pH values maintained in the individual test solutions ranged from 1.5 to 7.0. After incubation, the chloride content in the supernatant was measured by IC (ICS-1600; Thermo Scientific).

Equilibrium chloride binding to veverimer was calculated by determining the difference between the amount of chloride added during incubation and the amount remaining in the supernatant at the end of the study.

**Animal and Ethics Approval.** All animal experiments were conducted in accordance with the Guide for the Care and Use of Laboratory Animals from the Institute for Laboratory Animal Research, National Research Council and as adopted and promulgated by US National Institutes of Health and approved by the vendor.
Veverimer Proof-of-Concept Efficacy Study in Rats. Veverimer was evaluated in vivo in an adenine-induced rat model of CKD and metabolic acidosis (Diwan et al., 2018). The study was designed in two parts (Fig. 2). In both parts, male Sprague-Dawley rats (7 to 8 weeks, 260–280 g) were obtained from Charles River Laboratories (Hollister, CA) for the purposes of this study and used to evaluate the pharmacodynamics of veverimer in the CKD model with respect to fecal chloride binding and effect on serum bicarbonate. The animals were pair-housed in a temperature-controlled facility under a 12-hour light/12-hour dark regimen. Diet and water were provided ad libitum. Adult Wistar Han rats and Beagle dogs as well as juvenile Sprague-Dawley rats were used for bioavailability assessments of [14C]-veverimer absorption, distribution, metabolism, and excretion and were tested and housed at MPI/Charles River (Mattawan, MI).

Study diet

<table>
<thead>
<tr>
<th>Study group</th>
<th>Part 1</th>
<th>Part 2</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>No treatment</td>
<td>No treatment</td>
</tr>
<tr>
<td>2</td>
<td>1.5 wt% veverimer</td>
<td>No treatment</td>
</tr>
<tr>
<td>3</td>
<td>3.0 wt% veverimer</td>
<td>0.0 wt% veverimer</td>
</tr>
<tr>
<td>4</td>
<td>4.5 wt% veverimer</td>
<td>No treatment</td>
</tr>
</tbody>
</table>

Sample collection

- Blood draw for measurement of serum bicarbonate and blood pH
- Fecal collection for anion measurement

Study week

1 2 3 4 5 6 7 8 9 10 11 12 13

Part 1 of the study investigated the effect of early treatment with veverimer administered to rats on a casein diet with 0.25 wt% adenine for the 4 weeks after the 2-week induction period. Part 2 of the study investigated the effect of delayed treatment with veverimer; in this part, after the 2-week induction period, rats were maintained on a casein diet containing 0.25 wt% adenine for 5 weeks prior to initiation of veverimer administration and through the dosing phase with veverimer. During the veverimer treatment periods, rats were fed diets admixed with 0, 1.5, 3.0, and 4.5 wt% veverimer. After correcting for body weight and food consumption, the 1.5, 3.0, and 4.5 wt% veverimer treatment groups received mean veverimer doses of 0.8, 1.5, and 2.4 g/kg per day, respectively, in part 1 of the study and 0.7, 1.4, and 2.3 g/kg per day, respectively, in part 2 of the study.

Study parts 1 and 2 both assessed the effect of withdrawing veverimer after the end of the treatment phase, with a 2-week withdrawal phase in which veverimer was discontinued in the low (1.5 wt%) and high (4.5 wt%) veverimer dose groups but was continued in the mid-dose group (3.0 wt%). All animals received a casein diet with 0.25 wt% adenine during the withdrawal phase.

Animals were randomized based on serum bicarbonate levels at baseline (i.e., after adenine induction of nephropathy and before initiation of the dosing period) so that mean baseline serum bicarbonate levels were comparable across all dose groups. Body weights and food consumption were determined weekly, allowing calculation of actual administered dose. For both part 1 and part 2 of the study, blood samples were obtained from the rats before initiation of veverimer dosing, then weekly during the dosing and withdrawal phases. Samples were analyzed for serum bicarbonate levels and pH values. In addition, a 24-hour fecal sample was obtained from each animal in the control and high-dose groups during weeks 4 and 6 of part 1 of the study and weeks 9 and 11 of part 2 of the study. Anions were extracted from lyophilized and homogenized fecal samples by incubating the samples with NaOH for 18 hours at 37°C. Sample supernatants were analyzed for chloride and phosphate by IC (ICS-2100, Thermo Scientific).

Bioavailability Assessment. The bioavailability of veverimer was assessed in pharmacokinetic studies conducted in Wistar Han rats and Beagle dogs by administering [14C]-radiolabeled veverimer as a single oral dose to male and female animals at 0.0735 g/kg. The target radioactive dose was 100 μCi/kg. Absorption was characterized by measuring levels of radioactivity in the blood. The doses used in the absorption studies were selected to be representative of doses that will...
be administered to human subjects, including pediatric subjects. In addition, to determine whether bioavailability differed between juvenile and adult animals because of different stages of GI maturity an oral absorption study was conducted in juvenile Sprague-Dawley rats. Rat pups on postnatal days 7 and 11 (n = 18/sex per group) were administered [14C]-radiolabeled veverimer at a total labeled/unlabeled combined veverimer dose of 0.15 g/kg (target radioactive dose = 100 μCi/kg). Serial blood samples were taken from each animal before dosing and at 0.5, 1, 2, 4, 6, 24, and 48 hours postdose in the adult rat before dosing and at 0.5, 1, 2, 4, 6, 12, 24, 48, 72, 120, and 168 hours postdose in the dog and 1, 2, 4, 8, and 24 hours postdose in the juvenile rat. Absorption was characterized by measuring levels of radioactivity in whole blood (dog and juvenile rat) and/or plasma (dog, adult rat, and juvenile rat) using liquid scintillation counting (Packard 2300TR; PerkinElmer, Waltham, MA).

To examine distribution of veverimer, adult Wistar Han rats (n = 7/sex per group) received a single oral administration of [14C]-veverimer at a target dose level of 0.0735 g/kg (100 μCi/kg). At each time point evaluated (0.5, 1, 6, 24, 48, 120, and 168 hours postdose), rats (n = 1/sex per time point) were euthanized, and carcasses were subjected to quantitative whole-body autoradiography using a Fuji FLA-5100 fluorescent image analyzer (Fuji Photo Film Co., Ltd., Japan).

To determine excretion of veverimer, adult Wistar Han rats (n = 3/sex) and adult Beagle dogs (n = 3/sex) received a single oral administration (via gavage) of [14C]-veverimer at 0.0735 g/kg (100 μCi/kg). Animals were placed in metabolic cages immediately after dosing, and urine, feces, expired air (rat only), and cage washings were collected predose and at the following postdose time intervals:

- Urine (Rat and Dog): 0–6 (rat) or 0–12 (dog), 6–24 (rat) or 12–24 (dog), 24–48, 48–72, 72–96, 96–120, 120–144 and 144–168 hours.
- Expired air (rat): 0–24, 24–48, and 48–72 hours.
- Cage washings (dog): 24, 48, 72, and 168 hours.

Samples were analyzed for radioactivity, and additional samples for radioactivity measurements were taken from carcasses. All rat and dog samples were analyzed using liquid scintillation counting (Packard 2300TR; PerkinElmer).

Results

**Veverimer Particle Size Distribution and Solubility in Aqueous and Organic Solvents.** The physical form of veverimer is that of a free-flowing powder composed of low-swelling, spherical beads (Fig. 3). Each bead is a single, high–molecular weight, crosslinked polyamine molecule. The particle size of veverimer was primarily controlled by the stir rate during the suspension polymerization reaction in which the beads are first formed to generate an average particle diameter of approximately 100 μm. Particle size distribution analysis showed a monomodal particle size distribution, wherein greater than 90% of the volume was composed of particles with a diameter larger than 40 μm (Fig. 3).

The solubility of veverimer was evaluated in a variety of aqueous and organic solvents and expressed as the amount of veverimer (milligram) dissolved in 1 l of solvent (Table 1). Apart from the 0.1 N HCl (veverimer solubility of 3 mg/l) and 50 mM acetate buffer (pH 4.5) with 2 wt% CTAB (veverimer solubility of 2 mg/l) matrices, veverimer was uniformly insoluble in the aqueous and organic solvents tested. In accordance with USP (General Notices and Requirements), the descriptive term of insoluble or practically insoluble is applied to any material wherein 10,000 or more parts of solvent are required for one part of solute, and thus veverimer is determined to be insoluble in aqueous and organic solvents and in media encompassing the physiologic pH range with and without surfactant.

**Characterization of Veverimer Hydrochloric Acid Binding In Vitro at Physiologically Relevant pH Values.** Three in vitro assessments of the chloride binding capacity of veverimer were conducted in test matrices that mimicked various GI compartments with respect to pH and anion composition (Table 2). Competing anions (e.g., phosphate, citrate, and taurocholate) were included in the two matrices simulating intestinal fluid (SIB and SOB). Table 3 summarizes the matrices used, the incubation times, the competing anions, and the binding of chloride and competing anions by veverimer. Binding studies in the test matrices demonstrated that veverimer bound chloride with high capacity and selectivity. Veverimer bound 10.7 ± 0.4 mmol of chloride per gram of polymer in SIB. In the SIB matrix, veverimer preferentially bound chloride over phosphate with binding capacities of 4.3 ± 0.1 mmol of chloride and 1.5 ± 0.3 mmol of phosphate per gram of polymer, respectively. In the most complex SOB matrix, veverimer preferentially bound chloride (3.8 ± 0.3 mmol/g of polymer) in the presence of various competing inorganic and organic anions such that the polymer essentially did not bind phosphate, citrate or taurocholate.

An in vitro GCTA was used to measure the binding and retention of various anions to veverimer under conditions simulating the passage of the polymer through distinct compartments of the GI tract. Sequential incubation in three different matrices simulated the pH, ionic environment and residence times associated with transit through the stomach (SGF buffer), distal small intestine/cecum (SOB buffer), and colon (retention buffer). Veverimer bound an average of 8.1 mmol of chloride per gram of polymer under simulated gastric conditions (pH 1.2, 1 hour, 37°C) and retained most of the bound chloride (7 mmol/g of polymer; approximately 85%) after being challenged by conditions designed to mimic the environment of the distal small intestine and cecum (pH 6.2, 2 hours, 37°C). After the third incubation under conditions representing the colon (pH 7.0, 42 hours, 37°C), approximately 60% of initially bound chloride (approximately 5 mmol chloride/per gram of polymer) was retained. Release of chloride was attributed to the pH-dependent decrease in the total binding capacity of the veverimer polymer. Veverimer bound very little or no measurable amounts of the other competing anions (i.e., acetate, phosphate, citrate, taurocholate, oleic acid, sulfate, and bicarbonate) present in the simulated human distal small intestine/cecum and colon matrices.

The pH-dependent equilibrium chloride binding of veverimer (4 mg/ml) was determined in 100 mM sodium chloride solutions (i.e., test solution) in which the pH of the solution was maintained at a predetermined value by slowly adding 0.1 N HCl. The pH values maintained in the individual test solutions ranged from 1.5 to 7.0. Although the chloride binding capacity for veverimer decreased with increasing pH, which was considered related to a decrease in veverimer protonation, veverimer maintained significant chloride binding capacity (i.e., >5 mmol/g of polymer) over a range of pH values (1.5–7.0) that are physiologically relevant for the GI tract (Fig. 4).

**Veverimer Effects on Serum Bicarbonate and Fecal Chloride Excretion in Rats with CKD and Metabolic Acidosis.** Veverimer was evaluated in a two-part in vivo study in male Sprague-Dawley rats (n = 10/group) administered...
adenine (0.75 wt% in casein diet) for 2 weeks to induce CKD (Fig. 2). The veverimer-related effects on serum bicarbonate and fecal excretion of chloride and phosphate are summarized in Figs. 5 and 6. Animals were randomized based on serum bicarbonate levels at baseline (i.e., after adenine induction of CKD and before initiation of the dosing period) so that mean baseline serum bicarbonate levels were similar across all dose groups. Endpoints evaluated in the study included serum bicarbonate and pH (weekly measurements) and fecal chloride and phosphate (24-hour samples measured at weeks 4 and 6 of part 1 and weeks 9 and 11 of part 2). The part 1 and part 2 rat studies were designed to accommodate a withdrawal period, with one group (3.0 wt%) maintained on continuous dosing as an internal control for the withdrawal groups tested (1.5 and 4.5 wt%).

Serum bicarbonate was elevated to within the normal range (23.0–31.1 mEq/l based on values for five age-matched rats) in veverimer-dosed rats but remained below normal in untreated CKD control rats. Serum bicarbonate levels generally remained within the normal range with continued veverimer dosing, whereas withdrawal of veverimer was associated with a decline in serum bicarbonate within 2 weeks to values that were generally less than those observed in control animals. The effect of veverimer on blood pH reflected the results observed for serum bicarbonate (unpublished data). A significant increase in fecal chloride was observed in veverimer-dosed rats relative to untreated controls. Although veverimer also significantly increased fecal phosphate excretion, the effect was notably less than that observed for chloride. The study results support the proposed veverimer mechanism of action (i.e., selective binding, retention and removal of HCl from the body through the feces), and the CKD model provides proof-of-concept that veverimer administration results in restoration of serum bicarbonate to the normal range in a dose-dependent manner.

### Veverimer Absorption, Distribution, Metabolism, and Excretion

The physicochemical properties of veverimer, including its insolubility in aqueous and organic solvents and particle size distribution, suggested that the polymer would not be absorbed from the GI tract. To test this hypothesis, the absorption, distribution, metabolism, and excretion of an administered pharmaceutical compound within an organism of [14C]-veverimer after oral administration of a single dose in both Wistar Han rats and Beagle dogs was assessed.

Absorption of [14C]-veverimer into the blood was not detectable in any adult rat plasma, dog blood, or juvenile rat blood or plasma sample. Radioactivity was detected in only one dog plasma sample at a single time point but was not considered to indicate absorption since the detectable level was close to the limit of detection, fell within the range of background sample radioactivity values, and was not observed in this dog at time points bracketing the sample. The results of the radiolabeled absorption studies indicated that veverimer was not systematically absorbed from the GI tract after oral administration to the juvenile or adult rat or the dog.

Distribution of veverimer was assessed by quantitative whole-body autoradiography in the adult rat. Radioactivity was not detectable in any tissue or organ other than within the GI tract, and therefore, tissue distribution could not be quantitated. The intestinal transit of the radiolabel was effectively complete by approximately 120 hours after

### TABLE 3

Chloride binding by veverimer in matrices mimicking the GI tract

<table>
<thead>
<tr>
<th>Matrix</th>
<th>Incubation Time (h)</th>
<th>Chloride (mmol/g Polymer)</th>
<th>Phosphate (mmol/g Polymer)</th>
<th>Citrate (mmol/g Polymer)</th>
<th>Taurocholate (mmol/g Polymer)</th>
</tr>
</thead>
<tbody>
<tr>
<td>SGB*</td>
<td>16</td>
<td>10.7 ± 0.4</td>
<td>N/A</td>
<td>N/A</td>
<td>N/A</td>
</tr>
<tr>
<td>SIB†</td>
<td>1</td>
<td>4.3 ± 0.1</td>
<td>1.5 ± 0.3</td>
<td>N/A</td>
<td>N/A</td>
</tr>
<tr>
<td>SOB‡</td>
<td>24</td>
<td>3.8 ± 0.3</td>
<td>0.1 ± 0.1</td>
<td>&lt;0.1†</td>
<td>&lt;0.1†</td>
</tr>
</tbody>
</table>

N/A, not applicable.

*SOB, pH = 1.2; it mimics the acidic environment of the fasted stomach and reflects an optimal condition for binding of HCl.
†SIB, pH = 5.5; pH is representative of the human proximal small intestine (duodenum and early jejunum), and that contains chloride and a high conc. of phosphate as a potential competing anion to chloride. The solution was buffered by 2-(N-morpholino)ethanesulfonic acid.
‡SOB, pH = 6.2; pH and competing anion content (bile acid, phosphate, citrate, oleic acid, and acetate) represent components of the human distal small intestine (late jejunum, ileum) and cecum. The solution was buffered by N,N-bis(2-hydroxyethyl)-2-aminoethanesulfonic acid.
*Phosphate conc. used to calculate phosphate binding capacity of veverimer lots were estimated by calculating the change in the phosphate peak area before and after incubating the polymer in SOB.
†Citrate and taurocholate peaks for veverimer lots were not well separated in the IC chromatogram because of method limitations. Combined peak area was used to estimate conc. of these anions before and after incubating the polymer in SOB. Oleic acid was not determined because it was not detected by the IC method used.
Excretion of veverimer was predominantly through the feces (Tables 4 and 5). Approximately 95%–97% of the administered radioactive dose in the rat and 96%–99% in the dog was recovered in the feces. Excretion of radioactivity was relatively rapid with most of the radioactivity (approximately 90%) excreted within the first 24 hours postdose in both species. In general, radioactivity was not detectable in any fecal or urine sample after 72 hours postdose in the rat or after 120 hours in the dog. Very small amounts of radioactivity were detected over the first 24-hour postdose period in rat urine and expired air (mean ≤0.02% of the administered dose, each matrix) and dog urine (mean = 0.02% of the administered dose). The amount in urine and/or expired air for both the rat and dog corresponded within the variability of the study to the unincorporated radiolabel remaining in the administered material [14C]-veverimer. In addition, some degree of cross-contamination of urine by feces was likely to have contributed to the very low levels of radiolabel detected in the urine of both species because complete separation of urine and feces is rarely completely achieved despite the use of metabolic cages. Thus, excretion data together with blood and plasma data demonstrated that veverimer was not systemically absorbed in rats and dogs after oral administration, with almost 100% of the recovered radioactivity in the feces.

Evaluation of veverimer metabolism in the rat and dog was not conducted based on the absence of systemic exposure. The in vivo pharmacology studies (summarized above) demonstrating binding and removal of chloride after oral administration of [14C]-veverimer. The lack of radioactivity in tissues or organs outside of the GI tract demonstrated that radiolabeled veverimer was not absorbed from the gut after oral administration to rats.

Fig. 4. Veverimer chloride binding through a range of pH values. Equilibrium chloride binding was assessed at different pH values for veverimer as described. Veverimer had a chloride binding capacity of 10.1 mmol/g under highly acidic conditions (i.e., pH 1.5) in which all the available free amine groups (primary, secondary, and tertiary) were protonated and bound to chloride as a counterion. As the pH of the incubation solution was increased, the equilibrium chloride binding capacity of veverimer decreased but still bound ~5.0 mmol/g of chloride under physiologically relevant conditions (i.e., pH 5.0–7.0).

Fig. 5. Effect of veverimer on serum bicarbonate and fecal excretion of chloride and phosphate in part 1 of proof-of-concept rat study. (A) Effect of veverimer on serum bicarbonate in part 1. Mean ± S.D. from blood samples collected at week 6 (treatment phase) and week 8 (withdrawal phase). The 3.0 wt% veverimer treatment group remained on veverimer, whereas 1.5 and 4.5 wt% veverimer treatment groups were withdrawn from drug during the withdrawal phase (weeks 7 to 8). Horizontal dotted lines mark the normal serum bicarbonate range for male Sprague-Dawley rats of the same age. Statistical analysis: two-way ANOVA with Dunnett’s multiple comparisons test vs. untreated group; N = 10 rats per group. (B) Fecal chloride excretion in rats during part 1 of the study. Mean ± S.D. from fecal samples collected at week 4 and week 6. Statistical analysis: unpaired, two-tailed t test; N = 10 rats per group. (C) Fecal phosphate excretion in rats during part 1 of the study. Mean ± S.D. from fecal samples collected at week 4 and week 6. Statistical analysis: unpaired, two-tailed t test; N = 10 rats per group.
in fecal chloride but only a 2-fold increase in phosphate fecal excretion relative to controls in rats with adenine-induced nephropathy administered veverimer suggested that the polymer retained its functional integrity (i.e., high capacity, selective chloride binding) and, therefore, structural integrity during transit through the GI tract after oral administration, supporting a lack of veverimer metabolism or degradation within the GI tract.

Treatment with veverimer did not cause diarrhea in the rats or dogs used in the absorption, distribution, metabolism, and excretion of an administered pharmaceutical compound within an organism assessment or in the rat model of metabolic acidosis associated with CKD. In addition, analysis of serum data for sodium, potassium, and calcium showed no change of these serum cations during the animal studies. Serum magnesium was not measured during the animal studies.

**Discussion**

Veverimer is an orally administered, nonabsorbed, counterion-free HCl binder under development for the treatment of metabolic acidosis in patients with CKD. The mechanism of action of veverimer relies on highly specific binding and retention of proton and chloride as the polymer traverses the GI tract (Fig. 7). In the first step of this process, veverimer becomes protonated after ingestion, with high-capacity proton binding occurring across the pH range (pH = 1.5–7.0) found in

**TABLE 4**

Recovery of administered [14C]-veverimer in excreta and carcasses of male and female Wistar Han rats

<table>
<thead>
<tr>
<th>Sample</th>
<th>% of Administered Dose</th>
<th>Males (n = 3)</th>
<th>Females (n = 3)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Mean</td>
<td>S.D.</td>
<td>Mean</td>
</tr>
<tr>
<td>Total feces</td>
<td>97.09</td>
<td>1.72</td>
<td>95.05</td>
</tr>
<tr>
<td>Total urine</td>
<td>0.02</td>
<td>0.01</td>
<td>0.02</td>
</tr>
<tr>
<td>Total expired aira</td>
<td>0.02</td>
<td>0.01</td>
<td>0.01</td>
</tr>
<tr>
<td>Cage washb</td>
<td>ND</td>
<td>NC</td>
<td>ND</td>
</tr>
<tr>
<td>Carcassb</td>
<td>ND</td>
<td>NC</td>
<td>ND</td>
</tr>
<tr>
<td>Total recovery</td>
<td>97.13</td>
<td>1.72</td>
<td>95.08</td>
</tr>
</tbody>
</table>

NC, not calculable; ND, radioactivity not detectable.
aUp to 72 h postdose, as this time sampling was discontinued as per plan because of radioactivity not being detectable in the 24–48 and 48–72-h collections.
bAt 168 h postdose only.
Mechanism of Action of Veverimer

Recovery of administered $^{14}$C-veverimer in excreta and carcasses of male and female Beagle dogs

<table>
<thead>
<tr>
<th>Sample</th>
<th>% of Administered Dose</th>
<th>Sample</th>
<th>% of Administered Dose</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Males ($n = 3$)</td>
<td>Females ($n = 3$)</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Mean</td>
<td>S.D.</td>
<td>Mean</td>
</tr>
<tr>
<td>Total feces</td>
<td>98.98</td>
<td>3.35</td>
<td>96.72</td>
</tr>
<tr>
<td>Total urine</td>
<td>0.02</td>
<td>0.01</td>
<td>0.02</td>
</tr>
<tr>
<td>Total cage wash</td>
<td>0.01</td>
<td>0.01</td>
<td>0.01</td>
</tr>
<tr>
<td>Total recovery</td>
<td>99.01</td>
<td>3.36</td>
<td>96.74</td>
</tr>
</tbody>
</table>

the GI lumen (Fig. 4; Table 3). The concurrent binding and retention of chloride by the protonated polymer facilitated by the highly crosslinked polymer backbone that restricts the binding of larger anions effectively removes HCl from the GI tract, which is then eliminated with the polymer in the feces (Figs. 5 and 6). HCl removal in this manner results in an elevation of serum bicarbonate (Figs. 5 and 6). In patients with CKD and metabolic acidosis, in which kidney-mediated acid excretion is reduced resulting in acid retention (Wesson et al., 2020), veverimer can restore an important acid excretion capacity and so is distinguished from other metabolic acidosis interventions that either reduce acid intake or neutralize systemic acidity.

The physicochemical analysis of veverimer along with the nonclinical pharmacology studies reported here illustrate various aspects of the proposed mechanism of action for veverimer. Veverimer is a free-flowing powder composed of highly crosslinked, low-swelling, spherical polymeric beads (Fig. 3). Particle size analysis showed a monomodal particle size distribution, wherein greater than 90% of the volume was composed of particles with a diameter $>$40 $\mu$m and an average diameter of approximately 100 $\mu$m. These size parameters were carefully selected to prevent systemic absorption from the GI tract while being small enough to optimize oral administration and tolerability (Bushinsky et al., 2018). The particle size distribution of the polymer beads is primarily controlled by the stir rate during the suspension polymerization in which the beads are first formed. Solubility tests in a variety of organic and aqueous solvents showed no significant solubility of the polymer. In vitro measurement of chloride binding in a range of intestinal mimics and pHs demonstrated that the extensive cross-linking promoted high capacity and selective HCl binding, particularly in the most complex matrix mimicking the lower GI tract, in which veverimer preferentially bound chloride (3.8 mmol/g polymer) in the presence of various competing anions, some of which provide precursors for bicarbonate production (Table 3). Furthermore, the retention of chloride, which was evaluated with GCTA that simulated the passage of the polymer through various compartments of the GI tract, showed approximately 60% of initially bound chloride (approximately 5 mmol/g polymer) was retained. The highly crosslinked structure of veverimer confers a marked size-exclusion selectivity to the negatively charged moieties that bind to the protonated polymer, strongly favoring binding of the smallest anions and restricting binding of larger anions. Based on the physicochemical characteristics of veverimer and the results from in vitro and human studies, veverimer is unlikely to have significant drug-drug interactions (Shao et al., 2020).

Proof-of-concept data for veverimer were provided by a rat model of adenine-induced CKD (Diwan et al., 2018) that is characterized by low serum bicarbonate ($<23.0$ mEq/l) and acidemia (pH $<$ 7.4). Dosing with veverimer initiated immediately after the 2-week adenine CKD-induction period or 7 weeks after initiation of adenine exposure at doses of 0.7–2.4 g/kg per day by dietary admixture resulted in a significant, dose-dependent increase in serum bicarbonate within 1 week, a dose-dependent increase in blood pH, and a significant, concomitant increase in fecal chloride excretion relative to untreated controls (Figs. 5 and 6). The pharmacological activity of veverimer—specifically maintenance of serum bicarbonate levels within the normal range—was shown to be durable through at least 6 weeks of dosing. Discontinuation of dosing was associated with a decrease in serum bicarbonate and pH values that were below the normal range for age-matched rats, confirming the treatment effect of veverimer.

The results from the rat model of CKD are consistent with clinical findings in patients with CKD and metabolic acidosis treated with veverimer (Bushinsky et al., 2018; Wesson et al., 2019a,b). The veverimer doses at which an effect on serum bicarbonate was observed in both phase 3 clinical trials (0.05–0.15 g/kg per day) were lower than the doses used in the nonclinical study (0.7–2.4 g/kg per day) reported here, but rats produce approximately 7.5-fold more acid per kg of body weight than humans (Toto and Alpern, 1996; Lin et al., 1998). Correcting the rat dose by this factor (7.5) results in a dose (0.1–0.32 g/kg per day) that is consistent with the pharmacologically active dose in humans. In addition, it is noted that the limited phosphate binding detected in the in vitro matrix experiments (Table 3) and the in vivo treatment experiments (Figs. 5C and 6C) did not translate into physiologic changes in serum or urinary phosphate in either animal studies or human clinical studies (Bushinsky et al., 2018; Wesson et al., 2019a,b).

Lack of veverimer absorption from the GI tract was demonstrated in both rats and dogs administered a single oral dose of $^{14}$C-veverimer. The mean bead size of approximately 100 $\mu$m with a narrow particle size distribution likely restricted absorption of particles from the GI tracts of the test animals, which was consistent with literature reports showing that particles larger than 0.5 $\mu$m are not systemically absorbed (Jung et al., 2000). Because radioactivity was not observed in the plasma of either species, metabolism was not evaluated. The lack of absorption in conjunction with the pharmacological effects on serum bicarbonate and fecal chloride excretion in rats supports that veverimer is not metabolized or degraded but maintains functional and, therefore, structural integrity during transit through the GI tract after oral administration.

Veverimer has a distinct mechanism of action that removes rather than neutralizes endogenous acid as a means of...
restoring acid-base balance in the body under the conditions of metabolic acidosis. As kidney function declines, generation of metabolic acids continues, but their excretion is compromised, leading to acid retention. Current treatment options to attenuate this acid retention include dietary interventions with very low-protein diets (Garneata et al., 2016), base-producing fruits and vegetables (Goraya et al., 2013, 2014, 2019), or oral alkali supplementation (de Brito-Ashurst et al., 2009; Phisitkul et al., 2010; Di Iorio et al., 2019; Dubey et al., 2020). Although dietary interventions are effective in elevating serum bicarbonate and reducing kidney function decline, adherence to dietary interventions can be difficult to achieve (Garneata et al., 2016) and necessarily involves the consumption of foods high in dietary potassium (Goraya and Wesson, 2019). Oral alkali supplementation introduces a significant amount of sodium into the body, along with the neutralizing bicarbonate or citrate equivalents, increasing overall sodium load in patients with CKD. Indeed, to raise serum bicarbonate by 3–4 mEq/l in an 80-kg patient, 6–8 g/day of sodium bicarbonate is required, delivering 1.7–2.2 g/day of sodium in addition to normal dietary sodium intake (Abramowitz et al., 2013), thus exceeding the published guidelines for daily sodium intake in patients with CKD (Wright and Cavanaugh, 2010). This additional sodium has the potential to aggravate sodium-sensitive comorbidities, such as hypertension, congestive heart failure, and edema, in patients with CKD (Bushinsky, 2019; Navaneethan et al., 2019). The additional sodium load from sodium-based oral alkali may also diminish the effect of renoprotective agents, such as renin-angiotensin-aldosterone system inhibitors (Lambers Heerspink et al., 2012; Veger et al., 2012), and lead to worsening albuminuria (Raphael et al., 2020), thereby counteracting the benefits of treating metabolic acidosis.

To avoid these challenges, veverimer removes acid through the GI tract, thus reducing the overall acid accumulation that is the hallmark of chronic metabolic acidosis, and in so doing introduces bicarbonate into the circulation without the addition of deleterious counterions. This mechanism of acid removal has similarities to the physiologic response to prolonged vomiting, chloride-losing diarrhea, or nasogastric suction, each of which removes HCl from the stomach and leads to a rapid increase in serum bicarbonate levels that may result in hypochloremic metabolic alkalosis (Kassirer and Schwartz, 1966; Niv and Fraser, 2002; Khanna and Kurtzman, 2006; Valeur and Julsrud, 2013). It is fundamentally different from these responses in persons with normal kidney function.

![Fig. 7. Mechanism of action of veverimer in the gastrointestinal tract.](image-url)

The mechanism of action of veverimer is illustrated in the figure. Veverimer is administered orally as a non-absorbed, free-amine polymer. Within the parietal cell, HCl is converted into H+ and HCO₃⁻ under the influence of carbonic anhydrase (Yao and Forte, 2003). H+ is secreted from the gastric parietal cell into the GI lumen in exchange for K+ through the action of the H+/K⁺ ATPase (Shin et al., 2009). Chloride and potassium ions within the parietal cell are transported into the gastric lumen via apical membrane conductance channels (potassium voltage-gated channel subfamily E member 2 and potassium voltage-gated channel subfamily Q member 1 K⁺ channels, and chloride voltage-gated channel 2 Cl⁻ channels) (Heitzmann and Warth, 2007), so that the exchange of K⁺ and H⁺ effectively recycles the potassium cation. Veverimer binds H⁺ with high capacity in the GI lumen, and every bound H⁺ results in an HCO₃⁻ entering the blood via basolateral exchange with serum Cl⁻ (Petrovic et al., 2003). Veverimer binds Cl⁻ with high selectivity in the GI lumen, excluding larger organic anions from the protonated polymer. HCl-bound veverimer is excreted in the feces, and the resulting increase in HCO₃⁻ in the blood leads to a correction of metabolic acidosis.
however, in that veverimer only binds HCl and does not induce the loss of fluid, sodium, and potassium that are hallmarks of hypochloremic metabolic alkalosis (Khanha and Kurtzman, 2006).

The selective binding and removal of HCl by veverimer allows for the excretion of acid while generating bicarbonate in the blood by the mechanisms described in the process of parietal cell HCl secretion (summarized in Fig. 7). In the process of facilitating the excretion of HCl through the gut in this manner, veverimer can be seen as supplementing and replacing the deficiency of acid excretion in the diseased kidney, thus extending the known structural and functional aspects of the gut epithelium that often mimic epithelial cells of the kidney nephrons (Coudrier et al., 1988; Schultheis et al., 1998; Wang et al., 2002; Alper and Sharma, 2013; Soleimani, 2013).

In conclusion, the studies reported here demonstrate that the chemical structure and design of veverimer are translated into a potent and selective binding of HCl, as measured with in vitro matrices mimicking various aspects of the GI tract and in animal studies that measure chloride removal through the feces with a polymer that is nonabsorbable and stable throughout GI transit. The physiologic consequence of specific HCl removal is the elevation of serum bicarbonate. The polymer does not introduce counterions nor does it remove organic anions in the GI tract that provide sources of bicarbonate precursors once absorbed. Veverimer treatment in an animal model of CKD with metabolic acidosis demonstrated that HCl binding and removal by the polymer increased serum bicarbonate, which was consistent with completed studies in human subjects that demonstrate the potential for veverimer to treat chronic metabolic acidosis in patients with CKD.

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**Contributed new reagents or analytic tools:** Biyani, Kade, Kiersted, Gbur, Tabakman, Nguyen.

**Performed data analysis:** Klaerner, Shao, Biyani, Kade, Kiersted, Gbur, Tabakman, Nguyen, Buyssse.

**Wrote or contributed to the writing of the manuscript:** Klaerner, Shao, Biyani, Kiersted, Gbur, Tabakman, Nguyen, Buyssse.

References


Waldapfel (Tricida, Inc.) in the course of the work and in the preparation of this report.


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