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1	Effects of nitrate contamination and seasonal variation on the denitrification and
2	greenhouse gas production in La Rocina stream (Doñana National Park, SW Spain)
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4	Germán Tortosa* ^a , David Correa ^a , A. Juan Sánchez-Raya ^a , Antonio Delgado ^b , Miguel A.
5	Sánchez-Monedero ^c and Eulogio J. Bedmar ^a
6	^a Departamento de Microbiología del Suelo y Sistemas Simbióticos, and ^b Departamento de
7	Geoquímica Ambiental, Estación Experimental del Zaidín, CSIC, 18080-Granada, Spain.
8	^c Departamento de Conservación de Suelos y Agua y Manejo de Residuos Orgánicos, Centro de
9	Edafología y Biología Aplicada del Segura, CSIC, 30100-Murcia, Spain.
10	
11	*G. Tortosa and D. Correa contributed equally to this article.
12	
13	Correspondence:
14	Germán Tortosa
15	Departamento de Microbiología del Suelo y Sistemas Simbióticos
16	Estación Experimental del Zaidín, CSIC
17	P. O. Box 419
18	18080-Granada, Spain.
19	
20	E-mail: german.tortosa@eez.csic.es
21	Tel.: + 34-958181600 (ext. 286)
22	Fax: +34-958-181609
23	

26 Climatic influence (global warming and decreased rainfall) could lead to an increase in the ecological and toxicological effects of the pollution in aquatic ecosystems, especially 27 28 contamination from agricultural nitrate (NO₃) fertilizers. Physicochemical properties of the surface waters and sediments of four selected sites varying in NO3⁻ concentration along La 29 30 Rocina stream, which feeds Marisma del Rocio in Doñana National Park (South West, Spain), 31 were studied. Electrical conductivity, pH, content in macro and microelements, total organic 32 carbon and nitrogen, and dissolved carbon and nitrogen were affected by each the sampling site 33 and the sampling time. Contaminant NO_3^{-1} in surface water at the site with the highest NO_3^{-1} concentration (ranged in 61.6-106.6 mg L⁻¹) were of inorganic origin, most probably from 34 35 chemical fertilizers, as determined chemically (90% of the total dissolved nitrogen from NO_3^{-}) and by isotopic analysis of δ^{15} N-NO₃. Changes in seasonal weather conditions and hydrological 36 37 effects at the sampling sites were also responsible for variations in some biological activities (dehydrogenase, β-glucosidase, arylsulphatase, acid phosphatase and urease) in sediments, as 38 39 well as in production of the greenhouse gases CO₂, CH₄ and N₂O. Both organic matter and NO₃⁻ contents influenced rates of gas production. Increased NO₃⁻ concentration also resulted in 40 41 enhanced levels of potential denitrification measured as N₂O production. The denitrification 42 process was affected by NO_3^- contamination and the rainfall regimen, increasing the greenhouse 43 gases emissions (CO₂, CH₄ and especially N₂O) during the driest season in all sampling sites 44 studied.

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46 *Keywords*:

47 Doñana National Park, surface waters and sediments, nitrate contamination, greenhouse gases,

48 biological activities, denitrification

51 Anthropogenic influence on the biogeochemical N cycle can produce important alterations 52 of the cycle leading to concomitant environmental risks such as increased concentration of 53 greenhouse gases, acidification of soils, streams and lakes, transfer of nitrogen through rivers to 54 estuaries and coastal oceans, accelerated losses of biological diversity and human health and 55 economy problems (Vitousek et al., 1997; Galloway et al., 2008; Mulholland et al., 2008). In 56 aquatic ecosystems, water acidification, eutrophization, including occurrence of toxic algae, and 57 toxicity of ammonia (NH_3) , nitrite (NO_2) , and nitrate (NO_3) are the three major environmental 58 problems due to inorganic nitrogen pollution (Camargo and Alonso, 2006). Furthermore, 59 increasing global warming and decreased rainfall in some continental areas may increase 60 ecological and toxicological effects of this type of environmental contamination (Camargo and 61 Alonso, 2006). Abuse in utilization of nitrogenous chemical fertilizers have been shown to 62 enhance emission of carbon dioxide (CO₂), methane (CH₄), and nitrous oxide (N₂O) greenhouse 63 gases (Thornton and Valente, 1996; Merbach et al., 1996, 2001; Davidson et al., 2000; Liu and 64 Greaver, 2009). In addition to chemical fertilizers, release of greenhouse gases to the atmosphere can be induced by changes in precipitations, temperature, seasons, drought, regional 65 66 deforestation, global warming, and El Niño events (Christensen et al., 1990: Smith et al., 2003; 67 Davidson et al., 2004).

Wetlands are among the most important ecosystems on Earth because of their role in regulating global biogeochemical cycles. Climate change and anthropogenic effects may have significant impacts on coastal and inland wetlands (Mitsch and Gosselink, 2007; Olías et al., 2008). Accordingly, physicochemical and biological monitoring are needed for assessment of ecological risks due to freshwater pollution and to provide maximal information for adequate

73 protection of aquatic ecosystems (Camargo, 1994). Several authors and reports have shown that 74 NO₃⁻ contamination of soils and surface and groundwater is becoming more intense and frequent due to the great consume of inorganic nitrogen, mainly nitrate and ammonium salts from 75 76 agrochemicals, of the intensive farming (Spalding and Exner, 1993; European Comission, 2002). 77 Denitrification is the biological process by which NO_3^- can be transformed into molecular 78 nitrogen (N_2) via formation of NO_2^- and nitric oxide (NO). Thus, it represents the major pathway by which NO_3^- can be removed from soils and waters to avoid NO_3^- accumulation and 79 80 contamination. And yet, incomplete denitrification results in production of the greenhouse gases 81 NO and N_2O (Aulakh et al., 1992; Conrad, 1996; Groffman et al., 2006).

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83 The European directive 91/676/CEE concerning NO₃⁻ contamination from agricultural 84 sources defines the so called "nitrate vulnerable zones" as reference areas of special 85 environmental protection to prevent soil and water nitrate contaminations. An example is Doñana 86 National Park (DNP), one of the most important wetlands in Europe covering an area around 60,000ha in a marshy area of SW Spain, in the estuary of the Guadalquivir River. These water 87 88 flows are susceptible of NO_3^- contamination from small urban areas in the surrounding of the 89 park and agricultural practices allowed in the ecotone, where organic farming of strawberries and 90 rice is common. This area is the most fertile and productive zone of Doñana as a result of its 91 permanent humidity and of the fertilization it receives from the animals either living there or 92 crossing it (Suso and Llamas, 1993).

93 Several authors have noted that surface and groundwater of DNP wetland are becoming 94 polluted during the last 20 years. Suso and Llamas (1993) remarked that some wetlands and small 95 streams could be depleted by groundwater extraction for agricultural reclamation, affecting 96 negatively the quality of surface and groundwater. Olías et al. (2008) evaluated the water quality

of the Almonte-Marismas aquifer (upon which DNP is located) and showed that it was affected 97 98 by pollution of both agricultural and urban origins. They detected some shallow points located in the agricultural zones with high concentrations of NO_3^- and sulphates (SO_4^{2-}) from fertilizer 99 100 pollution. Finally, Serrano et al. (2006) reviewed the aquatic systems of DNP and they focused 101 on processes affecting water quality. They noted that there has been a considerable increase of 102 NO₃ concentration in the water-flows of La Rocina and El Partido streams during the past 103 decade, probably due to the increase in cultivated land and fertilizer applications. They advise 104 that the influence of this pollution on the eutrophication of the nearby marshes should not be 105 overlooked.

106 Our research aim was to evaluate the anthropogenic (especially from agriculture) and 107 seasonal influence in La Rocina stream and how it could be affected by the physicochemical and 108 biological characteristics of the surface water and its aquatic sediments, focusing in the NO_3^- 109 contamination and its influence on the greenhouse gas production (CO₂, CH₄ and N₂O) and the 110 denitrification process.

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112 **2. Materials and methods**

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	114	2.1. D	escription	and	selection	of san	npling	sites
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In 1982, DNP was declared a Reserve of the Biosphere by UNESCO and a Wetland of International Interest per RAMSAR Convention. DNP wetland has three important surface water inputs (Arambarri et al., 1996; Serrano et al., 2006). Two natural streams called La Rocina and El Partido, in the north edge of the park, and the Guadiamar River, wich represents the main water input of the wetlands and suffered the Aznalcollar mine spill in 1998 (Cabrera et al., 1999; 121 Grimalt et al., 1999; Sierra et al. 2003; Olías et al., 2005).

122 The study was performed on La Rocina stream, located in the north of the DNP wetland, 123 which is one of the main natural streams feeding El Rocío marsh (Fig. 1). Selection of sampling 124 sites was based on their NO₃⁻ content (in-situ measurements using a Nitrate Test Kit, CHEMetrics 125 Inc.) after screening of more than 25 points along the course of La Rocina basin (462 km²) at the 126 different sampling times. Four sampling sites along the course of La Rocina stream differing in 127 their NO_3^- concentration were selected (Fig. 1). The Universal Transverse Mercator (UTM) 128 coordinates for the sites were as follows: 29S 0718632, 4114294 for the lagoon of Palacio del 129 Acebrón (S1); 29S 0717797, 4113881 for the small stream Arroyo de la Cañada (S2); 29S 130 0722653, 4111704 for the junction between the stream and the marsh called Vado de la 131 Canariega (S3); and 29S 0723654, 4111088 for the El Rocío marsh (S4). S1 and S2 had the 132 lowest and the highest NO_3^- concentration, respectively, whereas the values detected in S3 and S4 133 were between those detected in S1 and S2. Also, differences on hydrological morphology (S1: a 134 small stream, S2: a lagoon, S3: the union between a stream and a marsh, S4: a marsh) and in 135 riparian vegetation (not present in S3 and S4) were noted.

Samples were taken in October 2008 (T1) and January (T2), April (T3) and July (T4) 2009
in order to represent the pluvial regimen (dry and wet). Rainfall, relative humidity and air
temperature were collected from the Manecorro RM1 meteorological station, which belongs to
the Singular Scientific and Technological Installation (<u>http://icts.ebd.csic.es/</u>) of Doñana National
Park located about 200 m away from S3, and from Estación Manual Palacio de Doñana (EM05,
<u>http://www-rbd.ebd.csic.es</u>).

- 143 2.2. Physicochemical properties
- 144

Four replicates of the surface waters (approximately 1-2 meters from the shore in streams and 3-4 meters in lagoons for each replicate) and semi disturbed sediments (0-10 cm from the upper layer using an EIJKELKAMP Peat sampler) were taken at each sampling site, placed in a portable fridge and processed in the laboratory within 24 h of sample collection. Subsets of samples from the sediments were lyophilized and kept frozen at -20 °C until use.

150 In water samples, pH and electrical conductivity (EC) were analyzed using a Basic 20 151 Crison pHmeter and a Basic 525 Crison conductimeter at the laboratory, respectively. After 152 filtration through 0.45 µm filters, dissolved organic carbon (DOC) and total dissolved nitrogen 153 (TDN) were determined by using an automatic Shimadzu TOC-VCSN analyser. NO_3^- and NO_2^- 154 concentration were estimated by ion chromatography (HPLC) using an IC-Pac anion HC (Waters) column at the facilities of Servicio de Instrumentación (EEZ-CSIC). NH_4^+ was 155 156 determined by a colorimetric method based on Berthelot's reaction (Kempers and Zweers, 1986; 157 Sommer et al. 1992), adding sodium citrate to complex divalent cations. Macro and 158 microelements (P, K, Ca, Mg, Na, S, Fe, Cu, Mn, and Zn) were determined by Inductively 159 Coupled Plasma Optical Emission Spectrometry (ICP-OES) using an IRIS Intrepid II XDL 160 (Thermo Fisher Scientific Inc.).

161 Texture of the sediments was determined in fresh samples according to the Spanish Official 162 Methods for Soils and Waters (MAPA, 1974). NH₄⁺, after 2 h extraction 1:20 (w:v) with 2 N 163 KCl, and water extracted (1:20, w/v) NO₃⁻ and NO₂⁻ were also analyzed in fresh samples as 164 indicated above. All other assays were performed in ground samples (0.2 mm) after 165 lyophilization. pH and EC were measured after water extraction (1:5, w:v) for 2 h. Total organic 166 carbon (TOC) and total nitrogen (TN) were determined using a LECO TruSpec CN Elemental 167 Analyser. DOC and TDN were obtained after 2 h water extraction 1:20 (w:v) and estimated as 168 indicated for surface waters. Macro and microelements were analyzed by ICP-OES after

- 169 microwave digestion with a mixture (1:1) HF:HCl. Organic nitrogen (N_{ORG}) was calculated as the 170 difference between either TDN in surface waters or TN in sediments, and the content in inorganic 171 nitrogen (N_{INORG}), considering N_{INORG} as $N-NO_3^- + N-NO_2^- + N-NH_4^+$.
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- 173 *2.3. Enzymatic analysis in sediments*
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A selection of some enzymatic activities related to the metabolic activity and the main 175 176 biogeochemical cycles were measured in the freeze-dried sediments. Dehydrogenase was used as 177 an estimation of overall microbial activity, β -Glucosidase as the enzyme that catalyses the 178 hydrolysis of disaccharides (C cycle), Arylsulphatase as a measure of the enzymes catalysing the 179 hydrolysis of organic sulphate esters (S cycle), Acid phosphatase as a measure of the enzymes 180 responsible for the hydrolysis of phosphate esters (P cycle), and Urease which catalyses the 181 hydrolysis of urea to CO₂ and NH₃ (N cycle) Dehydrogenase was determined according to Garcia 182 et al. (1997), β-Glucosidase, arylsulphatase and acid phosphatase, were determined as described 183 by Tabatabai (1982) and urease activity was determined according to Kandeler and Gerber et al., 184 (1988). Briefly, these techniques were based on a controlled incubation of the sediments after 185 2-*p*-iodophenyl-3-*p*-nitrophenyl-5-tetrazolium adding the initial substrate (INT: for 186 dehydrogenase, pNG: 4-nitrophenyl-beta-D-glucopyranoside for β-Glucosidase, PNPS: p-187 nitrophenyl sulphate for arylsulphatase, pNPP: 4-nitrophenyl phosphate for acid phosphatase and 188 urea for urease activity respectively) and measuring the ending product of each enzyme reaction 189 colorimetrically (INTF: iodonitrotetrazolium formazan for dehydrogenase, pNP: p-nitrophenol 190 for β -Glucosidase, arylsulphatase, acid phosphatase and NH₄⁺ (measured as described above for 191 water and sediment samples) for urease activity.

195 The emission of CO_2 , CH_4 and N_2O were measured after 24 hour aerobic incubation (25°C) 196 of 20-30 g of the freshly collected sediments in 125 mL glass bottles. Gas concentrations were 197 analysed in the headspace by a Varian 4900 Gas Chromatograph with a PoraPlot Q column (10 m 198 length, 0.15 mm internal diameter) and thermal conductivity detector (TCD). Denitrifying 199 Enzyme Activity (DEA) and Denitrification Potential (DP) were also determined in the fresh sediments using an acetylene inhibition technique adapted from Šimek et al., (2004). DEA is a 200 201 measure of denitryfing enzymes in the sediment and reflects wether the environmental conditions 202 of the sediments at the moment of sampling would induce the activity of the denitrifying bacteria, 203 whereas DP represents a long term denitrification potential, allowing the maximum regrowth of 204 denytrifying bacteria (Tiedje, 1994). DEA was determined using an anerobic slurry prepared by 205 mixing 25 g moist sediment and 25 ml of a solution containing 1 mM glucose, 1 mM KNO₃ and 1 g L⁻¹ chloramphenicol (to prevent protein sinthesis and growth) in a 125 mL glass bottle. The 206 207 headspace was evacuated and flushed four times with He and 10 ml of acetylene were added. The 208 samples were shaken at 25°C and the concentration of N₂O was measured in the headspace after 209 30 and 60 min of incubation by gas cromatography, as previously described. DEA was calculated 210 from the N₂O increase during a half an hour incubation (60–30 min) and using the Bunsen 211 coefficient for the N₂O dissolved in water. DP was determined by mixing 6 g moist sediment 212 with 5 mL of a solution containing 1 mM KNO₃ and 1 mM glucose in a 125 mL glass bottle. 213 After evacuating and flushing the headspace four times with He, 10 ml of acetylene were added 214 and the samples were incubated at 25°C during 48 hours. DP was calculated from the N₂O 215 increase in the headspace after the second day of incubation and using the Bunsen coefficient for 216 the N₂O dissolved in water.

218 2.5. Isotope measurements

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 δ^{15} N of NO₃ was determined following the methodology described by Silva et al., (2001) with 220 221 modifications. Water samples (10–30 l) were first filtered through Whatman filter paper and then 222 passed through 0.45 µm filters (High Capacity GWV, Groundwater Sample Filter). Possible 223 interferences from sulphate and phosphate in the samples were eliminated by adding an excess of BaCl₂, and dissolved CO₂ was removed by adding HCl and gentle heating. Water samples were 224 225 then eluted through a cation exchange resin (AG 50W X8 100-200 mesh, Bio-Rad) to remove dissolved organic matter and the excess of Ba²⁺, and passed through an anion exchange resin 226 227 (AG1 X8 100-200 mesh, Bio-Rad) to retain NO₃. Finally, nitrate was eluted from the column by 228 adding 1N HCl, and the solution containing HNO₃ and HCl was neutralized with Ag₂O (Merck). 229 The resulting AgCl precipitate was removed by filtration (0.45 mm membrane filter) leaving only Ag^+ and NO_3^- in solution. The solutions were freeze dried yielding a pure, dry AgNO₃ precipitate. 230 231 Nitrogen isotope ratios and total nitrogen contents of AgNO₃ precipitates were determined 232 by thermal de-composition in a Carlo Elba NC1500 elemental analyser on line with a Delta Plus 233 XL (ThermoQuest) mass spectrometer (EA-IRMS). The overall precision of analyses was ± 0.1 ‰ for $\delta^{15}N$. The stable composition is reported as δ values per mil: $\delta = (R_{sample}/R_{standard} - 1) x$ 234 1000, where $R = {}^{15}N/{}^{14}N$ for $\delta^{15}N$. Nitrogen contents of the AgNO₃ samples were typically ~ 8%. 235 indicating that no major contaminants were present in the precipitate. Commercial N₂ was used as 236 237 the internal standard for the nitrogen isotopic analyses, contrasted with the international standard. δ^{15} N values for all samples were normalized against internationally accepted reference materials 238 (IAEA N1, $\delta^{15}N = +0.4$ ‰, IAEA N2, $\delta^{15}N = +20.3$ ‰). The nitrogen isotope ratios of AgNO₃ 239

generated from dissolved IAEA-NO-3 potassium nitrate were within + 4.65 ‰ (n = 12), similar to the accepted value. Duplicate nitrogen isotope ratio determinations on AgNO₃ from laboratory nitrate samples were performed with a precision generally better than ± 0.2 ‰. δ^{15} N values are reported with respect to air.

245 2.6. Statistical analysis

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Differences between the different physicochemical parameters were checked out using the analysis of variance (ANOVA) and the Tukey post-hoc test at P<0.05. Pearson coefficients were calculated to obtained correlation between variables using the SSPS 17.0 program for Windows XP. A principal component analysis (PCA) was performed to analyse relationships among parameters concerning physicochemical characterization of the sediments, their enzymatic activities and gas production.

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254 **3. Results**

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256 3.1. Meteorological data

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Total rainfall at DNP during the period of study was 299.9 mm, a value which is lower than that of 477.5 mm, which represents the mean rainfall for the previous 5 years. Main rainfall was registered in October 2008 (119.25 mm) and March-April 2009 (49.49 and 30.90 mm) respectively, coinciding with T1 and T3 sampling times. T2 (January 2009) presented 9.3 mm and T4 (July 2009) 0.11 mm, being the driest season. This pluvial regimen affected water dynamic in the sampling places studied especially at T4, transforming the stream and lagoon waters sampled in swamps (especially in S3 that presented a high eutrophization rate). Relative
humidity was higher in autumn and winter seasons (T1: 73.79 % and T2: 79.69 %) than in spring
and summer seasons (T3: 65.38 % and T4: 48.68 %), decreasing with air temperature (T1:
18.04°C, T2: 9.73°C, T3: 14.55°C and T4: 25.31°C).

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269 3.2. Surface waters: Physicochemical characterization and isotopic analysis.

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Surface water showed in general a slightly basic pH values, especially in S2, S3 and S4 271 272 (average values of 7.94 and 7.83 in S2, S3 and S4 respectively, Table 1a). EC was related directly 273 to total K, Ca, Mg, Na and S concentration in waters, and also to the sampling season. EC values typically varied within the range from 0.12 and 1.22 dS cm⁻¹ depending on the sampling season. 274 275 The highest EC values were generally recorded during rainy the season (T1), with the exception of the large EC value (3.11 dS cm⁻¹) registered in S3 during the driest season (T4), due to the 276 277 eutrophization caused for the swamp water. In general, soluble organic matter was high at T1 (S1: 17.7, S2: 28.3 and S4: 77.7 mg L^{-1} of DOC) for all sampling sites studied with the exception 278 of S3, that presented 135.2 mg L^{-1} of DOC at T4, due to the high water eutrophication (the P 279 280 concentration at this location was the highest value for all sampling sites and seasons) which 281 produced an elevated suspended algae content (green water colour by visual observation).

S2 presented higher TDN concentrations (in the range 21.6 and 9.4 mg L^{-1}) than the other three locations S1, S3 and S4 that presented an overall of 0.5, 6.2 and 2.3 mg L^{-1} respectively (Table 1b). In S3 at the driest season (T4), TDN showed a value of 18.1 mg L^{-1} especially due to the high NH₄⁺ content (Table 1a and b). S2 presented a large NO₃⁻ concentration (61.6, 106.6, 101.6 and 68.8 mg L^{-1} at T1, T2, T3 and T4 respectively) respect to the other sampling sites studied (less that 6 mg L^{-1} , Table 1a). These NO₃⁻ concentrations represented between 90 and 97% of the TDN of the surface waters (Table 1b). Also, this fact was noticed in DOC/TDN ratio values, being smaller in S2 (average of 0.6) than in S1, S3 and S4 (23.7, 12.9 and 10.2 respectively). These sampling sites presented an important organic nitrogen fraction (average of 74.6, 68.9 and 50.1% respectively), not in S2 that was predominantly inorganic (average of 92.4% of NO₃⁻ respect to TDN content).

293 With the procedure used in this study, isotopic analysis of N-NO₃⁻ could be carried out only in S2, the site with the highest NO₃⁻ concentrations. Values of δ^{15} N ranged from -1.6 ‰ to +6 ‰ 294 (AIR) with an average of -0.78 ‰ (AIR). This relatively low value, closed to that of the 295 296 atmosphere air, indicates that contaminant NO₃⁻ were of inorganic origin because atmospheric air is used for their synthesis (Vitoria et al. 2004). Moreover, since mean average values of $\delta^{15}N$ for 297 298 most inorganic Spanish fertilizers vary between -1 and +2 ‰ (AIR), being the total range 299 between -4 and +6 ‰ (Otero et al. 2005), the stable isotopes of nitrogen indicate an origin related 300 with fertilizers used in agricultural practices.

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302 *3.3. Sediments: Physicochemical characterization, enzymatic activities and gas production*

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304 Sand constituted more than 85% of the components of the sediments. According to the 305 corresponding contents in clay and silt, S1 and S2 were classified as loamy sand sediments, those 306 from S3 as sand, and sandy loam for S4. Similar to surface waters, values of pH, EC and content 307 in macro and microelements in sediments from the four sampling sites varied both among the 308 sites and with the sampling time (Table 1b). Despite these differences, the values of TOC and TN 309 were always higher in S1 than in the remaining sampling sites (Table 2b). Regardless of the 310 sampling sites and sampling times, more than 90% of the nitrogen content in the sediments was 311 of organic origin and, accordingly, the greatest values of DOC and TDN were also found in S1 (Table 2b). Similarly, β -glucosidase, dehydrogenase, urease acid phosphatase and arylsulphatase activities varied greatly with both the sampling sites and the sampling times (Fig. 2). Whereas S4 showed the highest values of dehydrogenase (mean average value of 10.26 µg INTF g⁻¹ h⁻¹), acid phophatase (mean average value of 23.5 µg *p*NP g⁻¹ h⁻¹), urease (mean average value of 600 µg *p*NP g⁻¹ h⁻¹) and arylsulphatase (mean average value of 400 µg *p*NP g⁻¹ h⁻¹) activities, S1 was the site with the greatest values of β-glucosidase activity (mean average value of 90 µg *p*NP g⁻¹ h⁻¹) (Fig. 2).

319 The mean average value of CO_2 produced at sampling sites S1, S2, S3 and S4 were 112, 89, 13, and 41 μ g C-CO₂ g⁻¹ day⁻¹, respectively (Fig. 3). Methane production from the sediments was 320 also higher in S1 (mean average value of 9.7 μ g C-CH₄ g⁻¹ day⁻¹) than in S3 and S4 (mean 321 average values of 0.2 and 0.3 μ g C-CH₄ g⁻¹ day⁻¹, respectively). Methane production occurred in 322 S2 was in general low. Unexpectedly, values for S2 detected at T3 were the highest for all 323 samples analyzed (34.9 μ g C-CH₄ g⁻¹ day⁻¹) (Fig. 3). S2, the site with the highest NO₃⁻ contents in 324 325 its surface water, and S4 showed maximal values of N₂O production with mean average values of 500 and 310 ng N-N₂O g⁻¹ day⁻¹, respectively. Potential denitrification as assayed by DEA and 326 327 DP showed that S1 and S2 have the highest potential for denitrification compared to S3 and S4. 328 The highest values for DEA were obtained during the driest season (T4) with 1115, 2246 and 719 ng N-N₂O $g^{-1} h^{-1}$ in S1, S2 and S4 respectively. Despite fluctuations at the sampling times, S1 and 329 330 S2 also showed maximal values of DP with mean average values of 218, 164, 58 and 89 µg N- $N_2O g^{-1} d^{-1}$ for S1, S2, S3 and S4, respectively. 331

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Pearson correlation matrix revealed that TOC, TN, NH_4^+ , DOC and TDN were positive and

³³³ *3.4. Statistical analysis*

336	significantly ($p \le 0.01$) correlated with CO ₂ production and with β -glucosidase activity (Table 3).
337	After PCA analysis, except for NO_3^- and pH, the remaining parameters analyzed clustered in 3
338	main groups (Figure 4a). The first cluster contained most of the parameters related with the
339	organic fraction of the sediments (TOC, TN, DOC, TDN, NH_4^+ , CO ₂ , CH ₄ , DP and β -glucosidase
340	activity), the second cluster included the inorganic components (K, Ca, Mg, Fe, Cu, Mn and Zn),
341	and the third cluster was composed of the enzymatic activities dehydrogenase, acid phophatase,
342	urease and arylsulphatase. The first principal component (PC1) explained 36.79% of the total
343	variance of the data, whereas the second principal component (PC2) was responsible for 20.34%.
344	According to these two axes, the sampling sites were ordered as a function of the three clusters
345	mentioned above (Figure 4b). Consequently, S1 showed the highest values for the organic
346	fraction and S4 for the enzymatic fraction. Nevertheless, sizes of the clusters indicated that
347	seasonal variation affected much more to S1 and S4 than to S2 and S3.

349

350 **3. Discussion**

351

352 Although to a different extent depending on the sampling site, physicochemical properties 353 of the surface waters sampled along La Rocina stream were influenced by seasonal variation 354 (especially rainfall regime). These results agree with those of Espinar and Serrano (2009) which 355 indicate that development of temporary wetlands in DNP are influenced by climate and geology 356 of the region. This is especially important in wetlands located in semiarid areas such as the South 357 of Spain, where the climate is unpredictable and produces a wide range of hydrological 358 conditions (Serrano et al., 2006). Thus, it is possible that rainfall, evaporation, groundwater 359 discharge, biogeochemical interactions at the sediment-water interface affected chemical 360 composition of the surface waters along the course of La Rocina stream. Hydrological dynamic at 361 each sampling site was affected by the sampling time. That was clearly visible in S3 at T4, the 362 driest sampling time, where stream waters were transformed into swampy waters, and in S4 363 where desiccation almost emptied the El Rocio marsh.

364 Several authors have reported continuous increases in pollution (Suso and Llamas, 1993; 365 Olías et al., 2008), and more precisely in NO₃⁻ content, in surface- and ground-waters of DNP 366 during the last two decades (García-Quesada et al., 1987, Serrano et al., 2006). Recently, contamination due to NO_3^{-1} and SO_4^{-2-1} in the Doñana aquifer has been linked to utilization of 367 368 agrochemicals during the agricultural practices that take place in the ecotone of the Park (Olías et al., 2008). Values of NO₃⁻ content in S2 were higher than the 50 mg L⁻¹ defined by the European 369 370 directive 91/676/CEE as the upper limit for NO_3^- contamination from agricultural sources 371 (European Comission, 1991). At that site, concentration of inorganic N represented more than 372 90% of the TDN. In this study, based on isotopic analysis of the contaminant NO₃⁻ in surface 373 waters of La Rocina stream, we show that, at least in S2, they were of inorganic origin, more 374 probably from chemical fertilizers.

375 The interpretation of the nutrients dynamic in aquatic ecosystems could be biased by the 376 strong effects of hydrology on physicochemical (Espinar and Serrano, 2009). For that, 377 microbiological processes involved in the principal biogeochemical cycles are needed 378 (Faulwetter et al., 2009). Soil microorganisms mediate many processes that are of particular 379 interest in freshwater wetland ecosystems where nutrient cycling is highly responsive to 380 fluctuating hydrology and nutrients and soil gas releases may be sensitive to climate warming 381 (Gutknecht et al., 2006). Determination of enzymatic activities in sediments of La Rocina stream 382 varied both among sampling sites and sampling times. Although determinations of enzymatic 383 activities in sediments are relatively scarce, previous analyses have shown they vary widely

384 across the different wetland ecosystems examined (Gutknecht et al., 2006). In our study, dryness 385 and temperature positively affected dehydrogenase, as values of activity were always greater at 386 T4. Similarly, β -glucosidase activity correlated significantly with the content of the organic 387 matter fraction, as the highest values of activity were detected in S1 and S2, the sites with the 388 highest TOC concentrations. Similar results were reported by Williams and Jochem (2006) who 389 showed that, despite the complex relationships between biological and environmental parameters, 390 the kinectic of several ectoenzymes, among them β -glucosidase, were controlled by organic 391 matter availability.

392 Wetlands play an important role on carbon cycle and on global climate change. The 393 emission of greenhouse gases, especially CO₂, and CH₄, shows a large spatial and temporal 394 variation due to the complex interactions between environmental variables and the 395 microbiological processes leading to gas production. The carbon flux is related to many external 396 factors, including soil environment, hydrological conditions, vegetation type and exogenous 397 nitrogen (Ma and Lu, 2009). As revealed by Pearson correlation matrix (Table 3), CO₂ and CH₄ 398 fluxes showed a strong seasonal influence, especially at S1 and S2, the sampling sites with the 399 highest TOC concentrations. There is to note, however, that NO₃⁻ contamination increased 400 production not only of N₂O, but also of CO₂ and CH₄. These results agree with those which show 401 that alterations in the biogeochemical cycles in nature may lead to altered biogenic fluxes of CO₂,

402 CH₄ and N₂O, the three main gases contributing to global warming (Liu and Greaver, 2009).

In addition to NO_3^- , denitrification correlated positively with the content of organic matter in the sediments. Accordingly, increased potential denitrification was observed at sites with the highest TOC values. Similar results were obtained during studies on denitrification and its relationship with organic carbon quality in three coastal wetland soils (Dodla et al., 2008). Also, Sirivedhin and Gray (2006) found that the sediment denitrification potential showed a positive

408 relationship with the biodegradable organic carbon concentration produced by the periphytic 409 algae in wetlands. Denitrification was also affected by the pluvial regime, as the highest values of 410 DEA were registered during the driest season at each sampling time. Hernández and Mitsch 411 (2007) founded that soil temperature, flood frequency and nitrate availability were important 412 factors controlling denitrification in created wetlands. Davidson, (1991) observed an increase 413 N_2O production in dry season, and specially during drying and wetting cycles, caused by a 414 temporal accumulation of mineral nitrogen into soil surface, which will become rapidly available 415 to microbial biomass when dry soil is wetted.

- 416
- 417 **4. Conclusions**
- 418

419 The surface water of La Rocina stream showed NO₃⁻ contamination, probably to 420 agricultural sources. This contamination decreased along La Rocina basin and apparently, the 421 superficial water body of DNP wetland was not affected. More research is needed to evaluate how the NO_3^- pollution could affect DNP groundwater. The environmental conditions such as 422 423 precipitation rate, hydrological morphology and organic matter content greatly influenced the 424 physicochemical characteristics of the surface waters of DNP wetland. The biological activity 425 and greenhouse gas production in their aquatic sediments was also affected by these 426 environmental parameters, specially the hydrology which had a major effect during the driest 427 season. The denitrification process was affected by anthropogenic activity (nitrate contamination 428 from agricultural practices) and the rainfall regimen, increasing the GHG emissions (CO₂, CH₄ 429 and especially N₂O) during the driest season in all sampling sites studied.

430

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44	3
44	4 6. References
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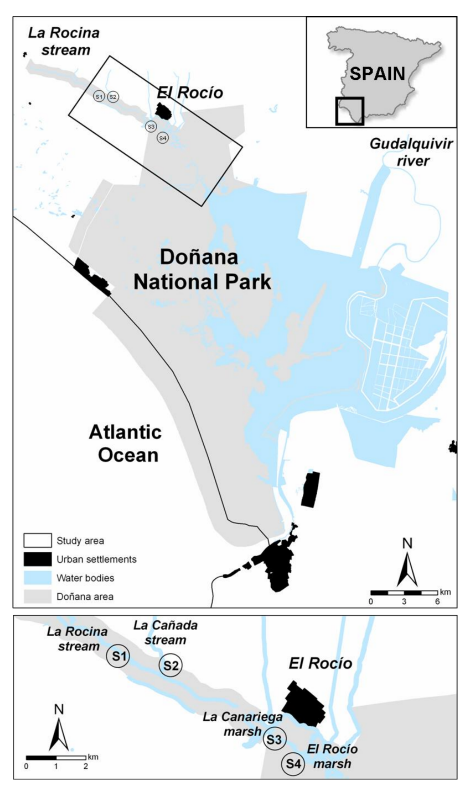


Fig 1. Geographical situation of Doñana National Park (DNP) and La Rocina stream. Sampling sites: Palacio del Acebrón (S1), Arroyo de la Cañada (S2), Vado de la Canariega (S3) and Marisma del Rocío (S4).

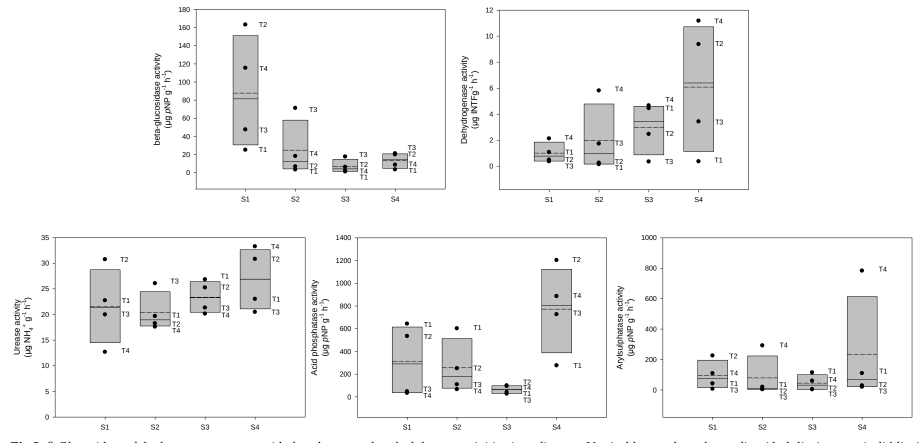


Fig 2. β-Glucosidase, dehydrogenase, urease, acid phosphatase and arylsulphatase activities in sediments. Vertical boxes show the median (dash line), mean (solid line) and the 5th/95th percentiles. Sampling sites: Palacio del Acebrón (S1), Arroyo de la Cañada (S2), Vado de la Canariega (S3) and Marisma del Rocío (S4). Sampling times: October 2008 (T1), January 2009 (T2), April 2009 (T3) and July 2009 (T4).

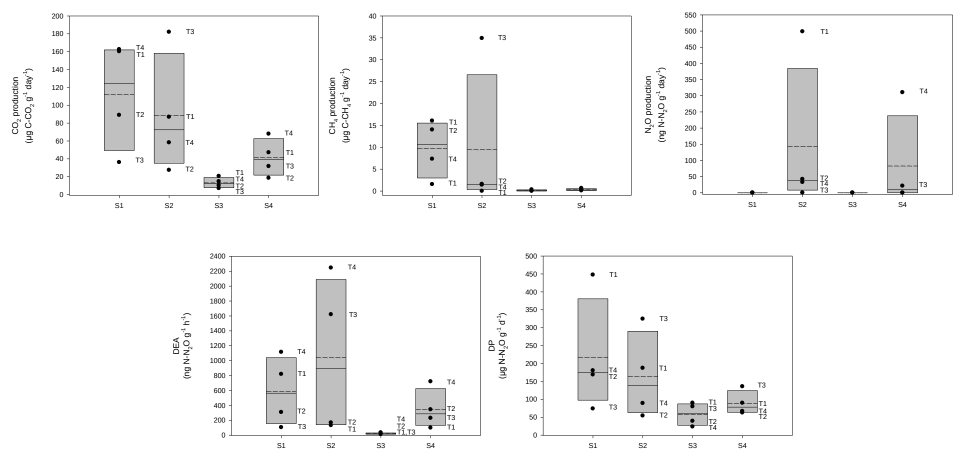


Fig 3. CO_2 , CH_4 and N_2O emissions, denitrifying enzymatic activity (DEA) and denitrification potential (DP) in sediments. The vertical boxes show the median (dash line), mean (solid line) and the 5th/95th percentiles. Sampling sites: Palacio del Acebrón (S1), Arroyo de la Cañada (S2), Vado de la Canariega (S3) and Marisma del Rocío (S4). Sampling times: October 2008 (T1), January 2009 (T2), April 2009 (T3) and July 2009 (T4).

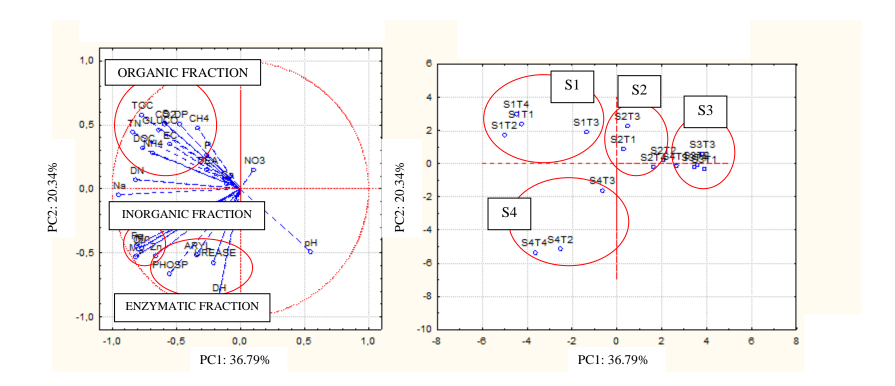


Fig 4. a. Principal Component Analysis performed on the whole set of measured sediments properties and b., the spatial and seasonal distribution of the parameters. Sampling sites: Palacio del Acebrón (S1), Arroyo de la Cañada (S2), Vado de la Canariega (S3) and Marisma del Rocío (S4). Sampling times: October 2008 (T1), January 2009 (T2), April 2009 (T3) and July 2009 (T4).

Sampling Site	Times	pН	¹ EC	² DOC	$^{2}NO_{3}$	$^{2}NO_{2}$	$^{2}NH_{4}^{+}$	$^{2}\mathbf{P}$	^{2}K	² Ca	² Mg	² Na	^{2}S	² Fe	² Cu	² Mn	² Zn
S1	T1	6.62b	0.63a	17.7a	0.2c	nd	0.1	<0.6	9.0a	49.3a	20.9a	66.8a	61.2a	0.0d	nd	0.9	< 0.01
	T2	6.57b	0.26c	16.7b	0.3b	0.6	nd	<0.6	4.8c	12.6b	6.0c	43.8b	7.0c	0.3a	0.1	< 0.01	0.3
	T3	6.99b	0.12d	5.7d	0.6a	nd	nd	< 0.6	2.0d	7.2c	3.3d	16.8c	2.4d	0.1c	0.1	< 0.01	0.2
	T4	7.60a	0.33b	8.9c	nd	nd	nd	<0.6	5.2b	3.4d	8.6b	8.6d	46.6b	0.2b	< 0.01	< 0.01	0.2
S2	T1	7.75c	0.52c	28.3a	61.6d	0.1c	0.1	<0.6	10.9c	38.5b	6.8d	20.5c	16.2c	nd	nd	< 0.01	< 0.01
	T2	7.15d	0.57b	3.6c	106.6a	nd	nd	1.9a	16.0b	36.5c	11.2c	28.7b	18.5c	< 0.01	0.1	0.1	< 0.01
	T3	8.89a	0.62a	6.7b	101.6b	0.5b	0.1	1.2b	27.7a	56.9a	20.3b	51.6a	30.0b	< 0.01	0.1	0.1	< 0.01
	T4	7.96b	0.50d	3.1c	68.8c	1.2a	nd	0.6c	14.4b	15.1d	58.7a	16.6c	46.2a	0.1	< 0.01	0.1	0.1
S3	T1	8.03b	1.22b	68.1b	1.9	0.5	0.3b	< 0.6	24.3b	71.2a	25.3b	144.2a	84.1b	nd	< 0.01	< 0.01	nd
	T2	7.30c	0.44c	12.3c	3.0	0.4	nd	< 0.6	5.5d	24.2d	7.9b	37.9c	14.9b	nd	0.1	< 0.01	0.1
	T3	8.38a	0.34d	21.0c	nd	nd	0.2b	<0.6	9.7c	28.8c	12.2b	53.1b	9.2b	0.52	0.3	< 0.01	0.5
	T4	8.03b	3.11a	135.2a	nd	nd	4.7a	2.3	41.3a	43.0b	30.1a	30.2d	581.3a	0.22	< 0.01	< 0.01	< 0.01
S4	T1	7.95a	0.90a	77.7a	1.1c	0.1a	0.8b	< 0.6	21.2a	70.3a	25.6b	103.4a	42.9b	< 0.01	< 0.01	0.2b	nd
	T2	7.54b	0.34d	6.8d	3.6b	0.2a	0.3d	< 0.6	4.5d	23.0c	6.3d	25.4c	9.6d	< 0.01	< 0.01	0.2b	< 0.01
	T3	8.32a	0.60c	12.9c	5.7a	nd	0.1c	<0.6	13.7c	51.4b	21.6c	79.4b	24.7c	< 0.01	< 0.01	< 0.01	< 0.01
	T4	7.51b	0.83b	16.3b	nd	nd	0.9a	<0.6	17.3b	14.1d	59.2a	26.0c	105.3a	< 0.01	< 0.01	1.2a	0.1
<u>s</u>		*	*	*	*	-	*	-	*	*	*	*	*		-	-	-
<u> </u>		*	*	*	*		*		*	*	*	*	*	····· · ·····		<u>.</u>	

Table 1a. Physicochemical properties of the surface waters at the sampling sites.

Sampling sites: Palacio del Acebrón (S1), Arroyo de la Cañada (S2), Vado de la Canariega (S3) and Marisma del Rocío (S4). Sampling times: October 2008 (T1), January 2009 (T2), April 2009 (T3) and July 2009 (T4). For each variable, at a given sampling site, values followed by the same letter are not statistically different according to Tukey's test at $p \le 0.05$. ¹Values of electrical conductivity (EC) are expressed in dS cm⁻¹. ²Values of dissolved organic carbon (DOC), NO₃⁻, NO₂⁻ and NH₄⁺ are expressed in mg L⁻¹. nd: no detected. *: $p \le 0.05$.

Sampling Site	Times	TDN (mg L^{-1})	DOC/DN	$N_{ORG}(\%)$	$N-NO_{3}(\%)$	$N-NO_{2}^{-}(\%)$	$N-NH_4^+(\%)$
S1	T1	0.6b	29.5	79.5	7.5	-	13.0
	T2	0.7a	23.9	64.2	9.7	26.1	-
	Т3	0.3d	19.0	54.8	45.2	-	-
	T4	0.4c	22.3	100.0	-	-	-
S2	T1	15.6b	1.8	10.1	89.2	0.2	0.5
	T2	15.3b	0.1	3.7	96.3	-	-
	Т3	21.6a	0.3	4.2	94.7	0.7	0.4
	T4	9.4c	0.2	8.6	89.3	2.1	-
S3	T1	4.4b	15.5	81.5	9.8	3.5	5.3
	T2	1.1b	11.2	27.3	61.6	11.1	-
	Т3	1.2b	17.5	87.0	-	-	13.0
	T4	18.1a	7.5	79.8	-	-	20.2
S4	T1	3.6a	21.6	75.0	6.9	0.8	17.3
	T2	1.1d	6.2	20.6	73.9	5.5	-
	Т3	2.1c	6.1	35.0	61.3	-	3.7
	T4	2.3b	7.1	69.6	-	-	30.4

Table 1b. Total dissolved nitrogen (TDN), dissolved organic carbon (DOC) and organic nitrogen (N_{ORG}) in the surface waters at the sampling sites.

Sampling sites: Palacio del Acebrón (S1), Arroyo de la Cañada (S2), Vado de la Canariega (S3) and Marisma del Rocío (S4). Sampling times: October 2008 (T1), January 2009 (T2), April 2009 (T3) and July 2009 (T4). For DN, at a given sampling site, values followed by the same letter are not statistically different according to Tukey's test at $p \le 0.05$. $N_{ORG} = DN - N_{INORG}$, where N_{INORG} (inorganic nitrogen) = N-NO₃⁻ + N-NO₂⁻ + N-NH₄⁺. nd, no detected.

Sampling Site	Times	pH (1:5)	$^{1}\text{EC}(1:5)$	^{2}P	^{2}K	² Ca	² Mg	² Na	2 S	² Fe	³ Cu	³ Mn	³ Zn
S1	T1	5.44	860	0.12d	5.07c	7.28a	3.09b	0.71b	7.75b	16.39a	11c	252a	25b
	T2	5.51	390	0.45b	5.51b	5.89b	2.88b	0.71b	3.34d	15.40b	25a	163c	27b
	Т3	3.98	565	0.83a	4.43d	5.28c	1.96c	0.55c	9.66a	13.47c	11c	98d	44a
	T4	5.90	178	0.32c	6.71a	6.89a	3.40a	0.82a	5.61c	13.42c	11c	189b	22b
S2	T1	5.05d	628a	0.12d	2.92b	5.04bc	1.72a	0.28a	3.80a	10.32b	8a	136a	16b
	T2	7.13c	82c	0.45a	3.21a	3.86c	1.79a	0.24b	0.65b	12.26a	4b	136a	17b
	Т3	7.59b	153b	0.21c	1.92c	5.75b	0.92c	0.21bc	0.48c	5.06d	10a	114b	16b
	T4	8.26a	113bc	0.27b	2.02c	36.83a	1.33b	0.21c	0.52c	5.81c	9a	107b	19a
S3	T1	8.58a	90b	0.12b	0.68b	0.44c	0.44bc	0.10b	0.16a	2.57ab	5a	58a	7c
	T2	7.72c	84c	0.45a	0.91b	0.96a	0.62b	0.10b	0.15a	4.11a	4a	82a	11a
	Т3	6.75d	39d	0.02c	0.57b	0.25c	0.28c	0.11b	0.14a	2.07b	2a	65a	17b
	T4	8.20b	101a	0.04c	1.29a	0.68b	0.76a	0.25a	0.12a	3.61ab	2a	69a	4d
S4	T1	7.36c	117c	< 0.01	1.92d	1.32d	1.08c	0.23a	0.20a	5.66d	3d	140c	13d
	T2	7.55b	165b	< 0.01	8.72b	5.58a	5.19a	0.48b	0.67a	26.60a	26a	313a	56a
	Т3	6.81d	115c	0.17b	7.61c	2.78c	3.51b	0.49b	0.59a	13.67c	19c	150c	34c
	T4	7.68a	221a	0.33a	10.95a	4.37b	5.12a	0.80a	0.65a	19.37b	23b	256b	46b
S		*	*	*	*	*	*	*	*	*	*	*	*
Т		*	*	*	*	*	*	*	*	*	*	*	*
S x T		*	*	*	*	*	*	*	*	*	*	*	*

Table 2a. Physicochemical properties of the sediments at the sampling sites.

Sampling sites: Palacio del Acebrón (S1), Arroyo de la Cañada (S2), Vado de la Canariega (S3) and Marisma del Rocío (S4). Sampling times: October 2008 (T1), January 2009 (T2), April 2009 (T3) and July 2009 (T4). For each variable, at a given sampling site, values followed by the same letter are not statistically different according to Tukey's test at $p \le 0.05$. ¹Values of EC (electrical conductivity) are expressed in μ S cm⁻¹. ²For each variable, values are expressed in g Kg⁻¹ (sediment dry weight). ³For each variable, values are expressed in mg Kg⁻¹ (sediment dry weight). *: $p \le 0.05$.

Sampling Site	Season	¹ TOC	¹ TN	TOC/TN	N _{ORG} (%)	$^{2}NO_{3}$	$^{2}NO_{2}$	$^{2}NH_{4}^{+}$	² DOC	² DN
S1	T1	183.4a	5.7b	31.9	99.4	nd	nd	41b	1137c	70b
	T2	154.2b	9.0a	17.1	99.5	34	nd	44b	2580a	144a
	T3	99.8c	3.7c	27.3	99.3	7	nd	30c	332d	30c
	T4	206.0a	6.9b	29.8	99.1	nd	nd	79a	1934b	138a
S2	T1	78.4a	3.1a	25.5	99.6	32	nd	6c	280b	22bc
	T2	18.3b	1.0c	18.8	97.5	54	nd	16b	258b	23b
	T3	38.4b	1.6bc	23.7	97.8	3	nd	45a	174c	16c
	T4	22.7b	0.5c	42.9	96.8	nd	nd	20b	412a	42a
S3	T1	3.1a	0.3a	11.7	98.5	32a	nd	6c	163b	15b
	T2	3.4a	0.3a	12.2	97.8	4b	nd	7c	147b	17b
	Т3	1.8b	0.2a	8.3	92.3	3b	nd	20b	25c	3c
	T4	1.7b	0.2a	11.0	89.9	nd	nd	26a	206a	33a
S4	T1	5.4d	0.4c	10.8	97.3	nd	nd	14c	310c	27c
	T2	16.6c	1.5b	10.9	98.6	4	nd	25b	382b	43b
	T3	25.2a	1.9a	12.6	97.7	nd	nd	55a	196d	19c
	T4	20.1b	1.9a	10.5	98.8	nd	nd	30b	669a	104a
S		*	*	*	*	-	-	*	*	*
Т	•	*	*	*	*	-	-	*	*	*
S x T		*	*	*	*	-	-	*	*	*

 Table 2b. Total organic carbon (TOC), total nitrogen (TN), organic nitrogen (N_{ORG}), dissolved organic carbon (DOC) and dissolved nitrogen (DN) in sediments at the sampling sites.

Sampling sites: Palacio del Acebrón (S1), Arroyo de la Cañada (S2), Vado de la Canariega (S3) and Marisma del Rocío (S4). Sampling times: October 2008 (T1), January 2009 (T2), April 2009 (T3) and July 2009 (T4). For each variable, at a given sampling site, values followed by the same letter are not statistically different according to Tukey's test at $P \le 0.05$. ¹Values of TOC and TN are expressed in g Kg⁻¹ (sediment dry weight). ²Values of NO₃⁻, NO₂⁻, NH₄⁺, DOC and DN are expressed in mg Kg⁻¹ (sediment dry weight). DOC and DN were obtained after 2 h water extraction (1:20, w:v) of the lyophilized sediments. N_{ORG} = TN - N_{INORG}, where N_{INORG} (inorganic nitrogen) = N-NO₃⁻ + N-NO₂⁻ + N-NH₄⁺. nd, no detected. *: $p \le 0.05$.

	pН	EC	TOC	TN	$\mathbf{NH_4^+}$	DOC	DN	DH	GC	AS	AP	UR	CO ₂	CH ₄	DEA	DP
pН																
EC	-0.770**															
тос	-0.720**	0.676**														
TN	-0.698**	0.601*	0.932**													
$\mathbf{NH_4}^+$	NS	NS	0.663**	0.644**												
DOC	NS	NS	0.815**	0.992**	0.612*											
DN	NS	NS	0.718**	0.815**	0.622*	0.932**										
DH	0.549*	NS	NS	NS	NS	NS	NS									
GC	NS	NS	0.707**	0.839**	0.689**	0.861**	0.736**	NS								
AS	NS	NS	NS	NS	NS	NS	0.521*	0.654**	NS							
AP	NS	NS	NS	NS	NS	NS	NS	0.509*	NS	NS						
UR	NS	NS	NS	NS	NS	NS	NS	0.499*	NS	NS	0.554*					
CO ₂	NS	NS	0.714**	0.605**	0.625**	0.516*	NS	NS	0.542*	NS	NS	NS				
CH ₄	NS	NS	NS	NS	NS	NS	NS	NS	0.539*	NS	NS	NS	0.804**			
DEA	NS	NS	NS	NS	NS	NS	NS	NS	NS	NS	NS	NS	0.606*	0.506*		
DP	NS	0.641**	0.632**	0.513**	NS	NS	NS	NS	NS	NS	NS	NS	0.854**	0.765**	NS	

Table 3. Pearson correlation matrix (n=16) between the physicochemical properties, enzymatic activities and greenhouse gas emissions in sediments at four sampling sitesalong La Rocina stream.

**, *: significant at p < 0.01 and 0.05, respectively, NS: not significant. EC: Electrical conductivity, TOC: total organic carbon, TN: total nitrogen, DOC: dissolved organic carbon, DN: dissolved nitrogen, DH: dehydrogenase activity; GC, β -glucosidase activity; AS, arylsulphatase activity; AP, acid phosphatase activity; UR: urease activity; DEA: denitrifying enzymatic activity; and DP, denitrification potential.