What is the P value of Siberian soils? Soil phosphorus status in south-western Siberia and comparison with a global dataset

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Abstract. Climate change is particularly strong in Northern Eurasia and substantial ecological changes are expected in this extensive region. The reshaping and migration northwards of bioclimatic zones may offer opportunities for agricultural development in western and central Siberia. However, the bioclimatic vegetation models currently employed for projections still do not consider soil fertility, in spite of being highly critical for plant growth. In the present study, we surveyed the phosphorus

- 5 (P) status in the south-west of Siberia where soils are developed on loess parent material. We selected six sites differing in pedoclimatic conditions and the soil was sampled at different depths down to one metre in aspen (*Populus tremula* L.) forest as well as in grassland areas. The P status was assessed by conventional methods and by isotope dilution kinetics. We found that P concentrations and stocks, as well as their distribution through the soil profile, were fairly homogeneous at the regional scale studied, although there were some differences between sites (particularly in organic P). The young age of the soils, together
- 10 with slow kinetics of soil formation processes have probably not yet resulted in a sufficiently wide range of soil physicochemical conditions to observe more diverging P status. The comparison of our dataset with similar vegetation contexts on the global scale revealed that the soils of south-western Siberia and more generally of Northern Eurasia, often have (very) high levels of total, organic and inorganic P. The amount of plant-available P in topsoils, estimated by the isotopically exchangeable phosphate ions, was not particularly high, but was intermediate at the global scale. However, large stocks of plant-available
- 15 P are stored in subsurface layers which currently have low fine-root exploration intensities. These results suggest that the P resource is unlikely to constrain vegetation growth and agricultural development under the present conditions or in the near future.

1 Introduction

Occupying about 10 million km⁻² (6.7% of global terrestrial land), Siberia has a paramount influence because ecological
 processes occurring here can have an impact at the global scale. Ranging from latitudes 45 to 75° N, it covers several bioclimatic zones, from south to north: steppe, forest-steppe, sub-taiga, southern taiga, middle taiga, northern taiga, forest tundra and

tundra. As the global climate change signal is particularly strong in Northern Eurasia (IPCC, 2013), substantial reshaping of ecosystems is ongoing in the region. The expected increase in average air temperatures will be responsible for longer vegetation growing seasons and frost-free periods, for the melt of permafrost in northern areas and for the modification of soil freeze-thaw cycles in southern areas (Groisman et al., 2012). The intensity and distribution of precipitation may change, resulting

- 5 in differences in fire and hydrological regimes (Shiklomanov and Lammers, 2009; Shkolnik et al., 2010; Soja et al., 2007). Due to these altered physical conditions, vegetation communities may be modified. Projections indicate a shift of the Siberian bioclimatic zones northwards and a variation of their relative sizes (Jiang et al., 2012; Shuman et al., 2015; Soja et al., 2007; Tchebakova et al., 2009, 2010). In particular, the area occupied by steppe and forest-steppe would increase at the expense of taiga zones. These modifications of ecosystem features may result in alternative land uses (Bergen et al., 2012; Kicklighter
- 10 et al., 2014). Notably, under future climatic conditions, cropping of new species will be possible, or existing species may be used on vaster zones than at present in the southern parts of Western and Central Siberia (Kicklighter et al., 2014; Tchebakova et al., 2011). Primary productivity may be enhanced due to a "fertilization" effect induced by higher CO₂ levels in the air (Mooney et al., 1991; Norby et al., 2005; Schimel, 1995) along with longer periods sustaining plant growth. However, such projections lack the consideration of other important drivers of plant productivity such as the availability of resources like
- 15 nutrients and water in soils (Fernández-Martínez et al., 2014; He and Dijkstra, 2014; Oren et al., 2001; Reich et al., 2006a, b, 2014; van Groenigen et al., 2006). Even though there is an increasing interest in the study of Siberian ecosystems, functional ecological data remain sparse in the international literature (Gordov and Vaganov, 2010; Groisman and Soja, 2009).

In the domain of biogeochemistry, knowledge about the status of the major nutrients is lacking and potentially misleading assumptions could be made for Siberia. Together with nitrogen (N), phosphorus (P) is frequently a limiting resource for primary

- 20 production at the ecosystem scale (Elser et al., 2007; Harpole et al., 2011), but it is often considered that P is not the main limiting factor in northern ecosystems (Hedin, 2004; Reich and Oleksyn, 2004). In addition, P fertilization in agriculture is barely practised in Siberia, or not at all. However, a reconsideration of such a paradigm might be necessary in the context of global change. In fact, the increase of atmospheric CO₂ concentrations and N deposition, which are in general stronger and faster than any P input, are modifying the CNP stoichiometry of ecosystems (Peñuelas et al., 2013). As a consequence,
- 25 a progressive shift from N limitation to P limitation or N-P co-limitation can occur (Ågren et al., 2012; Peñuelas et al., 2012; Vitousek et al., 2010). These modifications of biogeochemical cycling at global and regional scales will participate in the way ecosystem reshaping is driven, for example through the modification of plant communities as they adapt to new stoichiometric constraints (Güsewell, 2004). It may also have consequences on agricultural potential at these scales.
- In addition, the global resources of P that are used for mineral fertilizer production are limited (Cooper et al., 2011; Cordell 30 et al., 2009). Therefore, enhancing our understanding of P cycles and managing them appropriately at the global scale is highly relevant (Cordell et al., 2011; MacDonald et al., 2011) since modern terrestrial P cycling is dominated by human activities (Filippelli, 2008). One solution which would help to restrict the use of primary P resources, would be the development of cropping systems in areas where the soils contains sufficient plant-available P to prevent the (massive) use of P fertilizers. In this perspective, parts of Siberia are expected to become climatically more suitable for agriculture. Assessing the P status of
- 35 these Siberian regions is thus of relevance, and this was the main goal of our study.

In the present study, we aimed to identify the P status of the soils of SW Siberia, a region characterized by different types of soil along a North–South climatic gradient. We addressed the following questions: (1) how is the P stock structured in terms of pools and with depth, in the soils of SW Siberia? (2) Which environmental factors control this P status? (3) How can we qualify this P status in comparison with a panel of contrasting pedo-climatic conditions at the global scale? To do this, we selected

5 six sites with contrasting pedo-climatic conditions in SW Siberia and presenting two characteristic vegetation covers, aspen (*Populus tremula*) forests and grasslands. We quantified total P, organic P, phosphate ions in solution and diffusive phosphate ions as a function of time in the soils from these sites. Classical soil analysis methods were combined with an isotopic dilution kinetics method. The sizes of P pools assessed at Siberian sites were compared with a global dataset compiled from 236 references.

10 2 Materials and methods

2.1 Site description

Many soils of south-western (SW) Siberia have developed on loess deposits – the Eurasian loess belt covers a broad latitudinal zone between 40 and 60° N – and present favourable texture and mineralogy for plant growth (Chlachula, 2003; Muhs, 2007). Soil formation depends on climatic conditions, vegetation cover and can be further shaped by anthropogenic actions. All

- 15 of these climatic conditions, vegetation cover and human activities differ in intensity essentially along a gradient from south to north. Consequently, from the common origin of loess deposits, the soils in SW Siberia have undergone different types of development and are classified in the main groups of Chernozems, Phaeozems, and Luvisols. Water-table movements, leaching of carbonates and organic matter accumulation or distribution throughout the profile are the most striking features in these soils. The soil forming processes related to soil moisture levels and dynamics as well as the stability of organic matter
- 20 (mineralization rates in relation to temperature and moisture regimes) and the type of vegetation are known to influence the soil P status (Giesler et al., 2002; Miller et al., 2001; Sundqvist et al., 2014; Vincent et al., 2014). We selected six sites in SW Siberia covering a transition including forest-steppe and sub-taiga zones. The main site characteristics are given in Table 1 (see also Tables S1–S3 in the Supplement and in Brédoire et al., 2016).
- All the soil profiles studied were developed on a loess parent material and vegetation cover had comparable features in terms of dominant species composition, stand age and low human impact (i.e. no active management for the last few decades; Tables S2 and S3 in the Supplement). The main characteristics of the initial loess material are the predominance of coarse-silt particles and clay and the presence of CaCO₃; the latter having had different fates related to the different soil development processes. In Barnaul (BAR), Chebula (CHE), Krasnozerskoye (KRA) and Salair East (SAE), the main soil forming processes are the formation and accumulation of organic matter, leaching of carbonates in the topsoil and formation of secondary car-
- 30 bonates in deep soil layers. Soils belong to the Chernozems and Phaeozems soil groups. In Salair West (SAW) and Tomsk (TOM) soils experience watertable movements, with periodical saturation. Consequently, clays are washed from the topsoil and accumulate in the deeper layers and carbonates have disappeared from the first metre of the soil profile. In addition, the

litter decomposes faster than in forest-steppe and the accumulation of organic matter is very low at the soil surface. At these two sites, soils belong to the Luvisols group.

Five of the sites were almost pure aspen (*Populus tremula* L.; Table S2) forest stands together with nearby grassland areas. One site (SAW) only had forest cover with aspen. So there were six forest sites and five with grassland in our dataset for SW Siberia. All aspen stands had closed canopies.

2.2 Sampling and preparation of the samples

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For each type of vegetation cover at each site, we delimited three study plots (about 300 m^2) about 200-2000 m apart. Those three plots were considered as replicates. One sampling campaign permitted sampling at all sites within three weeks in July 2013. At each plot, we dug a soil pit (with a surface area of about 2 by 1 m) down to 120 cm, except in SAE grasslands where we reached a dense schist material around 80 cm which prevented us from going any deeper than 100 cm. In each soil pit, about 1 kg of soil was sampled horizontally with hand tools, at depths of 5, 15, 30, 60 and $100 \text{ cm} \pm 5 \text{ cm}$. Another sample was taken with a cylinder (97 cm⁻³) to assess soil density.

Litter was sampled over an area of $30 \,\mathrm{cm}$ by $40 \,\mathrm{cm}$ in the vicinity of each soil pit. In this study we defined all the dead plant material deposited on the soil surface as "litter" (senescing leaf litterfall, small branches and senescing understorey vegetation

15 in forests; senescing herbaceous vegetation in grasslands). Consequently, the material collected in July, 2013 resulted mostly from the dead material from the previous vegetation season (2012) and the residues of older seasons, i.e. mainly OL and OF horizons, and possibly OH (at BAR, CHE, KRA and SAE).

Bulk soil samples were air dried to constant weight. After drying, soil samples coming from the same site and the same vegetation cover (i.e. 3 samples per site and per vegetation cover) were pooled and sieved at 2 mm to remove stones and coarse

20 roots. Soil density samples were not pooled. They were oven dried at 105 °C for 48 h and stones were removed when present (i.e. only in deep horizons of SAE).

Bulk litter samples were oven dried at $60 \,^{\circ}$ C to constant weight. They were then pooled by site and by vegetation cover and the composite samples (3 per site and per vegetation cover) were ground before chemical analyses, except for the isotopic dilution.

- Such soil preparation (sieving and drying) has been reported to affect physico-chemical processes only at a low magnitude (Černohlávková et al., 2009; Chapman et al., 1997). However, it also permits the prevention of microbial processes between sampling and analysis. We were careful to limit drying effects (Achat et al., 2012) using a "soft method" (air drying to a constant weight instead of oven-drying at 60 °C, for example). In addition, air drying is a part of the usual protocol of the ³²P isotopic dilution method (along with sieving at 2 mm, the use of a biocide such as toluene and measurements at 20 °C, see
- 30 below). This allows the replication of measurements and comparison of the results with those in the literature.

2.3 Physico-chemical analyses

2.3.1 Main soil properties

The French standard methods (Association Française de NORmalisation; AFNOR, 1999) were used for most of the physicochemical soil analyses. For soil texture, the five size-fractions for clay ($< 2 \,\mu m$ diameter), fine loam (2–20 μm), coarse loam

5 (20–50 μm), fine sand (50–200 μm), and coarse sand (200–2000 μm) were assessed after decarbonation (NF X 31–107). Soil pH-H₂O was determined in a water/soil suspension with a mass-to volume ratio of 1 g : 2.5 mL (NF ISO 10390). Total organic C and N contents were determined by dry combustion with oxygen (NF ISO 10694 and NF ISO 13878, respectively). Total calcium carbonate contents were assessed using a volumetric method (NF X 31–105). Poorly crystalline aluminium (Al) and iron (Fe) oxides were extracted with an ammonium oxalate solution (McKeague and Day, 1966).

10 2.3.2 Total, organic and inorganic P

Total P concentrations (P_{tot} , in µg g⁻¹ soil) were determined, after grinding, by ICP following wet digestion with concentrated fluoric (HF) and perchloric acids after calcination at 450 °C based on a standardized procedure (AFNOR NF X 31–147; AFNOR, 1999). Total soil organic P concentrations (P_{org} , in µg g⁻¹ soil) were determined as the difference between P extracted with H₂SO₄ in ignited and non-ignited soil samples (2 g of dry soil for 70 mL of 0.2 N H₂SO₄; Saunders and Williams, 1955),

15 concentrations were determined with a green malachite colourimetric method (van Veldhoven and Mannaerts, 1987). Total inorganic P concentrations (P_{inorg} , in $\mu g g^{-1}$ soil) were subsequently calculated as the difference between P_{tot} and P_{org} .

2.3.3 Plant-available phosphate ions

Plants take up P as ions from the soil solution. Thus, a good way of estimating a realistic plant-available P pool in the soil is to quantify both the concentration of phosphate ions in solution and the capacity of the solid phase to maintain this concentration.
To do this, we quantified the phosphate ions in the soil solution (*Cp* in µg mL⁻¹ soil solution or *Qw* in µg g⁻¹ soil) and the diffusive phosphate ions at the solid–solution interface (*Pr* in µg g⁻¹ soil). *Pr* is the quantity of phosphate ions that can be exchanged between solid constituents (ions are adsorbed on soil particles) on a concentration gradient. *Pr* results from molecular agitation; it can be considered as a "P buffering capacity". The sum of *Pr* and *Qw* is the isotopically exchangeable phosphate ions (*E*) and is considered as a good proxy for the gross amount of plant-available P (Fardeau, 1996; Morel and Sinaj, 1997; Erossard et al. 2011) as described below.

1997; Frossard et al., 2011) as described below. For each litter or mineral soil sample, five suspensions (1 g of litter or soil, sieved at 2 mm, with 10 mL of deionized water) were equilibrated for 16 h on a roller (40 cycles min⁻¹) at 20 °C (this temperature is commonly reported in the literature

were equilibrated for 16 h on a roller (40 cycles min⁻¹) at 20 °C (this temperature is commonly reported in the literature and is close to the average temperature of the soil at 20 cm in our study sites in the summer, Table S1). Toluene (10μL
toluene mL⁻¹ soil solution; M. Lineres, unpublished results) was added to the suspension at the beginning in order to stop microbial activity. This biocide does not affect P physico-chemical processes (Bünemann et al., 2007). The phosphate ions in

solution of the pre-equilibrated suspensions were labelled by introducing carrier-free ³²P ions in negligible concentrations but with known amounts of the introduced radioactivity (*R*). Suspensions were then sampled with a plastic syringe after 4, 10, 40, 100 and 400 min and filtered on a membrane at 0.2 μ m. Then, both *Cp* and radioactivity remaining in the filtered solution at the time of sampling (*r*(*t*)) were quantified. *Cp* was determined using a green malachite colourimetric method (van Veldhoven

5 and Mannaerts, 1987) and Qw was calculated using the volume of water (V in mL) and the mass of litter or soil (m_s in g):

$$Qw = Cp \times \frac{V}{m_s} \tag{1}$$

For each sample, Cp was not impacted by the sampling time of the isotopic dilution method (Fig. S1 in the Supplement). The radioactivity remaining in the filtered solution (r(t)) was determined in a counter (Packard TR 1100) using a liquid scintillation cocktail. In the steady state conditions of the suspension (Cp constant), the gross transfer of phosphate ions from

10 the solid constituents to the solution is equal to the gross transfer of phosphate ions from the solution onto the solid constituents. We assumed that no isotopic discrimination occurs between the two P isotopes (³¹P ions and ³²P ions) during the transfers between the liquid and the solid phases. The amount of unlabelled phosphate ions newly transferred from the solid constituents to the solution (Pr(t)) was then calculated from Qw and r(t) values following the principle of isotopic dilution (R is diluted in E).

15
$$\frac{R}{E} = \frac{r(t)}{Qw} = \frac{R - r(t)}{Pr(t)}$$
(2)

Rearranging Eq. (2) gives:

$$Pr(t) = \frac{Qw \times (R - r(t))}{r(t)} = Qw \left(\frac{1}{r(t)/R} - 1\right)$$
(3)

where: $\frac{r(t)}{R}$ (dimensionless) is the isotopic dilution ratio.

The theoretical Eq. (4), adapted from Fardeau (1993, 1996) was used to fit the experimental values of $\frac{r(t)}{R}$ closely, as 20 a function of isotopic dilution time:

$$\frac{r(t)}{R} = m\left(t + m^{\frac{1}{n}}\right)^{-n} \quad \text{for} \quad \frac{r(t)}{R} \ge \frac{r(\infty)}{R} \tag{4}$$

where: m and n are fitting parameters and $\frac{r(\infty)}{R}$ corresponds to the maximum possible dilution of the isotope, considering that all inorganic P can take part in the isotopic dilution. The value of $\frac{r(\infty)}{R}$ tends towards $\frac{Q_W}{P_{\text{inorg}}}$ (Fardeau, 1993; Frossard et al., 2011). The parameter m, which is the fraction of radioactivity remaining in solution after $1 \min(\frac{r(1\min)}{R})$, accounts for the

25 immediate physico-chemical reactions, while the parameter *n* accounts for the slow ones (Fardeau et al., 1991; Fardeau, 1993). The quality of the fit for Eq. (4) is shown in Fig. S2 in the Supplement and the values of *m* and *n* are provided in Table S5 in the Supplement.

Combining Eqs. (3) and (4), we can derive the value of *Pr* over time, each value corresponding to a pool of P more or less rapidly available to plants. The number and the size of such pools can be defined considering plant functioning (Fardeau, 1993).

30 In this study, we computed the values of Pr for 1 day, 1 week, and 3 months. While 1 day is the mean duration for active root uptake, 3 months is approximately the duration of the vegetation season in south-western Siberia and we might expect this to fit with intense root activity.

2.4 Data handling and statistics

Five soil layers were defined between 0 and 120 cm according to the soil horizon description of each soil pit (horizons were merged or divided in order to have 5 layers corresponding to the 5 sample depths, the profiles studied presented between 4 and 7 horizons, the mean number of horizons of the 33 profiles studied is 5). Assuming that elemental concentrations and soil

5 densities measured in each of the five defined horizons were representative of the entire horizon, we computed the stock (in $Mg ha^{-1}$) of each P pool using mean soil densities and horizon thicknesses:

$$\operatorname{stock} = \frac{1}{10000} \times [P] \times d \times h \tag{5}$$

where: [P] is the concentration of the P pool (in ugg^{-1}), d the soil density (in ggm^{-3}) and h the thickness of the soil horizon in cm. In litter, the P stocks were computed (in Mg ha^{-1}) using the mass and the surface area sampled:

10 stock =
$$\frac{1}{10000} \times [P] \times m_{\text{litter}}$$
 (6)

where: [P] is the concentration of the P pool (in $\mu g g^{-1}$) and m_{litter} the mass of litter (in $g m^{-2}$). We used the limit of -20 cmto distinguish between top- and subsoil to fit with most agronomic studies, but also with the zone where most of the fine root exploration occurs in SW Siberia (Brédoire et al., 2016).

- In order to gain an idea of the structure of the P stocks in the soils studied, we computed the relative proportions of the different P pools measured, as a fraction of P_{tot}. P_{tot} is the sum of P_{org} and P_{inorg}. Since a biocide was added to the suspension, 15 mineralization was stopped and we only measured physico-chemical processes. Thus, all exchangeable P (the sum of Qw and Pr) is part of $P_{\text{inorg.}}$ Pr being calculated as a function of time and the maximum time considered in this study being 3 months, the potential remaining fraction of P_{inorg} is considered as non-diffusive, or diffusive in more than 3 months. We note that when considering two values of Pr computed at different times, the pool of exchangeable P computed at the longer time includes the
- 20 one computed at a shorter time.

Relations between P parameters and soil physico-chemical properties were investigated by computing Spearman's rank correlation coefficients, scatter plots and (non-)linear regressions. Soil physico-chemical properties varied with soil depth (Table S4 in the Supplement) as well as the P parameters investigated (Table 2). Thus, we looked for correlations in each soil layer separately in order to avoid covariation and interdependence issues (Table S6 in the Supplement). We also analyzed correlations with fine root (diameter $< 0.8 \,\mathrm{mm}$) length density (FRLD) and fine root mass density (FRMD) measured in the

25 same soil pits and at the same soil depths as for the soil physico-chemical properties (Brédoire et al., 2016).

Since analyses were made on composite samples we did not quantify the variability of our measurements at the site level for given vegetation cover and soil depth. Thus, differences between sites were not tested through formal statistical tests. However, we calculated the coefficient of variation (ratio of the standard deviation to the mean) for each layer and vegetation cover.

All data management, (non-)linear regressions and statistical analyses (correlation coefficients and their significance), were 30 performed with R 3.2.1 (R Core Team, 2015).

2.5 Comparison at the global scale

To compare the phosphorus status of our study sites with other grassland or forest ecosystems and with croplands, we compiled data on different P fractions in soils. In practice, we used different requests involving keywords such as "soil", "phosph*", "total content", "isotopic dilution", "isotopically exchangeable P", "grassland", "forest", "woodland", etc. These

- 5 requests were carried out both in Web of Science and Google Scholar. To derive the pools of diffusive and isotopically exchangeable phosphate ions, we selected all publications using the same isotopic dilution procedures as in the present study (i.e. Fardeau's procedure; Fardeau, 1993, 1996). Additional publications were selected in order to improve the geographical coverage for total and organic P. In particular, we examined all the tables of contents of the *Soviet Soil Science* and the *Eurasian Soil Science* journals to provide a better cover of Northern Eurasian ecosystems. Based on all the se-
- 10 lected references, we compiled a dataset of different P fractions (total P, organic P, phosphate ions in soil solution, diffusive phosphate ions and isotopically exchangeable phosphate ions) in soils of grasslands, forests, or croplands. This database contained P values for up to 373 distinct sites depending on the P fraction, the geographical scale and the vegetation type studied, which were collected from 236 references. This database was representative of soils throughout the world as shown by the geographical distribution of compiled sites (Fig. 1), even though the studies using the isotopic dilution ki-
- 15 netics method in forest were sparse. In particular, out of the 116 forest study sites present in the compilation with values for inorganic P, organic P, phosphate ions in solution, diffusive phosphate ions and isotopically exchangeable phosphate ions –, 106 are located in France. Nevertheless, France does have very diverse soils and geology. The most represented soil types are Podzols, Cambisols and Luvisols, but Planosols, Leptosols, Calcisols, Arenosols, Regosols and Andosols are also present in some sites (IUSS Working Group WRB, 2014). The main parent materials are calcareous formations, eruptive
- 20 and metamorphic rocks, sandstone, detritic and weathered formations. Of these 106 French sites, 50 are hardwood forests (2 species) and 56 are coniferous forests (5 species). Therefore, our dataset was representative of forests at the global scale for total soil P, and representative of very diverse temperate forests for isotopic P data. In addition to the concentrations in different P pools, we also collected data on other soil properties such as texture, pH-water, cation exchange capacity (CEC), base saturation, organic C, and total N. The references of the data compilation are provided in the Supplement.

25 3 Results

3.1 Quantification of P pools

The concentrations of total P (P_{tot} ; 694–1095 µg g⁻¹ at -5 cm, 319–694 µg g⁻¹ at -100 cm), organic P (P_{org} ; 389–774 µg g⁻¹ at -5 cm, 37–79 µg g⁻¹ at -100 cm) and phosphate ions in solution (Qw; 2–22 µg g⁻¹ at -5 cm, 0.1–0.4 µg g⁻¹ at -100 cm) decreased with depth in the 1 m profiles at all the studied sites for both forest and grassland (Table 2). The litter layer presented

30 the highest concentrations for these pools; Qw being one to two order(s) of magnitude more concentrated in the litter (223– 638 µg g⁻¹) than in the upper soil layer (2–22 µg g⁻¹ at -5 cm). No systematic variation with depth was found throughout the profile for inorganic P (P_{inorg} ; 296–626 µg g⁻¹ at -5 cm, 282–616 µg g⁻¹ at -100 cm) and diffusive phosphate ions (e.g. Pr(1 day); 16–56 µg g⁻¹ at -5 cm, 2–67 µg g⁻¹ at -100 cm; Table 2). However, Pr(1 day) decreased in the three first mineral soil layers, except for the grassland in SAE.

We computed stocks (Mg ha⁻¹) for the different P pools (Table 3). With the exception of *Qw*, the subsoil contributed the most to the total stocks (72–85% of *P*_{tot}, 64–73% of *P*_{org} excluding SAE, 82–90% of *P*_{inorg} and 67–94% of *Pr*(1 day)). The three layers considered (litter, topsoil and subsoil) contributed almost equally to the total stock of *Qw* (respectively, 10–56, 26–65 and 7–49%). All sites presented values in the same order of magnitude for a given P pool and a given layer. The values for forest and grassland were also close. One notable difference occurred at the site SAE, where soil P pools were lower in forest than in grassland and where the pools in forest were lower than in the other sites. KRA presented the highest stocks in litter for all P pools, however, its stocks in the topsoil were the lowest (except for *Qw* and *Pr*(1 day) in grassland) and they

10 were also relatively low in the subsoil.

For each layer, we calculated the relative contribution of each P pool to total P (Fig. 2). With the exception of one grassland site (TOM), P_{org} accounted for more than 50 % of P_{tot} in the litter layer. The concentration of phosphate ions in solution (*Qw*) represented 20–38 % of P_{tot} in the litter layer with similar values for forest and grassland at each site. One site (TOM) presented much higher values of *Qw* in litter, reaching 45 and 67 % of P_{tot} in forest and grassland respectively. All the sites

- studied, and whatever the vegetation cover, exhibited the same pattern along the mineral soil profile. The relative proportion of P_{org} decreased, while the relative proportion of P_{inorg} increased with depth. In the two upper soil layers the distribution of P pools was very comparable. P_{org} accounted for 34–71 % of P_{tot} and Pr (3 months) for 3–13 %. Qw represented up to 2 % of P_{tot} at -5 cm, dropped below 0.1 % at -30 cm and was about 0.01 % at -100 cm. The proportions of Pr tended to be higher in forest than in grassland. Below -15 cm, the proportion of P_{org} decreased to 6–19 % at -100 cm. In the deepest layers, the proportions of Pr tended to be higher than in the first two, with notable exceptions: extremely low values at -100 cm in SAE,
- and Pr (3 months) representing 100 % of P_{inorg} at -60 cm in TOM grassland.

3.2 Relations between P pools and environmental parameters

We tested the correlations between P parameters and the main soil physico-chemical properties (Table S6 and Fig. S3 in the Supplement). P_{inorg} was significantly correlated with P_{tot} (Spearman's rank correlation coefficients ranging between 0.627 and 0.989). This was also observed between n and pH at most depths except at -15 and -100 cm (-0.636 to -0.793). In the three top layers, P_{org} was significantly correlated with organic C (0.682 to 0.843) and m with Qw (0.609 to 0.855). In the two deepest layers, Pr and isotopically exchangeable phosphate ions (E) were significantly correlated with the clay fraction (0.782 and 0.852). They were also negatively correlated with CaCO₃ (-0.649) at -60 cm (Table S6) but this was driven by one point which was very depleted in CaCO₃ and with very high Pr and E (Fig. S3). A few correlations were found with Al and Fe oxides: with P_{org} at -30 cm (-0.636) and with n at -60 cm (0.718).

A few relationships between fine root densities and P pools were significant (Table S6 and Fig. S3). At -15 cm, fine root length density (FRLD) was significantly and negatively correlated with Qw (-0.636), m (-0.764), Pr (1 day) (-0.691) and E (1 day) (-0.736). At -30 cm, FRLD was significantly and negatively correlated with n (-0.773) and fine root mass density (FRMD) with n (-0.655), Pr (0.636) and E (0.618).

No relationship was found between the different variables of the P status and any of the climatic parameters presented in Table S1 (data not shown).

3.3 Comparison at the global scale

Total P concentrations in topsoil (the first 20 cm of the soil) ranged at the global scale between 62 and 2480 μ g g⁻¹ in croplands,

- 5 between 19 and $3090 \,\mu g \, g^{-1}$ in forests and between 32 and $3548 \,\mu g \, g^{-1}$ in grasslands (Fig. 3). Our measurements in SW Siberia ranged between 345 and 770 $\mu g \, g^{-1}$ in forests and between 481 and 741 $\mu g \, g^{-1}$ in grasslands; these values were close to and above the global upper quartile for forests and between the global median and the upper quartile for grasslands. Compared with global cropland values, these Siberian concentrations were mostly above the upper quartile. Restricting the domain of comparison to Northern Eurasia, SW Siberian soils ranged between the lower and the upper quartiles for forests and between
- 10 the median and the upper quartile for grasslands, indicating that our sites are representative of Northern Eurasia. In the subsoil (-20 to -100 cm), less points were available at the global and Northern Eurasian scales for comparison. However, our SW Siberian forest and grassland soils occupied wider ranges in comparison to the corresponding vegetation types: from below the median to above the upper quartile at the global scale and a similar range of values to Northern Eurasia.

The concentrations of P_{org} , P_{inorg} and phosphate ions in solution (*Cp*) in the topsoil of the SW Siberian forests studied were 15 generally above the upper quartile, in comparison essentially with French forests having contrasting species, soil and geology (Fig. 4). In grassland, the SW Siberian values were mostly comprised between the global median and upper quartile. Compared with global cropland ranges, our measurements were around and above the upper quartile for P_{tot} and P_{org} , mainly below the upper quartile for P_{inorg} and from the median to above the upper quartile for *Cp*. Interestingly, the proportion of measured P_{org} (% of P_{tot}) varied quite a lot in the range reported at the comparative scales, particularly in forests where SW Siberian values 20 varied from below the lower quartile to above the upper quartile. In contrast with the other P pools, the concentrations of *Pr* (1 day) and *E* (1 day) were more moderate: they ranged between the median and the upper quartile in forests and between the

lower quartile and the median in grasslands, these values being lower than the global cropland median.

Relative to the soils included in the comparison dataset, the SW Siberian topsoils studied presented high contents (often above the third quartile of the global distribution ranges for both forest and grassland) of the silt fraction (2–50 µm) and low contents (below the first quartile) of the sand fraction (50–2000 µm) (Fig. S4). Their pH-water (5.3–7.5) tended to be higher when compared with the global levels (4.7–6.3, first–third quartiles), particularly in forests, but the ranges of the SW Siberian soils were similar to those of Northern Eurasia. Our study soils also had particularly high CEC (12–46 cmol + kg⁻¹ as compared to the global first–third quartiles 3–15 cmol + kg⁻¹) and extremely high base saturation (93–99 % versus 21– 87 %). Their concentrations in organic C were medium to very high (27–83 g kg⁻¹ versus 15–52 g kg⁻¹ for the first–third

30 global quartiles), as were their concentrations in total N (2–6 g kg⁻¹ versus 1–4 g kg⁻¹). Finally, the studied Siberian forest soils presented lower C: N, C: P, and N: P mass ratios than the global levels (C: N ranged between 12 and 15 versus 12–19 for the first–third global quartiles; C: P was 35–83 versus 50–153; and N: P was 3–6 versus 4–10) while the SW Siberian grasslands presented ratios in the same ranges as our global data compilation (Fig. S4).

Discussion 4

4.1 A relatively homogeneous P status

All the sites, for both aspen forest and grassland, presented a similar distribution of the P pools throughout the soil profile. Total P concentrations decreased with depth, mainly in relation to the decrease in the concentration of organic P (Table 2). The

5 stock variation was of the same order of magnitude between and within sites, and between contrasting vegetation cover types. The concentrations and the stocks we computed were close to those reported by Achat et al. (2013a). These authors reported standard errors of 1-27% for the concentrations of total P, organic P, inorganic P, phosphate ions in soil solution and diffusive phosphate ions, with three replicates per condition in two sites of the same region. Assuming a similar spatial variability, the concentrations and stocks we measured or computed for given soil layers appeared relatively homogeneous (values in the same 10 order of magnitude without notable outliers) at the regional scale.

Despite lower concentrations of the P pools in the subsoil, this contributed the most to the total stocks computed over 1 m (Table 3) because of its greater thickness. For the same reason, at the soil profile scale, inorganic P represented far more than 50% of the total P stock (Table 3), while organic P concentrations represented a high proportion of total P in the litter and in the first three soil layers (Table 2). For the whole profile, the P stock in the soils studied can be qualified as predominantly mineral down to 1 m.

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4.2 Environmental factors controlling the regional P status

In spite of small variations in the current P status of the soils investigated, we found that this status was impacted by a set of variables. At first, we observed that the amount of P was highly dependent on the amount of inorganic P, particularly in the subsurface layers of the sites studied (Fig. 2 and Table 3). Biogeochemical cycling and soil development processes explain 20 the P status with depth. In the topsoil, organic P represented a large part of total P (Fig. 2) and was related to organic C (Table S6 and Fig. S3). This is a direct consequence of P uptake, immobilization in plant tissues, followed by litterfall and subsequent accumulation in the top horizons (Barber, 1995). Clay minerals (on their edges), carbonates and organic matter have surfaces presenting positive charges that are reactive with phosphate ions (Gérard, 2016; Hinsinger, 2001; Parfitt, 1978). In the topsoil, the preponderance of organic P suggests that microbial processes may play an important role in the plant-

- 25 availability of P through the release of phosphate ions by mineralization. Conversely, in deep horizons P plant-availability is principally explained by mineral phases such as the clay fraction (Table S6 and Fig. S3). Contrary to other case studies (Achat et al., 2011; do Carmo Horta and Torrent, 2007; Tran et al., 1988; Walbridge et al., 1991), we found only a few relationships with Al and Fe oxide concentrations. These oxides also present positive charges that are known to be reactive with phosphate ions (Achat et al., 2011; Regelink et al., 2015).
- 30 The inspection of the correlations involving parameters m and n provides further insights into the drivers of phosphate ions exchange at the solid-solution interface. Correlations with m, which is the fraction of radioactivity remaining after one minute in the isotopic dilution, give information about rapid processes, while correlations with n are considered to be indicative of processes driving slow exchange reactions. Both parameters were related to the phosphate concentration in solution (Qw).

Generally m increased with Qw while n decreased (Table S6 and Fig. S3), in agreement with previous studies (Achat et al., 2009, 2013a; Fardeau, 1993; Morel et al., 2000). The dynamics of phosphate ions (slow reactions in the present study) were also dependent on pH, which modifies the charge of reactive solid surfaces and the speciation of phosphate ions (Barrow, 1983; Hinsinger, 2001; Strauss et al., 1997a, b; Ziadi et al., 2013). In addition, we found slight secondary effects of Al and Fe oxides

5 on slow phosphate ion exchange reactions: residual values of the parameter n increased with the increase in oxide contents (data not shown). This is in accordance with a preliminary study in SW Siberia (see more details on the effects of Al and Fe oxides on parameter n in Achat et al., 2013a).

Nevertheless, at some sites, a few layers did not exhibit the general features of the P status described above. These "outliers" can be understood by considering the impact of soil forming processes on soil physico-chemical properties. At TOM, the

- 10 periodical water-table movements may be responsible for the relatively stronger accumulation of clays and oxides in deeper soil layers (Table S4). Since these elements are reactive with phosphate ions, they may contribute to the higher concentrations and proportions of diffusive phosphate ions in these soil layers (especially at -60 cm in grassland; Table S4). At KRA, the accumulation of CaCO₃ (Table S4) could be responsible for high levels of diffusive phosphate ions in the subsoil. In fact, in alkaline soils such as in KRA, phosphate ions tend to precipitate with Ca cations which have an increasing solubility at pHs
- 15 above 8 (Hinsinger, 2001; Kuo and Lotse, 1972). At SAE, the schist material underlying the loess deposit (below -80 cm) is probably responsible for low P pool concentrations (particularly in forests) and extremely low proportions of diffusive phosphate ions in the deep layers compared with the other sites.

The restricted number of significant correlations identified – between P pools or isotopic dilution parameters (m and n) and soil physico-chemical properties – is not necessarily indicative of an absence of control of the P status. It may simply reflect

- 20 that the soil variables tested and P pool fall within a restricted range (with differences only up to one order of magnitude; Tables 2 and S4). In fact, Al and Fe oxides, but also C, have noticeable effects when studying data compilations with wider ranges of values (Achat et al., 2016). This restricted range of variation in SW Siberia, despite the contrasting soil processes, probably has to be related to the relatively young age of the soils, which have developed on loess deposited during the two last glaciations in the Quaternary era (Chlachula, 2003; Muhs, 2007). Additionally, soil forming processes are expected to be related to be
- 25 relatively slow in such a dry and cold region (Jenny, 1941).

4.3 High levels of total P fractions but moderate ones for plant-available P

In general, the SW Siberian soils studied presented very high concentrations (above the third quartile) for total P, organic P, inorganic P and phosphate ions in soil solutions in forest and high concentrations (close to the third quartile) in grassland, when compared with our compilation of data at the global scale (or diverse soil and geological contexts mainly in France, for

30 all P pools except total P, in forests) (Figs. 3 and 4). In addition, it might be possible to generalize the high level of total P stocks to Northern Eurasia, at least for the soils developed in the loess belt (Fig. 3). Of course, more field measurements are required to verify this statement, particularly in the vast zone currently covered by taiga and which is not in the loess belt. This result, if confirmed, would be of primary importance in the context of global change and of tensions related to resources of P for agriculture.

However, we noted that these relatively high concentrations of total P in SW Siberian soils did not automatically indicate a high P availability for plant nutrition. In fact, an important parameter is the ability of the soil to refill a depleted soil solution (e.g. due to root uptake) with phosphate ions. This "P buffering capacity" assimilates the quantity of diffusive phosphate ions between the solid and the liquid phases of soil. Contrary to the other measured P pools in the SW Siberian soils studied, the

- 5 concentrations of diffusive phosphate ions in the topsoil were not that high in comparison with global levels (although not being very low; Fig. 4). Several hypotheses may explain such a phenomenon, notably: the early development stage of the soils studied, the relatively slow kinetics of the processes producing phosphate ions, and the relatively low contents in metal oxides coupled with high pH. These three hypotheses are developed in the next two paragraphs.
- Following the conceptual model of Walker and Syers (1976), which describes the changes in the forms and amounts of P 10 pools with time, together with the comments we made in the sections above about the regional homogeneity of the P status, we concluded that these SW Siberian soils are probably in the early stages of soil development. This stage is characterized by the build-up of an appreciable organic P stock, but also by a stock of primary inorganic P, which remains large and is made available by weathering. Thus, there may still be a high potential of primary mineral weathering in these soils. Moreover, the mineralization of organic matter is another source of phosphate ions available to refill the soil "P buffering capacity" (Achat
- 15 et al., 2013b; Bünemann, 2015). The kinetics of both mineral weathering and mineralization might be relatively slow in SW Siberia due to the relatively cold and dry climate conditions (Jenny, 1941). The study of the kinetics of these mechanisms is relatively difficult and was not carried out within the scope of this study. However, they could be of importance, as they are likely to be impacted by global change. Organic matter mineralization would mainly depend on temperature and moisture (Bengtson et al., 2005; Paul et al., 2002). Mineral alteration would mainly depend on temperature and pH (Augusto et al., 2000;
- 20 Drever, 1994).

In French forests, the sums of Al and Fe oxides range from 4.5 to $1157.7 \text{ mmol kg}^{-1}$ and pHs from 3.6 to 8.3 (data compilation of 106 sites; Achat et al. (2016)). In comparison, our SW Siberian soils have low sums of Al and Fe oxides (68.44–184.08 mmol kg⁻¹) and very high pHs (5.37–7.16, Table S4). When compared with our global data compilation, SW Siberian pH values can be qualified as high to very high (around and above the third quartile) in forests, but remain in the global range

- 25 (first-third quartile) for grasslands, which generally exhibit higher pH values than forests in our data compilation (Fig. S4). The narrow ranges of values measured in our study sites may explain why we found only a few significant correlations between P pools and oxides. Furthermore, the (very) high pHs of these Siberian soils are probably partly responsible for the low reactivity of phosphate ions, notably because the number of positive charges decreases with increasing pH (Barrow, 1983; Hinsinger, 2001; Ziadi et al., 2013). Coupled with a low quantity of oxides (i.e. fixation sites), this might explain, at least partly, the
- 30 average values of diffusive (Pr) and isotopically exchangeable (E) phosphate ions in the SW Siberian soils studied, while total pools were (very) high.

The absence of correlation between fine root densities and P pools (Table S6) suggests that root exploration is not related to a search for P. In addition, the richness of total P in the soils studied also impacted elemental stoichiometry, particularly in forests where the C:P (35–83) and the N:P (3–6) ratios were relatively low compared with global levels (Fig. S4). These findings

35 - low C: P and N: P ratios in SW Siberian soils - are in accordance with the global mean values reported for forests (212

and 14.6 for C:P and N:P, respectively) but also for grasslands (166 and 12.3 for C:P and N:P, respectively) by Cleveland and Liptzin (2007). The ranges reported by Xu et al. (2013) for boreal forests, temperate broadleaf forests and grasslands also confirm that the Siberian soils studied have high amounts of P in comparison with other elements such as C and N. The relatively low N:P ratios measured in the litter layers (9–14; Table S4), in the soil (at least in forest, Fig. S4) as well as in

- 5 green aspen leaves (9–12, data not shown) suggest that P is unlikely to be the primary limiting nutrient for plant nutrition and litter decomposition, and that it could be N which might be the limiting factor (Aerts and Chapin, 1999; Reich and Oleksyn, 2004; Güsewell and Gessner, 2009). However, compared with our global data compilation, the C : N ratios of the Siberian soils studied were relatively low, particularly in forests (Fig. S4), indicating that these soils may have high amounts of total N. In that case, N would only be limiting if mineralization processes – on which most of plants are dependent for N resources – were
- 10 not sufficient to respond to plant requirements, at least during the period of maximum requirements (for example in spring). In general, the availability of P probably does not constrain plant growth in the ecosystems studied to any great extent. This conclusion is in line with the review made by Smurygin (1974) about fertilization experiments in the former USSR.

Will this P status be sufficient to fulfil future plant requirements in the context of global change? In the speculative situation where topsoils are depleted by intense biomass exports, our results suggest that the large P stocks in deeper soil layers could

sustain the demand (Table 3). This would imply a deepening of plant fine root systems, which has already been observed with ongoing global change, related to the increase in atmospheric CO_2 concentrations (Iversen, 2010) or with the lengthening of vegetation growing seasons (Lempereur et al., 2015; Majdi and Öhrvik, 2004). In SW Siberia, we suggest that deeper fine root systems would be more likely to be driven by other resources, in particular water, in the steppe and forest-steppe zones (Brédoire et al., 2016).

20 5 Conclusions

This study revealed that the concentrations, the stocks of the different P pools measured and their distribution in the soil profile were relatively homogeneous at the scale of SW Siberia, although there were some differences between sites (mainly organic P) possibly due to varying microbial activity and slight differences in physico-chemical soil properties. In this region, we argue that the young age of the soils, developed on loess parent material, coupled with slow kinetics of pedogenesis have probably not yet resulted in a sufficiently wide range of soil physico-chemical conditions to observe more diverging P status. The comparison of these Siberian P levels with similar types of vegetation on the global scale revealed high to very high levels of total, organic and inorganic P in the topsoils. It would seem to be possible to generalize these results to Northern Eurasia, but additional measurements are required to verify this statement. The amount of plant-available P in topsoils, evaluated as isotopically exchangeable phosphate ions, was intermediate on the global scale. However, large stocks

30 of isotopically exchangeable phosphate ions are stored in the subsurface layers where fine root exploration is currently low. These results suggest that the P resource is unlikely to constrain vegetation growth and agricultural development under present conditions and in the near future.

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Table 1. Main characteristics of the study sites. Additional information are provided in Tables S1–S3.

Site ID	BAR	CHE	KRA	SAE	SAW	ТОМ					
Geographical characteristics											
Name ^a	Barnaul	Chebula	Krasnozerskoye	Salair East	Salair West	Tomsk					
Ecological zone	Forest-steppe	Forest-steppe	Steppe to	Sub-taiga to	"Blackish taiga"	Sub-taiga					
	(southern part)	(northern part)	Forest-steppe	Forest-steppe							
Latitude (° N)	53.41	55.55	53.59	54.39	54.18	56.30					
Longitude (° E)	83.47	84.00	79.14	85.75	85.17	85.43					
$Elevation \left(m a.s.l.\right)$	221	186	141	305	358	232					
Climate characteristics (annual mean 1981–2010)											
Air temperature (°C)	2.7	1.3	2.9	2.3	1.2	0.9					
Precipitation (mm)	431.5	509.8	324.5	432.3	453.0	566.5					
Soil classification											
Soil in forest	Haplic Phaeozem	Haplic Phaeozem	Phaeozem	Leptic Phaeozem	Haplic Luvisol	Albic Luvisol					
Soil in grassland	Calcic Chernozem	Haplic Phaeozem	Calcic Hortic Chernozem	Leptic Phaeozem		Albic Luvisol					
Forest stand characteristics (mean values)											
Density (tree ha^{-1})	1664	387	767	1883	1144	1139					
DBH ^b (cm)	14.9	33.9	26.3	13.7	22.8	21.4					
Height (m)	11.2	28.0	18.7	15.7	24.8	18.2					
Age (years)	27	62	51	21	47	56					

^a Name of the closest city or name of the local area. ^b Diameter at breast height (1.3 m).

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Table 2. Phosphorus concentrations of different pools measured in litter and soil layers of south-western Siberia. Concentrations are expressed in μ g P g⁻¹ soil (or litter), depth is in cm. "Litter" means all the dead plant material deposited on the soil surface (senescing leaf litterfall, small branches and senescing understorey vegetation in forests; senescing herbaceous vegetation in grasslands) i.e. mainly OL and OF horizons, and possibly OH (at BAR, CHE, KRA and SAE) at the date of sampling.

P pool	Depth				Forest		Grassland							
	(cm)	BAR	CHE	KRA	SAE	SAW	TOM	CV	BAR	CHE	KRA	SAE	TOM	CV
Total P	Litter	1235.6	1174.5	1318.6	1231.3	1515.1	1011.8	13	1165.8	1340.4	1122.1	1318.6	953.1	13
	$^{-5}$	1017.3	851.4	729.2	1095.9	1039.2	759.7	17	877.6	956.2	847.0	921.3	694.2	12
	-15	951.8	663.7	615.6	908.2	864.5	676.8	19	864.5	938.7	698.6	873.2	624.4	17
	-30	960.6	628.7	541.4	416.1	768.5	558.9	30	755.4	899.4	628.7	563.2	528.3	23
	-60	764.1	676.8	394.7	323.5	585.1	489.0	31	593.8	689.9	515.2	611.3	493.4	14
	-100	637.5	659.3	408.2	319.2	646.2	528.3	27	593.8	694.2	414.4	593.8	537.0	18
Organic P	Litter	817.3	760.1	839.5	896.2	910.8	758.9	8	800.5	690.4	640.7	855.5	660.0	13
	$^{-5}$	391.6	405.6	418.5	773.7	470.0	436.1	30	388.8	539.2	539.5	521.8	398.7	16
	-15	319.6	286.8	335.1	614.6	379.4	375.7	31	358.3	472.2	413.7	495.6	333.6	17
	-30	332.1	208.7	286.4	202.6	284.3	217.5	21	276.9	422.0	343.6	100.3	206.1	46
	-60	156.8	133.5	96.9	58.2	114.7	137.8	30	108.1	90.1	201.4	123.4	89.7	38
	-100	50.3	58.8	79.5	37.1	68.5	63.5	25	48.6	78.7	45.7	37.0	64.5	30
Inorganic P	Litter	418.3	414.4	479.1	335.1	604.3	253.0	29	365.3	650.0	481.4	463.1	293.1	30
-	$^{-5}$	625.7	445.8	310.7	322.2	569.2	323.7	32	488.8	417.0	307.6	399.4	295.5	21
	-15	632.2	376.9	280.5	293.6	485.1	301.1	35	506.3	466.5	284.9	377.7	290.8	26
	-30	628.5	420.1	255.1	213.5	484.2	341.4	39	478.5	477.4	285.2	462.9	322.2	23
	-60	607.3	543.3	297.8	265.3	470.4	351.2	33	485.7	599.7	313.9	487.9	403.7	23
	-100	587.1	600.5	328.8	282.1	577.7	464.8	29	545.2	615.5	368.7	556.8	472.6	19
Phosphate ions	Litter	353.2	397.6	406.1	343.8	500.4	457.1	15	393.7	520.7	223.2	271.2	637.8	42
in soil solution	-5	10.3	15.8	11.2	22.1	17.9	1.5	55	6.4	2.9	13.7	1.9	1.7	95
	-15	5.3	2.3	1.5	6.0	3.3	1.6	57	1.0	1.3	1.1	1.0	0.7	23
	-30	3.6	0.8	0.7	0.4	0.9	0.2	114	0.4	0.8	0.8	0.5	0.1	56
	-60	1.1	0.3	0.5	0.1	0.2	0.2	89	0.2	0.3	0.7	0.7	0.2	62
	-100	0.2	0.2	0.2	0.1	0.4	0.3	44	0.2	0.2	0.4	0.1	0.1	50
Diffusive	Litter	88.8	84.2	72.9	80.0	103.9	37.3	29	36.6	23.5	59.9	85.8	3.0	77
phosphate ions	-5	42.5	56.1	34.9	53.4	48.1	26.3	26	31.6	20.3	36.1	21.1	16.2	33
in 1 day	-15	33.1	25.0	17.1	26.7	35.4	26.2	24	21.5	15.8	27.3	22.8	18.7	20
	-30	29.2	15.3	26.6	15.6	24.6	11.9	35	16.6	13.5	17.7	40.8	13.7	56
	-60	39.6	38.4	30.5	14.7	25.4	29.9	31	20.0	48.5	20.4	13.4	89.4	82
	-100	24.3	40.4	32.3	2.2	66.7	57.4	62	17.5	35.3	47.4	2.5	36.3	64

CV: coefficient of variation (ratio of the standard deviation to the mean, in %).

Table 2. Continued.

P pool	Depth				Forest						Grass	land		
	(cm)	BAR	CHE	KRA	SAE	SAW	TOM	CV	BAR	CHE	KRA	SAE	TOM	CV
Diffusive	Litter	88.8	84.2	72.9	80.0	103.9	37.3	29	36.6	23.7	74.3	109.8	3.0	86
phosphate ions	$^{-5}$	64.7	92.7	52.0	79.8	74.5	57.3	22	47.8	33.1	53.8	33.7	29.4	27
in 1 week	-15	52.9	45.9	29.8	43.5	70.0	51.0	27	34.2	26.8	51.3	42.7	39.3	24
	-30	46.4	28.8	57.2	29.4	51.5	26.6	33	28.4	24.3	30.9	71.0	31.7	51
	-60	70.1	73.9	51.1	31.9	53.9	64.4	27	35.2	95.9	37.1	20.2	178.7	89
	-100	47.0	73.1	61.7	4.0	137.6	91.8	65	30.4	65.1	94.3	4.6	64.5	67
Diffusive	Litter	88.8	84.2	72.9	80.0	103.9	37.3	29	36.6	24.0	94.3	143.7	3.0	96
phosphate ions	$^{-5}$	108.7	170.4	84.5	129.2	125.8	155.5	24	80.1	61.0	87.1	61.6	62.4	17
in 3 months	-15	95.3	99.1	60.4	79.1	166.4	119.5	36	62.4	52.4	116.4	96.2	102.6	32
	-30	83.4	65.0	154.6	66.7	133.8	75.6	39	57.1	51.5	63.5	146.2	94.5	48
	-60	147.4	174.1	100.3	87.9	144.2	175.4	27	73.9	234.2	80.4	34.1	403.4	93
	-100	111.5	159.3	144.1	8.5	355.3	169.6	71	62.4	145.0	231.7	10.2	136.9	72

CV: coefficient of variation (ratio of the standard deviation to the mean, in %).

Table 3. Stocks of the different P pools computed in $Mg ha^{-1}$ for the litter, the topsoil (about 0 to -20 cm), the subsoil (about -20 to -120 cm) and for the whole profile for the different sites. "Litter" means all the dead plant material deposited on the soil surface (senescing leaf litterfall, small branches and senescing understorey vegetation in forests; senescing herbaceous vegetation in grasslands) i.e. mainly OL and OF horizons, and possibly OH (at BAR, CHE, KRA and SAE) at the date of sampling.

				Grassland									
Layer	BAR	CHE	KRA	SAE	SAW	TOM	CV	BAR	CHE	KRA	SAE	TOM	CV
Litter	0.037	0.028	0.057	0.024	0.014	0.012	58	0.008	0.003	0.026	0.012	0.002	95
Topsoil	2.549	1.749	1.121	1.462	1.729	1.433	29	2.338	2.030	1.464	2.089	1.432	22
Subsoil	9.728	8.865	6.247	3.887	10.190	7.287	31	9.541	9.682	6.848	7.740	7.366	16
Total	12.313	10.642	7.424	5.374	11.934	8.732	29	11.888	11.714	8.338	9.840	8.800	16
Litter	0.024	0.018	0.036	0.018	0.009	0.009	55	0.006	0.001	0.015	0.008	0.001	89
Topsoil	0.894	0.782	0.622	1.003	0.764	0.805	16	0.995	1.079	0.896	1.185	0.791	16
Subsoil	2.010	1.520	1.787	0.958	2.075	1.705	24	1.809	2.002	1.964	1.043	1.604	23
Total	2.928	2.320	2.445	1.978	2.848	2.518	14	2.809	3.082	2.875	2.235	2.396	13
Litter	0.012	0.010	0.021	0.007	0.006	0.003	65	0.003	0.001	0.011	0.004	0.001	107
Topsoil	1.655	0.967	0.498	0.459	0.965	0.628	52	1.344	0.951	0.568	0.904	0.641	35
Subsoil	7.718	7.345	4.460	2.929	8.116	5.582	34	7.732	7.680	4.883	6.696	5.762	19
Total	9.385	8.322	4.979	3.395	9.086	6.214	35	9.078	8.631	5.462	7.605	6.404	20
Litter	0.010	0.010	0.017	0.007	0.005	0.005	52	0.003	0.001	0.005	0.002	0.001	63
Topsoil	0.018	0.015	0.008	0.016	0.012	0.003	45	0.008	0.004	0.012	0.003	0.002	67
Subsoil	0.017	0.004	0.006	0.002	0.008	0.003	83	0.003	0.005	0.008	0.005	0.002	44
Total	0.045	0.029	0.031	0.024	0.025	0.012	40	0.014	0.010	0.025	0.011	0.006	54
	Layer Litter Topsoil Subsoil Total Litter Topsoil Subsoil Total Litter Topsoil Subsoil Total	Layer BAR Litter 0.037 Topsoil 2.549 Subsoil 9.728 Total 12.313 Litter 0.024 Topsoil 2.010 Total 2.010 Total 2.028 Litter 0.012 Topsoil 1.655 Subsoil 7.718 Total 9.385 Litter 0.010 Topsoil 0.018 Subsoil 0.010 Topsoil 0.018	Layer BAR CHE Litter 0.037 0.028 Topsoil 2.549 1.749 Subsoil 9.728 8.865 Total 12.313 10.642 Litter 0.024 0.018 Topsoil 0.894 0.782 Subsoil 2.010 1.520 Total 2.928 2.320 Litter 0.012 0.010 Topsoil 1.655 0.967 Subsoil 7.718 7.345 Total 9.28 8.322 Litter 0.010 0.010 Topsoil 0.018 0.015 Subsoil 0.018 0.015 Subsoil 0.018 0.015 Subsoil 0.017 0.004 Topsoil 0.018 0.015 Subsoil 0.017 0.004	Layer BAR CHE KRA Litter 0.037 0.028 0.057 Topsoil 2.549 1.749 1.121 Subsoil 9.728 8.865 6.247 Total 12.313 10.642 7.444 Litter 0.024 0.018 0.036 Topsoil 2.804 0.782 6.222 Subsoil 2.010 1.520 1.787 Total 2.928 2.320 2.445 Litter 0.012 0.010 0.021 Topsoil 1.655 0.967 0.498 Subsoil 7.718 7.345 4.460 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CV: coefficient of variation (ratio of the standard deviation to the mean, in %).

Table 3. Continued.

	Forest									Grassland						
P pool	Layer	BAR	CHE	KRA	SAE	SAW	TOM	CV	BAR	CHE	KRA	SAE	TOM	CV		
Diffusive	Litter	0.003	0.002	0.003	0.002	0.001	0.000	56	0.000	0.000	0.001	0.001	0.000	118		
phosphate ions	Topsoil	0.094	0.083	0.039	0.052	0.073	0.054	32	0.068	0.038	0.059	0.052	0.039	25		
in 1 day	Subsoil	0.397	0.480	0.447	0.110	0.741	0.532	45	0.280	0.464	0.496	0.187	0.588	41		
	Total	0.494	0.565	0.490	0.164	0.815	0.586	41	0.348	0.503	0.557	0.240	0.627	35		
Diffusive	Litter	0.003	0.002	0.003	0.002	0.001	0.000	56	0.000	0.000	0.002	0.001	0.000	122		
phosphate ions	Topsoil	0.148	0.144	0.064	0.081	0.136	0.108	31	0.106	0.064	0.101	0.093	0.077	20		
in 1 week	Subsoil	0.707	0.898	0.843	0.223	1.535	0.937	49	0.487	0.882	0.961	0.313	1.142	46		
	Total	0.858	1.044	0.910	0.306	1.673	1.045	45	0.593	0.945	1.064	0.407	1.219	40		
Diffusive	Litter	0.003	0.002	0.003	0.002	0.001	0.000	56	0.000	0.000	0.002	0.001	0.000	126		
phosphate ions	Topsoil	0.260	0.291	0.117	0.141	0.304	0.267	35	0.186	0.121	0.202	0.199	0.187	19		
in 3 months	Subsoil	1.506	2.037	1.951	0.563	3.979	2.032	56	1.005	2.045	2.286	0.616	2.623	50		
	Total	1.769	2.330	2.071	0.705	4.284	2.300	52	1.191	2.166	2.491	0.816	2.810	45		

CV: coefficient of variation (ratio of the standard deviation to the mean, in %).



Figure 1. Location of the study sites (south-western Siberia is highlighted in white, data points in beige) and the data points from a literature compilation. Distinction is made between the quantity of information available for each point (colour) and between vegetation cover (shape). "All P pools" stands for: total P, organic P, inorganic P, phosphate ions in soil solution, diffusive phosphate ions and isotopically exchangeable phosphate ions. Winkel-Tripel projection, graticules 15°.



Figure 2. Structure of total P (P_{tot}) in terms of P ions in the soil solution (Qw), diffusive P ions at different time scales (Pr) and non-diffusive, or diffusive in more than 3 months, P ions ($P_{non-diff}$). Each fraction of P is expressed as % of P_{tot} . Note that the diffusive fractions for the longer times are included in the diffusive fractions at longer time (e.g. Pr(1 day) is included in Pr(1 week) and they are both included in Pr(3 months)). Depth "1" is the litter. "Litter" means all the dead plant material deposited on the soil surface (senescing leaf litterfall, small branches and senescing understorey vegetation in forests; senescing herbaceous vegetation in grasslands) i.e. mainly OL and OF horizons, and possibly OH (at BAR, CHE, KRA and SAE) at the date of sampling.



Figure 3. Comparison of total P concentrations in topsoils (about 0 to -20 cm) and subsoils (-20 to -100 cm depth) of south-western Siberia (coloured dots) with similar vegetation contexts (CRO: croplands; FOR: forests; GRA: grasslands) at the global scale and at the Northern Eurasian scale (box- and violin-plots). The "*n*" provided indicates the number of individual points used to build the box- and the violin-plots.



Figure 4. Comparison of the different components of the P status in topsoils (about 0 to -20 cm) of south-western Siberia (coloured dots) and different vegetation types (CRO: croplands; FOR: forests; GRA: grasslands – box- and violin-plots) at the global scale (croplands and grasslands) or at the country scale (forests mostly located in France, along broad gradients of soil and geology). "Iso. exchangeable" is an abbreviation for "isotopically exchangeable". The "*n*" provided indicates the number of individual points used to build the box- and the violin-plots.