Investigating the synergistic potential Si and biochar to immobilize soil Ni in a contaminated calcareous soil after Zea mays L. cultivation

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Abstract. Silicon (Si) is a beneficial plant element that has been shown to mitigate the effects of 15 potentially toxic elements (PTEs) on crops. Biochar is a soil amendment that sequesters soil 16 17 carbon, and that can immobilize PTEs and enhance crop growth in soils. No previous studies have examined the potentially synergistic effect of Si and biochar on soil Ni chemical fractions and 18 immobilization. Therefore, the aim of this study was to examine the interaction effects of Si levels 19 20 and biochars, to alleviate soil Ni bioavailability and its corresponding uptake in corn (Zea Mays) in a calcareous soil. A 90-day factorial greenhouse study with corn was conducted. Si application 21 levels were applied at 0 (S_0), 250 (S_1) and 500 (S_2) mg Si kg⁻¹ soil and biochar treatments (3% wt.) 22 23 including rice husk (RH) and sheep manure (SM) biochars produced at 300°C and 500°C (SM300, 24 SM500, RH300 and RH500). At harvest, corn shoot Ni-concentrations, soil chemical Ni fractions and DPTA-release kinetics were determined. Simultaneous utilization of Si and SM biochars led 25 to a synergistic reduction (15-36%) of soluble and exchangeable soil Ni fractions compared to 26 application of Si (5-9%) and SM (5-7%) biochars separately. The application of the Si and biochars 27 also decreased DPTA-extractable Ni and corn Ni shoot concentration (by up to 57%), with the 28 combined application of $SM500+S_2$ being the most effective. These effects were attributed to the 29 transformation of Ni from more bioavailable fractions to more stable iron oxide bound fractions, 30 related to soil pH increase. The SM500 was likely the most effective biochar due to its higher 31 alkalinity and lower acidic functional group content which enhanced Ni sorption reactions with Si. 32 The study demonstrates the synergistic potential of Si and sheep manure biochar at immobilizing 33 Ni in contaminated calcareous soils. 34

35 **1 Introduction**

One of the most important ways for potentially toxic elements (PTEs) to enter the human food chain is through the consumption of plants grown in soils contaminated with PTEs. Potentially toxic elements pollute soil environments as a result of mining, metal smelting, using sewage sludge and domestic and industrial effluents in agriculture especially in developing countries (Liu et al., 2018). Potentially toxic elements in soils cannot undergo biodegradation by living organisms, so they possess great stability and longevity in the soil (Poznanović Spahić et al., 2019). Unlike other PTEs found in soils, such as mercury (Hg), cadmium (Cd) and lead (Pb),

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nickel (Ni) is essential for plant growth at very low concentrations. Nevertheless, at elevated 43 contents (>35 mg Ni kg⁻¹ soil), it causes many physiological and morphological malfunctions in 44 plants and severely stunts their growth (Shahzad et al., 2018; Antoniadis et al., 2017). In 45 46 agricultural soils of Iran in the vicinity of the industrial areas, the weighted average concentration of Ni is 349.8 mg kg⁻¹ soil. In these soils, the pollution index (the ratio of the element concentration 47 to the standard concentration) calculated for the Ni is greater than 5, which indicates a severe 48 degree of pollution from the point of view of environmental protection (Shahbazi et al., 2022). 49 50 Shahbazi et al. (2020) collected 711 agricultural soil samples from different climates of Iran and reported that the Ni content in the soils was between 2.79 mg kg⁻¹ and 770 mg kg⁻¹ with an average 51 of 68 mg kg⁻¹ soil. The results showed that the concentration of Ni in 11.3% of these soils was 52 higher than the threshold value. Removing PTEs from contaminated sites via traditional methods 53 such as pump and treat technologies, soil washing and excavation is very expensive and time-54 consuming, therefore, for plant cultivation in these areas, low-cost and effective methods should 55 be sought to stabilize PTEs in soils and prevent them from being transferred to the plant (Gao et 56 al., 2023). 57

Silicon (Si) is a valuable nutrient for plant growth, and it is only considered essential for 58 59 some plant species such as rice. Applying Si to the soil can enhance plant resistance against biological and non-biological tensions, including physiological stress caused by PTEs in soil (Bhat 60 et al., 2019; Yan et al., 2018). The use of Si to promote plant growth and mitigate PTEs toxicity is 61 becoming increasingly popular in agriculture (Li, 2019; Adrees et al., 2015). The application of Si 62 in soils contaminated with PTEs may reduce PTEs bioavailability by increasing soil pH, increasing 63 the secretion of organic ligands by the roots and forming insoluble compounds with PTEs, and 64 ultimately enhancing plant growth (Bhat et al., 2019; Xiao et al., 2021). The soil pH increase 65 associated with Si application is attributed to the hydrolysis reaction of the silicate anion in soil 66 solution which generates hydroxyl ions (Ma et al. 2021). 67

Biochar is an organic soil amendment that sequesters soil carbon (C) that has received 68 69 much attention in recent years to stabilize PTEs in polluted sites (El-Naggar et al., 2018). Biochar is a carbon-rich, porous organic material which is prepared in a limited or no oxygen conditions 70 by pyrolysis of organic wastes, including crop and animal residues, urban waste, wood byproduct 71 72 (Vickers, 2017; Ankita Rao et al., 2023). The organic surface functional groups of biochar such as carboxylic and phenolic groups provide cation exchange capacity in soils (Tomczyk et al., 2020). 73 74 Addition of biochar to the soil not only improves the soil chemical and physical properties, but also reduces the bioavailability of PTEs in contaminated soils through some physicochemical 75 76 processes such as sedimentation, complexation, and electrostatic adsorption (Bandara et al., 2020; Deng et al., 2019; Derakhshan Nejad et al., 2018). The complexation of Ni with oxygen-containing 77 functional groups on biochar surfaces including carboxyl, ether, carbonyl, and hydroxyl, has been 78 identified as a key mechanism for Ni immobilization in soil (Alam et al., 2018; El-Naggar et al., 79 2018). Electrostatic attraction of Ni by negatively charged functional groups on the surfaces of 80 biochar is another potential mechanism for soil Ni stabilization (Ahmad et al., 2014). Increasing 81 soil pH following the application of biochar also promotes Ni adsorption reactions (Uchimiya et 82 al., 2010). However, the efficiency of biochar prepared from different feedstocks and under 83 different production conditions in stabilizing PTEs in soils can vary significantly (Dey et al., 2023). 84

Potentially toxic elements in soil can exist in different chemical fractions such as water soluble and exchangeable (WsEx), bound to carbonates (CAR), organic materials (OM), iron and manganese oxides (FeMnOx) and residual (Res) (found in minerals) (Singh et al., 1988). The

bioavailability of these forms differs, the WsEx fraction has the highest bioavailability and the Res 88 form is considered unusable by plants. The other chemical fractions of PTEs in soils could be 89 potentially accessible for plant roots depending on soil characteristics such as soil texture, soil pH 90 91 and soil organic matter content (Kamali et al., 2011; Bharti et al., 2018). The diethylene triamine penta-acetic acid (DTPA) extraction is commonly employed for assessing Ni availability in 92 calcareous soils (Lindsay and Norvell, 1978). However, it is important to acknowledge that this 93 methodology solely assesses Ni availability for plants, while the quantity of released Ni may vary 94 95 across distinct stages of plant development. Consequently, the examination of alterations in extractable Ni levels over time using the DTPA solution can prove valuable in estimating soil Ni 96 bioavailability. The PTE desorption capacities of soils are anticipated to be contingent upon factors 97 such as soil pH, cation exchange capacity, the specific nature of metal ions, and the source of the 98 metals (Kandpal et al., 2005). Furthermore, the release kinetic parameters can provide insight into 99 the mechanisms of PTEs bonding in soils and their potential risk for leaching into groundwater or 100 surface water (El-Naggar et al., 2021). Therefore, sequential extraction methods and release 101 kinetics models have been employed to assess the efficacy of amendment materials in stabilizing 102 PTEs in contaminated soils. Xiao et al. (2021) found that addition of mineral Si fertilizer to a 103 contaminated paddy soil caused a significant decrease in the Cd and Pb fractions bound to 104 carbonates and iron-manganese oxides while the forms of residual and bound to organic matter 105 increased. In another study, application of cotton residue biochar (1.5 wt. %) to a calcareous soil 106 107 with a light texture containing different levels of Cd contamination was more efficacious than corn and wheat straw biochars in decreasing the WsEx-Cd and Car-Cd forms and enhancing the Res-108 Cd form. In addition, application of cotton residue biochar decreased EDTA-extractable Cd by 109 45–52% compared to the control (Boostani et al., 2023a). 110

As both biochars and Si are economical and effective soil amendments to reduce plant PTE 111 uptake and stress in contaminated soils, their potential synergistic effect on the immobilization of 112 PTEs in soils should be further investigated. Currently, no previous studies have examined the 113 combined application effects of Si and biochars on the chemical fractions and release kinetics of 114 Ni in calcareous soils. The primary objective of the present study was to elucidate the interaction 115 of biochars and Si levels, to alleviate soil Ni bioavailability and its corresponding accumulation in 116 corn (Zea Mays L. 604) plant. Additionally, the study sought to elucidate the underlying soil 117 chemical mechanisms that are likely to be responsible for such effects. 118

119 2 Materials and methods

120 2.1 Soil sampling, characterization and Ni treatment

121 A composite soil sample from the surface layer (0-30 cm) was collected with an auger at the research farm of the College of Agriculture and Natural Resources in Darab, southern Iran (28 122 $^{\circ}$ 45' 0.99" N 54° 26' 52.14" E, Elevation 1105 m). The soil sample was air-dried, sieved 123 through a 2 mm mesh, and then physicochemical properties were determined. Soil sand, silt and 124 clay content were determined by sieving and the hydrometer method (Gee and Bauder, 1986). Soil 125 pH and EC were determined using a saturated paste (Rhoades, 1996), while organic matter was 126 determined using Walkley-Black procedure (Nelson and Sommers, 1996). Calcium carbonate 127 equivalent (CCE) was determined by acid neutralization (Loeppert and Suarez, 1996), while cation 128 exchange capacity was determined using 1M ammonium acetate (Merck, 99%) method (Sumner 129 and Miller, 1996). Available Ni was determined using DPTA (Merck, 99%) extraction (Lindsay 130 and Norvell, 1978). Plastic containers were filled with two kilograms of soil and then 500 mL 131 NiCl₂ (Merck, 99%) solution was mixed into to them to achieve a Ni concentration of 300 mg Ni 132

 kg^{-1} soil. The Ni-treated soil samples were then allowed to dry out at room temperature, and then

rewetted to field capacity using deionized water and allowed to dry out again. The rewetting and room temperature drying cycle was repeated three times to allow the Ni to equilibrate with the soil

136 (Boostani et al., 2023c).

137 2.2 Production of biochar and its properties

138 The sheep manure and rice husk were respectively procured from an active animal husbandry and rice mill factory situated in the Darab region, Fars province, Iran. Subsequently, 139 140 the raw materials underwent a 1-week period of air-drying, followed by electrical milling and sieving through a 2 mm mesh. A slow pyrolysis procedure (2 h at 300 °C and 500 °C) in an oxygen-141 142 limited environment was carried out to generate biochars from feedstocks (Anand et al., 2023). The generated biochars were then cooled at ambient temperature and sieved with a 0.5 mm mesh 143 to ensure consistent particle size. The chemical characteristics of the biochars were assessed using 144 the following standard methods. Biochar pH and EC was determined in a 1:10 deionized water 145 suspension (Sun et al., 2014), while CEC was determined using the method of Abdelhafez et al. 146 (2014). Biochar total C, N and H contents was determined by elemental analyzer (ThermoFinnigan 147 148 Flash EA 1112 Series, Thermoscientific, USA). Biochar moisture and ash content were determined by heating in an oven, while the O+S content was calculated by subtraction of C, N, H, ash and 149 moisture content from total biochar mass (Keiluweit et al., 2010). Biochar total Ni content was 150 determined by combustion and dissolution of the ash in 2M HCl (Merck, 37%) (Boostani et al., 151 2018a). The Ni content in the acid solution was determine using atomic absorption spectroscopy 152 (AAS) (PG 990, PG Instruments Ltd., UK). The biochar surface functional groups were assessed 153 using Fourier Transform Infrared (FTIR) spectroscopy using a Shimadzu DR-8001 instrument and 154 KBr pellet transmission method. Biochar morphology was assessed using scanning electron 155 microscopy (SEM) (TESCAN-Vega3, Czech Republic). 156

157 2.3 Greenhouse experiment

A completely randomized factorial experiment was conducted in a greenhouse 158 environment with three replications. The first factor consisted of the biochar treatments including 159 160 rice husk (RH) and sheep manure (SM) generated at 300 °C and 500 °C (Control (C) (with no biochar), SM300, SM500, RH300 and RH500), each at the rate of 3% wt. The second factor 161 included Si application levels (0 (S_0), 250 (S_1) and 500 (S_2) mg Si kg⁻¹ soil) supplied as Na₂SiO₃ 162 (Sigma Aldrich, 98%) solution. Based on the experimental design, Si levels were added to the 2 163 kg of Ni-treated soil samples and after drying the soil and mixing it, the prepared biochars were 164 added to the required amount. Immediately after that, the treated soil samples were transferred to 165 plastic pots (45 pieces each containing 2 kg soil) and to facilitate the required reactions, the 166 moisture content of the samples was kept at field capacity level for a duration of two weeks. 167 Thereafter, 6 corn seeds (Zea mays L. 604) were planted in each pot, and at the 4-leaf stage, 2 168 plants were kept in each pot until the end of cultivation. During the growth of the plant, distilled 169 water was used to maintain the soil moisture content in the pots at field capacity. After 90 days, 170 the plants were harvested at the soil interface, rinsed with distilled water to remove contamination, 171 immediately air-dried and kept for Ni determination of plant shoots. After separating the roots 172 and air drying, the soil of the pots was sifted via a 2 mm mesh, and subsequently utilized for 173 performing Ni release kinetics experiment and determining the Ni chemical fractions. 174

175 2.4 Sequential extraction procedure

The present study employed a successive extraction technique (Singh et al., 1988) to 176 fractionate soil nickel (Ni) in the following chemical forms, namely water-soluble and 177 exchangeable (WsEx), carbonate-bound (Car), organic matter-bound (OM), manganese oxide-178 bound (MnOx), amorphous iron oxide-bound (AFeOx), crystalline iron oxide-bound (CFeOx), and 179 residual (Res). The methodological specifics are provided in Table 1. 180

Chemical speciation acronym Duration of agitation (h)			Extractants	Relative density (g.cm ⁻³)
Exchangeable and soluble	WsEx	2	1 M magnesium nitrate (Merck, 98%)	1.10
Carbonate	Car	5	1 M sodium acetate (Merck, 99%) (pH=5)	1.04
Organic	OM	0.5	0.7 M sodium hypocholoride (pH=8.5)	1.00
Mn oxide	MnOx	0.5	0.1 M hydroxyl amine hydrochloride (Merck, 98%) (pH=2 by nitric acid (Merck, 65%))	1.00
Amorphous Fe oxides	AFeOx	0.5	0.25 M hydroxyl amine hydrochloride (Merck, 98%) + 0.25 M choloridric acid (Merck, 37%)	1.01
Crystalline Fe oxides	CFeOx	0.5	0.2 M ammonium oxalate (Merck, 99%) + 0.2 M oxalic acid (Merck, 99%) + 0.1 M ascorbic acid (Merck, 99.7%)	1.02

с	(1, 1, 1, 2, 3, 3, 3, 3, 3, 3, 3, 3, 3, 3, 3, 3, 3,	1000
Successive extraction	technique of Singh et al. (1988)

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Table 1

2.5 Release kinetics experiment 182

A fifty milliliters centrifuge tube was filled with 10 g of soil. After that, 20 ml of DTPA 183 solution (0.005 M DTPA (Merck, 99%) + 0.1 M tri-ethanol amine (Merck, 99%) + 0.01 M calcium 184 chloride (Merck, 97%)) (pH: 7.3) (Lindsay and Norvell, 1978) was added to the soil. The soil-185 DTPA mixtures were stirred for specific periods of time, i.e. 5, 15, 30, 60, 120, 360, 720 and 1440 186 minutes at a constant temperature (25 \pm 2 °C). After each stirring time, the soil suspension was 187 centrifuged $(2683 \times g)$ to separate the soil particles from the liquid phase. Atomic absorption 188 spectroscopy (AAS) (PG 990, PG Instruments Ltd., UK) was used to analyze the Ni concentration 189 in the liquid phase. The Ni concentration in the liquid phase versus time was plotted to obtain a Ni 190 191 release kinetic curve. A total of seven kinetic models namely order models (zero, first, second and third), parabolic diffusion, power function and simple Elovich were assessed to fit the Ni release 192 data. The best models for describing the data were selected according to the maximum value of 193 the coefficient of determination (R^2) and the minimum amount of the standard error of estimate 194 (SEE)(Nasrabadi et al., 2022). 195

196 2.6 Data analysis

The ANOVA test was utilized to assess treatments effects in the individual and combined 197 biochar and silicon treatments. Additionally, a comparison of means was conducted using the 198 MSTATC computer program, applying Duncan's test with a significance level of 5%. Figures were 199 generated using Excel 2013 software. Pearson correlation coefficients among parameters in the 200 201 dataset were determined using SPSS 12.0.

202 **3 Results and Discussion**

203 3.1 Soil characteristics

204 The soil used in the study prior to experimental treatment, exhibited a sandy loam texture and possessed alkaline properties with significant calcium carbonate content, while not being 205 classified as saline (Table 2). The quantity of soil organic matter was extremely low, a distinct 206 characteristic of soils from arid and semi-arid regions (Okolo et al., 2023) (Table 2). The relatively 207 low levels of clay and organic matter present in the soil contributed to a correspondingly low soil 208 209 cation exchange capacity (CEC) (Table 2). The soils in Iran mainly originate from calcareous 210 alluvium under xeric, ustic or aridic and mesic, thermic or hyperthermic moisture and temperature regimes, respectively. These soils have varied properties such as calcium carbonate equivalent (1-211 81%), clay content (1-75%), EC (0.4-49.0 dS m⁻¹), organic matter (0.1-21.5%) and gypsum content 212 213 (0-91%) (Ghiri et al., 2011). Furthermore, it should be noted that the concentration of available soil Ni extractable by DTPA was very low (Table 2). 214

Table 2							
Certain physicochemical attributes of the soil prior to							
cultivation.							
Sand (%)	58.0						
Silt (%)	30.0						
Clay (%)	12.0						
Soil textural class	Sandy loam						
pH _(s)	7.59						
EC ($dS m^{-1}$)	2.60						
CCE (%)	55.0						
OM (%)	0.50						
CEC (cmol ₍₊₎ kg ⁻¹)	11.7						
Total Ni (mg kg ⁻¹)	28						
Ni-DTPA (mg kg ⁻¹)	0.39						
Notes: EC, electrical conduc	ctivity; OM, organic matter;						

CCE, calcium carbonate equivalent; CEC, cation exchange capacity.

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216 3.2 Chemical characteristics of the biochars

As the pyrolysis temperature rose from 300 °C to 500 °C, the SM biochars demonstrated 217 elevated pH and EC values, with the highest levels observed at the highest temperature (Table 3). 218 The elevated levels of alkali salts, which are reflected in the high ash content (Table 3), are the 219 contributing factor behind this observation in the SM biochars in comparison to the RH biochars. 220 Plant-based biochars commonly exhibit reduced levels of dissolved solids in comparison to 221 animal-based biochars (Sun et al., 2014). The SM300 biochar possessed the highest CEC value of 222 19.70 cmol+ kg⁻¹. The observed phenomenon may be attributed to the diminution of surface 223 functional groups, namely carboxyl and phenol, at elevated pyrolysis temperatures. These groups 224 are predominantly responsible for facilitating the cation exchange capacity (CEC) of biochars 225 (Tomczyk et al., 2020). As the pyrolysis temperature increased, there was an observed increase in 226 227 the C content of the biochars, and a corresponding decrease in the content of hydrogen, oxygen, and nitrogen (Table 3). The observed increase in the concentration of C as pyrolysis temperature 228 rises is consistent with a concomitant rise in the degree of carbonization. The observed reduction 229 in the levels of H and O might be attributed to the occurrence of dehydration reactions. 230 decomposition of oxygenated bonds, and the liberation of low molecular weight byproducts rich 231

in H and O, as recently noted by Zhao et al. (2017). Nitrogen compound volatilization explains the 232 diminished N content of the biochars at elevated pyrolysis temperatures. The ratios of H:C and 233 O:C are significant indicators of the aromaticity and polarity of biochars; the lower the ratios the 234 235 more condensed aromatic C the biochar contains (Chatterjee et al., 2020). The results shown in Table 3 indicated that the H:C and O:C mole ratios showed a gradual decrease as the pyrolysis 236 temperature was increased, which can be interpreted as a sign of improved carbonization of the 237 biochars (Zhao et al., 2017). The Ni content in the biochars derived from rice husk was below 238 detection, whereas a limited quantity of Ni was detected in the biochars produced from sheep 239 manure (Table 3). 240

Table	3
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Some physical and chemical properties of the biochars

Joine physical and chemical properties of the blochars.						
	SM300	SM500	RH300	RH500		
pH (1:20)	9.96	11.0	9.0	10.3		
EC (1:20) (dS m ⁻¹)	3.94	4.28	0.84	1.17		
$CEC (cmol_+ kg^{-1})$	19.70	18.94	18.94	15.33		
C (%)	25.4	31.8	45.0	50.0		
H (%)	1.85	0.8	2.28	1.06		
N (%)	2.10	1.57	1.30	1.10		
Ni (mg kg ⁻¹)	3.0	15.4	Nd	Nd		
Moisture content (%)	1.91	1.82	2.65	2.37		
Ash content (%)	53.8	60.0	34.2	44.8		
H:C mole ratio	0.87	0.30	0.60	0.25		
O+S:C mole ratio	0.44	0.09	0.24	0.01		

Notes: SM300, sheep manure biochar generated at 300 °C; SM500, sheep manure biochar generated at 500 °C; RH300, rice husk biochar produced at 300 °C; RH500, rice husk biochar produced at 500 °C; CEC, cation exchange capacity; EC, electrical conductivity; Nd, non-detectable.

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242 3.3 FTIR and SEM of the biochars

243 The FTIR spectra of the SM and RH biochars are shown in Figure 1. The SM and RH biochars produced at 300 °C contained a higher content of carboxyl groups (1700 cm⁻¹) (Keiluweit 244 et al., 2010) than the biochars produced at 500 °C, which is in agreement with the O:C values of 245 the biochars (Table 3). All of the biochars contained absorption bands associated with lignin (1430 246 cm⁻¹) and cellulose (1030 -1160 cm⁻¹) (Keiluweit et al., 2010). The SM biochar contained more 247 calcite than the RH biochar as indicated by the greater intensity of calcite characteristic peaks at 248 1432, 875, and 711cm⁻¹ (Myszka et al., 2019) in the SM biochars (Fig. 1a). There was also 249 evidence of the presence of Ca oxalate in the SM biochars, indicated by the characteristic peaks at 250 1618, 780 and 518 cm⁻¹ (Maruyama et al., 2023). All the biochars contained silica as indicated by 251 the intense silica absorption peaks at 1100, 800 and 470 cm⁻¹ (Zemnukhova et al., 2015). 252

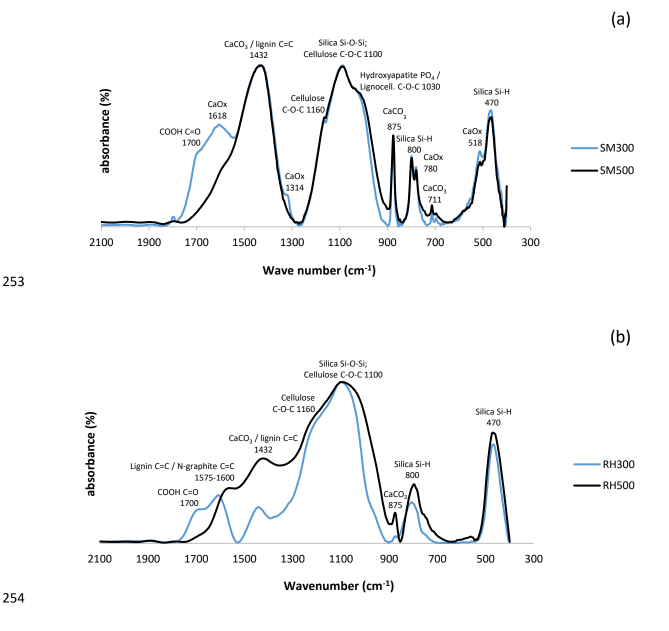


Fig. 1. FTIR of the biochars in the wave number range of 400-2000 cm⁻¹. Notes: SM300, sheep manure biochar produced at 300 °C; SM500, sheep manure biochar produced at 500 °C; RH300, rice husk biochar produced at 300 °C; RH500, rice husk biochar produced at 500 °C.

The SEM images of the SM and RH biochars are shown in Figure 2. The morphology of the biochars became more rigid and porous at higher temperatures, as evidenced by the cell wall shrinkage attributed to devolatilization of organic tissues (Claoston et al., 2014).

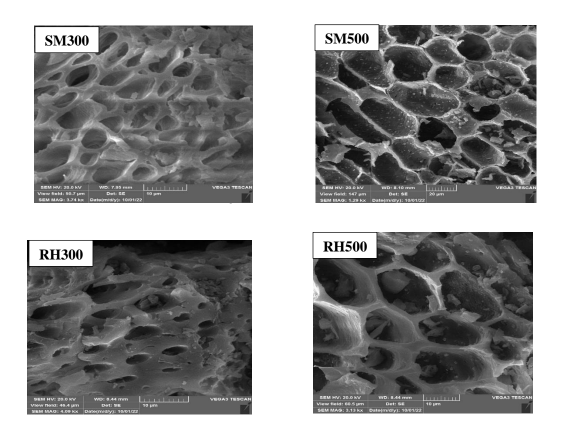
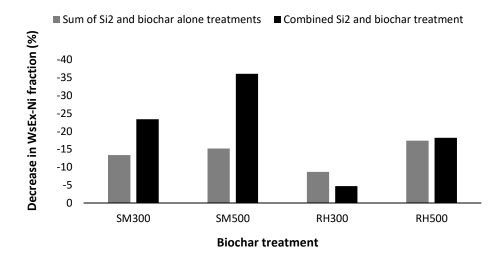


Fig. 2. SEM of the biochars. Notes: SM300, sheep manure biochar produced at 300 °C; SM500, sheep manure biochar produced at 500 °C; RH300, rice husk biochar produced at 300 °C; RH500, rice husk biochar produced at 500 °C.
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268 3.4 Soil Ni chemical fractions after the addition of silicon and biochars

The main effects of treatments (biochars and Si levels) and their interactions had a 269 statistically significant effect (P<0.01) on all the soil Ni chemical fractions, except for the Ni-Car 270 fraction, where only the main effects were significant. The soil Ni concentration in the WsEx 271 fraction was significantly reduced by the application of Si rates from S_0 to S_2 by 14.8% (Table 4). 272 Among the biochar treatments, the greatest decrease in WsEx-Ni fraction compared to the control 273 was due to SM500 by 17%, while the RH300 treatment had no significant effect (Table 4). The 274 interaction effect of treatments indicated that the lowest WsEx-Ni concentration was due to the 275 combined treatment of SM500+S₂ (4.04 mg Ni kg⁻¹ soil) (Table 4). The combined treatment of S₂ 276 and SM biochars had strong synergistic effect on reducing WsEx-Ni fraction (23-36% reduction) 277 compared to the sum of the treatments alone (13-15% reduction) (Fig. 3). Whereas this synergistic 278 effect of the combined treatments was not evident for the RH biochars (Fig. 3). There was a 279 280 negative correlation between soil WsEx fraction and soil pH (r = -0.66, p < 0.01) indicating that the reduction in WsEx fraction was strongly linked to the increase in soil pH due to the 281 amendments. Previous studies have also shown that application of biochars and silicates result in 282 increases in soil pH, thus reducing the bioavailability of PTEs and their conveyance to plant roots 283 (Shen et al., 2020; Ma et al., 2021). Among the applied biochars, the maximum pH and ash content 284 (Table 3) and calcite (lime) content (Fig. 1) were attributed to the SM500 biochar. Therefore, the 285 combined SM500+S₂ was most effective at reducing WsEx-Ni fraction, likely due to the higher 286

alkalinity and ash content of SM500 promoting Ni precipitation and adsorption (Sachdeva et al.,
2023).



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Fig. 3. Comparison of the effect of sum of the Si₂ and biochar alone treatment versus the combined
Si₂ and biochar treatments on the % reduction of the WsEx-Ni fraction. Notes: SM300, sheep manure
biochar produced at 300 °C; SM500, sheep manure biochar produced at 500 °C; RH300, rice husk biochar produced
at 300 °C; RH500, rice husk biochar produced at 500 °C.

The reduced effectiveness of biochars produced at 300 °C, as compared to those produced 294 295 at 500 °C, in decreasing soil Ni-WsEx content could probably be attributed to the lower rates of 296 microbial oxidation and mineralization of RH500 and SM500, which is indicated by their higher environmental stability (as reflected by lower H/C mole ratio values) (Table 3). Consequently, 297 298 biochar produced at 500 °C may not provide sufficient acidic carboxyl functional groups to the soil to stimulate SOM decomposition, leading to a greater increase in soil pH (Sun et al., 2023). 299 According to Zhu et al. (2015), the addition of wine lees-based biochar (a material from a wine 300 processing factory) to a heavy metal-contaminated soil (at rates of 0.5% and 1% w/w) resulted in 301 an increase in soil pH and a decrease in the soil Ni content in the WsEx fraction. Furthermore, the 302 increase in soil pH due to the increase in Si levels may lead to the precipitation of Ni in the forms 303 304 of Ni silicate and hydroxide. Due to the high solubility of Na metasilicate, the hydrolysis of silicate anion in the soil solution is intensified, leading to a high concentration of OH⁻ and a subsequent 305 306 increase in soil pH (Ma et al., 2021).

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Mean 6.07 A 5.60 B 5.17 C 9.40 A 8.76 B 7.82 C
5.60 B 5.17 C 9.40 A 8.76 B
5.60 B 5.17 C 9.40 A 8.76 B
5.17 C 9.40 A 8.76 B
9.40 A 8.76 B
8.76 B
8.76 B
8.76 B
7.82 C
7.15 A
6.48 B
5.89 C
11.21 C
11.96 B
12.74 A
80.67 C
83.03 B
85.88 A
200.3 A
199.5 A
199 A

Effects of biochars and silicon levels on the soil Ni chemical fractions (mg kg⁻¹) and Ni mobility factor (%) after corn cultivation.

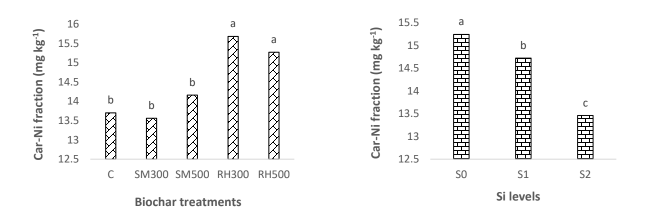
Table 4

Notes: C, control; SM300, sheep manure biochar produced at 300 °C; SM500, sheep manure biochar produced at 500 °C; RH300, rice husk biochar produced at 300 °C; RH500, rice husk biochar produced at 500 °C; S₀, without Si addition; S₁, application of 250 mg Si kg⁻¹ soil; S₂, application of 500 mg Si kg⁻¹ soil. WsEx, water soluble and exchangeable fraction; OM, organic fraction; MnOx, bound to manganese oxides; AFeOx, bound to amorphous iron oxides; CFeOx, bound to crystalline iron oxides; Res, residual fraction; MF, mobility factor. *Numbers followed by same letters in each column and rows, in each section, are not significantly (P<0.05) different

Application of Si rates from S₀ to S₂ significantly decreased the soil Ni content in the Car 314 fraction by 11.70% (from 15.24 mg Ni kg⁻¹ soil to 13.46 mg Ni kg⁻¹ soil) (Figure 4). The SM 315 biochars had no significant effect on the Car-Ni fraction whereas addition of RH biochars led to a 316 significant increase in this fraction (Figure 4). Ippolito et al. (2017) found that addition of two 317 biochars (pine [Pinus contorta] and tamarisk [Tamarix spp.]) to a mine contaminated soil caused 318 a significant increase in the soil Cd content bound to carbonates. They concluded that the reduction 319 320 in Cd bioavailability may have been due to the ability of biochar to raise soil pH levels and induce the precipitation of $CdCO_3$. Similarly, Yuan et al. (2011) proposed that the decrease in PTEs 321 bioavailability in soil might have been caused by the creation of metal-carbonate species and 322 carbonate-surface functional group reactions, which could function as a mechanism for 323 sequestration. The decrease in the concentration of Ni in the carbonate form with an increase in 324

the Si levels could potentially be explained by the competition between silicate (SiO₄⁻⁴) and carbonate ions for binding with Ni⁺² ions in the soil solution (Sparks et al., 2022).

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Fig. 4. Effects of (a) biochars and (b) silicon levels on the soil Ni concentration (mg kg⁻¹) in the carbonate-bound fraction after corn cultivation. Notes: C, control; SM300, sheep manure biochar produced at 300 °C; SM500, sheep manure biochar produced at 500 °C; RH300, rice husk biochar produced at 300 °C; RH500, rice husk biochar produced at 500 °C; S₀, without Si addition; S₁, application of 250 mg Si kg⁻¹ soil; S₂, application of 500 mg Si kg⁻¹ soil. * Numbers followed by same letters in each section, are not significantly (P<0.05) different.

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The biochars produced at 300 °C had no significant effect on the OM-Ni fraction compared 335 to control, while the biochars generated at 500 °C significantly decreased it (Table 4). The greatest 336 OM-Ni reduction (18.6%) was due to SM500. Lu et al. (2017) explored how the application of 337 338 bamboo and rice straw biochars with varying mesh sizes (0.25 and 1 mm) and at three different rates (0, 1, and 5% w/w) affected the distribution of Cd in a contaminated sandy loam soil, using 339 the BCR (Bureau Communautaire de Référence) sequential extraction method. In contrast to the 340 341 present study, they reported that the biochars increased the concentration of the Cd-OM fraction as affected, and that this was closely related to the increase in Cd immobilization. In another study, 342 the application of sheep manure biochar produced at 500 °C at the rate of 3% (w/w) to a Cd-343 contaminated calcareous soil resulted in a significant increase in the OM-Cd fraction, whereas the 344 addition of other biochar treatments (wheat straw, corn straw, rice husk, licorice root pulp) caused 345 a significant decrease in the concentration of Cd in the OM form when compared to the control 346 347 soil (Boostani et al., 2018b). In the study conducted by Boostani et al. (2018), the reduction in OM-Cd fraction as affected by application of rice husk biochar is in line with our results, however; 348 the increase in soil OM-Cd content with addition of sheep manure biochar is in conflict with the 349 result of the present study. According to the above-mentioned points, it seems that, in addition to 350 the characteristics of biochar and the level of its application (Lu et al., 2017), soil characteristics 351 (calcium carbonate percentage, soil texture, etc.) and the type of heavy metal can also have a 352 substantial role in the binding of PTEs to soil organic matter. By increasing the Si rates from S_0 to 353 S₂, the OM-Ni fraction was reduced by 16.8% (Table 4). It has been shown that the application of 354 Si to cultivated soils resulted in a reduction of soil organic matter content. This implies that Si 355

facilitates the decomposition and accessibility of organic matter to plants (Ma et al., 2021). The interaction effects of biochars and Si levels showed that the lowest OM-Ni concentration was due to the combined treatment of SM500+S₂ (7.12 mg Ni kg⁻¹ soil), which was equal to a 26.7% decrease compared to the combined treatment of C+S₀ (9.72 mg Ni kg⁻¹ soil) (Table 4).

360 All the biochar treatments caused a significant decrease in MnOx-Ni fraction compared to control, with the greatest reduction was attributed to the SM300 by 52.6% (Table 4). The lower 361 temperature biochars were more effective than the higher temperature biochars in decreasing the 362 363 MnOx-Ni fraction (Table 4). In agreement with the present study, Boostani et al. (2023c) observed that biochars produced from cow manure, municipal solid waste and licorice root pulp at lower 364 pyrolysis temperature (300 °C) decreased the soil Ni content in the MnOx fraction to a greater 365 366 extent than those prepared at higher temperature (600 °C). Hydrophobicity of biochar is decreased with increasing the pyrolysis temperature (Kameyama et al., 2019). At the same soil water content, 367 the water content of soil pores treated with biochars produced at low pyrolysis temperature is 368 higher due to lower absorption of water by the biochars. Therefore, in soils with high soil pore 369 water content and low oxygen conditions, the concentration of MnOx is decreased due to chemical 370 while concomitantly, the exchangeable and water-soluble Mn concentration are 371 reduction. 372 increased (Sparrow and Uren, 2014). Furthermore, increasing the Si concentration from S₀ to S₂ significantly decreased MnOx-Ni by 17.6% (Table 4). The interaction effect of treatments showed 373 that the highest and the lowest MnOx-Ni concentrations were found in the untreated control (11.58 374 mg Ni kg⁻¹ soil) and combined SM300+S₂ (2.98 mg Ni kg⁻¹ soil), respectively (Table 4). The 375 concentrations of soil Ni bound to AFeOx and CFeOx were significantly increased by application 376 of Si levels from S_0 to S_2 by 13.6% and 6.5%, respectively (Table 4). Belton et al. (2012) 377 demonstrated that exogenous silicon application resulted in the attachment of silicate to the surface 378 379 of iron oxide in the form of a polymer. Following the complexation of ferrosilicon, a significant number of negatively charged functional groups, including silanol, were formed. These groups 380 provided numerous adsorption sites for PTEs, ultimately reducing their bioavailability (Belton et 381 al., 2012). In general, all the biochars caused a significant increase in CFeOx-Ni fraction, and there 382 were no significant differences among the SM500, RH300 and RH500 treatments (Table 4). 383 However, the only the RH500 treatment increased the AFeOx-Ni concentration of soil compared 384 385 to control (Table 4). Among all the biochars, only the SMB300 resulted in a significant increase in the soil Ni concentration in the Res fraction compared to the control (Table 4). The application 386 of Si also did not significantly effect this form (Table 4). 387

Mailakeba and Bk (2021) studied the addition of kunai grass biochar (0.75%) to a soil with 388 different Ni contamination levels (0, 56, 100, and 180 mg Ni kg⁻¹ soil). They found that the 389 application of the grass biochar increased the Res-Ni fraction and reduced the WsEx and OM-Ni 390 fractions. In another study, Boostani et al. (2023c) demonstrated that the application of biochars 391 392 (cow manure, municipal compost and licorice root pulp each at 3%(w/w)) to a Ni-contaminated soil increased the concentrations of OM-bound and residual Ni fractions, and decreased the 393 concentrations of WsEx, Car, and Fe/Mn oxide-bound Ni fractions. Whereas, Boostani et al. 394 (2023b) found that the application of manure and compost biochars (3% w/w) to Pb-contaminated 395 soil did not significantly affect the Res-Pb fraction but did decrease the WsEx fraction. Therefore, 396 it seems that the effect of biochars on the transformation of soil PTE chemical fractions depends 397 398 on the raw materials and production conditions of the biochar, the soil application rates, type of PTEs, the degree of soil contamination with PTEs, the selection of sequential extraction procedure 399 and the soil properties (Mailakeba and Bk, 2021; Boostani et al., 2023a, b; Boostani et al., 2021). 400

In summary, the application of biochars in the present study resulted in the transformation of Ni in the soil from more bioavailable and mobile fractions (WsEx, MnOx, OM) to more stable forms (AFeOx and CFeOx). These changes were particularly evident in the WsEx fraction when SM biochar was applied in conjunction with silicon, indicating that the simultaneous use of these two substances was much more effective than applying them separately.

406 3.5 Shoot Ni concentration of *Zea mays L*. as affected by treatments

The main effects of biochars, Si rates and their interactions were statistically significant on the 407 shoot Ni concentration of the corn. Addition of Si levels from S₀ to S₂ resulted in 32% decrease in 408 shoot Ni concentration (Table 5). In addition, the shoot Ni concentration was significantly 409 decreased by application of all the biochar treatments compared to the control (with no biochar 410 addition) (Table 5). The interaction effects of treatments showed that the highest and lowest shoot 411 Ni concentration were due to the combined treatments of C+S₀ (10.4 mg Ni kg⁻¹ DM) and 412 SM500+S₂ (4.45 mg Ni kg⁻¹ DM), respectively (Table 5). The shoot Ni concentration had a 413 significant and positive correlation with the Ni-WsEx fraction (r = +0.62, P < 0.01) while there 414 were a significant and negative correlation between the soil pH (r = -0.60, P < 0.01) and Ni-415 CFeOx fraction (r = -0.50, P < 0.01). This indicates that the application of Si and biochar can 416 reduce the shoot Ni concentration by increasing soil pH and, as a result, reducing the amount of 417 Ni in the fraction of WsEx and increasing the Ni content attached to crystalline iron oxides. 418 Boostani et al. (2019a) reported the reduction of shoot Ni concentration of spinach (Spinacia 419 oleracea L.) due to the application rice husk and licorice root pulp biochars (2.5% w/w) application 420 in a Ni-contaminated calcareous soil. Additionally, they reported that the biochars produced at 350 421 °C were more effective at reducing crop Ni uptake and promoting plant growth than the biochars 422 produced at 550 °C. The most significant factors that contribute to the reduction of PTE-uptake by 423 plants in contaminated soils that have been amended with biochars include surface adsorption of 424 425 heavy metals, increased soil pH, altered redox conditions of PTEs, improved physical and biological properties of the soil, changes in the activity levels of antioxidant enzymes, and a 426 decrease in the transfer of PTEs to the plant shoots (Zeng et al., 2018; Rizwan et al., 2016). Several 427 428 studies have investigated the effect of Si application on shoot Ni concentration and other heavy metals in various plant species. Khaliq et al. (2016) observed a notable increase Ni concentration 429 430 and accumulation within the leaf, stem, and roots of cotton after Ni application. Whereas, Si application was observed to induce a significant reduction in Ni concentrations across these 431 respective plant components. In another study, Maryam et al. (2024) concluded that addition of Si 432 433 caused an increase in the growth indices of maze through reducing the Pb-shoot concentration. 434 One possible explanation for the reduction in shoot Ni concentration is that Si can compete with 435 Ni for uptake by plant roots. Silicon has a similar ionic radius to Ni, which means that it can occupy the same binding sites on root cell membranes and reduce the uptake of Ni. Additionally, Si can 436 437 induce the expression of genes that are involved in Ni transport and homeostasis, which may 438 contribute to the reduced shoot Ni concentration (Hossain et al., 2012; Liang et al., 2005).

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Shoot Ni concentration (mg Ni kg ⁻¹ DM) as affected by biochars and silicon application levels.						
	С	SM300	SM500	RH300	RH500	_
\mathbf{S}_0	10.4 a	7.35 bc	9.85 a	7.55 bc	7.65 b	8.56 A
\mathbf{S}_1	7.65 b	6.90 bc	6.60 cd	7.05 bc	7.35 bc	7.11 B
S_2	7.20 bc	5.05 ef	4.45 f	5.80 de	6.60 cd	5.82 C
Mean	8.41 A	6.43 C	6.96 BC	6.80 BC	7.20 B	

Table 5

 Shoot Ni concentration (mg Ni kg⁻¹ DM) as affected by biochars and silicon application level

Notes: C, control; SM300, sheep manure biochar generated at 300 °C; SM500, sheep manure biochar generated at 500 °C; RH300, rice husk biochar produced at 300 °C; RH500, rice husk biochar produced at 500 °C; S₀, without Si application; S₁, addition of 250 mg Si kg⁻¹ soil; S₂, addition of 500 mg Si kg⁻¹ soil. Numbers followed by same letters in each section, are not significantly (P<0.05) different.

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443 3.6 Soil Ni desorption as affected by Silicon levels and biochars

The cumulative soil Ni desorption (extracted by DTPA) as a function of time are shown in 444 Fig. 5. The release of Ni from the soil initially proceeded at a much higher rate during the first 445 446 hour, and then proceeded at a much slower rate during the next 24 hours, as illustrated by the trendline in Fig. 5. This two-stage process of releasing heavy metals from soil has also been reported 447 by other researchers (Sajadi Tabar and Jalali, 2013; Boostani et al., 2023a). It is likely that the 448 first stage of release is related to forms of Ni that are less strongly attached to soil particles, 449 including WsEx and Car, while the second stage of desorption is likely from fractions of Ni with 450 451 less bioavailability, such as FeOx and Res (Saffari et al., 2015). In general, the amount of soil Ni 452 desorption was reduced by addition of biochars and Si levels (Fig. 5). In addition, the effects of biochars produced at higher pyrolysis temperature (500 °C) on reducing the soil Ni release was 453 more than those generated at lower pyrolysis temperature (300 °C). The highest amount of soil Ni 454 release was due to the combined treatment of C+S₀ (37.84 mg Ni kg⁻¹ soil) while the lowest was 455 observed in the combined application of SM500 and S₂ (31.13 mg Ni kg⁻¹ soil) treatment. 456

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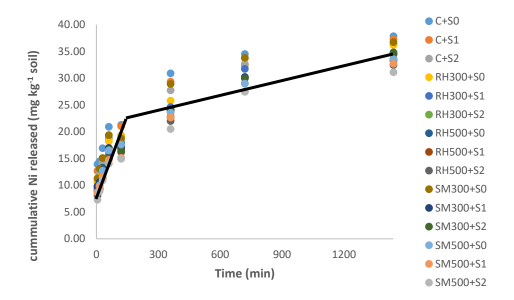


Fig. 5. Cumulative soil Ni desorption (extracted by DTPA) (mg kg⁻¹) as affected by different treatments. Notes: C, control; SM300, sheep manure biochar produced at 300°C; SM500, sheep manure biochar produced at 500°C; RH300, rice husk biochar produced at 300°C; RH500, rice husk biochar produced at 500°C; S₀, without Si addition; S₁, application of 250 mg Si kg⁻¹ soil; S₂, application of 500 mg Si kg⁻¹ soil.

464 3.7 Fitting of Ni release data to kinetics models

The soil Ni release data during 24 hours for all the biochar and Si treatments were evaluated 465 by seven different kinetic models (Table 6). The effectiveness of the various kinetic models to 466 describe the observed soil Ni desorption was analyzed by considering the coefficient of 467 determination (R^2) and standard error of estimate (SEE), so that the highest value of the R^2 and the 468 lowest value of the SEE were set as the criteria. As seen in Table 6, the order kinetic models did 469 not adequately describe soil Ni release, and with the increase in the order of the kinetic model 470 (from zero to third), the value of the R^2 decreased. This has also been found by other researchers 471 for the release of heavy metals from soil (Boostani et al., 2019b; Ghasemi-Fasaei et al., 2006). 472 Whereas, the non-order kinetic models, including power function, parabolic diffusion and simple 473 474 Elovich, acceptably described the soil Ni release of the various treatments (Table 6). Among them, the power function model was the best according to the highest value of R^2 (0.98) and the lowest 475 value of SEE (0.055). Boostani et al. (2018b) also reported that the power function was the best 476 477 kinetic model to describe soil Cd desorption from a Cd-contaminated soil treated with biochars 478 and zeolite.

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Table 6

	\mathbb{R}^2		SEE		
Kinetic models	Range	Mean	Range	Mean	
Zero order	0.79-0.87	0.80	3.36-4.67	3.67	
First order	0.69-0.75	0.75	0.22-0.29	0.25	
Second order	0.53-0.61	0.52	0.011-0.026	0.0018	
Third order	0.39-0.51	0.41	0.0013-0.0052	0.0030	
Parabolic diffusion	0.94-0.98	0.96	1.26-2.44	1.85	
Power function	0.97-0.99	0.98	0.054-0.057	0.055	
Simple Elovich	0.92-0.97	0.95	2.04-2.78	2.50	

The range of coefficients of determination (R^2) and standard error of estimate (SEE) of applied kinetic models to all the soil treatments.

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3.8 Using the parameters of power function model to investigate the effect of treatments on soilNi desorption

As the power function model $(q = at^b)$ described the soil Ni release data the best, its 488 parameters (a and b) were used to investigate the effect of biochar application and Si levels on the 489 490 release of Ni from the Ni-contaminated soil (Table 7). The main effects of biochars and Si levels and their interactions on the 'a' and 'b' parameters were significant (P < 0.01). As Dang et al. 491 (1994) reported, in this kinetics model, a decrease in parameter 'a' and an increase in parameter 'b' 492 indicates a decrease in the rate of heavy metals desorption from the soil. The main effects of 493 treatments showed that addition of all the biochar treatments caused a significant decrease in the 494 'a' parameter compared to the control while the 'b' parameter was significantly increased (Table 495 496 7). The same trend was observed for all the Si treatment rates (Table 7). Therefore, it can be concluded that the use of all the biochars and Si levels has caused a decrease in the rate of Ni 497 release from the Ni-contaminated soil. Generally, there was a greater decrease in Ni desorption in 498 biochar treatments prepared at the higher temperature (Table 7). The interaction effects indicated 499 that the most effective combined treatment in reducing the rate of Ni release from the soil was 500 $SM500+S_2$ which had the lowest value of parameter a (4.52) and the highest value of parameter 501 b'(0.264) among the treatments. 502

If it is differentiated from the power function equation $(q = at^b)$ with respect to time (t) 503 $(dq/dt = ab t^{b-1})$, when t = 1 s = 0, the ratio of dq/dt becomes 'ab'. This parameter indicates the 504 amount of heavy metal desorption in the initial time (Dalal, 1985). The *ab* parameter was affected 505 by the application of Si levels and biochars, so that this parameter was significantly decreased 506 compared to the control with addition of all the biochars (12.4%, 24.2%, 15.4% and 21.3% for the 507 508 SM300, SM500, RH300 and RH500, respectively) and Si rates (13% from S_0 to S_2), (Table 7). This finding also confirmed the effect of applied treatments in reducing the amount of Ni release. 509 510 The greatest reduction was observed in the combined treatment of $SM500+S_2$ by 33.5% compared to the control (Table 7). 511

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atter corn cult	ivation.					
	С	SM300	SM500	RH300	RH500	
		8	(mg Ni kg ⁻¹ h ⁻¹)) ^b		
\mathbf{S}_0	9.15 a	7.39 c	5.56 gh	6.49 e	5.95 f	6.91 A
\mathbf{S}_1	7.92 b	6.01 f	5.23 i	5.66 g	5.21 i	6.00 B
\mathbf{S}_2	6.90 d	5.39 hi	4.52 k	5.22 i	4.84 j	5.38 C
Mean	7.99 A	6.27 B	5.11 E	5.80 C	5.34 D	
			b (mg Ni kg ⁻¹) ⁻¹			
\mathbf{S}_0	0.196 i	0.222 g	0.247 d	0.238 e	0.237 e	0.228 C
\mathbf{S}_1	0.212 h	0.238 e	0.246 d	0.250 cd	0.254 bc	0.240 B
\mathbf{S}_2	0.230 f	0.254 bc	0.264 a	0.256 b	0.262 a	0.253 A
Mean	0.212 E	0.238 D	0.252 A	0.248 B	0.251 AB	
			ab			
\mathbf{S}_0	1.79 a	1.68 c	1.37 h	1.54 e	1.41 g	1.55 A
\mathbf{S}_1	1.69 b	1.43 f	1.29 k	1.41 fg	1.32 j	1.42 B
\mathbf{S}_2	1.59 d	1.37 h	1.19 m	1.34 i	1.271	1.35 C
Mean	1.69 A	1.48 B	1.28 E	1.43 C	1.33 D	
T G	1 (1) (2000 1		1 1 20	0°0 01 (500 1		

Table 7
The coefficients of power function model as affected by biochars and silicon levels in a Ni-polluted calcareous soil
after corn cultivation.

Notes: C, control; SM300, sheep manure biochar produced at 300 °C; SM500, sheep manure biochar produced at 500 °C; RH300, rice husk biochar produced at 300 °C; RH500, rice husk biochar produced at 500 °C; S₀, without Si addition; S₁, application of 250 mg Si kg⁻¹ soil; S₂, application of 500 mg Si kg⁻¹ soil.

* Numbers followed by same letters in each column and rows, in each section, are not significantly (P<0.05) different

517 The correlation between the parameters of the fitted power function model with soil Nichemical fractions, shoot Ni content and soil pH are shown in Table 8. The 'a' and 'ab' parameters 518 had a positive correlation with the soil WsEx, OM and MnOx Ni fractions, while there was a 519 negative correlation among the 'a' and 'ab' parameters the AFeOx and CFeOx Ni fractions. This 520 trend was inverse for the 'b' parameter of the power function model. These correlations verified 521 that the application of silicon and biochar to the Ni-contaminated calcareous soil led to a decrease 522 523 in the rate and amount of Ni release from the soil by reducing the Ni concentration in chemical forms with higher bioavailability including WsEx, OM and MnOx. Furthermore, the 'a' and 'ab' 524 parameters were negatively correlated with soil pH. Whereas there were positive correlations 525 526 between these parameters and shoot Ni concentration (Table 8). These findings once again 527 confirmed that the increase in soil pH due to the application of silicon and biochar can cause a decrease in the bioavailability of soil Ni and, as a result, a decrease in the concentration of Ni in 528 529 aerial parts of the plant.

Table 8

The correlation coefficients (r) between the power function model parameters (a, b, ab) and soil Ni chemical fractions, shoot Ni concentration and soil pH.

	WsEx	Cor	ОМ	MnOx	AFeOx	CFeOx	Res	Shoot Ni	Soil
	WSEX	Car	OM	MIIOX	AFEOX	CreOx	Res	Concentration	pН
а	0.63**	0.02 ^{ns}	0.70^{**}	0.53^{**}	-0.44**	-0.80**	0.27 ^{ns}	0.62^{**}	-0.52**
b	-0.59**	0.03 ^{ns}	-0.68**	-0.54**	0.46^{**}	0.83**	-0.28 ^{ns}	-0.63**	0.51^{**}
ab	0.68^{**}	0.04 ^{ns}	0.74^{**}	0.46^{**}	-0.46**	-0.80**	0.29 ^{ns}	0.06^{**}	-0.51**

Notes: WsEx, water soluble and exchangeable fraction; OM, organic fraction; MnOx, bound to manganese oxides; AFeOx, bound to amorphous iron oxides; CFeOx, bound to crystalline iron oxides; Res, residual fraction.

** and ^{ns} indicate significance at the 0.01 probability level and non-significant, respectively.

531 4 Conclusions

The application of biochars and Si in the present study resulted in the transformation of Ni 532 in the soil from more bioavailable and mobile fractions (WsEx, MnOx, OM) to more stable forms 533 (AFeOx and CFeOx). These changes were particularly evident in the WsEx fraction when SM 534 biochars were applied in conjunction with silicon, indicating a strong synergistic effect related to 535 536 soil pH increase. Application of all biochars and Si reduced DPTA-extractable Ni release from the soil, which was most strongly associated with the increase in CFeOx fraction. Application of all 537 biochars and Si decreased corn Ni uptake, with the combined SM500+S₂ being the most effective. 538 539 The decrease in corn uptake was correlated with the decrease in the WsEx-Ni fraction and increase in CFeOx fraction. SM500 was likely the most effective biochar due to its higher alkalinity and 540 ash content, and lower acidic functional group content which enhanced Ni sorption reactions with 541 Si. Future research is needed to better understand the mechanisms underlying the interaction 542 effects of Si and biochar application on the distribution of soil Ni chemical forms and to optimize 543 Si application strategies for sustainable Ni management in agricultural and natural ecosystems. 544

Authors' Contributions H.R.B. Conceptualization, Formal analysis, Methodology, Investigation,
Validation A.G.H. Writing - Review & Editing M.N. Project administration, Visualization E.B.
Review & Editing E.F. Laboratory analyses.

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