



User Name: =

Date and Time: = 2025-10-25

Job Number: = 266245909

Documents (1)

Client/Matter: -None-

Search Terms: Headline(Vulnerability of mineral-associated soil organic carbon to climate across global drylands)

Search Type: bool

Content Type

news

Narrowed by

Source: Nature Climate Change

1. Vulnerability of mineral-associated soil organic carbon to climate across global drylands



Vulnerability of mineral-associated soil organic carbon to climate across global drylands

Nature Climate Change

July 30, 2024

Copyright 2024 The Author(s), under exclusive licence to Springer Nature Limited All Rights Reserved

Section: Pg. 976-982; Vol. 14; No. 9; ISSN: 1758-678X,1758-6798

Length: 4781 words

Byline: fernando.maestregil@kaust.edu.sa

eduardo.moreno@uam.es

cesar.plaza@csic.es

Body

Main

Soils in drylands—the largest set of biomes on the planet—store 646 Pg organic carbon (C), more than all living vegetation on Earth,. This vast soil organic C pool supports essential ecosystem services, including food provision and water and climate regulation for more than 2.5 billion people,. Yet temperature increases and precipitation reductions forecasted for many dryland regions are expected to disrupt the balance of soil organic C, accelerating microbial decomposition, reducing plant C inputs into the soil and resulting in more CO₂ emissions to the atmosphere,.

The sensitivity of organic C in soils (*sensu ref.*) to temperature and precipitation at timescales relevant to climate change mitigation is thought to be controlled largely by interactions with soil minerals, which restrict the accessibility of microbial decomposers by encapsulating and adsorbing organic matter–. Plant-derived materials at early stages of decomposition are the main constituents of the mineral-unprotected, particulate organic C (POC) fraction of soil organic matter. The POC fraction is thus directly affected by changes in plant C inputs into the soil and is more exposed to microbial decomposition than the organic component of the mineral-associated organic C (MAOC) fraction, which has, therefore, a lower turnover rate,. As a result, large-scale meta-analyses and observational studies suggest that POC is more sensitive to changes in climate, and particularly to warming, than is MAOC,–. Because of the typically large ratio of soil minerals to organic matter in drylands, MAOC is expected to dominate over POC, potentially driving a high persistence of soil organic C in these ecosystems,,. However, no studies to date have examined the relationship of POC and MAOC with climate across the diverse environmental gradients that characterize global drylands. Investigating this relationship is particularly timely and relevant as it would

Vulnerability of mineral-associated soil organic carbon to climate across global drylands

substantially reduce the uncertainty surrounding the land carbon–climate feedback. In addition, it would provide valuable insights for adapting soil carbon-related ecosystem services to ongoing climate change.

Here we evaluated how mean annual temperature and precipitation relate to POC and MAOC contents across global drylands after accounting for major biotic (net primary productivity, vegetation type, woody cover, plant and herbivore richness and grazing pressure) and soil biogeochemistry (clay and silt contents, pH, chemical index of alteration, exchangeable Ca, non-crystalline Al and Fe, available N and P and microbial biomass C) factors known to potentially affect soil organic C content by regulating C inputs and stabilization processes,. To do so, we surveyed in situ 326 plots from 98 dryland ecosystems located in 25 countries from 6 continents (Extended Data Fig.). Our survey spans the broad gradients of temperature, precipitation, aridity, soil properties, vegetation types and grazing pressures that can be found across drylands worldwide (Extended Data Tables and),. At each site, we collected topsoil samples (0–7.5 cm) from areas both covered (322) and not covered (326) by perennial vegetation from 2–4 plots located across a local gradient of extensive grazing pressure (648 samples in total;). We subjected all samples to a size fractionation procedure to separate and quantify C content in POC and MAOC pools,. Using these data, we tested the hypothesis that MAOC, being protected by minerals, is less sensitive than POC to increases in temperature and decreases in precipitation,,,. We also hypothesize that the presence of vegetation mitigates declines in soil C, particularly POC, by increasing soil C inputs.

MAOC dominates soil organic C and is sensitive to climate

Our results show that MAOC was the dominant soil organic C fraction in drylands globally (Fig.). In particular, median MAOC content was 5.2 gC kg⁻¹ soil, equivalent to 66% of the total soil organic C content, whereas median POC content was 2.3 gC kg⁻¹ soil. This quantification falls within the range of soil organic C content (MAOC and POC) commonly found in drylands and is relevant to improve the performance of emerging models of soil organic C formation and persistence using POC and MAOC frameworks,–.

Distribution of soil organic C contents in POC and MAOC fractions and their relationships with climate in global drylands.

a, POC and MAOC contents. Box, first and third quartiles; central horizontal line, median; upper vertical line end, largest value smaller than 1.5 times the interquartile range; lower vertical line end, smallest value larger than 1.5 times the interquartile range (n = 326 plots). b,c, Relationships between POC and MAOC contents and mean annual temperature (MAT; b) and mean annual precipitation (MAP; c). Lines and shading represent linear regressions and 95% confidence intervals, respectively. d, Summary of a linear mixed-effects model, controlling for biotic factors and soil biogeochemistry (). The panel shows coefficients (circles) and 95% CI (bars) for main and interaction effects of C fraction type (binary variable, either POC or MAOC) and climate (MAT and MAP) on POC and MAOC contents. The variance explained (R²) by the fixed and random effects relative to the total variance was 77% and 12%, respectively (n = 634 POC and MAOC observations). Carbon fraction contents were natural-logarithm transformed, and all the predictors were standardized. The positive coefficient of C fraction type (MAOC versus POC) indicates that MAOC contents are significantly greater than POC contents (P < 0.001). For the observed negative association of MAT and positive association of MAP with C content (P < 0.001 and P = 0.039, respectively), negative coefficients for the interaction of C fraction type with MAT and MAP indicate that increasing MAT has a stronger negative effect on MAOC than on POC (P = 0.053) contents, while decreasing MAP has a stronger negative effect on POC than on MAOC (P = 0.181).

Contrary to our hypothesis, we found that MAOC and POC were equally sensitive to differences in climate across global drylands. In particular, both MAOC and POC were negatively associated with increasing temperature and decreasing precipitation to a similar extent, as indicated by the similar slopes of the associations (Fig.). These results were supported by the lack of a significant interaction between the effects of temperature and precipitation and the type of fraction (MAOC versus POC) tested by a linear mixed-effects model (Fig. ;). On the basis of the results from this model, we estimated that POC and MAOC contents significantly declined with temperature at an average rate of 3.2% per °C (95% confidence interval (CI): 1.8, 4.6) and increased with precipitation at an average rate of 6.6% per 100 mm (95% CI: 0.6, 12.6).

Vulnerability of mineral-associated soil organic carbon to climate across global drylands

Warming accelerates the microbial decomposition of soil organic matter, and precipitation reduction constrains plant production and organic matter inputs into the soil,. Our results are, therefore, consistent with previously reported reductions in soil organic C content with increasing temperature and reducing precipitation across terrestrial ecosystems—. However, and contrary to expectations of smaller sensitivity of MAOC versus POC to changes in climate observed in more mesic systems,, our findings based on a space-for-time substitution highlight that the MAOC and POC fractions may decrease at similar rates in response to climate warming and precipitation reduction across global drylands. Therefore, they suggest that the current paradigm of mineral protection may not determine soil C persistence in dryland ecosystems,—. The apparent lack of protection by minerals, which contrasts with what was observed in mesic systems richer in organic matter, was consistent across the range of soil organic C content found in drylands (Extended Data Fig.). There is recent evidence that MAOC is controlled not only by C stabilization in soil organo-mineral complexes, but also by changes in C inputs driven by climate. In drylands, not only precipitation reduction but also warming may increase water deficit, which may decrease plant productivity, C inputs into the soil and C accumulation into the MAOC fraction. These is also evidence that dryland soils maintain a high oxidative potential during dry periods, mainly through the stabilization of enzymes, which results in a rapid organic matter decomposition in wet periods, and may further limit C inputs to the MAOC fraction.

Vegetation buffers soil C declines with warming

Both POC and MAOC contents were higher in soil beneath perennial vegetation (Fig.). We further observed that as mean annual temperature increased, POC and MAOC contents decreased, but to a lesser extent, beneath vegetation. Conversely, as mean annual precipitation increased, both contents increased in a similar manner in open areas and in areas under the canopy of perennial vegetation (Fig.). These results are important because they suggest that the presence of vegetation buffers, but does not fully compensate for, the negative effects of higher temperature on soil C fractions. While the buffering effect of vegetation did not completely counteract the vulnerability of organic C pools to increasing temperatures, our findings indicate that management practices aimed at protecting vegetation in drylands may help to maintain soil organic C stocks in global drylands and reduce their losses in response to a changing climate.

Relationships between climate and POC and MAOC contents in soils under the canopy of the dominant perennial vegetation and in open areas across global drylands.

a–d, Relationships between POC and MAT (a) and MAP (c), and between MAOC and MAT (b) and MAP (d) in both open areas (O) and perennial vegetation (V) microsites. Lines and shading represent linear regressions and 95% CIs ($n = 326$ and 322 for O and V, respectively). e, Coefficients (dots) and 95% CIs (bars) of linear mixed-effects model illustrating the fixed main and interaction effects of MAT, MAP and the presence of vegetation cover (V versus O) on POC and MAOC contents ($n = 648$ V and O areas). The variance explained (R^2) by the fixed and random effects relative to the total variance was 30% and 55%, respectively, for POC, and 32% and 61%, respectively, for MAOC.

Coupling of POC and MAOC in drylands

We found that POC and MAOC contents were strongly correlated across global drylands ($r = 0.83$, $n = 326$, $P < 0.001$; Fig.). These results strongly suggest that both fractions remain highly coupled in drylands despite their different levels of putative protection against decomposition by microorganisms.

Coupling and drivers of POC and MAOC in global drylands.

a, Relationship between POC and MAOC contents. Dots represent individual dryland plots, with the colours of the dots illustrating their aridity ($1 - \text{annual precipitation/potential evapotranspiration}$) values. The line and shading represent the fitted linear regression and 95% confidence interval, respectively. b, Variance explained (R^2) by linear mixed-effects models for POC and MAOC contents partitioned into the fraction attributable to unique and shared among groups of drivers (climate: mean annual temperature and mean annual precipitation; biotic factors: net primary productivity, type of vegetation, woody cover, plant richness, grazing pressure and herbivore richness; and soil biogeochemistry: clay and silt, pH, chemical index of alteration, exchangeable Ca, non-crystalline Al and Fe, available N and P and microbial biomass carbon). The variance explained (R^2) by the fixed and random effects

Vulnerability of mineral-associated soil organic carbon to climate across global drylands

relative to the total variance was 69% and 20% for POC ($n = 317$) and 84% and 11% for MAOC ($n = 317$), respectively.

Variance partitioning of linear mixed-effects models and random-forest analysis showed that the order of importance of the group of factors that explained most of the variation of POC and MAOC across global drylands was essentially the same for both organic C fractions (Fig. and Extended Data Fig.). Soil biogeochemistry, above climate and biotic factors, was the most important predictor of both POC and MAOC contents. Both C fractions were negatively associated with soil pH and positively associated with exchangeable Ca, available N and P and microbial biomass C contents; in addition, MAOC was associated positively with clay and silt and non-crystalline Al and Fe contents (Extended Data Fig.). Slightly acidic to neutral soils generally feature higher nutrient availability and more fertility than alkaline soils, which may thus favour soil organic C accumulation in drylands through increased plant-derived C inputs and microbial activity. The prevalent role of soil fine texture and non-crystalline Al and Fe in MAOC formation has been widely documented in the literature. Sorption of organic matter to mineral surfaces is known to be promoted by the relatively high specific surface area and charge of clay and silt, while non-crystalline Fe and Al phases are also known to form strong associations with organic matter.

The coupling of POC and MAOC observed here for drylands may be, however, disrupted in more productive terrestrial ecosystems, where higher plant inputs may result in larger POC contents—. In contrast to experimental manipulation studies, our work addresses the vulnerability of soil C fractions using a space-for-time substitution. Further research into the pace of the climate-induced changes and the causality of the associations found in our study is thus warranted.

Concluding remarks

By using a global standardized field study and by focusing exclusively on dryland ecosystems, our work expands previous efforts to understand abiotic and biotic drivers of POC and MAOC along large geographical gradients, which either have been based on literature syntheses, which use datasets that are inherently heterogeneous, or have focused on ecosystems other than drylands. Our study generated highly standardized field data on the POC and MAOC fractions of dryland soils worldwide, along with their major predictors. These data substantially expand existing global databases and can be used to refine current soil organic C models.

Our findings suggest that ongoing changes in climate, particularly warming, may adversely affect both unprotected and mineral-protected soil C content in drylands to a similar extent. The results obtained also indicate that maintaining vegetation cover can mitigate, but not fully counteract, the negative impacts of rising temperatures on soil organic C fractions. Our study enhances our understanding of how POC and MAOC contents in soil respond to key abiotic and biotic drivers, revealing that mineral protection has limited potential to sustain organic C storage in dryland soils in the face of ongoing global warming. The novel insights provided here about dryland soil C pools and their sensitivity could facilitate much-needed advances in our model representation of dryland ecosystems and their response to climate change.

Methods

Global field survey and soil sampling

Fieldwork was conducted from January 2016 to September 2019. A total of 326 plots distributed across 98 study sites in 25 countries from all continents except Antarctica (Algeria, Argentina, Australia, Botswana, Brazil, Canada, Chile, China, Ecuador, Hungary, Iran, Israel, Kazakhstan, Kenya, Mexico, Mongolia, Namibia, Niger, Palestine, Peru, Portugal, South Africa, Spain, Tunisia and the United States of America) and encompassing the wide range of vegetation, soil, climate and grazing-pressure levels found in drylands worldwide were surveyed using a common and standardized protocol,.

At each site, we gathered field data within multiple 45 m × 45 m plots situated along a gradient of grazing pressure, encompassing high ($n = 98$), medium ($n = 97$) and low ($n = 88$) pressure levels, as well as ungrazed areas ($n = 43$). To establish the grazing gradients, in 90 out of the 98 sites surveyed, we strategically positioned these plots at varying distances from artificial watering points, which are usually created in drylands to supply introduced livestock

Vulnerability of mineral-associated soil organic carbon to climate across global drylands

with permanent water sources. The closer the plot to the permanent water source, the more intense the grazing,. In the remaining eight sites, local variations in grazing-pressure gradients were ascertained by observing different paddocks featuring varying grazing intensities. See ref. for additional details on the characterization and validation of the local grazing-pressure gradients established.

A portable Global Positioning System was used to record the coordinates and elevation of each plot, which were standardized to the World Geodetic System 1984 ellipsoid for visualization and analyses. During the dry season at each site, four soil cores (145 cm³) from 0 to 7.5 cm depth (topsoil) were collected from five 50 × 50 cm quadrats randomly placed in areas under the canopy of the dominant perennial vegetation and five placed in open areas not covered by perennial vegetation. The soil cores were homogenized and composited to form a sample representative of the soil under the dominant vegetation and a sample representative of the soil in open areas within each plot. The soil samples were passed through a 2 mm sieve. A portion of each soil sample was air dried and used for organic matter fractionation and texture and pH analysis, and another portion was stored at -20 °C and used for microbial biomass C analysis. A portion of the air-dried soil samples was ground with a ball mill for additional chemical analysis.

Soil organic carbon fractionation and quantification

All the soil samples, a total of 648 (326 from open areas and 322 from under the canopy of the dominant vegetation), were subjected to a size fractionation method, to separate the POC (not protected by minerals from microbial decomposition) and MAOC (protected by minerals) fractions. Aggregates were dispersed by adding 30 ml of sodium hexametaphosphate (5 g L⁻¹) to 10 g of soil and shaking with an overhead shaker for 18 h. After dispersion, the mixture was thoroughly rinsed through a 53 µm sieve to separate the POC (>53 µm) and MAOC (<53 µm) fractions using an automated wet sieving system. The isolated fractions were oven dried at 60 °C, weighed and ground with a ball mill. The whole soil samples and the POC and MAOC fractions were analysed for organic C contents by dry combustion and gas chromatography using a ThermoFlash 2000 NC Soil Analyzer (Thermo Fisher Scientific) after removing carbonates by acid fumigation.

Climate data

Mean annual temperature and mean annual precipitation data were obtained from WorldClim 2.0 (ref.), a high-resolution (30 arcsec, or ~1 km at the Equator) database based on a large number of climate observations and topographical data for the 1970–2000 period. Aridity index (ratio of average annual precipitation to potential evapotranspiration) data were obtained from the Global Aridity Index and Potential Evapotranspiration Climate Database v.3 (ref.). Aridity was calculated as 1 – aridity index.

Vegetation and herbivore richness survey

Each plot was classified as grassland, shrubland or forest by identifying the dominant type of vegetation. Net primary productivity was estimated using the mean annual Normalized Difference Vegetation Index averaged monthly values between 1999 and 2019 at a resolution of 30 m from Landsat 7 Enhanced Thematic Mapper Plus. The cover of perennial vascular plants (plant cover) was measured along four parallel 45 m transects separated by 10 m and oriented downslope during the peak of the growing season using the line–intercept method,. Woody cover was measured in 25 contiguous quadrats (1.5 m × 1.5 m) placed in each transect (100 quadrats per plot). Plant richness was the total number of unique perennial species found along the quadrats and transects surveyed. The richness of herbivores was quantified at each plot using dung data collected systematically in situ along the four 45 m transects established as described in ref. .

Soil analyses

All the bulk soil samples were analysed as follows. Clay and silt contents were determined by sieving and sedimentation. Soil pH was measured in a water suspension at a soil-to-water ratio of 1.0:2.5 (ref.). The chemical index of alteration, which is an indicator of the degree of weathering, was calculated as the molecular proportion of Al₂O₃ versus Al₂O₃ + CaO + Na₂O + K₂O (ref.), using total Al, Ca, Na and K contents and after correcting Ca for soils with carbonates; total Al, Ca, Na and K contents were determined by inductively coupled plasma atomic

Vulnerability of mineral-associated soil organic carbon to climate across global drylands

emission spectroscopy (ICP-AES) after digestion in nitric and perchloric acids,. Exchangeable Ca content was determined by ICP-AES after extraction with ammonium acetate at pH 7.0 (refs.). Non-crystalline Fe and Al contents were determined by ICP-AES after extraction with acid ammonium oxalate. Available N (ammonium and nitrate) content was determined by extraction with 0.5 M K₂SO₄ and the indophenol blue method using a microplate reader. Available P content was determined by the Olsen method. Microbial biomass C was determined by substrate-induced respiration using an automated microrespirometer.

Statistical analyses

We compared the content of MAOC with that of POC in global dryland soils controlling for confounding factors, and tested the hypothesis that the effects of climate (mean annual temperature and precipitation) on POC and MAOC contents depends on (interacts with) the C fraction type. For these analyses, we aggregated soil data for open and vegetation-covered areas by plot using plant cover area as a weighting factor, and fitted a linear mixed-effects model on the response of C content with C fraction type as a binary categorical predictor (either MAOC or POC). In the fixed-effects term of the model, we also included mean annual temperature, mean annual precipitation and the interactions of mean annual temperature and mean annual precipitation with C fraction type, as well as key biotic (net primary productivity, type of vegetation, woody cover, plant richness, grazing pressure and herbivore richness) and soil biogeochemical (clay and silt, pH, chemical index of alteration, exchangeable Ca, non-crystalline Al and Fe, available N and P and microbial biomass C) covariates to control for confounding factors. In the random term of the model, we incorporated an intercept structure with plot nested within site as a categorical variable to account for the lack of independence in the residuals due to the paired POC and MAOC separation and the plot sampling design. We checked whether the fit of this linear mixed-effects model improved by including quadratic terms of mean annual temperature, mean annual precipitation, and both mean annual temperature and precipitation, using the Akaike information criterion and likelihood ratio tests. None of the quadratic models tested was a significantly better fit to the data ($\chi^2(1) < 1.0$, $P > 0.3$) than the linear model (lowest Akaike information criterion).

To examine separately the variance of POC and MAOC contents explained by the groups of predictors (climate: mean annual temperature and mean annual precipitation; biotic factors: net primary productivity, type of vegetation, woody cover, plant richness, grazing pressure and herbivore richness; soil biogeochemistry: clay and silt, pH, chemical index of alteration, exchangeable Ca, non-crystalline Al and Fe, available N and P and microbial biomass C), we built two linear mixed-effects models (one for POC and another for MAOC) with site as a random categorical variable. These two separate models were used to assess the importance of the different groups of predictors in explaining either POC or MAOC, and not to test statistically for differences in the size of the effects of the predictors between POC and MAOC. To support the linear mixed-effects models, we tested the importance of the same groups of predictors of POC and MAOC using random-forest regression modelling. In particular, we built two random-forest models, one for POC and one for MAOC, combining 500 trees, and quantified the importance of each predictor by computing the increase in mean squared error across trees when the predictor was permuted.

We tested whether the presence of vegetation cover interacted with the effects of temperature and precipitation also by linear mixed-effects modelling. For this purpose, we built two linear mixed-effects models, one for POC content and another for MAOC content in areas under the canopy of the dominant perennial vegetation and open areas, with vegetation cover as a binary predictor and plot nested within site in the random term.

For all the linear mixed-effects models, POC, MAOC, exchangeable Ca, non-crystalline Al and Fe, available N and P and microbial biomass C were natural-logarithm transformed to reduce the skewness of the data. To compare effect sizes, all the numeric predictors were standardized by subtracting the mean and dividing by two standard deviations, and the binary variables (C fraction type and vegetated versus open areas) were rescaled to -0.5 and 0.5 (ref.). The coefficients of the models were estimated by the restricted maximum likelihood approach, 95% CIs were calculated, and P values were computed on the basis of the Satterthwaite approximation. The validity of the assumptions of normality, homoscedasticity and linearity were examined using residual plots. The generalized variance inflation factors were computed to check for multicollinearity among predictors (the values were <3 in all cases, suggesting that multicollinearity was low). All statistical analyses were performed using R version 4.3.0 (ref.) and the R packages arm version 1.13 (ref.), ggplot2 version 3.4.4 (ref.), lme4 version 1.1 (ref.), lmerTest version

3.1 (ref.), partR2 version 0-9-1 (ref.), patchwork version 1.1.3 (ref.), rnatualearth version 0.3.2 (ref.), randomForest version 4.7 (ref.), sf version 1.0 (ref.), terra version 1.7 (ref.) and viridis version 0.6.3 (ref.).

Reporting summary

Further information on research design is available in the [linked to this article](#).

Online content

Any methods, additional references, Nature Portfolio reporting summaries, source data, extended data, supplementary information, acknowledgements, peer review information; details of author contributions and competing interests; and statements of data and code availability are available at <https://doi.org/10.1038/s41558-024-02087-y>.

Acknowledgements

This research was funded by the European Research Council (ERC Grant agreement 647038, BIODESERT), the Spanish Ministry of Science and Innovation (PID2020-116578RB-I00) and Generalitat Valenciana (CIDEAGENT/2018/041), with additional support by the University of Alicante (UADIF22-74 and VIGROB22-350). F.T.M. acknowledges support from the King Abdullah University of Science and Technology (KAUST) and the KAUST Climate and Livability Initiative. D.J.E. is supported by the Hermon Slade Foundation. H.S. is supported by a María Zambrano fellowship funded by the Ministry of Universities and European Union-Next Generation plan. L.W. acknowledges support from the US National Science Foundation (EAR 1554894). B.B. and S.S. were supported by the Taylor Family–Asia Foundation Endowed Chair in Ecology and Conservation Biology. M.B. acknowledges support from a Ramón y Cajal grant from the Spanish Ministry of Science (RYC2021-031797-I). A.L. and L.K. acknowledge support from the German Research Foundation, DFG (grant CRC TRR228) and German Federal Government for Science and Education, BMBF (grants 01LL1802C and 01LC1821A). L.K. acknowledges travel funds from the Hans Merensky Foundation. A.N. and C. Branquinho acknowledge support from FCT—Fundação para a Ciência e a Tecnologia (CEECIND/02453/2018/CP1534/CT0001, PTDC/ASP-SIL/7743/2020, UIDB/00329/2020), from AdaptForGrazing project (PRR-C05-i03-I-000035) and from LTsER Montado platform (LTER_EU_PT_001). S.C.R. was supported by NASA (NNH22OB92A) and is grateful to E. Geiger, A. Howell, R. Reibold, N. Melone and M. Starbuck for field support. Any use of trade, firm or product names is for descriptive purposes only and does not imply endorsement by the US Government. We thank the landowners for granting access to the sites and many people and their institutions for supporting our fieldwork activities: L. Eloff, J. J. Jordaan, E. Mudongo, V. Mokoka, B. Mokhou, T. Maphanga, D. Thompson (SAEON), A. S. K. Frank, R. Matjea, F. Hoffmann, C. Goebel, the University of Limpopo, South African Environmental Observation Network (SAEON), the South African Military and the Scientific Services Kruger National Park.

Notes

Extended datais available for this paper at <https://doi.org/10.1038/s41558-024-02087-y.Supplementary> informationThe online version contains supplementary material available at <https://doi.org/10.1038/s41558-024-02087-y.Publisher's> note Springer Nature remains neutral with regard to jurisdictional claims in published maps and institutional affiliations.

Load-Date: September 6, 2024