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Short- and medium- term effects of three fire fighting chemicals on the properties of a burnt soil

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Abstract

The impact of three fire fighting chemicals (FFC) on 11 chemical soil properties and on soil recovery (0-2 cm depth) was evaluated 1, 30, 90 and 365 d after a prescribed fire. Five treatments were considered: unburnt soil (US) and burnt soil with 21 m⁻² of water alone (BS) or mixed with the foaming agent Auxquímica RFC-88 at 1% (BS+Fo), Firesorb at 1.5% (BS+Fi) and FR Cross ammonium polyphosphate at 20% (BS+Ap). At t=1 d, soil pH increases in the order US < BS ≈ BS+Fo, BS+Fi < BS+Ap, which was most likely due to the accumulation of ashes, the reduction of organic acids and the cations supplied by FFC. In all burnt treatments, soil pH remained significantly higher than in US up until t=90 d. SOM richness remained similar and constant until t=90 d in all plots, but, probably due to fire-triggered erosion, at t=365 d it was significantly lower in BS+Ap (C, N), BS and BS+Fo (C) than in US. Immediately after the fire, soil δ ¹⁵N decreased in all burnt soils (significatively in BS+Ap) due to the inputs of ¹⁵N depleted ashes from leguminous vegetation. Compared with US, soil δ ¹⁵N increased significantly in all burnt plots between t=90 d (30 d in BS+Ap) and t=365 d, suggesting a medium-term fire-triggered increment of N outputs (¹⁵N depleted). As is habitually the case, there was a transient post-fire increase of NH4+-N levels (significative for BS+FFC plots) that lasted for 30 (BS, BS+Fo and BS+Fi) to 90 d (BS+Ap). The high initial NH_4^+ -N levels in BS+Ap (200x that of US; 9-18x those of BS, BS+Fo and BS+Fi), and its persistence can delay the post-fire vegetation recovery due to the toxicity of NH₄⁺ to seeds and seedlings. NO₃-N levels changed significantly only in BS+Ap between t=30 and t=90 d due to the nitrification of its large NH_4^+ -N pool. Except in BS+Ap, whose soil P levels were 70-140x (t=1 d) and 10-20x (t=365 d) higher than in the other treatments, available P content in BS and BS+FFC was not significatively higher than in US. The concentrations of available cations in BS and BS+FFC were higher (not always significatively, except for K) than in US until t=90 d, likely due to ashes- and FFC-derived cations. Contrarily to divalent cations, monovalent cations (more soluble and easily leached) decreased slowly until t=90 d.

Key words: Available nutrients; burnt soil; fire fighting chemicals; flame retardants; δ ¹⁵N; prescribed fire; wildfire.

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1. Introduction

Wildfires, which frequently degrade ecosystems from their base (the soil) by increasing nutrient and soil losses through leaching and erosion (Chandler et al., 1983; Carballas et al., 1993), are one of the most widespread factors of ecosystem degradation around the world; consequently, the improvement of wildfire fighting techniques is a global concern. The addition of substances to water to increase its effectiveness as a forest fire extinguishing agent has been a common practice since the beginning of the 1930s (Giménez et al., 2004). Studies on these substances, generically called fire fighting chemicals, have been traditionally focussed on the evaluation of their effectiveness as fire extinguishers when mixed with water, usually under laboratory conditions, although there are also field experiments (see reviews by George et al., 1976, and Giménez et al., 2004). Since most fire-fighting chemicals are typically applied in environmentally sensitive areas, natural areas and areas set aside for wildlife, there is a need to determine their potential effects on terrestrial and aquatic ecosystems (Basanta et al., 2002). There is a great deal of information regarding the effects of fire-fighting chemicals on plants and animals, both aquatic and terrestrial (Larson and Duncan, 1982; Bradstock et al., 1987; Gaikowski et al., 1996; MacDonald et al., 1996, 1997; Adams and Simmons, 1999; Larson et al., 2000; Giménez et al., 2004; Hartskeerl, et al., 2004; Bell et al., 2005). Conversely, in spite of its interest, the effects of these compounds on the soil have scarcely been studied (see Giménez et al., 2004). However, since the beginning of 1980s there has been concern regarding the possible consequences of N and P enrichment due to fire fighting chemicals application in ecosystems usually poor in these nutrients (Neary and Currier, 1982). More recently, Basanta et al. (2002) studied under laboratory conditions the effects of a fire fighting chemical over some biochemical properties of two heated and two unheated soils, whereas Hopmans and Bickford (2003) studied, for the period of one year under field conditions, the impact of another fire-fighting compound on the chemical properties of two unburnt soils. The last two studies focussed on two scenarios, unburnt and burnt soils plus retardant unaffected by the high temperatures associated with wildfire, probably due to the difficulties for studying the third possible scenario: soils in the fire line that receive retardant, and which are affected by both the heating and oxidation caused by the fire. On the other hand, the ratios of stable nitrogen isotopes expressed as δ ¹⁵N values can indicate the openness of nitrogen cycles in ecosystems, and they are useful tools in evaluating the characteristics of N transformation in an ecosystem, as well as the past and current N status of forested ecosystems (Garten, 1993; Garten and Van Miegroet, 1995; Hogberg, 1991; Hogberg and Johannisson, 1993; Cook, 2001; Robinson, 2001; Koba et al. 2003). Moreover, soil isotopic signatures have been previously used to infer patterns of fire history (Schmidt and Stewart, 2003; Aranibar et al., 2003).

The aim of this paper is to evaluate, under field conditions, the impact of three fire fighting chemicals (foaming agent RFC-88 from Auxquímica, terpolymer Firesorb from Stockhausen and ammonium polyphosphate FR from Cross), applied immediately after a prescribed fire (i.e. with the fire extinguished but with the soil still warm), on 11 chemical properties of a forest soil in order to determine their effects on the post-fire recovery of the soil.

2. Material and methods

The experimental field, with UTM coordinates 29T⁰⁵182 - ⁴⁶509, is located at Alto da Pedrada (Tomiño, Galicia, NW Spain) at an altitude of 455 m a.s.l. The soil, developed over a parent material of paragneises and with a slope of 18-19%, has a vegetation cover dominated by *Ulex*, *Chamaespartium* and *Erica* 50-60 cm heigh.

Five treatments were considered: a) unburnt soil (US) as a control; b) burnt soil with 2 lm^{-2} of water (BS); c) burnt soil with 2 lm^{-2} of water plus foaming agent Auxquímica RFC-88 at 1%

(BS+Fo); d) burnt soil with 21 m⁻² of water plus Firesorb at 1.5% (BS+Fi); and e) burnt soil with 21 m⁻² of water plus FR Cross ammonium polyphosphate at 20% (BS+Ap). After a prescribed fire (i.e. with the fire extinguished but with the soil still warm), burnt soil treatments were arranged in a fully randomized design with four replications and 1 m separation around each plot (4 x 4 m), whereas the four unburnt soil replicates were established along the slope and adjacent to the burnt ones (see Fig. 1).

Soil samples were taken from the A horizon (0-2 cm depth) immediately before and after burning and 1, 3 and 12 months after the prescribed fire. Five 15 x 15 cm squares, uniformly distributed around each plot, were sampled, mixed and thoroughly homogenized after sieving at 4 mm. Before the prescribed fire, plant litter was also sampled in all plots, dried at 60°C, finely ground (< 100 μ m) and kept for analysis.

The dry matter content of fresh soils and litter was assessed by oven-drying aliquots of them at 110 °C for 5 h. Soil pH was measured with a pH-meter (Metröhm, Switzerland) in 0.1 M KCl employing a soil: solution ratio of 1:2.5.

Available Na, K, Ca, Mg and P were extracted with acetic acid 0.5 M (soil:extractant ratio 1:40). All available nutrients were measured by inductively coupled plasma optical emission spectrometry (Varian Vista Pro simultaneous ICP-OES).

The C and N contents of soils and litter, as well as their δ^{15} N values, were measured on finely ground samples (< 100 µm) with an elemental analyzer (EA) coupled on-line with an isotopic ratio mass spectrometer (Finnigan Mat, delta C, Bremen, Germany).

For the inorganic N analysis (NH_4^+ and NO_3^-) an extraction-diffusion method, at room temperature, was used. Fresh soil samples (20 g), in duplicate, were shaken with 100 ml of 2 M KCl for 1 h and the extracts were filtered through glass microfibre filters (Whatman GF/A, 125 mm diameter). In order to sequentially liberate NH_4 -N and $NO_2 + NO_3$ -N in the form of NH_3 25 ml aliquots of 2 M KCl soil extracts, placed in a 500 ml wide-mouth glass jar, were treated successively (5 d periods) with MgO and MgO plus Devarda's alloy. Gaseous NH_3 was trapped in 10 ml of 0.005 M H_2SO_4 in a Teflon bottle attached to the glass jar and measured by back titration of the excess of H_2SO_4 with 0.01 M NaOH.

The nutrient content of the fire fighting agents, measured by ICP-OES (Na, K, Ca, Mg and P) and AE-IRMS (N, δ ¹⁵N), as well as their densities are shown in Table 1.

Data of soil variables analysed at several dates after the fire were statistically analysed by two-way ANOVA (with treatment and sampling time as factors) and the Levene's test was used for verifying the equality of variances among groups. In the case of homocedasticity, significant differences among the mean groups were established at P < 0.05 using the Bonferroni test for multiple comparisons. In the case of unequal variances, the original data were subjected to Cox-Box transformations to obtain equality of variances, and significant differences among the mean groups were then established at P < 0.05 using the Bonferroni test for multiple comparisons.

3. Results

Before the prescribed fire there were no significative differences among the plots assigned to the different treatments for any of the variables studied, which shows acceptable spatial homogeneity of the experimental field.

Compared with that of the unburnt soil (pH_{KCI} 3,36), the pH_{KCI} of the burnt soil was 0.4 units higher (P < 0.05), and those of the burnt soil with fire fighting chemicals were 0.5 (BS+Fo and BS+Fi; P < 0.05) to 0.9 units (BS+Ap) higher, the pH_{KCI} in the latter being significantly higher (P < 0.05) than in US and BS (Fig. 2). In the burnt soils, with or without fire fighting chemicals, the

pH remained significantly higher than in the US for at least three months after the prescribed fire, although there was a clear decreasing tendency in BS+Ap during this period. One year after the fire the burnt soils pH decreased to values close to US; the same was true for the burnt soils treated with fire fighting chemicals.

The SOM richness in the different plots remained similar and constant (total C: 149-207 g kg⁻¹; total N: 6.7-8.7 g kg⁻¹ ds; Fig. 3) up to three months after the fire. However, one year later, the SOM content in US (total C and N) was higher than in all the other treatments, the differences with BS+Ap (C and N) and with BS and BS+Fo (C) being significant.

In the first year after the fire, δ ¹⁵N values for US rose from +1.45 to +1.75 δ , with no clear temporal tendency. However, immediately after the prescribed fire, there was a decrease of 0.2-0.5 δ ¹⁵N in BS, BS+Fo, BS+Fi and BS+Ap in comparison with US, although the difference was only significative in the case of BS+Ap (Fig. 4). One month later, a slight decrease in BS remained, but δ ¹⁵N in BS+Fo, BS+Fi and BS+Ap started to rise, especially in BS+Ap, the mean value of which did not differ from US; the same was true for BS, BS+Fo and BS+Fi at t=90 d. This tendency to rise was still evident after 365 d, and all BS treatments reached values (+2.22 to +2.57 δ) significatively higher than US.

Compared with US, immediately after burning there was an increase in NH_4^+ -N in all BS treatments, although differences were only significant in those with fire fighting chemicals. At t=1 d, NH_4^+ -N concentration increased in the order US \leq BS \leq BS+Fo, BS+Fi \leq BS+Ap, with the NH_4^+ -N content in BS+Ap being 200 times higher than that in US, and 9-18 times higher than those in BS, BS+Fo, BS+Fi (Fig. 5). The post-fire increase of NH_4^+ -N levels was transient in BS, BS+Fo and BS+Fi, their values being similar to that of US at t=30 d; conversely, in BS+Ap the NH_4^+ -N content remained significantly higher than in the other treatments for at least three months, although differences were not significant at t=365 d.

Contrary to NH_4^+ -N, NO_3^- -N concentrations did not differ significantly among treatments at t=1 d (Fig. 5). The same was true one month later, except for BS+Ap, for which NO_3^- -N content was higher (P < 0.05) than in the other treatments. Irrespective of soil treatment, at t=90 d there was a generalized increase of NO_3^- -N content and, like at t=30 d, maximum NO_3^- -N levels (P < 0.05) were found in BS+Ap. At t=365 d, NO_3^- -N concentration was similar in all treatments and comparable to those data obtained before burning.

Available P content in BS was higher than in US, but the difference was not significative (Fig. 6). The same was true for the treatments with fire fighting chemicals, except for BS+Ap which showed a huge increase in soil available P (817 mg P kg⁻¹ ds at t=1 d; 70-140 times higher than the concentration in the other treatments; Fig. 6). Despite the sustained decrease in P levels throughout the entire study period, even after a year the BS+Ap values (51.9 mg P kg⁻¹ ds) were still significatively higher than those of the other treatments (10-20 times).

The concentration of available cations in the burnt plots, with or without fire fighting chemicals, was higher than in the unburnt plots up until 90 d after the prescribed fire, although differences were not always significative, except for K (Fig. 7). Monovalent cation concentrations decreased slowly during the first three months after the burning; conversely, after a transitory decrease at t=30 d, and the divalent cation content at t=90 d increased to levels that were close to those measured immediately after burning.

4. Discussion

The significant increase of soil pH_{KCI} observed after the prescribed fire was in all likelihood due to the accumulation of ashes rich in oxides and carbonates of basic ions and the reduction of organic acids that have been reported by several authors (Chandler et al., 1983; Kutiel et al.,

1990; Carballas, 1997). The additional increment of soil pH_{KCl} in burnt soils treated with flame suppression chemicals could be due to the cations supplied by these chemicals, whose water solutions at the concentration used in fire-fighting have a neutral pH (foaming agent at 1% pH 7.2; Firesorb at 1.5% pH 7.0; ammonium polyphosphate at 20% pH 6.2). Although initially (the first 3 months of warm and dry summer) the soil pH remained significantly higher in BS than in US, one year after the fire the burnt soil pH decreases to values close to US, following a common trend in burnt soils, which is mainly due to the leaching of basic ions during the wet season (Viro, 1974; Pritchett and Fisher, 1987); the same was true for the burnt soils treated with fire fighting chemicals, as was also found by Hopmans and Bickford (2003) in one of the two unburnt soils they treated with Phos-Check.

The lack of significant differences in SOM richness among the soil treatments up until three months after the fire showed that neither the prescribed fire nor the addition of fire fighting chemicals influenced the soil C and N content in the short term; similarly, no significant differences were found by Hopmans and Bickford (2003) in SOM content of unburnt soils treated with Phos-Check. On the other hand, the SOM depletion observed one year later in all treatments with burnt soil may be the medium-term consequence of the fire-triggered erosion processes on the surface of burnt plots.

The δ^{15} N decrease in burnt soils just after the fire was without doubt due to the N added to the soil surface by the ashes from plant material (either living plants or litter), which were impossible to differentiate from the upper soil layer ashes and, consequently, jointly sampled. The natural abundance of ¹⁵N in the litter (-1.1 δ) showed that the leguminous plants (*Ulex europaeus, Chamaespartium tridentatum*) that dominated the plant community of the plots were also depleted in ¹⁵N (as is usual in plants that actively fix N₂; Shearer and Kohl, 1993) and, therefore, the contribution of ashes from litter and living plants provoked the decrease in δ^{15} N in the 0-2 cm soil layer. The increment in δ^{15} N that was evident between three months (one month in BS+Ap) and one year after the prescribed fire in all burnt plots, compared with US, clearly suggests that the fire had a medium-term influence on N balance, promoting the outputs, and which was undoubtedly due to an increase in lixiviation of nitrates or in volatilization of ammonia, which are ¹⁵N depleted (Högberg et al., 1995).

The NH_4^+ -N increase that takes place just after the fire in all burnt soil treatments is a common tendency of recently heated or burnt soils (DeBano et al., 1979; Chandler et al., 1983; Prieto-Fernández et al., 1993). As has previously been mentioned, differences in NH_4^+ -N content compared with the US were significant for all soils treated with fire fighting chemicals, but the extremely high concentration of BS+Ap was very likely due to the NH_4^+ -N supplied by the ammonium polyphosphate. Similarly, in field experiments with unburnt soils, Hopmans and Bickford (2003) also observed that Phos-Check (which contains ammonium polyphosphates and sulphates) stimulates a three to ten-fold increment in available N over initial values in the 0-20 cm soil layer. Contrary to the results of Hopmans and Bickford (2003), who found no significant differences between control and Phos-Check - treated soils after 2 months, in BS+Ap the NH_4^+ -N levels remained significantly higher than in the other treatments for at least 3 months. The very high NH₄⁺-N levels in BS+Ap, and its relative persistence over time, are features of high ecological significance for the post-fire vegetation recovery, as seed germination and seedling establishment can be inhibited by NH_4^+ toxicity (see Britto and Kronzucker, 2002). In fact, in a greenhouse experiment with a burnt soil that received ammonium polyphosphate at the same dose as in the field, the germination of Lolium multiflorum and Agrostis tenuis seeds was reduced by 86-100% compared with the control burnt soil, and seedling establishment was completely inhibited (data not shown). However, in the medium-term the NH₄⁺-N supplied by the ammonium polyphosphate could have a fertilizing effect.

The initial lack of significant differences among treatments in the NO₃-N levels disappeared for BS+Ap between one and three months after the prescribed fire due to the active nitrification

of its large NH_4^+ -N pool, which is unquestionably the cause of the significant soil acidification observed in the BS+Ap plots during this period (see Fig. 2). On the other hand, the general increase in the NO₃-N concentration recorded at t=90 d in all soil treatments reflected a rise in the nitrifier microorganism activity, which was favoured by the higher soil humidity during the autumn, the season in which this sampling was carried out.

The increment in available P in BS compared with US was probably due to the strong mineralizing effect of fire on organic P reported by Saá et al. (1993), although in our case differences did not reach significant levels (Fig. 6). No fire fighting chemical had any influence on available P except ammonium polyphosphate, which provoked an increment in soil P levels that remained significantly higher (10-20 times) than for the other treatments, even one year after the fire. This result is in accordance with that of Hopmans and Bickford (2003) who, subsequent to the application of Phos-Check, also found a noticeable increase in available P (6-60 times in the 0-20 cm soil layer), which also remained over control soil levels over the following year. The medium-term (\geq 1 year) persistence of high P levels may acquire great importance for the post-fire vegetation recovery as: a) at high levels, P could have an antagonistic effect on Fe and Zn plant uptake (Marschner, 1995); and b) P is a common limiting nutrient. Moreover, the enhanced nutrient losses due to leaching and erosion that have habitually been reported in soils affected by wildfires (Chandler et al., 1983; Carballas et al., 1993) may increase risks of water eutrophication due to higher soil P outputs from the burnt soils treated with P-rich fire fighting chemicals.

The fire-induced increase of available cations observed in the burnt plots were in accordance with the results of DeBano et al. (1979) and Trabaud (1983) with burnt soils, and also with those of Fassbender (1975) for Ca, Mg and Na in oven-heated soils. This effect is probably due to the accumulation of ashes rich in oxides and carbonates of basic ions that have been reported by several authors (Chandler et al., 1983; Kutiel et al., 1990; Carballas, 1997), and also to the cations (especially Na) supplied by the fire fighting chemicals. While in all treatments divalent cation concentrations did not follow a clear temporal trend, due perhaps to seasonal changes in their availability, those of the monovalent cations decreased slowly during the first three months after the burning, which, in all probatility, was due to the greater solubility and propensity to leaching of the latter (Duchaufour, 1987).

Conclusions

Main consequences of soil burning are a transient (until 30-90 d) increase in pH and, not always significatively, in available nutrients (exchangeable NH_4^+ , available P, Na, K, Mg and Ca). The prescribed fire also provoked an initial decrease in soil δ ¹⁵N followed by a steady increase, which, together with a medium-term (from 90 to 365 d) reduction in SOM content, suggest a fire-triggered increment of soil erosion and N outputs (¹⁵N depleted).

The transient effects of prescribed fire on soil pH, exchangeable NH_4^+ and $\delta^{15}N$ were increased by the three fire fighting chemicals (slightly by foaming agent and Firesorb and more strongly by ammonium polyphosphate), whereas only the ammonium polyphosphate had effects on available P. The foaming agent increased the effects of fire on all available cations, whereas the addition of Firesorb only affected the Na and, to a lesser extent, Ca contents, and the application of ammonium polyphosphate only had minor effects on Mg levels. The extremely high initial levels of NH_4^+ -N and available P in BS+Ap can delay the post-fire recovery of vegetation due to the toxicity of NH_4^+ to seeds and seedlings and the antagonistic effect of P on Fe and Zn plant uptake. However, in the medium-term the NH_4^+ -N and available P supplied by the ammonium polyphosphate could have a fertilizing effect on vegetation growth. The enhanced nutrient losses due to leaching and erosion that are usually reported in soils affected

by wildfires may increase risks of water eutrophication due to higher soil N and P outputs from the burnt soils treated with N- and P-rich fire fighting chemicals.

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	Foaming agent Auxquímica RFC-88	Firesorb	Ammonium polyphosphate FR Cross	
Density $(a 1^{-1})$	1012 + 2	1024 + 2	2710 + 10	
Density (g 1)	1013 ± 2	1024 ± 2	2710±10	
Na (g l ⁻¹)	19.7 ± 0.5	8.7 ± 0.1	10.2 ± 0.1	
K (mg l ⁻¹)	22.8 ± 0.2	0.0 ± 0.0	1588 ± 11	
Ca (mg l ⁻¹)	10.8 ± 0.4	38.5 ± 0.6	1210 ± 19	
Mg (mg l ⁻¹)	0.0 ± 0.0	0.0 ± 0.0	3124 ± 77	
P (g l ⁻¹)	0.0 ± 0.0	0.2 ± 0.1	933 ± 5	
N (g l ⁻¹)	3.7 ± 0.3	12.6 ± 0.6	683 ± 21	
$\delta^{\ {}^{15}}N$	6.5 ± 1.4	6.5 ± 1.4	-0.9 ± 0.5	

Table 1. Density, total nutrient content and $\delta^{15}N$ (mean ± s.d.; n=3) in the three fire fighting chemicals used.

		\rightarrow	North		
BS+Ap	BS+Fi	BS+Fi	BS+Fo	US	
BS	BS	BS+Fo	BS+Fi	US	
BS+Fi	BS+Ap	BS	BS	US	
BS+Ap	BS+Ap	BS+Fo	BS+Fo	US	
4 m Slope					

Fig. 1.Plot distribution on the study area. Key: US, unburnt soil; BS, burnt soil; BS+Fo, burnt soil +
foam agent; BS+Fi, burnt soil + Firesorb; BS+Ap, burnt soil + ammonium polyphosphate.



Fig. 2. Evolution of pH in the 0-2 cm soil layer along a year after the prescribed fire. Key: US, unburnt soil; BS, burnt soil; BS+Fo, burnt soil + foam agent; BS+Fi, burnt soil + Firesorb; BS+Ap, burnt soil + ammonium polyphosphate. At each sampling date, different letters (a, b, c,...) indicate significant differences (P<0.05).</p>



Fig. 3. Evolution of C and N content in the 0-2 cm soil layer along a year after the prescribed fire. Key: US, unburnt soil; BS, burnt soil; BS+Fo, burnt soil + foam agent; BS+Fi, burnt soil + Firesorb; BS+Ap, burnt soil + ammonium polyphosphate. At each sampling date, different letters (a, b, c,...) indicate significant differences (P<0.05).</p>



Fig. 4. Evolution of δ^{15} N in the 0-2 cm soil layer along a year after the prescribed fire. Key: US, unburnt soil; BS, burnt soil; BS+Fo, burnt soil + foam agent; BS+Fi, burnt soil + Firesorb; BS+Ap, burnt soil + ammonium polyphosphate. At each sampling date, different letters (a, b, c,...) indicate significant differences (P<0.05).



Fig. 5. Evolution of NH₄⁺-N and NO₃-N content in the 0-2 cm soil layer along a year after the prescribed fire. Key: US, unburnt soil; BS, burnt soil; BS+Fo, burnt soil + foam agent; BS+Fi, burnt soil + Firesorb; BS+Ap, burnt soil + ammonium polyphosphate. At each sampling date, different letters (a, b, c,...) indicate significant differences (P<0.05).



Fig. 6. Evolution of available P content in the 0-2 cm soil layer along a year after the prescribed fire.
Key: US, unburnt soil; BS, burnt soil; BS+Fo, burnt soil + foam agent; BS+Fi, burnt soil + Firesorb; BS+Ap, burnt soil + ammonium polyphosphate. At each sampling date, different letters (a, b, c,...) indicate significant differences (P<0.05).



Fig. 7. Evolution of available Na, K, Mg and Ca content in the 0-2 cm soil layer along a year after the prescribed fire. Key: US, unburnt soil; BS, burnt soil; BS+Fo, burnt soil + foam agent; BS+Fi, burnt soil + Firesorb; BS+Ap, burnt soil + ammonium polyphosphate. At each sampling date, different letters (a, b, c,...) indicate significant differences (P<0.05).