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EXTENDED Halogen enrichment on the continental surface: a perspective from loess

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OPEN_C **ACCESS** Abstract <https://doi.org/10.7185/geochemlet.2442>

Halogen (F, Cl, Br, and I) concentrations for 129 loess samples from worldwide localities yield geometric means of 517 ± 53 μg/g F, 150 ± 20 μg/g Cl, 1.58 ± 0.16 μg/g Br, 1.16 ± 0.11 μg/g I (2 standard errors). These concentrations, notably for Br and I, are substantially higher than previous estimates for the average upper continental crystalline bedrocks, with enrichment factors of $1.3^{+0.7}_{-0.4}$ (F), $1.8^{+2.4}_{-0.8}$ (Cl), $3.8^{+1.3}_{-1.0}$ (Br), and 39^{+71}_{-16} (I) (95 % confidence), documenting enrichment of halogens on the continental surface. These surface halogens are likely sourced from the oceans and may be influenced by climate fluctuations. Halogen ratios (Br/Cl, I/Cl, and Br/I) in loess are similar to those of organic-rich soils/sediments from both terrigenous and marine settings,

La Ce Pr Nd Sm Eu Gd Tb Dy Ho Er Tm Yb Lu F Cl Br I suggesting that terrigenous and marine organic matter have indistinguishable halogen ratios. The Br/I ratios differ from those in the fine grained matrix of glacial diamictites, indicating that another process (beyond biological influence) is responsible for fractionating halogens in the upper continental crust. Using a mixing model, we calculate that over 80–90 % of loess originates from crystalline bedrocks, while the remainder (<10–20 %) derives from the halogen- and organic-rich sedimentary cover or other sources (e.g., marine aerosols).

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Introduction

Halogens (F, Cl, Br, and I), an important group of lithophile and volatile elements, are primarily concentrated in Earth's surface reservoirs (e.g., crust, seawater, and sediments) and play a critical role in modulating Earth's habitability ([Kendrick, 2024](#page-4-0) and references therein). To quantify the halogen cycle operating on Earth, it is necessary to have an accurate estimate of halogen distributions in different terrestrial reservoirs. However, such estimates are still lacking for much of the continental crust. This lack of knowledge regarding the geochemical behaviour of halogens during terrestrial geological processes, such as chemical weathering, metamorphic dehydration, and geobiological activity, limits our understanding of the role of the continental crust in the global halogen cycle, and also hinders our ability to utilise halogen concentrations and elemental ratios to trace geologic processes ([Hanley and Koga, 2018\)](#page-4-0).

To date, there are only five independent estimates of halogen concentrations in the upper continental crust (UCC), and these estimates can differ by up to a factor of∼50 ([Shaw](#page-5-0) et al., [1967](#page-5-0); [Wedepohl, 1995;](#page-5-0) Gao et al.[, 1998;](#page-4-0) [Muramatsu and](#page-4-0) [Wedepohl, 1998;](#page-4-0) Han et al.[, 2023](#page-4-0)). Poorly constrained halogen concentrations in continental rocks are partly due to heterogeneous halogen distributions, and partly due to the analytical challenges of obtaining precise and accurate halogen concentrations at low abundances, especially for Br and I. With recent advances in analytical techniques, Han et al. [\(2023\)](#page-4-0) reported

a comprehensive set of high quality F-Cl-Br-I data for 24 composite samples of the fine grained matrix of ancient glacial diamictites, considered a proxy for upper continental crust ([Gaschnig](#page-4-0) et al., 2016), and used these data to derive halogen estimates for the crystalline and weathered UCC. However, because the diamictites record chemical weathering in their provenance, the new estimates may represent minima for the crystalline UCC (Han et al.[, 2023\)](#page-4-0). Moreover, significant fractionation of Br/Cl and I/Cl ratios is observed in the glacial diamictites relative to pelagic sediments, and the reasons for this are unknown. These differences may reflect organic matter enrichment in the sea floor sediments relative to the UCC, chemical weathering of the UCC, or the effects of diagenesis/metamorphism on the diamictites (Han et al.[, 2023](#page-4-0)). To better constrain the average halogen composition of the continental surface and to explore the mechanism(s) responsible for the fractionation of halogen ratios in sediments, we analysed 129 loess samples from worldwide localities.

Loess is a silt-sized, terrestrial aeolian sediment that covers about 10 % of the continental surface ([Pye, 1995\)](#page-4-0). Previous studies have used loess as a proxy for the average composition of the UCC because of its wide scale sampling of the continental surface, limited chemical weathering (compared to shales) and minor mineralogical and elemental fractionation during sedi-mentary transportation and deposition (e.g., [Taylor](#page-5-0) et al., 1983; Gallet et al.[, 1998;](#page-4-0) [Chauvel](#page-4-0) et al., 2014; [Sauzéat](#page-4-0) et al., 2015). However, previous research on loess mainly focused on

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insoluble elements like rare earth elements, with little attention paid to halogens. Only two previous studies have reported halogen concentrations in loess, each focusing on just one or two halogens (F and Cl by Liu et al.[, 1981](#page-4-0); I by Fan [et al.](#page-4-0), [2021](#page-4-0)) in loess-palaeosol profiles to track palaeoclimate. Thus, there have been no systematic studies of the behaviour of halogens in loess. Here, we report F-Cl-Br-I concentrations, along with major and rare earth element abundances, for 129 loess samples from Germany, Switzerland, the United States, Argentina, Kazakhstan, and China [\(Table S-1](https://www.geochemicalperspectivesletters.org/article2442/#Supplementary-Information) and [Fig. S-1](https://www.geochemicalperspectivesletters.org/article2442/#Supplementary-Information)), and discuss the behaviour of halogens on the continental surface and implications for global halogen recycling.

Halogen Concentrations and Mineralogical Hosts in Loess

Halogen concentrations were analysed by two methods: (1) combustion ion chromatography analysis (C-IC) for F, and (2) $NH₄HF₂$ digestion and ICPMS analysis (N-ICPMS) for Cl, Br, and I ([Supplementary Information](https://www.geochemicalperspectivesletters.org/article2442/#Supplementary-Information)). Halogen concentrations of loess are reported in [Table S-1](https://www.geochemicalperspectivesletters.org/article2442/#Supplementary-Information) and plotted in Figure 1. Each halogen has distinctive geochemical properties, making each likely to be controlled by different phases in loess. Loess is mainly composed of quartz, feldspar, and various amounts of phyllosilicates and carbonates [\(Pye, 1995](#page-4-0)). Fluorine, with the smallest anionic radius of the group, may readily replace the hydroxyl anion and be incorporated into phyllosilicates. This is supported by a good correlation between \overline{F} and \overline{Al}_2O_3

 $(R^2 = 0.84)$ and K_2O $(R^2 = 0.78)$ in loess, the latter two elements being enriched in phyllosilicates [\(Fig. 2a,b](#page-2-0)). Another mineral in loess capable of hosting F is carbonate, as two F[−] can substitute for CO_3^{2-} (Feng *et al.,* 2021). However, F and CaO show a slight negative correlation [\(Fig. S-3\)](https://www.geochemicalperspectivesletters.org/article2442/#Supplementary-Information), suggesting that F in loess is mainly controlled by phyllosilicates, rather than carbonates. The anti-correlation between F and CaO contrasts with the relatively high F content observed in some marine carbonates ([Rude and Aller, 1991\)](#page-4-0). The reason for a lack of carbonate control on F in the loess is unclear but may indicate that F substitution within carbonate occurs only in certain geological environments, which, in turn, might relate to the salinity, $pCO₂$, and/or temperature of the water body from which the carbonates precipitate (e.g., [Rude and Aller, 1991](#page-4-0); [Ramos](#page-4-0) et al., 2005; Feng et al.[, 2021\)](#page-4-0).

The increasing anionic radii of Cl, Br, and I compared to F make them less likely to be incorporated into the crystal lattices of silicate minerals, particularly for Br and I. Instead, they may be absorbed on mineral surfaces or concentrated in fluid inclusions, evaporite minerals, and/or organic matter. We conducted a leaching experiment with MilliQ water on 14 samples to determine the location of these elements in loess ([Table S-1](https://www.geochemicalperspectivesletters.org/article2442/#Supplementary-Information) and [Fig. S-4](https://www.geochemicalperspectivesletters.org/article2442/#Supplementary-Information)). The water soluble component of halogens in loess gradually decreases from Cl (as high as ∼85 %) to Br (∼50 %), to I (∼25 %) ([Fig. S-4](https://www.geochemicalperspectivesletters.org/article2442/#Supplementary-Information)), suggesting that most of the Cl in loess exists in soluble form, while Br and I are mainly present in water-insoluble phases. The soluble halogens in loess (Cl and Br) may originate from a range of sources: (1) soluble inorganic halogens incorporated or absorbed onto the silicate phases

Figure 1 Halogen (F, Cl, Br, and I) concentrations of loess samples. The blue dash-dotted and the red dashed lines represent the geometric and arithmetic means of the loess data set, respectively, along with uncertainties of 2 standard errors (s.e., light-shaded bars; see [Fig. S-5](https://www.geochemicalperspectivesletters.org/article2442/#Supplementary-Information) for more details on the calculation). Symbols above each histogram represent previous estimates of halogen abundances in the UCC.

Figure 2 Fluorine versus (a) Al_2O_3 and (b) K_2O contents of loess. (c) Normalised rare earth element (REE) and halogen concentrations of loess samples relative to the composition of the crystalline UCC (UCC $_{X|t}$): REE estimates for UCC are from [Rudnick and](#page-4-0) [Gao \(2003\)](#page-4-0) and halogens are from Han et al. [\(2023\)](#page-4-0). The cyanshaded area shows the compositional field of glacial diamictite composites.

within loess source materials, (2) soluble halogens in organic matter, (3) evaporite minerals like halite, and (4) post-depositional contamination from either anthropogenic sources and/or meteoric/ground water. The first three sources may reflect halogen distributions on the continental surface when loess deposits formed, while the fourth suggests potential disturbances of halogen contents after deposition. Since no significant regional differences are observed in the Cl and Br contents of samples from different areas ([Fig. 1](#page-1-0)), and considering that their concentrations in meteoric water and groundwater are typically 10–100 times lower than those of loess [\(Worden, 2018](#page-5-0)), any enrichment from localised post-depositional processes, if they occurred, is likely minimal.

The concentrations of residual Br and I (after leaching) in loess are significantly higher than their averages in glacial diamictites ([Fig. S-4](https://www.geochemicalperspectivesletters.org/article2442/#Supplementary-Information)). Because both elements have a strongly biophilic affinity, they may primarily be hosted in organic matter. Loess contains a significantly greater amount of organic matter than crystalline bedrocks, which are the main source of glacial diamictites (Han et al.[, 2023](#page-4-0)). A possible reason for this difference might be the colder environment prevailing during the formation of glacial diamictites, leading to lower biomass on land surfaces and consequently less enrichment of halogens in the diamictites. This explanation also aligns with the proposition put forward by Fan et al. [\(2021\)](#page-4-0) regarding climate fluctuations and I content in loess: in warmer periods, more halogens will be transported from the ocean to land and retained by surface biota, while during colder periods, both halogen transport efficiency and retention on land will decrease. In other words, halogen concentrations in terrigenous sediments, particularly for the less soluble Br and I, may be used as a proxy to monitor past temperature changes and to trace patterns of halogen movement between land and sea.

As a whole, the geometric and arithmetic means calculated from the data set are systematically higher than the recent estimates for crystalline UCC derived from data for ancient glacial diamictite composites (Han et al.[, 2023](#page-4-0)). Using a bootstrap approach, we estimate the degree of halogen enrichment in loess

relative to crystalline UCC to be $1.3^{+0.7}_{-0.4}$ times for F, $1.8^{+2.4}_{-0.8}$ times for Cl, $3.8^{+1.3}_{-1.0}$ times for Br, and 39^{+71}_{-16} times for I (95 % confidence, see [Fig. S-6\)](https://www.geochemicalperspectivesletters.org/article2442/#Supplementary-Information), with an exponential increase in enrichment from F to I (i.e. $F < Cl < Br << I$; Fig. 2c and [Fig. S-7\)](https://www.geochemicalperspectivesletters.org/article2442/#Supplementary-Information). There are two non-mutually exclusive factors that could explain the higher halogen contents in loess relative to glacial diamictites: (1) a lower degree of chemical weathering in the provenance of loess compared to that of glacial diamictites, and (2) greater potential for incorporation and preservation of organic matter in loess than in glacial diamictites ([Fig. S-8\)](https://www.geochemicalperspectivesletters.org/article2442/#Supplementary-Information). On the other hand, the mean halogen concentrations in loess are comparable to previous estimates of halogens in the UCC based on either large scale sampling (Shaw et al.[, 1967;](#page-5-0) Gao et al.[, 1998](#page-4-0)) or models based on lithologies ([Wedepohl, 1995](#page-5-0); [Muramatsu and](#page-4-0) [Wedepohl, 1998\)](#page-4-0) [\(Fig. 1\)](#page-1-0), both of which incorporate data not only for crystalline bedrocks, but also for many halogen-rich lithologies such as shales, carbonates, and evaporites from the overlying sedimentary cover. The similarity between mean halogen concentrations in loess and these previous estimates suggests that halogens in loess may provide a useful average of halogen concentrations across various types of rocks and sediments at the continental surface. Halogen enrichment on the continental surface may derive from the oceans, a major halogen reservoir [\(Hanley and Koga, 2018\)](#page-4-0), with halogen fluxes from oceans to continents potentially regulated by climate fluctuations, as suggested by Fan et al. [\(2021\)](#page-4-0).

Halogen Ratios in Loess

The Br/Cl and I/Cl ratios of loess, as well as the leached residues of loess samples, are significantly fractionated relative to those in igneous rocks like MORB/OIB and arc/backarc lavas, closely tracking the compositional range seen in pelagic sediments, whose halogen concentrations are primarily controlled by marine organic matter (John et al.[, 2011](#page-4-0)) ([Fig. 3\)](#page-3-0). This supports the inference above, based on leaching experiments [\(Fig. S-4\)](https://www.geochemicalperspectivesletters.org/article2442/#Supplementary-Information), that Br and I primarily reside in water-insoluble organic matter in loess. It also suggests that both terrigenous and marine organic matter have indistinguishable halogen ratios. This notion is further supported by the consistently narrow range of the Br/I ratio observed in organic-rich soils and sediments from Japan [\(Yamasaki](#page-5-0) et al., 2015) and China (He et al.[, 2018](#page-4-0)) ([Fig. S-9\)](https://www.geochemicalperspectivesletters.org/article2442/#Supplementary-Information). However, no correlation is observed between Br (or I) and TOC in either the loess or Japanese soil samples [\(Fig. S-10\)](https://www.geochemicalperspectivesletters.org/article2442/#Supplementary-Information), which is somewhat counterintuitive and may result from varying Br/TOC (or I/TOC) ratios in different types of organic matter ([Muramatsu and Wedepohl, 1998;](#page-4-0) [Mayer](#page-4-0) et al., 2007). Therefore, great caution should be exercised when using the correlation of Br and I with TOC to infer the presence of organohalogens in sediments.

The halogen ratios observed in loess are also similar to those found in blueschists and serpentinites from subduction zone settings [\(Fig. 3\)](#page-3-0), indicating that the halogens in these metamorphic rocks may be overprinted by fluids derived from subducted terrigenous organic-bearing sediments, not just marine sediments (e.g., John et al.[, 2011;](#page-4-0) [Kendrick](#page-4-0) et al., 2013; Pagé et al.[, 2016\)](#page-4-0). Most glacial diamictite composites (excluding five carbonate and terrigenous sediment-rich samples from Namibia), however, have significantly higher Br/I ratios (7–74) compared to loess (0.25–6.3) ([Fig. 3](#page-3-0)). This difference suggests that halogen ratios are significantly fractionated within the upper continental crust by an additional process, such as continental weathering and/or diagenesis/metamorphism. We note that the diamictites generally carry a stronger chemical weathering signature than loess ([Fig. S-8](https://www.geochemicalperspectivesletters.org/article2442/#Supplementary-Information)), and most experienced greenschist facies metamorphism (Han et al.[, 2023\)](#page-4-0).

Figure 3 Br/Cl versus I/Cl ratios of loess (symbols as in [Fig. 2](#page-2-0); this study) and glacial diamictite composites (cyan diamonds; Han [et al.](#page-4-0), [2023\)](#page-4-0). Note that (1) the leached loess residues are plotted as deepred dashed symbols, and (2) five carbonate-rich and terrigenous sediment-rich glacial diamictites from Namibia have notably elevated I content and relatively low Br/I ratios and are shown with open diamonds (see Han [et al.,](#page-4-0) 2023 for discussion). Halogen ratios of seawater (grey triangle), bulk silicate Earth (BSE, grey plus sign), mid-ocean ridge and ocean island basalts (MORB/OIB, grey rectangle, [Kendrick](#page-4-0) et al., 2017), arc/backarc lavas (grey outline, [Kendrick](#page-4-0) et al.[, 2020\)](#page-4-0), pelagic sediments (grey dashed outline, John [et al.](#page-4-0), [2011\)](#page-4-0), and marine pore fluids (green field, Fehn et al.[, 2006;](#page-4-0) [Muramatsu](#page-4-0) et al., 2007) are plotted for comparison. Thin grey dashed lines are constant Br/I ratio contours, ranging from 0.01 to 10,000. The inset shows samples from the subduction zone setting: pelagic sediments, marine pore fluids, serpentinites (John [et al.](#page-4-0), [2011;](#page-4-0) [Kendrick](#page-4-0) et al., 2013), and blueschists (Pagé et al.[, 2016\)](#page-4-0).

Implications for Loess Provenance

Previous studies suggested that loess derives from rocks that have generally experienced moderate chemical weathering during multiple cycles of sedimentation (Gallet et al.[, 1998;](#page-4-0) [Sauzéat](#page-4-0) et al., 2015). Based on the Li isotopic composition of loess, [Sauzéat](#page-4-0) et al. (2015) estimated that the proportion of chemically weathered material at Earth's surface was 37^{+17}_{-10} %. However, because Li isotopes are not sensitive to the presence of organic matter, it remains unclear what proportion of material within loess is derived from pulverised unweathered crystalline bedrocks versus mature sediments that have extensively interacted with the biosphere. As Br and I are strongly enriched in organic-bearing sediments and their concentrations will be sensitive to the amount of such sediment in the loess provenance; we used a mixing model to calculate the proportions of the above sources in loess. In this calculation, the Br and I contents in loess are assumed to result from a mixture of (1) clastic sediments derived from glacially pulverised unweathered crystalline bedrocks, (2) an overlying halogen-rich sedimentary cover, and (3) inputs from other sources, such as precipitation (marine aerosols) and evaporite minerals (Fig. 4). Although the composition of the third end member is difficult to define and may vary between samples and locations, we assume that the first end member is well represented by the glacial diamictites [\(Han](#page-4-0) et al., [2023\)](#page-4-0), and the second by the Japanese soil samples ([Yamasaki](#page-5-0) et al.[, 2015\)](#page-5-0). When projecting the geometric or arithmetic mean of Japanese soils ([Fig. S-11\)](https://www.geochemicalperspectivesletters.org/article2442/#Supplementary-Information) onto the mixing line for either Br or I, assuming no other inputs, the calculated mass proportions of the UCC crystalline bedrocks in loess are consistently >80–90 % (Fig. 4). This proportion would be even higher if other inputs were considered. Overall, this suggests that the sediments that have extensively interacted with the biosphere in the loess provenance is <10–20 %, which is somewhat lower than the sum proportion of chemically weathered components, 37^{+17}_{-10} %, at

Figure 4 (a) Mixing model for the source of halogens in loess. In the equations, C_{Xlt} , C_{Hrl} , and C_{Loess} represent halogen concentrations of the UCC crystalline bedrocks, halogen-rich sedimentary layer, and loess, respectively; D_{Other} represents other potential inputs from precipitation and/or evaporites that could contribute significant halogens without substantially affecting the system's mass; m_{Xlt} and m_{Hrl} are assumed mass proportions of the UCC crystalline bedrocks and the sedimentary halogen-rich component in loess. (b) and (c) show the simulated relationship between $m_{X/t}$ and C_{Hrl} from Equation 3 in (a) when $D_{Other} = 0$, for mixing of Br and I, respectively. The red and blue curves are calculated using either the geometric (blue) or arithmetic (red) means of the loess samples and the adjacent pale red/blue circles represent related errors (modelled from Monte Carlo re-sampling; 10,000 times). The vertical dashed lines are geometric (in blue, 2 s.e.) and arithmetic (in red, 2 s.e.) means of Japanese soils. The calculated mass proportion of crystalline rocks in loess is shown as horizontal dashed lines (with 2σ uncertainties), which are obtained when projecting the geometric or arithmetic mean of Japanese soils [\(Fig. S-10](https://www.geochemicalperspectivesletters.org/article2442/#Supplementary-Information)) to the red or blue curves.

the continental surface as estimated by Sauzéat et al. (2015). Thus, it can be concluded that the primary source of loess material is the crystalline bedrocks of the UCC, which validates its use to infer the average composition of the present day UCC [\(Taylor](#page-5-0) et al., 1983; Gallet et al., 1998; Chauvel et al., 2014).

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Additional Information

Supplementary Information accompanies this letter at [https://](https://www.geochemicalperspectivesletters.org/article2442) [www.geochemicalperspectivesletters.org/article2442.](https://www.geochemicalperspectivesletters.org/article2442)

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Halogen enrichment on the continental surface: a perspective from loess

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Supplementary Information

The Supplementary Information includes:

- ➢ Samples and Analytical Methods
- ➢ Tables S-1
- \triangleright Figures S-1 to S-11
- ➢ Supplementary Information References

Samples and Analytical Methods

The 129 loess samples investigated in this study were collected from four continents: Europe (Germany and Switzerland), North America (United States), South America (Argentina), and Asia (Kazakhstan and China). The latitude and longitude of the loess sections, along with references to the original investigations of these units, are given in Table S-1 and plotted in Figure S-1. Data for major elements, rare earth elements (REEs), total organic carbon (TOC), and halogens are reported in this study.

Major Elements and REE Analyses

The samples were powdered in an agate mortar mill (RS200, Retsch, Germany) prior to analysis. Major element concentrations were determined by X-ray fluorescence (RIX2100, Japan) on fused glass disks at Northwest University in Xi'an, China. Reference materials BCR-2 (basalt, USGS) and GSR-3 (basalt, Chinese National Standard) indicate that precision and accuracy are better than 5 %. Rare earth elements were analysed by ICP-MS (Agilent 7900) after high-pressure acid digestion of samples in Teflon bombs at the State Key Laboratory of Geological Processes and Mineral Resources (GPMR), China University of Geosciences, Wuhan. About 50 mg of sample powder was weighed into a Teflon bomb, and then 1 ml of concentrated $HNO₃$ and 1 ml of concentrated HF were added. The sealed bomb was heated at 190 °C in the oven for 72 hours. After cooling, the solution was evaporated to dryness at ~120 °C. This was followed by adding 1 ml of concentrated HNO₃ and evaporating to dryness again. The resultant salt was re-dissolved by adding 1 ml of HNO₃, 1 ml of Milli-Q water (18.2 M Ω), and 1 ml of 1µg/ml In internal standard solution. The solution was then resealed and heated in the bomb at 190 °C for ~24 hours. The final solution was diluted to ∼100 g with 2 %

HNO³ for ICP-MS analysis. Total procedure blanks are below detection limits. The results for reference materials (BCR-2, BHVO-2, AGV-2, and SCo-1) agree well with the recommended values, within 10 % uncertainties. Results for the reference materials are in Park *et al.* (2012).

Total Organic Carbon Analysis

Organic C measurements were conducted at the State Key Laboratory of Biogeology and Environmental Geology, China University of Geosciences, Wuhan. About 2 g of powdered sample was acidified with 50 % HCl to remove inorganic carbon from carbonates. The residue was rinsed with deionised water to neutralise pH, then centrifuged and dried for 48 hours, before being analysed using the 902 T C–S analyser (Beijing Wanliandaxinke Instruments Co. Ltd.). Based on multiple analyses of AR-4007 (total carbon: 7.62 %, Alpha Resources Inc.), AR-4017 (total carbon: 0.50 ± 0.03 %, Alpha Resources Inc.) and B2152 (total carbon: 1.30 ± 0.09 %, Elemental Microanalysis Ltd.), TOC data are reported with a precision better than \pm 0.1 %.

Halogen Analyses and Leaching Experiment

Halogen concentrations of loess are obtained using two different methods: (1) F-Cl-Br analysis by combustion ion chromatography analysis (C-IC) conducted at Fachbereich Geowissenschaften, Universität Tübingen, and (2) Cl-Br-I analysis by NH4HF² digestion and ICPMS analysis (N-ICPMS) conducted at State Key Laboratory of Geological Processes and Mineral Resources, China University of Geosciences, Wuhan. More detailed descriptions of these two methods can be found in Han *et al*. (2023). Although Cl and Br are both analysed in the above two methods, the measured Cl and Br content in loess by C-IC are often 25-50 % lower than those obtained by N-ICPMS, which is consistent with the results obtained by Han *et al*. (2023) on glacial diamictites (Fig. S-2). This discrepancy could result from incomplete halogen recovery by pyrohydrolysis (*e.g.*, Marks *et al*., 2017; Kendrick *et al*., 2018; Han *et al*., 2023). Therefore, we used the F data from C-IC and the Cl, Br, I data from N-ICPMS in this paper to investigate the behaviour of halogens in loess (Table S-1 and Fig. 1).

Halogen leaching experiments were performed on 14 loess samples from various localities (five from Germany, one from the United States, one from Kazakhstan, and seven from China) at State Key Laboratory of Geological Processes and Mineral Resources, China University of Geosciences, Wuhan. For each sample, ~150 mg of sample powder was first mixed with 5 ml Milli-Q water (18.2 M Ω), then sonicated for 5 minutes using ultrasound, and subsequently centrifuged for 7 minutes in a centrifuge. The solid residue was then separated and dried in an oven at 50 ºC for 48 hours. Finally, the residual sample powders were dissolved and analysed for Cl-Br-I content using the above N-ICPMS method.

Supplementary Table

Table S-1 Major elements, total organic carbon (TOC), and rare earth elements of loess samples.

Table S-1 is available for download (.xlsx) from the online version of this article at [https://doi.org/10.7185/geochemlet.2442.](https://doi.org/10.7185/geochemlet.2442)

Supplementary Figures

Figure S-1 Distribution of loess samples from this study. The number in parentheses indicates the quantity of samples analysed in each section.

Figure S-2 Comparison of Cl **(a-b)** and Br **(c-d)** results for loess (this study) and glacial diamictite composites (Han *et al*., 2023) by C-IC and N-ICPMS, respectively. Error bars represent 1σ.

Figure S-3 (a) Loss of Ignition (LOI) of loess increases as the CaO content increases. The data follow the trends produced by having variable amounts of calcite (red-dashed line) and dolomite (blue dashed line) in the samples, suggesting that the CaO in loess is mainly controlled by carbonate. **(b)** Fluorine versus CaO content. There is a negative trend between F and CaO, likely due to the dilution of F content in loess samples by carbonate. Symbols as Fig. 2.

Figure S-4 Leaching experiments on loess samples for Cl, Br, and I. **(a-c)** show the Cl, Br, and I contents of the samples (on the x-axis) both before (original) and after (residual) leaching. The blue dashed lines represent the average Cl-Br-I content of glacial diamictites from Han et al. (2023). The inset figure in **(a)** is in log-scale. In **(d-f)**, the proportion of residual halogen components in the analysed loess samples (y-axis) is plotted against the original halogen concentrations (x-axis).

Figure S-5 Bootstrapped geometric **(a-d**, upper panel**)** and arithmetic **(d-g**, lower panel**)** means of loess F-Cl-Br-I data, which are computed using MATLAB, employing 10,000 trials. Each trial represents the geometric or arithmetic mean of a randomly resampled set of the loess data, with replacement allowed. The geometric/arithmetic means for the loess samples (Fig. 1) are determined by averaging the results of the 10,000 bootstrapped trials, shown as vertical black lines (2 standard errors indicated by black dashed lines). The arithmetic means consistently surpass the geometric means due to the skewed distribution of halogen concentrations in loess (Fig. 1).

Figure S-6 Enrichment factor of halogens in loess relative to the estimates of the crystalline UCC (394 \pm 67 µg/g F, 83 ± 24 μ g/g Cl, 0.41 ± 0.04 μ g/g Br, and 0.03 ± 0.01 μ g/g I, with 2 standard deviations, from Han *et al.*, 2023), calculated by bootstrapping, with 10,000 trials. Each trial represents the median of randomly sampled loess compared to a random composition of the crystalline UCC within its 2 standard deviations. The black line represents the median of the 10,000 bootstrapped trials, and the dashed lines are 95 % confidence interval.

Figure S-7 The halogen enrichment factors in loess increase exponentially with their atomic masses. Error bars of halogen enrichment factors are at 95 % confidence level, see Figure S-6 for further details.

Figure S-8 Comparison of CIA value (chemical index of alteration, CIA = molar Al₂O₃ / (Al₂O₃ + CaO^{*}+ K₂O + Na₂O), where CaO^{*} is corrected for carbonate and apatite (Nesbitt and Young, 1982) and the carbonate correction is based on the approach of McLennan (1993), **a** and **c)** and TOC results **(b** and **d)** for loess (this study) and glacial diamictite composites, respectively. The CIA values of glacial diamictites are from Gaschnig *et al.* (2016) for 141 individual glacial diamictite samples, and the TOC data are from Greaney *et al.* (2020) for 24 glacial diamictite composites. The dashed line in each plot represents the median value for each population, and loess samples generally have limited and lower CIA values but higher TOC than glacial diamictites.

Figure S-9 Br versus I content of Japanese soils (Yamasaki *et al.*, 2015), Chinese soils/sediments (He *et al*., 2018), loess (this study), pelagic sediments (John *et al*., 2011), and glacial diamictites (Han et al., 2023). Within the glacial diamictite composites, five samples from Namibia exhibit notably elevated I content and relatively low Br/I ratios (the elevated I content of the Namibian samples is likely due to the presence of carbonate and abundant terrigenous sedimentary rocks in their provenance, see discussion in Han *et al*. (2023)).

Figure S-10 Br and I versus TOC content of loess samples **(a-b**, this study**)** and Japanese soils **(c-d**, Yamasaki *et al.*, 2015**)**. In **(a)** and **(b)**, the correlation of Br and I with TOC of the analysed loess samples is indicated by the black dashed line, both showing very poor correlation (\mathbb{R}^2 value ≤ 0.1). Br/C ratios for organic matter (OM) of terrigenous and marine sources $(0.43*10⁻³$ and $7.7*10⁻³$, respectively, from Mayer *et al.* (2007)) are plotted as grey dash-dotted lines in (a) and (c) for comparison. In b and d, the grey dash-dotted line represents the I/C ratio (1.0*10⁻³) of typical marine plankton (Elderfield and Truesdale, 1980).

Figure S-11 Bromine and I content in various types of Japanese soil samples (Yamasaki *et al.*, 2015). The blue dashdotted and the red dashed lines represent the geometric and arithmetic means of the dataset, respectively, along with 2 standard errors. Their calculations are the same as those for the loess data, see Figure S-5 for further details.

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