**Submarine Ground-Water Discharge: Nutrient Loading and Nitrogen Transformations**

Eutrophication of coastal waters due to nonpoint source land-derived nitrogen (N) loads is a worldwide phenomenon and perhaps the greatest agent of change altering coastal ecology (National Research Council, 2000; Howarth and others, 2000). Within the United States, a majority of estuaries have been determined to be moderately to severely impaired by eutrophication associated with increasing nutrient loads (Bricker and others, 1999).

In coastal watersheds with soils of high hydraulic conductivity and permeable coastal sediments, ground water is a major route of transport of freshwater and its solutes from land to sea. Freshwater flowing downgradient from aquifers may either discharge from a seepage face near the intertidal zone, or flow directly into the sea as submarine ground-water discharge (SGD) (fig. 1). In the coastal aquifer, entrainment of saline pore water occurs prior to discharge, producing a gradient in ground-water salinity from land to sea, referred to as a subterranean estuary (Moore, 1999). In addition, processes including density-driven flow and tidal pumping create brackish and saline ground-water circulation. Hence, submarine ground-water discharge often consists of a substantial amount of recirculating seawater. Mixing of fresh and saline ground waters in the context of coastal sediments may alter the chemical composition of the discharging fluid. Depending on the biogeochemical setting, removal of fixed N due to processes leading to N₂ (dinitrogen gas) production in the nearshore aquifer and subterranean estuary may significantly attenuate land-derived N loads; or, processes such as ion exchange and tidal pumping in the subterranean estuary may substantially accelerate the transport of both land-derived and sediment re-mineralized N to estuarine water columns.

As emphasized by Burnett and others (2001, 2002), a fundamental problem in evaluating the importance of ground-water discharge in marine geochemical budgets is the difficulty of collecting samples across the salinity gradients of coastal aquifers. In addition, locating and quantifying rates of submarine ground-water discharge remains a challenge due to the diffuse and spatially and temporally heterogeneous nature of discharge. As a result, with regard to the study of biogeochemical cycles and chemical loads to coastal waters, the seepage face and subterranean estuary are relatively new and under-studied zones in the aquatic cascade from watershed to sea. Processes occurring in those zones must be understood and considered for proper modeling and management of coastal water resources.

**Approaches to Investigating Transformations**

To investigate controls on nutrient loads carried by submarine ground-water discharge, the U.S. Geological Survey (USGS) and Woods Hole Oceanographic Institution are examining ground-water flow, nutrient transport, and biogeochemistry in nearshore portions of coastal aquifers in a range of settings that contrast in terms of hydrogeology, land uses, and climate. This fact sheet describes research at sites in Waquoit Bay, Massachusetts, and in Tampa Bay, Florida. The biogeochemical investigations of nitro-

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*Figure 1. Cross-section of a submarine ground-water discharge zone (from Swarzenski, Bratton, and others, 2004).*
gen are being carried out in concert with examinations of radiochemical ground-water discharge tracers, seepage meter studies, and streaming resistivity surveys to identify locations and sources of submarine ground-water discharge and to quantify fluxes (Swarzenski and others, 2001; Talbot and others, 2003; Abraham and others, 2003; Swarzenski, Burnett and others, 2004; Swarzenski, Charette, and Langevin, 2004; Swarzenski, Bratton, and Crusius, 2004; and Crusius and others, 2005).

Our three primary areas of research with regard to N biogeochemistry in submarine ground-water discharge zones are:

1. Using naturally occurring N stable isotope ratios to trace fixed N loss.
2. Application of isotope pairing experiments to develop new understanding of the N cycle in coastal aquifers.
3. Using noble (inert) gases as tracers of N₂ production.

**Nitrogen Stable Isotopic Ratios**

At the Waquoit Bay site the research team is using a stainless-steel drive point piezometer system to collect pore water samples in high-resolution vertical profiles. The samples are analyzed for a suite of biogeochemically cycled elements and tracers (Charette and Sholkovitz, 2006; Kroeger and Charette, unpublished data). The piezometer system allows profiles to be collected to a depth of 9 meters (m) or more with a sampling interval as small as 0.15 m. The study is examining three major subsurface water masses with distinct biogeochemical characteristics: the nearshore freshwater aquifer, and two fresh and saline water mixing zones (referred to as shallow and deep subterranean estuaries).

The dominant pattern in nitrogen concentrations in the nearshore freshwater aquifer at the head of Waquoit Bay is the occurrence of both nitrate and ammonium plumes (fig. 2A) with concentrations exceeding 200 micromolar (µM) in some locations. The nitrate and ammonium plumes were constant features during 3 years of sampling, but concentrations do vary over time.

Ammonium is the dominant form of nitrogen within the deep subterranean estuary, with maximum concentrations in individual profiles (40 to 70 µM) occurring at salinities of 26 to 28, and at 5- to 8-m depths within the sediment (fig. 2A and 2C). Brackish to saline pore waters are generally devoid of nitrate, have dissolved oxygen concentration from 0 to 3 milligrams per liter, and a low oxidation/reduction potential. During transport through the subterranean estuary, ammonium concentration generally suggests conservative behavior. The greatest concentrations of ammonium are at the highest salinity. The apparent conservative behavior of ammonium along the salinity gradient suggests that the preponderance of ammonification occurs at highest salinity, and that mixing and dilution are the dominant processes controlling ammonium concentration in the deep subterranean estuary. Based on radiochemical and hydrological estimates of fresh and saline ground-water discharge and on nitrogen concentrations in those zones, regenerated ammonium transport to the bay by advection of brackish to saline pore water is estimated to be in the range of 25 percent of the land-derived dissolved inorganic nitrogen load (Talbot and others, 2003; Abraham and others, 2003).

The co-occurrence of nitrate and ammonium is uncommon throughout the nearshore aquifer and subterranean estuary. The largely mutually exclusive distributions of nitrate and ammonium may be in part due to the loss of both N species in locations where advection brings nitrate-bearing waters with a low

![Figure 2](image-url)

**Figure 2.** Graph showing (A) nitrate and ammonium concentrations; (B) nitrogen stable isotopic ratios; (C) salinity and dissolved oxygen concentration, in a depth profile through a freshwater aquifer and into a subterranean estuary. Blue shaded areas indicate regions of co-occurrence of nitrate and ammonium. Abbreviations and symbols are as follows: δ¹⁵N (per mil), the stable isotope ratio calculated as \([(^{15}N/^{14}N \text{ sample})/^{15}N/^{14}N \text{ atmospheric nitrogen} \] - 1) x 1000; m, meters; µM, micromoles per liter; D.O., dissolved oxygen; mg/L, milligrams per liter.
dissolved-oxygen concentration into contact with relatively reducing ammonium-rich waters (fig. 2, blue shaded areas). In those mixing zones, a decrease in concentration and an increase in the nitrogen stable isotopic ratio in both nitrate and ammonium suggests concurrent loss of ammonium and nitrate, perhaps due to coupled nitrification/denitrification, or to other N₂-producing microbial processes (fig. 2A and 2B).

The pattern of nitrogen loss shown in figure 2 is a consistent one, observed in a number of locations in both fresh and saline portions of the nearshore aquifer. Mixing of the nitrate and ammonium plumes in the freshwater zone may be related in part to compression of groundwater flow lines associated with thinning of the freshwater aquifer near the seepage face. Denitrification of land-derived nitrate at low salinity within the subterranean estuary (fig. 2, blue shaded area at 5.5-m depth) may (1) provide an explanation for the absence of nitrate in brackish and saline subterranean waters and (2) suggest that transformations occurring within subterranean estuaries do have the potential to modify land-derived nitrogen loads prior to discharge to the sea.

**Isotope Pairing**

There has been a dramatic increase in recent years in the number of microbial N transformation pathways that have been shown or proposed to occur in marine environments (Luther and others, 1997; Hulth and others, 1999; Zehr and Ward, 2002). Of particular importance among the “new” N transformations is anaerobic ammonium oxidation (anammox) (fig. 3), a microbial process that in some settings occurs at significant rates at the ecosystem scale (Dalsgaard and others, 2003; Kuyper and others, 2003). Anammox has not been previously investigated in submarine ground-water discharge zones, and yet preliminary evidence from isotope pairing experiments at Waquoit Bay suggests that it may be an important pathway for N₂ formation in those environments.

With academic collaborators at the Woods Hole Oceanographic Institution, the study team is developing methods to investigate the consequences of advection through permeable sediments on N transport and transformations via a series of batch incubation experiments and flow-through experiments designed to simulate water transport through permeable marine sediments. The study is particularly focused on loss of fixed nitrogen due to production of N₂, and will involve a combined approach of isotope pairing and membrane inlet mass spectrometry with molecular biology to quantify and identify pathways of N₂ production.

**Noble Gases as Tracers**

With collaborators at the Marine Biological Laboratory, Woods Hole, and the Woods Hole Oceanographic Institution, the USGS study team is estimating the magnitude of N₂-producing processes within the range of biogeochemical zones occurring within freshwater, brackish, and saline aquifers. Membrane inlet mass spectrometry (MIMS) is used for rapid analysis of N₂ and argon concentrations in ground water. Measured N₂ concentrations will be compared to concentrations expected in the absence of in-aquifer N₂ production to calculate concentration of excess N₂. Complications in the interpretation of dissolved gas concentrations in ground water are (1) uncertainty regarding water temperature at the time of recharge to the aquifer, which will determine the inherited N₂/argon (Ar) ratio and (2) the presence of excess air due to entrapment of air bubbles at the time of recharge (Heaton and Vogel, 1981), followed by partial or complete dissolution of the trapped gases (Peeters and others, 2002). After recharge, hydrostatic pressure and bubble surface tension tend to cause bubble dissolution and oversaturation of gases. Since N₂ concentration/Ar concentration in air is greater than in water at saturation, bubble entrapment would tend to increase measured N₂/Ar in ground water and may result in overestimation of denitrification. To estimate concentrations of dissolved gases due to bubble entrapment and to constrain estimates of temperature at recharge, a subset of samples will be analyzed in the laboratory at the Woods Hole Oceanographic Institution for the full suite of noble gas concentrations by peak-height manometry using combined magnetic sector and quadrupole mass spectrometry. For those analyses, a set of samples will be obtained in stainless steel transfer vessels, and vacuum extracted in the laboratory for storage in glass ampoules. The gas samples will be chemically purified and cryogenically separated for mass spectrometric analysis. Based on the results of those analyses, recharge temperature, bubble entrapment, and the extent of bubble dissolution can be modeled.

Estimates of N₂ production in freshwater aquifers are being compared to estimates of land-derived, ground-water transported nitrogen loads to coastal waters. Those comparisons will provide an important test of hypotheses regard-
ing the behavior of N during transport through watersheds and of N loading models used to manage coastal watersheds and coastal waters. In addition, the study will test for N2 production in mixing zones between fresh and saline ground water.

References Cited


Related Links

USGS Submarine Groundwater Discharge Program: http://coastal.er.usgs.gov/sgd/

Coastal Groundwater Geochemistry (WHOI): http://www.whoi.edu/science/MCG/groundwater/

Submarine Ground-Water Discharge and Its Influence on Coastal Processes and Ecosystems: http://soundwaves.usgs.gov/2004/06/research4.html


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