Reactive and Mixing Processes Governing Ammonium and Nitrate Coexistence in a Polluted Coastal Aquifer

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Abstract: A comprehensive hydrochemical, stable isotope and microbial analyses characterisation has been performed to evaluate the sources of groundwater, nitrogen pollution and degradation processes occurring in an industrial polluted coastal aquifer in the framework of a complex hydrodynamic system. The coexistence of ammonium and nitrate has been observed in almost all the investigated monitoring wells, reaching maximum values of 100 and 200 mg/L for both species. Chloride and potassium concentration coupled with groundwater stable isotopes data show the influence of local and urban recharge and the occurrence of seawater intrusion in areas near the coastline. δ15N–NH4+ values ranging between −4.9 and +14.9% suggest that different processes such as partial nitrification of ammonium, probably anammox activities and sorption, are occurring at the site. The isotope data for NH4+ also showed the existence of the remnant of an old fertilizer plume in the downgradient area. The nitrate isotope data ranging between +9 and +46% and +6 and +26% for δ15N–NO3− and δ18O–NO3−, respectively, suggest that nitrate content is attenuated by denitrification and probably anammox. The fast groundwater flow field is one of the reasons for the coexistence of NH4+ and NO3− in groundwater, since both compounds can penetrate the reducing zone of the aquifer. The influence of leakage of sewage pipelines on the aquifer cannot be discerned due to the complexities of the nitrogen attenuation processes, also influenced by pumping activities.

Keywords: nitrification; denitrification; stable isotopes; coastal aquifer; contamination sources

1. Introduction

Nitrate (NO3−) and ammonium (NH4+) are often recognised as major contaminants in both surface water and groundwater [1–3]. The causes can be multiple, including fertilizers leaching in agricultural environments [4–7], storm and sewage water spillages in urban environments [8–10], natural elevated background concentrations [11,12] and spillages of complex chemical mixtures below industrial plants [13,14]. Moreover, the nitrification of NH4+ into NO3− in oxic environments often leads to an increased NO3− concentration along the flow paths, up to values well beyond to the maximum water quality admissible levels [15]. Accordingly, the origin and transformation of NH4+ in groundwater is a key issue in water quality assessments. The nitrification process is kinetically
driven and depends on many factors, such as the availability of both electron donors and acceptors, salinity, pH and Eh [16]. Such peculiarities are exacerbated in aquifers, where partial/incomplete mixing between reactants often leads to a slowdown of the overall reaction kinetics [17] with respect to well-mixed bioreactors [18]. Indeed, partial/incomplete mixing can lead to the coexistence of thermodynamically unstable redox couples like \( \text{NH}_4^+ \) and \( \text{NO}_3^- \), especially in industrial areas where both these reactive nitrogen species have been or are still used and thus various pollution sources can be active at the same site [13,19]. Such a coexistence can trigger or slow down nitrogen transformation processes like nitrification or denitrification, and the degree of partial/incomplete mixing could vary in different parts of the site, for example when “pump and treat” remediation schemes are active. Thus, it is important to understand the processes that induce \( \text{NH}_4^+ \) and \( \text{NO}_3^- \) coexistence or the prevalence of one species over another. In addition, it is worth mentioning that \( \text{NO}_2^- \), an intermediate product of nitrification and denitrification, is present at a very small concentration in groundwater compared to nitrate and ammonium [20]. In recent years, nitrogen-stable isotopes are more often employed to assess different sources of pollution and nitrogen transformation in aquifers [21–26]. Although, in the abovementioned studies, \( \text{NO}_3^- \) was found to be the dominant nitrogen species in the oxic zone while \( \text{NH}_4^+ \) was the dominant nitrogen species in the anoxic zone, their coexistence has been often limited to narrow reactive fringes. However, few studies have been done on the mechanisms that control \( \text{NH}_4^+ \) and \( \text{NO}_3^- \) coexistence in large portions of the aquifer at the site scale [13,27].

This study aims to contribute to the understanding of nitrogen processes at industrial sites affected by different sources of nitrogen pollutants. The site under investigation is a coastal aquifer having an active hydraulic barrier designed to control the extent of contaminants plumes [19,28,29]. The approach included an evaluation of sources, distribution and degradation processes concerning reactive nitrogen species in the framework of a complex hydrodynamic system. The research tools include major ions and environmental isotopes that have been used to evaluate sources of groundwater and the sources and fate of nitrogen species. Those data have been coupled with total coliforms analysis, useful to evaluate the role of sewage leakage. The novelty of this study is to present a set of bio-geochemical analyses that provide information about the pollution sources of \( \text{NH}_4^+ \) and \( \text{NO}_3^- \) and to constrain their major biogeochemical reactions in a coastal aquifer.

2. Geological and Hydrogeological Setting

The geological and hydrogeological setting of a portion of an industrial site is the framework for the nitrogen investigation (Figure 1). The different nitrogen pollution sources, which include a fertilizer plant, an ammonia (\( \text{NH}_3 \)) production plant, and a sewage line, are also identified in Figure 1. Briefly, the unconfined aquifer is highly polluted by a range of organic and inorganic contaminants [14,19,28]. In this portion of the site, \( \text{NH}_4^+ \) and \( \text{NO}_3^- \) exhibited historical concentrations up to 3000 and 500 mg/L, respectively. The unconfined aquifer is formed by fine to coarse Quaternary sandy sediments, interbedded by silty clay lenses. The aquifer is characterised by a groundwater flow system influenced by regional and local recharge and also by seawater intrusion and leakage from a sewer pipeline near the coastline. The natural groundwater flow is perpendicular to the coastline, with an average hydraulic gradient of 0.6%. A complete description of the industrial site and its hydrogeological modelling and pollutants characterisation can be found elsewhere [19,28,29]. The presence of a “pump and treat” remediation system close to the shoreline, here called for simplicity the “hydraulic barrier,” highly affects the groundwater flow system, creating large capture zones and increasing the saltwater encroachment [14]. The pumping rate of the hydraulic barrier in this area is approximately 27 L/s and the hydraulic gradient reaches a maximum of 1%. Due to the site complexity, elevated salinity, and chemical gradients present in the unconfined aquifer, a detailed vertical characterisation of the aquifer via multi-level sampling (MLS) devices was also carried out to avoid cross-contamination and misleading results.

The investigated monitoring wells have been classified, for simplicity, into three zones: MLS located upgradient (UG) with respect to the hydraulic barrier, near (HB) the hydraulic barrier, and
downgradient (DG) with respect to the hydraulic barrier. This configuration allows us to identify the location and the composition of the sources of pollutants that affect the aquifer.

Figure 1. Top panel: Location map of the long screen monitoring wells used in the study including: MLS located upgradient with respect to the hydraulic barrier (diamonds), near the hydraulic barrier (circles), and downgradient with respect to the pumping wells (brown crosses), and related shallow monitoring wells sampled for coliforms analyses (brown circles with crosses). Pumping wells equivalent freshwater heads contours above sea level (blue lines). Various N pollution sources creating NH$_4^+$ and NO$_3^-$ plumes (red shaded areas): sewer pipeline, Fertilizer storage buildings and NH$_3$ plant. Red line shows the industrial site boundary. Bottom panel: Cross section A-A' (dashed line on the top panel) representing a simplified geological profile of the unconfined aquifer. Vertical exaggeration is 1:10.
3. Materials and Methods

3.1. Groundwater Sampling Technique and Chemical Analysis

Groundwater has been pumped to obtain depth–discrete samples, mimicking an MLS system, from standard fully-screened monitoring wells equipped with straddle inflatable packers (800-l Solinst) to obtain ion concentrations and nitrogen isotopes depth profiles. Samples were collected in November 2011 and April 2012, during groundwater sampling surveys. The packers' assemblage consisted of a 30 cm long sampling port, set in the middle of two inflatable packers, and a submersible centrifuge pump used for low flow purging. From each monitoring well, three to five samples have been collected at different depths, as described in Mastrocicco et al. [29] and Sbarbati et al. [19]. Samples were filtered through a 0.45 \( \mu \)m filter, stored in a cool box at 4 \( ^\circ \)C, and immediately transported in the laboratory. Groundwater samples collected for ion analyses have been analysed for the major ions (Cl\(^-\), K\(^+\), NH\(_4^+\), NO\(_3^−\), NO\(_2^−\)) using an isocratic dual pump ion chromatography ICS-2000 Dionex. The uncertainty of concentration results is generally lower than the detection limits. A flow-through cell has been employed to measure the in situ physical–chemical parameters (pH, electrical conductivity, Eh, dissolved oxygen) for each sampling depth, connected with a multi-parameter probe WTW Multi 340i (Weilheim, Germany), which includes: a SentiTix 41 pH combined electrode with a built-in temperature sensor for pH measurement, a combined AgCl–Pt electrode for Eh measurement and a Tetracon 325 4-electrode conductivity cell for EC measurements.

3.2. Groundwater Isotopes Analyses

The stable isotope analyses in water samples (\( \delta^{18} \)O and \( \delta^{2} \)H) have been performed in the Isotopes Laboratory of the University of Parma. The oxygen isotopic composition has been measured by the water–CO\(_2\) equilibration technique suggested by Epstein and Mayeda [30], either by manual preparation of the samples or using the Finnigan GLF 1086 (Burgwedel, Germany) automatic equilibration device [31]. The hydrogen measurements have been carried out by reducing the water sample to hydrogen by passage over hot zinc at about 520 \( ^\circ \)C according to the procedure described by Coleman et al. [32]. The isotopic measurements have been carried out using the Finnigan Delta S, Fisons Optima and Finnigan Delta Plus mass spectrometers. The isotopic results obtained are reported in terms of \( \delta \) units (% deviation of the isotope ratio from the international standard VSMOW, which is 0.0); the analytical error was \( \pm 0.1 \)% and \( \pm 1.0 \)% respectively for \( \delta^{18} \)O and \( \delta^{2} \)H [31].

3.3. Nitrogen Isotope Analyses

Nitrogen isotope analyses have been performed in the Environmental Isotope Laboratory of the University of Waterloo and in the Isotope Science Laboratory of the University of Calgary for NO\(_3^−\) isotope samples with relatively high salinity (more than 300 mg/L of Cl\(^−\)). Concerning \( \delta^{15} \)N–NH\(_4^+\) determination, a modified version of the acidified disk diffusion method [33,34] was adopted. Samples were placed in a 60 mL jar with 4 M (molar) solution of KCl (potassium chloride) and a phenolthalein indicator was added. Samples were basified with NaOH (sodium hydroxide) until the solution turned bright pink. A buffer solution of sodium tetraBorate was added and a Teflon packet containing an acidified quartz disc was added and sealed in the jar. After a period of 10 days all the ammonium in the sample should have volatilized into ammonia and been absorbed onto the acidified quartz disc as the Teflon is permeable to gas. Three internal \(^{15}\)NH\(_4^+\) isotope standards (\( \delta^{15} \)N = +0.6\%, +20.9\%, +35.2\%) were included in each diffusion run. All standards and samples have been run in duplicate or triplicate. Once diffusion was complete, the acidified disks were frozen and freeze-dried overnight and then stored in air-tight vials. The N isotopic composition of the diffusion disks containing the NH\(_4^+\) as (NH\(_4^+\))\(_2\)SO\(_4\) have been analysed using a Carlo Erba 1108 Elemental Analyser interfaced with a Delta plus–IRMS (Thermo Fisher Scientific, Milan, Italy). NH\(_4^+\)–N isotope ratios are reported in delta (\( \delta \)) notation relative to atmospheric N\(_2\). The uncertainty associated with the \( \delta^{15} \)N–NH\(_4^+\) analysis was generally lower than \( \pm 0.3 \)%.
the analyses for $\delta^{15}$N–NO$_3^-$ and $\delta^{18}$O–NO$_3^-$ have been performed in two different laboratories. Groundwater samples were frozen until arrival at laboratories and analysed approximately 20 days after their collection. The samples having relative low salinity were analysed using the silver nitrate (AgNO$_3$) method [34] adopted in the Environmental Isotope Laboratory of the University of Waterloo. Extraction of NO$_3^-$ was performed by passing samples through an ion exchange column, followed by HCl desorption and further precipitation of AgNO$_3$ by the addition of Ag$_2$O. The obtained AgNO$_3$ solution was filtered in two steps at 11 and 0.45 µm to remove the precipitated AgCl and then freeze-dried to obtain solid AgNO$_3$. Nitrogen gas for $^{15}$N analysis was produced by combustion in the presence of CuO, Cu, and CaO. For $^{18}$O analysis, nitrate oxygen was converted to CO$_2$ by combusting AgNO$_3$ with excess graphite. $^{15}$N and $^{18}$O were analysed in a VG Micromass spectrometer and in a VG Prism Micromass spectrometer respectively. The denitrifier method” [35–38] was used at the Isotope Science Laboratory of Calgary University for high-salinity samples; the method is based on the bacterial reduction of nitrate (NO$_3^-$) to nitrous oxide (N$_2$O) via a bacterium that lacks nitrous oxide reductase, so further reduction to N$_2$ does not occur. This enables simultaneous determination of both $\delta^{15}$N and $\delta^{18}$O values of the sample NO$_3^-$ by measuring the $\delta^{15}$N and $\delta^{18}$O values of the produced N$_2$. The bacterial strain Pseudomonas aureofaciens is used to reduce NO$_3^-$ to N$_2$O. The system is comprised of a home-made, 24-vial auto-sampler interfaced to a HP 6890 gas chromatogram with a PreCon® device (Wilmington, DE, USA) interfaced to a Finnigan Mat Delta + XL mass spectrometer. The accuracy and precision of $\delta^{15}$N and $\delta^{18}$O determinations based on a long-term record of analyses of ISL in-house reference (ISL-KNO3) are 0.3% and 0.7%, respectively.

3.4. Total Coliform Analysis

For total coliform analysis, groundwater samples were collected directly from pumping and monitoring wells in duplicate sterile 100-mL bottles containing sodium thiosulfate and stored at 4 °C in the dark until incubation for measuring the total coliforms. Samples were analysed for total coliforms in the laboratory of the Department of Life Sciences and Biotechnology of University of Ferrara, using the MPN-based Colilert™ assay (IDEXX Laboratories, Inc., Westbrook, ME, USA). The MPN solution of Hurley and Roscoe [39] was used to determine an MPN/100 mL and respective 95% confidence intervals. The detection limit based on duplicate measurements is 0.5 MPN/100 mL. For undiluted samples, the range of the method (>2400 MPN/100 mL) have been exceeded by two for total coliforms, thus appropriate dilution was needed to correctly quantify the total coliforms. Two sterile water blanks were analysed during each sampling period and were always negative for total coliforms.

4. Results and Discussion

4.1. In Situ Measurements

Electric conductivity (EC) and Eh measurements were done during sampling activities (Table 1). Electrical conductivity, which provides information about salinity patterns, varied from minimum values of 159 µS/cm in the upgradient areas to values greater than 50,000 µS/cm near the coastline. Some salinity stratification toward high EC values with depth is observed in the monitoring wells DG4 and DG5 (Table 1). The highest EC values are associated with the influence of seawater and potentially the contribution of salts from fertilizer plants. Eh values, which give information about aquifer redox conditions, range between −185 mV and 300 mV and most of the values are in the negative range, indicating that the groundwater is under anaerobic and reducing conditions.
Table 1. Physical–chemical parameters, major ion concentration values, groundwater and nitrogen-stable isotopes results for groundwater sampled via MLS monitoring wells (November 2011–April 2012), rainfall and urban water. Alphabetical order indicates increasing sampling depth.

| Samples ID | Elevation (m a.s.l.) | EC (µS/cm) | Eh (mV) | Cl (mg/L) | K (mg/L) | δ¹⁸O–H₂O | δD–H₂O | NH₄ (mg/L) | NO₃ (mg/L) | NO₂ (µg/L) | δ¹⁵N–NH₄ | δ¹⁵N–NO₃ | δ¹⁸O–NO₃ |
|------------|----------------------|------------|---------|-----------|----------|-----------|---------|-----------|-----------|------------|----------|-----------|-----------|
| UG1a       | −3.02                | 7800       | −24     | 880.0     | 498.0    | -         | -       | 293.8     | 214.0     | 52.12      | 13.8     | -          | -         |
| UG1b       | −7.02                | 10,200     | −60     | 1510.0    | 383.0    | -         | -       | 245.0     | 93.0      | 29.00      | 10.9     | -          | -         |
| UG1c       | −11.02               | 27,400     | −137    | 4300.0    | 395.0    | -         | -       | 239.0     | 233.0     | >10        | 3.5      | -          | -         |
| UG2a       | −0.61                | 168        | 3       | 452.0     | 23.6     | −5.9      | −36.8   | >0.1      | 17.0      | >10        | -        | 9.4       | 6.4       |
| UG2b       | −4.61                | 159        | 2       | 580.0     | 23.8     | −5.2      | −32.2   | >0.1      | 8.5       | >10        | -        | 13.7      | 6.7       |
| UG2c       | −10.61               | 313        | 2       | 580.0     | 21.3     | −6.5      | −42.3   | >0.1      | 6.9       | >10        | -        | 14.4      | 8.9       |
| HB1a       | −4.47                | 35,887     | −180    | 1730.0    | 66.0     | -         | -       | 1.0       | 7.4       | 22.20      | -        | -          | -         |
| HB1b       | −12.47               | 39,516     | −280    | 18,200.0  | 431.0    | -         | -       | 245.0     | 93.0      | 29.00      | -        | -          | -         |
| HB1c       | −11.02               | 27,400     | −137    | 4300.0    | 395.0    | -         | -       | 239.0     | 233.0     | >10        | 3.5      | -          | -         |
| HG1a       | −3.12                | 2245       | 276     | 386.0     | 23.0     | −5.2      | −32.2   | 6.5       | 2.4       | 25.10      | 14.9     | -          | -         |
| HG1b       | −7.12                | 2378       | 241     | 347.0     | 25.9     | -         | -       | 0.4       | 16.4      | >10        | -        | 45.8      | 26.3      |
| HG1c       | −11.12               | 2368       | 226     | 369.0     | 24.9     | −5.3      | −33     | 0.3       | 20.6      | >10        | -        | 39.9      | 22.3      |
| HG1d       | −19.12               | 2369       | 208     | 368.0     | 25.5     | -         | -       | 0.2       | 21.6      | >10        | -        | 39.8      | 20.7      |
| HG2a       | −7.52                | 36,523     | −68     | 3060.0    | 65.0     | -         | -       | 205.0     | 17.5      | >10        | −1.0     | -          | -         |
| HG2b       | −11.52               | 37,412     | −83     | 4680.0    | 119.0    | -         | -       | 569.0     | 1.2       | 17.30      | -        | -          | -         |
| HG2c       | −15.52               | 42,631     | −96     | 5000.0    | 131.9    | -         | -       | 627.0     | 1.9       | 15.00      | -        | -          | -         |
| HG2d       | −19.52               | 43,203     | −101    | 5600.0    | 142.4    | -         | -       | 769.0     | 1.2       | <10        | -        | -          | -         |
| HG3a       | −5.60                | 43,388     | −73     | 12,000.0  | 160.0    | −2.8      | −17.1   | 12.6      | >0.1      | 53.80      | −3.9     | -          | -         |
| HG3b       | −9.60                | 50,832     | −96     | 11,100.0  | 216.0    | -         | -       | 25.4      | >0.1      | >10        | −3.8     | -          | -         |
| HG3c       | −13.60               | 50,822     | −101    | 21,200.0  | 301.0    | 0.3       | 2.2     | 13.3      | >0.1      | >10        | −3.9     | -          | -         |
| HG3d       | −17.60               | 50,864     | −104    | 19,200.0  | 318.0    | -         | -       | 12.2      | >0.1      | >10        | −4.9     | -          | -         |
| HG3e       | −21.60               | 49,653     | −93     | 16,000.0  | 315.0    | 0.4       | 2.6     | 10.8      | >0.1      | >10        | −3.2     | -          | -         |
| HG4a       | −2.10                | 2441       | 123     | 344.0     | 33.9     | −4.8      | −27.8   | 19.3      | 14.9      | 3040.00    | 11.6     | 32.1       | 18.9      |
| HG4b       | −5.10                | 2506       | 125     | 377.0     | 37.9     | -         | -       | 20.3      | 20.5      | 1520.00    | 8.1      | 29.5       | 15.4      |
| HG4c       | −8.10                | 2504       | 129     | 2000.0    | 53.0     | −2.9      | −16.9   | 1.8       | 12.9      | 1070.00    | -        | 19.2       | 14.4      |
| HG4d       | −11.10               | 10,614     | 64      | 17,600.0  | 351.0    | -         | -       | 20.9      | 1.6       | >10        | −4.9     | -          | -         |
| HG4e       | −15.10               | 48,052     | −55     | 800.0     | 152.0    | -         | -       | 31.5      | 4.4       | >10        | 1.4      | -          | -         |
Table 1. Cont.

| Samples ID | Elevation (m a.s.l.) | EC (µS/cm) | Eh (mV) | Cl (mg/L) | K (mg/L) | δ¹⁸O–H₂O | δD–H₂O | NH₄ (mg/L) | NO₃ (mg/L) | NO₂ (µg/L) | δ¹⁵N–NH₄ | δ¹⁵N–NO₃ | δ¹⁸O–NO₃ |
|------------|----------------------|------------|---------|-----------|----------|-----------|---------|------------|------------|------------|-----------|-----------|-----------|
| Detection limit | 0.1   | 0.4       | 0.1     | 0.1       | 10       | 10        | 10      | 10         | 10         | 10         | 10        | 10        | 10        |
| DG4f       | −19.10        | 50,375     | −70     | 17,500.0  | 392.0    | 0.2       | 0.1     | 5.9        | 2.0        | >10        | −2.8      | −10       | −2.8      |
| DG5a       | −4.76         | 4074       | 301     | 180.0     | 15.0     | −5.2      | −31.5   | 189.0      | 31.7       | 123.30     | −0.2      | 15.9      | 9.5       |
| DG5b       | −8.76         | 4572       | 299     | 306.0     | 15.5     | −         | −       | 206.0      | 31.5       | 144.00     | 3.4       | 17.3      | 12.3      |
| DG5c       | −12.76        | 5132       | 263     | 199.0     | 15.3     | −         | −       | 192.0      | 35.4       | 142.21     | 2.3       | 13.4      | 8.2       |
| DG5d       | −16.76        | 28,115     | 68      | 840.0     | 31.5     | −5.1      | −30.2   | 107.9      | 30.7       | 135.07     | −1.0      | 15.3      | 11.8      |
| DG5e       | −20.76        | 36,629     | −25     | 493.0     | 23.4     | −5.1      | −29.9   | 60.4       | 22.4       | 149.00     | −0.9      | 24.8      | 14.6      |
| Rainfall   | −            | −          | −       | −         | −        | −5.0      | −28.0   | −          | −          | −          | −         | −         | −         |
| Urban water | −            | −          | −       | −         | −        | −8.6      | −51.0   | −          | −          | −          | −         | −         | −         |
4.2. Cl⁻ and K⁺ Concentration Patterns

Cl⁻ concentration ranges between 180.0 mg/L and 20,000.0 mg/L, and K⁺ between 15.0 mg/L and 500.0 mg/L. The high concentrations of K⁺ and Cl⁻ are associated with fertilizers and seawater, respectively, so the relationship between K⁺ and Cl⁻ concentrations for the selected MLS (Figure 2) has been used for a preliminary evaluation of the influence of different contributions (local aquifer, seawater and sewage leaking) on groundwater hydrochemistry. In Figure 2, the mixing area between the background groundwater and seawater is also plotted to visualise the unreactive mixing between the two end-members; a mixing area is depicted instead of a mixing line to account for the variability of both K⁺ and Cl⁻ in the end-members.

The upgradient samples, representing the background conditions (UG2), correspond to the lower end member; the higher end-members are represented by some samples of monitoring wells HBI and DG4. Most of the samples fall inside the mixing area, underlining the great variability in groundwater composition for this part of the aquifer, as confirmed by the wide range of electrical conductivity values. Inside the mixing area it is also possible to identify two different clusters of downgradient monitoring wells: the first one represented by DG3 and partially DG4 (deep depth), and the second one by DG1, DG5 and the remaining samples of DG4 (shallow depth). The first cluster is characterised by relatively high concentrations of both ions, suggesting a clear influence of seawater intrusion, confirmed by the electrical conductivity values; instead, the second cluster of downgradient MLS has relatively low K⁺ and Cl⁻ concentrations and lower electrical conductivity values, suggesting that the aquifer is locally recharged by leaking sewer pipelines. In addition, the flow and transport model simulations for this area of the site required a recharge amount of 200 mm/year near the sewer pipelines [14], as opposed to the rest of the site, where approximately 80 mm/year were necessary to calibrate the numerical model [28]. The most significant exception from the mixing pattern was observed in monitoring well UG1, which is characterised by a high K⁺ concentration, marking a distinct trend that could be associated with fertilizer leaching (possibly N–P–K fertilizers) (see Figure 1).

**Figure 2.** Plot of Cl⁻ versus K⁺ for MLS located upgradient (UG1, UG2), near (HBI, HB2) and downgradient (DG1 ÷ DG5) with respect to the hydraulic barrier (see Figure 1 for sampling site location). The grey shaded area shows the mixing between the background groundwater concentration and seawater.
4.3. Stable Isotope Data

Groundwater isotope data have been used to further illustrate the effect of seawater intrusion and local and urban recharge. Figure 3 (left panel) shows that most of the collected samples fall along the Global Meteoric Water Line (GMWL). Monitoring wells UG2, DG1 and DG5 are characterised by an isotopic signal similar to rainfall (local recharge), with values ranging between $-6.5\%$ and $-5.1\%$ for $\delta^{18}O$ and between $-42.3\%$ and $-29.9\%$ for $\delta^D$. Groundwater with isotope values over $-20.0\%$ for $\delta^D$ and $-2.0\%$ for $\delta^{18}O$ is separate from the GMWL and follows a mixing line with the seawater composition. Monitoring wells DG3 and DG4 clearly show this enrichment trend in both $\delta^{18}O$ and $\delta^D$, confirming the influence of seawater intrusion, as already suggested by $Cl^-$ concentration and electrical conductivity values. The effect of seawater intrusion on groundwater isotopic composition is also highlighted in Figure 3 (right plot), showing the vertical profiles of $\delta^{18}O$ for the same monitoring wells. For DG3 and DG4, it is evident that the isotopic signal changes with depth, starting from depleted $\delta^{18}O$ values in the shallower part of the aquifer and moving to more enriched values, having an isotopic signal like seawater, in the deeper part.

![Figure 3. $\delta^{18}O$–$\delta^D$ (left) and $\delta^{18}O$-depth (right) plots in samples analysed from MLS, located upgradient (UG2) and downgradient (DG1, DG3, DG4, DG5), respectively, with respect to the hydraulic barrier (see Figure 1 for sampling site location).]

4.4. Nitrogen Pollution Sources and Degradation Processes Characterisation

4.4.1. Concentration Patterns for Nitrogen Species and Coliforms

The concentration data show values ranging between 0.1 mg/L and 769.0 mg/L for $NH_4^+$ and 0.1 mg/L and 233 mg/L for $NO_3^-$ (Table 1). $NO_2^-$ has negligible concentration in most of the samples (<10 $\mu$g/L), except in well DG4, which shows values between 1.07 to 3.04 mg/L. These relative high nitrite concentrations could be associated with the partial nitrification of ammonium, associated with leakage of the sewage pipeline. The highest $NH_4^+$ values are observed in well DG2 (516.0–769.0 mg/L) and could be associated with leakage of the sewage line and/or could be part of the fertilizer plume. This monitoring well is characterised by a low $NO_3^-$ concentration (1.2–2.2 mg/L). The highest $NH_4^+$ and $NO_3^-$ concentration values are observed in well UG1, located downgradient to the fertilizer storage building and clearly related to fertilizer leakage from this facility. Background concentrations of $NO_3^-$ are usually very low, with an average value of 2.5 mg/L, while $NH_4^+$ is usually present at higher concentrations, with an average value of 6.5 mg/L [14].
The relationship between NH$_4^+$ and NO$_3^-$ concentrations in the selected MLS monitoring wells (Figure 4) highlights the coexistence of both nitrogen species in almost all the investigated monitoring wells. The existence of ammonium is associated with the partial nitrification of ammonium in the shallow unsaturated and saturated aerobic zone and the general reducing conditions of the groundwater in the aquifer. The coexistence of NH$_4^+$ and NO$_3^-$ in groundwater is well represented by MLS-UG1, which has high concentrations of both species between 100 and 200 mg/L, and negative Eh values (reducing conditions) recorded all along the depth profile (Table 1). UG1 is located close to the source areas (see Figure 1). The coexistence of NH$_4^+$ and NO$_3^-$ is also observed in other MLS samples, with a general and steep NO$_3^-$ concentration decrease with respect to NH$_4^+$ in MLS located near and downgradient the hydraulic barrier; this could be associated with degradation processes or mixing with other water sources depleted in NO$_3^-$.

![Figure 4](image_url)  
**Figure 4.** Plot of NH$_4^+$ versus NO$_3^-$ in samples analysed from MLS located upgradient (UG1, UG2), near the hydraulic barrier (HB1, HB2) and downgradient (DG1 ÷ DG5) with respect to the hydraulic barrier (see Figure 1 for sampling site location).

The only instances of the non-coexistence of nitrogen species are represented by monitoring wells UG2, DG2 and DG3. Specifically, monitoring well UG2, located at the fringe of the NH$_4^+$ plume, shows relatively high NO$_3^-$ concentrations (6.3 to 17.9 mg/L) with NH$_4^+$ below detection limits, associated with positive Eh values. DG2 and DG3, located downgradient of the hydraulic barrier, show high NH$_4^+$ (516.0–769.0 mg/L for DG2; 10.8–25.4 mg/L for DG3) and negligible NO$_3^-$ concentrations under reducing conditions. Those samples could belong to an old plume that has already migrated downgradient with respect to the actual source area, before the installation of the pump and treat system [14]. A comparison of NH$_4^+$ and NO$_3^-$ concentrations does not clearly identify the different sources of pollution and reactive processes occurring at the site, but highlights the co-existence of both NH$_4^+$ and NO$_3^-$ in large parts of the contaminant plume.
In addition, total coliform analyses, sampled in additional monitoring wells near the hydraulic barrier (distance lower than 3 m), highlight that sewer raw waters are present in the monitoring wells located near (distance between 10 and 20 m) the sewer pipeline (Table 2). The locations of sampling point are shown in Figure 1. The three corresponding pumping wells have coliform content below the detection limits due to the large amount of groundwater gathered from the lower layers of the aquifer where the pumps are located. The upgradient monitoring well, as expected, displays total coliforms below the detection limits.

Table 2. Total coliforms detection via Most Probable Number analysis, from Sbarbati et al. [19]. Monitoring wells and pumping wells locations are showed in Figure 1 (except the upgradient monitoring well).

| Sample Name                              | Value (MPN/100 mL) |
|------------------------------------------|--------------------|
| Upgradient monitoring well               | Below detection limits |
| Sewer pipeline                           | <100,000           |
| Monitoring well 1 at the hydraulic barrier | 145                |
| Monitoring well 2 at the hydraulic barrier | 300                |
| Monitoring well 3 at the hydraulic barrier | 8719               |
| Pumping well 1 of the hydraulic barrier  | Below detection limits |
| Pumping well 2 of the hydraulic barrier  | Below detection limits |
| Pumping well 3 of the hydraulic barrier  | Below detection limits |

Based on the information gained by major ion concentration values, groundwater isotopes analyses and total coliform analyses, it is evident that different water sources and nitrogen sources are active at the site. Additional insights regarding nitrogen sources and processes that affect nitrogen species in groundwater can be provided by the nitrogen isotopes analysis, performed on seven MLS (UG1, UG2, HB2, DG1, DG3, DG4, DG5).

4.4.2. Isotope Data on NH$_4^+$ and NO$_3^-$

The isotope composition of the NH$_4^+$ and NO$_3^-$ is controlled by the isotopic composition of the sources and transformation processes such as nitrification, denitrification and anammox [5,25,40,41]. The NO$_3^-$ isotopes data at the site show a wide range of values ranging between +9.4 and +45.8% and +6.4 and +26.3% for $\delta^{15}$N–NO$_3^-$ and $\delta^{18}$O–NO$_3^-$, respectively (Table 1). These values are much higher than the literature values for fertilizer and sewage [40] at the site, indicating that NO$_3^-$ is been affected by an attenuation process. The enrichment trend observed for $\delta^{15}$N–NO$_3^-$ and $\delta^{18}$O–NO$_3^-$ (Figure 5) is typically associated with denitrification [42], which is possibly due to the reducing condition of the groundwater. Data from UG2, DG1, DG4 and DG5 monitoring wells show that NO$_3^-$ in the aquifer is affected by different degrees of denitrification. The enriched $\delta^{15}$N pattern in residual NO$_3^-$ cannot rule out anammox also being involved in degradation processes at the site [8,19,42]. NH$_4^+$ also shows extremely variable $\delta^{15}$N–NH$_4^+$ values, ranging from −4.9 to +14.9% (Table 1). The most enriched (positive) $\delta^{15}$N–NH$_4^+$ values are observed downgradient of the fertilizer storage building (UG1), which is also characterised by the highest concentration of NH$_4^+$ and NO$_3^-$, and in the downgradient area near the coastline at the shallow depth in DG1 and DG4 wells, which are characterised by NH$_4^+$ and NO$_3^-$ concentrations less than 20.0 mg/L (Table 1). The most depleted $\delta^{15}$N–NH$_4^+$ concentrations are observed in some downgradient wells, HB2, DG3 and DG5, which showed comparable NH$_4^+$ concentrations to UG1 in the case of the DG5 well (Table 1).
Figure 5. Diagram of $\delta^{15}$N–NO$_3^-$ versus $\delta^{18}$O–NO$_3^-$ in samples analysed from MLS located upgradient (UG2) and downgradient (DG1, DG4 and DG5) with respect to the hydraulic barrier (see Figure 1 for sampling site location). Modified from Clark and Fritz [43].

The relationship between $\delta^{15}$N–NH$_4^+$ and NH$_4^+$ concentrations has been used to evaluate the origin and fate of NH$_4^+$ in the aquifer (Figure 5). Considering the upgradient area and taking into account that NH$_4^+$ fertilizers range between −7.4 and +6.6% [40], the highly enriched $\delta^{15}$N–NH$_4^+$ values correlated with high concentrations of NH$_4^+$ and NO$_3^-$ at UG1 indicate that partial nitrification should be responsible for the enriched $\delta^{15}$N values in NH$_4^+$. These data also show a vertical stratification, with enriched $\delta^{15}$N–NH$_4^+$ values (+10.8 and +13.8%) observed at shallow depths and more depleted values (+3.5%) at the deepest depths. It is well documented that nitrification tends to enrich the remaining NH$_4^+$ in $\delta^{15}$N [40,44]. The effect of partial nitrification on the isotopic composition of the NH$_4^+$ was also observed in the shallow part of a septic system plume [8]. The other process that can isotopically enrich NH$_4^+$ is anammox [13,41], but, unfortunately, the isotope data for NO$_3^-$, which could also provide evidence about the role of anammox in NH$_4^+$ transformation in one of the source zones, are not available at UG1.

Highly enriched $\delta^{15}$N–NH$_4^+$ values are also observed in the shallow part of the aquifer at wells DG1 and DG4, located below the hydraulic barrier and the sewage line (see Table 1 and Figure 6). Low values of 6.5 and 2.4 mg/L for NH$_4^+$ and NO$_3^-$ are observed in DG1. Comparable concentrations of NH$_4^+$ and NO$_3^-$ in the range of 14 to 20 mg/L are observed at DG4 and NO$_3^-$ is also characterised by enriched $\delta^{15}$N–NO$_3^-$ and $\delta^{18}$O–NO$_3^-$ values, associated with denitrification and anammox processes [42]. Concerning the most depleted $\delta^{15}$N–NH$_4^+$ values (−4.9 and +1.4%), these are observed at all depths in DG3 and DG5 and in the deepest depth at DG4 in the downgradient area. The groundwater was characterised by less than 25.4 mg/L of NH$_4^+$ and no NO$_3^-$ in the case of DG3, 1.8 to 20.9 mg/L for NH$_4^+$ and less than 20.5 mg/L of NO$_3^-$ in DG4. DG5 is characterised by high NH$_4^+$ (60.4 and 206.0 mg/L) and relatively high NO$_3^-$ (22.4 and 35.4 mg/L). The high NH$_4^+$ concentration observed at DG5 is comparable to the high concentration observed in UG1 in the source zone. These depleted $\delta^{15}$N–NH$_4^+$ values cannot represent sewage nitrogen since this nitrogen is characterised by $\delta^{15}$N values between +4.0 and +6.0% [8]. This type of $\delta^{15}$N–NH$_4^+$ value is
characteristic of NH$_4^+$ fertilizers and could be part of an old plume that was already present before the installation of the pump and treat system about 20 years ago. One process that could be partly responsible for the depleted $\delta^{15}$N-NH$_4^+$ values and low NH$_4^+$ concentration could be adsorption. It is well documented that during this process the remaining NH$_4^+$ gets depleted in $\delta^{15}$N [45]. Looking at the data in Figure 6, decreasing NH$_4^+$ concentrations seem to be correlated with a trend toward negative $\delta^{15}$N-NH$_4^+$ values (grey dashed arrow). The occurrence of adsorption at the study site is supported by the high cation exchange capacity of the aquifer sediments [14]. Dilution and dispersion processes could also be partially responsible for the decreasing trend in NH$_4^+$ concentrations. However, it is difficult to explain the depleted $\delta^{15}$N-NH$_4^+$ values accompanied by a high concentration of NH$_4^+$ observed at DG5 based only on adsorption. It is important to highlight that NO$_3^-$ coexists with NH$_4^+$ at DG5 and NO$_3^-$ samples were very enriched in $\delta^{15}$N-NO$_3^-$, indicating that NO$_3^-$ was affected by denitrification. Thus, a more plausible theory to explain the depleted $\delta^{15}$N-NH$_4^+$ values is that the NH$_4^+$ and NO$_3^-$ were both part of an old fertilizer plume that was already well developed before the installation of the pump and treat system at the site.

The other group, which is plotted in a different field to the rest of the samples, is formed by three samples (shallow depths at DG1 and DG4, Table 1) with NH$_4^+$ concentrations of 6.5 mg/L (DG1) and around 20.9 mg/L (DG4) and highly positive $\delta^{15}$N-NH$_4^+$ values (grey box in Figure 6) that could be ascribable to the partial nitrification of NH$_4^+$ and also to anammox [46]. NO$_3^-$ is also present at concentrations similar to NH$_4^+$ in these monitoring wells (Table 1). The NO$_3^-$ in DG4 also showed highly enriched $\delta^{15}$N-NO$_3^-$ and $\delta^{18}$O-NO$_3^-$ values (see Table 1 and Figure 5), indicating that NO$_3^-$ has been affected by denitrification and was involved in anammox processes [46]. These monitoring wells are located downgradient to the sewer pipeline and the groundwater stable isotopes data clearly

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**Figure 6.** Plot of NH$_4^+$ versus $\delta^{15}$N-NH$_4^+$ in samples analysed from MLS located upgradient (UG1), near (HB2) and downgradient (DG1, DG3, DG4, DG5), respectively, with respect to the hydraulic barrier (see Figure 1 for sampling site location).
show the influence of local recharge (Table 1), indicating that the nitrogen species can be associated with the leakage of sewage pipelines. Therefore, from nitrogen-stable isotopes analyses we have identified the main degradation processes acting at the site, namely partial nitrification, denitrification, adsorption and anammox. In addition, the coexistence of NH$_4^+$ and NO$_3^−$ at elevated concentrations in most of the analysed MLS monitoring wells is also favoured by the general reducing condition of the aquifer and the fast groundwater flow field. The latter allows the transport of oxidised species like NO$_3^−$ from the source (located in the vadose zone) into reducing zones within the aquifer, further triggering denitrification reactions.

5. Conclusions

The results of this study show that the combined use of hydrochemical, stable isotopes data and microbial analyses have provided valuable information to evaluate the pollutant sources and processes that affect reactive nitrogen species in an industrial coastal aquifer. The installation of the pump and treat system has significantly changed the natural flow system and the behaviour of contaminant plumes in the aquifer. The main focus of this study was to understand the processes that govern the nitrogen dynamics in the aquifer. The chemical and stable isotope data in groundwater show the influence of recharge and the occurrence of seawater intrusion in areas near the coastline. A key feature of the study site is the coexistence of NH$_4^+$ and NO$_3^−$. Redox parameter data (Eh) show that groundwater is under reducing conditions in most parts of the aquifer and flow velocity is elevated due to the pumping wells’ remediation system. The fast groundwater flow field is one of the reasons for the coexistence of NH$_4^+$ and NO$_3^−$ in groundwater, since both compounds can be mobilised in the reducing zone of the aquifer. The highly enriched $\delta^{15}$N–NH$_4^+$ values (+13.8 and +14.9%) observed at the site are associated with partial nitrification of NH$_4^+$, but anammox could also be partially responsible for these values. The NO$_3^−$ isotopes data showed that NO$_3^−$ is attenuated by denitrification, but anammox is also involved as an attenuation process for NO$_3^−$. The other feature of the site is the existence of NH$_4^+$, characterised by very depleted $\delta^{15}$N–NH$_4^+$ values (+1.4 and −4.9%). Based on the isotope and concentration patterns at the site, it is postulated that the depleted $\delta^{15}$N–NH$_4^+$ values represent the remnant of an old plume that was developed before the initiation of pumping by the hydraulic barrier about 20 years ago. The effect of adsorption on the isotopic composition of NH$_4^+$ also cannot be overlooked since there is evidence of NH$_4^+$ adsorption in sediments at the site. Due to the complexities of the nitrogen cycle disturbed by pumping wells, the effect of leakage from septic pipelines, in spite of the presence of coliforms, could not be resolved based on the isotope data on nitrogen compounds, except for one spot where the groundwater stable isotopes indicated local recharge, probably associated with leakage of the sewage pipelines. Summarising, the chemical and isotopic data have shown that partial nitrification of NH$_4^+$, denitrification of NO$_3^−$ and anammox have been involved in the transformation and attenuation of NH$_4^+$ and NO$_3^−$ at the study site. Evidence of the existence of an old fertilizer plume was also found. This study has clearly shown that a multidisciplinary approach is required to evaluate nitrogen cycling in industrial sites affected by multiple sources of contamination and seawater intrusion.

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