

1 **Cr(VI) sorption/desorption on untreated and mussel shell-
2 treated soil materials: fractionation and effects of pH and
3 chromium concentration**

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12

13 **Abstract**

14 We used batch-type experiments to study Cr(VI) sorption/desorption on granitic material,
15 forest soil, pyritic material, mussel shell, and on forest soil and granitic material amended
16 with 12 t ha⁻¹ (1.2 kg m⁻²) shell, considering the effects of varying Cr(VI) concentration and
17 pH. Sequential extractions were carried out to fractionate adsorbed Cr(VI) and to determine
18 the stability of Cr(VI) retention. The pyritic material had the highest Cr(VI) retention
19 capacity, whereas the granitic material showed the lowest retention potential. When high Cr
20 concentrations were added, some saturation of the adsorbent surfaces became apparent, but Cr
21 release remained low. The highest Cr retention was achieved at very acid pH value, with
22 release progressively increasing as a function of increasing pH. The amendment with 12 t ha⁻¹
23 mussel shell did not cause marked changes in Cr(VI) retention. Sorption data were
24 satisfactory adjusted to the Freundlich model. Regarding Cr(VI) fractionation, the soluble
25 fraction (weakly bound) was the dominant in mussel shell and in the un-amended and
26 amended granitic material, whereas more stable fractions dominated in the pyritic material
27 (residual fraction) and in the forest soil (oxidizable fraction). In conclusion, the pyritic
28 material presented the highest Cr(VI) retention capacity, while the retention was low and
29 weak on the granitic material; mussel shell was characterized by not marked Cr(VI) retention

1 potential, and it did not cause remarkable increase in Cr(VI) retention when used to amend the
2 granitic material or the forest soil.

3

4 **1 Introduction**

5 Mining, industrial and agricultural activities are the main sources of chromium pollution
6 affecting the environment, notably the water and soil compartments (Alves et al. 1993; Di et
7 al. 2006). Cr(III) is the chemically most stable form of chromium, whereas Cr(VI) is highly
8 toxic and more easily mobilized. **Mobilization of Cr(VI), and then risks of water pollution and**
9 **even of transfer to the food chain, are strongly related to retention processes affecting the**
10 **pollutant (Lilli et al. 2015).**

11 Different **bio-adsorbents have** been tried to remove Cr(VI) from polluted environments, as
12 was the case for some microorganisms and other natural sorbents (Schiever and Volesky
13 1995). Schmuhl et al. (2001) found high Cr(VI) **sorption** on chitosan, with best results at pH
14 5. Blázquez et al. (2009) obtained Cr(VI) **sorption** >80% on olive waste **at pH <2**, although
15 **sorption** clearly diminished when **pH increased**. Good Cr(VI) **sorption** results were achieved
16 using algae and cyanobacteria (Park et al. 2006; Gupta and Rastogi 2008 a, b), as well as
17 using waste from the coffee and tea industries (Fiol et al. 2008; Duran et al. 2011).

18 **Globally, it is necessary to increase the knowledge on Cr(VI) retention processes by sorbent**
19 **materials.** In this way, Fernández-Pazos et al. (2013) studied quantitative and kinetic aspects
20 regarding Cr(VI) **sorption**/desorption on various solid media (fine and coarse mussel shell,
21 un-amended and mussel shell-amended forest and vineyard soils, slate processing fines and
22 pyritic material). In addition to the kinetic characterization, it would be interesting to
23 elucidate complementary aspects, such as the effects on Cr(VI) retention caused by changing
24 pH, or the fractions where the retained Cr(VI) was bound, which can aid to estimate the
25 degree of stability of that retention.

26 **Therefore**, the main objectives of this work are: **a)** to determine Cr(VI) **sorption**/desorption
27 when different Cr(VI) concentrations are added to a granitic material, a forest soil, a pyritic
28 material, and fine mussel shell, as well as to the granitic material and the forest soil amended
29 with 12 t ha⁻¹ (1.2 kg m⁻²) fine mussel shell; **b)** to elucidate the influence of varying pH as
30 regards Cr(VI) **sorption** on such materials; **and, finally, c)** to determine the fractions where

1 Cr(VI) is retained in the various solid materials investigated, which affect Cr(VI) release and
2 then to overall risks of pollution.

3 **2 Materials and methods**

4 **2.1 Materials**

5 The materials used in this study are indicated in Table 1. The granitic material (GM) was
6 sampled in Santa Cristina (Ribadavia, Ourense Province, Spain), and resulted from the
7 evolution of a rocky substrate, similar to a C horizon, nowadays exposed to the atmosphere
8 after the elimination of the upper horizons, then needing organic matter and nutrients to be
9 restored, as happens with granitic mine spoils. The forest soil (FS) was an A horizon, with
10 dominance of *Eucalyptus globulus* as tree species, and it was sampled in the vicinity of the
11 aluminum industry Alcoa (San Cibrao, Lugo Province, Spain). The pyritic material (PM) was
12 from a copper mine spoil (Touro, A Coruña Province, Spain). The mussel shell (Sh) (<1 mm)
13 was from the factory Abonomar S.L (Illa de Arousa, Pontevedra Province, Spain). The two
14 mixtures (granitic material + 12 t ha⁻¹ mussel shell -GM + Sh-, and forest soil + 12 t ha⁻¹
15 mussel shell -FS + Sh-, were shaken for 48 h to achieve homogenization. The unity t ha⁻¹ is
16 widely used, but note that the dose 12 t ha⁻¹ mussel shell is equivalent to 1.2 kg m⁻². The FS,
17 PM and Sh here used were previously characterized by Fernández-Pazos et al. (2013) in a
18 study focusing on kinetics of Cr(VI) retention, as well as in evaluating the effects of adding
19 different concentrations of the pollutant to various sorbent materials. Fernández-Pazos et al.
20 (2013) found that the amendment of pyritic material with mussel shell had not positive effect
21 on Cr(VI) retention, so this combination was discarded in the present study.

22 FS, PM and GM were sampled in a zigzag manner (20 cm depth), taken 10 subsamples to
23 perform each of the composite FS, PM and GM final samples. These samples were
24 transported to the laboratory to be air dried and sieved through 2 mm. Finally, chemical
25 determinations and trials were carried out on the <2 mm fraction.

26 **2.2 Methods**

27 **2.2.1 Characterization of the solid materials used**

28 The particle-size distribution of the materials was determined by using the Robinson pipette
29 procedure. A pH-meter (model 2001, Crison, Spain) was used to measure pH in water and in

1 KCl (solid:liquid ratio 1:2.5). Total C and N were quantified by means of the elemental Tru
2 Spec CHNS auto-analyzer (LECO, USA). Available P was determined according to Olsen
3 and Sommers (1982). The exchangeable cations were displaced using NH₄Cl 1M solution,
4 then quantifying Ca, Mg and Al by atomic absorption spectroscopy, and Na and K by atomic
5 emission spectroscopy (AAnalyst 200, Perkin Elmer, USA); the effective cationic exchange
6 capacity (eCEC) was calculated as the sum of all these cations (Kamprath 1970). Total
7 concentrations of Na, K, Ca, Mg, Al, Fe, Mn, as well as As, Cd, Co, Cr, Cu, Ni and Zn, were
8 determined by means of ICP-mass (820-NS, Varian, USA), after nitric acid (65%) microwave
9 assisted digestion. Different selective solutions were used to obtain the following Al and Fe
10 fractions (Álvarez et al. 2013): total non-crystalline Al and Fe (Al_o, Fe_o), total Al and Fe
11 bound to organic matter (Al_p, Fe_p), non-crystalline inorganic Al and Fe (Al_{op}, Fe_{op}), Al bound
12 to organic matter in medium and low stability complexes (Al_{cu}), Al bound to organic matter in
13 high stability complexes (Al_{pcu}), Al bound to organic matter in medium stability complexes
14 (Al_{cula}), Al bound to organic matter in low stability complexes (Al_{la}).

15 **2.2.2 Sorption/desorption when different Cr(VI) concentrations are added**

16 Cr(VI) sorption and desorption as a function of the added concentration of the pollutant were
17 studied as per Arnesen and Krogstrand (1998).

18 The adsorbents used were the materials previously commented (Table 1). Fernández-Pazos et
19 al. (2013) found that the amendment of pyritic material with mussel shell had not positive
20 effect on Cr(VI) retention, so this combination was discarded in the present study.

21 As in Fernández-Pazos et al. (2013), 3 g of each solid sample were added with 30 mL NaNO₃
22 0.01M dissolution containing 0, 0.5, 5, 10, 25, 50 or 100 mg L⁻¹ of Cr(VI), prepared from
23 analytical grade K₂Cr₂O₇ (Panreac, Spain). The resulting suspensions were shaken for 24 h,
24 centrifuged at 4000 rpm (6167×g) for 15 min, and finally filtered using acid-washed paper. In
25 the equilibrium dissolution, pH was measured using a glass electrode (Crison, Spain),
26 dissolved organic carbon (DOC) was determined by means of UV-visible spectroscopy (UV-
27 1201, Shimadzu, Japan), and Cr(VI) using ICP-mass (Varian 800-NS, USA). All trials were
28 performed by triplicate.

29 Immediately after finalizing each batch experiment corresponding to the sorption trials, each
30 sample was added with 30 mL of NaNO₃ 0.01M solution to desorb Cr(VI), then it was shaken

1 during 24 h, **centrifuged** and **filtered** as in the **sorption** trials (Arnesen and Krogstad 1998).
2 Desorbed Cr(VI), DOC and pH were determined in all samples.

3 **2.2.3 Cr(VI) sorption/desorption as a function of pH**

4 To study **sorption**, triplicate samples (1 g each) **of the sorbent materials (Table 1)** were added
5 with 10 mL of solutions containing 5 mg L⁻¹ Cr(VI) and different concentrations of HNO₃
6 (0.0025 M, 0.0038 M, 0.005 M, 0.0075 M) or NaOH (0.0025 M, 0.0038 M, 0.005 M, 0.0075
7 M), also including NaNO₃ 0.01M as background electrolyte. Control samples were
8 constituted by each of the solid materials added with 10 mL of solutions containing NaNO₃
9 0.01M and 5 mg L⁻¹ Cr(VI), but without HNO₃ or NaOH. All samples were **shaken** for 24 h,
10 centrifuged for 15 min at 4000 rpm (**6167×g**), and filtered through acid-washed paper. The
11 resulting liquid was analyzed for pH, DOC and Cr(VI). Adsorbed Cr(VI) was calculated as
12 the difference between added Cr(VI) concentration and that remaining in the equilibrium
13 solution.

14 Desorption was studied using triplicate samples (1 g each) of the same solid materials than in
15 the **sorption** trials, that were added with 10 mL of solutions containing 100 mg L⁻¹ Cr(VI),
16 also including NaNO₃ 0.01 M as background electrolyte. All samples were **shaken** for 24 h,
17 centrifuged for 15 min at 4000 rpm (**6167×g**), and filtered through acid-washed paper,
18 discarding the liquid phase. The remaining solid phase was then subjected to a desorption
19 procedure, adding 30 mL of solutions containing NaNO₃ 0.01 M and diverse HNO₃ or NaOH
20 concentrations aiming to provide a wide pH range, then being different for the various solid
21 samples. All samples were **shaken** for 24 h, centrifuged for 15 min at 4000 rpm (**6167×g**), and
22 filtered through acid-washed paper. The resulting liquid was analyzed for pH, DOC and
23 Cr(VI). Desorbed Cr(VI) was calculated as the difference between the amount retained in the
24 **sorption** phase and that released to the equilibrium solution in the desorption phase, and it was
25 expressed as percentage of the total amount adsorbed.

26 **2.2.4 Fractionation of the Cr(VI) adsorbed at three different incubation times**

27 Samples corresponding to **the sorbent materials (Table 1)** were added with a NaNO₃ 0.01 M
28 solution containing 100 mg L⁻¹ Cr(VI) (1:10 solid:solution ratio), then **they were shaken** for
29 24 h and **filtered**. The resulting liquid was analyzed for pH, DOC and Cr(VI). Finally,
30 fractionation of the adsorbed Cr(VI) was carried out using the BCR procedure modified by
31 Rauret et al. (1999). The fractionation was performed **after** 24 h, 1 week and 1 month. The

1 resulting fractions were: acid-soluble fraction, reducible fraction, oxidizable fraction, and
2 residual fraction (all of them measured, not estimated).

3 **2.2.5 Data analysis**

4 The statistical package SPSS 19.0 (IBM, USA) was used to check data for normality. Then,
5 Pearson correlations were calculated.

6 Finally, Cr(VI) sorption data were fitted to the Freundlich model (Eq. 1). Fitting to the
7 Langmuir model was not possible due to estimation errors being too high.

8 The formulation of the Freundlich equation is as follows:

9 $q_e = K_F \cdot C_e^n$ (Eq. 1)

10 where q_e (mg kg⁻¹) is the ion sorption per unit of mass for the adsorbent, C_e (mg L⁻¹) is the
11 equilibrium concentration of the dissolved Cr, K_F (Lⁿ g⁻¹ mg⁽¹⁻ⁿ⁾) is a constant related to the
12 sorption capacity, and n (dimensionless) is a constant related to the sorption intensity.

13

14 **3 Results and discussion**

15 **3.1 Characterization of the solid materials used**

16 Table 2 shows data of some basic characteristics of the solid materials used in this study.

17 **3.2 Sorption/desorption when different Cr(VI) concentrations were added**

18 **3.2.1 Sorption**

19 Figure 1 shows that Cr(VI) sorption increased with Cr(VI) concentration in the equilibrium
20 solution, which was directly related to the increase in the Cr(VI) concentration added. The
21 same behavior was observed by Fernández-Pazos et al. (2013) using mussel shell, pyritic
22 material, forest soil and slate processing fines, by Rawajfih and Nsour (2008) using
23 *Phragmites australis* biomass, and by Vinodhini and Nilanjana (2009) using various bio-
24 adsorbent materials. In the present work, the maximum sorption corresponded to the pyritic
25 material (between 97.1 and 98.7% of the initial amount added), significantly higher ($p <$
26 0.005) than that achieved by forest soil (between 22.1 and 86%), mussel shell (between 20.9
27 and 31.2%), and the granitic material (between 19.5 and 31.2%). The mussel shell amendment

(12 t ha⁻¹, equivalent to 1.2 kg m⁻²) caused that Cr(VI) sorption reached values between 21 and 86% on forest soil, whereas Cr(VI) sorption achieved up to 50% on granitic material. In a previous work, Fernández-Pazos et al. (2013) found slightly lower Cr(VI) sorption on pyritic material and mussel shell, whereas sorption was very similar on their forest soil samples.

Significant correlations ($p < 0.005$) were found between the amounts of Cr(VI) sorbed (in mg kg⁻¹) and total Fe ($r=0.995$), Fe extracted with ammonium oxalate ($r=0.993$), non-crystalline inorganic Fe ($r=0.992$), and pH in water ($r= -0.900$), when chromium concentrations of 100 mg L⁻¹ were added to the sorbent materials. Previously, Martin and Kempton (2000) observed that Cr(VI) sorption increased as a function of Fe oxides content, whereas Mesuere and Fish (1992) and Weerasooriya and Tobschall (2000) indicated that Cr(VI) has slow to moderate affinity for Fe and Al oxy-hydroxides. These facts are in relation with the high sorption capacity of our pyritic material, characterized by having high Fe contents and very acid pH; furthermore, a step-wise regression test corroborated the importance of the total Fe content on Cr(VI) sorption, explicating 99% of the variance. In our study, the pyritic material was the one with the highest Cr(VI) sorption capacity, as well as the lowest pH (2.97), and high concentration of amorphous minerals. It is remarkable that this pyritic material had high specific surface and pH-dependent exchange capacity, developing positive charge at acid pH, while chromium was in anionic form (HCrO_4^- , CrO_4^{2-} and $\text{Cr}_2\text{O}_7^{2-}$). The granitic material showed the lowest Cr(VI) sorption, which can be in relation with its low total and non-crystalline Fe contents (Table 2). Correlations ($p < 0.005$) between adsorbed Cr(VI) and pH of the solution were positive for un-amended ($r = 0.701$) and shell-amended granitic material ($r = 0.770$), and for un-amended ($r = 0.672$) and shell-amended forest soil ($r = 0.819$), whereas correlations were negative for mussel shell by itself ($r = -0.994$) and for pyritic material ($r = -0.424$). These differences could be due to different mechanisms acting when Cr(VI) sorption takes place on the various materials: electrostatic bindings, then including the possibility of OH⁻ release and consequent pH increase when chromium anions adsorb (Arnesen and Krogstad 1998; Bower and Hatcher 1967; Gago et al. 2012), or other mechanisms not including OH⁻ release, such as Van der Waals and H bindings (Boddu et al. 2003). Furthermore, in the present study DOC values increased as a function of adsorbed Cr(VI), with significant correlations ($p < 0.005$) for granitic material by itself ($r = 0.978$) or mussel shell-treated ($r = 0.983$), forest soil by itself ($r = 0.905$) or mussel shell-treated ($r = 0.984$), mussel shell ($r = 0.978$), and pyritic material ($r = 0.973$), which can be in relation with release of organic ions when Cr(VI) sorption takes place. When added Cr(VI) concentrations

1 increased, both granitic material and forest soil showed decreasing **sorption** percentages, then
2 suggesting progressive saturation of the adsorbent surfaces (Nameni et al. 2008). Maximum
3 **sorption** was clearly higher for forest soil (86% **of the initial amount added**) than for granitic
4 material (31%), which can be due to the higher organic matter, organo-aluminum complexes,
5 and Alo and Feo contents in the forest soil. At this respect, Kantar et al. (2008) indicate that
6 some ferric organic compounds facilitate the stabilization of Cr(VI) in acid to slightly alkaline
7 soils due to the catalytic reduction of Cr(VI) to Cr(III).

8 **Sorption** data were satisfactory fitted to the Freundlich model **through non-linear regression**
9 (Table 3), as other authors found for various bio-sorbents (Cetinkaya-Donmez et al. 1999;
10 Prakasham et al. 1999). **Due to the fact that the Freunlich model considers that, theoretically,**
11 **sorption could be infinite, the fitting to this equation means** that **sorption** maximum would not
12 be easily predictable for these materials.

13 Significant correlations ($p < 0.005$) were found between the Freundlich constant K_F and Fe_T (r
14 = 1), Fe_{op} ($r = 1$), and Fe_o ($r = 0.999$), making evident the importance of amorphous minerals
15 in Cr(VI) **sorption**.

16 3.2.2 Desorption

17 Table 4 shows that the lowest Cr(VI) desorption corresponded to the pyritic material (0.4-
18 0.8%), whereas mussel shell by itself released between 17 and 26% of the amounts previously
19 adsorbed. When 12 t ha⁻¹ mussel shell were added to forest soil, Cr(VI) desorption increased
20 to between 8 and 44%, however desorption decreased to between 29 and 40% when the
21 granitic material was amended.

22 3.3 Cr(VI) **sorption/desorption as a function of pH**

23 3.3.1 **Sorption**

24 **Except that in panel a),** Figure 2 shows an overall increase in Cr(VI) **sorption** as a function of
25 decreasing pH values in the equilibrium solutions. Similarly, different authors indicated that
26 optimum pH values for Cr(VI) **sorption** are between 1 and 2.5 (Huang and Wu 1977; Boddu
27 et al. 2003; Mohanty et al. 2006; Rawajfih and Nsour 2008; Vinodhini and Nilanjana 2009;
28 Wang et al. 2009), due to a higher density of positive charges on the adsorbent surface, then
29 facilitating the binding to chromium anions that dominate at these very acid pH values
30 ($HCrO_4^-$, CrO_4^{2-} and $Cr_2O_7^{2-}$) (Boddu et al. 2003; Gupta et al. 2001; Ucun et al. 2002).

1 Rawajfih and Nsour (2008), as well as Wang et al. (2009), found that increasing pH values
2 cause competition between chromium oxyanions and OH⁻, then decreasing Cr(VI) **sorption**.
3 **The pyritic material showed the maximum Cr(VI) sorption**, concretely 50 mg kg⁻¹ (equivalent
4 to 99% **of the initial amount added**) at pH 3.3 (Figure 2a), remaining high for the whole pH
5 range. The mussel shell retained around 50% **of the initial amount of Cr(VI) added** when it
6 was treated with acid and the solution pH approached 7; however, the addition of base caused
7 that **sorption** diminished to 2.3 mg kg⁻¹ (equivalent to 4.5%) when pH increased to 10-12
8 (Figure 2b). The granitic material showed increased Cr(VI) retention at pH <3 (Figure 2c).
9 The forest soil adsorbed 45.7 mg kg⁻¹ Cr(VI) (equivalent to 91% **of initial amount added**) at
10 pH 2.97, but it decreased to 19.7 (39%) after being treated with base, reaching pH 5.7 (Figure
11 2d). Although positive charges on variable charge surfaces can explain maximum Cr(VI)
12 **sorption** when pH <3, the fact that the pyritic material showed high **sorption** even when pH
13 value was increased can be in relation with its abundant oxy-hydroxides content, some of
14 them with elevated point of zero charge (Bradl 2004), as well as in relation with Cr(VI)
15 reduction to Cr(III) in presence of S²⁻ and Fe²⁺, gave that Cr(III) may be precipitated at pH
16 values between 6 and 11 (Eary and Rai 1988; Sass and Rai 1987; Weng et al. 1994). Mussel
17 shell amendment did not cause great changes in the **sorbent** behavior of the amended
18 materials (Figure 1), however it provoked an slight pH increase, and slight lowering in Cr(VI)
19 **sorption**. But, in the case of the forest soil and the granitic material, this amendment increased
20 Cr(VI) **sorption** if compared with samples having similar pH values, which can be attributed
21 to precipitation with carbonates, as signaled by Aziz et al. (2008) studying chromium
22 retention on limestone.

23 **3.3.2 Desorption**

24 Figure 3 shows chromium desorption percentage for the various materials after being added
25 with 100 mg L⁻¹ Cr(VI). Desorption from the pyritic material was very low at pH <4 (Figure
26 3a), increasing with pH up to **51.7% (478.8 mg kg⁻¹)** at pH 11, similarly to that found by
27 Muthukrishnan and Guha (2008). Low crystallinity Fe and Al oxy-hydroxides can be
28 positively charged at pH <7 (Parfitt 1978; Richard and Bourg 1991), then retaining CrO₄²⁻
29 (Rai et al. 1989). As pH decrease, HCrO₄⁻ concentration increases, and this anion adsorbs
30 strongly both to low crystallinity Fe and Al oxy-hydroxides and to crystalline Fe and Al
31 oxides (Zachara et al. 1989). Cr(VI) desorption from mussel shell was **minimum** at pH close
32 to 5.5 (Figure 3b), clearly increasing at higher and (largely) at lower pH values. Cr(VI)

1 released from the granitic material hardly changed as a function of pH (Figure 3c), and was
2 always lower than 15%. Cr(VI) was **almost completely** retained in forest soil at pH < 7 (Figure
3d), increasing release with pH up to a maximum attained at pH 10.8. Chrysochoou et al.
4 (2010) **indicated** that the presence of organic matter (as in forest soil) or sulfides (as in pyritic
5 material) **facilitates** Cr(VI) reduction to Cr(III), which can precipitate at pH > 5.
6 Avundainayagam et al. (2001) **found** that Ca^{2+} can favor Cr(VI) retention in calcareous soils,
7 acting as cationic bridge on negatively charged surfaces, or by means of CaCrO_4 formation
8 (Perkins and Palmer 2000). **Although without statistical significance**, the mussel shell
9 amendment increased Cr(VI) desorption from the granitic material (Figure 3c), similarly to
10 that found by Yolcubal et al. (2007) in a calcareous soil. However, the shell amendment had
11 not effect on Cr(VI) release from our forest soil (Figure 3d).

12 **3.4 Fractionation of the Cr(VI) adsorbed at three different incubation times**

13 Figure 4 shows the results corresponding to the fractionation of the adsorbed Cr(VI), after 24
14 h (Figure 4a), 1 week (Figure 4b), and 1 month of incubation (Figure 4c). The soluble fraction
15 (the most labile -Gleyzes et al. 2002-, constituted by exchangeable and carbonate-bound
16 forms) was 95% of the adsorbed Cr in mussel shell, and 80% in granitic material, after 24 h of
17 incubation. The mussel shell amendment caused that the soluble fraction in the granitic
18 material increased to 95%, with parallel diminution of other more stable fractions, probably
19 due to Cr binding to carbonates present in the shell. Mussel shell and the granitic material
20 (un-amended or amended) did not show relevant modifications in the percentage of the
21 soluble fraction for more extended periods of incubation (1 week and 1 month). At 24 h of
22 incubation, the soluble fraction was 35% for forest soil, and 7% for the pyritic material. The
23 value did not suffer relevant changes with time for the latter, but in the case of forest soil it
24 decreased to 17% and to 11% when incubation time was 1 week and 1 month, respectively,
25 due to the increase of a more stable fraction (the oxidizable one, related with organic matter).
26 The mussel shell amendment did not cause remarkable changes in the content of the soluble
27 fraction of forest soil. At 24 h of incubation, the reducible fraction (Cr bound to Fe and Al
28 oxides and oxy-hydroxides) represented less than 12% in mussel shell, as well as in amended
29 and un-amended forest soil and granitic material, but more than 35% in the pyritic material,
30 which can be due to its Fe_T and amorphous Fe contents (Reddy et al. 1997; Nieto et al. 2008).
31 In fact, in the present study, **considering all values**, significant ($p < 0.005$) **partial** correlations
32 were found between Cr(VI) in the reducible fraction and Fe_{op} ($r = 0.999$), Fe_T ($r = 0.998$), Fe_o

1 (r = 0.997), and Mn_T (r = 0.964). When the incubation time increased to 1 week and 1 month,
2 the reducible fraction decreased in the pyritic material, increasing in parallel the residual
3 fraction (that corresponding to Cr incorporated to minerals). The mussel shell amendment did
4 not cause relevant changes in the reducible fraction contents. At 24 h of incubation, the
5 oxidizable fraction (Cr bound to organic matter) represented between 3 and 15% in mussel
6 shell and the pyritic and granitic materials, however it was 55% in forest soil, which had
7 higher organic matter content. When incubation time increased, Cr(VI) in the oxidizable
8 fraction also increased in the forest soil, reaching 80%, whereas that in the soluble fraction
9 (the most labile one) decreased. At 24 h of incubation, the residual fraction was quantitatively
10 the most important in the pyritic material, representing 40% of the adsorbed Cr(VI), and it
11 increased to 50% when incubation lapsed 1 week.

12

13 **4 Conclusions**

14 The pyritic material showed the highest Cr(VI) retention capacity among the solid substrates
15 studied, while the lowest corresponded to the granitic material. The forest soil presented high
16 **sorption** potential when pH was acid and the Cr(VI) concentration added was < 10 mg L⁻¹.
17 When the concentrations added were high (50-100 mg L⁻¹), certain saturation of the adsorbent
18 surfaces became apparent, although Cr(VI) release was low in the wide pH range studied.
19 Cr(VI) retention was more pronounced at very acid pH, while increasing pH values favored
20 its release. Mussel shell showed limited Cr(VI) retention capacity, and it did not cause marked
21 changes in Cr(VI) **sorption** when used as amendment on forest soil and granitic material.
22 **Sorption** curves fitted satisfactory to the Freundlich model. The soluble Cr(VI) fraction was
23 the dominant in the mussel shell and in the un-amended and amended granitic material,
24 whereas more stable fractions dominate in the pyritic material (residual fraction), and in the
25 forest soil (oxidizable fraction).

26

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3

1 Table 1. Materials investigated and abbreviations used to designate it.

Granitic material	Forest soil	Pyritic material	Mussel shell	Granitic material + 12 t ha ⁻¹ mussel shell	Forest soil + 12 t ha ⁻¹ mussel shell
GM	FS	PM	Sh	GM + Sh	FS + Sh

2
3

1 Table 2. General characteristics of the solid materials (average values for 3 replicates, with
 2 coefficients of variation always <5%). (Element_c: Concentration in the exchange complex;
 3 Element_T: Total concentration; Al_o, Fe_o: extracted with ammonium oxalate; Al_p, Fe_p:
 4 extracted with sodium piro-phosphate; Al_{cu}: extracted with copper chloride; Al_{la}: extracted
 5 with lanthanum chloride; Al_{op}: Al_o-Al_p; Al_{pcu}: Al_p-Al_{cu}; Al_{cula}: Al_{cu}-Al_{la}; Fe_{op}: Fe_o-Fe_p)

	Granitic material	Forest soil	Mussel shell	Pyritic material
Sand (%)	60.00	65.00	99.53	67.00
Silt (%)	17.00	20.00	0.34	14.00
Clay (%)	23.00	15.00	0.13	19.00
C (%)	0.11	4.22	11.43	0.26
N (%)	0.04	0.33	0.21	0.04
pH _{H2O}	5.72	5.65	9.39	2.97
pH _{KCl}	3.69	4.70	9.04	2.58
Ca _c (cmol kg ⁻¹)	0.18	4.37	24.75	0.36
Mg _c (cmol kg ⁻¹)	0.13	0.66	0.72	0.29
Na _c (cmol kg ⁻¹)	0.27	0.33	4.37	0.14
K _c (cmol kg ⁻¹)	0.31	0.60	0.38	0.24
Al _c (cmol kg ⁻¹)	1.63	1.92	0.03	2.86
e-CEC (cmol kg ⁻¹)	2.53	7.88	30.26	3.89
Al-saturation (%)	64.55	24.41	0.11	73.68
P _{Olsen} (mg kg ⁻¹)	2.56	28.80	54.17	8.80

6

1 Table 2 (continuation).

	Granitic material	Forest soil	Mussel shell	Pyritic material
Ca _T (mg kg ⁻¹)	<0.01	709	280168	603
Mg _T (mg kg ⁻¹)	355	831	981	8384
Na _T (mg kg ⁻¹)	102	515	5174	412
K _T (mg kg ⁻¹)	1434	1544	202	3186
Mn _T (mg kg ⁻¹)	24	93	34	296
Cu _T (mg kg ⁻¹)	7	16	7	773
Zn _T (mg kg ⁻¹)	18	37	8	58
Ni _T (mg kg ⁻¹)	1	11	8	5
Cd _T (mg kg ⁻¹)	<0.001	0.43	0.07	0.08
Cr _T (mg kg ⁻¹)	3	18	5	99
Co _T (mg kg ⁻¹)	0.4	1.4	1.0	3.1
As _T (mg kg ⁻¹)	3	4	1	7
Al _T (mg kg ⁻¹)	5981	19660	433	9624
Fe _T (mg kg ⁻¹)	3505	9486	3535	135157
Al _o (mg kg ⁻¹)	1425	4275	178	563
Al _p (mg kg ⁻¹)	463	4163	78.7	229
Al _{op} (mg kg ⁻¹)	963	112	99.7	335
Al _{cu} (mg kg ⁻¹)	150	868	22.9	186
Al _{pcu} (mg kg ⁻¹)	312	3295	55.8	42.7
Al _{la} (mg kg ⁻¹)	137	146	2.6	91.1
Al _{cula} (mg kg ⁻¹)	12.8	722	20.3	134
Fe _o (mg kg ⁻¹)	224	2333	171	41860
Fe _p (mg kg ⁻¹)	54.3	2246	37.7	625
Fe _{op} (mg kg ⁻¹)	170	86.9	133	41235

1 Table 3. Fitting of the various materials to the Freundlich model ($p < 0.005$). (GM: Granitic
 2 material; FS: Forest soil; Sh: Mussel shell; PM: Pyritic material).

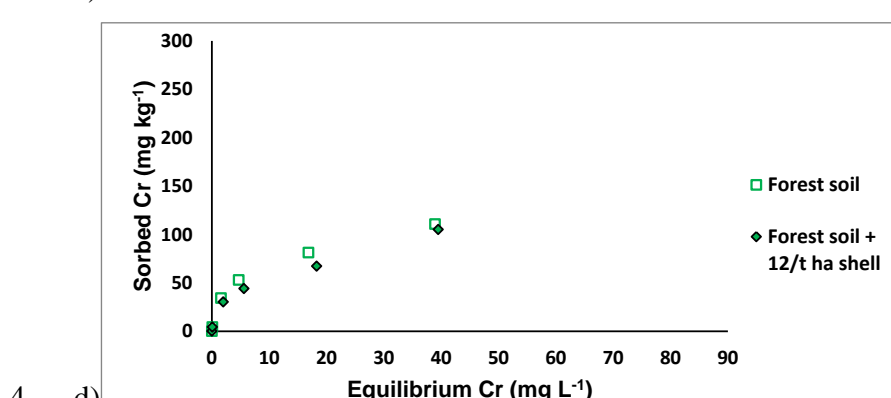
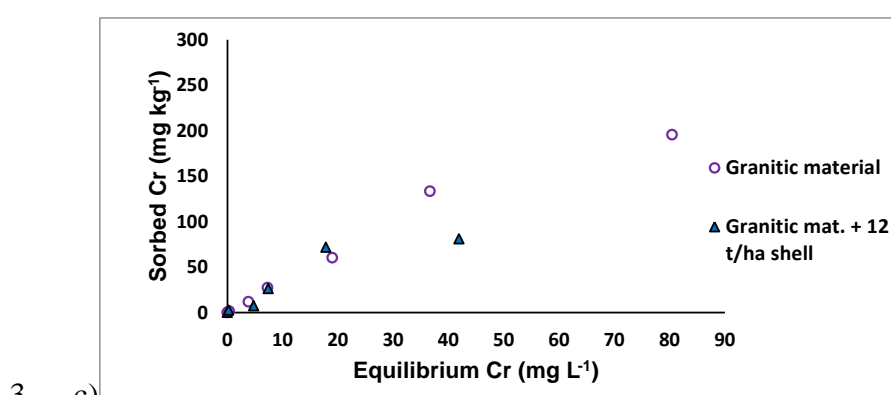
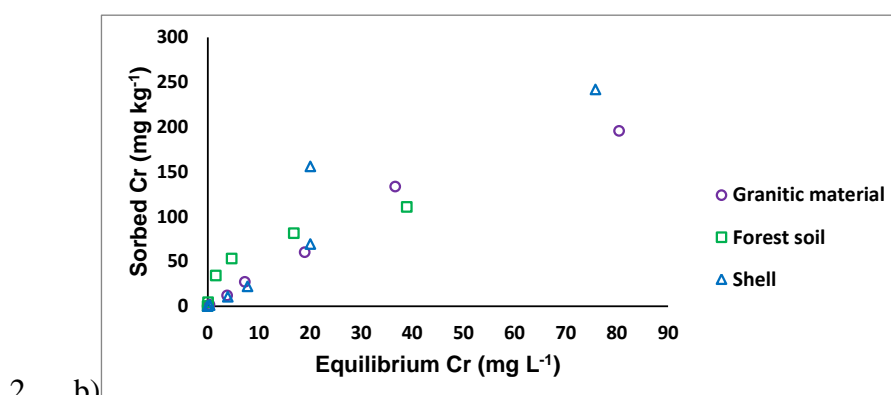
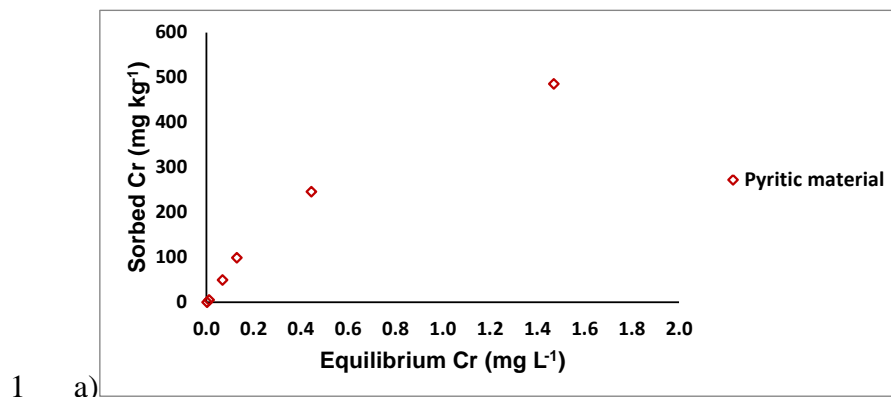
	Freundlich		
	K_F ($L^n g^{-1} mg^{(1-n)}$)	n	R^2
GM	7.5±2.5	0.75±0.08	0.980
FS	28.1±2.0	0.38±0.02	0.995
Sh	10.7±7.1	0.73±0.17	0.895
PM	381.6±11.7	0.66±0.04	0.994
GM + Sh	8.3±5.2	0.63±0.19	0.891
FS + Sh	19.9±2.1	0.45±0.03	0.993

3

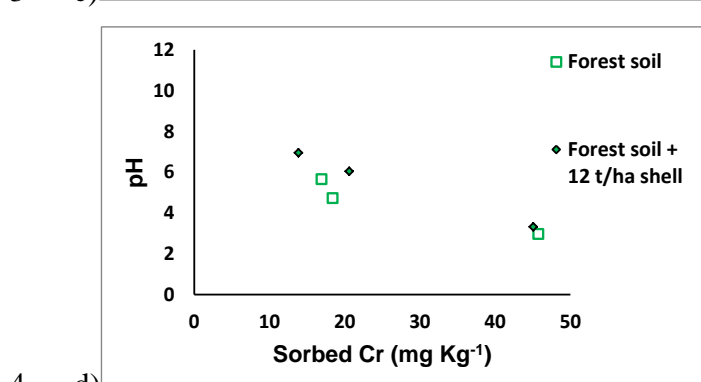
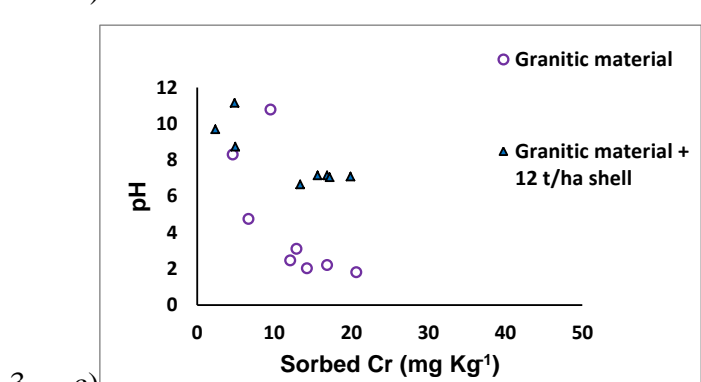
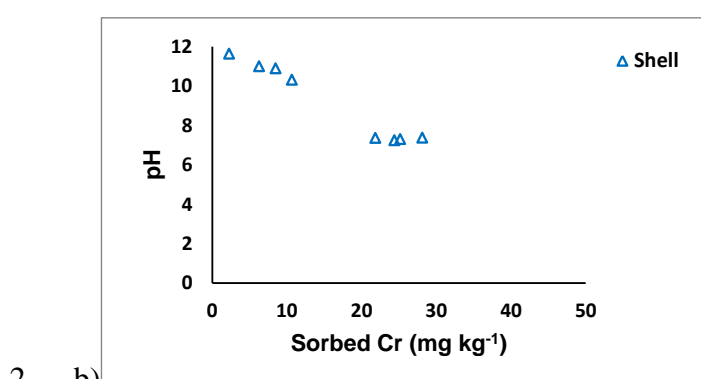
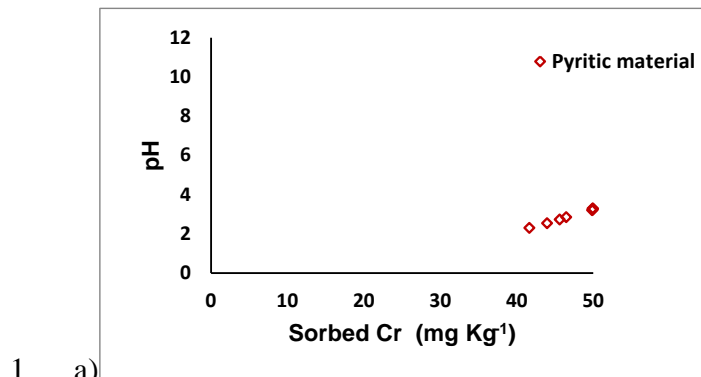
1 Table 4. Desorbed Cr (mg kg⁻¹ and % of the amount previously retained) when 0, 0.5, 5, 10,
 2 25, 50 or 100 mg L⁻¹ Cr were previously added to the various materials. (GM: Granitic
 3 material; FS: Forest soil; PM: Pyritic material; Sh: Mussel shell).

		Desorbed Cr	Added Cr (mg L ⁻¹)						
			0	0.5	5	10	25	50	100
GM	mg kg ⁻¹	0.003	0.943	10.249	16.893	37.799	68.150	128.289	
	%	0	61	92	65.5	66	55	67	
FS	mg kg ⁻¹	0.000	0.634	0.905	1.314	4.773	17.711	45.115	
	%	0	15	2.6	2.5	6	16	21	
PM	mg kg ⁻¹	0.018	0.040	0.213	0.401	1.335	4.015	4.722	
	%	0	0.8	0.4	0.4	0.5	0.8	0.5	
Sh	mg kg ⁻¹	0.038	0.178	2.301	4.565	11.784	29.580	52.747	
	%	0	17	23	26	22	20	23	
GM+12t ha ⁻¹ Sh	mg kg ⁻¹	0.000	0.787	2.943	9.093	21.312	30.976	61.636	
	%	0	31	41	35	33	40	29	
FS+12t ha ⁻¹ Sh	mg kg ⁻¹	0.000	0.626	2.276	4.560	16.198	37.978	85.178	
	%	0	15	8	10	24	36	44	

4
 5



5 Figure 1. **Sorption points** corresponding to the pyritic material (a), granitic material, forest soil
6 and mussel shell (b), granitic material with or without mussel shell (c), and forest soil with or
7 without mussel shell (d). **Mean values; triplicate samples with coefficient of variation <5%.**



5 Figure 2. Relation between pH and sorbed Cr (mg kg^{-1}) for pyritic material (a), mussel shell
6 (b), amended and un-amended granitic material (c), and amended and un-amended forest soil
7 (d), after being added with 100 mg L^{-1} Cr. Mean values; triplicate samples with coefficient of
8 variation $<5\%$.

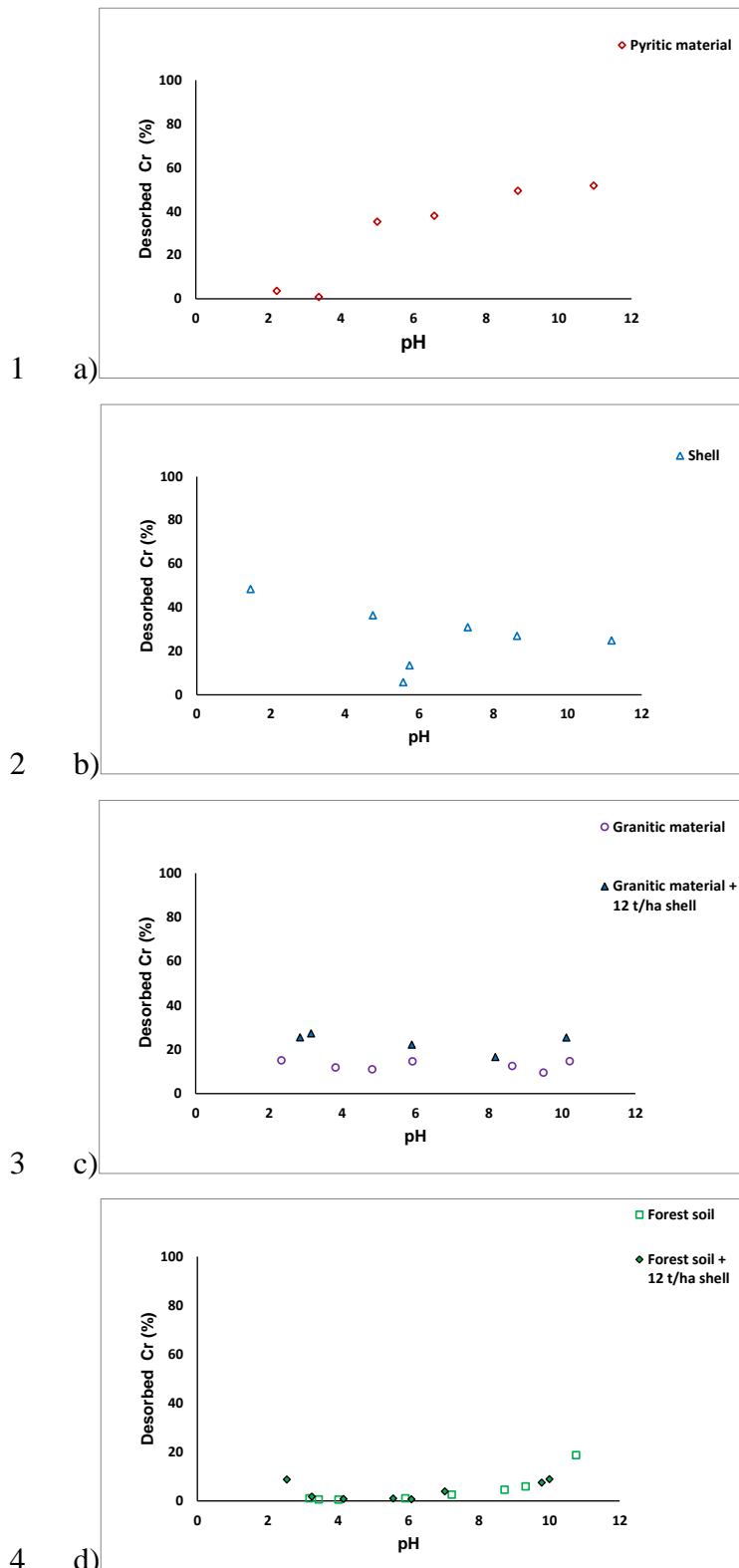
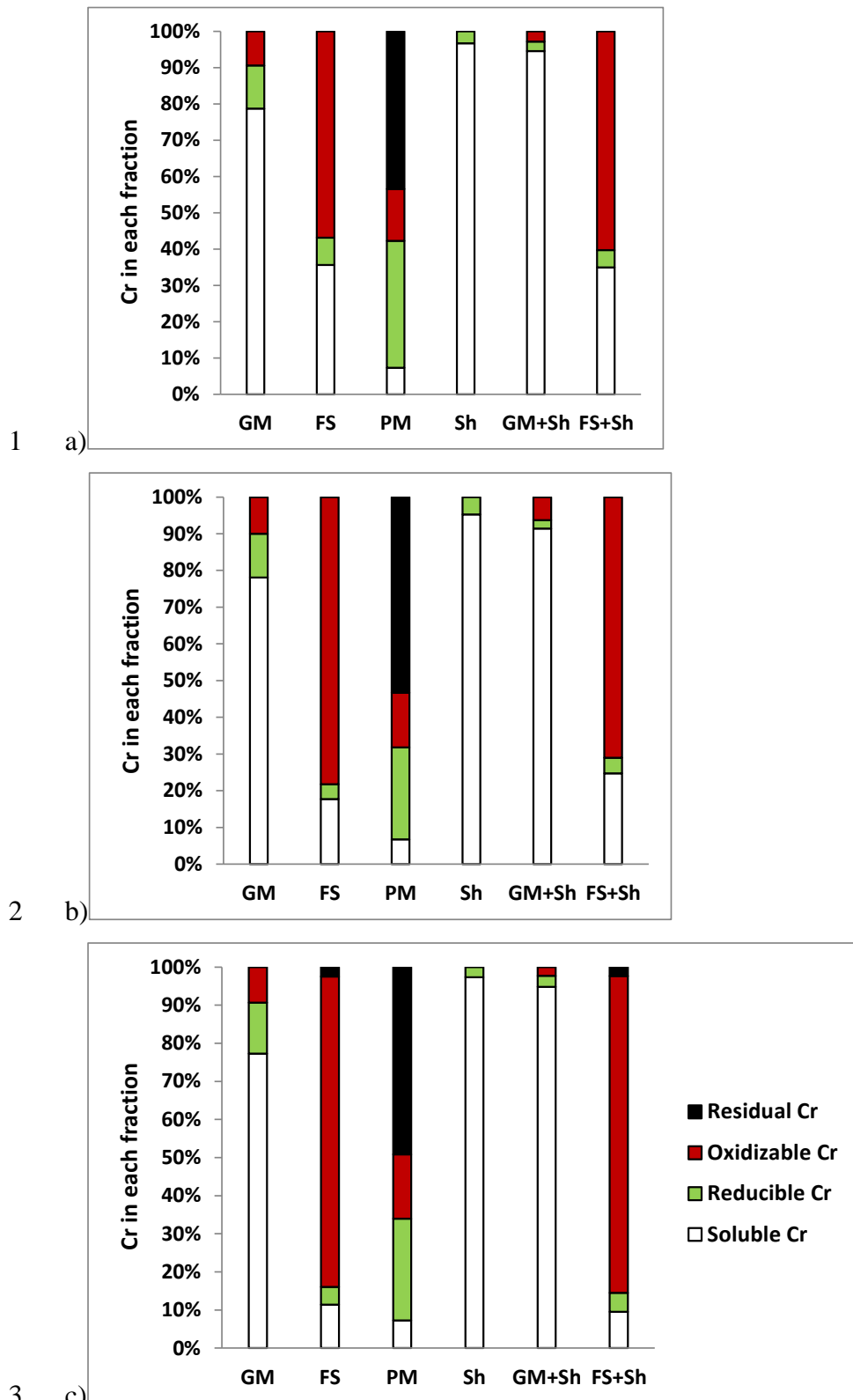


Figure 3. Relation between desorbed Cr(VI) (%) and pH for pyritic material (a), mussel shell (b), amended and un-amended granitic material (c), and amended and un-amended forest soil (d), after being added with 100 mg L^{-1} Cr. Mean values; triplicate samples with coefficient of variation <5%.



4 Figure 4. Percentages of the various fractions of chromium sorbed after the addition of 100
5 mg L^{-1} Cr at time 0, and taken 24 h (a), 1 week (b) and 1 month (c) of incubation. GM:
6 granitic material; FS: forest soil; PM: pyritic material; Sh: mussel shell. Mean values;
7 triplicate samples with coefficient of variation <5%.