Submarine groundwater discharge is an important net source of light and middle REEs to coastal waters of the Indian River Lagoon, Florida, USA

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Abstract

Porewater (i.e., groundwater) samples were collected from multi-level piezometers across the freshwater–saltwater seepage face within the Indian River Lagoon subterranean estuary along Florida’s (USA) Atlantic coast for analysis of the rare earth elements (REE). Surface water samples for REE analysis were also collected from the water column of the Indian River Lagoon as well as two local rivers (Eau Gallie River, Crane Creek) that flow into the lagoon within the study area. Concentrations of REEs in porewaters from the subterranean estuary are 10–100 times higher than typical seawater values (e.g., Nd ranges from 217 to 2409 pmol kg$^{-1}$), with submarine groundwater discharge (SGD) at the freshwater–saltwater seepage face exhibiting the highest REE concentrations. The elevated REE concentrations for SGD at the seepage face are too high to be the result of simple, binary mixing between a seawater end-member and local terrestrial SGD. Instead, the high REE concentrations indicate that geochemical reactions occurring within the subterranean estuary contribute substantially to the REE cycle. A simple mass balance model is used to investigate the cycling of REEs in the Indian River Lagoon and its underlying subterranean estuary. Mass balance modeling reveals that the Indian River Lagoon is approximately at steady-state with respect to the REE fluxes into and out of the lagoon. However, the subterranean estuary is not at steady-state with respect to the REE fluxes. Specifically, the model suggests that the SGD Nd flux, for example, exported from the subterranean estuary to the overlying lagoon waters exceeds the combined input to the subterranean estuary from terrestrial SGD and recirculating marine SGD by, on average, ~100 mmol day$^{-1}$. The mass balance model also reveals that the subterranean estuary is a net source of light REEs (LREE) and middle REEs (MREE) to the overlying lagoon waters, but acts as a sink for the heavy REEs (HREE). Geochemical modeling and statistical analysis further suggests that this fractionation occurs, in part, due to the coupling between REE cycling and iron redox cycling within the Indian River Lagoon subterranean estuary. The net SGD flux of Nd to the Indian River Lagoon is ~7-fold larger than the local effective river flux to these coastal waters. This previously unrecognized source of Nd to the coastal ocean could conceivably be important to the global oceanic Nd budget, and help to resolve the oceanic “Nd paradox” by accounting for a substantial fraction of the hypothesized missing Nd flux to the ocean.

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1. INTRODUCTION

Understanding the sources and sinks of trace elements such as the rare earth elements (REE) in the oceans has important implications for quantifying their global geochemical cycles, their application as palaeoceanographic tracers, and in discerning the geochemical reactions that mobilize, sequester, and fractionate REEs in the environment. This is especially important for neodymium (Nd) as radiogenic Nd isotopes are commonly used in palaeoceanographic investigations over glacial-interglacial to million year time scales (e.g., von Blanckenburg, 1999; Rutberg et al., 2000; Frank, 2002; Scher and Martin, 2004; Piotrowski et al., 2005; Frank et al., 2006; Foster et al., 2007; MacLeod et al., 2008; Palinke et al., 2008).

However, our ability to confidently use Nd isotopes to trace oceanic water mass circulation in the modern ocean as well as in palaeoceanographic studies is hampered, in part, by the relatively poor constraints on the sources and sinks of Nd to and from the global ocean (von Blanckenburg, 1999; Lacan and Jeandel, 2001; Frank, 2002; Goldstein and Hemming, 2003; Piotrowski et al., 2008; also see Johannesson and Burdige, 2007, for a general review of this problem). Related to this is the so-called “Nd paradox” (Bertram and Elderfield, 1993; Jeandel et al., 1995, 1998; Tachikawa et al., 1997, 1999a,b; Lacan and Jeandel, 2001; Goldstein and Hemming, 2003; Siddall et al., 2008; Arrouse et al., 2009), which is the apparent decoupling of dissolved Nd concentrations (nutrient-like water column profiles suggestive of vertical cycling and relatively long, \(10^8\) years, oceanic residence times) and Nd isotope ratios (inter- and intra-ocean differences, which presumably exclude vertical cycling and supports an oceanic residence time of Nd that must be similar to, or less than the \(500-1500\) years oceanic mixing time).

More specifically, estimates of the atmospheric and riverine fluxes, which are considered the chief sources of Nd to the ocean, are insufficient by roughly a factor of 10 to balance the oceanic Nd budget and preserve the observed inter- and intra-oceanic Nd isotope values. The difference, which amounts to \(5.6 \times 10^{-9}\) moles of Nd per year, is known as the “missing Nd flux” and has been the subject of substantial speculation (Tachikawa et al., 2003; Lacan and Jeandel, 2005a; Arrouse et al., 2009). However, other than the broadly defined “boundary exchange” occurring along continental margins (e.g., Lacan and Jeandel, 2005b), none of the other specific processes discussed in the literature – redissolution of particulate- or sediment-associated Nd in mid- to high-salinity regions of estuaries (Sholkovitz and Szymczak, 2000), diffusive fluxes of porewater REEs to the overlying water column (Elderfield and Sholkovitz, 1987), or dissolution of atmospheric particle derived Nd (i.e., the atmospheric Nd flux) in the oceans (Zhang et al., 2008; Arrouse et al., 2009) – appear to have the proper magnitude and Nd isotope values to balance the ocean Nd budget.

Using average Nd concentrations and isotope values for groundwater from the literature, and an estimate of the global, terrestrial submarine groundwater discharge (SGD) flux, Johannesson and Burdige (2007) speculated that SGD may constitute an important component of this missing Nd flux. However, Johannesson and Burdige (2007) did not measure Nd in actual SGD samples and only hypothesized that the marine component of the SGD Nd flux may be considerable. Submarine groundwater discharge includes all flow of water from the seafloor into the overlying marine water column within continental margin settings notwithstanding the physical driving mechanism and chemical composition (Church, 1996; Burnett et al., 2003; Moore, 2010). Consequently, SGD consist of both “fresh submarine groundwater discharge” originating from meteoric recharge of terrestrial aquifers and “recirculated saline submarine groundwater discharge” (i.e., marine waters) that cycles through coastal aquifers due to processes such as tidal pumping, wave set-up, bioirrigation, and/or geothermal or density gradients (Taniuchi et al., 2002). Hereafter, these components of SGD are referred to as “terrestrial SGD” and “marine SGD”, respectively, and the combination of the two as “total SGD” (Martin et al., 2007).

A number of previous studies have shown that SGD represents a significant flux of water, nutrients, carbon, and metals to the oceans (Moore, 1996, 1997, 2010; Cai et al., 2003; Charette and Buesseler, 2004; Slomp and Van Cappellen, 2004; Kim et al., 2005; Windom et al., 2006; Charette, 2007; Moore et al., 2008). In this contribution, we present REE concentration data for porewaters (i.e., groundwaters) and surface waters of the Indian River Lagoon system, and show using a simple mass balance approach that, to a first approximation, total SGD is a substantial net source of light REEs (LREE) and middle REEs (MREE) to these coastal waters.

2. STUDY SITE

The field site is located along the Indian River Lagoon near the city of Melbourne, Florida (Fig. 1). The Indian River Lagoon system occupies a large portion of Florida’s Atlantic coast, extending for 250 km, and includes the Mosquito and Banana River Lagoons to the north. The lagoon is characterized as microtidal as the amplitudes of tides are typically less than 10 cm. Nonetheless, the shallow water column (mean depth of 1.5 m, with a maximum depth of 5 m) is generally well mixed owing to changes in water level largely driven by wind.

Pleistocene beach ridges surround the Indian River Lagoon in the study site. The lagoon is underlain by the Holocene Anastasia Formation, which makes up the Surficial Aquifer in the study area. The Anastasia Formation consists of quartz sand that is interbedded with coquina, along with minor amounts of silt, clay, and carbonate rock fragments (Martin et al., 2007). Porosity of the Surficial Aquifer within the Indian River Lagoon subterranean estuary (Moore, 1999) ranges from 37% to 45%, whereas the hydraulic conductivity ranges between \(10^{-2}\) and \(10^{-5}\) m s\(^{-1}\) within the upper 300 cm, but exhibits a more uniform range of \(10^{-4}-10^{-8}\) m s\(^{-1}\) in the top 70 cm (Martin et al., 2004; Hartl, 2006; Smith et al., 2008a; Roy et al., 2010). The Surficial Aquifer is separated from the underlying Floridan Aquifer system (Paleocene to Oligocene...
limestones) by approximately 30 m of the confining Hawthorne Group (Miocene sand, marl, clay, limestone, dolostone, and phosphatic materials; Miller, 1986). The thickness of the Hawthorne Group rocks in the study region indicates that essentially all of the terrestrial SGD to the Indian River Lagoon subterranean estuary originates within the Surficial Aquifer (Martin et al., 2007). Our study specifically focuses on a portion of the Indian River Lagoon that is located 4.5 km north of Melbourne, Florida, and 0.5 km south of Eau Gallie, Florida (Fig. 1b and c). Freshwater is delivered to the studied portion of the Indian River Lagoon via direct precipitation, urban runoff, discharge of the Eau Gallie River and Crane Creek, and terrestrial SGD (Martin et al., 2007). Fluxes of SGD were previously determined at the study site using a combination of methods that included seepage meters, chemical and thermal tracers, and models of chemical profiles (excess $^{222}$Rn, Ra isotope decay, Cl concentrations; Cable et al., 2004, 2006; Martin et al., 2004, 2006, 2007; Smith et al., 2008a,b; Roy et al., 2010). The flux of terrestrial SGD decreases with distance offshore becoming negligible near the freshwater–saltwater seepage face (Fig. 2; Smith et al., 2008a; Roy et al., 2010). Further offshore beyond the freshwater–saltwater seepage face (e.g., in the vicinity of CIRL-39, 250 m offshore), bioirrigation results in rapid, daily exchange of lagoon waters with porewaters within the upper 70 cm of the sediments, leading to flow rates as large as 150 cm day$^{-1}$ (Martin et al., 2004, 2006). Consequently, marine SGD is chiefly Indian River Lagoon water that is cycled through the lagoon bottom sediments by bioirrigation, whereas the total SGD consists of both a relatively small component of terrestrial SGD and this substantially larger component (~2 orders of magnitude larger) of marine SGD (Cable et al., 2004; Martin et al., 2004, 2006, 2007; Fig. 2). Exchange of Indian River Lagoon waters with the ocean occur via three inlets located within the southern part of the lagoon. Consequently, the residence time of water in the Indian River Lagoon can be as much as a year in the far north, but is generally on the order of 18 days in the vicinity of the study site (Smith, 1993).
Sediments within the Indian River Lagoon subterranean estuary are described in detail in Hartl (2006) and Roy et al. (2010). Four sediment cores recovered in the vicinity of CIRL-39 (Figs. 1 and 2) consisted of 2 m of black (sulfide-rich), marine sediments overlying orange, Fe(III) oxide/oxyhydroxide coated quartz sands of terrestrial origin (Hartl, 2006). The thickness of the orange sediments is currently unknown as they extended to the base of these sediment cores. Roy et al. (2010) report similar lithostratigraphy for 6 cores collected along a near-shore transect shown in Fig. 1c. Specifically, these sediment cores were collected approximately 1 m north of sites EGN-0, EGN-10, EGN-22.5, EGN-30, and CIRL-39 (Fig. 1c; Roy et al., 2010). The thickness of the black, sulfide-rich marine sediments is ~17 cm at the shoreline (EGN-0), but increases in thickness with distance from shore, reaching 68 cm at EGN-30, and more than 250 cm at CIRL-39 (Hartl, 2006; Roy et al., 2010). Because sedimentation rates vary little within the Indian River Lagoon (Hartl, 2006), the increase in the thickness of the sulfide-rich marine sediments with distance offshore is interpreted to be the result of sea-level rise and the consequent increase in Fe–sulfide accumulation times accommodated by progressive seawater inundation (Roy et al., 2010).

3. METHODS

3.1. Sample collection

Methods employed to determine volumetric fluxes of SGD that are used here in the mass balance model were previously described (Cable et al., 2004; Martin et al., 2004, 2006, 2007). Porewater (i.e., groundwater) samples for REE analysis were collected in April 2007 from three multilevel piezometers (i.e., multisamplers; Martin et al., 2003) along a near-shore transect that extends 30 m from...
the Indian River Lagoon mainland shore (i.e., 0 m; Fig. 1c) by pumping water from the different multisampler ports into an overflow cup using a peristaltic pump (Martin et al., 2007; Roy et al., 2010). Before porewater samples were collected, the specific conductivity, dissolved oxygen, temperature, and salinity of pumped waters was monitored until these parameters stabilized. In each case, less than 2 L was pumped in order to minimize the possibility of induced vertical flow during sampling. Once these parameters stabilized, porewater (~60–100 mL) was collected (via pumping) and subsequently filtered through 0.45 μm in-line (GEO-TECH, polyether sulfone membrane) filter cartridges directly into precleaned (acid washed) HDPE sample bottles. Sample bottles were rinsed three times with the filtered porewater before filling. The filtered porewater samples were immediately acidified with ultra-pure HNO₃ (Seastar Chemicals, Baseline, sub-boiling, distilled in quartz) and subsequently filtered through 0.45 μm in-line (GEO-TECH, polyether sulfone membrane) filter cartridges directly into precleaned (acid washed) HDPE sample bottles. Sample bottles confirmed with a series of REE calibration standards of known concentrations (1, 2, 5, 10, 100, 500, and 12 μg L⁻¹). In addition to the REEs, solutions were analyzed for 115In as an internal standard and for 56Fe (Roy et al., 2010) to separate the REEs from the terrestrial component of SGD and the inflow streams. The REEs were then eluted from the columns using 8 M ultra-pure HNO₃ (Seastar Chemicals, Baseline, sub-boiling, distilled in quartz) and collected in Teflon® beakers. The eluant was subsequently taken to dryness on a hot-plate, and the residue then redissolved in 10 mL of a 1% v/v ultra-pure HNO₃ solution. Each 10 mL sample was then spiked with 135La and 137Cs as an internal standard to the measured REE values for SLEW-3 reported by Lawrence and Kamber (2007). Analytical precision of the REE analyses was always better than 4% relative standard deviation (RSD), and generally better than 2% RSD.

### 3.2. Sample analysis

Major solutes (Ca²⁺, Mg²⁺, Na⁺, K⁺, Cl⁻, SO₄²⁻) were determined in waters from the Indian River Lagoon system by ion chromatography (Dionex DX500) at the University of Florida. Alkalinity was determined by standard titration techniques. Chloride was measured in the Eau Gallie River of Florida. Alkalinity was determined by standard titration by ion chromatography (Dionex DX500) at the University of Florida by monitoring the Eu isotopes and the heavy REEs (HREE) where also additional check for potential interferences. In addition, the Eu isotopes and the heavy REEs (HREE) were also monitored in high-resolution mode, which allowed us to resolve interferences on 151Eu and 153Eu from BaO⁺ species formed in the plasma stream, and LREEO⁺ and middle REEO⁺ (MREEO⁺) species on the HREEs. The HR-ICP-MS was calibrated and the concentrations of REEs in the samples confirmed with a series of REE calibration standards of known concentrations (1, 2, 5, 10, 100, 500, and 1000 ng kg⁻¹). The calibration standards were prepared from NIST traceable High Purity Standards (Charleston, SC). In addition, check standards for the REEs were prepared using Perkin–Elmer multi-element solutions. The National Research Council Canada (Ottawa, Ontario, Canada) Standard Reference Material (SRM) for estuarine waters (SLEW-3) was analyzed to check for accuracy by comparison with the measured REE values for SLEW-3 reported by Lawrence and Kamber (2007). Analytical precision of the REE analyses was always better than 4% relative standard deviation (RSD), and generally better than 2% RSD.

### 3.3. REE solution complexation modeling

Waters from the Indian River Lagoon system range in concentration from dilute freshwaters that characterize the terrestrial component of SGD and the inflow streams to the saline waters of the lagoon itself (salinity of lagoon water near our field site ranges from 22 to 29; Martin et al., 2007). Consequently, solution complexation of the REEs in waters from the Indian River Lagoon system was evaluated using a combined specific ion interaction and ion pairing model initially developed for the REEs by Millero (1992). The model links the specific ion interaction approach (e.g., Pitzer, 1979) with and ion pairing model (Garrels and Thompson, 1962; Millero and Schreiber, 1982), thus allowing for the evaluation of REE complexation with inorganic ligands in dilute to highly saline natural waters (Johannesson and Lyons, 1994; Johannesson et al., 1996a,b). The model was updated by adding the most recently determined stability constants for REEs complexation with inorganic ligands (e.g., Lee and Byrne, 1992; Schijf and Byrne, 1999, 2004; Klinkness and Byrne, 2000;
Luo and Byrne, 2001, 2004). Free concentrations of inorganic ligands (e.g., $\left[CO_2\right]_v$, $\left[S0_2\right]_v$, $\left[F\right]_v$) used in the REE complexation modeling were computed from the major solute composition of Indian River Lagoon waters via the SpecE8 program of the Geochemist’s Workbench (release 7.0; Bethke, 2008) using the thermodynamic data base from PHRQPITZ (Plummer et al., 1989), and following the approach outlined by Millero and Schreiber (1982).

3.4. Mass balance model

Mass balance calculations were performed using a standard box model approach, which assumes steady-state flow conditions and includes separate boxes (i.e., reservoirs) for: (1) the water column portion of the Indian River Lagoon system; (2) the subterranean estuary beneath the lagoon; and (3) the coastal ocean (Fig. 3). The section of the Indian River Lagoon considered in the box model is shown in Fig. 1b, and includes the portion of the Indian River Lagoon between the Eau Gallie River and Crane Creek. Because the distance between the Eau Gallie River and Crane Creek is 5 km, the width of the Indian River Lagoon in the study region (i.e., distance from the mainland to the barrier island) is 1.8 km, and the average depth of the lagoon is 1.5 m, the volume of the lagoon considered in our model is $1.35 \times 10^7$ m$^3$.

The box model was initially employed to evaluate the hydrologic budget and Cl mass balance for the studied portion of the Indian River Lagoon. To examine the hydrologic budget we used previous estimates of terrestrial and marine SGD (Martin et al., 2007) and stream gauge data (U.S. Geological Survey) for the Eau Gallie River and Crane Creek. The total SGD to the studied portion of the Indian River Lagoon was computed by summing the previously determined estimates of terrestrial and marine SGD. The volume of seawater exchange between the lagoon and the coastal ocean was estimated by assuming that Cl is conservative and that the hydrologic budget is at steady-state. The salinity of the coastal ocean end-member was assumed to be 35‰, which is consistent with seawater off Florida’s east coast (e.g., Smith, 1993; Swarzenski et al., 2006). The mass balance model was subsequently employed to investigate the cycling of the REEs within the studied portion of the Indian River Lagoon. A major objective was to evaluate whether geochemical reactions occurring in the subterranean estuary beneath the Indian River Lagoon influence the SGD flux of REEs to the overlying lagoon waters (e.g., see Duncan and Shaw, 2003). It is important to note, however, that the relatively simple box modeling approach used here only represents a first attempt to quantify REE fluxes in the Indian River Lagoon system. Additional sampling and data collection will no doubt help to constrain the model (e.g., see Roy et al., 2010).

4. RESULTS

4.1. Rare earth element concentrations

Rare earth element concentrations for waters from the Indian River Lagoon system waters are presented in Table 1, which also summarizes the shale-normalized Yb/Nd ratios for these waters, as well as presents mean REE concentrations for seawater taken from the literature. Concentrations of REEs in surface waters within the Indian River Lagoon (i.e., water column) decrease with increasing distance offshore. For example, at a distance of 5 m offshore, the Nd concentration of the lagoon water is 638 pmol kg$^{-1}$, whereas at a distance of 250 m from shore (i.e., water column station at CIRL-39) the Nd concentration is 164 pmol kg$^{-1}$. We note, however, that the general decrease in REE concentration with increasing distance from shore is punctuated by at least two local concentration maxima at 17.5 and 22.5 m offshore, where Nd concentrations are 389 and 354 pmol kg$^{-1}$, respectively (Table 1).

The same general decrease in REE concentration with increasing distance from shore is also observed for the terrestrial component of SGD, such that porewaters from multisampler EGN-0 exhibit higher REE concentrations.

![Fig. 3. Box model used to evaluate REE fluxes within the Indian River Lagoon system.](image_url)
Table 1
Rare earth element concentrations (in pmol kg\(^{-1}\)) in waters from the Indian River Lagoon system, and average seawater computed with data from the literature. For the SGD samples, cmbsf = centimeters below sea floor.

<table>
<thead>
<tr>
<th>Water Column</th>
<th>La</th>
<th>Ce</th>
<th>Pr</th>
<th>Nd</th>
<th>Sm</th>
<th>Eu</th>
<th>Gd</th>
<th>Tb</th>
<th>Dy</th>
<th>Ho</th>
<th>Er</th>
<th>Tm</th>
<th>Yb</th>
<th>Lu</th>
<th>((Yb/\text{Nd})_a)</th>
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<tbody>
<tr>
<td>EGN-5, WC</td>
<td>559</td>
<td>1047</td>
<td>151</td>
<td>638</td>
<td>121</td>
<td>34.8</td>
<td>127</td>
<td>23.8</td>
<td>181</td>
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<td>523</td>
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<td>369</td>
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<td>102</td>
<td>21.3</td>
<td>109</td>
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<td>EGN-15, WC</td>
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<td>71.4</td>
<td>21</td>
<td>91</td>
<td>17</td>
<td>114</td>
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<td>131</td>
<td>28.3</td>
<td>239</td>
<td>52.7</td>
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<td>EGN-17.5, WC</td>
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<td>8.01</td>
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<td>EGN-30, WC</td>
<td>141</td>
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<td>164</td>
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<td>7.13</td>
<td>42.2</td>
<td>8.31</td>
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Mean ± SD  
256 ± 51.3| 498 ± 97.9| 58.2 ± 15.6| 315 ± 58| 56.8 ± 11.4| 12.7 ± 4.28| 69 ± 10.6| 12.4 ± 2.03| 98.4 ± 16.2| 27 ± 7.5| 145 ± 55.2| 33.5 ± 15.3| 293 ± 158| 66.5 ± 36.1

Terrestrial SGD  
<table>
<thead>
<tr>
<th>Water Column</th>
<th>La</th>
<th>Ce</th>
<th>Pr</th>
<th>Nd</th>
<th>Sm</th>
<th>Eu</th>
<th>Gd</th>
<th>Tb</th>
<th>Dy</th>
<th>Ho</th>
<th>Er</th>
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<th>Yb</th>
<th>Lu</th>
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<tr>
<td>EGN-0, 15 cmbsf</td>
<td>167</td>
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<td>417</td>
<td>83</td>
<td>22.1</td>
<td>117</td>
<td>21.4</td>
<td>242</td>
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<td>141</td>
<td>312</td>
<td>26.6</td>
<td>227</td>
<td>35</td>
<td>3.73</td>
<td>41.1</td>
<td>8.13</td>
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<td>55.8</td>
<td>12.2</td>
<td>120</td>
<td>18.8</td>
<td>7.23</td>
</tr>
<tr>
<td>EGN-10, 115 cmbsf</td>
<td>137</td>
<td>307</td>
<td>27.5</td>
<td>217</td>
<td>28.1</td>
<td>3</td>
<td>30.2</td>
<td>5.36</td>
<td>47.9</td>
<td>9.06</td>
<td>42.7</td>
<td>9.71</td>
<td>84.4</td>
<td>14.5</td>
<td>5.32</td>
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</table>

Mean ± SD  
187 ± 41.2| 427 ± 84.5| 47.4 ± 12.3| 350 ± 59.6| 66.3 ± 16.5| 15.1 ± 5.02| 82 ± 21.1| 14.6 ± 3.89| 139 ± 36.8| 43.3 ± 16.2| 268 ± 109| 70.9 ± 31.2| 716 ± 343| 155 ± 82.9

Total SGD  
<table>
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<th>Water Column</th>
<th>La</th>
<th>Ce</th>
<th>Pr</th>
<th>Nd</th>
<th>Sm</th>
<th>Eu</th>
<th>Gd</th>
<th>Tb</th>
<th>Dy</th>
<th>Ho</th>
<th>Er</th>
<th>Tm</th>
<th>Yb</th>
<th>Lu</th>
<th>((Yb/\text{Nd})_a)</th>
</tr>
</thead>
<tbody>
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<td>EGN-10, 15 cmbsf</td>
<td>171</td>
<td>367</td>
<td>32.7</td>
<td>277</td>
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<td>70.2</td>
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<td>99.8</td>
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<td>170</td>
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<td>289</td>
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<td>258</td>
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<td>45.9</td>
<td>8.69</td>
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<td>12.3</td>
<td>4.96</td>
</tr>
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<td>523</td>
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<td>401</td>
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<td>99.1</td>
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<td>88.6</td>
<td>16.8</td>
<td>104</td>
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<td>1.85</td>
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<td>715</td>
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<td>28.4</td>
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<td>112</td>
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<td>135</td>
<td>22.3</td>
<td>188</td>
<td>24.8</td>
<td>3.6</td>
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<td>2879</td>
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<td>1703</td>
<td>361</td>
<td>2488</td>
<td>434</td>
<td>14.2</td>
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</table>

Mean ± SD  
395 ± 110| 772 ± 237| 88.4 ± 31| 485 ± 109| 82.6 ± 17.3| 16.4 ± 4.37| 99.3 ± 17.6| 13.2 ± 1.84| 105 ± 11.4| 25.4 ± 4.42| 104 ± 16.3| 17.7 ± 2.2| 145 ± 19.1| 22.5 ± 3.76

* Rivers  
* Seawater  
* Mean ± SD  

---

a PAAS normalized ratio.
b Does not include the EGN-22, 5, 186 cmbsf sample (see text for discussion).
c Discharge-weighted mean and standard deviation.
than those from EGN-10 (Table 1). More specifically, Nd concentrations range from 417 to 568 pmol kg$^{-1}$ in porewaters collected from multisampler EGN-0 and from 217 to 277 pmol kg$^{-1}$ in porewaters from EGN-10. However, the highest REE concentrations in porewaters from the Indian River Lagoon subterranean estuary occur at the freshwater–saltwater seepage face where terrestrial SGD mixes with marine SGD forming total SGD (Table 1). Here, porewater Nd concentrations reach values as high as 2409 pmol kg$^{-1}$.

Based on Cl$^{-}$ concentrations and the position of the freshwater–saltwater seepage face (Tables 1 and 2; Fig. 2), porewaters from multisamplers EGN-0 and EGN-10 (i.e., 25, 55, and 115 cmbsf, where embfs = centimeters below sea floor) are employed in the mass balance model to represent the terrestrial SGD component, whereas porewaters from EGN-22.5, and the 15 cmbsf depth from EGN-10, reflect the total SGD (i.e., terrestrial SGD + marine SGD; Martin et al., 2007; Roy et al., 2010) in the model. Thus, Nd concentrations for terrestrial SGD range from 217 to 568 pmol kg$^{-1}$, exhibiting a mean (±SD) of 350 ± 59.6 pmol kg$^{-1}$, whereas for total SGD, Nd ranges from 258 to 2409 pmol kg$^{-1}$ with a mean (±SD) of 806 ± 333 pmol kg$^{-1}$. The deepest SGD sample from EGN-22.5 is the presumed outlier as its Nd value is more than a factor of 3 greater than the next highest Nd concentration (i.e., 773 pmol kg$^{-1}$ for EGN-22.5, 106 cmbsf; Table 1). Thus, the mean (±SD) Nd concentration of total SGD (i.e., 485 ± 109 pmol kg$^{-1}$; Table 1) that we employ in our mass balance calculations excludes this presumed outlier.

Shale-normalized REE plots for porewaters and surface waters from the Indian River Lagoon system are all enriched in HREEs relative to LREEs (Fig. 4). Excluding the presumed outlier (EGN-22.5, 186 cmbsf), porewater and lagoon water nearest to shore (i.e., multisampler EGN-0, water column sample EGN-5, WC) exhibit the highest HREE enrichments (mean ± SD (Yb/Nd)$_{Shale}$ = 58 ± 12; Table 1; Fig. 4). By comparison, shale-normalized Yb/Nd ratios for seawater are typically around 4 (Table 1). The shale-normalized REE patterns further demonstrate that porewater within the Indian River Lagoon subterranean estuary consists of both the terrestrial component of SGD (e.g., Fig. 4b and c), as well as recirculated Indian River Lagoon water (Fig. 4a).

4.2. REE solution complexation

The results of the solution complexation modeling for the REEs in waters of the Indian River Lagoon system are shown in Fig. 5. The average major solute compositions of terrestrial SGD, total SGD, and the lagoon water column (Table 2) were used in the model calculations. The model predicts that the REEs chiefly occur in solution within Indian River Lagoon waters complexed with carbonate ions (Fig. 5). For example, the percentage of each REE complexed with carbonate ions (i.e., LnCO$_3^-$ + Ln(CO$_3$)$_2^{2-}$, where Ln represents the individual REE) ranges from 76% for La to 98% for Lu in terrestrial SGD, from 58% for La to 97% for Lu in total SGD, and from 89% for La to 99% for Lu in the lagoon water column. Carbonato complexes (LnCO$_3^-$) predominate in the neutral pH, terrestrial SGD accounting for 60–80% of each REE in solution, whereas in the higher pH water column samples, the dicarbonato complex (Ln(CO$_3$)$_2^{2-}$) is the principal dissolved form for all REEs except La (Fig. 5). The model predicts that REE carbonate complexes also predominate in the total SGD, except for Yb and Lu, which occur chiefly as carbonato complexes (Fig. 5).

4.3. Hydrologic and chloride mass balance model

United States Geological Survey stream gauge data for the Eau Gallie River (USGS 02249007) between January 1991 and September 2004 (average 3.46 × 10$^8$ m$^3$ day$^{-1}$; n = 4555) and Crane Creek (USGS 02249518) between January 1987 and September 2004 (average 1.04 × 10$^8$ m$^3$ day$^{-1}$; n = 6377) provides an estimate of the mean discharge of the two rivers ($Q_{river} = 1.39 \times 10^8$ m$^3$ day$^{-1}$; Table 3) into the studied portion of the Indian River Lagoon. For the terrestrial and marine components of SGD to the Indian River Lagoon, the estimates of Martin et al. (2007) of 1.15 and 117 m$^3$ day$^{-1}$ per meter of shoreline, respectively, are employed. The terrestrial and marine SGD fluxes (i.e., $Q_{terrestrial SGD}$, $Q_{marine SGD}$) are recalculated for the entire 5 km seepage front between the Eau Gallie River and Crane Creek giving 5.75 × 10$^8$ and 5.85 × 10$^8$ m$^3$ day$^{-1}$, respectively (Table 3). Hence, total SGD to the Indian River Lagoon (i.e., $Q_{total SGD} = Q_{terrestrial SGD} + Q_{marine SGD}$) exceeds river discharge by a factor of four. Assuming that Cl is conservative, the Cl mass balance for the Indian River Lagoon subterranean estuary can be written as:

\[
[Cl^-]_{terrestrial SGD} + [Cl^-]_{lagoon} = [Cl^-]_{total SGD} \quad (1)
\]

where $[Cl^-]_{terrestrial SGD}$ and $[Cl^-]_{lagoon}$ are the mean Cl concentrations of terrestrial SGD and the Indian River Lagoon water column, respectively (Table 3; Fig. 3). The Cl concentration of total SGD, $[Cl^-]_{total SGD}$, computed by solving Eq. (1) (i.e., 335 mmol kg$^{-1}$) is identical to the mean (±SD) of the Cl concentrations measured in the lagoon water column (338 ± 15 mmol kg$^{-1}$; Table 3), as well as measured in porewater samples from multisamplers EGN-30 and CIRL-39 (335 ± 8.55 mmol kg$^{-1}$; data not shown). These observations further support that recirculating lagoon water is the principal source of SGD within the Indian River Lagoon subterranean estuary.

Eq. (1) can be recast into terms of the Cl fluxes as:

\[
J_{Cl}^{terrestrial SGD} + J_{Cl}^{marine SGD} = J_{Cl}^{total SGD} \quad (2)
\]

where, for example, $J_{Cl}^{terrestrial SGD} = [Cl^-]_{terrestrial SGD}Q_{terrestrial SGD}$. The respective Cl fluxes are listed in Table 3 for terrestrial SGD, total SGD, and marine SGD (i.e., recirculated lagoon water). The computed Cl fluxes also indicate that the terrestrial component of SGD has a negligible impact on the salt budget in the subterranean estuary beneath the Indian River Lagoon because $J_{Cl}^{terrestrial SGD}$ is almost four orders of magnitude smaller than $J_{Cl}^{marine SGD}$. Hence, the amount of Cl entering the bottom sediments of the Indian River Lagoon as marine SGD (i.e., recirculating lagoon water; 198(±8.78) × 10$^8$ mol day$^{-1}$) is essentially equal to that leaving the subterranean estuary as total SGD (198(±8.8) × 10$^8$ mol day$^{-1}$).
Table 2
Geochemical composition of waters from the Indian River Lagoon system. Major solute compositions are in mmol kg\(^{-1}\), and dissolved oxygen (DO), Fe, and sulfide concentrations are in l mol kg\(^{-1}\). Ionic strength is reported in mol kg\(^{-1}\).

<table>
<thead>
<tr>
<th>Water column</th>
<th>pH</th>
<th>Temp. (°C)</th>
<th>DO (l mol kg(^{-1}))</th>
<th>Ca (mmol kg(^{-1}))</th>
<th>Mg (mmol kg(^{-1}))</th>
<th>Na (mmol kg(^{-1}))</th>
<th>K (mmol kg(^{-1}))</th>
<th>Cl (mmol kg(^{-1}))</th>
<th>SO(_4) (mmol kg(^{-1}))</th>
<th>Alk (mmol kg(^{-1}))</th>
<th>Fe (l mol kg(^{-1}))</th>
<th>S(-II) (l mol kg(^{-1}))</th>
<th>Ionic strength (mol kg(^{-1}))</th>
<th>SIb (goethite)</th>
</tr>
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<tbody>
<tr>
<td>EGN-5, WC</td>
<td>8.63</td>
<td>28.9</td>
<td>337</td>
<td>6.47</td>
<td>204</td>
<td>4.46</td>
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<td>17.3</td>
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<td>0.376</td>
<td>8</td>
<td>0.267</td>
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<td>3.17</td>
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<td>0.376</td>
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<td>Mean ± SD(a)</td>
<td>8.41 ± 0.06</td>
<td>23.6 ± 1.6</td>
<td>329 ± 23.8</td>
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<td>22.5</td>
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<td>1.72 ± 0.34</td>
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<td>26.5</td>
<td>14.1</td>
<td>10.5</td>
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<td>11.9</td>
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<tr>
<td>Mean ± SD(a)</td>
<td>7.19 ± 0.05</td>
<td>24.2 ± 0.69</td>
<td>38.9 ± 11.9</td>
<td>10.6 ± 1.26</td>
<td>200 ± 27.4</td>
<td>3.45 ± 0.66</td>
<td>335 ± 149</td>
<td>11.7 ± 1.64</td>
<td>3.87 ± 0.27</td>
<td>127 ± 55.5</td>
<td>162 ± 74.6</td>
<td>0.266 ± 0.031</td>
<td>-0.48 ± 0.001</td>
<td></td>
</tr>
<tr>
<td>Rivers</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Eau Gallie River</td>
<td>7.73</td>
<td>231</td>
<td>9.97</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Crane Creek</td>
<td>7.62</td>
<td>224</td>
<td>10.7</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

\(a\) Mean values were used in the REE solution complexation model.

\(b\) SI = saturation index (log \(IAP/K_{sp}\)) computed with the React program of the Geochemist's Workbench\(^{a}\) (version 7; Bethke, 2008).

\(c\) Calculated using Eq. (1).
Because the discharge weighted mean Cl concentration for the two rivers entering the studied portion of the Indian River Lagoon is more than 30 times lower than the average Cl concentration of the lagoon water (Table 3), most of the Cl within the Indian River Lagoon comes from exchange with coastal seawater. Seawater exchange between the coastal ocean and the Indian River Lagoon is estimated using Cl and water mass balances for the Indian River Lagoon (Fig. 3). Chloride mass balance for the lagoon is expressed as:

$$Q_{\text{seawater}} = \frac{\left[ Cl \right]_{\text{lagoon}} (Q_{\text{RL}} + Q_{\text{rivers}} + Q_{\text{Terrestrial SGD}}) - \left[ Cl \right]_{\text{rivers}} Q_{\text{rivers}} - \left[ Cl \right]_{\text{Terrestrial SGD}} Q_{\text{Terrestrial SGD}}}{\left[ Cl \right]_{\text{seawater}} - \left[ Cl \right]_{\text{lagoon}}}.$$  \hspace{1cm} (7)

$$\left[ Cl \right]_{\text{Terrestrial SGD}} = \left[ Cl \right]_{\text{lagoon}} Q_{\text{Marine SGD}} + \left[ Cl \right]_{\text{lagoon}} Q_{\text{out}}$$  \hspace{1cm} (3)

where $\left[ Cl \right]_{\text{seawater}}$ is the Cl concentration of seawater (Table 1; Fig. 3), and $Q_{\text{seawater}}$ is the volumetric exchange of seawater with the studied portion of the Indian River Lagoon. Water mass balance in the Indian River Lagoon implies,

$$Q_{\text{rivers}} + Q_{\text{seawater}} + Q_{\text{Terrestrial SGD}} = Q_{\text{out}} + Q_{\text{Marine SGD}}.$$  \hspace{1cm} (4)

Using the values in Table 3 for mean chloride concentrations and the corresponding volumetric discharges, $Q_v$ we estimate that seawater exchange with the studied portion of the Indian River Lagoon is on the order of $2.24(\pm0.69) \times 10^5$ m$^3$ day$^{-1}$ (Fig. 3; Table 3). To evaluate whether this amount of seawater exchange is reasonable, we calculate the residence time of water and Cl in the studied portion of the Indian River Lagoon using our estimated volumetric discharges and Cl fluxes. For water, we obtain a

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Fig. 4. Shale-normalized REE plots of waters from the Indian River Lagoon system showing: (a) Indian River Lagoon water column, Eau Gallie River, and Crane Creek samples; (b) terrestrial component of SGD beneath the Indian River Lagoon; (c) the total SGD (terrestrial SGD + recirculated marine SGD) at the freshwater–saltwater seepage face; and (d) the mean seawater data (±1 SD) presented in Table 1. Note, the scale of the ordinate for the mean seawater sample differs from the Indian River Lagoon water samples by a factor of 100 lower. Rare earth elements concentrations for all water samples are normalized to the Post Archean Australian Shale (PAAS) composite (Nance and Taylor, 1976). See on-line version for colors.
residence time of 14.2 ± 1 days, whereas the estimated residence time for Cl is 14.2 ± 1.1 days. These residence times are identical, and similar to the 18 days flushing time for this portion of the Indian River Lagoon computed with a 1-D numerical model (Smith, 1993), which lends credibility to the applicability of the mass balance model presented here for the Indian River Lagoon system.

4.4. REE mass balance

Using the calculated, mean Nd concentrations measured in terrestrial SGD, total SGD, the Eau Gallie River and Crane Creek, and the lagoon water column as an example for the other REEs (Table 1), along with the volumetric water fluxes (Table 3), we calculate Nd fluxes for the studied portion of the Indian River Lagoon system (Table 4). It is important to note, however, that because our REE data are for a single point in time, concentrations and REE fluxes may vary seasonally. For example, residence times of fresh groundwater in the Surficial Aquifer range between 5 and 7 months, indicating that wet season subtropical precipitation is reflected in SGD a half year later (Smith et al., 2008a). Nonetheless, long-term head-level monitoring and SGD measurements demonstrate that in the absence of tropical cyclones during the wet season, terrestrial SGD from the Surficial Aquifers varies by less than a factor of 4 between the wet and dry seasons (Smith et al., 2008b). The mean Nd concentration for the seawater component is computed using values published in the literature (Table 1; Fig. 5).

\[
J_{\text{Nd}}^{\text{in}} = J_{\text{Nd}}^{\text{out}};
\]

which upon expansion gives the following expression:

\[
J_{\text{Nd}}^{\text{rivers-eff}} + J_{\text{Nd}}^{\text{seawater}} + J_{\text{Nd}}^{\text{TotSGD}} = J_{\text{Nd}}^{\text{MarineSGD}} + [Nd]_{\text{lagoon}}Q_{\text{out}}
\]

where \(J_{\text{Nd}}^{\text{rivers-eff}}\) is the effective river flux of Nd into the Indian River Lagoon following 70% removal of Nd (i.e., \([Nd]_{\text{rivers-eff}}\)) in the estuarine portions of the Eau Gallie River and Crane Creek, and \([Nd]_{\text{lagoon}}\) is the mean Nd concentration of the lagoon waters (Table 4). The effective river flux accounts for REE removal by salt-induced,}

### Table 3

Mean chloride concentrations (± 1 SD of the mean) and fluxes used in the mass balance model.

<table>
<thead>
<tr>
<th></th>
<th>CI (mmol kg(^{-1}))</th>
<th>(Q) (m(^3) day(^{-1}))</th>
<th>(J_{\text{CI}}) (mol day(^{-1}))</th>
</tr>
</thead>
<tbody>
<tr>
<td>Terrestrial SGD</td>
<td>4.67 ± 1.74</td>
<td>5.75 × 10(^5)</td>
<td>26.9(± 10) × 10(^3)</td>
</tr>
<tr>
<td>Marine SGD</td>
<td>338 ± 15(^b)</td>
<td>5.85 × 10(^5)</td>
<td>198(± 8.78) × 10(^6)</td>
</tr>
<tr>
<td>Total SGD</td>
<td>335 ± 14.9(^c)</td>
<td>5.91 × 10(^5)</td>
<td>198(±8.8) × 10(^6)</td>
</tr>
<tr>
<td>Rivers</td>
<td>10.5 ± 0.23(^d)</td>
<td>1.38 × 10(^5)</td>
<td>1.45(± 0.03) × 10(^6)</td>
</tr>
<tr>
<td>Lagoon</td>
<td>338 ± 15</td>
<td>545.88(^e)</td>
<td>2.24(±0.69) × 10(^6)</td>
</tr>
<tr>
<td>Seawater</td>
<td>458 ± 8</td>
<td>2.24(±0.69) × 10(^6)</td>
<td>122(±37.7) × 10(^6)</td>
</tr>
</tbody>
</table>

\(^a\) Measured variability of SGD fluxes is estimated to be ±50%. The total SGD discharging from the subterranean estuary is \(Q_{\text{TotSGD}} = Q_{\text{TSGD}} + Q_{\text{MarineSGD}}\), which is equivalent to \(Q_{\text{TSGD}} + Q_{\text{RL}}\), where RL is recirculated Indian River Lagoon water (see Martin et al., 2007 and Fig. 3).

\(^b\) Mean ± SD CI concentration of IRL water column samples (see Table 1).

\(^c\) Calculated using Eq (1).

\(^d\) Discharge weighted mean ± SD for the Eau Gallie River and Crane Creek (see Martin et al., 2007).

\(^e\) Assumes a salinity of 35\(\%\)_o for the coastal ocean (see text for details).
colloid coagulation as fresh river water mixes with brackish estuarine waters, which, on average, removes ~70% of the dissolved river-borne Nd load (Goldstein and Jacobsen, 1987; Sholkovitz, 1993, 1995; Sholkovitz and Szymczak, 2000; Lawrence and Kamber, 2006). Removal of REEs in the subterranean estuary by this process is, however, not significant as the marine component of SGD is already saline (Duncan and Shaw, 2003). Neodymium inputs to the Indian River Lagoon from the effective river flux, seawater exchange, and total SGD sum to 304 ± 64.7 mmol day\(^{-1}\), whereas the Nd flux out of the lagoon via input to the subterranean estuary by marine SGD (i.e., recirculated lagoon water) and export to the coastal ocean is estimated to be 105 ± 287(±64.4) mmol Nd day\(^{-1}\) (Fig. 3; Table 4). These values indicate that the Nd concentrations within the Indian River Lagoon water column are roughly at steady-state with respect to the Nd fluxes into and out of the lagoon. We estimate a residence time for Nd of 14.1 ± 3.2 days for the studied portion of the lagoon, which is identical to the estimated residence time for Nd of 14.1 ± 3.2 days for the studied portion of the Indian River Lagoon (Duncan and Shaw, 2003). Neodymium inputs to the Indian River Lagoon subterranean estuary are presented in Table 5. As with Nd, fluxes of REEs into the subterranean estuary, \(J_{\text{in}}^{\text{REE}}\), include terrestrial SGD and marine SGD (i.e., recirculated lagoon water), whereas the REEs are exported from the subterranean estuary to the overlying lagoon water, \(J_{\text{out}}^{\text{REE}}\), via total SGD (Fig. 3). The difference between \(J_{\text{in}}^{\text{REE}}\) and \(J_{\text{out}}^{\text{REE}}\) again represents the net flux of each REE from the subterranean estuary to the overlying coastal waters. The computed net fluxes reveal that more of each LREE and MREE is exported from the subterranean estuary to the overlying coastal waters of the Indian River Lagoon than imported into the subterranean estuary via terrestrial SGD and recirculating lagoon waters (Table 5). However, the opposite is observed for REEs heavier than Ho such that the amount of each of these HREEs imported into the subterranean estuary exceeds that which is exported to the coastal lagoon waters via total SGD. Therefore, the subterranean estuary beneath the Indian River Lagoon appears to act as a net source of LREEs and MREEs to the overlying coastal waters and as a sink for the HREEs.

### 5. DISCUSSION

#### 5.1. SGD fluxes of REEs in the Indian River Lagoon subterranean estuary

Our current best estimate of the total SGD Nd flux to the studied portion of the Indian River Lagoon is 287 ± 64.4 mmol day\(^{-1}\), which is ~150-fold larger than

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### Table 4

Mean Nd concentrations (± 1 SD of the mean), mean discharge of groundwater and surface waters, and neodymium fluxes for the studied portion of the Indian River Lagoon.

<table>
<thead>
<tr>
<th>Source</th>
<th>Nd (pmol kg(^{-1}))</th>
<th>(Q) (m(^3) day(^{-1}))</th>
<th>(J_{\text{out}}^{\text{Nd}}) (mol day(^{-1}))</th>
</tr>
</thead>
<tbody>
<tr>
<td>Terrestrial SGD</td>
<td>350 ± 59.6</td>
<td>5.75 × 10(^3)</td>
<td>2.01(± 0.34) × 10(^{-3})</td>
</tr>
<tr>
<td>Marine SGD</td>
<td>315 ± 58</td>
<td>5.85 × 10(^3)</td>
<td>184(± 33.9) × 10(^{-3})</td>
</tr>
<tr>
<td>Total SGD</td>
<td>485 ± 109</td>
<td>5.91 × 10(^3)</td>
<td>287(± 64.4) × 10(^{-3})</td>
</tr>
<tr>
<td>Rivers</td>
<td>311 ± 129(^a)</td>
<td>1.38 × 10(^3)</td>
<td>(12.7(± 5.27) × 10(^{-3})</td>
</tr>
<tr>
<td>Lagoon</td>
<td>315 ± 58</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Seawater</td>
<td>17.9 ± 1.16</td>
<td>2.24(± 0.69) × 10(^3)</td>
<td>4.01(± 1.26) × 10(^{-3})</td>
</tr>
<tr>
<td>Lagoon</td>
<td>304(±64.7) × 10(^{-3})</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Sediments</td>
<td>186(±33.9) × 10(^{-3})</td>
<td></td>
<td>100(±72.8) × 10(^{-3})</td>
</tr>
</tbody>
</table>

\(^a\) Discharge weighted mean Nd concentration of Eau Gallie River and Crane Creek.

\(^b\) Effective river flux that assumes that 70% of the river borne Nd is removed in these estuaries.

\(^c\) \(J_{\text{in}}^{\text{Nd}}\) \(=\) \(J_{\text{Terrestrial SGD}}^{\text{Nd}} + J_{\text{Marine SGD}}^{\text{Nd}} + J_{\text{TSGD}}^{\text{Nd}}\).

\(^d\) \(J_{\text{Terrestrial SGD}}^{\text{Nd}} = \frac{\left(Q_{\text{Terrestrial SGD}} \times [Nd]_{\text{Terrestrial SGD}}\right)}{Q_{\text{Terrestrial SGD}}}\).

\(^e\) \(J_{\text{Marine SGD}}^{\text{Nd}} = \frac{\left(Q_{\text{Marine SGD}} \times [Nd]_{\text{Marine SGD}}\right)}{Q_{\text{Marine SGD}}}\).

\(^f\) \(J_{\text{TSGD}}^{\text{Nd}} = \frac{\left(Q_{\text{TSGD}} \times [Nd]_{\text{TSGD}}\right)}{Q_{\text{TSGD}}}\).

\(^g\) \(J_{\text{TSGD}}^{\text{Nd}} = \frac{\left(Q_{\text{TSGD}} \times [Nd]_{\text{TSGD}}\right)}{Q_{\text{TSGD}}}\).

\(^h\) \(J_{\text{in}}^{\text{Nd}}\) \(=\) \(J_{\text{Terrestrial SGD}}^{\text{Nd}} + J_{\text{Marine SGD}}^{\text{Nd}}\).
Rare earth elements in submarine groundwater discharge

Table 5

<table>
<thead>
<tr>
<th>Element</th>
<th>$J_{REE}^{TOT}$</th>
<th>$J_{REE}^{MARINE}$</th>
<th>$J_{REE}^{SGD}$</th>
<th>$J_{REE}^{in}$</th>
<th>$J_{REE}^{out}$</th>
<th>$J_{REE}^{net}$</th>
</tr>
</thead>
<tbody>
<tr>
<td>La</td>
<td>1.08 ± 0.24</td>
<td>150 ± 30</td>
<td>233 ± 65</td>
<td>151 ± 30</td>
<td>223 ± 65</td>
<td>82.6 ± 71.6</td>
</tr>
<tr>
<td>Ce</td>
<td>2.46 ± 0.49</td>
<td>291 ± 57.3</td>
<td>456 ± 140</td>
<td>294 ± 57.3</td>
<td>456 ± 140</td>
<td>162 ± 151</td>
</tr>
<tr>
<td>Pr</td>
<td>0.27 ± 0.07</td>
<td>34 ± 9.13</td>
<td>52.2 ± 18.3</td>
<td>34.3 ± 9.13</td>
<td>52.2 ± 18.3</td>
<td>17.9 ± 20.5</td>
</tr>
<tr>
<td>Nd</td>
<td>2.01 ± 0.34</td>
<td>184 ± 33.9</td>
<td>287 ± 64.4</td>
<td>186 ± 33.9</td>
<td>287 ± 64.4</td>
<td>100 ± 72.8</td>
</tr>
<tr>
<td>Sm</td>
<td>0.38 ± 0.09</td>
<td>33.2 ± 5.67</td>
<td>48.8 ± 10.2</td>
<td>33.6 ± 6.67</td>
<td>48.8 ± 10.2</td>
<td>15.2 ± 12.2</td>
</tr>
<tr>
<td>Eu</td>
<td>0.09 ± 0.03</td>
<td>7.43 ± 2.5</td>
<td>9.69 ± 2.58</td>
<td>7.5 ± 2.5</td>
<td>9.69 ± 2.58</td>
<td>2.18 ± 3.6</td>
</tr>
<tr>
<td>Gd</td>
<td>0.47 ± 0.12</td>
<td>40.4 ± 1.2</td>
<td>58.7 ± 10.4</td>
<td>40.8 ± 6.2</td>
<td>58.7 ± 10.4</td>
<td>17.8 ± 12.1</td>
</tr>
<tr>
<td>Tb</td>
<td>0.08 ± 0.02</td>
<td>7.25 ± 1.19</td>
<td>7.8 ± 1.09</td>
<td>7.34 ± 1.19</td>
<td>7.8 ± 1.09</td>
<td>0.46 ± 1.61</td>
</tr>
<tr>
<td>Dy</td>
<td>0.80 ± 0.21</td>
<td>57.6 ± 9.48</td>
<td>62.1 ± 6.74</td>
<td>58.4 ± 9.48</td>
<td>62.1 ± 6.74</td>
<td>3.69 ± 11.6</td>
</tr>
<tr>
<td>Ho</td>
<td>0.25 ± 0.09</td>
<td>15.8 ± 4.39</td>
<td>15 ± 2.61</td>
<td>16.4 ± 3.9</td>
<td>15 ± 2.61</td>
<td>-1.03 ± 5.11</td>
</tr>
<tr>
<td>Er</td>
<td>1.54 ± 0.63</td>
<td>84.8 ± 32.3</td>
<td>61.5 ± 9.63</td>
<td>86.4 ± 32.3</td>
<td>61.5 ± 9.63</td>
<td>-24.9 ± 33.7</td>
</tr>
<tr>
<td>Tm</td>
<td>0.41 ± 0.18</td>
<td>19.6 ± 8.95</td>
<td>10.5 ± 1.3</td>
<td>20 ± 8.95</td>
<td>10.5 ± 1.3</td>
<td>-9.54 ± 9.05</td>
</tr>
<tr>
<td>Yb</td>
<td>4.12 ± 1.97</td>
<td>171 ± 92.4</td>
<td>85.7 ± 11.3</td>
<td>176 ± 92.5</td>
<td>85.7 ± 11.3</td>
<td>-89.8 ± 93.1</td>
</tr>
<tr>
<td>Lu</td>
<td>0.89 ± 0.48</td>
<td>38.9 ± 21.1</td>
<td>33.2 ± 12.2</td>
<td>39.8 ± 21.1</td>
<td>33.2 ± 12.2</td>
<td>-26.5 ± 21.2</td>
</tr>
</tbody>
</table>

* $J_{REE}^{TOT} = J_{REE}^{SGD} + J_{REE}^{MARINE}$
* $J_{REE}^{out} = J_{REE}^{SGD}$
* $J_{REE}^{net} = J_{REE}^{SGD} - (J_{REE}^{SGD} + J_{REE}^{MARINE})$

The results of these calculations are schematically shown in Fig. 6, which presents the mean, discharge-weighted, shale-normalized REE pattern of the flux into the Indian River Lagoon subterranean estuary (i.e., terrestrial SGD + recirculating lagoon water) along with the mean, shale-normalized REE pattern of the outflow from the subterranean estuary (i.e., total SGD). Fig. 6 demonstrates that the mean, shale-normalized REE signature of water entering the Indian River Lagoon subterranean estuary is more enriched in HREEs compared to LREEs than the flux of water leaving the subterranean estuary as total SGD. These observations can be further quantified by comparison of the shale-normalized Yb/Nd ratio (i.e., (Yb/Nd)$_{SN}$), which is 12.9 for the inflow water (terrestrial SGD + recirculated marine SGD) and 4.1 for the outflow water (total SGD). We note that the (Yb/Nd)$_{SN}$ ratio of total SGD is similar to that of seawater (i.e., ~4; Table 1; Fig. 4). The difference between the shale-normalized REE patterns of the inflow and outflow illustrates the changes in REE concentrations and fractionation patterns that occur as groundwater flows into and through the subterranean estuary beneath the Indian River Lagoon. Specifically, geochemical reactions affecting REEs within the Indian River Lagoon subterranean estuary lead to a general flattening of shale-normalized REE patterns as LREEs and MREEs are added to groundwater flowing through these sediments and HREEs are retained in the sediments. These observations are remarkable as geochemical reactions that fractionate REEs in surface estuaries (i.e., removal in low-salinity regions by salt-induced, colloid coagulation; mobilization in mid- to high-salinity regions) tend to further enrich HREEs in solution relative to LREEs and MREEs compared to shale (e.g., Elderfield et al., 1990; Sholkovitz, 1992, 1993, 1995; Sholkovitz and Szczepczak, 2000; Lawrence and Kamber, 2006).

5.2. REE cycling in the Indian River Lagoon subterranean estuary

Rare earth element concentrations in porewaters from the Indian River Lagoon subterranean estuary are more than 10–100 times greater than typical open-ocean seawater REE concentrations (e.g., Table 1). The highest REE concentrations occur in SGD from the vicinity of the freshwater–saltwater seepage face (i.e., EGN-22.5; Figs. 2 and 4).
These REE concentrations are too high to originate from binary mixing of seawater with terrestrial SGD, and instead point to geochemical reactions occurring within the subterranean estuary as an important source of REEs to SGD, and hence, the overlying lagoon waters. This interpretation is consistent with the observations of Duncan and Shaw (2003) who argued that geochemical reactions occurring in a subterranean estuary along the coast of South Carolina, USA, appear to add REEs to local coastal waters.

A number of geochemical reactions could play important roles in mobilizing REEs within the Indian River Lagoon subterranean estuary. For example, mobilization of REEs may in part reflect the increase in salinity across the seepage face within the subterranean estuary. It is well documented that REEs are released from sediments in the mid- to high-salinity regions of surface estuaries (Sholkovitz, 1993, 1995; Sholkovitz and Szymbczak, 2000). Furthermore, Tang and Johannesson (2010) demonstrated in laboratory experiments that an increase in ionic strength from 0.01 to 0.1 mol kg\(^{-1}\) dramatically suppresses REE adsorption onto Fe(III) oxide/oxhydroxide-coated quartz sands. These results imply that mixing of recirculating marine SGD with terrestrial SGD in subterranean estuaries could induce REE desorption from aquifer mineral surface sites, thereby contributing to the REE flux to coastal waters (Tang and Johannesson, 2010). However, REE concentrations are, at best, only weakly associated with ionic strength in SGD from the Indian River Lagoon subterranean estuary (r = 0.35, Nd; r = 0.31, Gd; r = −0.12, Yb), and none of these relationships are statistically significant. Hence, although salinity-related REE release to SGD in the Indian River Lagoon subterranean estuary likely contributes to the overall net flux of LREEs and MREEs exported to the overlying coastal lagoon waters, it does not appear to be the predominant geochemical reaction mobilizing REEs in this subterranean estuary. Alternatively, because Fe diagenesis in the Indian River Lagoon subterranean estuary also chiefly occurs at the freshwater–saltwater seepage face similar to the REEs (Roy et al., 2010), geochemical reactions controlling the Fe cycle in the subterranean estuary represent another possible mechanism for REE mobilization.

The Fe cycle in the Indian River Lagoon subterranean estuary has recently been discussed in detail (Roy et al., 2010). As mentioned above Fe(III) oxide/oxhydroxide coatings on the subterranean estuary sediments are ubiquitous, clearly indicating precipitation of Fe(III) oxides/oxhydroxides under sub-aerial conditions when sea-level was lower and the subterranean estuary was further offshore (e.g., during the Last Glacial Maximum; Hartl, 2006; Roy et al., 2010). With rising sea-level, the freshwater–saltwater seepage face has moved landward, flooding these orange, terrestrial sediments and consequently impacting the Fe cycle such that it now involves three distinct steps (Roy et al., 2010). First, reductive dissolution of Fe(III) oxides/oxhydroxides occurs within these orange sediments of the subterranean estuary when hypoxic terrestrial SGD mixes with recirculating marine SGD. Reduction of Fe(III) is likely driven by the introduction of labile dissolved organic carbon (DOC) entrained within the DOC-rich marine SGD (e.g., Cherrier et al., 2007; Roy et al., 2010). Second, the resulting dissolved Fe(II) is transported upwards with advecting groundwater into the sulfate reduction zone at the base of the overlying black, marine sediments, where it is subsequently precipitated as Fe-sulfides (Roy et al., 2010). Third, some of these Fe-sulfides are redissolved near the sediment–water interface where bioturbation provides dissolved oxygen from the overlying,oxic lagoon waters (Roy et al., 2010).

Iron cycling within the Indian River Lagoon subterranean estuary differs from that of Waquoit Bay on Cape Cod, Massachusetts, where Fe oxidation and subsequent precipitation of Fe(III) oxides/oxhydroxides predominates (Charette and Sholkovitz, 2002; Testa et al., 2002; Charette et al., 2005; Spiteri et al., 2006, 2008). At Waquoit Bay, the large tidal range (~1.5 m) regularly exposes the subterranean estuary to the atmosphere. Consequently, dissolved Fe(II) in hypoxic, terrestrial SGD oxidizes and precipitates within the subterranean estuary as terrestrial SGD mixes with oxic marine SGD (Charette and Sholkovitz, 2002). Because the Indian River Lagoon is microtidal, the corresponding subterranean estuary is not regularly exposed to the atmosphere, and hence, remains submerged and anoxic (Roy et al., 2010). Furthermore, Fe(III) oxides/oxhydroxide precipitation in the subterranean estuary beneath Waquoit Bay is also apparently driven by an increase in pH from 5.5 nearshore, to ~8 at the freshwater–saltwater seepage face (Spiteri et al., 2006). In contrast, pH changes little within the Indian River Lagoon subterranean estuary and thus does not affect dissolved Fe concentrations in the same manner as in the Waquoit Bay subterranean estuary (Table 2; Roy et al., 2010). Therefore, Fe cycling in subterranean estuaries can be dominated by oxidation (Waquoit Bay, Massachusetts, USA; Patos Lagoon, Brazil) or reduction (Indian River Lagoon, Florida; North Inlet, South Carolina; Moses Hammock, Georgia, USA), which depends on the composition of the sediments and porewaters, as well as the “structure” (e.g., large tidal range vs. microtidal) of the subterranean estuary (Charette and Sholkovitz, 2002; Duncan and Shaw, 2003; Snyder et al., 2004; Windom et al., 2006; Roy et al., 2010).

The high dissolved Fe and low dissolved oxygen concentrations for the SGD samples presented in Table 2 further supports the Fe cycling model of Roy et al. (2010). That Fe(III) reduction predominates in the studied portion of the subterranean estuary is also supported by the low NO\(_3^−\) concentrations (~0.3 \(\mu\)mol kg\(^{-1}\)) measured in SGD seaward of the EGN-0 multisampler (Cherrier et al., 2007). Furthermore, all of the SGD from the Indian River Lagoon sampled in this study are undersaturated with respect to goethite (Table 2), indicating that reductive dissolution of goethite is thermodynamically favorable. Taken together, these data point to Fe(III) reduction as the chief redox buffering reaction controlling the redox conditions within the portion of the subterranean estuary sampled.

Similar to the REEs, dissolved Fe concentrations at the freshwater–saltwater seepage face are also substantially higher than can be explained by simple mixing of seawater and terrestrial SGD, indicating \textit{in situ} reactions are responsible for the elevated Fe concentrations (Roy et al., 2010). Because REEs strongly adsorb onto Fe(III) oxides/oxhydroxides under sub-aerial conditions when sea-level was lower and the subterranean estuary was further offshore (e.g., during the Last Glacial Maximum; Hartl, 2006; Roy et al., 2010). With rising sea-level, the freshwater–saltwater seepage face has moved landward, flooding these orange, terrestrial sediments and consequently impacting the Fe cycle such that it now involves three distinct steps (Roy et al., 2010). First, reductive dissolution of Fe(III) oxides/oxhydroxides occurs within these orange sediments of the subterranean estuary when hypoxic terrestrial SGD mixes with recirculating marine SGD. Reduction of Fe(III) is likely driven by the introduction of labile dissolved organic carbon (DOC) entrained within the DOC-rich marine SGD (e.g., Cherrier et al., 2007; Roy et al., 2010). Second, the resulting dissolved Fe(II) is transported upwards with advecting groundwater into the sulfate reduction zone at the base of the overlying black, marine sediments, where it is subsequently precipitated as Fe-sulfides (Roy et al., 2010). Third, some of these Fe-sulfides are redissolved near the sediment–water interface where bioturbation provides dissolved oxygen from the overlying,oxic lagoon waters (Roy et al., 2010). Iron cycling within the Indian River Lagoon subterranean estuary differs from that of Waquoit Bay on Cape Cod, Massachusetts, where Fe oxidation and subsequent precipitation of Fe(III) oxides/oxhydroxides predominates (Charette and Sholkovitz, 2002; Testa et al., 2002; Charette et al., 2005; Spiteri et al., 2006, 2008). At Waquoit Bay, the large tidal range (~1.5 m) regularly exposes the subterranean estuary to the atmosphere. Consequently, dissolved Fe(II) in hypoxic, terrestrial SGD oxidizes and precipitates within the subterranean estuary as terrestrial SGD mixes with oxic marine SGD (Charette and Sholkovitz, 2002). Because the Indian River Lagoon is microtidal, the corresponding subterranean estuary is not regularly exposed to the atmosphere, and hence, remains submerged and anoxic (Roy et al., 2010). Furthermore, Fe(III) oxides/oxhydroxide precipitation in the subterranean estuary beneath Waquoit Bay is also apparently driven by an increase in pH from 5.5 nearshore, to ~8 at the freshwater–saltwater seepage face (Spiteri et al., 2006). In contrast, pH changes little within the Indian River Lagoon subterranean estuary and thus does not affect dissolved Fe concentrations in the same manner as in the Waquoit Bay subterranean estuary (Table 2; Roy et al., 2010). Therefore, Fe cycling in subterranean estuaries can be dominated by oxidation (Waquoit Bay, Massachusetts, USA; Patos Lagoon, Brazil) or reduction (Indian River Lagoon, Florida; North Inlet, South Carolina; Moses Hammock, Georgia, USA), which depends on the composition of the sediments and porewaters, as well as the “structure” (e.g., large tidal range vs. microtidal) of the subterranean estuary (Charette and Sholkovitz, 2002; Duncan and Shaw, 2003; Snyder et al., 2004; Windom et al., 2006; Roy et al., 2010).
Sediments, such as Fe(III) oxides/oxyhydroxides, are solubilizing sediment diagenesis as minor phases within the waters relative to HREEs has previously been reported during similar preferential release of LREEs and MREEs to porewaters (Sholkovitz et al., 1987; Sholkovitz et al., 1989, 1992).

The overlying lagoon waters (Table 5; Fig. 6). We note that the magnitude of the net SGD fluxes for LREEs and MREEs to the overlying lagoon waters (Table 5: Fig. 6). We note that similar preferential release of LREEs and MREEs to porewaters relative to HREEs has previously been reported during sediment diagenesis as minor phases within the sediments, such as Fe(III) oxides/oxyhydroxides, and that the order of release follows: LREE > MREE > HREE. This order of release is broadly consistent with the relative magnitude of the net SGD fluxes for LREEs and MREEs to the overlying lagoon waters (Table 5; Fig. 6). We note that similar preferential release of LREEs and MREEs to porewaters relative to HREEs has previously been reported during sediment diagenesis as minor phases within the sediments, such as Fe(III) oxides/oxyhydroxides, and solubilized by reducing conditions (Elderfield and Sholkovitz, 1987; Sholkovitz et al., 1989, 1992).

All of the REEs exhibit weak, inverse relationships with dissolved sulfide in Indian River Lagoon SGD, none of which are statistically significant: Nd ($r = -0.32$); Gd ($r = -0.22$); and Yb ($r = -0.32$). These relationships suggest that REEs are not strongly associated with sulfur cycling in the subterranean estuary. We interpret this as preliminary evidence that REEs are not removed from advecting SGD to the same degree as Fe(II) by precipitating Fe-sulfides at the base of the black, marine sediments within the Indian River Lagoon subterranean estuary. This is not surprising as convincing evidence currently does not exist implicating sulfide mineral formation with REE removal from marine waters (Naveau et al., 2006; also see Schijf et al., 1995; DeCarlo et al., 1998; Chailou et al., 2006). Nevertheless, without sediment data for the REEs, we cannot unequivocally address whether REEs are also removed with precipitating metal sulfides in Indian River Lagoon sediments. Future investigations will involve examination of REE speciation in these sediments.

Consequently, the data and mass balance calculations presented here indicate that advective SGD fluxes, in combination with geochemical reactions occurring in the subterranean estuary, act in concert to deliver a substantial net flux of LREEs and MREEs to the overlying coastal waters of the Indian River Lagoon. Specifically, we suggest that hypoxic, terrestrial SGD mixes with DOC-rich, recirculating lagoon water in the Indian River Lagoon subterranean estuary, promoting reductive dissolution of Fe(III) oxides/oxyhydroxides that were precipitated within the aquifer sediments under sub-aerial conditions when sea-level was lower during the last glacial maximum (Roy et al., 2010). During the previous sea-level low-stand, REEs in percolating vadose waters were scavenged by precipitating Fe(III) oxides/oxyhydroxides that are now ubiquitous in these sediments. During the Holocene the former vadose zone was flooded as the water table rose in response to rising sea-level (Roy et al., 2010). Because LREEs and MREEs preferentially sorb to amorphous Fe(III) oxide/oxyhydroxides compared to HREEs in the presence of strong complexing ligands like carbonate ions (Quinn et al., 2006b; Tang and Johannesson, 2010), subsequent reductive dissolution of the Fe(III) oxides/oxyhydroxides releases relatively more LREEs and MREEs to the porewater of the subterranean estuary than HREEs. Upon release from the sediments, REEs form strong solution complexes with carbonate ions, with LREEs and MREEs predominantly forming LnCO$_3^-$ complexes at these pH’s (6.8 $\leq$ pH $\leq$ 7.4), and the HREEs chiefly forming Ln(CO$_3$)$_2^-$ complexes (Fig. 5). Because amorphous Fe(III) oxides/oxyhydroxides exhibit a positive surface charge for pH $< 8$ (Dzombak and Morel, 1990), re-adsorption of negatively charged Ln(CO$_3$)$_2^-$ complexes of the HREEs to Fe(III) oxide/oxyhydroxides sequesters HREEs in the subterranean estuary. However, positively charged LnCO$_3^-$ complexes of LREEs and MREEs are stable in solution (Tang and Johannesson, 2005) and thus transported with advecting groundwater to overlying coastal waters. Heavy REEs also exhibit greater affinity for quartz than lighter REEs (Byrne and Kim, 1990), and hence, may be preferentially sorbed onto the predominant quartz grains that make up the Surficial Aquifer. However, this is unlikely to be significant as quartz grains within the aquifer sediments are typically coated with Fe(III) oxides/oxyhydroxides (Hartl, 2006; Roy et al., 2010). The net effect of these geochemical reactions on REE cycling in the Indian River Lagoon subterranean estuary is that of progressive flattening of the shale-normalized fractionation patterns as LREEs and MREEs are added to recirculating marine SGD, and hence the coastal lagoon waters and the HREEs are removed within the subterranean estuary.

5.3. Global implications

The global implications of SGD fluxes of REEs, and Nd in particular, to the coastal oceans await additional investigations at more sites around the world. It is important to note that owing to its shallow depth and microtidal characteristics, the Indian River Lagoon and its underlying subterranean estuary may be unique and thus not representative of coastal aquifers and subterranean estuaries in general. Nonetheless, the net flux of Nd to these coastal waters of Florida represents a previously unrecognized source of Nd to the coastal ocean that could conceivably be important to the global oceanic Nd budget. More importantly, our interpretation of the factors controlling SGD Nd fluxes implies that on glacial-interglacial time scales these fluxes could vary significantly due to changes in sea-level (Morrissey et al., 2010; Severmann et al., 2010; Roy et al., 2010). This could be complicated by SGD directly to the deep ocean along the shelf-slope break during sea-level low stands (Robb, 1984; Person et al., 2003, 2007; Cohen et al., 2010). As a result the use of Nd concentrations and isotopes in biogenic sediments as tracers of changes in ocean.
circulation on these time scales (Frank, 2002; Goldstein and Hemming, 2003) may be more complex than previously thought. Clearly, additional investigations are warranted to constrain the spatial and temporal variations in SGD Nd fluxes to the coastal oceans.

6. CONCLUSIONS

Concentration variations of the REEs in porewaters collected across the freshwater–saltwater seepage face within the Indian River Lagoon subterranean estuary are too high to be explained by simple, binary mixing between terrestrial SGD and seawater. Furthermore, these high REE concentrations can not be explained by mixing of terrestrial SGD with recirculated Indian River Lagoon waters. Instead, the high REE concentrations at the seepage face point to in situ geochemical reactions occurring in the sediments of the subterranean estuary as an important source of REEs to SGD and hence, the overlying coastal lagoon waters. Specifically, geochemical reactions occurring within the subterranean estuary (e.g., reductive dissolution of Fe(III) oxides/oxyhydroxides) preferentially mobilize LREEs and MREEs from the subterranean estuary sediments. These trace elements form strong, positively charged carbonato complexes, LnCO$_3^-$, in solution and are subsequently advectively transported to the overlying coastal waters, accounting for a net flux of LREEs and MREEs from the subterranean estuary to the coastal lagoon waters. The net SGD flux of Nd to the overlying lagoon waters is ~7 times larger than the local effective river flux to the Indian River Lagoon. This previously unrecognized source of Nd to the coastal ocean could be important to the global oceanic Nd budget, and help to resolve the oceanic “Nd paradox” by accounting for a substantial fraction of the hypothesized missing Nd flux to the ocean.

The subterranean estuary beneath the Indian River Lagoon acts a sink for HREEs owing to their re-adsorption onto Fe(III) oxides/oxyhydroxides. Unlike the LREEs and MREEs, significant fractions of the HREEs occur in positively charged surface sites on Fe(III) oxides/oxyhydroxides that predominate in these circumneutral pH porewaters. The overall effect of these geochemical reactions occurring within the subterranean estuary is progressive flattening of the shale-normalized fractionation patterns as LREEs and MREEs are added to recirculating marine SGD, and thus the coastal lagoon waters and HREEs are sequestered within the sediments of the subterranean estuary. Future investigation will involve more extensive sampling for both analysis of REE concentrations and the Nd isotope compositions of groundwater and surface waters of the Indian River Lagoon system.

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