

1 **Title:** Elephant megacarcasses increase local nutrient pools in African savanna soils and plants

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25 **Abstract**

26 African elephants (*Loxodonta africana*) are the largest extant terrestrial mammals, with bodies
27 containing enormous quantities of nutrients. Yet we know little about how these nutrients move
28 through the ecosystem after an elephant dies. Here, we investigated the initial effects (1-26
29 months postmortem) of elephant megacarcasses on savanna soil and plant nutrient pools in
30 Kruger National Park, South Africa. We hypothesized that: (H1) elephant megacarcass
31 decomposition would release nutrients into soil, resulting in higher concentrations of soil
32 nitrogen (N), phosphorus (P), and micronutrients near the center of carcass sites; (H2) carbon (C)
33 inputs to the soil would stimulate microbial activity, resulting in increased soil respiration
34 potential near the center of carcass sites; and (H3) carcass-derived nutrients would be absorbed
35 by plants, resulting in higher foliar nutrient concentrations near the center of carcass sites. To test
36 our hypotheses, we identified 10 elephant carcass sites split evenly between nutrient-poor
37 granitic and nutrient-rich basaltic soils. At each site, we ran transects in the four cardinal
38 directions from the center of the carcass site, collecting soil and grass (*Urochloa trichopus*,
39 formerly *U. mosambicensis*) samples at 0, 2.5, 5, 10, and 15 m. We then analyzed samples for
40 CNP and micronutrient concentrations and quantified soil microbial respiration potential. We
41 found that concentrations of soil nitrate, ammonium, $\delta^{15}\text{N}$, phosphate, and sodium were elevated
42 closer to the center of carcass sites (H1). Microbial respiration potentials were positively
43 correlated with soil organic C, and both respiration and organic C decreased with distance from
44 the carcass (H2). Finally, we found evidence that plants were readily absorbing carcass-derived
45 nutrients from the soil, with foliar %N, $\delta^{15}\text{N}$, iron, potassium, magnesium, and sodium
46 significantly elevated closer to the center of carcass sites (H3). Together, these results indicate
47 that elephant megacarcasses release ecologically consequential pulses of nutrients into the soil

48 that influence soil microbial activity and are absorbed by plants into the above-ground nutrient
49 pools. These localized nutrient pulses may drive spatiotemporal heterogeneity in plant diversity,
50 herbivore behavior, and ecosystem processes.

51 **Sect. 1 Introduction**

52 Living animals affect nutrient flows through ecosystems (Schmitz et al. 2018), but we have only
53 recently acknowledged that the nutrients from animal carcasses could also influence ecosystem
54 processes (Barton et al. 2013; Monk et al. 2024). In marine ecosystems, whale carcasses function
55 as unique hotspots of nutrient cycling, biodiversity, and ecosystem processes (Roman et al.
56 2014). In terrestrial systems, mass mortality events (e.g., wildebeest, cicadas) create nutrient
57 hotspots (Yang, 2004; Subalusky et al. 2020), while individual small and medium-sized
58 carcasses release pulses of nutrients into the soil (Town, 2000; Barton et al. 2016; Olea et al.
59 2019). Yet, terrestrial ecosystem ecology lacks knowledge about the role of megacarcasses
60 (carcasses of animals such as elephants and rhinoceros that are >1000 kg at death) as potential
61 drivers of spatiotemporal heterogeneity in nutrient cycling and ecosystem processes. Importantly,
62 these megacarcasses may be functionally different than smaller carcasses due to the
63 extraordinarily high concentration of nutrients and residence time of the decomposing animal
64 (see reviews by Barton et al. 2013; Barton, 2016; Barton & Bump 2019). This question around
65 the role of megacarcasses is particularly relevant given the megaherbivore losses that occurred
66 during the Pleistocene extinctions and that are still occurring today (Ripple et al. 2015). We are
67 only beginning to understand how the ‘extinction aftershock’ of losing the largest species
68 impacts ecosystems (Owen-Smith, 1989; Flannery, 1990), and no study has yet investigated how
69 the loss of megacarcasses might influence the dynamics of terrestrial ecosystems (Doughty et al.
70 2013; Doughty et al. 2016).

71 We can only evaluate the importance of terrestrial megacarcasses for nutrient cycling in
72 ecosystems where megaherbivores still exist, such as African savannas. The African savanna
73 elephant (*Loxodonta africana*) is the largest extant land animal and is known for its key

74 ecological effects in savannas while alive (e.g., dispersing seeds, creating plant refuges,
75 preventing woody encroachment) (Skarpe et al. 2004; Asner et al. 2009; Campos-Arceiz &
76 Blake, 2011; Coverdale et al. 2016; Guy et al. 2021). Yet, the elephant's large body mass may
77 mean that it also has an outsized impact in these ecosystems even after death. A 4000-kg
78 elephant megacarcass likely represents ~2000 kg carbon (C), ~300 kg nitrogen (N), and ~125 kg
79 phosphorus (P) deposited in the savanna landscape (estimated from stoichiometry of elephants
80 and other mammals in Sterner & Elser, 2002). The N deposition from one elephant megacarcass
81 (in a 700 m² impact zone assuming a 15 m disturbance radius) is roughly equivalent to the N
82 delivered to 10,000 m² of savanna from ~100 years from atmospheric deposition (Mphepya et al.
83 2006).

84 If megacarcasses provide large nutrient pulses, then they likely create hotspots of
85 important below- and aboveground processes. Belowground, soil respiration and organic matter
86 decomposition might increase with nutrient inputs from carcasses (Risch et al. 2020).
87 Concentrations of C, N, P, and potassium (K) are often elevated near carcasses of medium-sized
88 animals (e.g., bison, moose, kangaroo, vicuña) (Towne, 2000; Bump et al. 2009a; Macdonald et
89 al. 2014; Risch et al. 2020; Monk et al. 2024), and nutrients such as P and calcium (Ca) continue
90 leaching from bones even after soft tissues have been consumed or degraded (Coe, 1978; Keenan
91 & Beeler, 2023). Aboveground, plant growth in African savannas is limited by nutrient
92 availability, most commonly N and P (Ries & Shugart, 2008; Pellegrini, 2016), and
93 micronutrients such as sodium (Na) and potassium (K) may co-limit plant productivity as well
94 (Epron et al. 2012; Chen et al. 2024). Thus, the large influx of nutrients released from
95 megacarcasses might increase the mobilization of nutrients by plants, potentially increasing
96 nutrient accessibility for vertebrate and invertebrate herbivores (Yang, 2008; Grant & Scholes,

97 2006; Anderson et al. 2010; Joern et al. 2012). Indeed, carcasses of smaller vertebrates (e.g.,
98 salmon, deer) can increase the proportions of nitrogen and $\delta^{15}\text{N}$ (an indicator of animal-driven N)
99 in plants within just a few months postmortem (Hocking & Reynolds, 2012; van Klink et al.
100 2020).

101 To assess the effects of megacarcasses on local nutrient pools (Figure 1), we measured
102 the initial contributions of elephant carcasses (1-26 months postmortem) to soil and plant
103 nutrients in the Kruger National Park (KNP), South Africa. Further, we examined the effects of
104 elephant carcasses on the two main soil types in KNP: sandy, relatively nutrient-poor granitic
105 soils and clayey, relatively nutrient-rich basaltic soils (Venter et al. 2003). At each site, we ran
106 transects in each cardinal direction from the center of the site where an elephant died, collecting
107 samples of soil and a palatable grass species (*Urochloa trichopus*) at 0, 2.5, 5, 10, and 15 m. We
108 then analyzed soil samples for CNP and micronutrient content, quantified soil microbial
109 respiration potential, and measured %N, $\delta^{15}\text{N}$, and macro- and micronutrient content in grass
110 tissue. We hypothesized that: (H1) elephant megacarcass decomposition would release nutrients
111 into soil, resulting in higher concentrations of soil N, P, and micronutrients near the center of
112 carcass sites; (H2) C inputs to the soil would stimulate microbial activity, resulting in increased
113 soil respiration potential near the center of carcass sites; and (H3) carcass-derived nutrients
114 would move from soil into plants, resulting in higher foliar nutrient concentrations near the
115 center of carcass sites. We predicted that enrichment effects from megacarcasses would be
116 greater on sites with fresher carcasses relative to older carcasses and on nutrient-poor granitic
117 sites compared to nutrient-rich basaltic sites.

118

119 **Sect. 2 Methods**

120 2.1 Study system and sample collection

121 We performed this research in the southern part of the Kruger National Park (KNP), South
122 Africa (24.996 S, 31.592 E, ~275m elevation). The two dominant soil types in KNP are granitic
123 soils (Inceptisols) and basaltic soils (Versitols or Andisols) (Khomo et al. 2017). The clay-rich
124 basaltic soils have relatively large surface area, enabling them to retain larger quantities of water
125 than granitic soils, which drain water more quickly and therefore are lower in water-soluble
126 nutrients (Buitenwerf, Kulmatiski, & Higgins, 2014; Rughöft et al. 2016). The landscape of
127 KNP is a mix of savanna grasslands and broadleaf woodlands, with an overstory dominated by
128 trees from the genus *Combretum* (red bushwillow, *C. apiculatum*; russet bushwillow, *C.*
129 *hereroense*; leadwood, *C. imberbe*) and trees formerly known as acacias (knobthorn,
130 *Senegalensis nigrescens*; umbrella thorn, *Vachellia tortillis*). The park hosts a full suite of
131 African savanna animals, including ~30,000 elephants (*Loxodonta africana*) (Coetsee &
132 Ferreira, 2023), with a mortality rate of ~2% (~600 elephants per year). The targeted region of
133 KNP has a high density of scavengers and predators, including white-backed vultures (*Gyps*
134 *africanus*), spotted hyenas (*Crocuta crocuta*), and lions (*Panthera leo*) (Owen-Smith & Mills,
135 2007).

136 During the wet season in March 2023, we identified ten elephant carcass sites (1-26
137 months postmortem), five on relatively nutrient-rich basaltic soil and five on nutrient-poor
138 granitic soil. KNP section rangers provided precise GPS locations of where elephant carcasses
139 had been found. Most elephants died of old age, illness, injury, or, in the case of one young bull,
140 fighting over territory. Carcass sites were recognizable *in situ* by a persistent bonefield,
141 undigested gut contents, and an absence of herbaceous vegetation. At each site, we hammered a
142 rebar post into the center of the megacarcass disturbance and ran 15 m transects out from the

143 post in each of the four cardinal directions. We collected green leaf material from *U. trichopus*, a
144 common and abundant palatable grass species, and used an auger to collect soil samples to a
145 depth of 10 cm at five points along each transect (0.5, 2.5, 5, 10, and 15 m) (Bump, Peterson, &
146 Vucetich, 2009; Holdo & Mack, 2014; Gray & Bond, 2015; Monk et al. 2024). We treated the
147 10-15m distances as representative of background concentrations of nutrients based on pilot data
148 showing that the effect of elephant carcasses on soil nutrient concentrations was undetectable at
149 this distance away from the carcass site, similar to studies on the carcasses of other large
150 vertebrates (e.g., Towne, 2000; Bump et al. 2009). We pooled and homogenized the samples to
151 yield one composite leaf and one composite soil sample per sampling distance from each carcass
152 site. Soil samples were sieved in a 5-mm metal sieve which was cleaned in between samples
153 with 70% ethanol. Soil samples were stored in a cooler during fieldwork. On the day they were
154 collected, we used 5 g of each soil sample for soil respiration measurements (described below).
155 The rest of each sample was placed in a plastic bag on the day of collection and stored in a -20°C
156 freezer for up to 1 month; samples were stored in coolers with ice blocks during the transition
157 from the freezer at the field site to the lab for analysis. We chose to freeze samples rather than
158 storing at room temperature based on literature demonstrating that the impacts of freezing on soil
159 nitrate and ammonium concentrations are fairly minimal, except in specific cases of high soil
160 acidity or peaty soils that were not present at our field site (Esala, 1995; Turner & Romero, 2009;
161 Sollen-Norrin & Rintoul-Hynes, 2024). Leaf samples were stored in paper bags at room
162 temperature until dried for analyses (see below).

163

164 **2.3 Hypothesis testing**

165 We tested our first hypothesis that elephant carcass decomposition would release nutrients into
166 the soil by performing soil nutrient analyses. We sent 250 g of each soil sample to Eco-Analytica
167 laboratory at the North-West University in Potchefstroom, South Africa for measurements of soil
168 concentrations of ammonium $[\text{NH}_4]^+$, nitrate $[\text{NO}_3]^-$, phosphate $[\text{PO}_4]^{3-}$, and plant-available P.
169 Samples were air-dried and sieved through $< 2\text{mm}$ mesh prior to chemical analysis. Plant-
170 available P was extracted from 4 g of soil and 30 ml extraction fluid (1:7.5 ratio) using an acid-
171 fluoride solution (P Bray-1) (FAO, 2021), measured colorimetrically using a Systea
172 EasyChem200 analyser, and expressed as mg/kg. The detection limit was 0.5 mg/kg, and plant
173 available P measurements < 0.5 mg/kg were replaced with half the detection limit (0.25 mg/kg)
174 (Croghan & Egeghy, 2003). Water-soluble nitrate and phosphate anions were extracted from
175 volume on volume 100 mg soil and 200 ml deionized water (Sonneveld & van den Ende, 1981),
176 analyzed by ion chromatography on a Metrohm 930 Compact Flex System, and measured as
177 mg/L. Ammonium (also 1:2 water extract) was analyzed colorimetrically using a Systea
178 EasyChem200 analyzer and measured as mg/L. Detection limits for soil ions were 0.01 mg/L,
179 and soil ion concentrations measured as < 0.01 mg/L were replaced with half the detection limit
180 (0.005 mg/L). To convert the nitrate, ammonium, and phosphate units from mg/L to mg/kg, we
181 multiplied by 2, based on the 1:2 soil to water extraction ratio.

182 To determine whether soil micronutrients were distinct and elevated at the center of
183 carcass sites relative to soil further from the center, we measured concentrations of sodium (Na),
184 magnesium (Mg), iron (Fe), calcium (Ca), potassium (K), and phosphorus (P). Air-dried and
185 sieved (> 2 mm) soil samples, weighed to 0.2 g, were microwaved in 9 ml 65% nitric acid
186 (HNO_3) and 3 ml 32% hydrochloric acid (HCl) according to EPA 3051b in a Milestone, Ethos
187 microwave digester with UP, Maxi 44 rotor. A period of 20 minutes allowed the system to reach

188 1800 MW at a temperature of 200 °C which was maintained for 15 minutes. After cooling, the
189 samples were brought up to a final volume of 50 ml and analyzed on an Agilent 7500 CE ICP-
190 MS fitted with CRC (Collision Reaction Cell) technology for interference removal. The
191 instrument is optimized using a solution containing Li, Y, Ce, and Tl (1 ppb) for standard low-
192 oxide/low interference levels ($\leq 1.5\%$) while maintaining high sensitivity across the mass range.
193 The instrument was calibrated using ULTRASPEC® certified custom mixed multi-element stock
194 standard solutions containing all the elements of interest (De Bruyn Spectroscopic Solutions,
195 South Africa). Calibrations spanned the range of 0 – 30 ppm for the mineral elements Ca, Mg,
196 Na, and K and 0 – 0.3 ppm for the rest of the trace elements. Elemental concentrations were
197 expressed as mg/kg.

198 Finally, to determine whether elevated N levels in soils were derived from the carcass, we
199 sent 10 g of each sample to the BIOGRIP laboratory within the Central Analytical Facility at
200 Stellenbosch University for measurements of soil %N and $\delta^{15}\text{N}$, obtained using a Vario Isotope
201 Select Elemental Analyzer connected to a thermal conductivity detector and an Isoprime
202 precisions isotope ratio mass spectrometer (IRMS). Samples were oven-dried at 60°C for 48
203 hours and milled to a fine powder using a Retsch MM400 mill (Germany). The powdered
204 samples were weighed (2 – 60 mg) prior to combustion at 950°C. The gasses were reduced to N_2
205 (undiluted) in the reduction column, which was held at 600°C. A high organic carbon (HOC) soil
206 standard ($0.52 \pm 0.02\%$ N), along with two international reference standards (USGS40 ($\delta^{15}\text{N} -$
207 4.52% AIR) and USGS41 ($\delta^{15}\text{N} +47.57\%$ AIR)) were used for calibration. The N elemental
208 content was expressed relative to atmospheric N as N_2 $\delta^{15}\text{NAIR}$ (‰). The quantification limit for
209 $\delta^{15}\text{N}$ on the IRMS is 1 nA (nanoAmp), and the quantification limit for %N is 0.06%. The

210 precision for %N was 0.02% and for $\delta^{15}\text{N}$ is $\pm 0.11\%$, determined using the HOC standard, which
211 was run multiple times throughout the analysis.

212 To test our second hypothesis that nutrient inputs to the soil would stimulate microbial
213 activity, we measured soil organic C, water content, and microbial respiration potential. We sent
214 10 g of each sample to the BIOGRIP laboratory for measurements of soil organic C using a
215 Vario TOC Cube (Elementar, Germany). Samples (dried and milled as above) were weighed (10
216 – 60 mg), acidified using 10% HCl to remove the total inorganic C (carbonates), and dried
217 overnight at 60°C. All samples were analyzed through combustion at 950°C. The released CO₂
218 was measured by a non-dispersive infrared (NDIR) sensor. A high organic C ($7.45 \pm 0.14\%$ C)
219 soil standard from Elemental Microanalysis Ltd (UK) was included during the analysis. The
220 quantification limit for %C is 0.14%. The precision for the %C was 0.09% and was determined
221 using the low organic C (LOC) standard ($1.86 \pm 0.14\%$ C), which was run multiple times
222 throughout the analysis.

223 To quantify soil respiration and water content, we used an incubation method (Lemoine
224 et al. 2023) in which 5 g (± 0.2 g) of each sample was placed into a 100 ml clear glass bottle,
225 sealed, and flushed with CO₂-free air. Following flushing, we incubated the bottles for one hour
226 at 25°C. We then recorded CO₂ concentrations using an LI-850 CO₂/H₂O infrared gas analyzer.
227 After soil respiration measurements, we determined sample dry weight by drying each sample at
228 60°C for 24-48 hours until stable mass was achieved. We subtracted dry weight from starting
229 weight to obtain soil water content. Finally, we used the dry weights and the Ideal Gas Law to
230 standardize all respiration measurements to CO₂ $\mu\text{g h}^{-1}\text{g dry soil}^{-1}$.

231 To test our third hypothesis that carcass-derived nutrients would be incorporated by
232 plants, we measured foliar nutrient concentrations in *U. trichopus*. Two grams of each dried leaf

233 sample was sent to the BIOGRIP laboratory for preparation and measurements of %N and $\delta^{15}\text{N}$
234 via stable isotope analysis as described above. A Sorghum flour standard (1.47 ± 0.25 %N) from
235 Elemental Microanalysis Ltd (UK) was used for calibration, along with two international
236 reference standards (USGS40 and USGS41). The quantification limit for $\delta^{15}\text{N}$ on the IRMS is 1
237 nA, and the quantification limit for %N is 1.3%. The precision for the %N was 0.02% and for
238 $\delta^{15}\text{N}$ is $\pm 0.08\%$. Limits were determined using the sorghum flour standard, which was run
239 multiple times throughout the analysis. Additionally, we sent 5 g per sample to Cedara
240 Analytical Services Laboratory to quantify micronutrients in grass tissue (P, Na, Mg, K, Ca, and
241 Fe) using Inductively Coupled Plasma Optical Emission Spectroscopy (ICP-OES 5800, Agilent,
242 USA). Samples were dried (110°C overnight) and milled to a fine powder. Subsamples (0.5 g)
243 were ashed at 450°C for 4 hours, and the ash was re-wet using 2 mL conc. HCl (32%). Samples
244 were evaporated to dryness then re-suspended in 25 mL 1M HCl before filtering. Lastly, the
245 filtrate was diluted with de-ionized water in a ratio of 5:20 filtrate to water. To calibrate the ICP-
246 OES, solutions containing known amounts of each element were measured (10-20 ppm for Na
247 and C, 200-1500 ppm for Fe, 0.5-3.75% for K, and 0.125-0.5% for P), prepared from 1000 ppm
248 primary single standards. At three of the ten sites, we did not find sufficient plant material at the
249 central point for analysis, resulting in a sample size of $N = 7$ for the center (distance = 0.5m)
250 measurement for leaf nutrient analyses.

251 To test whether each response variable for the three hypotheses was significantly
252 associated with soil type and/or distance from the carcass center, we performed a model selection
253 procedure. For each response variable, we ran five generalized linear mixed models using the
254 gamma family (link = log) in the package *lme4* (Bates et al. 2015): (i) soil type + distance + soil
255 type \times distance interaction, (ii) soil type + distance, (iii) soil type, (iv) distance, and (v) a null

256 model indicating no significant difference in slope or intercept after accounting for carcass site.
257 All models included carcass site as a random effect to account for individual variation. Each
258 model included 50 observations (10 sites x 5 distances per site). For samples in which the
259 nutrient level was listed as 0 or undetectable, we accounted for the uncertainty by using half the
260 detection level as described above. The narrow distribution of ages (1-26 months since death)
261 with the sample size of $N = 10$ sites made testing for the effect of age challenging, so we did not
262 include carcass age in the models. We compared the models for each response variable using
263 Akaike Information Criterion (AICc). Models with a $\Delta AICc \leq 2$ were considered roughly
264 equivalent in fit (Burnham and Anderson, 2002).

265 In addition to these models, for our second hypothesis we regressed soil respiration
266 potential against soil organic C, expecting that the two would be positively correlated. We ran a
267 generalized linear mixed model with soil respiration potential as the response variable. The
268 model included soil organic C + distance + soil type, with carcass site as a random effect. We did
269 not include an interaction with soil type in this model due to sample size restrictions. Respiration
270 potential and organic C were both log-transformed to achieve normality.

271 To determine whether leaf and soil micronutrient composition differed with distance and
272 soil type, we ran permutational analysis of variance (perMANOVA) in *vegan* (Oksanen et al.
273 2022). We ran the same model separately for soil and leaf micronutrient composition (soil type +
274 distance). To determine which micronutrients contributed most to compositional differences
275 across distances and soil types, we calculated samplewise Bray-Curtis dissimilarity and
276 performed principal component analysis. We also tested for differences in variance in
277 micronutrient composition across distances and soil types using “betadisper” in *vegan* (Oksanen

278 et al. 2022). We ran linear models to test for correlations between leaf and soil concentrations of
279 each micronutrient. Each model included distance as a covariate and site as a random effect.

280 Finally, to test the impact of carcass age on key soil metrics, we ran exponential decay
281 functions for soil ammonium, nitrate, phosphate, and respiration verses carcass age for samples
282 from the center of the carcass site (0.5m sampling location). We also performed a t-test to verify
283 that there was no difference in mean carcass age across soil types.

284 All statistical analyses were performed in R version 4.2.1 (R Core Team, 2022).

285

286 **Sect. 3 Results**

287 **3.1 Hypothesis 1: Effects of megacarcasses on soil nutrient pools**

288 We found partial support for our first hypothesis that soil N and P concentrations would be
289 higher closer to the center of carcass sites (Table S1). Soil %N (Figure 2A) was overall greater in
290 basaltic soils, and it decreased with distance from the carcass site on granitic soils. Soil nitrate
291 (Figure 2B) decreased with distance from the carcass site but did not differ between soil types.
292 Ammonium (Figure 2C) also decreased with distance, but only in granitic soils. $\delta^{15}\text{N}$ (Figure 2D)
293 was greater in granitic soils and decreased with distance in both soil types, indicating that the
294 proportion of animal-sourced N in the soil was greater near the center of the carcass site. Soil
295 phosphate, plant available P, and mineral P (Figure 2E-G) all exhibited significant soil \times distance
296 interactions. Phosphate (Figure 2E) was highly elevated at the center of carcass sites and
297 decreased steeply with distance, but only in granitic soils. Plant-available P (Figure 2F)
298 decreased with distance in both soil types, but the effect was strongest in granitic soils. Finally,
299 mineral P (Figure 2G) was greater in basaltic soils, and there was a small decrease with distance
300 in granitic soils but not in basaltic soils.

301 Contrary to our first hypothesis, soil micronutrient composition did not differ
302 significantly with distance from the carcass center; nor did most individual micronutrients (Table
303 S1). The perMANOVA results showed that soil micronutrient composition did not differ
304 significantly with distance ($R^2 = 0.00$, $F_{4,44} = 0.0$, $P = 1.000$) (Figure S2A), but it did differ
305 significantly with soil type ($R^2 = 0.71$, $F_{1,44} = 108.8$, $P = 0.001$) (Figure S2B). There was no
306 significant difference in variance with distance ($F_{4,45} = 0.0$, $P = 0.996$) or soil type ($F_{1,48} = 2.6$, P
307 $= 0.115$). Principal components analysis showed that dimension 1 explained 53.6% of the
308 variation between soil types and was driven primarily by differences in Mg, Ca, and Fe.
309 Dimension 2 explained 25.9% of variation and was driven primarily by differences in K. Soil Na
310 (Figure S3A) was marginally greater in granitic soils and decreased with distance from the
311 carcass, with the effect greater in granitic soils. Soil K (Figure S3B) was greater in basaltic soils
312 and decreased marginally with distance. Soil Fe, Mg, and Ca (Figure S3C-E) were greater in
313 basaltic soils, with minimal effects of distance.

314

315 **3.2 Hypothesis 2: Effects of megacarcasses on soil carbon and respiration**

316 Consistent with our second hypothesis, soil respiration potential was marginally positively
317 correlated with soil organic carbon concentration and decreased significantly with distance but
318 did not differ with soil type (Figure 3). We found no evidence for differences in soil water
319 content (Figure S4A) or soil pH (Figure S4B) with distance or soil type. In both cases, the null
320 ranked among the set of top models (Table S1).

321

322 **3.3 Hypothesis 3: Effects of megacarcasses on plant nutrient pools**

323 Consistent with our third hypothesis, we found elevated foliar nutrient concentrations in *U.*
324 *trichopus* at elephant carcass sites. Leaf %N (Figure 4A) and $\delta^{15}\text{N}$ (Figure 4B) both decreased
325 with distance from the carcass center. $\delta^{15}\text{N}$ exhibited a significant soil \times distance interaction in
326 which it was overall greater in basaltic soils, but the difference between the two soil types was
327 greater closer to the carcass site. Foliar P was greater in basaltic soils and decreased only
328 marginally with distance in the granite soils. Finally, the foliar N:P ratio was greater in granitic
329 soils and decreased with distance in the basaltic soils.

330 Leaf micronutrient composition did not differ significantly with distance ($R^2 = 0.13$, $F_{4,40}$
331 $= 1.9$, $P = 0.062$; Figure S5A) but did differ with soil type ($R^2 = 0.17$, $F_{1,40} = 9.7$, $P = 0.001$;
332 Figure S5B). There was no significant difference in variance with distance ($F_{4,41} = 0.5$, $P =$
333 0.713) or soil type ($F_{1,44} = 1.9$, $P = 0.173$). Dimension 1 explained 42.8% of the variance across
334 soil types and was primarily driven by Mg, Na, and P. Dimension 2 explained 26.6% of the
335 variance and was driven mainly by K, Ca, and Fe. Foliar Na (Figure S6A) and Mg (Figure S6B)
336 were both greater in basaltic soils and decreased with distance from the carcass center. Foliar K
337 (Figure S6C) and Fe (Figure S6D) both decreased with distance as well but did not differ with
338 soil type. The null model was in the top set for foliar Ca, indicating no significant relationship
339 between foliar Ca concentrations and soil type or distance from the carcass center. Individual
340 micronutrients (K, Ca, Mg, Fe) were not correlated between leaf and soil samples, with the
341 exception of Na (Table S3).

342

343 **3.4 Effects of carcass age on soil ions and respiration potential**

344 Soil ammonium ($\alpha = 0.018$, $P < 0.001$), phosphate ($\alpha = 0.023$, $P < 0.001$), and respiration
345 potential ($\alpha = 0.058$, $P < 0.001$) all decreased significantly with carcass age (Figure 5A-C). The

346 exponential decay model for nitrate failed to converge due to an outlier with extremely high soil
347 nitrate (1454 mg/kg) at 258 days postmortem (Figure 5D). We ran a t-test to test for a difference
348 in mean carcass age between soil types and found no significant difference between the two
349 groups ($P = 0.294$).

350

351 **Sect. 4 Discussion**

352 Here, we show that elephant megacarcasses influence soil and foliar nutrients during at least the
353 first two years following mortality. Consistent with our hypotheses, soil nitrate (Figure 2B),
354 ammonium (Figure 2C), $\delta^{15}\text{N}$ (Figure 2D), and P (Figure 2E-F) concentrations were all elevated
355 at the center of carcass sites and decreased with distance from the center. Soil %N, nitrate,
356 ammonium, and plant-available P concentrations at the 15m point were consistent with those
357 found in other studies of soil nutrient content in Kruger (Aranibar et al. 2003; Rughöft et al.
358 2016), confirming that the 15m point serves as an effective baseline control in this experiment.
359 Microbial respiration potential was also elevated towards the center of carcass sites and was
360 correlated with the abundance of organic C (Figure 3). Finally, %N (Figure 4A) and $\delta^{15}\text{N}$ in a
361 common grass (Figure 4B) were both elevated closer to the centers of carcass sites compared to
362 grass farther from carcasses. Together, these results indicate that carcass-derived nutrients move
363 into soil and subsequently get absorbed by plants over relatively short time scales, cycling
364 essential nutrients such as N from carrion into the soil and then back into aboveground nutrient
365 pools.

366 The initial influx of ammonium from elephant carcasses may have time-dependent
367 impacts on plant abundance at elephant carcass sites. The mean ammonium level at the center of
368 carcass sites (34.8 mg/kg) was higher than the level generally considered toxic to plants (Britto

369 & Kronzucker, 2002). Yet, we found living grass—typically *U. trichopus*—in the center of the
370 carcass site at seven out of ten of our sites (ammonium range 10-172 mg/kg) and at the 2.5m
371 distance for all sites (ammonium range 0-72 mg/kg). The three sites without vegetation in the
372 center had the highest ammonium levels (70-144 mg/kg), suggesting that *U. trichopus* has a
373 higher degree of ammonium tolerance than some sympatric grass species but may still be limited
374 by the high ammonium levels at the centers of these three relatively fresh carcass sites. However,
375 the recentness of the disturbance from the carcass likely also plays a role in determining plant
376 abundance near the center of the carcass. Because of the elephant carcass site age distribution,
377 (mean 350 days postmortem; range 24-811 days), this study may not have captured the full
378 impact of ammonium release from carcasses during the early stages of decomposition. Soil
379 ammonium spiked early and decreased rapidly (Figure 5A), and future research on carcasses
380 within the first few weeks postmortem would enhance our understanding of these early nutrient
381 dynamics.

382 Soil nitrate (Figure 2B) and soil respiration potential (Figure 3) were also elevated near
383 the center of carcass sites, indicating higher activity rates for nitrifying bacteria and heterotrophic
384 microbes (Prosser, 2011). These results are consistent with other work on carrion, where
385 microbial activity tends to be greater in soils near carcasses as compared to surrounding soil
386 (Bump et al. 2009b). However, carcass effects on soil microbial respiration exhibit a high degree
387 of intra-system variation (Risch et al. 2020), and the potentially short window during which
388 increased respiration occurs may make capturing these variations challenging. For example, soil
389 respiration potential at the center of the three youngest carcass sites was on average 2x higher
390 than the seven older sites (18.43 and 9.62 $\mu\text{g CO}_2$ per hour per gram of dry soil, respectively;
391 Figure 5D). Thus, the impact of increased organic C on soil microbial processes may be

392 relatively short lived and only last a matter of months (Keenan et al. 2018; Keenan, Schaeffer, &
393 DeBruyn, 2019). These trends are consistent with soil ammonium and phosphate, both of which
394 are highest at the youngest carcass sites (<200 days postmortem; Figure 5A-B). Soil microbial
395 respiration rate is also highly elevated early on, but it decreases at a faster rate over time than soil
396 ions (Figure 5C). Thus, soil dynamics during the first several months after death may play a
397 crucial role in determining the long-term impacts of megacarcasses on savannas and therefore
398 provides a promising avenue for future research.

399 Elevated soil phosphate (Figure 2E) and plant-available P (Figure 2F) at the center of
400 carcass sites were also consistent with expectations from the literature (Bump et al. 2009a;
401 Parmenter & MacMahon, 2009). However, elevated P levels in soil did not translate to elevated
402 P in grass leaves (Figure 4C), which could suggest a lag between trends in soil and plants that is
403 longer for P than for N. This lag could occur because phosphate easily forms chemical bonds
404 with other soil ions (e.g., iron and aluminum in acidic soils and calcium in basic soils). Nitrate
405 does not form these bonds and therefore has greater water solubility and mobility in soils and
406 may be more readily taken up by plants (Wiersum, 1962; Arai & Sparks, 2007). However, it is
407 also possible that P limitation in Kruger is not as strong as it is in some other African savanna
408 systems (Pellegrini, 2016). The foliar N:P ratios measured in this experiment were higher closer
409 to the center of the carcass site (median 9.38 at 0 m and 4.83 at 15 m), indicating that N
410 limitation may be relatively stronger further from the carcass site, and P limitation may be
411 relatively stronger closer to the center (Figure 4D, Table S2). These relatively high foliar N:P
412 ratios at the center of carcass sites are similar to those found in N fertilization studies in Kruger
413 (Craine et al. 2008), further supporting the idea that the influx of N from megacarcasses may
414 shift the soil from relatively more N limited to more P limited.

415 The elevated plant-available P at the center of carcass sites likely came primarily from
416 phosphate released from decomposing tissue (Yong et al. 2019). Bone decomposition, which is
417 also likely a major source of P from animal carcasses (Subalusky et al. 2020), occurs over long
418 time scales (Coe, 1978; Subalusky et al. 2020) and therefore should result in the slow release of
419 P and a gradual decrease in the N:P ratio (Parmenter & MacMahon, 2009; Quaggiotto et al.
420 2019). Indeed, initial inorganic N influxes to the Mara River in Kenya from mass wildebeest die-
421 offs are 10-fold greater than concurrent increases in P, which instead releases slowly over about
422 seven years of bone decomposition (Subalusky et al. 2017). Research following megacarcasses
423 over longer timeframes postmortem is needed to clarify when P from enriched soil is absorbed
424 by plants and at what stage megacarcass bones begin contributing to soil P dynamics. It is also
425 possible that bone dispersal by scavengers may result in less P leaching from bones close to
426 where the elephant died and more P being distributed across the landscape at distances far from
427 the carcass site.

428 The contributions of megacarcasses to soil nutrient pools were strongly associated with
429 soil type. Our results confirmed the previously-established trend that basaltic soils are overall
430 more cation rich than granitic soils, with greater concentrations of P, K, Fe, Mg, and Ca (Figure
431 2G; Figure S3B-E; Gertenbach, 1983; Craine, Morrow, & Stock, 2008; Wigley et al. 2014).
432 However, soil ammonium, $\delta^{15}\text{N}$, and phosphate were all higher in the granitic soils towards the
433 center of carcass sites, decreasing steeply to be similar to basaltic soils about 10 m from the
434 carcass center (Figure 2C-E). These results indicate that the impact of organic matter from
435 megacarcasses may be stronger in relatively nutrient-poor and sandy granitic soil compared with
436 nutrient-rich and clayey basaltic soil. We were surprised that grass on basaltic soil did not
437 consistently exhibit greater nutrient concentrations. One potential explanation is that grass may

438 primarily be limited by macronutrients like N and P on both soil types (Craine et al. 2008;
439 Holdo, 2013) rather than by micronutrients. Thus, even with increased micronutrient availability
440 their actual uptake may not differ substantially. Studies on ungulate carcasses (e.g., muskoxen,
441 moose, zebra) have shown increased foliar N at carcass sites (Danell et al. 2002; Bump et al.
442 2009b; Turner et al. 2014), but to date there is little research on the flow of micronutrients from
443 carrion to plants and none on the pipeline from megacarcasses to plants. Moreover, it remains to
444 be seen whether increases in foliar N and other nutrients affect herbivory rates at carcass sites
445 and how long such effects may last.

446 The magnitude of nitrogen inputs from megacarcasses, as well as the substantial size and
447 duration of their impact zones, means their impacts on ecosystem processes may be functionally
448 distinct from smaller carrion. Soil nitrate concentrations at elephant carcass sites are orders of
449 magnitude higher than at carcass sites of smaller carrion (e.g., rabbits, white-tailed deer,
450 kangaroo) (Quaggiato et al. 2019; Bump et al. 2009; Barton et al. 2016). Even for large ungulates
451 such as moose, total soil inorganic nitrogen (ammonium + nitrate) at carcass sites is a mean 300
452 mg/kg (Bump, Peterson, & Vucetich, 2009), substantially lower than the mean total soil
453 inorganic nitrogen at elephant carcass sites (2.5m distance; 473 mg/kg). Termite mounds,
454 another long-lasting source of savanna nutrient heterogeneity, have mean soil nitrate
455 concentrations (197 mg/kg) lower than elephant carcass sites, but maximum nitrate
456 concentrations that are on par with them (974 mg/kg) (Seymour et al. 2014), again indicating that
457 elephant carcasses are one of the strongest known individual contributors of soil nitrogen in
458 African savanna ecosystems, which may have important implications for savanna ecology.
459 Indeed, there is evidence that carcass size strongly impacts scavenger food web structure
460 (Moleón et al. 2015; Morris et al. 2023). Moreover, the attraction of animals to carcasses via

461 scavenging, predation, or mourning (Goldenberg & Wittemyer, 2020) could have positive
462 feedbacks on nutrient cycling (Bump, Peterson, & Vucetich, 2009; Monk et al. 2024), which
463 may be magnified by carcass size. Thus, the impacts of megacarcasses on savanna ecosystem
464 processes may be dissimilar to the effects of small carrion and more similar to other more
465 persistent contributors to savanna ecosystem processes, such as termite mounds (Davies et al.
466 2016), cattle bomas (Augustine, 2003), and even mass animal mortality events (Subalusky et al.
467 2017, 2020).

468

469 **Sect. 5 Conclusions**

470 This study is an important first step in understanding the ecological legacies of megacarcasses on
471 savanna ecosystem processes. During the first two years postmortem, elephant carcasses released
472 pulses of nitrogen and phosphate, which influence savanna primary productivity. These nutrients
473 stimulated soil microbial activity and enriched foliar N, and the effects were strongest in
474 nutrient-poor soil, with potential long-term impacts on savanna nutrient heterogeneity. These
475 carcass-derived nutrient hotspots represent a previously unstudied function of megaherbivores on
476 savannas—one that we need to better understand in order to comprehend the full impacts of
477 megaherbivore population declines in modern ecosystems.

478

479 **Data and Code Availability:** Data and computer code are archived on Dryad Digital Repository
480 (<https://doi.org/10.5061/dryad.wpzgmsbwm>).

481

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486 Burkepile collected samples. Courtney G. Reed, Nathan P. Lemoine, Dave I. Thompson, and
487 Deron E. Burkepile analyzed the data. Courtney G. Reed drafted the manuscript, and all authors
488 contributed to editing.

489

490 **Competing Interests:** The authors declare that they have no conflict of interest.

491

492 **Acknowledgments:** Funding for this research was provided by the National Science Foundation
493 (#s 2128092, 2128093, and 2128094) and the University of California Santa Barbara Academic
494 Senate. All research was completed under permits from South African National Parks (SS554).
495 We thank the field assistants of SANParks for guiding and protection in the field, as well as the
496 section rangers and Sandra Snelling for GPS locations and ages of carcasses. We thank Mr.
497 Lucky Sithole (Cedara Analytical Services Laboratory), Ms. Terina Vermeulen (Eco-Analytica
498 Laboratory), and Dr. Janine Colling (BIOGRIP laboratory) for extensive laboratory support.
499 Select data used in this research paper were generated using equipment in the DSI funded
500 BIOGRIP soil and water node at Stellenbosch University.

501

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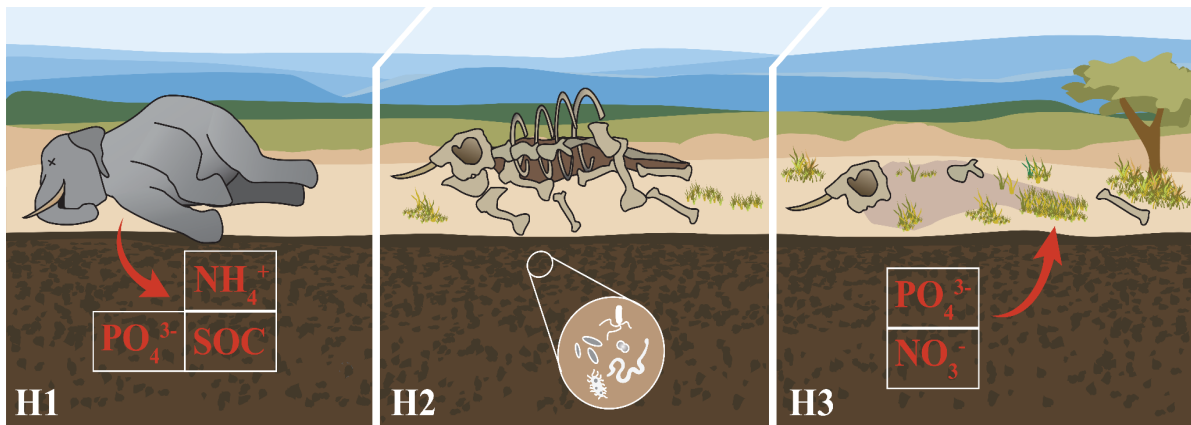
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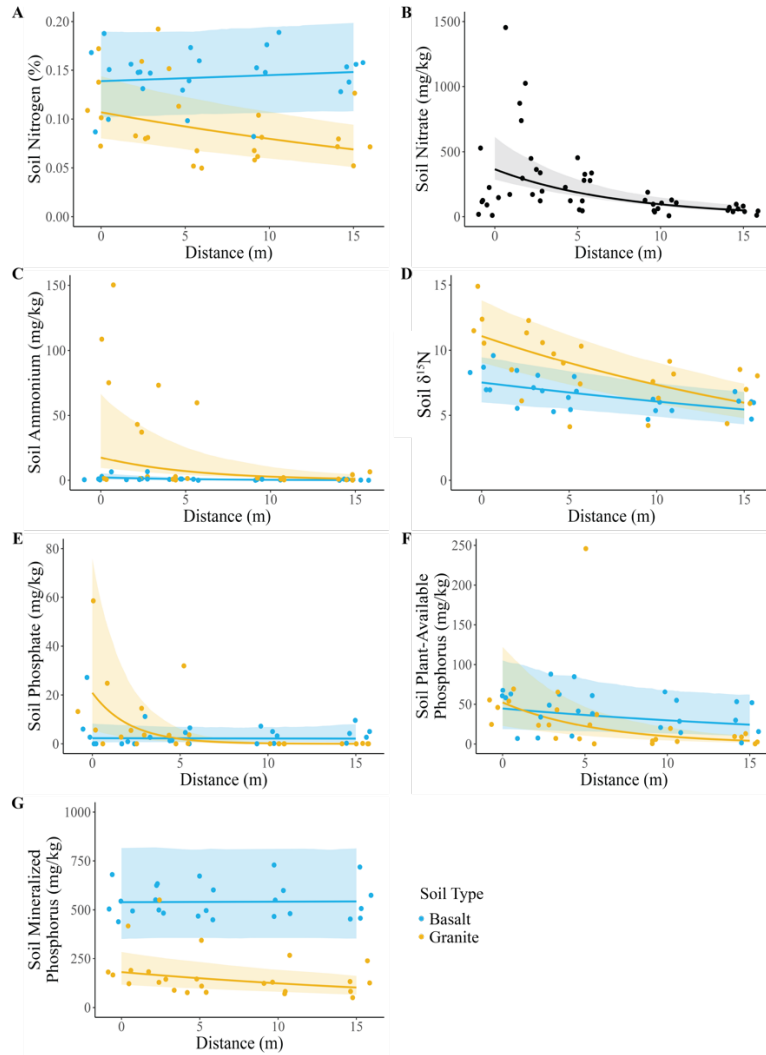
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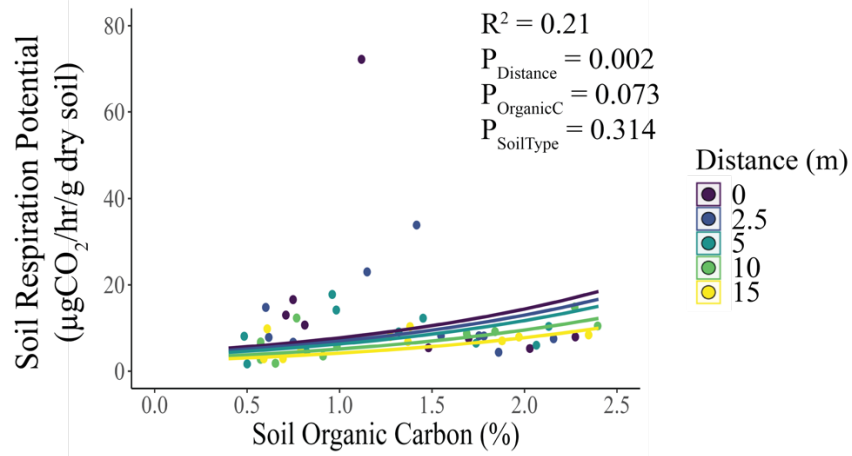
762 **Figure 1.** Hypothesized impacts of elephant megacarcasses on soil and plant nutrients. First
 763 (H1), we hypothesized that elephant carcasses would release pulses of nutrients into the soil,
 764 resulting in higher concentrations of soil ions such as nitrogen (ammonium, $[\text{NH}_4]^+$), phosphorus
 765 (phosphate, $[\text{PO}_4]^{3-}$), and soil organic C. Second (H2), we hypothesized that C inputs from the
 766 carcass would result in increased soil microbial respiration potential. Third (H3), we
 767 hypothesized that plants would take up nutrients from the carcass soil, resulting in plants with
 768 distinct nutrient profiles and increased concentrations of key limiting nutrients such as N and P.

769 Image credit: Kirsten Boeh.



770

771 **Figure 2.** Soil N and P responses to elephant carcasses. (A) Soil N (%) was greater in basaltic
 772 soils, and in granitic soils it decreased with distance from the carcass site. (B) Soil nitrate
 773 nitrogen decreased with distance but did not differ with soil type. (C) Soil ammonium nitrogen
 774 and (D) $\delta^{15}\text{N}$ were both greater in granitic soils and decreased with distance from the carcass. (E)
 775 Soil phosphate, (F) plant-available P, and (G) mineralized P decreased with distance in granitic
 776 soils but not basaltic soils. Points represent individual measurements taken at 0, 2.5, 5, 10, and
 777 15m and are offset to be visible when they would otherwise overlap. Lines show predictions
 778 calculated from the top model (see Table S1). Shading indicates the 95% confidence interval.



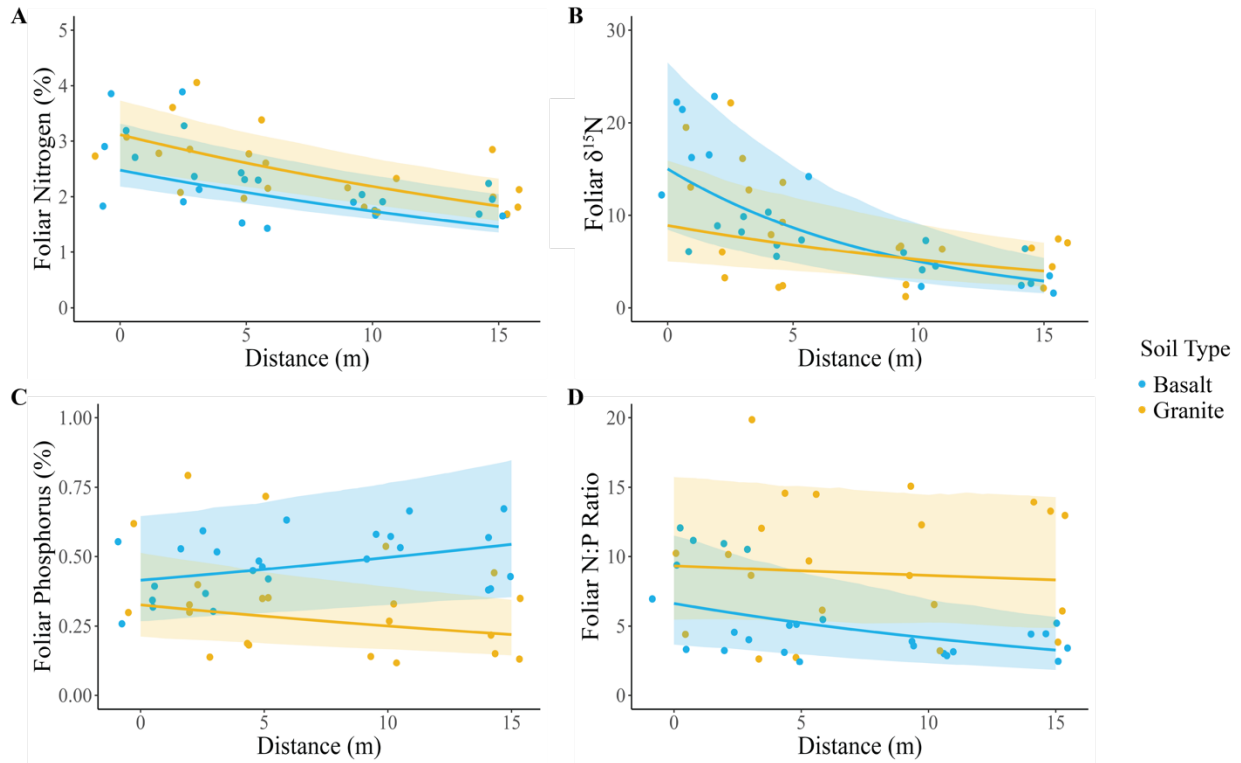
779

780 **Figure 3.** Soil respiration potential was marginally positively correlated with soil organic C (%)

781 and decreased significantly with distance from the carcass. Points represent individual

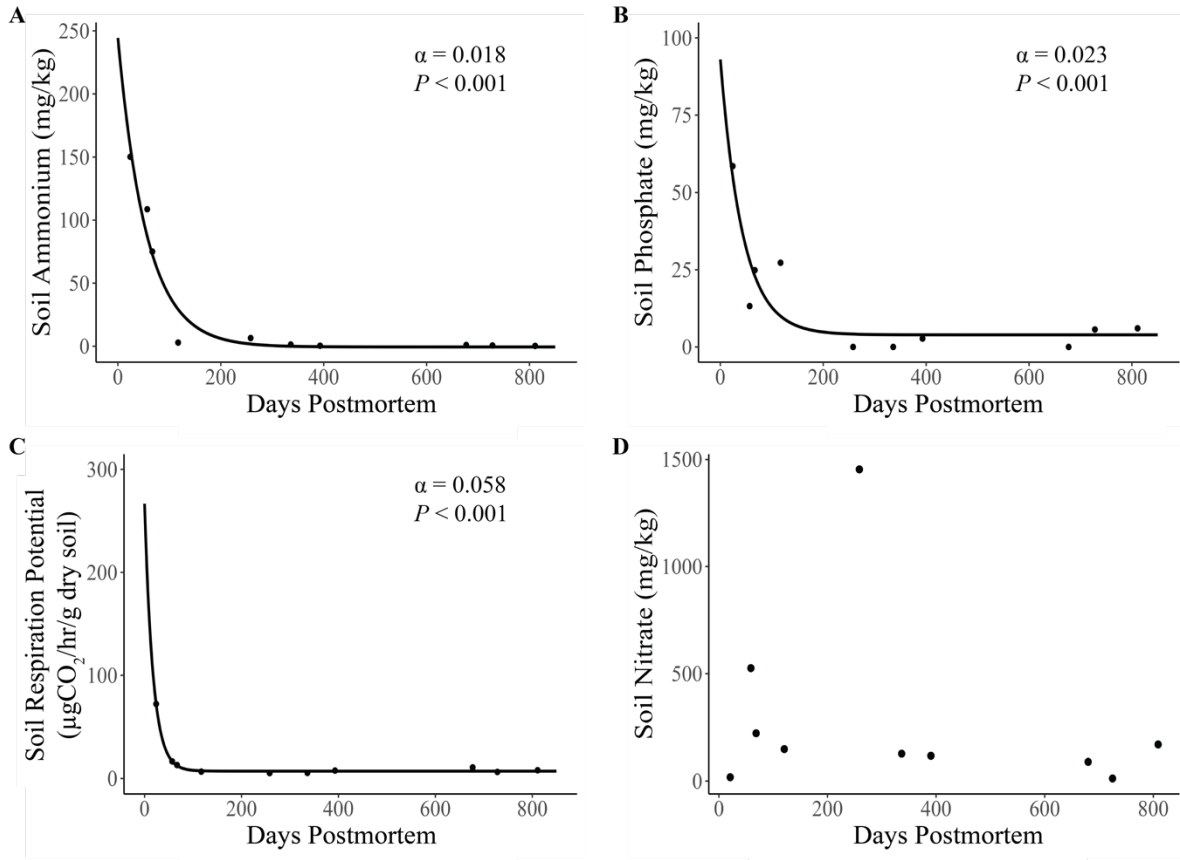
782 measurements taken at 0, 2.5, 5, 10, and 15m and are offset to be visible when they would

783 otherwise overlap. Lines represent model predictions.



784

785 **Figure 4.** Foliar N and P responses to elephant carcasses. (A) Foliar %N and (B) $\delta^{15}\text{N}$ both
 786 decreased with distance from the carcass center. (C) Foliar P was greater in basaltic soils and
 787 decreased with distance in granitic soils. (D) Foliar N:P ratio was greater in granitic soils and
 788 decreased with distance from the carcass center. Points represent individual measurements taken
 789 at 0, 2.5, 5, 10, and 15m and are offset to be visible when they would otherwise overlap. Lines
 790 show predictions calculated from the top model (see Table S2). Shading indicates the 95%
 791 confidence interval. Three of the ten sites had bare ground at the 0 m distance, resulting in a
 792 sample size of 7 sites for that distance and 10 for the other distances.



793

794 **Figure 5.** Relationship between carcass age and key soil metrics (soil ion concentrations and
 795 respiration potential). Lines represent predictions from exponential decay models, with α equal
 796 to the rate of decay. (A) Soil ammonium, (B) phosphate, and (C) respiration potential all
 797 decreased significantly with carcass age. The model for (D) soil nitrate failed to converge. Points
 798 represent individual measurements taken at the center of the carcass site (distance = 0.5m).