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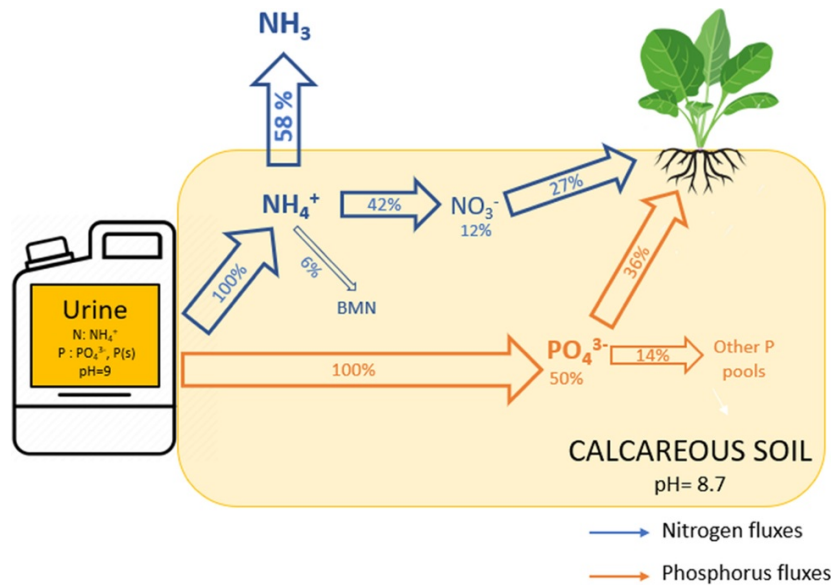
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Corresponding Author	FamilyName	<b>Rumeau</b>
	Particle	
	Given Name	<b>Manon</b>
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	Organization	Eco&SolsUniv Montpellier, CIRAD, INRAE, Institut Agro, IRD
	Address	Montpellier, France
	Division	School of Geography, Earth and Environmental Sciences
	Organization	University of Birmingham
	Address	Birmingham, UK
	Phone	
	Fax	
	Email	MLR094@student.bham.ac.uk
	URL	
	ORCID	<a href="http://orcid.org/0000-0001-7731-6247">http://orcid.org/0000-0001-7731-6247</a>
Author	FamilyName	<b>Marsden</b>
	Particle	
	Given Name	<b>Claire</b>
	Suffix	
	Division	
	Organization	Eco&SolsUniv Montpellier, CIRAD, INRAE, Institut Agro, IRD
	Address	Montpellier, France
	Phone	
	Fax	
	Email	
	URL	
	ORCID	
Author	FamilyName	<b>Mouheb</b>
	Particle	
	Given Name	<b>Nassim Ait</b>
	Suffix	
	Division	
	Organization	INRAE, UMR GEAU, Univ Montpellier
	Address	Montpellier, France
	Phone	
	Fax	
	Email	
	URL	
	ORCID	
Author	FamilyName	<b>Crevoisier</b>
	Particle	
	Given Name	<b>David</b>

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Organization	LISAH, Univ Montpellier, INRAE, Institut Agro, IRD	
Address	Montpellier, France	
Phone		
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Author	FamilyName Particle Given Name Suffix Division Organization Address Phone Fax Email URL ORCID	<b>Pistocchi</b> <b>Chiara</b>
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Abstract	<p>Human urine concentrates 88% of the nitrogen and 50% of the phosphorus excreted by humans, making it a potential alternative crop fertilizer. However, knowledge gaps remain on the fate of nitrogen in situations favouring <math>\text{NH}_3</math> volatilization and on the availability of P from urine in soils. This study aimed at identifying the fate of nitrogen and phosphorus supplied by human urine from source separation toilets in a calcareous soil. To this end, a spinach crop was fertilized with 2 different doses of human urine (<math>170 \text{ kgN ha}^{-1} + 8.5 \text{ kgP ha}^{-1}</math> and <math>510 \text{ kgN ha}^{-1} + 25.5 \text{ kgP ha}^{-1}</math>) and compared with a synthetic fertilizer treatment (<math>170 \text{ kgN ha}^{-1} + 8.5 \text{ kgP ha}^{-1}</math>) and an unfertilized control. The experiment was conducted in 4 soil tanks (50-cm depth) in greenhouse conditions, according to a randomized block scheme. We monitored soil mineral nitrogen over time and simulated nitrogen volatilization using Hydrus-1D and Visual Minteq software. We also monitored soil phosphorus pools, carbon, nitrogen and phosphorus (CNP) in microbial biomass, soil pH and electrical conductivity. Only an excessive input of urine affected soil pH (decreasing it by 0.2 units) and soil conductivity (increasing it by 183%). The phosphorus supplied was either taken up by the crop or remained mostly in the available P pool, as demonstrated by a net increase of the resin and bicarbonate extractable P. Ammonium seemed to be nitrified within about 10 days after application. However, both Visual Minteq and Hydrus models estimated that more than 50% of the nitrogen supplied was lost by ammonia volatilization. Overall, our results indicate that direct application of urine to a calcareous soil provides available nutrients for plant growth, but that heavy losses of volatilized nitrogen are to be expected. Our results also question whether long-term application could affect soil pH and salinity.</p> <p><i>Graphical Abstract:</i></p>	



Keywords (separated by '-') Source separation - Fertilization - Hydrus - Ammonia volatilization

Footnote Information

Responsible Editor: Kitae Baek • Nitrogen from urine was bioavailable for plants and microbes, but half of the N applied could be lost by volatilization in a calcareous soil. • Phosphorus from urine was either taken up by plants or remained mostly in available pools in a calcareous soil. • Only excessive doses of urine application affected soil pH and conductivity in the short term. The online version contains supplementary material available at <https://doi.org/10.1007/s11356-023-26895-5>.



## 2 **Fate of nitrogen and phosphorus from source-separated human urine 3 in a calcareous soil**

4 **Manon Rumeau<sup>1,2</sup> · Claire Marsden<sup>1</sup> · Nassim Ait Mouheb<sup>3</sup> · David Crevoisier<sup>4</sup> · Chiara Pistocchi<sup>1</sup>**

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### 7 **Abstract**

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23 **Keywords** Source separation · Fertilization · Hydrus · Ammonia volatilization

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### A2 **Highlights**

- Nitrogen from urine was bioavailable for plants and microbes, but half of the N applied could be lost by volatilization in a calcareous soil.
- Phosphorus from urine was either taken up by plants or remained mostly in available pools in a calcareous soil.
- Only excessive doses of urine application affected soil pH and conductivity in the short term.

A10 Manon Rumeau  
A11 MLR094@student.bham.ac.uk

## Introduction

The global nitrogen (N) cycle has been massively altered for decades, by the synthesis of reactive N from atmospheric N<sub>2</sub> for fertilizer production and by the conversion of reactive N into N<sub>2</sub>O or N<sub>2</sub> during wastewater treatment (Gruber and Galloway 2008; Steffen et al. 2015). In these two opposite processes greenhouse gases are emitted and fossil fuel energy is used, while nitrogen pollution is caused as a side effect (Kampschreuer et al. 2009). The phosphorus (P) cycle has also been heavily disrupted, with the additional issue that phosphate rock, from which P fertilizers are sourced, is becoming scarce (Desmidt et al. 2015). Wastewater treatment only removes a part of the P, which is however little recycled, and the rest is discharged into surface waters. Therefore, re-looping N and P fluxes appears to be a promising solution to reduce wastewater pollution and synthetic fertilizer dependency. Human urine is of particular interest because it concentrates 88% of the nitrogen and 50% of the phosphorus

A12 <sup>1</sup> Eco&SolsUniv Montpellier, CIRAD, INRAE, Institut Agro, IRD, Montpellier, France

25

26

A13 <sup>2</sup> Present Address: School of Geography, Earth  
A14 and Environmental Sciences, University of Birmingham,  
A15 Birmingham, UK

27

A16 <sup>3</sup> INRAE, UMR GEAU, Univ Montpellier, Montpellier, France

28

A17 <sup>4</sup> LISAH, Univ Montpellier, INRAE, Institut Agro, IRD,  
A18 Montpellier, France

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42 excreted by humans (Martin et al. 2022). If recycled, it could  
 43 account for more than 13% of the global agricultural fertilizer  
 44 demand (Wald 2022). In addition, urine represents less than  
 45 1% of the volume of wastewater but 79% of the nitrogen and  
 46 47% of the phosphorus treated by sewage treatment plants  
 47 (Larsen et al. 2013). Hence, diverting it would reduce by  
 48 more than half the nutrient pollution from wastewater (Wald  
 49 2022). Urine can be considered sterile in most cases and does  
 50 not present risks of disease transmission if not contaminated  
 51 with faeces. This is possible in source separation systems  
 52 (toilets with separated outlets or urinals) (Lienert and Larsen  
 53 2010) which allow the safe collection of urine.

54 Each human produces 1 to 1.5L of urine per day (Karak  
 55 and Bhattacharyya 2011). Urine is composed of 95% water,  
 56 and the remaining 5% consists of amino compounds (such  
 57 as urea or creatinine), organic anions and inorganic salts  
 58 (Maggi and Daly 2006). After urea hydrolysis, nitrogen is  
 59 mainly in ammonium form and phosphorus is either dis-  
 60 solved in solution or precipitated as struvite (magnesium  
 61 ammonium phosphate) (Udert et al. 2006). The fertilizing  
 62 ability of animal urine has been known for a long time (Di  
 63 and Cameron 2007; Fanjaniaina et al. 2022), while that of  
 64 human urine has been proven in recent years (Akpan-Idiok  
 65 et al. 2012; Martin et al. 2022; Pradhan et al. 2009). How-  
 66 ever, the fate and dynamics of nitrogen and phosphorus from  
 67 human urine are highly uncertain as soil biotic and abiotic  
 68 processes involving N and P could be affected by the other  
 69 compounds and nutrients present in urine. For instance, hip-  
 70 puric acid in urine can inhibit denitrification (Kool et al.  
 71 2006), and the formation of ammonium bicarbonate can  
 72 inhibit nitrification (Clough et al. 2003; Somers et al. 2019).

73 Calcareous soils are common agricultural land in France.  
 74 These soils are prone to ammonia ( $\text{NH}_3$ ) volatilization because  
 75 of their alkaline pH. Therefore, urine application on such soils  
 76 is likely to result in high  $\text{NH}_3$  losses by volatilization. However,  
 77 published volatilization rates range from 0 to 63% of the am-  
 78 monium supplied (Mills et al. 1974; Powlson and Dawson 2022).  
 79 Phosphorus availability is also an important issue in alkaline  
 80 soils and could be improved by fertilization with urine: firstly  
 81 because urine is a source of P, and secondly because the appli-  
 82 cation of ammonium can decrease soil pH by enhancing acidifi-  
 83 fying processes such as ammonia volatilization, nitrification  
 84 and subsequent nitrate leaching (Bolan et al. 1991; Raza et al.  
 85 2021). Hence, a decrease in soil pH could increase the availabil-  
 86 ity of P from urine (i.e. dissolved as phosphate or precipitated  
 87 as struvite) which largely depends on pH (Frossard et al. 2000;  
 88 Helfenstein et al. 2020; Meyer et al. 2018).

89 Aside from the effect on P availability, soil acidification is  
 90 an important process to monitor as it can generate negative  
 91 feedbacks on soil fertility through a reduction of the cation  
 92 exchange capacity (Barak et al. 1997) and an increase in soil  
 93  $\text{CO}_2$  emissions from the dissolution of carbonate in calcareous  
 94 soils (Raza et al. 2021).

Moreover, urine is a multi-component solution: as well as  
 N and P, it contains  $\text{K}^+$ ,  $\text{S}$ ,  $\text{Ca}^{2+}$ ,  $\text{Mg}^{2+}$ ,  $\text{Na}^+$ ,  $\text{Cl}^-$  and other  
 micronutrients. Thus, it could represent a complete fertilizing  
 solution even though the bioavailability of urine micro-  
 nutrients has not been well documented (Olivia et al. 2015). Yet,  
 the high concentrations in  $\text{Na}^+$ ,  $\text{Cl}^-$  and  $\text{SO}_4^{2-}$  could  
 also cause soluble salt accumulation in soils (Boh and Sauerborn  
 2014; Mnkeni et al. 2008; Shingiro et al. 2019).

The objectives of this study were to determine the fates of  
 N and P from urine fertilization on a calcareous soil and the  
 effect on soil pH and salinity. The main hypotheses were: (1)  
 N supplied with urine is readily bioavailable, but a signifi-  
 cant amount of N is lost by volatilization potentially causing  
 a decrease in soil pH; (2) urine application and associated  
 decrease in soil pH increase phosphorus availability in soils;  
 (3) urine increases soil salinity because of its soluble salt  
 concentration. To address these hypotheses, we conducted  
 a fertilization trial on a spinach crop (*Spinacia oleracea*  
 L.) where we compared the effect of two different doses  
 of source separated human urine with that of a synthetic  
 fertilizer with equivalent N and P concentrations and an  
 unfertilized control.

## Materials and methods

### Site and experiment description

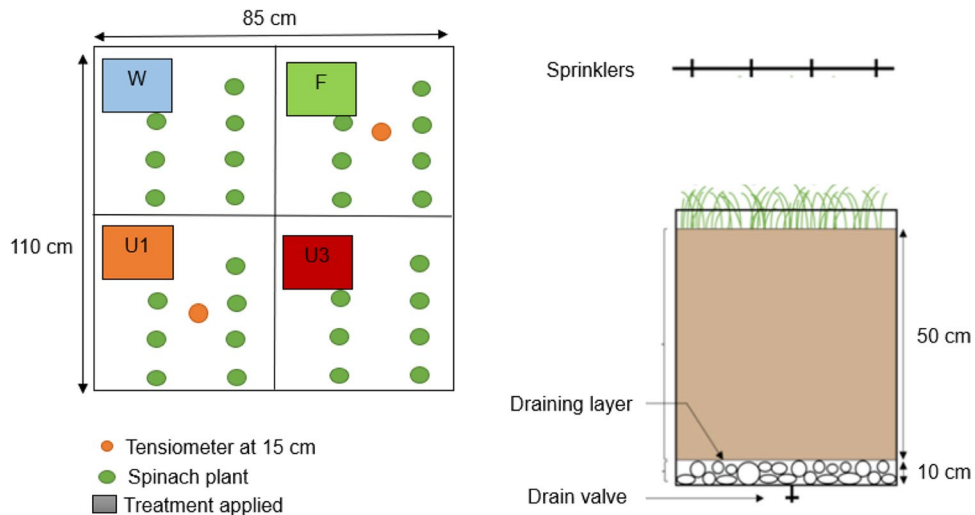
The experiment was carried out in a greenhouse of the UMR  
 G-EAU in Montpellier between May 30<sup>th</sup> 2020 and July 5<sup>th</sup>  
 2020. Meteorological variables were measured by a weather  
 station located at the experimental site. Air temperature and  
 relative humidity in the greenhouse were measured by a tem-  
 perature and relative humidity probe (model CS215, CAMP-  
 BELL SCIENTIFIC), and global radiation was measured by  
 a pyranometer (model SP1110, CAMPBELL SCIENTIFIC)  
 (Table 1).

The experiment was conducted in 4 soil tanks (soil sur-  
 face equal to 0.935 m<sup>2</sup> and 50-cm soil depth) (Fig. 1). Each  
 tank was filled with approximatively 0.53 tonnes of air-dried  
 loamy clay soil (24% clay, 25.6% silt, 19.5% very fine sand,  
 16.4% fine sand, 14.4% coarse sand) with 45% of carbonate  
 and a pH of 8.7. A spinach crop (*Spinacia oleracea* L.) was

**Table 1** Climatic parameters inside the greenhouse from June 17<sup>th</sup> to July 2<sup>nd</sup>, 2020

	Air tempera- ture (°C)	Air relative humidity (%)	Global radiation (kW m <sup>-2</sup> )
Mean	24.3	63.7	0.203
Minimum	9.5	23.7	0
Maximum	36.0	99.7	1.041

**Fig. 1** Scheme of the experimental design (representation of one tank, from above (left) and from the side (right)) with crops and sensor positions. For each tank, the position of the different treatments was randomized

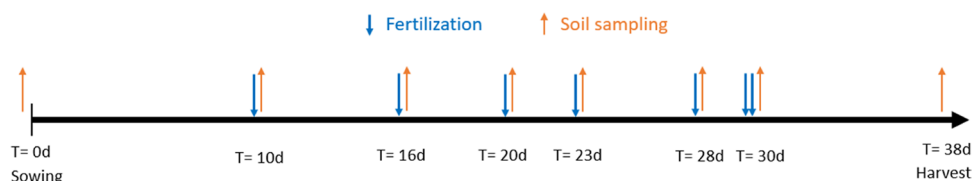


134 sown directly in the tanks on May 30<sup>th</sup> with a plant den-  
 135 sity of 17 plant m<sup>-2</sup> and the growing cycle lasted 37 days.  
 136 On July 5<sup>th</sup>, the above-ground biomass was collected and  
 137 dried for biomass and nutrient content analysis. Spinach was  
 138 chosen because of its relatively high N requirements (170  
 139 kgN ha<sup>-1</sup> under optimal growth conditions) (Frerichs et al.  
 140 2022) and short growing cycle. Irrigation was conducted  
 141 with sprinklers located above the tanks and controlled with  
 142 tensiometers placed at 15-cm depth in the soil to maintain  
 143 soil moisture around field capacity. Approximately 200 mm  
 144 of water was supplied over the duration of the experiment.  
 145 Each tank was divided into 4 quarters using vertical alu-  
 146 minium sheets driven 30 cm into the soil (below maximum  
 147 root depth to avoid transfers of nutrients between quarters).  
 148 The experimental treatments were assigned according to a  
 149 randomized block scheme in which the experimental unit  
 150 was a quarter of a tank and the tank was the block. We com-  
 151 pared two different doses of human urine with a synthetic  
 152 fertilizer and a water control. The four treatments applied  
 153 were: U1 = Urine dose  $\times$  1 (170 kgN ha<sup>-1</sup> + 8.5 kgP ha<sup>-1</sup>  
 154 supplied), U3 = Urine dose  $\times$  3 (510 kgN ha<sup>-1</sup> + 25.5 kgP  
 155 ha<sup>-1</sup> supplied), F = Synthetic fertilizer dose  $\times$  1 (170 kgN  
 156 ha<sup>-1</sup> + 8.5 kgP ha<sup>-1</sup> supplied) and W = Water. Treatments U1  
 157 and F represent the recommended doses of N for the spin-  
 158 ach crop, while treatment U3 represents 3 times this dose  
 159 and exacerbates the effects of urine as well as approximates

the recommended dose of phosphorus. Nitrogen and phosphorus in the F treatment were added as ammonium nitrate (NH<sub>4</sub>NO<sub>3</sub>) and potassium phosphate (KH<sub>2</sub>PO<sub>4</sub>) respectively. Potassium was not limiting in soils (8 mg kg<sup>-1</sup> at T0, data not shown) so we assumed that the slight difference in potassium concentration between the urine and the synthetic fertilizer would not affect plant growth.

The treatments were fractionated into 6 applications, and each one was diluted by 11.8 for U1 and F and by 3.8 for U3 (Fig. 2).

The urine used in this experiment was collected by the start-up EcoSec (Montpellier, France, <https://ecosec.fr/>), which manufactures and sells source separating toilets. These toilets allow the selective collection of urine and faeces thanks to a gravity system, but do not prevent cross contamination with faeces. Prior to the experiment, the urine was stored for a year in an opaque and airtight container in order to sanitize the effluent according to World Health Organization (WHO) recommendations (Schöning and Stenström 2004). The urine's chemical composition is summarized in Table 2. As expected, P was only in inorganic form and nitrogen was mostly present in ammonium form. However, ammonium concentration in urine decreased by 0.58 gN L<sup>-1</sup> between the beginning (T0) and the end of the experiment (TF) due to ammonia volatilization during container openings; this was considered in the N budget. Organic carbon concentration was very low (57 mgC L<sup>-1</sup>) despite potential cross contamination with faeces. Furthermore, the high concentration



**Fig. 2** Timeline of the fertilization treatments. The scale is in day (d) from sowing (day 0) to harvest (day 38). Each fertilization and soil sampling are represented by a blue and an orange respectively. Double dose of fertilization was supplied on day 30

**Table 2** Chemical composition at T0 (start of the experiment) of the undiluted urine used in this experiment after 1 year of storage in an air-tight opaque tank

Parameters measured	Values	Method
Electrical conductivity at 25 °C (mS cm <sup>-1</sup> )	40.5	NF EN 27,888
pH water	9	NF EN ISO 10523
N-NH <sub>4</sub> <sup>+</sup> (mg L <sup>-1</sup> )	4341	Filtered at 0.45 µm, ISO 7150-1
N-NO <sub>3</sub> <sup>-</sup> (mg L <sup>-1</sup> )	10.7	Filtered at 0.45 µm, ISO 7890-1-2-1986
Ptotal (mg L <sup>-1</sup> )	206	Filtered at 0.45 µm, ICP-AES
Porganic (mg L <sup>-1</sup> )	Not detected	
K <sup>+</sup> (mg L <sup>-1</sup> )	1107.8	Filtered at 0.45 µm, ICP-AES
Mg <sup>2+</sup> (mg L <sup>-1</sup> )	0.758	Filtered at 0.45 µm, ICP-AES
Ca <sup>2+</sup> (mg L <sup>-1</sup> )	3.49	Filtered at 0.45 µm, ICP-AES
Na <sup>+</sup> (mg L <sup>-1</sup> )	1245.6	Filtered at 0.45 µm, ICP-AES
SO <sub>4</sub> <sup>2-</sup> (mg L <sup>-1</sup> )	1228.81	ISO 11885
Cl <sup>-</sup> (mg L <sup>-1</sup> )	3574.85	Water extraction 1/5, NF EN 9297
DBO <sub>5</sub> (mg L <sup>-1</sup> ): 5-day biochemical oxygen demand	3670	NF EN 1899-1
ST-DCO (mg L <sup>-1</sup> ): chemical oxygen demand	6790	NF T90-101
Labile carbon (mgC L <sup>-1</sup> )	11.9	Filtered at 0.45 µm, POXC
Total dissolved carbon/inorganic carbon/organic carbon (mgC L <sup>-1</sup> )	2400/2343/57	Filtered at 0.45 µm, TOC-TN analyser
Cu (mg L <sup>-1</sup> )	0.264	Filtered at 0.45 µm, ICP-AES
Fe (mg L <sup>-1</sup> )	0.126	
Mn (mg L <sup>-1</sup> )	<0.008	
Zn (mg L <sup>-1</sup> )	0.198	
Bo (mg L <sup>-1</sup> )	0.556	
Al (mg L <sup>-1</sup> )	0.056	
SAR (sodium absorption ratio)	157.6	
- Solution U1	45.8	
- Solution U3	80.8	

Abbreviations: ICP-AES inductively coupled plasma atomic emission spectroscopy, POXC permanganate-oxidizable carbon, TOC-TN total organic carbon/total nitrogen

of Na<sup>+</sup> (1289 mg L<sup>-1</sup>) and Cl<sup>-</sup> (3574 mg L<sup>-1</sup>) along with the conductivity (40.6 mS cm<sup>-1</sup>) showed that the urine had high salinity (Table 2). Additionally, the sodium absorption ratio (SAR, indicating the potential sodium hazard for irrigation or fertirrigation) of pure urine, solution U1 and solution U3 was calculated with the following formula, and SAR values are detailed in Table 2:

$$193 \quad SAR = \frac{Na^+}{\sqrt{\frac{1}{2}(Ca^{2+} + Mg^{2+})}} \quad (1)$$

where concentrations of cations (Na<sup>+</sup>, Ca<sup>2+</sup>, Mg<sup>2+</sup>) are expressed in meq L<sup>-1</sup>.

## 197 Parameter monitoring and measurement

### 198 Soil sampling

199 Bulk soil samples at 0–10-cm depth composed of 2 sub-  
200 samples were taken in every quarter of a tank at the begin-  
201 ning of the experiment before sowing (T0), right after every

fertilization (T1 to T6) and at the end of the experiment (TF). For every sample, the gravimetric water content was measured by drying the soil at 105 °C for 48 h. Mineral nitrogen (NO<sub>3</sub><sup>-</sup> and NH<sub>4</sub><sup>+</sup>) was measured in all the samples, whereas pH, conductivity, organic and inorganic P concentration and microbial biomass were quantified only in the T0 and TF samples.

### 209 Soil bulk density

210 In each tank, 3 undisturbed soil cores of 5-cm depth were  
211 taken. Soil cores were then oven dried at 105 °C, and the  
212 bulk density was calculated for each core by dividing the  
213 dry mass of soil by the volume of the core.

### 214 Mineral nitrogen measurement

215 Soils were extracted with 1 M KCl (soil to extractant ratio  
216 of 1:3) on the day of sampling. The extracts were filtered  
217 at 0.45 µm and frozen until analysis. Then, samples were

218 analysed by continuous flow colorimetry (Skalar SA 3000  
 219 analyser). This method measures nitrite and nitrate species  
 220 together and ammonium separately.

221 **pH**

222 pH measurement was carried out according to the ISO stand-  
 223 ard using a pH probe (AFNOR, 2005). Soils were sieved,  
 224 air dried and extracted with a soil to water ratio of 1:5, and  
 225 measurements were performed after 1 h of agitation.

226 **Electrical conductivity**

227 Soils were sieved, air dried and extracted with a soil to water  
 228 ratio of 1:5. Then, the solution was agitated for 30 min, cen-  
 229 trifuged and filtered. Electrical conductivity was then mea-  
 230 sured on the aqueous extract with a conductometer probe.

231 **Organic and inorganic P pool**

232 To quantify soil phosphorus pools, we performed a Hedley  
 233 sequential fractionation as modified by Tiessen and Moir  
 234 (1993). This method operationally identifies organic and  
 235 inorganic P pools. Soil samples were sequentially extracted  
 236 with 4 different reagents in the following order: anionic  
 237 exchange resin membranes (BDH #55,164, 6 cm × 4 cm,  
 238 named P resin pool), 0.5 M sodium bicarbonate (NaHCO<sub>3</sub>,  
 239 Pi and Po bicarbonate pools), 0.5 M NaOH (Pi and Po NaOH  
 240 pools) and 1 M HCl (P HCl pool). To quantify organic P,  
 241 aliquots of bicarbonate and NaOH extracts were mineralized  
 242 by acid digestion, and the organic P concentration was cal-  
 243 culated as the difference between inorganic P in the digested  
 244 extract (corresponding to the total P in the extract) and inor-  
 245 ganic P in the non-digested extract. Inorganic phosphorus  
 246 concentration in each extract was measured by the malachite  
 247 green colorimetric method (Ohno and Zibilske 1991).  
 248 The P resin and the Pi and Po bicarbonate are commonly  
 249 considered the available P pool (Tiessen and Moir 1993).  
 250 The exchange times of sequentially extracted P pools with  
 251 the soil solution increase with the strength of the extractant  
 252 (Helfenstein et al. 2020). Hence, the P pools extracted with  
 253 the stronger extractants (i.e. 0.5 M NaOH and 1 M HCl) are  
 254 most likely less available.

255 **C N P in microbial biomass**

256 The fumigation extraction method was applied to determine  
 257 C, N and P in microbial biomass (AFNOR, 1997). For each  
 258 soil sample, 4 subsamples were weighed, and 2 were fumi-  
 259 gated with chloroform overnight. Blanks without soil were  
 260 included. Then, two fumigated/non-fumigated subsamples

underwent potassium sulphate (K<sub>2</sub>SO<sub>4</sub>, 0.025 M) extrac-  
 261 tion for C and N quantification. The two other fumigated/  
 262 non-fumigated subsamples underwent sodium bicarbonate  
 263 (NaHCO<sub>3</sub> 0.5 M) extraction for P quantification. C and N  
 264 were determined in the filtered extracts using a TOC-TN  
 265 analyser (VCPh Shimadzu + TN module). Phosphorus in  
 266 the fumigated/non-fumigated extracts was measured with  
 267 the malachite green method. Microbial C, N and P (MBC,  
 268 MBN and MBP) were calculated as the difference in C, N  
 269 and P concentration between the fumigated and the non-  
 270 fumigated samples and divided by a conversion factor of  
 271 0.45 for C and N.

273 **Soil micronutrient concentrations**

274 Soil concentration of K<sup>+</sup>, Na<sup>+</sup>, PO<sub>4</sub><sup>2-</sup>, SO<sub>4</sub><sup>2--</sup>, Mg<sup>2+</sup>, Ca<sup>2+</sup>  
 275 were measured at T0 and TF by the laboratory Aurea Agro-  
 276 Sciences (<https://www.aurea.eu/>) following the method  
 277 ISO 11885 for the determination of selected elements by  
 278 inductively coupled plasma optical emission spectrometry  
 279 (ICP-OES).

280 **Crop nutrient uptake**

281 The above-ground biomass was dried at 60 °C for 48 h,  
 282 weighed and ground. Carbon and nitrogen concentrations were  
 283 determined by a CHN elemental analyser (Thermo Fisher Sci-  
 284 entific Flash 2000). P, K, Ca, Mg, Na, Cu, Fe, Mn, Zn, Bo and  
 285 Al concentrations were measured using ICP-AES spectros-  
 286 copy. Nutrient uptake was calculated as the product of tissue  
 287 concentration and dry biomass and expressed in kg ha<sup>-1</sup>.

288 Nitrogen fertilizer use efficiency was calculated in all fer-  
 289 tilized treatments as the difference between the amount of N  
 290 taken up by the crops in the fertilized treatments (NupFT)  
 291 and N taken up by the crops in the control treatment (NupW)  
 292 divided by the total amount of N supplied by each treatment  
 293 (NsupFT).

$$294 NUE(\%) = \frac{NupFT - NupW}{NsupFT} \times 100 \quad (2) \\ 295$$

296 Phosphorus fertilizer use efficiency (PUE) was calculated  
 297 in the same way as NUE.

298 **Modelling of ammonia volatilization**

299 Two models were used to estimate ammonia volatilization in  
 300 our experiment: Hydrus 1D (PC progress, version 4.17) and  
 301 Visual Minteq (version 3.1). Each model was able to give  
 302 an estimation of the ammonia volatilized using a different  
 303 approach. Visual Minteq is an equilibrium speciation model

304 and simulated the concentration of ammonia in the soil solution  
 305 by calculating chemical equilibria using the pH, the  
 306 ionic strength and the concentrations of different chemical  
 307 species. Hydrus 1D, on the other hand, is a reactive transport  
 308 model and simulated ammonia volatilization by a first-order  
 309 reaction process in function of the continuous concentration  
 310 of ammonium in soils (calibrated with the observed mea-  
 311 surements). A comparison of the parameters used for the two  
 312 models is detailed in Table 5S, Supplementary Data. Finally,  
 313 the rates of ammonia volatilization simulated by the two dif-  
 314 ferent models were compared.

### 315 Hydrus 1D model

316 **Model description** Hydrus 1D (Šimůnek et al. 2008) is a  
 317 free software able to simulate water flow and solute transport  
 318 in one dimension. All the following reactions and processes  
 319 were considered in the simulation: ammonia volatilization,  
 320 nitrification, denitrification,  $\text{NO}_3^-$  leaching, N mineraliza-  
 321 tion and  $\text{NH}_4^+$  and  $\text{NO}_3^-$  root uptake.

322 **Input parameters** A homogeneous soil profile of 0–10-cm  
 323 depth was considered.

324 **Water flow parameters** Soil water dynamics were modelled  
 325 according to the Van Genuchten model (van Genuchten  
 326 1980), and the hydraulic parameters were derived from  
 327 the soil texture (sandy loam). An atmospheric bound-  
 328 ary condition (BC) with the surface layer was set for the  
 329 upper BC, and free drainage was set for the lower BC. The  
 330 reference evapotranspiration (ETo) was calculated using  
 331 the Hargreaves formula using temperature and relative  
 332 humidity data. Crop evapotranspiration was calculated as  
 333  $\text{ETP} = \text{Kc} * \text{ETo}$  (with Kc estimated from the spinach growing  
 334 stages) accordingly to the FAO 56 (Allan and Smith 1998).  
 335 Root water uptake was modelled with Feddes parameters  
 336 (Feddes et al. 1978) using the lettuce parameters embedded  
 337 in the software. Finally, the average soil moisture measured  
 338 at T0 was used to set the initial water content.

339 **Solute transport parameters** Ammonium and nitrate were  
 340 the 2 solutes modelled in Hydrus 1D; their molecular dif-  
 341 fusion coefficients in water were 1.52 and 1.64  $\text{cm}^2 \text{ day}^{-1}$   
 342 respectively (Li et al. 2015). The Henry's law constant for  
 343  $\text{NH}_4^+$  was  $2.95 \times 10^{-4}$  (Li et al. 2015), and the adsorption  
 344 coefficient for  $\text{NH}_4^+$  (Kd) was set at 3.5  $\text{L mg}^{-1}$  (Hanson  
 345 et al. 2006). The nitrification and denitrification rates were  
 346 assumed to be 0.2  $\text{day}^{-1}$  and 0.04  $\text{day}^{-1}$  respectively (Castaldelli  
 347 et al. 2018; Li et al. 2015). N mineralization was assumed  
 348 to follow a zero-order reaction process and was calibrated  
 349 with N content from the control treatment. The rate was set at  
 350  $1.5 \times 10^{-6} \text{ g cm}^{-3} \text{ day}^{-1}$ , falling in the same range as the one  
 351 used by Tao et al. (2021) for agricultural soils. The boundary

352 condition at the top was set as "stagnant for volatile solutes"  
 353 on a 1-cm layer to allow gaseous diffusion of solutes (Jury  
 354 et al. 1983). A "zero concentration gradient" was used at the  
 355 bottom of the soil profile to allow N leaching fluxes. Root sol-  
 356ute uptake was set as passive for both solutes with the highest  
 357 soil concentrations as the maximum uptake allowed. Addi-  
 358 tionally, initial soil concentrations in  $\text{NO}_3^-$  and  $\text{NH}_4^+$  were set  
 359 using the average concentration measured per treatment at T0.

360 **Determination of the volatilization rate** The volatilization rate  
 361 was calibrated against soil ammonium and nitrate concentrations  
 362 with the inverse solution model using the U3 treatment configura-  
 363 tion. U3 was chosen because early simulations showed a better  
 364 fitting between simulated and measured  $\text{NH}_4^+$  soil content for  
 365 this treatment. The inverse solution analysis returned a rate of  
 366 1.4  $\text{day}^{-1}$  for ammonia volatilization. This value fits the upper  
 367 range of rates found in the literature (Castaldelli et al. 2018), and  
 368 we considered that it represented well the optimal conditions for  
 369 ammonia volatilization in this experiment (i.e. high soil pH and  
 370 high temperature). This rate was then used to model the nitrogen  
 371 dynamics in the other treatments.

372 **Model evaluation** Simulated ammonium and nitrate soil  
 373 concentrations in the three fertilized treatments (F, U1 and  
 374 U3) were compared to measurements to validate the model.  
 375 The discrepancy between simulated and observed data was  
 376 evaluated by calculating the coefficient of correlation ( $r^2$ )  
 377 and the root mean square error (RMSE). AQ2 7

$$378 \text{RMSE} (\text{kgN.ha}^{-1}) = \sqrt{\frac{1}{n} \sum_i^n (S_i - O_i)^2} \quad (3)$$

379 where  $S_i$  ( $\text{kg ha}^{-1}$ ) and  $O_i$  ( $\text{kg ha}^{-1}$ ) are respectively simu-  
 380 lated and observed nitrogen concentration and  $n$  the number  
 381 of measurements ( $n=8$ ). The observed nitrogen concentra-  
 382 tion represents the mean value per treatment ( $n=4$ ). 383

### 384 Visual Minteq model

385 Visual Minteq is a free software modelling chemical equilib-  
 386 ria (Gustafsson 2011). In our study, it was used to calculate  
 387 the theoretical amount of  $\text{NH}_3$  produced in solution after urine  
 388 application. The input parameters were: soil pH, ionic strength  
 389 and soil solution concentration of major ions, i.e.  $\text{NH}_4^+$ ,  $\text{NO}_3^-$ ,  
 390  $\text{K}^+$ ,  $\text{Na}^+$ ,  $\text{PO}_4^{3-}$ ,  $\text{SO}_4^{2-}$ ,  $\text{Mg}^{2+}$ ,  $\text{Ca}^{2+}$  after each application.  
 391 Except for  $\text{NH}_4^+$  and  $\text{NO}_3^-$ , the other concentrations were  
 392 measured only at the beginning and the end of the experiment.  
 393 Therefore, they were considered equal to their concentration  
 394 at T0 until mid-experiment and then equal to their concentra-  
 395 tion at TF until the end. Regarding phosphorus, the amount of  
 396 P contained in urine was added to the  $\text{PO}_4^{3-}$  concentrations.

397 The other inputs of major ions added with urine additions were  
 398 considered negligible for this specific model application. The  
 399 model outputs give all chemical species in solution likely to  
 400 precipitate at thermodynamic equilibrium and the distribution  
 401 of each element among its different species. The percentage of  
 402  $\text{NH}_3$  after each urine application calculated by the model was  
 403 multiplied by the initial  $\text{NH}_4^+$  concentration to estimate the  
 404 total amount of  $\text{NH}_3$  produced in  $\text{kg ha}^{-1}$  for each treatment. To  
 405 compare Visual Minteq and Hydrus 1D outputs, we assumed  
 406 that all the  $\text{NH}_3$  produced in solution was volatilized.

## 407 N and P budget

### 408 Nitrogen

409 A nitrogen budget approach was used to compare the main  
 410 nitrogen fluxes in our experiment and give an estimation of  
 411 the unaccounted losses (i.e. denitrification and leaching). The  
 412 N budget was calculated for the 0–10-cm depth soil layer  
 413 using the N applied by fertilization ( $\text{Ferti}_N$ ), the measured soil  
 414 concentration of  $\text{NH}_4^+$  and  $\text{NO}_3^-$  at T0 and TF to calculate  
 415  $\Delta\text{Nsoil}$ , the measured crop N uptake between 0 and 10 cm  
 416 ( $\text{Nupt}$ ), the measured microbial N pool ( $\Delta\text{MBN}$ ) and the mea-  
 417 sured soil bulk density to convert values from  $\text{mg N kg}^{-1}$  to  
 418  $\text{kg N ha}^{-1}$ . Ammonia volatilization (Vol) was fitted according  
 419 to the Hydrus model outputs. Nitrogen mineralization (Min)  
 420 was calculated as the only input of N in the water treatment.  
 421 The nitrogen budget error was then calculated (as described  
 422 below) for each treatment; it can be interpreted as a measure  
 423 of the unaccounted losses and of the experimental error. We  
 424 assumed that a budget error below 10% of N supplied means  
 425 that unaccounted losses were minimal.

$$426 \text{Nitrogen budget error } (\delta N) (\text{kg.ha}^{-1}) = \text{Ferti}_N - \Delta\text{Nsoil} - \Delta\text{MBN} \\ - \text{Nupt} - \text{Vol} + \text{Min} \quad (4)$$

427 where

$$429 \Delta\text{Nsoil} = (\text{NH}_4^+ + \text{NO}_3^-)T_F - (\text{NH}_4^+ + \text{NO}_3^-)T_0 \quad (5)$$

$$431 \Delta\text{MBN} = \text{MBNT}_F - \text{MBNT}_0 \quad (6)$$

$$433 \text{Nupt} = \text{N}_{\text{uptake}} \times \% \text{root biomass}_{0-10\text{cm}} \text{ (estimated at 44\% according to) } \quad (7) \\ \text{And Heinrich et al. (2013)}$$

### 435 Phosphorus

436 The P budget was calculated for the 0–10-cm depth soil  
 437 layer excluding the pool of P extractable with HCl. As the  
 438 HCl-extractable P pool is very large in our calcareous soil,  
 439 variations occurring during the experiment were not detect-  
 440 able against the analytical error. Hence, the P budget error  
 441 can be interpreted as a net variation of the HCl-extractable

442 P pool size. Similarly, a net change in a P pool size gives an  
 443 indication of the net flux involving the pool; however, this  
 444 variation also includes the experimental error. The P budget  
 445 error is calculated with the amount of P supplied by ferti-  
 446 lization ( $\text{Ferti}_P$ ), the P taken up by the crops between 0 and  
 447 10 cm ( $\text{Pupt}$ ) and the concentration of P in the different soil  
 448 pools ( $\text{Pi}_{\text{resin}}$ ,  $\text{Pi}_{\text{HCO}_3}$ ,  $\text{Pi}_{\text{NaOH}}$ ,  $\text{Po}_{\text{HCO}_3}$ ,  $\text{Po}_{\text{NaOH}}$ ). Microbial P is  
 449 already included in the organic P pools due to the extraction  
 450 method; therefore, it does not appear in the budget calculation.

$$451 \text{P budget error } (\delta P) (\text{kg.ha}^{-1}) = \text{Ferti}_P - \Delta\text{Psoil} - \text{Pupt} \quad (8)$$

453 where

$$454 \Delta\text{Psoil} = (\text{Pi}_{\text{resin}} + \text{Pi}_{\text{HCO}_3} + \text{Pi}_{\text{NaOH}} + \text{Po}_{\text{HCO}_3} + \text{Po}_{\text{NaOH}})T_F \\ - (\text{Pi}_{\text{resin}} + \text{Pi}_{\text{HCO}_3} + \text{Pi}_{\text{NaOH}} + \text{Po}_{\text{HCO}_3} + \text{Po}_{\text{NaOH}})T_0 \quad (9)$$

455 P crop uptake between 0 and 10 cm is calculated the same  
 456 way as the N crop uptake on 0–10 cm (see above). 457

## 458 Statistical analysis

459 Statistical analyses were carried out with Rstudio software (ver-  
 460 sion 3.6.1) (R Core Team 2017). To test the homogeneity of ini-  
 461 tial variable values at T0 across blocks (tanks) and treatments,  
 462 we used a linear model with 2 factors: block ( $n=4$ ) and treat-  
 463 ment ( $n=4$ ) (16 samples in total), and significance was deter-  
 464 mined with the ANOVA (analysis of variance) function. Signif-  
 465 icant differences were only found for the variables Pi resin, and  
 466 MBN with differences of up to 41% and 52% respectively in the  
 467 mean pool size between the richest and the poorest tank. On T1  
 468 to TF values, ANOVAs were performed to assess the effect of  
 469 treatments on each variable and date separately. For each vari-  
 470 able measured, a linear mixed model (“lmer” function, lme4  
 471 package, (Bates et al. 2015, p. 4)) was produced with “treat-  
 472 ment” ( $n=4$ ) as fixed factor and “block” ( $n=4$ ) as random fac-  
 473 tor to account for the initial differences between tanks observed  
 474 for certain variables. A Tukey’s multi-comparison test was per-  
 475 formed when the treatments had a significant effect on the vari-  
 476 able (significance level:  $p$  value  $< 0.05$ ). Correlations between  
 477 variables were assessed using Pearson correlation tests.

## 478 Results

### 479 Global result analysis

#### 480 Soil analysis

481 At the end of the experiment, soil pH was lower in the U3  
 482 treatment (urine at  $510 \text{ kg N ha}^{-1}$ ) than in all other treat-  
 483 ments, reaching 8.5, while it was over 8.8 in all other

484 treatments. At T3, shortly after a fertilizer application, soil  
 485 conductivity was higher in all the fertilized treatments than  
 486 in the water treatment, but at the end of the experiment,  
 487 only the U3 treatment still had higher conductivity (+ 187%  
 488 compared to the water treatment,  $p = 0.002$ ) (Table 3). In  
 489 the U3 treatment, soil concentrations in  $\text{Na}^+$  and  $\text{Cl}^-$  were  
 490 also twice as high at TF as in the other treatments (Table 2S,  
 491 Supplementary data). MBC and MBP showed no significant  
 492 response, but MBN showed a significant difference between  
 493 U3 and the control W at TF (+ 370% in U3 compared to W)  
 494 (Table 3). Soil nitrate and ammonium concentrations and  
 495 soil P pool concentrations are commented in Sect. 2.1 and  
 496 3 respectively.

#### 497 Biomass and plant nutrient uptake

498 Aboveground biomass at the end of the experiment was  
 499 significantly different only between the control treatment  
 500 (W) and the synthetic fertilizer treatment (F). N and P  
 501 uptake by plants were significantly higher ( $p < 0.05$ ) in the  
 502 F and U3 treatments than in the W treatment, and inter-  
 503 mediate in the U1 treatment, i.e. not significantly differ-  
 504 ent from both W and U3 and F treatments. In addition,  
 505 the spinach crops in the tanks 1 and 8 took up more N  
 506 and P than the ones in the tanks 4 and 5 (up to 138% and  
 507 210% more for N and P respectively, data not shown). N  
 508 and P uptake were highly correlated (Pearson coefficient  
 509  $r = 0.97$ ,  $p < 0.0001$ ). N uptake was correlated with soil  
 510 nitrate content from day 23 to day 28 ( $p = 0.03$ ,  $p = 0.04$ ,  
 511  $p = 0.01$  for day 23, 26 and 28 respectively) but was not  
 512 correlated with soil ammonium content. P uptake was cor-  
 513 related with none of the P pools at the end of the exper-  
 514 iment ( $p > 0.05$ ) (Table 4).

#### 515 Nitrogen stock evolution and losses

##### 516 $\text{N-NH}_4^+$ , $\text{N-NO}_3^-$ soil concentrations

517 In the water treatment, soil ammonium and nitrate contents  
 518 were very close to 0  $\text{kgN ha}^{-1}$  throughout the experiment  
 519 (Fig. 3). In the synthetic fertilizer treatment (F), both nitrate  
 520 and ammonium increased after the first application (day 10),  
 521 but, while ammonium content stayed relatively stable under  
 522 100  $\text{kgN ha}^{-1}$ , nitrate content increased up to 150  $\text{kgN ha}^{-1}$   
 523 at day 30. In U3 and U1 treatments, ammonium reached a  
 524 peak during the experiment (at day 30 and day 20 respec-  
 525 tively) and then decreased to 0  $\text{kgN ha}^{-1}$  at the end (Fig. 3,  
 526 left panel). Nitrate concentration in U1 and U3 showed a  
 527 slow increase between day 10 and 20 and a sharp increase  
 528 between day 20 and 30 (Fig. 2). Overall, the nitrate and  
 529 ammonium curves of U1 and F were very similar, although  
 530 in the U1 treatment twice more ammonium was added than  
 531 in the F treatment.

#### 532 Estimation of ammonia volatilization

533 In order to validate the N fluxes simulated by Hydrus 1D,  
 534 simulated and observed ammonium and nitrate soil concen-  
 535 trations were compared (Fig. 4).

536 Modelled  $\text{NH}_4^+$  soil concentrations agreed relatively well  
 537 with measurements in the 3 treatments ( $\text{RMSE} < 69 \text{ kg ha}^{-1}$ ),  
 538 although they were slightly underestimated for U3. However,  
 539  $\text{NO}_3^-$  dynamics were poorly simulated by Hydrus 1D espe-  
 540 cially in the U3 treatment ( $\text{RMSE} = 142 \text{ kg ha}^{-1}$ ). As soil  
 541  $\text{NO}_3^-$  is not used in the calculation of ammonia volatilization,  
 542 this poor fitting should not affect the estimation of  $\text{NH}_3$  volatil-  
 543 ization. It prevents, however, a good estimation of  $\text{NO}_3^-$  leach-  
 544 ing and denitrification. Therefore, we did not use the Hydrus  
 545 1D model outputs to estimate these losses in the N budget.

546 As expected, ammonia volatilization simulated by  
 547 Hydrus 1D was especially high in the U3 treatment reach-  
 548 ing 260  $\text{kg ha}^{-1}$ , and it was almost two times higher in the  
 549 U1 treatment than in the F treatment (Table 5).

550 Visual Minteq estimated that 21% of  $\text{NH}_4^+$  in soils would  
 551 be in  $\text{NH}_3$  form for a soil at  $\text{pH} = 8.7$ ; however, this pro-  
 552 portion decreases to 13.9% when the pH is at 8.5 as in the  
 553 treatment U3 at the end of the experiment (Table 1S, Sup-  
 554 plimentary data).

555 Despite the models' dissimilarities, their output for the flux  
 556 of  $\text{NH}_3$  is similar (Table 5). Both models agree that more than  
 557 half of the nitrogen applied with the urine fertilizer was lost  
 558 by ammonia volatilization, whereas only approximately 30%  
 559 was lost with the  $\text{NH}_4\text{NO}_3$  synthetic fertilizer.

#### 560 Fate of phosphorus from urine

561 P was similarly distributed in the different P pools in F,  
 562 U1 and W treatments, whereas in U3 treatment, the three  
 563 inorganic phosphorus pools were larger than in all others  
 564 at the end of the experiment, in particular Pi resin (+ 179%  
 565 compared to the W treatment) ( $p = 0.00005$ ) and Pi NaOH  
 566 (+ 38%) ( $p = 0.0015$ ). (Fig. 5).

567 In contrast, the HCl-extractable Pi (Pi HCl) did not vary  
 568 significantly among treatments (Table 3). The P budget  
 569 calculation ( $\delta_P$ ) was negative in the U1 and U3 treatments  
 570 suggesting a potential decrease of approximatively 10  $\text{kgP}$   
 571  $\text{ha}^{-1}$  of the Pi HCl pool, which was not detectable against  
 572 replicate variability (Table 1S, Supplementary data).

## 573 Discussion

### 574 Effect of urine fertilization on soil pH and salinity

575 In our study, only the excessive dose of urine (U3) lowered  
 576 the pH of the calcareous soil. Previous studies on acidic soils  
 577 found a decrease in soil pH even with the appropriate urine

**Table 3** Results of ANOVA (p value) and post hoc Tukey tests on the experiment variables. Values in the table are the means of the 4 replicates in each treatment. Treatments are noted with letters, W=control, F=synthetic fertilizer, U1=urine dosed at 170 kgN ha<sup>-1</sup>, U3=urine in excess at 510 kgN ha<sup>-1</sup>. Different letters (a, b, c) indicate significant differences between treatments (p<0.05) for a same date (T0 at the beginning of the experiment before sowing, T1 to T6 right after every fertilization event and TF at the end of the experiment)

Variables	Time	W	F	U1	U3	p value treatment effect	p value tank effect
pH	T0	8.73 a	8.76 a	8.73 a	8.74 a	0.3	0.4
	TF	9.01 a	8.84 b	8.88 ab	8.52 c	<0.0001 ***	Nd
Conductivity (μS cm <sup>-1</sup> )	T0	160 a	135 a	155 a	149 a	0.23	0.33
	T3	150 c	248 bc	341 b	528 a	<0.0001	Nd
	TF	147 b	154 b	177 b	423 a	0.002*	Nd
NO <sub>3</sub> <sup>-</sup> +NO <sub>2</sub> <sup>-</sup> (mgN kg <sup>-1</sup> )	T0	7.6 a	5.0 a	8.2 a	6.3 a	0.21	0.6
	T1	2.4 a	19.1 a	4.8 a	3.9 a	0.06	Nd
	T2	8.8 b	47.0 a	23.1 ab	42.5 a	0.002 *	Nd
	T3	6.6 b	44.3 a	26.5 ab	39.7 a	0.01 *	Nd
	T4	10.4 c	88.8 b	80.1 bc	226.1 a	0.00004 ***	Nd
	T5	13 c	135.9 b	104.8 b	221.3 a	0.00001 ***	Nd
	T6	6.7 c	102.7 b	57.1 bc	214.3 a	<0.00001 ***	Nd
	TF	4.1 b	22 b	22 b	121 a	0.003 **	Nd
NH <sub>4</sub> <sup>+</sup> (mgN kg <sup>-1</sup> )	T0	0 a	0 a	0 a	0 a	Nd	Nd
	T1	0 b	25 ab	22 ab	58 a	0.01	Nd
	T2	2.9 c	35 bc	66 b	145 a	0.0001	Nd
	T3	1.1 c	38 c	86 b	187 a	<0.00001 ***	Nd
	T4	0 b	35 b	76 b	206 a	0.0001	Nd
	T5	0.3 b	35 b	51 b	260 a	<0.00001 ***	Nd
	T6	0.8 b	51 b	52 b	315 a	0.0001	Nd
	TF	0.6 a	0.9 a	0.8 a	1 a	0.5	Nd
Pi Resin (mgP kg <sup>-1</sup> )	T0	10.5 a	9.7 a	9.7 a	10.8 a	0.4	0.009 **
	TF	9.4 b	11.7 b	11.7 b	26.2 a	0.00005 **	Nd
Pi Bicarbonate (mgP kg <sup>-1</sup> )	T0	7.8 a	11.6 a	9.3 a	8.9 a	0.65	0.28
	TF	11.5 a	11.5 a	11.1 a	17.0 a	0.05	Nd
Pi NaOH (mgP kg <sup>-1</sup> )	T0	11.5 a	9.9 a	9.5 a	9.58 a	0.13	0.02 *
	TF	9.8 b	10.9 b	9.9 b	13.5 a	0.0015 **	Nd
Pi HCl (mgP kg <sup>-1</sup> )	T0	224 a	209 a	206 a	221 a	0.45	0.10
	TF	225 a	217 a	219 a	230 a	0.42	Nd
Po Bicarbonate (mgP kg <sup>-1</sup> )	T0	10.1 a	5.7 a	7.5 a	7.7 a	0.33	0.07
	TF	6.9 a	12.8 a	12.1 a	8.1 a	0.43	Nd
Po NaOH (mgP kg <sup>-1</sup> )	T0	24.5 a	22.9 a	20.5 a	22.5 a	0.81	0.09
	TF	23.9 a	21.3 a	23.8 a	20.9 a	0.6	Nd
P total (mgP kg <sup>-1</sup> )	T0	281.3 a	269.9 a	254.3 a	289.3 a	0.21	0.07
	TF	286.9 a	285.8 a	288.5 a	316.3 a	0.03 *	Nd
MBC (mgC kg <sup>-1</sup> )	T0	127 a	134 a	140 a	140 a	0.67	0.03
	TF	96 a	122 a	126 a	134 a	0.10	Nd
MBN (mgN kg <sup>-1</sup> )	T0	9.8 a	11.3 a	11.1 a	10.5 a	0.17	0.001 **
	TF	8.4 b	20.1 ab	21.3 ab	39.5 a	0.05 *	Nd
MBP (mgP kg <sup>-1</sup> )	T0	7.1 a	7.1 a	8.8 a	7.6 a	0.9	0.8
	TF	6.3 a	5.1 a	7.8 a	10.5 a	0.08	Nd

578 dosage (Mnkeni et al. 2008; Sangare et al. 2015). However,  
579 calcareous soils have a stronger pH buffering capacity (Mag-  
580 dodd and Bartlett 1985; Raza et al. 2021); they are probably

more resistant to the acidifying effect of urine in the short term. The effect of long-term urine application on soil pH is uncertain and is potentially much greater for non-calcareous

581  
582  
583

**Table 4** Results of ANOVA (p value) and post hoc Tukey tests on plant biomass and plant nutrient uptake variables assessed at the end of the experiment. NUE and PUE (respectively nitrogen and phosphorus use efficiency) were assessed as the efficiency of N and P fertilizer, i.e. the difference in nutrient uptake between a fertilized and non-fertilized treatment, divided by the amount of applied nutrient. Values in the table are the means of the 4 replicates. Different letters (a, b, c) indicate significant differences between treatments ( $p < 0.05$ ) for a same date

Variables	W	F	U1	U3	<i>p</i> value treatment effect
Aboveground biomass (g)	15.2 b	27.8 a	22.3 ab	23.2 ab	0.02 *
Nuptake (kg/ha)	23.6 b	52.9 a	41.8 ab	49.4 a	0.02 *
Puptake (kg/ha)	2.2 b	4.3 a	3.1 ab	4.1 a	0.03 *
C/N	10.19 a	8.03 b	8.21 b	7.57 b	0.005 **
C/P	130 a	106 a	119 a	101 a	0.17
NUE	/	17 a	10 a	5 a	0.06
PUE	/	24 a	10 a	7 a	0.05

584 than for calcareous soils. Soil acidification can nevertheless  
 585 be reduced by applying correct amounts and most importantly by reducing urine-derived ammonia volatilization,  
 586 which exacerbates soil acidification.

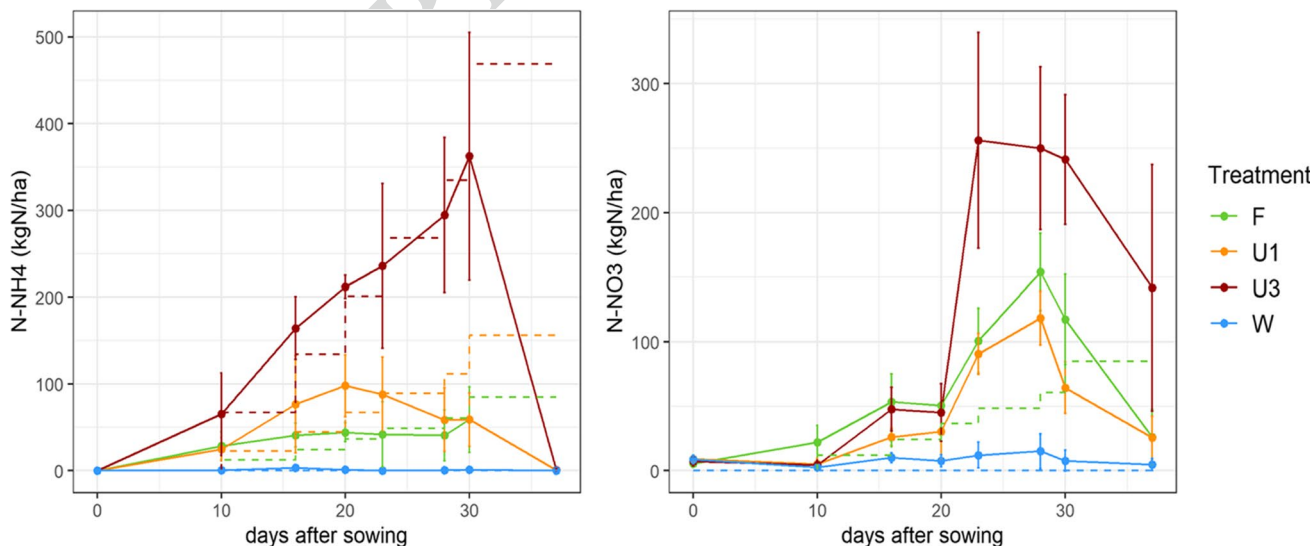
587 Similarly, urine caused a buildup of soluble salt concentration and specifically NaCl only in the U3 treatment.  
 588 However, crops under the U3 treatment did not take up more  
 589 sodium than under the other treatments (Table 3S, Supplementary data), and the soil salinity was still below the  
 590 591 spinach salt tolerance threshold established at  $9.4 \text{ dS m}^{-1}$   
 592 which exacerbates soil acidification.

593 by Ferreira et al. (2018); so, it is unlikely that the spinach crops underwent a saline stress. Other studies on urine fertilization confirm that critical values of soil conductivity are only observed when urine is applied in excess (up to  $13 \text{ dS m}^{-1}$  with  $800 \text{ kgN ha}^{-1}$ ) (Boh and Sauerborn 2014; Mnkeni et al. 2008; Neina 2013) and that the crop response depends on its salt tolerance threshold (Mnkeni et al. 2008). Nevertheless, the effects of urine fertilization on soil salinity have not been investigated so far for longer than one or two cropping seasons. The sodium adsorption ratio (SAR) of diluted urine being high (Table 2), the potential effect of urine on the buildup of harmful concentrations of soil exchangeable sodium should be assessed in the long term, and suitable solutions can be envisaged to avoid such a risk, such as the addition of Ca and Mg amendments (Ayers and Westcot 1985).

### Considerable losses of nitrogen by volatilization

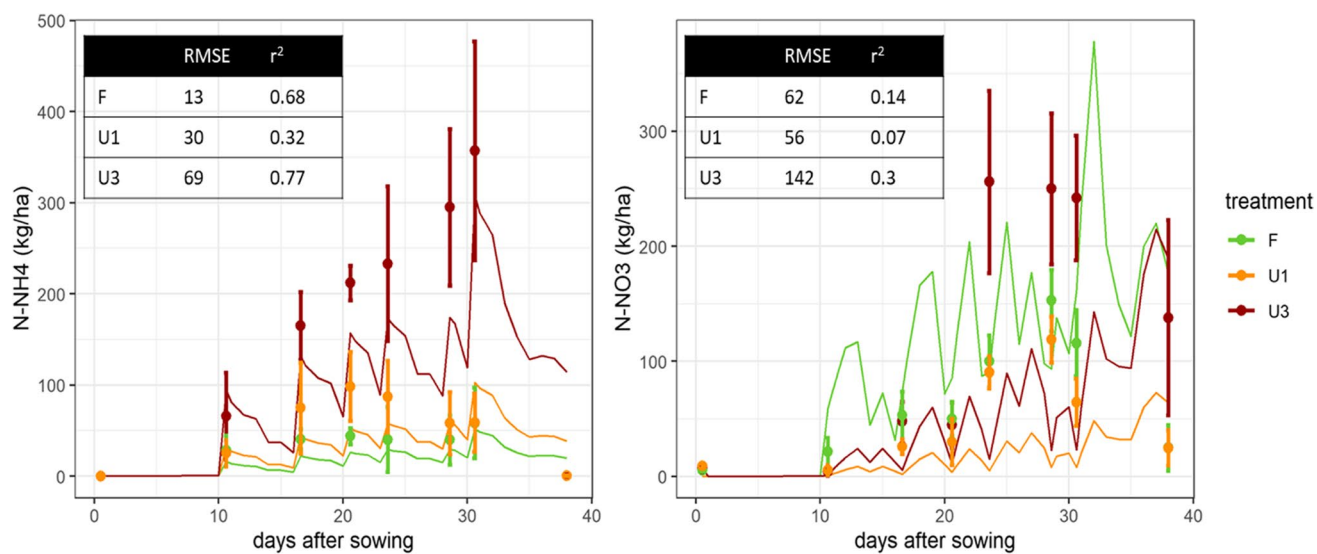
594 More than half of the nitrogen applied was estimated to be  
 595 lost by volatilization with urine application. Similar rates of  
 596 volatilization were obtained with liquid ammonium fertilizer  
 597 on calcareous soils (Hargrove et al. 1977; Powlson and Dawson 2022; Whitehead and Raistrick 1990). In addition, volatilization is almost double with urine than with ammonium  
 598 nitrate fertilizer; similar values were found when comparing  
 599 urea and ammonium nitrate (Eckard et al. 2003).

600 According to Visual Minteq simulations, a small shift  
 601 in soil pH can considerably affect the  $\text{NH}_4/\text{NH}_3$  chemical  
 602 equilibrium and so the potential of our soil for ammonia  
 603 volatilization. This highlights the interaction between  
 604



**Fig. 3** Evolution of soil mineral nitrogen stocks during the experiment as a function of treatment: F: synthetic fertilizer dose  $\times 1$ , U1: urine dose  $\times 1$ , U3: urine dose  $\times 3$ , W: unfertilized control (ammonium on the left panel and nitrate on the right panel). The dashed lines represent the cumulative fertilization inputs and the errors bars represent the standard error between the four replicates

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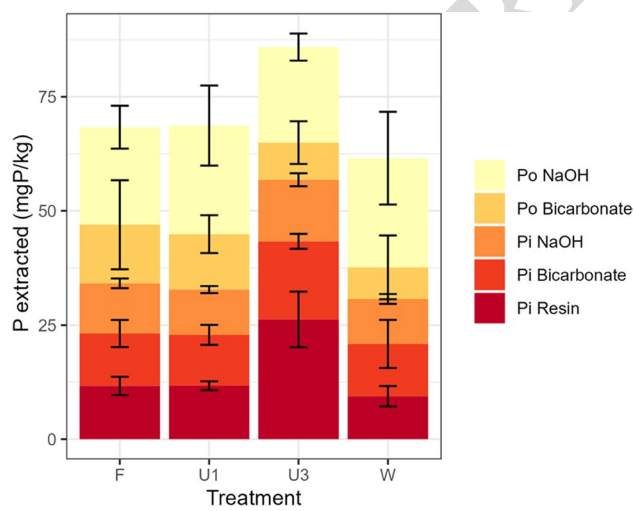


**Fig. 4** Comparison between concentration of ammonium and nitrate in soils, as simulated by Hydrus 1D (lines) and observed (points) for which the errors bars represent the standard error between the four

replicates. Tables in the figures report the RMSE (in  $\text{kg ha}^{-1}$ ) and  $r^2$  calculated for each treatment

**Table 5** Nitrogen lost by ammonia volatilization, as modelled by Visual Minteq and Hydrus 1D models

Treatment	Hydrus	Visual Minteq
U1	57% (90 $\text{kg ha}^{-1}$ )	67% (105 $\text{kg ha}^{-1}$ )
U3	55% (260 $\text{kg ha}^{-1}$ )	58% (270 $\text{kg ha}^{-1}$ )
F	28% (48 $\text{kg ha}^{-1}$ )	38% (66 $\text{kg ha}^{-1}$ )



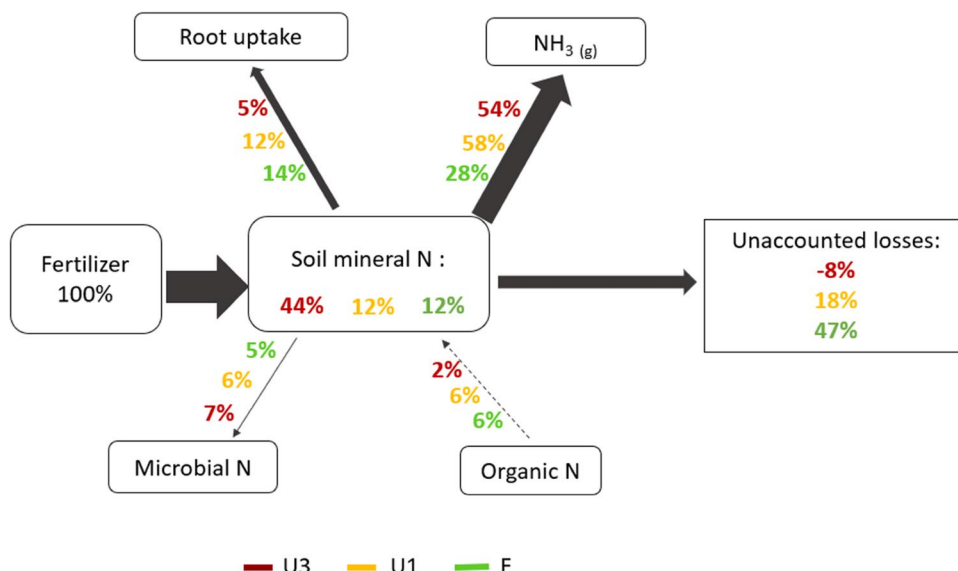
**Fig. 5** Distribution of phosphorus in different soil pools at the end of experiment. The errors bars represent the standard error of four replicates. Pi Resin=resin extractable inorganic P; Pi Bicarbonate=sodium bicarbonate extractable inorganic P; Po Bicarbonate=sodium bicarbonate extractable organic P; Pi NaOH=sodium hydroxide extractable inorganic P; Po NaOH=sodium hydroxide extractable organic P

fertilization, soil pH and ammonia volatilization that needs to be considered when studying nitrogen fluxes. It is likely that a high load of urine or long-term urine fertilization by causing high losses through ammonia volatilization and high nitrification rates decreases the pH of calcareous soils, thus decreasing the potential of the soil for subsequent ammonia volatilization.

The nitrogen budget error was especially high in the F treatment (Fig. 6 and Table 4S, Supplementary data), suggesting that unaccounted losses, most probably leaching and denitrification, had a higher contribution in the N budget of this treatment. Hence, nitrate leaching and denitrification were probably higher in the fertilizer treatment than in the urine treatments. Most likely, the  $\text{NH}_4\text{NO}_3$  fertilizer induced a smaller volatilization rate but higher leaching and denitrification rates because of its partition between nitrate and ammonium (Eckard et al. 2003; Fernández-Escobar et al. 2004). In addition, such high levels of excess N were not expected as the U1 and F fertilization rate had supposedly been adjusted to meet crop requirements, but crop growth was limited in our experiment possibly because of excessive heat during that summer. In conditions of adequate crop uptake, lower total N losses are to be expected.

In this experimental setup, soil conditions were optimal for ammonia volatilization (pH at 8.7 and air temperature ranging between 20 and 30 °C), so the high rates of volatilization simulated in the urine treatments can be considered an upper limit for urine fertilization. Although these high ammonia emissions raise concern, it is possible to reduce them with appropriate application techniques, timing and dosage (Mencaroni et al. 2021; Rodhe et al. 2004).

**Fig. 6** Nitrogen fate in each fertilized treatment on a 0–10-cm soil profile. Percentages represent the estimated percentage of the N supplied by fertilizer that had followed each pathway at the end of the experiment. Unaccounted losses are calculated by the nitrogen budget error. A small budget error (< 10% of N supplied) suggests that unaccounted losses are close to 0



## 654 Bioavailability of N from urine fertilizer for plants 655 and microbes

656 In our experiment, most of the ammonium supplied by urine  
657 was readily nitrified in the soil. This result is consistent with  
658 other studies with urine fertilization showing that the nitrogen  
659 applied was in nitrate form at the end of their experiment  
660 (Cuttle et al. 2001). Ledgard and Saunders (1982) observed  
661 a nitrate peak 10 days after fertilization, which is consistent  
662 with our peak 13 days after the first urine application.  
663 Ammonium oxidation into nitrite is faster than nitrite oxidation  
664 into nitrate (Monaghan and Barraclough 1992). As  
665 with our analytical method we could not distinguish nitrate  
666 and nitrite, we cannot exclude that nitrite might have been  
667 present along with nitrate. This could have caused nitrogen  
668 stress in the urine treatments, especially at the start of  
669 vegetative growth, given that nitrite is not bioavailable and  
670 that spinach prefers nitrate over ammonium (Okazaki et al.  
671 2009). The lower crop uptake in the U1 treatment (Table 4)  
672 could support the hypothesis of a nitrogen limitation due to  
673 incomplete nitrification at the beginning of the experiment.  
674 Another explanation for the lower N uptake in the U1 treatment  
675 could lie in the high rate of ammonia volatilization,  
676 reducing the amount of N available in the soil. In this case,  
677 the nitrogen stress should have occurred at the end of the  
678 growth cycle where the N demand was higher. However, the  
679 plant growth data are not sufficiently detailed to determine  
680 whether or when nitrogen stress occurred in the U1 treatment  
681 (Fig. 2S, Supplementary data).

682 The microbial N pool appears to have increased in the  
683 fertilized treatments between the beginning and end of  
684 the experiment, with a stronger response in the U3 treatment.  
685 Microbes were, therefore, able to immobilize N from  
686 urine, making it unlikely that urine had a negative effect on

687 microbial activity. However, this increase was never accom-  
688 panied by an increase in microbial C, causing a shift in the  
689 microbial C to N ratio. Mason-Jones et al. (2022) recently  
690 highlighted that soil microbes have the capacity to store  
691 surplus nutrients to reduce their loss and release them later  
692 upon microbial death. Thus, under urine application, part  
693 of the added ammonium was probably stored in the micro-  
694 bial biomass forming a readily available N pool. Similarly,  
695 Zaman et al. (2006) found an increase of only microbial N  
696 with NH<sub>4</sub>Cl fertilizer, while both N and C microbial pools  
697 increased with C-rich dairy shed effluent. Urine alone is  
698 relatively poor in dissolved organic carbon (Table 2) and,  
699 therefore, does not stimulate microbial growth.

## 700 Bioavailability of P from urine fertilizer

701 N and P uptake were highly correlated (Pearson coefficient  
702  $r=0.97$ ). Therefore, the lower P uptake observed for the U1  
703 treatment might be a consequence of the nitrogen stress that  
704 likely occurred under U1.

705 At low doses, the fate of phosphorus from urine and  
706 synthetic fertilizer seemed relatively similar as there was  
707 no difference in concentrations of the different P pools or  
708 total P between U1 and F. The fate of P from urine was  
709 clearly detectable in the U3 treatment, where the high dose  
710 of urine supplied significantly increased the available P and  
711 secondarily the less available P pools (Fig. 5). Therefore,  
712 phosphorus from urine distributed mostly into available and  
713 slightly available inorganic P (Pi Bicarbonate) with a resid-  
714 ual effect that was still detectable after 5 weeks of cropping.  
715 This result is in line with a study by Pandorf et al. (2018)  
716 finding that P from urine can be taken up by snap beans and  
717 turnips. In another experiment, Bonvin et al. (2015) applied

718 synthetic and nitrified urine and found a similar P recovery  
 719 rate in the crops as under mineral fertilizer.

720 In contrast, the application of urine did not affect the size  
 721 of organic P pools, including microbial P. Again, this might  
 722 be explained by the low concentration of dissolved organic C  
 723 in urine, which does not foster microbial activity or growth.  
 724 However, unlike nitrogen, there was no storage of P in the  
 725 microbial pool. According to Chen et al. (2019), the more  
 726 the element is limiting the more it is stored once it becomes  
 727 available. At T0, the soil concentration in mineral nitrogen  
 728 was around  $8 \text{ mgN kg}^{-1}$ , while the available phosphorus (Pi  
 729 resin + Pi Bicarbonate) was around  $19 \text{ mgP kg}^{-1}$ . Hence,  
 730 at the start of the experiment N was more limiting than P,  
 731 explaining the stronger N than P storage in the microbial  
 732 biomass.

733 Additionally, in the U3 treatment, P availability was  
 734 likely increased by the decrease in soil pH (0.2 units) causing  
 735 calcium phosphate dissolution. In our experiment, it is  
 736 likely that the subsequent mobilized phosphate was partially  
 737 adsorbed on the soil exchange complex (Frossard et al. 1995)  
 738 explaining the increase of iron and aluminium bonded P  
 739 (Pi NaOH pool) (Adhami et al. 2006; Prietzel et al. 2016)  
 740 and partially remained in solution increasing the P available  
 741 pool.

## 742 Conclusion

743 In this experiment, urine supplied in appropriate doses  
 744 had a very similar behaviour to that of a synthetic fertil-  
 745 izer in a calcareous soil suggesting that other compounds  
 746 present in urine did not affect N and P uptake by the crop  
 747 and their fate in the soil in the short term. Appropriate  
 748 doses of urine altered neither soil pH nor soil conductivity,  
 749 and provided nitrogen and phosphorus in bioavailable  
 750 forms. Supplied phosphorus was either taken up by the  
 751 crop or dissolved in the soil, and the ammonium was nitrified  
 752 within about 10 days after application. However, this  
 753 study raises awareness on the amount of ammonia that  
 754 can be lost by volatilization in a calcareous soil. Indeed,  
 755 chemical equilibrium and solute transfer models agreed on  
 756 the estimation that about half of the N applied with urine  
 757 could be lost by volatilization. Therefore, it is imperative  
 758 to find more suitable fertilization techniques to mitigate  
 759 ammonia volatilization on alkaline soils. Overall, our  
 760 results demonstrate that although direct usage of human  
 761 urine for fertilization is possible, timing, dosage and soil  
 762 type need to be carefully considered for this practice to be  
 763 environmentally sustainable. Future studies are encour-  
 764 aged to focus on the long-term effect of urine fertilization  
 765 especially on soil pH and salinity which are key factors  
 766 of soil quality and disentangle the various indirect effects  
 767 that human urine could have on soil and plant health.

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 774 odology, resources, funding acquisition, writing—review and editing.  
 775 CM: conceptualization, methodology, resources, funding acquisition,  
 776 writing—review and editing. NAM: resources, funding acquisition,  
 777 writing—review and editing. DC: formal analysis, writing—review  
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## 783 Declarations

784 **Ethical approval** Not applicable.

785 **Consent to participate** All authors gave their consent to participate in  
 786 the conception of this paper.

787 **Consent for publication** All authors read and approved the final manu-  
 788 script.

789 **Competing interests** The authors declare no competing interests.

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