

## Can long-term fertilization accelerate pedogenesis? Depicting soil processes boosted by annual NPK-inputs since 1928 on bare loess Luvisol (INRAE-Versailles)

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#### 2 annual NPK-inputs since 1928 on bare loess Luvisol (INRAE-Versailles)

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17 Abstract. Human activities worldwide menace beneficial soil ecosystem services, but long-term 18 anthropogenic impacts on soil properties and processes are often difficult to assess in field 19 conditions. Here we exploited INRAE's patrimonial '42-plots' bare-fallow experiment, an unique long 20 term experiment in the world created in 1928 in Versailles (France), to emphasize long-term impacts 21 of annual inputs of NPK fertilizers (ammonium, phosphate, potassium salts) and basic amendments 22 (lime, basic slag) on loess Luvisols. We selected plots receiving monovalent (Na<sup>+</sup>, K<sup>+</sup>), acid (NH<sub>4</sub><sup>+</sup>), 23 basic (Ca<sup>2+</sup>) and non-amended (reference) plots, thus embracing the today widely diverging 24 physicochemical surface soils conditions. Temporal changes of soil characteristics were studied on 25 historical archived topsoil samples, whereas soil-depth impacts were studied on samples from 26 subsurface horizons collected in 2015 until 120-cm depth. Bare-fallow management caused a rapid

1 organic matter (OM) decay, soil acidification, CEC reduction and lixiviation of cations. With reduced 2 OM-buffering capacities, specific fertilizer-induced physicochemical conditions enhanced the 3 development of several soil-forming processes.  $NH_4$ -fertilizers amplified soil acidification (pH < 4), 4 lixiviation (i.e. 1.5 kg m<sup>-2</sup> of Ca), aluminization of the exchange complex, and weathering of 5 ferromagnesian minerals and plagioclase feldspars. Under (NH<sub>4</sub>)<sub>2</sub>HPO<sub>4</sub> fertilization, a quasi-total 6 dissolution of chlorite and hornblende occurred, pointing to an acidocomplexolysis process in which 7 the PO<sub>4</sub><sup>3-</sup> anion likely plays the role of complexing organic acids in Podzols. Ammonium fertilizers also 8 affected E, B and C horizons. In reference plots, similar but lower effects remained restricted to the 9 surface horizon. Na/K-fertilizers favoured substantial clay translocation (*i.e.* 10-15 kg m<sup>-2</sup>) from the 10 Ap to underlying E and E/Bt horizons. Liming amendments counteracted acidifying effects of OM-11 depletion, and raised the pH to 8-8.5 and exchangeable Ca to > 95 %. It may be clear that the initial 12 design specifications do not allow a direct comparison with current conditions of agricultural soils. 13 However, in the view of global climate change, foreseeing a lowering of organic carbon contents in 14 soils, the 42-plots trial acts like an "alert launcher", forecasting risks of soil degradation with respect 15 to mineral soil phases, parameters and processes, generally buffered and masked by the presence of 16 organic matter. The 42-plots experiment forms a high-valued playing field for experimental research, 17 offering a unique centennial time-span of differing physicochemical properties in a soil context with 18 close initial pedogenetic connexion.

# Keywords: long-term bare-fallow experiment; fertilization; loess soil; clay leaching; acidification; mineral weathering

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22 Highlights

- 85 years fertilization forced soil forming processes in a bare-fallow loess Luvisol
- ammonium fertilizers promoted acidification, aluminization and mineral weathering
- (NH<sub>4</sub>)<sub>2</sub>HPO<sub>4</sub> caused acidocomplexolysis and loss of primary ferromagnesian minerals

- top horizon of non-amended reference plots showed notable pedogeochemical changes

#### 1 1. INTRODUCTION

2 Worldwide, agricultural intensification during the second half of the 20th century led to 3 decreasing organic matter (OM) contents and soil structure stability and to growing risks of soil 4 erosion (Montanarella, 2010; Lal, 2015). In northern France, loess soils are widely used for intensive 5 cereal cropping, but they are sensitive to acidification, loss of SOM, and erosion (Bresson and Boiffin, 6 1990; Le Bissonnais et al., 1996; Arrouays et al., 2002; Gis Sol, 2011). Anthropogenic pressure 7 represents major threats for sustainable land use and soil ecosystem services. Therefore, it is urgent 8 to identify, understand and anticipate the risks of soil degradation caused by human activities (Lal, 9 2015).

10 Recurring application of mineral fertilizers and amendments is increasingly questioned with 11 regard to potentially harmful impacts on physical, chemical and biological soil properties (Haney, 12 2017), as well as their role as a source for micropollutants (i.e. Kabata-Pendias, 2011; van Oort et al., 13 2017a). One main effect of mineral fertilizers on soil properties lies in the input of dominant cations, 14 either as target (K<sup>+</sup>, Ca<sup>2+</sup>, Mg<sup>2+</sup>) or accompanying elements (Na<sup>+</sup>, H<sup>+</sup>). By deeply modifying the cation 15 composition of the soil's exchange complex, fertilizers may significantly change soil properties and 16 processes (van Breemen and Buurman, 2013), particularly in soils with low SOM contents (Paradelo 17 et al., 2016). Thereby, ammonium fertilizers introduce substantial amounts of protons, causing soil 18 acidification and formation of protonated hydrogen clay. Such H-saturated clays are unstable since 19 protons enter the octahedral layer and free large amount of aluminium, and evolve rapidly into Al-20 saturated clay, finally leading to clay destruction (Eeckman and Laudelout, 1961; Bolt and 21 Bruggenwert, 1976). With time, the prolonged use of ammonium fertilizers generate large amounts 22 of exchangeable AI on the exchange complex (Sposito, 1989), highly toxic for plants. Monovalent Na, 23 and to a lesser extend K fertilizers promote clay dispersion and leaching, leading to soil densification 24 and periodically water logging (Levy and Torrento, 1995; Sparks, 2003; Paradelo et al. 2013). 25 Intensive soil liming by adding basic amendments may lead to high amounts of exchangeable Ca,

unbalancing the soil's Ca/Mg ratios and Fe availability, inducing chlorosis in plants due to iron
 deficiency (Filipek, 2011). In field conditions however, the effects of long-term fertilization are often
 difficult to assess due to varying soil nature, land use and lacking historical information on fertilizer
 input management.

5 Long-term agricultural experiments offer unique opportunities to study the cumulative effects of 6 fertilizers and amendments on crops and soils. Many historical experiments initiated since the middle 7 of the 19<sup>th</sup> century, primarily focusing on the benefits of fertilization on increasing crop production 8 and quality: *i.e.* Rothamsted, England (1843); Sachsen, Germany (1851); Gembloux, Belgium (1872); 9 Grignon, France (1875); Urbana-Illinois, United States (1876); Wageningen, The Netherlands (1877); 10 Askov, Denmark (1894). However, little attention was paid at that time to potential negative impacts 11 of chemical fertilizers on soil composition and properties. Therefore, when at the end of World-War 12 1, broad need raised for reviving the French agriculture, notably by introducing a massive use of 13 chemical fertilizers, a long-term bare fallow (LTBF) experiment was initiated in Versailles in 1928: the 14 42-plots trial giving special attention to the mineral soil fraction. Its initial objectives were to 15 "examine the impacts of prolonged applications of main N, P and K-fertilizers, as well as organic and 16 basic amendments on the composition and properties of loam soils" (Burgevin and Hénin, 1939). The 17 experiment was established on silt-loam textured loess soils, representative of large areas under 18 conventional cereal production in northern France and NW Europe, and under bare fallow 19 management (no vegetation, no cultivation). With its historical archive of soil samples, collected 20 since March 1929, the 42-plots trial is currently the oldest bare-fallow field experiment in the world. 21 Bare soil management was presumed to exacerbate the impacts of fertilizers on the soil (Burgevin 22 and Hénin, 1939). Although not virtually reflecting real field conditions of agricultural soils, bare-23 fallow trials represent a playground for experimental pedology under local climatic conditions, 24 offering exceptional opportunities to unravel long-term aspects of intensive fertilization impacts on 25 soil ecosystem services. Thus, the 42-plots trial gives access to the speed, chronology and limits of

soil processes that either may develop or amplify under long-term fertilization practices and corollary
 contrasting physicochemical conditions, offering an exceptional, centenary, retrospective view. It
 represents a valuable tool to answer questions whether chemical fertilizers can influence soil
 formation at a centenary time-span.

5 The present work assesses both temporal and soil-depth-impacts on major soil parameters, 6 organic carbon and clay contents, soil pH and cation exchange complex, in a bare-fallow Loess 7 Luvisol, constrained by almost 90 years of fertilization. Based on exhaustive previous 8 pedogeochemical work (van Oort et al., 2016, 2017a, 2018, 2020a,b), we selected a series of 9 treatments of nitrogen, phosphorus and potassium fertilizers as well as basic amendments and non-10 amended reference plot, representative for current contrasting physicochemical properties in the 11 soils of the experiment. For each treatment, we analysed surface samples from the historical 12 collection and compared them with analyses from subsurface E, Bt and C horizons, collected in 2015. 13 The aims were to depict the extent of differentiation of the main soil physicochemical properties 14 during almost 9 decades in the surface horizon, and to assess the magnitude and depth of impacts in 15 subsurface horizons. Considering the great number and wide panel of results, we employed a close 16 'Results-Discussion' canvas, linking inputs of fertilizers to effects in terms of accelerated soil forming 17 processes and pedogenesis.

NB. Limited amounts of historical samples can be provided on demand, for original complementary research
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#### 21 **2.** MATERIALS AND METHODS

22 2.1 Experimental design

The LTBF experiment of INRAE started in 1928 in the gardens of the Versailles Chateau on *in-situ*,
 silt loam textured Haplic Luvisol (IUSS, 2014), developed in Late Weichselian calcareous loess

1 (Antoine et al., 2003), initially containing 15-25% CaCO<sub>3</sub>. The experiment included 42 plots of 2 × 2.5 2 m within a total surface of 200 m<sup>2</sup>, a small area with a flat topography where limited edaphic and 3 geological heterogeneity is expected, thus favouring an initially close pedogenetic connection 4 between soils of each plot, with respect to soil nature, behaviour, and physical and geochemical 5 properties. Of course, this does not preclude some local variability of rapidly changing parameters 6 such as the soil organic matter contents and related exchange properties. Since the start, the 7 experimental design remained unchanged for more than 90 years. Sixteen fertilization treatments 8 were manually applied at the soil's surface, in duplicate, and incorporated in the 0-25-cm soil layer. 9 Equivalent NPK-inputs per ha/year are shown in Table 1. Ten reference plots without inputs 10 complete the 42-plots trial. The plots were kept free from weeds and the surface layer was turned-11 over by hand twice a year, in spring and autumn, to a depth of 25 cm. Samples have been collected 12 annually in the 0-25-cm-layer, nowadays every three to five years, air-dried, ground, sieved at 2 mm 13 and stored in the dark at room temperature. Five samples were taken in each plots, homogenized 14 and then about 800 g were stored. The soil archive currently includes  $\approx$  3000 bare-soil surface 15 samples, covering more than 9 decades, a world's unique collection (Figure 1).

#### 16 2.2 Sample selection

17 For the study of temporal changes of soil parameters, series of about 15 historical samples were 18 analysed for the 1929–2014 period with an average interval of 6 years, closer during the first decades 19 (3 years) where changes were supposed to happen faster, and more spaced in recent decades. The 20 first samples in the collection are from March 1929, i.e. 5 months after the start of the experiment in 21 November 1928 (Burgevin and Hénin, 1939). We selected different treatments accounting for major 22 contrasting physicochemical surface soil conditions, following previous detailed pedogeochemical 23 inventory work (van Oort et al., 2016, 2017a), in order to examine the principal fertilization-induced 24 soil processes. Our plot selection included two types of fertilizers with acidifying effects (ammonium 25 sulphate, ammonium phosphate), three types of fertilizers with clay-dispersive effects (sodium 1 nitrate, sylvinite, potassium chloride), two types of amendments with pH-increasing effects (basic 2 slag, calcium carbonate), as well as two non-amended reference plots (Table 2). In the same selected 3 plots, additional sampling was performed in 2015 in subsurface horizons to 120 cm soil depth, using 4 a root auger device ( $\emptyset$  8cm, h 15 cm, Eijkelkamp, Giesbeek, The Netherlands). The core samples 5 were separated according to soil horizon limits. Selected undisturbed parts were air-dried and resin-6 impregnated for thin-section preparation; the rest was ground, sieved to < 2 mm and stored for 7 chemical analyses.

8 2.3 Analyses

9 Analyses were performed according international standard protocols at INRAE's National Soil 10 Analysis Laboratory (LAS): pH in water (NF ISO 10390), grain size distribution (NF X 31-107), organic 11 carbon (NF ISO 10694) and nitrogen (NF ISO 13878). The cation exchange capacity (CEC) and 12 exchangeable cations was using the hexamminecobalt trichloride extraction method (NF ISO 23470), 13 noted noted CEC<sub>Cohex</sub>.), working at low concentrations, close to in situ soil field conditions (Ciesielski 14 and Sterckeman, 1997a,b). This reagent limits the development of additional variable charge 15 exchange sites with respect to methods using buffered solutions (e.g. ammonium acetate at pH 7, 16 Metson) and the release of elements from non-exchangeable sources (Ca from carbonates in alkaline 17 conditions, AI from aluminium oxyhydroxydes in acidic conditions, K from lattice positions). For the 18 42-plots trial, Pernes-Debuyser and Tessier (2002) modelled the differences between Cohex and 19 Metson as a function of soil pH. These authors showed that in 1999 the Cohex data for acid and 20 alkaline treatments varied from single to double (6.9-13.6 cmol<sup>+</sup> kg<sup>-1</sup>), whereas they differed less than 21 15 % when using the Metson method. Moreover, they highlighted a minimum CEC<sub>Cohex</sub> value of 22 8 cmol<sup>+</sup> kg<sup>-1</sup>, accredited to the intrinsic permanent charge of the mineral clay fraction for 1 kg of soil. 23 Selected undisturbed parts of the centre of core samples were transferred into hard cardboard

boxes (7.5 cm × 7.5 cm × 4 cm), air-dried and impregnated with polyester resin, before cutting and

polishing to 30-µm thickness for the production of thin sections (FitzPatrick, 1993). The nature, morphology and organization of coarse soil constituents and the fine plasma were studied with a petrographic Nikon Eclipse E400 polarizing light microscope, using the terminology currently employed in soil micromorphology (Stoops, 2003).

5 The silty-loam textured Luvisol contained about 40% of coarse silt (20-50 µm) and 20% of fine silt 6 (2-20  $\mu$ m). The latter fraction includes the main part of weatherable primary minerals whose 7 weatherability increases with decreasing particle size (van Oort, 2018). Consequently, impacts of 8 infra-centennial fertilization practices on the soil's solid phase will be visible first on fine silt fractions. 9 Hence, after preliminary standard granulometric extractions of clay, fine and coarse silt and sand 10 fractions, we separated the 2-20- $\mu$ m fine-silt fraction into 2-5  $\mu$ m, 5-10  $\mu$ m, and 10-20  $\mu$ m 11 subfractions by exhaustive repetitive sedimentation extractions in glass cylinders. X-ray diffraction 12 (XRD) diagrams were made on oriented deposits of fine silt subfraction suspensions (25 mg/ mL) with 13 a Siemens D-5000 Kristalloflex diffractometer (Co Ka radiation, 30 mA, 40 kV). Total element 14 concentrations were determined at LAS on fine silt subfractions after tri-acid (HF, HNO<sub>3</sub>, HClO<sub>4</sub>) 15 digestion (ISO NF X31-147) for Ca, K, Mg, Na, Al, Fe, and scandium (Sc) by ICP-AES spectrometry (NF 16 ISO 22036).

#### 17 2.4 Data presentation and statistics

Results of temporal and soil-depth changes of major soil parameters, sensitive to anthropogenic pressure on soils (organic carbon contents, pH, CEC, clay contents) are simultaneously presented for each selected fertilization treatment. For the reference plots, we presented average values and max/min bars. Specific marks used to identify different treatments are detailed in Table 2. Soil-depth concentration profiles were compiled using data from surface horizons sampled in 2014, and from subsurface horizons sampled in 2015.

ANOVA was used to assess the influence of treatment on the main soil properties (pH, OC, CEC, clay content, Table 3). The homogeneity of variance was tested using the Levene test and the

normality of data was checked using the Shapiro-Wilk test. When a significant effect of land use at a
level of significance of P<0.05 was found, the Tukey's multiple range test was used to separate</li>
groups. Data that did not pass the normality test were log-transformed for ANOVA. All statistical
analyses were performed using the R statistical package for MacOSX version R 3.1.3 (R Core Team,
2019) and the package R Commander version 2.6-1 (Fox and Bouchet-Valat, 2019).

6

#### 7 **3. R**ESULTS & DISCUSSION

8 3.1 PHYSICOCHEMICAL PROPERTIES

9 3.1.1 Organic carbon

10 After its use as extensive grassland, the organic carbon contents in the surface horizon decreased 11 rapidly under bare fallow management (Figure 2a), first by about 30% from 14-18 g kg<sup>-1</sup> to 10-12 g kg<sup>-1</sup> 12 <sup>1</sup> in only one decade (1929-1940), then more progressively reaching a content of 4-6 g kg<sup>-1</sup> in 2008. 13 Initial variability of OM contents during the first years of the experiments express a typical 14 divergence related to previous land use. The data from 2014 showed slightly increasing discrepancy 15 between the organic C data, for unknown reasons. Today, the organic matter (OM) pool is quasi-16 exclusively formed by persistent organic carbon, with a plural centennial residence time, and 17 intimately bound to the mineral fraction (Barré et al., 2010). More details on OM dynamics and 18 modelling on 42-plots historical samples, can be found in Barré et al. (2010, 2016, 2018), Lefèvre et 19 al. (2014), Menichetti et al. (2015), Paradelo et al. (2013, 2016), Lutfalla et al. (2017), and Cécillon et 20 al. (2018).

Bare fallow conditions lead to OM depletion in all treatments (except for horse manure plots). Since the last four decades (Table 3, Figure 2a), diverging trends of OM contents emerged: for acid and basic treatments, OC contents were higher than those of the reference plots, whereas for the monovalent treatments, they were either fairly similar (NaNO<sub>3</sub> plot) or noticeably lower (potassium

1 fertilizers, sylvinite, KCl). Concordant findings were reported by Paradelo et al. (2016) who examined 2 the relation between physical protection of organic matter and aggregate stability in surface samples 3 of KCl, CaCO<sub>3</sub> and reference plots. Such diverging tendencies illustrate the driving control of a marked 4 physicochemical ambiance on soil OM (Kögel-Knabner and Kleber, 2011). Usually, the role of Ca<sup>2+</sup> is 5 well recognized for its stabilizing effect on OM (e.g. Six et al., 2004; Wuddivira and Camps-Roach, 6 2006; Paradelo et al., 2015; Rowley et al, 2018). In acid soils of temperate regions, and in many tropical volcanic soils, the stabilizing role of Al<sup>3+</sup> was ascribed to the formation of Al-hydroxyl 7 8 complexes with OM (e.g. Scheel et al., 2008; Heckman et al., 2013; Takahashi and Dahlgren, 2016). 9 Monovalent Na<sup>+</sup> and K<sup>+</sup> produce a clay dispersive effect, leading to soil structure degradation and an 10 enhanced accessibility of microorganisms to OM (Levy and Torrento, 1995; Paradelo et al., 2013, 11 2016). The lowest OC contents were observed under K-fertilizers. Such different behaviour toward 12 OM dynamics of  $K^+$  with respect to Na<sup>+</sup> may be ascribed to smectite-illite transformation by K-13 retrogradation (Pernes-Debuyser et al., 2003), lowering the clays reactivity, specific surface and 14 exchange capacity (Andreoli, 1989; Andreoli et al., 1989), thus easing the decomplexation of OM.

Effects of centennial OM depletion in this experiment were mainly restricted to the surface horizon. In subsurface horizons (Figure 2b), the organic C content profiles showed little divergence between the treatments: below the E/Bt horizon, the range of variation between the different treatments was reduced to  $\approx 1 \text{ g kg}^{-1}$ , and total organic C content decreased from  $\approx 3 \text{ to} \approx 1 \text{ g kg}^{-1}$  in the calcareous loess.

This short overview emphasised the high-added value of the 42-plots LTBF experiment for unravelling complex mineral-organic associations that prevail in different pedogenetic environments (Kleber et al., 2015; Kögel-Knabner and Amelung, 2021). It offers indeed an exceptional historical sample set covering 92-years of OM decomposition, in a soil context with currently wide varying physicochemical conditions, but initially close pedogenetic connexion.

25 3.1.2 Soil pH

1 With time, pH ranges in the surface horizons significantly differed between the acid, basic and 2 monovalent Na/K and reference treatments (Table 3). For the reference plots and monovalent 3 treatments, the pH dropped by about 1 unit during the first 10 to 15 years (Figure 2c), for a great part 4 due to decomposing OM: CH<sub>2</sub>O (OM) +  $O_2 \rightarrow CO_2 + H_2O \rightarrow HCO_3^- + H^+$  (Bolt and Bruggenwert, 1976). 5 Since the 1940's, the pH in the reference plots stabilized around 5 (4.7-5.4); under potassium 6 fertilizers (KCl), the pH remained fairly close to those in the reference plots (Figure 2c), whereas 7 sodium-based fertilization (NaNO<sub>3</sub>) caused a slight pH increase with time, to 6.7-6.8. The input of a 8 mixed Na-K fertilizer (sylvinite) led to pH values intermediate between the pure K and Na treatments. 9 Ammonium fertilizers exert strong acidifying effects on soils (Bouman et al., 1995), due to the 10 production of protons by nitrification:  $NH_4^+ + 2O_2 \rightarrow NO_3^- + 2H^+ H_2O$ . Thus, di-ammonium phosphate 11 (NH<sub>4</sub>)<sub>2</sub>HPO<sub>4</sub> introduces one more proton with respect to ammonium sulphate (NH<sub>4</sub>)<sub>2</sub>SO<sub>4</sub>. Between 12 1929 and 1940, the pH in the ammonium fertilizer treatments dropped by about two units from 6.4 13 in 1928 (Burgevin and Hénin, 1939) to 4.4-4.6 today, as a conjugated result from nitrification and OM 14 decomposition. These data were close to pH determinations reported as soon as 1939 (Burgevin and 15 Hénin, 1939). A "plateau-value" around pH 4 occurred between ≈ 1960 and 2000, but since, a new 16 pH decrease was noted for the ammonium treatments, currently around pH 3.5. Such stepwise pH 17 decrease pointed to a changed soil acid neutralizing capacity (van Breemen et al., 1983), announcing 18 the alteration of soil constituents. For the basic amendments, the pH rapidly increased during the 19 two first decades of the experiment, in spite of the acidity produced by OM decomposition. The rate 20 of increase was higher for CaCO<sub>3</sub> than for basic slag. Since the 1950's, the pH continued to increase 21 more progressively, reaching currently 8.2-8.5. For both treatments, about 2 % of calcium carbonate 22 was found in the surface horizon in 2014 (van Oort et al., 2016).

In subsurface horizons of reference plots and K-based treatments, the soil pH evolved quite similarly, increasing gradually from pH 6 in the E horizon, to pH  $\approx$  7 in the E/Bt or upper Bt horizons and reaching pH 8 in the carbonate-rich Ck horizon (Figure 2d). Nevertheless, such pH-depth profiles

1 suggested a moderate acidification of the upper 50 to 60 cm, when comparing them with pH 2 gradients in similar, nearby, soils under cultivation with pH > 7, down from the plough layer 3 (Isambert, 1979; Moni, 2008). The Na-based fertilizers demonstrated such a pH-depth profile with pH 4 > 7. As to the long-term input of NH<sub>4</sub>-fertilizers, they strongly marked the soil with pH values by 1.5 5 to 3 units lower than observed for the reference plots, until the calcareous loess parent material 6 (Figure 2d). In the NH<sub>4</sub>-sulfate plot, the pH was < 4 in the upper 60 cm, and < 4.5 in the Bt horizon, 7 but under NH<sub>4</sub>-phosphate, the acidifying impacts diminished earlier in the soil profile, reaching pH 8  $\approx$  6 in the Bt horizon. Basic amendments had an opposite effect, with pH  $\geq$  8 in all soil horizons.

9 3.1.3 Clay contents

10 Temporal evolution of clay contents was determined for 9 sampling periods for each series of 11 samples (Figure 3a). During the first decades of the experiment, the interval of clay contents 12 remained rather low, between 15-20 g kg<sup>-1</sup> for all treatments. However, as organic matter depletion 13 proceeds, specific effects of contrasting cation compositions on clay dispersion/aggregation 14 (Talibudeen, 1981) become more clearly expressed. Thus, in the Na and K treatments, clay contents 15 decreased more rapidly than observed in the reference plots (Table 3) and are clearly lowest since 16 the 1990s (130-150 g kg<sup>-1</sup>) with respect to all other treatments. Such low contents point to a process 17 of dispersion and leaching of clay from the surface to subsurface horizons, favoured by Na and K salts 18 (Wilding and Tessier, 1988; Tessier, 1990). For the 42-plots experiment, clay losses between 1929 19 and 2014 from the 0-25-cm Ap horizon were estimated at  $\approx$  15 kg m<sup>-2</sup> for Na-based fertilization and 20 at  $\approx 8 \text{ kg m}^{-2}$  under K-fertilizers (van Oort et al., 2018). Surprisingly, in the acid treatments, clay 21 contents in the surface horizon tend to increase in comparison to reference plots, particularly since 22 the end of the 1990s. More detailed clay extraction studies are needed to confirm such tendency. 23 Increasing clay contents with time under strong acidic conditions was previously hypothesized to 24 result from the weathering of silt-sized primary minerals (van Oort et al., 2016). Such hypothesis was 25 supported by positive geochemical budgets of clay-constitutive or clay-associated major (Fe, AI) and

minor (Sc, Tl, As) elements in the surface horizon of ammonium fertilizer plots between 1929 and
 2014 (van Oort et al., 2018). Such effect was stronger for ammonium phosphate than for ammonium
 sulphate.

At depth, clay contents in E horizons were highest for Na and K fertilizers (Figure 3b). Strong clay gradients between Ap and E horizons argue for contemporary clay illuviation process, directly under the Ap horizon. At greater depth, clay contents under NaNO<sub>3</sub> inputs were clearly higher than for the other treatments, stressing the role of sodium for amplifying clay illuviation.

8 *3.1.4 Cation exchange capacity* 

9 In the reference plots, the CEC<sub>cohex</sub> decreased steadily between 1929 and 2014 (Figure 3c) as the 10 result of a twofold mechanism: a net loss of exchange sites due to OM decomposition, and a loss of 11 variable negative charges on organic and mineral components, neutralized by H<sup>+</sup> produced from 12 decaying OM (Bolt and Bruggenwert, 1976; Julien and Turpin, 1999). The CEC<sub>cohex</sub> values in soils 13 under ammonium-based fertilization decreased more rapidly than in the reference plots between 14 1929 and the early 1940's. Obviously, the additional input of protons by nitrification of ammonium 15 resulted in a rapid lowering of soil variable charges (Kunhikrishnan et al., 2016).

16 In monovalent Na/K treatments, the CEC<sub>cohex</sub> also rapidly decreased during the first decade of the 17 experiment, which can be attributed to the conjugated effects of disaggregation, enhancing and OM 18 mineralization and possibly favouring clay leaching. As shown in Figure 2a, the organic carbon 19 contents are lowest for the Na/K treatments as soon as the first decades of the experiment. Besides, 20 a marked soil structure degradation and decreased macroporosity for in the surface horizon for soils 21 under Na and K-fertilizers were already noted at the end of the first 10 years of experiment (Burgevin 22 and Hénin, 1939). For both acid and monovalent Na/K treatments, the CEC<sub>Cohex</sub> decrease lessened 23 and values remained between approximately 7 and 9 cmol<sup>+</sup> kg<sup>-1</sup>, close to the estimated permanent 24 charge of 8 cmol<sup>+</sup> kg<sup>-1</sup> (Pernes-Debuyser and Tessier, 2002). Such plateau level was more pronounced 25 for ammonium sulphate than for di-ammonium phosphate, and lower for K than for Na treatments.

1 Moreover, since the 1950's, CEC<sub>Cohex</sub> data under inputs of (NH<sub>4</sub>)<sub>2</sub>HPO<sub>4</sub> are obviously lower than in the 2 case of (NH<sub>4</sub>)<sub>2</sub>SO<sub>4</sub>. A receivable explanation refers to the formation of aluminium phosphates in acid 3 soil conditions and their great affinity for clay surfaces (Sposito, 1989). Thus, phosphates breaking 4 external OH<sup>-</sup> sites would results in lowered net charge (Parfitt, 1978), and preferential adsorption of 5 Al-phosphates on clay particles favours strong aggregation, and reduce clay surface area and 6 accessible exchange sites (Edzwald et al., 1976). In addition, adsorption of hydroxyl-Al in clay 7 interlayer also causes a CEC decrease (Barnhisel and Bertsch, 1989). Since 2000 however, the CEC<sub>cohex</sub> 8 for acid and Na/K treatments dropped again, attaining about 5-7 cmol<sup>+</sup> kg<sup>-1</sup> in 2014. Such low values 9 suggest recent modifications in the mineral soil composition. By contrast, under basic amendments, 10 the CEC<sub>Cohex</sub> varied little, decreasing from  $\approx$  16.5 to  $\approx$  13 in 85 years (Figure 3c). Liming improves soil 11 structure by increasing and strengthening clay-OM bonds (Oades, 1984; Kunhikrishnan et al., 2016), 12 thereby lowering OM accessible to microbial decomposition.

13 In subsurface horizons, CEC<sub>cohex</sub> values in soils under NH<sub>4</sub><sup>+</sup> fertilizers were clearly lower than in the 14 reference plots (Figure 3d), indicating that acidifying soil conditions led to a loss of variable charge to 15 considerable depths. The effect was greater for  $(NH_4)_2HPO_4$  than for  $(NH_4)_2SO_4$  in the E horizon, but 16 inversed in the Bt horizon, in agreement with a lower pH (6.1 vs. 4.2, resp. Figure 2b). In potassium 17 treatments, CEC<sub>Cohex</sub> data were lower than in the reference plots, until the Bt horizon, suggesting an 18 effect of K-retrogradation by vermiculite-smectite clay minerals (Pernes-Debuyser et al., 2003; 19 Huang, 2005). Among all treatments, the highest CEC<sub>Cohex</sub> values in subsurface horizons were 20 observed under Na-based fertilizers, suggesting clay enrichment by enhanced leaching. Under basic 21 amendments, the CEC<sub>cohex</sub> were higher than in reference plots, notably in E and E/Bt horizons (Figure 22 3d). At high pH, the CEC comprise both permanent and variable charges (Julien et Turpin, 1999).

23 3.1.5 Exchangeable cation compositions

Since 1928, soil acidification in the ammonium treatments produced a virtually perfect inversely
 proportional replacement of Ca<sub>exch</sub> by Al<sub>exch</sub> (Figure 4). Replacement of Ca by Al was more rapid under

NH<sub>4</sub>-sulphate (P2) than under NH<sub>4</sub>-phosphate (P14) fertilization: at the end of the 1950s, after only 3
 decades, the first exhibited already more than 60 % of Al<sub>exch</sub> versus < 25 % for NH<sub>4</sub>-phosphate, but
 such divergence diminished rapidly since the 1990s.

Today, Al<sub>exch</sub> occupies  $\approx$  95-98 % of the cation composition in both treatments. Until the 2000s, Mg<sub>exch</sub> represented  $\approx$  10 % for NH<sub>4</sub>-sulphate treatment, versus  $\approx$  20 % for NH<sub>4</sub>-phosphate, but its presence strongly decreased in 2014. In reference plots, significant amounts of Al<sub>exch</sub> appeared in the 1960s as the pH reached  $\approx$  5; Al<sub>exch</sub> proportions increased since the 2000s reaching  $\approx$  20 % in 2014.

8 Basic amendments (P35, P39) progressively chased all exchangeable cations other than Ca, Caexch 9 currently occupying 96-98 % of the exchange complex. Long-term input of K-fertilizers progressively 10 increased the proportion of Al<sub>exch</sub>, up to  $\approx$  30 % of the CEC for KCl (P23), and to 15-20 % of the CEC for 11 the mixed Na/K fertilizer sylvinite (P29). Under sodium-based fertilization (NaNO<sub>3</sub>, P4), the 12 proportion of  $Na_{exch}$  increased up to 10-20% of the CEC and little or no  $Al_{exch}$  was observed. 13 Remarkable amounts, up to 5 % of the CEC for Mnexch, were detected notably under inputs of 14 ammonium (P2, P14) and under Na-K-fertilizers (P23, P4). For NH<sub>4</sub>-treatments, the presence of Mn<sub>exch</sub> 15 highlighted a process of Mn lixiviation to depth under acid soil conditions; for Na/K-treatments, 16 higher Mnexch amounts were ascribed to clay leaching, aggregate dispersion, soil densification and 17 temporary anaerobic conditions as demonstrated by the typical mottling observed during profile 18 sampling.

At depth, high proportions of exchangeable aluminium in subsurface horizons emphasized the impacts of soil acidification due to repeated inputs of ammonium. For NH<sub>4</sub>-sulphate, Al<sub>exch</sub> exceeded 90 % of the cation composition as deep as the Bt1 horizon (70 cm) and still reached 25 % in the Bt2 horizon until 80 cm depth (Figure 5a). For NH<sub>4</sub>-phosphate, high Al<sub>exch</sub> rates (> 90 %) were observed until 40 cm depth in the upper horizons Ap and E, lowering to  $\approx$  50 % in E/Bt and to < 5 % in the Bt1 horizon (Figure 5b). Interestingly, in both soils, Mn<sub>exch</sub> reached maximum values ( $\approx$  5 %) as the pH raised above 4.5 in the Bt2 and E/Bt horizons under ammonium sulphate and -phosphate respectively, consistent with Mn-lixiviation in strongly acid conditions. Such detection of exchangeable Mn at depth in the acid treatment plots was remarkable, since for all other treatments, the exchangeable Mn contents were generally below the detection limit (< 0.005 cmol<sup>+</sup> kg<sup>-1</sup>). In reference plots, the effects of soil acidification under local climatic conditions remained mainly restricted to the Ap surface horizon, Al<sub>exch</sub> in the underlying E horizons was < 1% (Figure 5c).</p>

Basic amendments enhanced > 90 % Ca<sub>exch</sub> saturation of the CEC in all horizons, allowing a maximum of 5-10% of other exchangeable cations to occur, mainly magnesium (Figure 5d,e). Longterm input of K-fertilizers enhanced proportions of K<sub>exch</sub> up to 10-13 % until the B1 horizon (Figure 5f), but for sylvinite such abundance was hampered by appreciable proportions of Na<sub>exch</sub> (Figure 5g), occurring as deep as in the carbonated loess at 120-cm depth. Under Na-based fertilizer, about 10 % of Na<sub>exch</sub> was present in subsurface horizons down to the BC horizon (Figure 5h).

12 3.2 SOIL MICROSTRUCTURE AND MINERALOGY

#### 13 3.2.1 Micromorphology of contemporary pedofeatures

14 Pedofeatures are discrete soil fabric units recognizable from adjacent groundmass material by a 15 different concentration in one or more components, or by a different internal fabric (Stoops, 2003). 16 They are key indicators for current or past soil forming processes, changing land use or soil 17 management practices (Stoops et al., 2010), as well as for metal pollutant transfer in soils (van Oort 18 et al., 2007, 2017b, 2018). Thus, pedofeatures observed under specific fertilization treatments, 19 demonstrating occurrence, abundance or extension that distinctly differed from other treatments, as 20 well as from pedofeatures issued by local loess Luvisol pedogenesis, attest for contemporaneous 21 fabrics deriving from the 42-plots experimental conditions (Figure 6).

Distinctively under inputs of Na- and K-based fertilization, numerous layered and compoundlayered clay-silt coatings were observed covering the surface of large voids in subsurface horizons, with a thickness up to several hundreds of microns (Figure 6a-d). They mainly occurred in E and E/Bt

horizons, between 30 and 50 cm depth, but also in the upper parts of the Bt horizon, although of
lesser thickness. Their abundance and extent emphasized the amplitudes of estimations for claylosses from surface horizons, notably under Na–based fertilization, around 10-15 kg.m<sup>-2</sup> (10-15t.ha<sup>-1</sup>)
in 85 years (van Oort et al., 2018).

5 In reference plots and under acid ammonium-based fertilization, dark matrix coatings (matrans) 6 build up by a mixture of fine particles with various sizes and compositions (clay, silt, oxides and fine 7 organic matter) frequently occurred close below the plough layer (Figure 6e,f). Such matrans are 8 characteristic for surface horizons of agricultural soils with a low aggregate stability (Stoops et al., 9 2010). In acid soils, under input of ammonium fertilizers, black precipitates frequently occurred at 10 the surface of large pores (channels, vughs) or in the groundmass nearby pores (Figure 6g-i). Such 11 precipitates primarily contained manganese with minor contents of lead, zinc or nickel, as revealed 12 by micro-X-ray-fluorescence work (van Oort et al., 2018). Under ammonium sulphate, manganese 13 coatings (mangans) were present as deep as 80-85 cm, coinciding with low pH values (4-4.5, Figure 14 2d). Under di-ammonium phosphate, fine dendritic manganese nodules occurred in the proximity of 15 pores in the E/Bt horizon (pH  $\approx$  4.5). Such localized Mn-accumulations give evidence for strongly acid 16 conditions enhancing lixiviation of mobile elements.

#### 17 3.2.2 Mineralogy of fine silt fractions

18 We assumed that the impacts of infra-centennial fertilization practices would primarily affect low-19 resistant primary minerals in the finest silt fractions (cf. section 2.3). In order to substantiate the 20 impacts of acidifying ammonium treatments, hypothesized to lead to mineral dissolution, we 21 compared mineralogical compositions of fine silt fractions extracted from the 2014 surface horizon 22 samples of  $(NH_4)$ -sulphate,  $(NH_4)$ -phosphate, reference and calcium carbonate treatments (Figure 7). 23 Considering its rapid pH increase from the start of the experiment, restraining the evolution of 24 aluminosilicate minerals, we assumed that the CaCO<sub>3</sub> treatment might represent a better mineralogy 25 control than the reference plots. X-ray diffraction (XRD) diagrams were made on oriented fine silt

1 deposits, in order to favour the detection of basal  $d(00\ell)$  reflexions of (ferromagnesian)-2 phyllosilicates, primarily affected in loess soils under acid soil conditions (Hardy et al., 1999) as well 3 as the cleavage plan d(110) of ferromagnesian amphiboles (Oberti et al., 2018)

4 For the basic, reference, and ammonium sulphate treatments, the XRD diagrams of the oriented 5 fine silt (2-20- $\mu$ m) fractions (Figure 7a) showed distinct basal spacing reflexions for mica (d(001), 6 d(002)) at 1.0 and 0.5nm, respectively, and for chlorite (d(001), d(002), d(003) and d(004)), at 1.41, 7 0.707, 0.471 and 0.353 nm, respectively. The latter spacings point to a trioctahedral ferromagnesian 8 chlorite structure. Interestingly, the broad 1.4 nm reflexion in the CaCO<sub>3</sub> treatment suggested the 9 presence of vermiculite. Its absence in the other treatments suggested a local pedogenetic origin by 10 alteration from biotite/chlorite, but disappearing in the silt fractions in acidified soil conditions. 11 Several hkl reflexions at about 16°20 (0.649, 0.638 nm) and in the range of 25.5-29.0°20 (0.402, 12 0.378, 0.365 nm) indicated the occurrence of potassium feldspars and plagioclase (F). Amphibole 13 (d(110) at 0.842 nm) was poorly detectable on global 2-20- $\mu$ m-fractions. Strikingly however, the 14 basal reflexions of chlorite strongly diminished or were absent for the (NH<sub>4</sub>)<sub>2</sub>HPO<sub>4</sub>, treatment, 15 highlighting a process of dissolution of ferromagnesian phyllosilicates.

16 Additional size-fractionation of fine silt was performed to improve insight into mineral weathering 17 features. Figures 7b-d show X-ray diffraction patterns (5-15°2 $\theta$ ) for 10-20, 5-10 and 2-5  $\mu$ m-fine silt 18 subfractions, respectively. In the 10-20 µm fraction, basal reflections of chlorite ((001), (002)) and of 19 mica (001) occurred distinctly for the basic and reference treatments. In the acid treatments, the 20 intensity of chlorite peaks only little decreased for ammonium sulphate with respect to the 21 reference, but virtually disappeared for di-ammonium phosphate. The d(110) peak of amphibole 22 (hornblende), clearly detected for basic and reference plots, strongly diminished for (NH<sub>4</sub>)<sub>2</sub>SO<sub>4</sub>, and 23 missed for  $(NH_4)_2HPO_4$ . Interestingly, although not shown here, acidification by ammonium 24 fertilization affected more particularly Na/Ca-feldspar peaks, in comparison to XRD patterns of the 25 CaCO<sub>3</sub> treatment, notably 0.402 and 0.365 nm reflexions typical for plagioclase (Moore and Reynolds,

1 1997). In the 5-10 and 2-5-µm fractions, the basal reflexions of chlorite were also strongly reduced or 2 absent for  $(NH_4)_2HPO_4$ . Amphibole peaks were not detected for all treatments (Figure 7c,d). By 3 contrast, for  $(NH_4)_2$ HPO<sub>4</sub>, the intensity of the d(001) reflexion of mica was notably stronger in the 2-5-4 μm-fraction than in the 5-10 and 10-20-μm fractions, suggesting a process of mineral particle 5 breakdown under strong acidified soil conditions. Such findings were consistent with a hypothesis of 6 recharge of the soil clay fraction by primary phyllosilicate minerals (chlorite, mica) due to their 7 microdivision in the fine silt fraction (Hardy et al., 1999). The difference between partial ((NH<sub>4</sub>)<sub>2</sub>SO<sub>4</sub>) 8 or total  $((NH_4)_2HPO_4)$  mineral dissolution was ascribed to the complexation action by phosphate of Al, 9 liberated by weathering, favouring complete mineral dissolution; for ammonium sulphate, liberated 10 Al may occupy phyllosilicate-interlayers and favour the formation of secondary chlorite-like minerals. 11 Nevertheless, such processes of mineral breakdown and particle transfer from coarser to finer silt 12 fractions, and in fine, to the clay fraction, are difficult to unravel by means of X-ray diffraction.

13 3.3 INDICATORS FOR ACCELERATED SOIL FORMING PROCESSES AND PEDOGENESIS

#### 14 3.3.1 Natural soil chronosequences on loess under temperate climatic conditions

15 After an initial stage of carbonate dissolution followed by decalcification, the clay migration 16 process primarily characterizes the soil evolution on aeolian loess formations in Europe (Jamagne, 17 1973, 2011). With time, this process generates growing textural and structural contrast between the 18 eluvial E and illuvial Bt horizons, enhancing temporary water logging. Such hydromorphic, and 19 attending acidified, soil conditions produce clay destruction in the upper part of the Bt horizon, and 20 downward tonguing of the E horizon (Jamagne and Begon, 1984; Jamagne et al., 1984) with a related 21 soil chronosequence Eutric Cambisol  $\rightarrow$  Haplic Luvisol  $\rightarrow$  Retisol. Further intense soil acidification may 22 engender podsolization processes at the soils surface (Brahy et al., 2000), driving a Retisol-'incipient 23 micropodzol' development.

24 3.3.2 Indicators of amplified clay translocation under prolonged inputs of Na/K fertilizers

1 In France, contemporary impacts of human activity on clay composition and distribution patterns 2 in Luvisols were recently studied in case of changing land use (Sauzet et al., 2016), drainage 3 conditions (Montagne et al., 2013; Cornu et al., 2007, 2017), or centenary wastewater discharge 4 (Thiry et al., 2013; van Oort et al., 2008, 2017b). However, all these works referred to preliminary 5 comprehensive field surveys in soil pits or trenches, a basic study-step proscribed in the case of the 6 42-plots experiment with respect to the small size of the plots. In our work, micromorphology study 7 provided first qualitative indication for contemporary clay illuviation in subsurface horizons under 8 Na- and K-based fertilization (Figure 6). Quantitative evidence for an amplification of the clay 9 leaching process can be obtained by calculating textural differentiation indexes (TDI) (Figure 8). 10 Although such texture contrasts in Luvisols are generally established with respect to the eluvial E 11 horizon (Baize, 2018), we here calculated TDI with reference to the Ap horizon, the surface layer 12 being primarily impacted by fertilization and related modified exchangeable cation compositions 13 (Figure 4). At the scale of the solum, the TDI between the Ap and the most clay-rich Bt horizons 14 showed values close to 1.5 for the basic, ammonium and reference plots (Figure 8a), referring in the 15 French 'Référentiel Pédologique' (RPF) typology (AFES, 2009) to 'Néoluvisols', in the FAO 16 classification system to Eutric Luvisols (IUSS Working Group WRB. 2015). For monovalent Na & K 17 treatments, the TDI increased to 1.7 for KCl, 1.8 for sylvinite and to 2.2 for NaNO<sub>3</sub>. In the RPF system, 18 a TDI > 1.8 and 10-30% clay in the eluvial horizon refers to 'LUVISOLS TYPIQUES' (AFES, 2009) or Haplic 19 Luvisols (IUSS Working Group WRB. 2015). Moreover, additional field observation outlined 20 periodically water stagnation at the surface of the Na and K plots following each significant rainfall 21 event, leading to clay and soil structure degradation, and a premise for the development of Retisols 22 and some Planosols.

Between the surface Ap and the underlying E horizon, strong increasing textural differentiation was observed for monovalent treatments, to about 1.5 for K-based fertilizers and to 1.8 for NaNO<sub>3</sub> (Figure 8b), in comparison to the TDI values of  $\approx$  1.1-1.2 for basic and acid NH<sub>4</sub>- based treatments.

1 Reference plots showed a TDI close to 1.3. These findings clearly attested for a contemporary and 2 short-range intensification of clay migration out from the surface horizon. Interestingly, when 3 compiling TDI data conventionally, *i.e.* comparing the clay contents of the eluvial E and the most clay-4 rich Bt horizon, little or no difference was found between the different fertilization treatments 5 (Figure 8c). Our findings on notable increasing TDI's since 4 or 5 decades (Figure 3a) gave clear 6 evidence for an amplified clay transfer, boosted by prolonged inputs of Na/K-based fertilizers. So far 7 these transfers concern primarily clay eluviation from the Ap and its illuviation in the E horizon, but 8 also such clay translocation also already affects deeper E/Bt and Bt horizons, as shown by the 9 presence of contemporary thick clay coatings (Figure 6).

#### 10 3.3.3 Indicators of soil acidification and corollary mineral weathering

11 Late-Weichselian loess formations in northern France contain predominantly guartz (60-70%), and 12 notable amounts of Na-Ca plagioclase, K-feldspars, mica (muscovite, biotite), chlorite, and clay 13 minerals (smectite, vermiculite, illite, and kaolinite). In addition, they include trace amounts of heavy 14 minerals (epidote, amphibole, garnet), with proportions depending on the geological source basins of 15 Pleistocene aeolian silt and sand covers in north-western Europe (Jamagne, 1973; Antoine, 2002; 16 Antoine et al., 2003). According to Goldich (1938), ranking the resistance of mineral groups to 17 weathering, ferromagnesian amphiboles (hornblende) and phyllosilicates (biotite, chlorite) weather 18 more rapidly than dioctahedral mica (muscovite), Ca-rich feldspars more rapidly than Na-rich ones 19 and plagioclase more rapidly than potassium feldspars (Figure 9).

20 Mineral weathering depends on a wide panel of physicochemical, biochemical and hydrodynamic 21 soil parameters (Wilson, 2004). Although weathering rates are arduous to generalize, orders of 22 magnitude have been established for ferromagnesian silicates and feldspars under controlled 23 experimental conditions assumed to mimic real field conditions. Dissolution times of decades to one 24 century have been calculated for chlorite, biotite and for the calcium feldspar anorthite (Kodama et 25 al., 1983; Lasaga, 1984; Wilson, 2004), a time span compatible with the duration of the 42-plots

1 experiment. Many workers stressed the role of low pH and organic acids in intensifying mineral 2 weathering. Thus, vermiculitization was mentioned as a "simple form of weathering of biotite and 3 chlorite" (Wilson, 2004), but under more intensive weathering conditions, they may convert directly 4 to kaolinite. However, under strong acidic conditions and favoured by complexing organic acids, 5 ferromagnesian minerals may be completely dissolved (Robert et al., 1980a,b; Drever and Stillings, 6 1997), a major soil forming process in Podzols (van Breemen and Buurman, 2013). In our work, under 7 di-ammonium phosphate additions, phosphate apparently replaces the role of complexing organic 8 agents in strong acidified soil conditions, favouring a quasi-complete dissolution of ferromagnesian 9 minerals, as was demonstrated by XRD work (Figure 7). These findings evoke conditions close to 10 those of the acidocomplexolysis process (Pédro, 2018), typical for Podzols.

11 During experimental weathering of ferromagnesian minerals, substantial amounts of constitutive 12 Mg, Fe, and Al were liberated (Kodama et al., 1983). In Late Weichselian loess formations in north-13 western Europe, the main reserve of weatherable primary minerals is located in the silt fraction 14 (Pésci, 1990, Antoine et al., 2003), notably present in fine silt (Hardy et al., 1999). We therefore 15 analysed the major element compositions in the silt fractions (Figure 10) to support XRD findings on 16 mineralogical alterations issued by prolonged ammonium fertilizers (Figure 7). The concentrations of 17 Mg, Fe and Al were clearly lower in A, E and Bt horizons under NH4-additions, compared to the 18 reference and CaCO<sub>3</sub> plots. Their concentration patterns showed remarkable mutual similarities 19 (Figure 10), but also regarding their respective pH-profiles (Figure 2d). Moreover, both 20 concentration-deviations and total concentrations decreased from the finest to the coarsest fine silt 21 fractions (Figure 10 a-c). Although Mg, Fe, and Al are not exclusively constitutive of ferromagnesian 22 phyllosilicates and amphiboles, their concentration patterns strongly advocate for an increased 23 weathering intensity under acidified soil conditions within an infra-decennial time-span, 24 preferentially affecting the finest fine-silt fractions. Interestingly, the concentration profile of 25 scandium (Sc), showed remarkable similarities with the distribution patterns of Mg, Fe and Al

1 concentrations (Figure 10). Scandium, a rare earth trace element, was used in previous work to 2 validate geochemical budget calculations for major and trace elements in surface horizons of all 42 3 plots since the start of the experiment (van Oort et al., 2017a, 2018). In loess soils, scandium was 4 mentioned to occur quasi-exclusively in ferromagnesian amphibole-hornblende, chlorite and biotite 5 (Mitchell, 1964; Das et al., 1971; Horovitz et al., 1975; Hocquard, 2003; Aide et al., 2009), due to its 6 frequent substitution with iron, eased by comparable ion radius (Sc: 0.075 nm, Fe<sup>2+</sup>: 0.076 nm, Fe<sup>3+</sup>: 7 0.064 nm). Therefore, scandium concentrations may represent a highly valuable indicator for 8 weathering of ferromagnesian minerals. Moreover, varying Sc contents in Luvisols was linked to 9 translocation of phyllosilicate clays (Horovitz et al., 1975), and correlated with structural iron (van 10 Oort, 2018). Hence, scandium may also represent a useful indicator for the clay leaching process.

11 By contrast, the soil-depth distribution profiles of sodium and potassium presented somewhat 12 higher Na and K concentrations in acid ammonium treatments than in reference and CaCO<sub>3</sub> plots 13 (Figure 10). This tendency may indicate a relative accumulation of more resistant Na/K-feldspars in 14 acid soil conditions, at the expense of calcium-rich feldspars, in accordance with Goldich's mineral 15 stability series (Figure 9). The effect was more distinct for Na than for K, as beyond potassium 16 feldspars, a notable amount of K occurs in micas, including resistant muscovite and more 17 weatherable biotite. The strong increase of Ca concentrations in the finest silt fractions at the surface 18 horizon under liming amendments denoted the precipitation of calcium carbonate, promoting soil 19 aggregation and increasing soil structure stability (Paradelo et al., 2013, 2016). However, lower Ca 20 concentrations in the ammonium treatments with respect to the reference plot also suggested an 21 increased dissolution of Ca-rich feldspars. In this respect, geochemical budgets established for the 22 1928-2014 period (van Oort et al., 2018) showed a net loss of Ca from the surface horizons as high as 23 -1.5 kg m<sup>-2</sup> for the ammonium fertilization treatments and -1.2 kg m<sup>-2</sup> for the reference plots. These 24 losses represent almost twice the amounts of calcium removed by lixiviation of exchangeable Caexch 25 from the CEC, thus attesting for a process of mineral dissolution of Ca-rich plagioclase.

1

#### 2 4. CONCLUSIONS

3 The large historical sample collection of INRAE's long-term bare fallow experiment allowed us to 4 depict accurately the temporal changes of soil physicochemical properties constrained by prolonged 5 fertilization in the surface horizon of a loess Luvisol. They were compared with current soil-depth 6 parameter profiles, studied on subsurface samples from E, B and C -horizons. The options taken at 7 the start of the 42-plots trial, a continuously application of NPK inputs on the same plots, and a bare 8 fallow management, exacerbated the effects of fertilization treatments on major soil characteristics, 9 in accordance with the initial objectives reported in 1939 by Burgevin and Hénin. It may be clear that 10 a straightforward comparison between bare-fallow soils and agricultural soils in real field conditions 11 is precarious. However, the experimental design of the 42-plots trial offered valuable insight into 12 hazards of intensive agriculture on soil forming processes and soil development in loess Luvisols 13 under intensive agriculture with strongly lowered organic carbon contents at a centenary timespan. 14 Such findings are relevant for agricultural areas under intensive crop production, where 15 anthropogenic impacts on the solid soil phases are yet partially buffered and masked by the low 16 reactive OM stocks.

During the first decades of the experiment, decreasing OM contents and rationally the associated biological activity, rapidly engendered soil acidification, loss of exchangeable charges and divalent alkaline earth cations. A diminishing OM-buffering capacity facilitated an increasing control of diverging physicochemical conditions on soil characteristics and properties: fertilization led to growing contrasting exchangeable cation compositions, promoting the acceleration of soil-forming processes.

Thus, Na/K-based fertilizers enhanced clay eluviation from the surface to deeper E and E/Bt horizons, amplifying the Luvisol texture contrast, causing temporal anaerobic conditions and water logging, evolving to a Stagnic Luvisol and on the long-term, to a Retisol. The NH<sub>4</sub>-based fertilizers

1 promoted strong soil acidification (current pH ≈3.5), a total loss of exchangeable calcium and quasi-2 complete saturation of the soil's exchange complex by Al. X-ray-diffraction attested for weathering in 3 fine silt fractions of ferromagnesian minerals (chlorite, hornblende) and geochemical analyses for 4 weathering of Ca-rich plagioclase. Particularly under inputs of di-ammonium phosphate, acidification 5 caused a quasi-total dissolution of ferromagnesian minerals, the phosphate anion likely playing the 6 role of free Al-complexing organic acids in the acidocomplexolysis process in Podzols. In this view, 85 7 years of fertilization with (NH<sub>4</sub>)<sub>2</sub>HPO<sub>4</sub> promoted the development of a podsolization-like process in 8 the surface horizon. Moreover, acidification also affected subsurface horizons, sometimes as deep as 9 the carbonated loess parent material. By contrast, basic liming treatments counteracted processes of 10 acidification and clay translocation, and apparently stopped soil development.

11 In plots without fertilizer inputs, initially defined as 'reference plots', a steady process of 12 alteration developed under temperate climatic conditions of northern France. The current state of 13 these reference plots neither reflect the initial soil conditions at the start of the experiment, nor 14 those of soils under intensive agricultural use. Nevertheless, they were very useful for the study of 15 organic matter decay rates and speeds, the modelling the persistent organic carbon pool, or 16 estimating total amounts of atmospheric trace metal deposition in the Paris suburban region since 17 the 1930s. Effects of climatic acidification affected mainly the surface horizon of the reference plot 18 soil, with  $\approx$  20-25 % of the exchange complex occupied by Al. However, although suspected by 19 geochemical analyses, clear evidence for weathering of primary ferromagnesian minerals and Ca-rich 20 feldspar was not supported by XRD analyses.

21 Conclusively, one might consider INRAE's 42-plots LTBF experiment to perform as an 'alert 22 launcher' tool, forecasting risks of soil degradation in OM-depleted soils under intensive agriculture 23 land use. With reverse reasoning, the results of this experiment strongly advocate for preserving and 24 restore organic matter stocks in soils, in terms of quantity, quality and dynamics so to limit risks of

soil degradation and to guarantee the sustainability of invaluable ecosystem services delivered by
 soils to humankind.

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- 15

## 1 FIGURE CAPTION

Fig. 1. Left, an overview of the 42-plots experiment; middle, a sample collected on July 22, 1929 from plot n°15;
 right, view of archived samples (photos Sébastien Breuil).

4 Fig. 2. Temporal evolution in surface horizons of organic carbon (OC) contents (a) and pH (c) from 1929 to 2014 5 and related soil-depth concentration profiles in 2015 (b,d). Fertilization treatments: ammonium sulphate (P2), 6 di-ammonium phosphate (P14), calcium carbonate (P39), basic slag (P35), sodium nitrate (P4), sylvinite (P29), 7 potassium chloride (P23), ref: means of reference plots P9 and P34 with maximum/minimum value bars. 8 Fig. 3. Temporal evolution in surface horizons of cation exchange capacities (CECCohex) (a) and clay contents 9 (c) from 1929 to 2014 and related soil-depth concentration profiles in 2015 (b,d). Fertilization treatments: 10 ammonium sulphate (P2), di-ammonium phosphate (P14), calcium carbonate (P39), basic slag (P35), sodium 11 nitrate (P4), sylvinite (P29), potassium chloride (P23), ref: means of reference plots P9 and P34 with 12 maximum/minimum value bars. 13 Fig. 4. Temporal evolution in surface horizons of exchangeable cations. Fertilization treatments: ammonium 14 sulphate (P2), di-ammonium phosphate (P14), ref: means of reference plots P9 and P34, calcium carbonate 15 (P39), basic slag (P35), potassium chloride (P23), sylvinite (P29), sodium nitrate (P4). 16 Fig. 5. Soil-depth evolution of exchangeable cation compositions. Fertilization treatments: a) ammonium 17 sulphate (P2), b) di-ammonium phosphate (P14), c) ref: means of reference plots P9 and P34,d) calcium 18 carbonate (P39), e) basic slag (P35), f) potassium chloride (P23), g) sylvinite (P29), h) sodium nitrate (P4). 19 Fig. 6. Optical microscopy photographs of contemporary pedofeatures in subsurface soil horizons under 20 selected fertilization treatments. (a) laminated and layered clay coatings, NaNO<sub>3</sub> plot, 40-50 cm, (b) idem, 70-21 80-cm depth; (c) compound layered dusty clay-silt coatings, NaNO<sub>3</sub> plot, 30-40-cm depth, (d) idem, sylvinite 22 plot, 30-40-cm depth; (e) dusty, poorly sorted clay-silt coatings (matrans, arrows), reference plot, 30-40-cm

23 depth, (f) idem, ammonium sulphate plot 30-40-cm depth; (g) manganese impregnation of pore-walls

24 (hypocoatings), ammonium sulphate plot, 60-70-cm depth, (h) idem, 70-80-cm depth; (i) dendritic

25 manganese nodules (black arrow), di-ammonium phosphate plot, 45-55-cm depth. V: voids. Photos in crossed

26 polarized light (a, b, d), half crossed polarized light (e-h), or plain polarized light (c, i).

1	Fig. 7. X-ray diffraction diagrams of oriented fine silt fractions form 2014 surface horizon samples, after 85-year
2	inputs of di-ammonium phosphate (P14) and ammonium sulphate (P2), no-inputs (reference plot P9), or
3	liming amendments with calcium carbonate (P39). (a) global 2-20- $\mu$ m fraction and fine-silt subfractions 2-5
4	μm (b), 5-10 μm (c), and 10-20 μm (d). V: Vermiculite, Ch: chlorite, M: mica, Am: amphibole, K: kaolinite.
5	Fig. 8. Textural Differentiation Index (TDI) calculated for the eight fertilization treatments. a) at the scale of the
6	solum (horizons with the highest/lowest clay contents); b) surface soil contrast between the surface Ap and
7	underlying E horizon; c) conventional TDI between illuvial/eluvial horizons.
8	Fig. 9. Mineral stability series after Goldich (1938), Jackson and Sherman (1953), setting the resistance to
9	weathering of ferromagnesian amphibole (hornblende) and phyllosilicates (biotite and chlorite) present in
10	loess formations of north-western Europe.
11	Fig. 10. Soil-depth total concentration profiles of major elements Mg, Fe, Al, Ca, Na, and K and of the trace
12	element Scandium (Sc) determined on sub-fine-silt fractions: 2-5 μm (a), 5-10 μm (b), and 10-20 μm (c) for
13	four fertilization treatments ammonium sulphate (P2), di-ammonium phosphate (P14), non-amended
14	reference (P9), and calcium carbonate (P39).
15	

fertilization	composition	number of plots	equivalent input (ha-1.y-1)
	ammonium	8	
nitrogen	nitrate (Na, Ca)	4	150 kg N
	dried blood	2	
	K-based	4	
potassium	K/Na-based (sylvinite)	2	250 kg K <sub>2</sub> O
	phosphate	4	
phosphorus	basic slag	2	200 kg P <sub>2</sub> O <sub>5</sub>
	quick lime	2	
oasic amendments	calcium-carbonate	2	1000 kg CaO
organic matter	horse manure	2	100 t
reference	no-input	10	-

1 Table 1. Details of fertilization design in the 42-plots experiment and equivalent annual input-rates per hectare

plot-n°	designation	fertilization	composition	exch. cations	action	graphic marks
0.24	roforonoo	no input			evolution under local	
9, 34	reference	no-input	-	-	climatic conditions	•
2	acid	ammonium sulphate	(NH4)2SO4	Al³+ (← H+)	acidification, mineral	•
14		di-ammonium phosphate	(NH4)2HPO4		weathering	•
23		potassium chloride	KCI	K⁺	clay dispersion and	
29	monovalent	sylvinite	(K,Na)Cl	K⁺, Na⁺	leaching, aggregate	
4		sodium nitrate	NaNO <sub>3</sub>	Na⁺	breakdown, densification	$\bigcirc$
35	haaia	basic slag	PO4, CaO	Ca <sup>2+</sup>	liming, increase of pH and	
39	basic	calcium carbonate	CaCO₃		aggregate stability	0

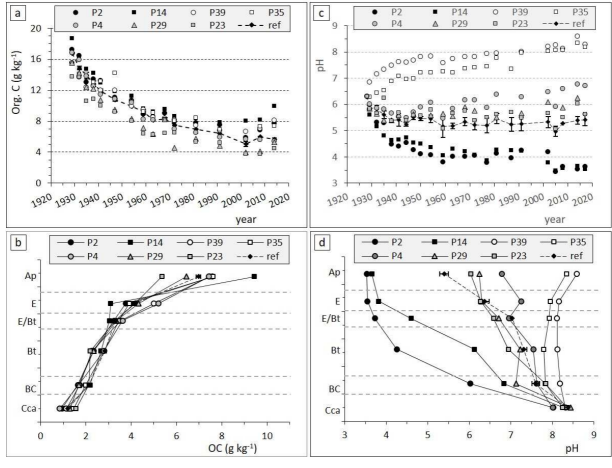
Table 2. Identification, composition and main effects of selected treatments, legend marks in graphs.

- 1 Table 3. Results of ANOVA for the effect of treatment on soil properties. Significance of the effect of factors is indicated as
- 2 follows: \* significant at a P-value of 0.05; \*\* significant at a P-value of 0.01; \*\*\* significant at a P-value of 0.001. Different
- 3 letters within each column indicate statistically significant differences between treatments in the Tukey test at p<0.05.
- 4

		pH	CEC	OC (1972-2014)	Clay
Treatment	F	129.9	16.2	6.9	5.0
	Р	<0.001***	<0.001***	<0.001***	<0.001***
no-input		b	bc	ab	С
ammonium sulphate		а	ab	bc	bc
di-ammonium phosphate		а	а	С	cd
potassium chloride		bc	а	а	ac
sylvinite		bc	а	а	ab
sodium nitrate		с	ab	ab	а
basic slag		d	d	bc	ac
calcium carbonate		d	cd	bc	ас







2 Fig. 2.

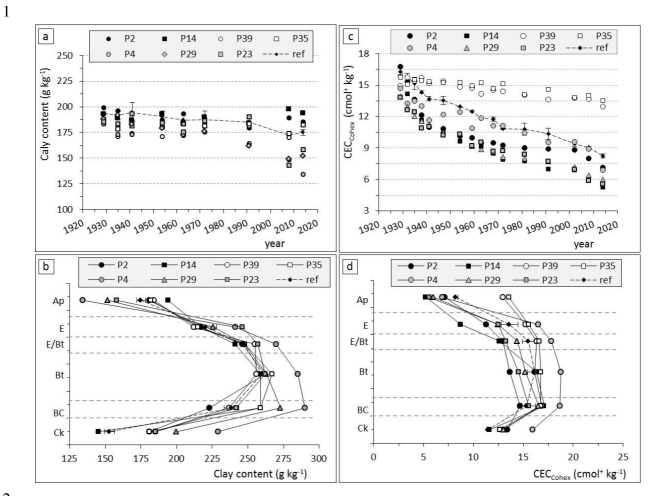
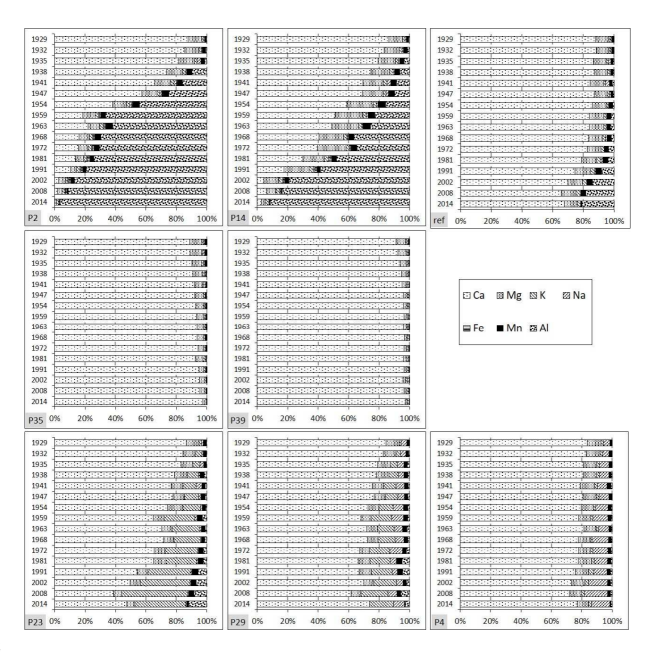




Fig. 3.

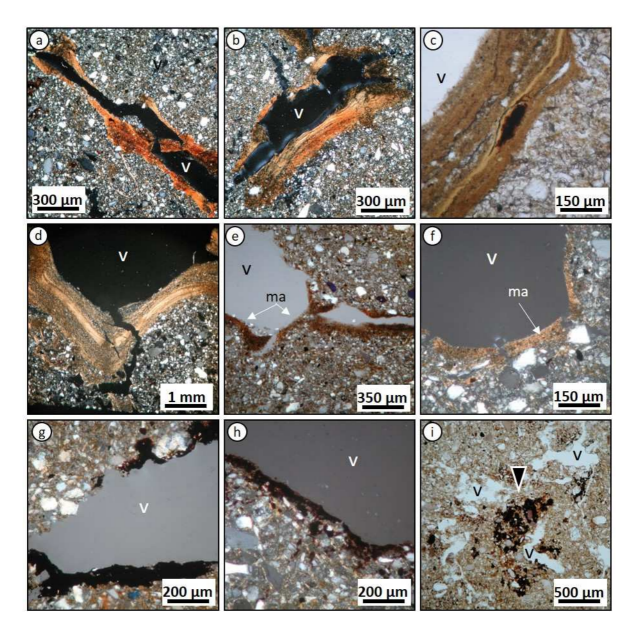




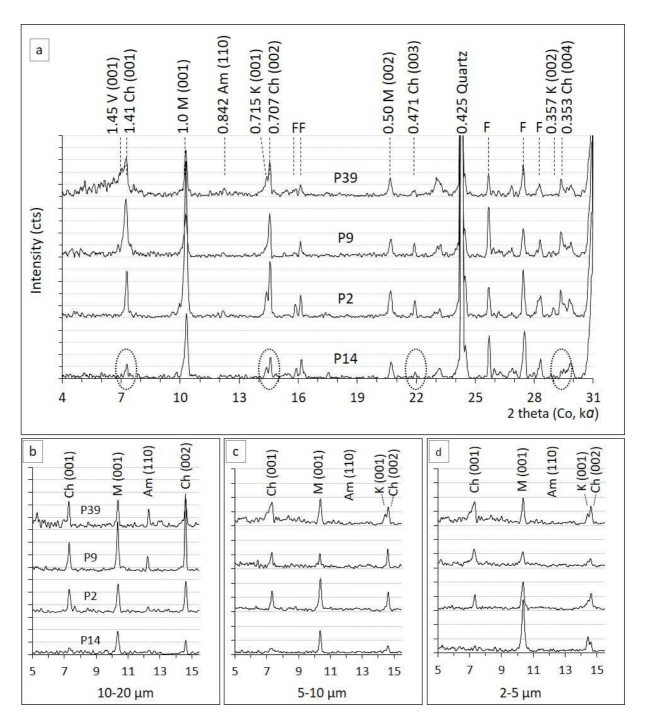
a ( P2	0% 20% 40% 60% 80% 100%	b (	0% 20% 40% 60% 80% 100%	с ( ref	0% 20% 40% 60% 80% 100%
Ap		Ар		Ар	
E		E		E	
E/Bt		E/Bt		E/Bt	
Bt1		Bt1		Bt1	
Bt2		Bt2		Bt2	
BC		BC		BC	
Ck		Ck		Ck	
Second State	0% 20% 40% 60% 80% 100%		0% 20% 40% 60% 80% 100%		
<b>P35</b> Ap		<b>P39</b> Ap			
E		E		<u> </u>	
E/Bt		E/Bt			Ca 🖩 Mg 🖾 K 🖾 Na
Bt1		Bt1			Fe ∎Mn 🖾 Al
Bt2		Bt2			
BC		BC			
Ck		Ck			
20 10 N	0% 20% 40% 60% 80% 100%	1.5	0% 20% 40% 60% 80% 100%		0% 20% 40% 60% 80% 100%
<b>P23</b> Ap		<b>P29</b> Ap		P4 Ap	
E		E		E	
E/Bt		E/Bt		E/Bt	
Bt1		Bt1		Bt1	
Bt2		Bt2		Bt2	
BC		BC		BC	
Ck		Ck		Ck	

3 4

Fig. 5.



2 



3 Fig. 7.

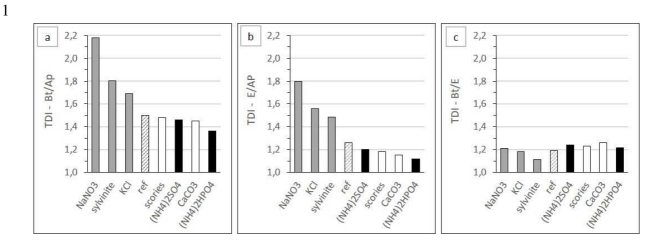


Fig. 8.

