

# Functional regimes define the response of the soil microbiome to environmental change

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## 17 Abstract

18 **The metabolic activity of soil microbiomes plays a central role in carbon and ni-**  
19 **trogen cycling. Given the changing climate, it is important to understand how the**  
20 **metabolism of natural communities responds to environmental change. However,**  
21 **the ecological, spatial, and chemical complexity of soils makes understanding the**  
22 **mechanisms governing the response of these communities to perturbations chal-**  
23 **lenging. Here, we overcome this complexity by using dynamic measurements of**  
24 **metabolism in microcosms and modeling to reveal regimes where a few key mech-**  
25 **anisms govern the response of soils to environmental change. We sample soils**  
26 **along a natural pH gradient, construct >1500 microcosms to perturb the pH, and**  
27 **quantify the dynamics of respiratory nitrate utilization, a key process in the nitro-**  
28 **gen cycle. Despite the complexity of the soil microbiome, a minimal mathematical**  
29 **model with two variables, the quantity of active biomass in the community and**  
30 **the availability of a growth-limiting nutrient, quantifies observed nitrate utiliza-**  
31 **tion dynamics across soils and pH perturbations. Across environmental perturba-**  
32 **tions, changes in these two variables give rise to three functional regimes each with**  
33 **qualitatively distinct dynamics of nitrate utilization over time: a regime where**  
34 **acidic perturbations induce cell death that limits metabolic activity, a nutrient-**  
35 **limiting regime where nitrate uptake is performed by dominant taxa that utilize**  
36 **nutrients released from the soil matrix, and a resurgent growth regime in basic**  
37 **conditions, where excess nutrients enable growth of initially rare taxa. The un-**  
38 **derlying mechanism of each regime is predicted by our interpretable model and**  
39 **tested via amendment experiments, nutrient measurements, and sequencing. Fur-**  
40 **ther, our data suggest that the long-term history of environmental variation in the**  
41 **wild influences the transitions between functional regimes. Therefore, quantita-**  
42 **tive measurements and a mathematical model reveal the existence of qualitative**  
43 **regimes that capture the mechanisms and dynamics of a community responding**  
44 **to environmental change.**

## 45 **Introduction**

46 The metabolic activity of soil, marine, and freshwater microbiomes drives carbon and nitrogen  
47 transformations that sustain biogeochemical cycles and life in the biosphere [1–3]. These micro-  
48 biomes are also subjected to environmental perturbations including changes in temperature, pH,  
49 moisture, oxygen, and nutrients stemming from natural and anthropogenic events. As such, in or-  
50 der to predict the effect of climate change on global nutrient cycles, it is necessary to understand  
51 how microbiome metabolism responds to environmental change in nature.

52 Determining how environmental change impacts community metabolism has proven vexing be-  
53 cause of the complexity of natural microbiomes. This complexity is perhaps most apparent in soils,  
54 which possess immense taxonomic diversity [4], spatial heterogeneity [5], and chemically diverse  
55 environments [6]. As a result, environmental perturbations can modify collective metabolic activity  
56 in many ways, from direct changes in microbial composition, physiology [7], and ecological inter-  
57 actions [8, 9] to indirect modification of nutrient availability [10–12] and spatial organization [5,  
58 13]. Thus, a key question arises: which mechanisms are important for determining the metabolic  
59 response of complex microbiomes to environmental change?

60 Large-scale surveys approach this question by quantifying correlations between environmental  
61 variation, community composition, and metabolic processes in the wild [14–25]. Although surveys  
62 have revealed robust correlations, they face two challenges in uncovering the mechanisms deter-  
63 mining community response to environmental change. First, and most importantly, surveys do not  
64 allow control for confounding factors, such as correlated environmental variables, rendering any  
65 causal inference infeasible. Second, it is difficult to quantify metabolic dynamics in situ on a large  
66 scale in the wild. As a result, surveys have limited power to determine the mechanisms that govern  
67 the metabolic response to environmental change in natural communities.

68 To control for confounding factors and gain mechanistic insights, we use soil microcosms,  
69 which remove correlated environmental fluctuations and permit controlled perturbations in the lab.  
70 To further control for confounding factors, these soils are sourced from a single site [26, 27] that

71 exhibits large natural pH variation but minimal variability in other environmental factors (e.g.,  
72 climate, moisture, soil texture, C/N ratio). Leveraging insights from global surveys, we focus on  
73 pH – the environmental variable that shows a strong correlation with soil microbiome composition  
74 and features of metabolism. [9, 15, 25, 28, 29]. Second, soil microcosms enable high-throughput  
75 quantification of metabolic time series in response to environmental perturbations. Our metabolic  
76 measurements focus on a key functional process in nitrogen cycling, the anaerobic respiration of  
77 nitrate which is ubiquitously performed by complex communities of soil bacteria, in response to  
78 natural and applied changes in pH.

79 Here, we measure nitrate utilization dynamics in >1500 microcosms across a wide range of  
80 natural and laboratory-induced pH changes. Next, we develop a judicious mathematical framework  
81 that accurately describes nitrate utilization dynamics across all microcosms. Our model shows that  
82 changes in functional dynamics in response to pH perturbations can be mechanistically understood  
83 by considering just two variables: the quantity of biomass in the community actively utilizing  
84 nitrate and the availability of growth-limiting nutrients. These two parameters emerge naturally  
85 from our mathematical model using only the community-level nitrate uptake data. The model  
86 predicts that changes in pH alter nitrate utilization dynamics by differentially affecting both the  
87 quantity of active biomass and the availability of nutrients that limit its growth.

88 As a result, despite the ecological, chemical, and spatial complexity of soils, we find that the  
89 functional response of the soil microbiome to changes in pH can be categorized into three mecha-  
90 nistically distinct regimes demarcated by the levels of these two variables. Each functional regime  
91 is defined by which of the two variables exerts greater control over nitrate utilization rates. During  
92 moderate pH perturbations, metabolic rates are set by the pH-mediated release of nutrients from  
93 soil particles that limit the growth of a large metabolically active biomass (Nutrient-limiting regime,  
94 Regime II). When soils are subjected to large basic perturbations, massive nutrient release relieves  
95 the nutrient limitation, but the dominant taxa are no longer metabolically active, and metabolism is  
96 set by the rapid growth of initially rare taxa (Resurgent growth regime, Regime III). During large

97 acidic perturbations, functional responses are limited by the pervasive death of the active biomass  
98 in the community (Acidic death regime, Regime I). The transition between functional regimes can  
99 be abrupt (from Regime II to III) or smooth (from Regime I to II) as pH is varied and depends  
100 on the long-term pH history of the soil. Thus, while the dynamics and mechanisms of each func-  
101 tional regime are conserved across soils, the transitions between regimes depend on environmental  
102 history and community composition. Our study demonstrates a generalizable approach wherein  
103 high-throughput soil microcosm experiments coupled with mathematical models can overcome the  
104 complexity of natural ecosystems to mechanistically reveal the specific microscopic processes that  
105 contribute to the microbiome's response to environmental change.

## 106 **Results**

107 Nitrate ( $\text{NO}_3^-$ ), which has critical implications for agriculture and climate, is reduced in soils when  
108 bacteria use it as an electron acceptor during anaerobic respiration in the absence of oxygen. Both  
109 denitrification ( $\text{NO}_3^- \rightarrow \text{NO}_2^- \rightarrow \dots \rightarrow \text{N}_2$ ) or dissimilatory nitrate reduction to ammonia (DNRA,  
110  $\text{NO}_3^- \rightarrow \text{NO}_2^- \rightarrow \text{NH}_4^+$ ) reduce nitrate to nitrite ( $\text{NO}_2^-$ ) while consuming organic carbon. Due to  
111 the importance of pH in microbial physiology and soil chemistry, decades of studies have examined  
112 how pH affects nitrate reduction [30]. However, discrepancies in experimental methods (Table S1)  
113 and limited modeling, have made it difficult to find principles governing metabolic responses to pH  
114 perturbations [17] (Table S2).

### 115 **Metabolite dynamics in soils after short and long-term pH perturbations**

116 To address this problem we measured nitrate utilization dynamics in soil microcosms across a  
117 range of native and perturbed pH levels. We sampled 20 top soils with pH from 4.7 to 8.3 (Fig. 1A,  
118 Table S3) at the Long-term Agricultural Research Cook Agronomy Farm (CAF) (Pullman, WA,  
119 USA). Sampled soils had similar characteristics (Table S3) which minimized the effects of con-  
120 founding factors that might alter metabolic responses to perturbations. At this site, long-term vari-

121 ation in soil pH arises from local agricultural practices and erosion.

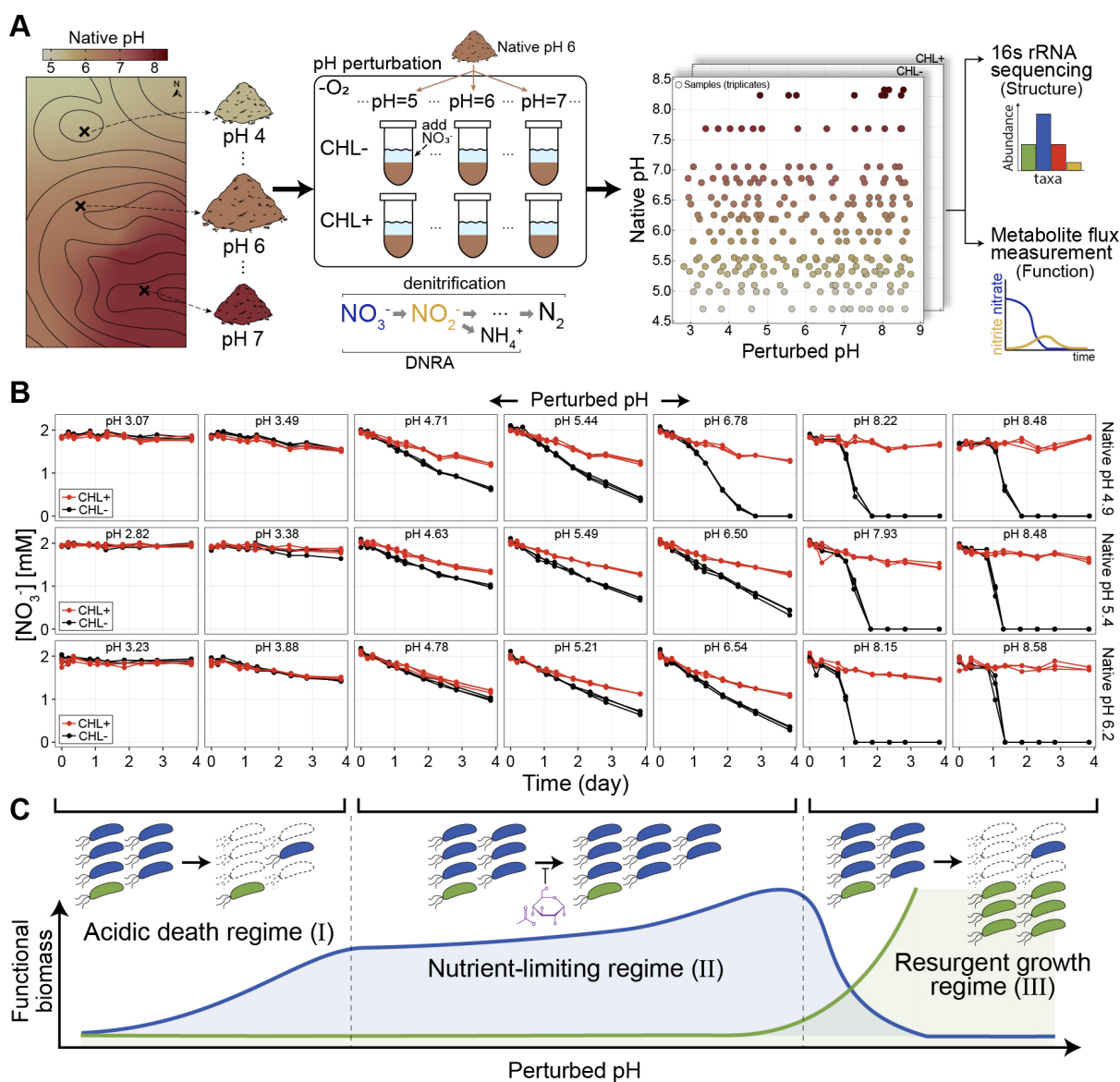
122 For each soil sample, we created mixtures of soil and water (slurries) with 2mM nitrate and  
123 varying levels of strong acid or base to perturb each soil's native pH to 13 values between 3 and 9  
124 (Fig. 1A). Therefore, our experiment quantifies the effects of short-term pH perturbations, while the  
125 differences between soils can inform us about the effects of long-term exposure to high or low pH.  
126 We employed slurries to make amendments easier, limit the effects of differential water content,  
127 and mimic rain events when most of the anaerobic respiratory nitrate utilization occurs [31, 32].

128 Soil slurries retained much, but not all, of the complexity of the natural context, including  
129 the diversity of the communities, the soil nutrient composition, and the spatial structure due to  
130 intact soil grains. The metabolic activity we observed relied only on the natively available carbon  
131 (electron donor for nitrate reduction), and thus we did not enrich for specific taxa beyond the nitrate  
132 added to the system. To separate the activity of pre-existing nitrate utilizers from growth in each  
133 condition [33], we included controls in every pH perturbation treated with chloramphenicol which  
134 inhibits protein synthesis (Fig. 1A). The dataset comprised 20 soils, at 13 distinct pH levels, with  
135 and without chloramphenicol, each in triplicate.

136 We measured the dynamics of the relevant metabolites (nitrate, nitrite, ammonium, and water-  
137 soluble organic carbon) during the 4-day incubation in anaerobic conditions (Fig. 1A). Focusing on  
138 non-gaseous metabolites enabled us to perform high temporal resolution measurements of metabo-  
139 lite dynamics across the ~1500 microcosms. For 10 of 20 soils, we performed 16S rRNA amplicon  
140 sequencing before and after incubations.

141 We observed three types of dynamics across pH perturbations and soils (Fig. 1B). First, all  
142 chloramphenicol-treated (CHL+) conditions exhibited linear nitrate ( $\text{NO}_3^-$ ) utilization dynamics  
143 (red lines, Fig. 1B, Figs. S1, S2). This is expected because, with chloramphenicol, nitrate reducers  
144 are unable to grow, and the rate of nitrate reduction remains constant [31]. The slope of nitrate in  
145 time in CHL+ conditions quantifies the activity of the pre-existing functional biomass. For large  
146 acidic or basic perturbations, we observe little or no nitrate reduction in the CHL+ condition (flat red

147 lines, far left/right columns, Fig. 1B) indicating that there is little pre-existing functional biomass  
148 that can reduce nitrate under large pH changes. Second, we observed linear nitrate/nitrite reduction  
149 dynamics even in samples without chloramphenicol (CHL-) for pH perturbations around the native  
150 pH (black lines, Fig. 1B, Figs. S2, S3). Thus, near the native pH, after some early growth, the  
151 functional biomass stays constant even without the growth-inhibiting drug (CHL-), suggesting that  
152 the growth of the functional biomass is limited by nutrients other than nitrate (schematic, Fig. 1C).  
153 Third, when we perturb the pH above 8, we observe an initial lag of nitrate reduction, followed  
154 by an exponential increase in reduction rate (black lines, far right, Fig. 1B). This indicates that an  
155 initially rare population grows rapidly reducing all available nitrate.



**Figure 1: Soil microbiome metabolite and abundance dynamics under long and short-term pH variation.** (A) Schematic of the field sampling for soils with long-term pH variations (20 soils, pH 4.7 to 8.3, Cook Agronomy Farm, Pullman, WA, USA) and the experimental setup for imposing short-term pH perturbations in laboratory conditions. With each of the 20 soils, we created slurries (1:2 soil:water) amended with 2mM nitrate, adjusted to 13 different pH levels, and treated with (CHL+, no growth) or without chloramphenicol (CHL-, growth). ~1,500 microcosms are depicted in a grid of different pH conditions (perturbed pH vs. native pH) each condition in triplicates. Microcosms were incubated anaerobically for a 4-days while nitrate and nitrite were quantified colorimetrically. For metabolic dynamics, we measured nitrate ( $\text{NO}_3^-$ ) and nitrite ( $\text{NO}_2^-$ ) flux, the first two intermediates in denitrification and dissimilatory nitrate reduction to ammonium (DNRA). Communities were quantified by 16S rRNA amplicon sequencing before and after slurry incubation. (Continued)



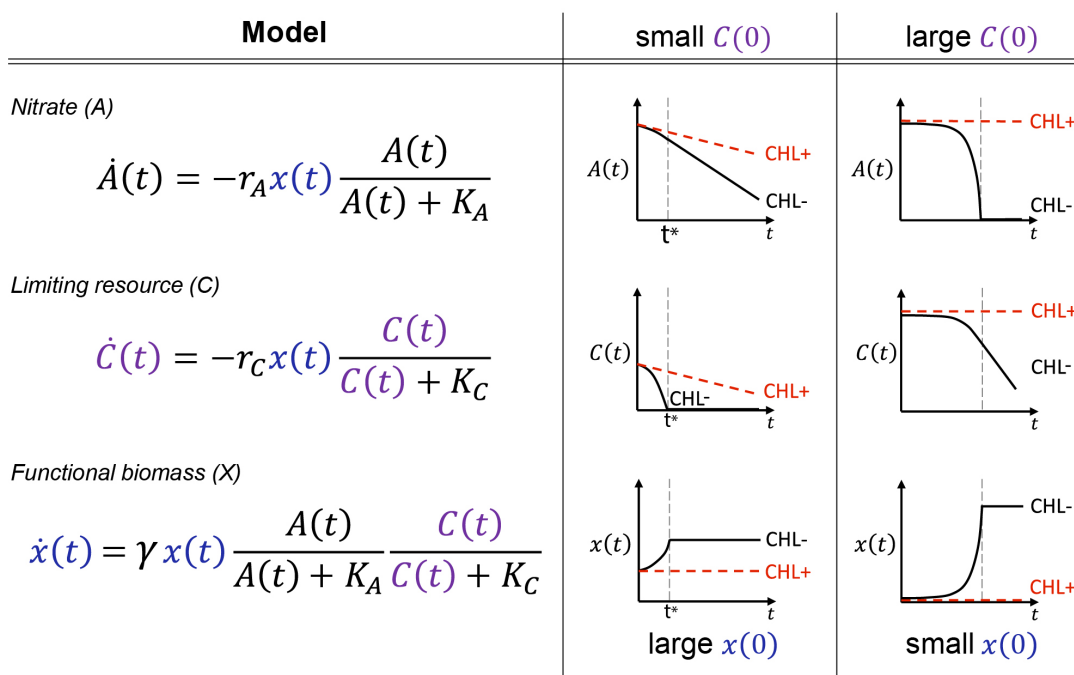
Figure 1: (Continued from the previous page) **(B)** A subset of nitrate concentration dynamics (function) during the 4-day anaerobic incubation: three topsoils with different native pH levels (rows) were perturbed to either acidic or basic pH (columns) at the start of the incubation ( $T_0$ ), all with (CHL+, red) and without chloramphenicol (CHL-, black) treatments in triplicates (see Methods). The pH indicated inside the panels is the stabilized end-point pH to which the slurries were perturbed (Methods). **(C)** Schematic depicting three different functional regimes that capture how the soil community responds to pH perturbations. With moderate pH perturbations, the functional response can be characterized as the Nutrient-limiting regime (Regime II), where nitrate utilization is performed by dominant taxa (blue) that utilize nutrients released from the soil matrix due to perturbation. Growth is limited by the amount of available growth-limiting nutrients (purple). During strong basic perturbations, growth-limiting nutrients are in excess, and rare taxa (green) rapidly outgrow dominant populations that cannot perform nitrate reduction in basic conditions, hence the Resurgent growth regime (Regime III). Strong acidic perturbations induce cell death that limits metabolic activity, resulting in an inactive state (Acidic death regime, Regime I). Functional biomass of the dominant (blue) and rare (green) taxa are shown by the lines below.

## 156 **Simple consumer-resource model captures metabolite dynamics across all pH** 157 **perturbations**

158 To describe the nitrate dynamics, we used the consumer-resource model presented in Fig. 2. Cru-  
159 cially, this model subsumes the ecological complexity of the soil microbiome into a single effective  
160 biomass rather than explicitly considering the multitude of possible interactions between taxa. The  
161 model has three variables: the functional nitrate-utilizing biomass ( $x$ ), nitrate concentration ( $A$ ),  
162 and the second growth-limiting nutrient ( $C$ ) whose existence we hypothesized above. The five  
163 model parameters include: consumption rates ( $r_A$  and  $r_C$ ), growth rate ( $\gamma$ ), and affinities ( $K_A$  and  
164  $K_C$ ). The consumption rate of a resource is determined by the amount of functional biomass ( $x$ )  
165 and per-biomass consumption rates. The biomass growth rate ( $\dot{x}$ ) is set to zero in CHL+ conditions  
166 due to chloramphenicol inhibition ( $\gamma = 0$ ,  $\dot{x} = 0$ ).

167 If the initial nutrient concentration  $C(0)$  is small (Fig. 2, middle column), the nutrient  $C$  runs  
168 out quickly, arresting biomass growth and resulting in  $A$  being consumed at a constant rate from  
169  $t^*$  onwards (dashed line). This recapitulates the late-time linear dynamics in CHL- conditions for  
170 moderate pH perturbations (Fig. 1B). In contrast, when the initial nutrient concentration  $C(0)$  is  
171 large (Fig. 2, right column), it is nitrate ( $A$ ) that runs out first. In this regime, the initial rate of  
172 nitrate utilization (determined by  $x(0)$ , the initial functional biomass) grows exponentially until  $A$   
173 runs out. Therefore, a small  $x(0)$  and a large  $C(0)$  recapitulates the initially slow but exponentially  
174 growing dynamics observed for large basic pH perturbations (Fig. 1B).

175 Our consumer-resource model provided a good fit to the observed nitrate dynamics in all soils  
176 (<10% error per data point, Fig. S7). To perform this fitting, we fixed the growth rate  $\gamma$  and  
177 the affinity parameters ( $K_A$ ,  $K_C$ ), and varied just two rescaled parameters:  $\tilde{x}(0) = x(0)r_A$  and  
178  $\gamma\tilde{C}(0) = \gamma C(0)r_A/r_C$  (see Methods). These parameters retain the same interpretation:  $\tilde{x}(0)$  re-  
179 flects the initial functional biomass, and  $\gamma\tilde{C}(0)$  the available limiting nutrient, the rescaling corre-  
180 sponds to measuring these quantities in terms of nitrate utilization rates (see Methods).



**Figure 2: Consumer-resource model captures metabolite dynamics** A mathematical representation of the consumer-resource model to fit the nitrate reduction dynamics of the community (**Model** column). The model describes the community through the total functional biomass ( $x$ , biomass) which describes the aggregated biomass of species that perform nitrate reduction, nitrate concentration ( $A$ , mM), and a limiting resource concentration ( $C$ , mM). Nitrate consumption rate ( $\dot{A}(t)$ ) takes a Monod [34] form with a reduction rate parameter ( $r_A$ , mM/biomass/day) and an affinity parameter ( $K_A$ , mM). Nitrite, which is reduced from nitrate, is not modeled. To capture linear nitrate dynamics, we include a non-substitutable resource that limits growth ( $C$ ) with Monod consumption function and parameters  $r_C$  (mM/biomass/day) and affinity parameter  $K_C$  (mM). Growth of functional biomass ( $\dot{x}(t)$ ) is determined by concentrations of nitrate ( $A$ ) and limiting nutrient ( $C$ ) with biomass growth rate ( $\gamma$ , 1/day). Plots in the right two columns show dynamics of  $x(t)$ ,  $A(t)$ , and  $C(t)$  at small  $C(0)$  and large  $x(0)$  (middle) and large  $C(0)$  with small  $x(0)$  (right). Red and black traces show dynamics with and without growth-inhibiting ( $\dot{x}(t) = 0$ ) chloramphenicol respectively. Without growth, the nitrate reduction rate is constant and proportional to the functional biomass  $x(0)$  (red lines, top row for large/small  $x(0)$ ). The **small C(0)** column illustrates how the model captures linear nitrate dynamics in chloramphenicol untreated (CHL-) conditions (black lines). With the small amount of initial limiting resource  $C(0)$ , functional biomass will stay constant after the limiting nutrient is depleted at  $t^*$ , which produces a constant nitrate reduction rate (linear  $\text{NO}_3^-$  dynamics, black line, top). The **large C(0)** column shows exponential nitrate depletion dynamics in CHL- conditions (black lines) when there is excess  $C(0)$  and  $x(0)$  is small. Functional biomass grows exponentially, resulting in exponential nitrate utilization dynamics (black line, top). The affinity parameters ( $K_A$ ,  $K_C$ ) and yield parameter ( $\gamma$ ) were fixed for all samples (see Methods for rationale and Fig. S8).

## 181 **Model reveals functional regimes**

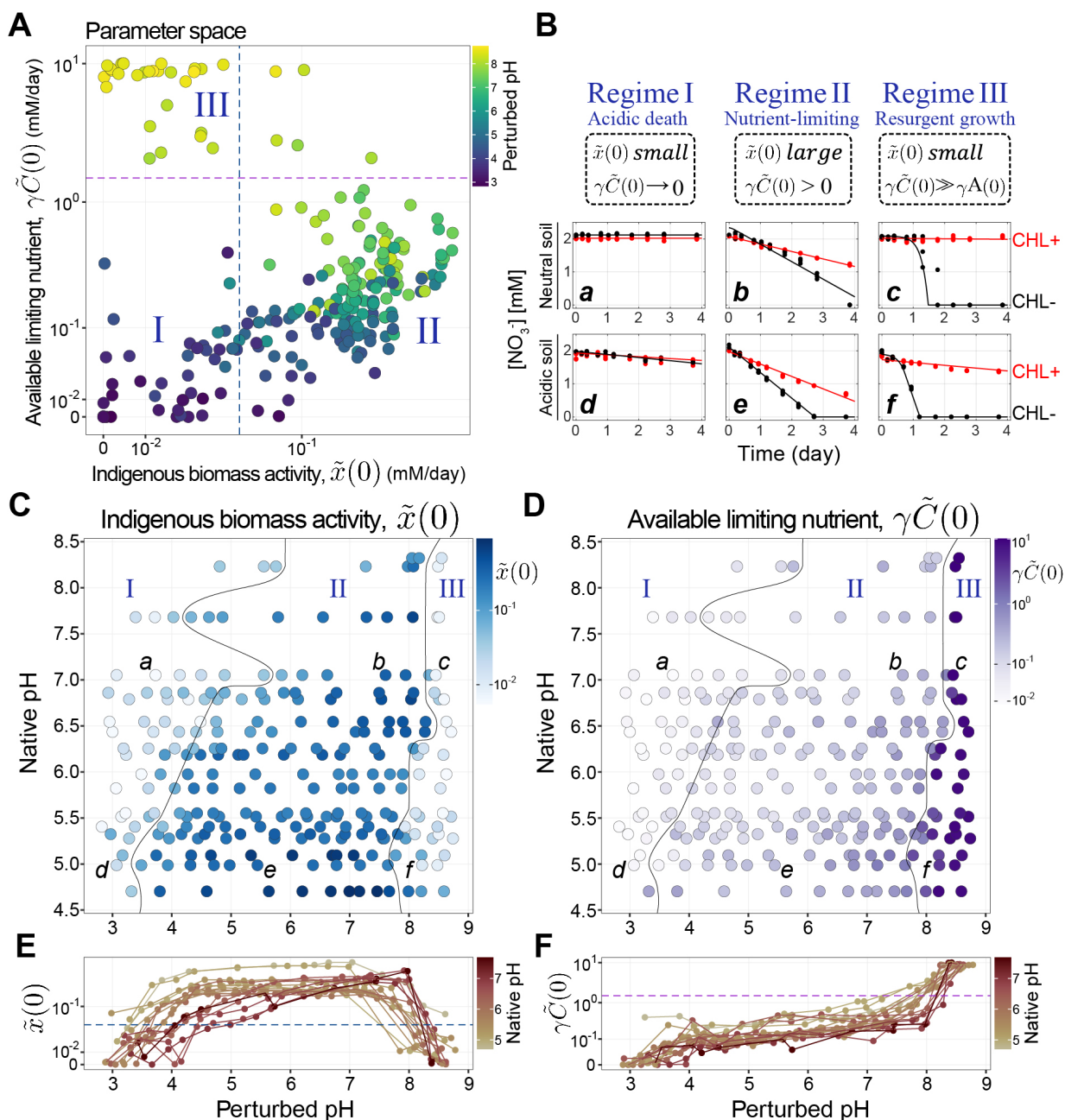
182 We plotted  $\tilde{x}(0)$  (pre-existing functional biomass) against  $\gamma\tilde{C}(0)$  (available limiting nutrient, Fig. 3A)  
183 and identified three regimes of nitrate utilization dynamics (Methods, Fig. S9). Regime I, the Acidic  
184 death regime (both  $\tilde{x}(0)$  and  $\gamma\tilde{C}(0)$  are low) is observed for  $\text{pH} \lesssim 4$ , and shows little to no nitrate  
185 reduction (Fig. 3B, (a) and (d)). Regime II, the Nutrient-limiting regime ( $\tilde{x}(0)$  is large and  $\gamma\tilde{C}(0)$   
186 is small) is observed for  $4 \lesssim \text{pH} \lesssim 8$ , and exhibits a relatively large initial nitrate reduction rate  
187 that transiently increases and then remains constant (Fig. 3B, (b) and (e)). Finally, Regime III, the  
188 Resurgent growth regime (small  $\tilde{x}(0)$ , large  $\gamma\tilde{C}(0)$ ) is observed for  $\text{pH} \gtrsim 8$ , and displays a close-  
189 to-zero initial utilization rate, followed by an exponential speed-up that continues until nitrate is  
190 depleted (Fig. 3B, (c) and (f)).

191 We observe all three functional regimes across all soils, but the pH at which a transition from  
192 one regime to another occurs depends on the native pH of the soil. Figure 3C&D shows the inferred  
193 initial functional biomass ( $\tilde{x}(0)$ ) and limiting nutrient ( $\gamma\tilde{C}(0)$ ) across soils of varying native pH (y-  
194 axis) and laboratory perturbed pH (x-axis). We next harnessed our model to identify mechanisms  
195 underlying these regimes.

## 196 **Metabolite dynamics in Regime II are governed by carbon release**

197 In Regime II (the Nutrient-limiting regime), the nitrate reduction rate increases with pH (Fig. 1B).  
198 Our model proposes that the mechanism behind this increase is the increasing availability of the  
199 growth-limiting nutrient (Fig. 3F), which translates into larger growth of active biomass and hence  
200 the increased nitrate reduction rate (Fig. 2). Here, we investigate whether this model prediction  
201 is valid by examining how increasing pH leads to higher levels of growth-limiting nutrients and  
202 identifying these nutrients.

203 Previous studies have observed that increasing pH can enhance the availability of organic car-  
204 bon in soils [35–37]. Studies indicate that this release of nutrients from soil is based on a substitu-  
205 tion mechanism at the ion exchange sites within the soil clay particles [38, 39] (Fig. 4B, detailed



**Figure 3: Conserved regimes capture soil's functional response to pH perturbations.** (A) Scatterplot of the two model parameters (functional biomass  $\tilde{x}(0)$ , limiting nutrient concentration  $\gamma\tilde{C}(0)$ ) inferred from nitrate dynamics across all samples. See text and Methods for details of model fitting. Note log-scale. The color of points indicates each sample's perturbed pH. Three regions separated by dashed lines indicate the distinct regimes of functional response against pH perturbations: the Acidic death regime (Regime I), Nutrient-limiting regime (Regime II), and Resurgent growth regime (Regime III). The locations of the dashed lines were determined by thresholding distributions of  $\tilde{x}(0)$  and  $\gamma\tilde{C}(0)$  (see Methods). (B) Example nitrate dynamics for each of the three regimes for a neutral soil (top row) and an acidic soil (bottom row). (Continued)

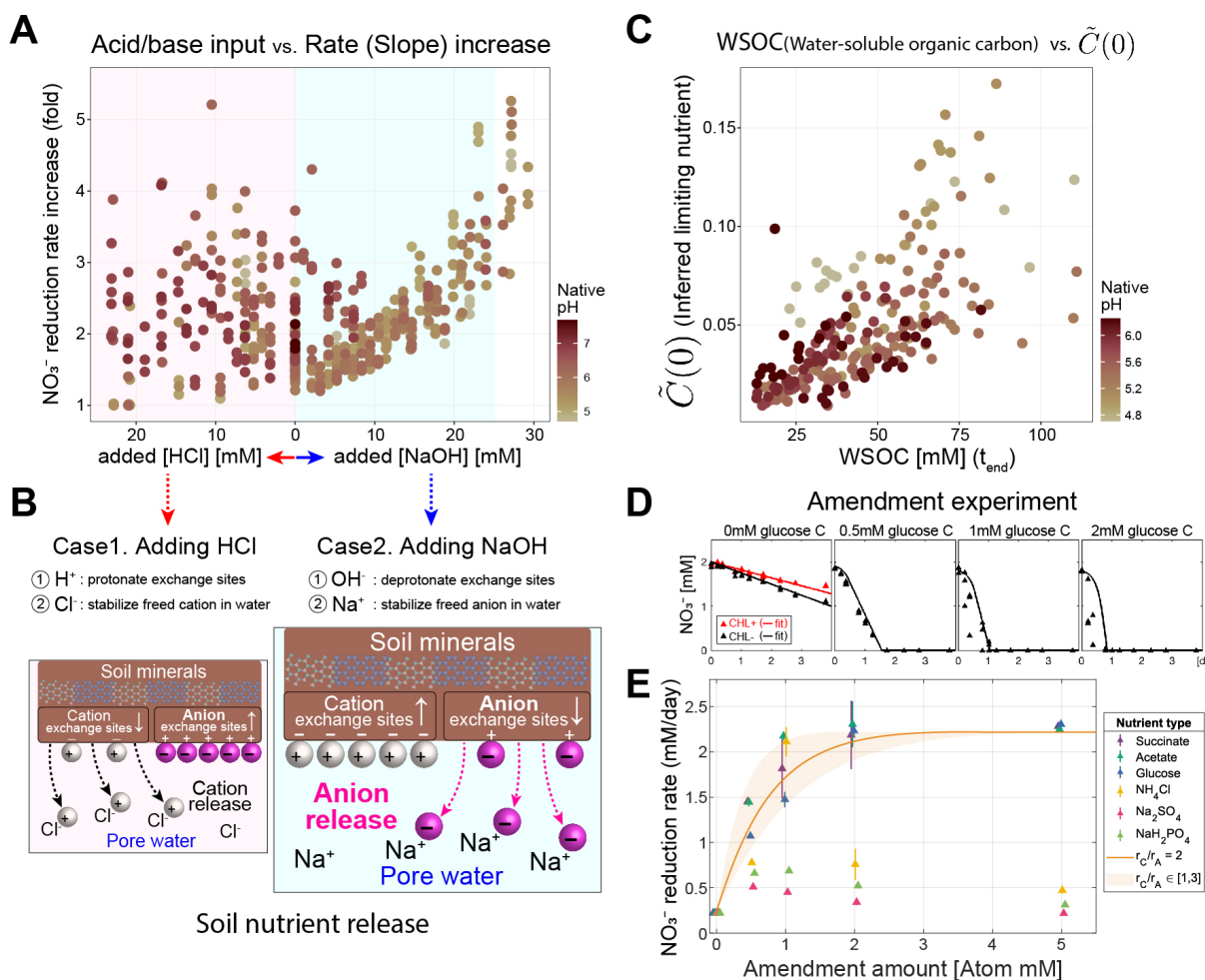
Figure 3: (Continued from the previous page) Red lines are with growth-arresting chloramphenicol and black without. (*a, b, c, d, e, f* correspond to perturbed conditions indicated in panels **C and D**). *a, d* show little nitrate reduction, *b, e* show linear nitrate dynamics with slopes that increase without chloramphenicol (see Fig. 2, middle column), and *c, f* show no activity without growth (red) but exponential nitrate utilization in the absence of the drug (Fig. 2, right column). (**C and D**) pH affects indigenous biomass activity  $\tilde{x}(0)$  (blue) and available limiting nutrient  $\gamma\tilde{C}(0)$  (purple). Fitted parameter values are shown with the color (log-scale) in the grid of long-term pH variation (y-axis, Native pH) and short-term pH perturbation (x-axis, Perturbed pH). In all soils from different native pH levels, we observe a conserved set of responses to short-term pH perturbations: Nutrient-limiting regime (region indicated by II) near the native pH, then transitioning to the Acidic death regime (region indicated by I) during acidic perturbation (black line), also transitioning to the Resurgent growth regime (III) for basic perturbations (black line). Long-term (native) pH dictates the pH thresholds of regime boundaries (black line). (**E**) Trends of  $\tilde{x}(0)$  (log-scale) across varying perturbed pH values for soils with different native pH levels (native pH indicated by line color), demonstrating consistent transition between regimes and a plateau of high activity within the mid-range pH (Regime II) across all soils. (**F**)  $\gamma\tilde{C}(0)$  (log-scale) with perturbed pH, showing a rise in limiting nutrients induced by short-term pH increases. Colors indicate native pH. We used the median fitted value of the three biological replicates for all data points of  $\tilde{x}(0)$  and  $\gamma\tilde{C}(0)$ .

206 mechanism in SM). Therefore, we hypothesized that the amount of nutrients released would be  
207 proportional to the quantity of either base (NaOH) or acid (HCl) added to the slurry. Based on  
208 this assumption, the fold change in nitrate reduction rate, reflecting the growth of active biomass  
209 limited by this nutrient, should be proportional to the quantity of acid or base added to the system.  
210 In Fig. 4A, we observe precisely this trend across all soils, as evidenced by a data collapse in the  
211 increase in nitrate reduction rate with NaOH (light blue region). The trend is specific to Regime II  
212 (Fig. S11A), and if the data are plotted against pH, the correlation becomes much weaker (Fig. S10).  
213 As further evidence supporting our hypothesis, we measured increases in the absolute abundances  
214 via 16S rRNA amplicon sequencing with internal standards (see Methods). Increasing sequencing  
215 reads reflect increases in biomass, both at a coarse level (fold change in total biomass, Fig. S11B)  
216 and fine level (individual ASVs that responded to the amendment of nitrate, Fig. S11C, see SM  
217 for details). Corroborating our hypothesis, we observe a linear relationship between the increase in  
218 absolute abundances and the amount of NaOH added to the system (Fig. S11).

219 The asymmetric response of the change in nitrate reduction rate upon the addition of NaOH  
220 rather than HCl (blue versus pink shaded regions Fig. 4A) provides insight into the identity of the  
221 released nutrient. Under the mechanism of ion-exchange-mediated nutrient release, adding ions

222 releases nutrients adsorbed to the clay particles into the pore water, making them accessible to  
223 microbes (Fig. 4B). HCl and NaOH will release cationic and anionic nutrients respectively (SM  
224 for details, Fig. S13B). Our observation that the limiting nutrient governing Regime II dynamics is  
225 released in proportion to the amount of NaOH indicates that the growth-limiting nutrient is anionic,  
226 with likely candidates including phosphates, sulfate, or carbon. Notably, measurements of water-  
227 soluble organic carbon (WSOC) at the endpoint increased linearly with NaOH added (Fig. 4C).  
228 This suggests that some WSOC is negatively charged (anionic) and that the growth-limiting nutrient  
229 might be WSOC, or concomitantly released nitrogen (N), sulfur (S), or phosphorus (P).

230 To further identify the limiting anionic nutrient, we performed an amendment experiment on  
231 a representative soil (Soil 6 (pH 5.4), see Methods). We amended a soil slurry without perturb-  
232 ing pH with glucose (neutral), succinate (anion when  $\text{pH} > \text{pK}_a = 4.2$ ), acetate (anion when  
233  $\text{pH} > \text{pK}_a = 4.75$ ), phosphate (anion), ammonium (cation), and sulfate (anion) added in varying  
234 concentrations (Methods, Fig. S14). We found that the amendment of carbon, but not other N,  
235 S, and P sources, immediately increased the nitrate reduction rate, changing the linear dynamics  
236 to exponential (Fig. 4D & E), indicating that carbon was the limiting nutrient. With a single free  
237 parameter, our model predicted the nitrate utilization dynamics in a soil amended with glucose  
238 (Fig. 4D). Similar results are found for other carbon sources, but not sources of N, S, or P (Fig. 4E).  
239 The single free parameter is the ratio  $r_C/r_A$ , which can be interpreted as a stoichiometry of carbon  
240 to nitrate utilization (Fig. 2). We find this ratio to be highest for glucose (2.5) and lowest for ac-  
241 etate (1), suggesting carbon is utilized more quickly relative to nitrate in glucose amendments. The  
242 relatively more rapid utilization of glucose may be because glucose can be consumed by anaerobic  
243 respiration (requiring nitrate) and fermentation, whereas acetate is not fermentable. The amend-  
244 ment experiment confirms the mechanism predicted by our model, that a nutrient other than nitrate  
245 limits reduction dynamics for modest pH perturbations. Critically, this insight emerged naturally  
246 from our mathematical description of the nitrate utilization dynamics across pH perturbations.



**Figure 4: Carbon limits growth in Regime II and is released by ion exchange mechanism** (A) The amount of NaOH added to the soil plotted against the fold-change of nitrate reduction rate (ratio of rate with growth (CHL-) and with no growth (CHL+)). Base additions from 0mM to 25mM NaOH correspond to soils in the Nutrient-limiting regime (Regime II, Fig. 3) (light blue background in A). No increase in growth was observed for acidic perturbations (>0 mM HCl addition, pink region, Fig. S11A). (B) Cartoon illustrating the mechanism of soil nutrient release hypothesis; NaOH results in the release of anionic nutrients (magenta-colored spheres) from soil particles (brown region), while the addition of HCl would release cationic nutrients (white spheres) and adsorbs anionic nutrients. Microbes cannot access the nutrients adsorbed in the soil particles but can access the nutrients dissolved in pore water. Added OH<sup>-</sup> ions decrease the number of anion exchange sites in the soil particles, releasing anionic nutrients. In concert, Na<sup>+</sup> ions stabilize the released anions (see text and Fig. S13B for additional details). A and B suggest the growth-limiting nutrients are anionic (negatively charged). (C) Scatterplot of model-inferred  $\tilde{C}(0)$  (available limiting nutrient) and measured water-soluble organic carbon (WSOC) measured via a chromate oxidation assay (Methods) that is not chemically specific and WSOC likely contains different C compounds, N, P, etc. Data points are from Soil 1–12 samples where there are enough number of points per soil to observe a linear relationship in the light blue region in A (0mM–25mM NaOH, ). (Continued)



Figure 4: (Continued from the previous page) **(D) and (E)** Amendment experiments for soil in the Nutrient-limiting regime (Regime II) at unperturbed pH. **(D)** Panels show nitrate dynamics with different levels of glucose amendments (red: CHL+, black: CHL-, points: data, lines: model fit), where linear dynamics (at 0mM C) transition into exponential dynamics ( $\geq 0.5\text{mM C}$ ) supports carbon limitation of nitrate utilization. Lines are model predictions. **(E)** Nitrate reduction rates after amending soils with different concentrations of nutrients (three carbon sources, ammonium, sulfate, and phosphate). Points are the mean rates, estimated by linear regression, of triplicates with error bars indicating standard deviation. Carbon (succinate ( $\text{pK}_a = 4.2$  and  $5.6$ ), acetate ( $\text{pK}_a = 4.75$ ), and glucose) amendments increased the nitrate reduction rates starting from low concentrations ( $0.5\text{CmM}$ ). Carbon compounds are negatively charged when  $\text{pK}_a < \text{pH}$  (here, the pH of soil 6 is 5.4). Ammonium, sulfate, and phosphate did not result in a similar increase in nitrate reduction. We cannot independently infer the ratio  $r_A/r_C$  (Methods), model predictions are shown for  $1 < r_A/r_C < 3$  (shaded region) with a line for  $r_A/r_C = 2$  (best fit). This ratio can be interpreted as the nitrate:carbon utilization ratio.

## 247 **Regime III arises from the rapid growth of rare taxa**

248 Under large basic pH perturbations, all soils exhibited a sharp transition from linear to exponential  
249 nitrate consumption dynamics. Our model fits suggest interpreting these metabolite dynamics as re-  
250 sulting from a small initial functional biomass ( $\tilde{x}(0)$ ) undergoing exponential growth due to excess  
251 nutrient  $\gamma\tilde{C}(0)$  (Fig. 3C-F). To test this interpretation, we used the sequencing data to investigate  
252 the compositional changes that occur after large basic perturbations (Regime III).

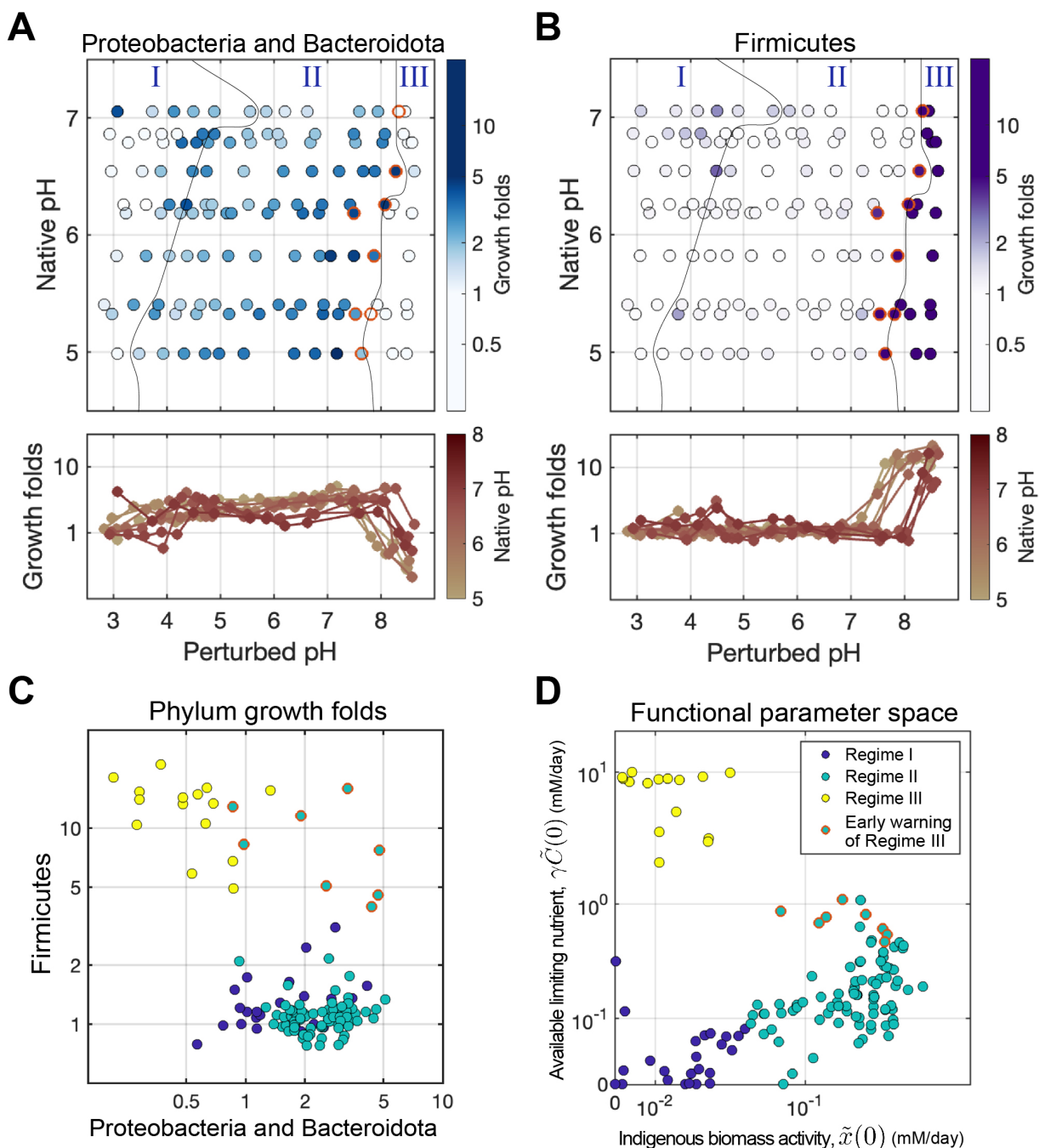
253 Sequencing measurements corroborate our model predictions by revealing a group of rare taxa  
254 enriched in Regime III. These are especially clear if ASVs are grouped at the phylum level, re-  
255 vealing that Firmicutes undergo explosive growth in this regime (10-fold enrichment at the aggre-  
256 gate phylum level, and several hundred-fold for individual ASVs, particularly in the Bacilli genus;  
257 Fig. S17). We computed the fold change of each phylum's absolute abundance across treatments  
258 relative to the no-growth CHL+ control. Non-negative matrix factorization (NMF) analysis of the  
259 growth fold values revealed that most of the variation in these data could be captured with just two  
260 axes of variation (Fig. S15B, Methods). Each of these axes was composed of one or two phyla, one  
261 included Proteobacteria and Bacteroidota, and the other Firmicutes.

262 Fig. 5A-B shows growth-folds for the two groups of phyla identified by NMF that dominate  
263 growth across all soils and pH conditions. In the Nutrient-limiting regime (Regime II), Proteobac-  
264 teria and Bacteroidota increased their growth with increased pH, then decreased towards the start of  
265 Regime III. This matches the growth behavior of indigenous functional biomass ( $\tilde{x}(0)$ ) revealed by  
266 the model in Regime II (Fig. 3C). Conversely, Firmicutes did not grow until a critical pH threshold  
267 between 7-8.5, which matches the onset of exponential nitrate utilization dynamics in Regime III  
268 (Fig. 3F). Importantly, the boundary between Regime II and III derived from the functional dy-  
269 namics data (Fig. 3 C & D), aligns with the shifts in growth responses of Firmicutes (Fig. 5B) and  
270 Proteobacteria/Bacteroidota (Fig. 5A). These growth patterns suggest that the changes in the iden-  
271 tity of the phyla responsible for nitrate reduction reflect the functional regimes. A more detailed  
272 analysis of the likely metabolic traits of these strains [40] suggests that the transition from Regime

273 II to III is also accompanied by a shift from denitrification to DNRA which agrees with the fact that  
274 excess carbon favors DNRA [41](Figs. S18 S19).

### 275 **Growth is an early-warning indicator of a transition between regimes**

276 Intriguingly, we found that Firmicutes begin increasing at pH levels just below the transition from  
277 Regime II to III, thereby acting as ‘early warning indicators’ for the impending transition (red cir-  
278 cles, Fig. 5D). Specifically, when we plot the growth folds of the Firmicutes versus Proteobacteria  
279 and Bacteriodota, we find that Firmicutes abundances begin to rise prior to the system entering  
280 Regime III as defined by nitrate utilization dynamics (Fig. 5C). This finding indicates that com-  
281 positional data can be used to predict impending functional state transitions during environmental  
282 perturbations.

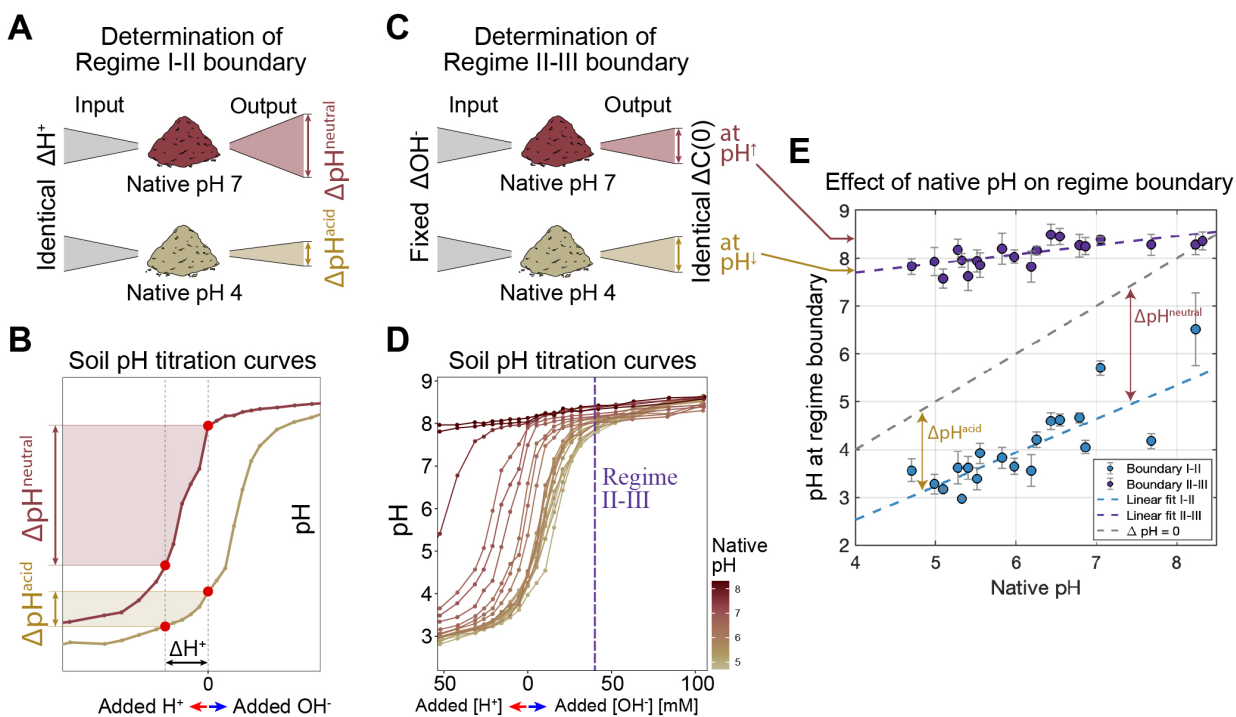


**Figure 5: Regime III: Resurgent growth emerges from native population decline and rare taxa expansion.** Global trends of growth in the phylum level across perturbed pH levels reveal taxonomic origins of the rapid growth in the resurgent growth regime (Regime III). 16S amplicon sequencing at the end of each incubation was used to identify amplicon sequence variants (ASVs) in CHL+/- conditions (Methods). ASVs were aggregated at the phylum level. For each phylum, a growth fold was computed as the ratio of abundances with/without growth ( $Abs_{CHL-}/Abs_{CHL+}$ ). A statistical decomposition across all conditions identified three phyla that dominated abundance changes due to growth: Proteobacteria and Bacteroidota with similar changes, and Firmicutes. (see main text and Fig. S15). (Continued)

Figure 5: (Continued from the previous page) **(A)** Growth folds for the phyla Proteobacteria + Bacteroidota (combined abundance) indicated by the color for each native and perturbed pH condition. The growth declines at a basic pH threshold, mirroring the patterns observed in the fitted model parameter  $\tilde{x}(0)$  (indigenous biomass activity, Fig 3C) at the Regime II-III boundary). Regime boundary lines are those determined in Fig. 3). Line plots (lower panel) growth folds were plotted in log-scale, color indicating native pH given in color bar. **(B)** Identical to (A) but showing growth folds (color) of Firmicutes increasing during the transition from Regime II to III, where pH perturbations are strongly basic. This mirrors the increase in inferred carbon concentrations  $\gamma\tilde{C}(0)$  (Fig. 3). **(C)** Scatter plot of growth folds of Proteobacteria + Bacteroidota against Firmicutes. Points marked in red, associated with Regime II (also red in (D)), exhibit high growth of both Proteobacteria + Bacteroidota and Firmicutes. For red points, Firmicutes abundances are an early-warning indicator of a transition between regimes. **(D)** Same plot as Fig. 3A of  $\tilde{x}(0)$  versus  $\gamma\tilde{C}(0)$  with points marked by the regime they belong to and red points indicating Regime II conditions near the boundary between Regime II and III. Note these red circles are in Regime II, but the Firmicutes abundances are high (panel (C)).

## 283 **Acidic perturbations in Regime I reduce functional biomass via death**

284 In response to a short-term decreases in pH, the model indicates a reduction in indigenous func-  
285 tional biomass ( $\tilde{x}(0)$ ) and a decrease in the availability of limiting nutrients (Fig. 3). Below a pH  
286 value of 3–5, depending on the soil’s native pH, nitrate reduction ceases (Regime I). We tested  
287 whether the sequencing data reflects the decreasing trend of functional biomass ( $\tilde{x}(0)$ ) with pH.  
288 We computed the fold-change in each Phylum’s endpoint absolute abundance in CHL+ conditions  
289 relative to abundances at the initial time point  $T_0$  (‘survival fold’; Fig. S20A). This ratio reflects  
290 the change in abundance in the absence of growth, hence we regard this as a proxy for death.  
291 For all phyla except the Firmicutes, we observed a consistent drop of survival folds during acidic  
292 perturbations (Fig. S20A). Furthermore, we confirmed that the survival folds exhibited an approxi-  
293 mately linear relationship with the  $\tilde{x}(0)$  values (Fig. S20B). These observations confirm the decline  
294 of biomass in acidic conditions, likely via death and DNA degradation, except in taxa tolerant to  
295 short-term pH changes (Firmicutes, Fig. S20A). Thus, we conclude that acidic perturbations lead  
296 to widespread death, while basic conditions lead to selective growth (Fig. S15C).



**Figure 6: Regime boundaries are determined by long-term pH and history of pH variation.** (A) Cartoon for how native pH impacts the pH of the Regime I and II boundary. The cartoon depicts how the identical amount of acid perturbations ( $\Delta H^+$ ) gives rise to larger changes in pH for neutral soils ( $\Delta pH^{neutral}$ ) than acidic soils ( $\Delta pH^{acidic}$ ) which arises due to differences in the location of the native pH on the titration curve shown in (B). (B) Shows titration curves where the pH (y-axis) is measured after adding different amounts of acid or base (Methods) for neutral soil (dark brown) and acidic soil (light brown). The dashed vertical line at 0 indicates the pH with no acid/base perturbation. Due to the shape of these curves, if both soils are subjected to the same  $\Delta H^+$  (bottom) the neutral soil experiences a larger change in pH (shaded regions). This suggests that acidic soils experience smaller pH fluctuations and therefore transition to Regime I from II after smaller pH perturbations as shown in (E). (C) Cartoon for how native pH determines the pH of the Regime II and III boundary. The cartoon depicts how the fixed amount of added base (NaOH) results in an identical amount of released carbon ( $\Delta C(0)$ , Fig. 4). Large  $C(0)$  drives the transition from Regime II to III. For a fixed addition of NaOH, more neutral soils reach higher pH again due to the shape of the titration curves as shown in (D). (Continued)

Figure 6: (Continued from the previous page) **(D)** Soil pH titration curves (identical to **C**) for all soils with different native pH levels. The vertical dashed line indicates the quantity of NaOH added to move from Regime II-III. More neutral soils (darker colors) reach higher pH values for this fixed quantity of added NaOH. This correlates with the increasing pH at the Regime II-III boundary (purple points, **(E)**). **(E)** pH levels (y-axis) when transitions between functional regimes occur from Regime II to I (blue points) and from Regime II to III (purple points) for soils from different native pH levels (x-axis). Regime boundaries are determined as the midpoint between the last pH perturbation in Regime I and the first in Regime II. Error bars represent the pH difference between these conditions. An identical strategy was used for Regime II-III. The dashed blue line (Regime I-II boundary) and dashed purple line (Regime II-III boundary) are weighted least squares fits, with the weights inversely proportional to the error of each point. The dashed black line is slope 1, where the change in pH from native to the regime boundary is constant for all soils. Lines with a slope different from 1 indicate that the difference between native pH and pH at the regime boundary depends on the native pH of the soil. The slope of the blue dashed line is 0.7 (95% confidence interval: [0.44, 0.97]).



## 297 **Long-term soil pH defines regime boundaries**

298 Next, we sought to understand what properties determine the pH at which soils transition between  
299 regimes. We observed that the native pH of the soil (long-term pH) determined the pH at which  
300 any given soil transitioned between functional regimes (Fig. 3C & D). This result suggests that the  
301 soil communities are adapted to their long-term pH conditions [19, 33, 42].

302 One key property of soils that impacts the pH variation the microbiome experiences is the  
303 soil's pH titration curve: how soil pH changes in response to acid/base additions. The shape of  
304 the titration curve was similar across all soils (Fig. 6D, Fig. S23A), showing a plateau at low and  
305 high pH with a nonlinearity in between. As a result, acidic soils with native pH near the lower  
306 plateau were more strongly pH-buffered than the neutral soils (with native pH around the steepest  
307 portion of the nonlinearity; Fig. 6B, Fig. S23A). This observation indicates that at similar levels of  
308 acid addition, neutral soils would experience *a larger drop in pH* than acidic soils ( $\Delta\text{pH}^{\text{acidic}} <$   
309  $\Delta\text{pH}^{\text{neutral}}$ , Fig. 6A, B). We speculate that this makes communities in acidic soils less tolerant of  
310 acidic pH fluctuations, as they are less likely to experience large reductions in pH. This reasoning  
311 would help explain the observation that acidic soils transition from regime II to I after a smaller  
312 perturbation in pH than neutral soils (Fig. 6E,  $\Delta\text{pH}^{\text{acidic}} < \Delta\text{pH}^{\text{neutral}}$ ). As a result, plotting the  
313 pH at the regime II to I transition against the native pH gives a line with slope  $< 1$  (Fig. 6E, bottom  
314 dashed line), where a line of slope 1 would indicate that entry into Regime I requires an acidic pH  
315 shift of a constant magnitude.

316 In contrast, we find that soils transition from Regime II to III when carbon is in excess. From  
317 Fig. 4, we know that carbon is released in proportion to the NaOH added to the slurry (Fig. 6C).  
318 Accordingly, we find that a constant addition of NaOH drives the transition from Regime III to II  
319 (Fig. S24). However, due to the shape of the titration curves as seen in Fig. 6D, for a constant base  
320 amendment, more neutral soils reach higher pH (dashed line Fig. 6E). Therefore, as expected from  
321 the titration curves, more basic soils transition to Regime III at higher pH (Fig. 6E).

322 Our sequencing data support the idea that variation in regime boundaries with native pH has a

323 basis in the taxonomic composition of the microbiome. In more acidic soils, the Proteobacteria and  
324 Bacteroidota show better survival at lower pH (Fig.S21). In contrast, the pH at which Firmicutes  
325 begin to grow in Regime III rises with the soil's native pH (Fig. 5B), in line with the Regime II to  
326 III transition observed in functional measurements (Fig.6A). In addition, the native pH of the soil is  
327 predictable from the identity of the strains that exhibit growth in regime III (Fig. S17A, S22). These  
328 findings suggest that prolonged exposure to a specific pH likely selects for specific taxa, thereby  
329 influencing the pH at which the community transitions between functional regimes.

## 330 **Discussion**

331 We showed that a simple mathematical model derived from quantitative measurements of metabo-  
332 lite fluxes delineates which mechanisms are relevant for understanding the functional response of  
333 the soil to perturbations. Remarkably, the model does not attempt to account for all processes in  
334 the soils and instead captures the behavior of the entire community using a single effective biomass  
335 subjected to nutrient limitation. From this perspective, we identified functional regimes demar-  
336 cated by whether active biomass or available nutrients dictate the metabolism of the system. For  
337 example, we discovered a nutrient-limiting regime (Regime II), where the indigenous biomass is  
338 robust to moderate pH changes [43] and metabolism is governed by carbon limitation. In contrast,  
339 metabolism in Regime III is governed by the growth of initially rare taxa perturbations [44].

## 340 **Limitations of the study**

341 Our study has several limitations. First, our soil slurries do not capture the full complexity of natural  
342 soils. Microcosms experience fixed anaerobic conditions, but nitrate utilization in the wild occurs  
343 during fluctuations between aerobic and anaerobic conditions [45]. Second, our more extreme pH  
344 perturbations ( $\Delta pH > 2$ ) are larger than is routinely experienced in natural systems. Third, unlike  
345 previous studies, we do not quantify nitrous oxide and nitrogen gas (Table S2) production both  
346 downstream products of denitrification. Nitrous oxide is of critical interest given its importance as  
347 a greenhouse gas, so it will be important to understand how its production varies across functional  
348 regimes.

349 Finally, the simplicity of our model, which describes a single effective biomass, leaves open  
350 the question of what role ecological interactions play in determining community metabolism. It is  
351 unclear within a given regime whether there are strong interactions between responding taxa or not.  
352 At the transition between regimes II and III, we cannot determine if the Firmicutes outcompete the  
353 Regime II taxa for carbon or nitrate or whether the physiology of taxa that dominate in Regime II  
354 does not permit their growth in more alkaline conditions.

## 355 **The significance of functional regimes and their generalizability**

356 Our study establishes the existence of functional regimes where specific chemical, physiological,  
357 or ecological processes govern system response. This demonstrates that understanding the commu-  
358 nity response to perturbation may not require grappling with every metabolic process or interaction  
359 in the community, but only with a handful of key features. Our demonstration comes in the context  
360 of nitrate utilization and soil pH. However, this study opens the door to asking whether similar  
361 functional regimes describe community response to a suite of key perturbations including temper-  
362 ature or xenobiotics. A previous study of the response of soils to temperature revealed dynamics  
363 strikingly similar to Regime III at high temperatures and the asymmetric response in the Acidic  
364 death regime (Regime I) at low temperatures [46].

## 365 **Functional regimes as guides for understanding complex omics data**

366 Sequencing measurements of complex microbiomes result in datasets with thousands of variables -  
367 taxa, genes, or transcripts. Distilling some understanding from these data presents a huge challenge.  
368 The existence of regimes guided our understanding of the dynamics of the >30,000 ASVs in our  
369 dataset by directing us to look for specific responses.

370 More broadly, the last decade has seen an explosion of methods for quantifying community dy-  
371 namics and metabolism from transcriptomic and metagenomic measurements [47–49] to single-cell  
372 metabolomics [50] and quantitative stable isotope probes [51–53]. The challenge is to synthesize  
373 these data to achieve insights into dynamics and function. Our work illustrates the promise of an  
374 approach where we acquire large-scale quantitative measurements of metabolism at the whole com-  
375 munity level, describe these dynamics mathematically, either phenomenologically or potentially  
376 new AI-driven methods [54], and then interpret the resulting model mechanistically. For example,  
377 in Regime III, we expect native taxa to exhibit stress response and declining metabolic activity, and  
378 the converse for Firmicutes. Thus the framework of regimes suggests a route for leveraging new  
379 technologies for a deeper understanding of mechanisms in complex microbiomes.

## 380 **Physiological insights from constant utilization rates in nutrient-limited envi-** 381 **ronments**

382 The linear dynamics of nitrate utilization observed in Regime II have been previously observed [11,  
383 31, 55–57] and attributed to carbon limitation [31]. Moreover, previous work supports the result  
384 that available organic carbon can be the limiting factor for nitrate utilization [10, 36, 55, 58–61].  
385 How can limited carbon lead to a constant rate of nitrate reduction? Carbon is the electron donor  
386 for anaerobic respiration of nitrate which is the terminal electron acceptor. If carbon runs out we  
387 might expect that cells will run out of reductant to convert nitrate to nitrite, but this is not what we  
388 observe. One hypothesis is that cells internally store carbon to regenerate reductant [62]. To test this  
389 hypothesis, we incubated individual denitrifying bacterial strains in minimal media supplemented  
390 with 2mM  $\text{NO}_3^-$  in the absence of exogenous carbon. Similar to the linear dynamics observed in the  
391 soil microcosm, we observed linear nitrate reduction dynamics in the carbon-limited monocultures  
392 (Fig. S6), revealing that the metabolism of a single strain can mirror the metabolism of the soil  
393 microbiome. The energy obtained from a constant rate of nitrate reduction is likely channeled  
394 to maintenance rather than growth. More broadly, the discovery of the three functional regimes,  
395 including the nutrient-limiting regime, is notable because it reflects a potential duality between  
396 the physiology of an ecosystem and the three phases of a cell: growth (Regime III), stationary  
397 (Regime II), and stress (Regime I). This duality suggests the possibility that cellular physiology  
398 might provide a conceptual framework for understanding the ecosystem.

## 399 **Unifying decades of prior work in a quantitative framework**

400 To limit confounding factors, our study focused on 20 samples from a single agricultural site. How-  
401 ever, the features of the three functional regimes defined here are present in many previous studies  
402 performed on soils from other sites, suggesting that these regimes are a general feature of nitrate  
403 utilization and pH perturbations. For example, Nömmik observed metabolite dynamics consistent  
404 with a transition from Regime II to III [63]. Parkin *et al.* observed a native pH dependent  $\tilde{x}(0)$  with

405 chloramphenicol applied [33], consistent with all three regimes. Simek observed increasing nitrate  
406 utilization rates with time as pH increased, another Regime II to III transition [64]. Anderson *et*  
407 *al.* observed increasing rates of nitrate utilization with increasing pH, and the recruitment of Fir-  
408 micutes in very basic conditions [35]. These results show that the regimes are potentially general  
409 and not specific to our study site or protocol.

#### 410 **Direct and indirect effects of pH perturbations**

411 In the context of community metabolism, it has been debated whether the indirect effect of pH on  
412 nutrient availability is as important as the direct effect of pH on microbial physiology [17]. Our  
413 results answer this question because the model enables us to quantify both the pH's indirect effect  
414 on the growth-limiting nutrient (changes in  $\tilde{C}(0)$ ) and its direct effect on indigenous functional  
415 activity that reflects physiology (changes in  $\tilde{x}(0)$ ). In the Nutrient-limiting regime (Regime II), the  
416 indirect pH effect (changes in  $\tilde{C}(0)$ ) is more important in determining the nitrate reduction rate  
417 because the indigenous functional biomass ( $\tilde{x}(0)$ ) is stable in this regime. In Regime III the model  
418 suggests physiological responses might be most important since nutrient limitation is relieved but  
419 only a small fraction of taxa grow.

#### 420 **Optimal pH and long-term adaptation**

421 Due to the agricultural importance of nitrate utilization, it has been debated whether soils exhibit an  
422 optimal pH for denitrification [64]. Previous studies have demonstrated the pH level associated with  
423 the highest rate of denitrification closely aligns with the native soil pH [33, 64] over short timescales  
424 (<3 hours, Table S2). Other studies observed a shift of optimum pH to more neutral values on  
425 longer timescales [64]. Our study reconciles these outcomes and elucidates the underlying cause.  
426 The fastest nitrate utilization occurs near the native pH of the soil on short timescales (Fig. 1B).  
427 This is consistent with our results because the pre-existing functional biomass ( $\tilde{x}(0)$ ) is the largest  
428 near the native pH (Fig. 3C&E). However, basic pH perturbations release carbon ( $C(0)$ ), driving  
429 growth and faster reduction in alkaline conditions. Furthermore, with pH perturbations over 8 and

430 long enough timescales (>12 hours), the growth of rare taxa drives fast nitrate reduction. As a  
431 result, the optimal pH depends on the timescale of the measurement.

## 432 **Functional regimes and environmental fluctuations: origins of microbial di-** 433 **versity in nature**

434 For large basic perturbations, the abundant native taxa could no longer perform nitrate reduction,  
435 while the rare biosphere grew rapidly to reduce nitrate (Regime III), acting as the source of func-  
436 tional resilience in the community. The adaptation of rare taxa in extreme environments suggests  
437 that there might be a trade-off between stress resistance and fast growth [65]. Rare taxa may spe-  
438 cialize in surviving under extreme stress conditions (e.g., Firmicutes, Fig. S20), but perform little  
439 metabolic activity when the environment is near its native state. Conversely, dominant taxa near  
440 native environmental conditions (e.g., Proteobacteria) may specialize in faster growth rates when  
441 the nutrient becomes available but have limited ability to persist in stressful environments. These  
442 observations give rise to a picture where rare taxa are sustained by the presence of environmental  
443 fluctuations that transiently provide an opportunity to exploit resources [66].

444 Soil pH can change daily due to plant exudates (shifts of 0.4 in 12 hrs) [67], seasonally due to  
445 changes in rainfall and temperature (1–1.5 pH units) [68], and through agricultural practices [69].  
446 The titration curves gave insights into the amplitude of pH fluctuations the community experiences  
447 in nature (Fig. 6B). These observations place taxa's distinct physiology and environmental fluctua-  
448 tions at the center of understanding the origin and structure of regimes and therefore the metabolism  
449 of natural microbial communities. While physiology has experienced a renaissance of late, with  
450 quantitative approaches providing key insights [70], we know comparatively little about the role  
451 of natural environmental fluctuations in the wild. Our results suggest that understanding the dy-  
452 namics and origins of these fluctuations could provide deep insights into the responses of complex  
453 communities to environmental change.

## 454 **Methods**

### 455 **Sample collection, site description, and soil characterization**

456 Twenty topsoils were sampled across a range of pH values (4.7–8.32) from the Cook Agronomy  
457 Farm (Table S3). The Cook Agronomy Farm (CAF, 46.78°N, 117.09°W, 800 m above sea level) is a  
458 long-term agricultural research site located in Pullman, Washington, USA. CAF was established in  
459 1998 as part of the Long-Term Agroecosystem Research (LTAR) network supported by the United  
460 States Department of Agriculture. Before being converted to an agricultural field, the site was zonal  
461 xeric grassland or steppe. CAF operates on a continuous dryland-crop rotation system comprising  
462 winter wheat and spring crops. CAF is located in the high rainfall zone of the Pacific Northwest  
463 region and the soil type is classified as Mollisol (Naff, Thatuna and Palouse Series) [71]. Sampling  
464 occurred from September 8-12, 2022, post-harvest of spring crops, to reduce plant's impact on soil  
465 microbial communities. This period was during the dry season preceding the concentrated autumn  
466 rainfall.

467 Topsoils were collected from the eastern region of the CAF at a depth of 10–20 cm, other than  
468 Soil 1 & 2 (depth of 0–10 cm). Eastern CAF practices no-tillage which eliminates soil inversion and  
469 mixing of the soil surface to 20 cm. The N fertilizer in this field has been primarily deep banded to  
470 depths of approximately 7 to 10 cm during the time of application, which creates a spike of nitrate  
471 resource in the soil depth we sampled. Each soil sample was obtained by cutting down through the  
472 hardened dry soil with a spade in a circular motion to create a cylindrical cake of soil of radius 10–  
473 20 cm until the desired depth. Each soil sample was not merged from sampling multiple replicates  
474 due to differences in pH in different locations. Samples were collected within a diameter of 500 m  
475 within the CAF to minimize the variation of edaphic factors other than pH. The large variation in  
476 soil pH comes from the long-term use of ammoniacal fertilizers and associated N transformations,  
477 which may undergo nitrification resulting in the release of H ions. In combination, spatial pH  
478 variation increases with field-scale hydrologic processes that occur under continuous no-tillage



479 superimposed over a landscape that has experienced long-term soil erosion.

480 To maximize the coverage of sampled native pH, we used a portable pH meter (HI99121, Hanna  
481 Instruments, Smithfield, RI, USA) to directly measure and estimate the soil pH without having to  
482 make slurry on site to determine whether to sample the soil before sampling. For accurate pH val-  
483 ues, pH was measured in the laboratory using a glass electrode in a 1:5 (soil to water w/w) suspen-  
484 sion of soil in water (protocol of International organization for standardization, ISO 10390:2021),  
485 where 7g of soil was vortexed with 35ml of Milli-Q filtered water, spun down, and filtered with  
486 (0.22  $\mu\text{m}$ ) pore size. With these pH values, we selected 20 topsoil samples that are well spread  
487 across a range of pH from 4.7–8.32 with intervals of 0.1–0.6. Twenty soil samples were sieved  
488 (<2 mm), removed of apparent roots and stones, and gravimetric water content was determined (  
489 105 °C, 24h). The sieved samples were stored in the fridge for 0-3 months until the incubation  
490 experiment. For sequencing the initial community, subsamples were stored in –80 °C until the  
491 DNA extraction. The twenty soils were sent to the Research Analytical Laboratory (University  
492 of Minnesota, USA) to measure soil texture (soil particle analysis; sand, silt, clay composition),  
493 total carbon and nitrogen, and cation exchange capacity. The soils were also sent to Brookside  
494 Laboratories, Inc. (New Bremen, OH, USA) for a standard soil analysis package (Standard Soil  
495 with Bray I phosphorus). Twenty soils had relatively similar edaphic properties: 5–9% gravimetric  
496 water content (g/drysoilg), soil texture of silty clay or silty clay loam with 0% sand and 32–43%  
497 clay, and C:N ratio of 12–16 with 1–1.9% total carbon (wt/wt) (Table S3).

## 498 **Soil rewetting, constructing soil pH titration curves, and pH perturbation ex-** 499 **periments**

500 To mimic the autumn rainfall in the Pacific Northwest region and minimize the effect of spiking  
501 microbial activity by rewetting dry soils [72], we rewetted the sieved soil for 2 weeks before the  
502 perturbation experiments at room temperature with sterile Milli-Q water at 40% of each soil's water  
503 holding capacity. After resetting, a soil slurry was made by adding 2mM sodium nitrate solution  
504 to the soil (2:1 w/w ratio of water to soil). The slurry was then transferred to 48-deep well plates

505 (2.35ml of slurry per well) for incubation under anaerobic conditions (950 RPM, 30 °C) for 4 days.  
506 Anaerobic incubation was performed in a vinyl glove box (Coy Laboratory Products 7601-110/220)  
507 purged of oxygen with a 99%/1% N<sub>2</sub>/CO<sub>2</sub> gas mixture, where the gaseous oxygen concentration  
508 was maintained below 50 ppm to prevent aerobic respiration [73].

509 To perturb the soil pH to desired levels, we constructed each soil's pH titration curve for the 20  
510 soils with varying native pH to know exactly how much acid or base to add to each soil sample.  
511 To do so, separate from the main pH perturbation experiment, we added 23 different levels of HCl  
512 (acid) or NaOH (base) in the slurry, final concentrations ranging from 0–100 mM HCl or NaOH.  
513 We additionally tested whether the anion of acid (Cl<sup>-</sup>) or the cation of base (Na<sup>+</sup>) had a distinctive  
514 effect on the nitrate reduction dynamics, which was not the case (for results, see Fig. S12 and SM).  
515 We colorimetrically measured the pH (see section below) immediately after and 4 days after adding  
516 each well's designated amount of acid/base. Due to natural soil's buffering capacity, it takes 1-2  
517 days to stabilize its pH level. Thus, we used the endpoint (after 4 days) pH measurements for all  
518 pH perturbations. We did a spline interpolation on the titration data points, plotting endpoint pH  
519 (y-axis) against acid/base input (x-axis), to compute how much HCl and NaOH needs to be added  
520 to the soil to obtain 13 different levels of pH with  $\approx 0.4$  intervals ranging from pH 3 to 9, including  
521 the pH level without the addition of any acid or base. For Soil 19 and Soil 20, we had only 7 and  
522 3 perturbed pH levels respectively, because the strong buffering capacity of these soils (native pH  
523 over 8) limited the range of pH perturbation.

524 For the main pH perturbation experiment, the computed levels of concentrated HCl or NaOH  
525 were added to the slurry in the 48-deep well plate with and without chloramphenicol treatment  
526 for each perturbed pH level in triplicates. The plates were immediately transferred to the shaking  
527 incubator (950 RPM in Fisherbrand Incubating Microplate Shakers 02-217-759, 3 mm orbital ra-  
528 dius, 30 °C) inside the anaerobic glove box and incubated for 4 days. For chloramphenicol-treated  
529 (CHL+) samples, we added concentrated chloramphenicol solution to the slurry to obtain a final  
530 concentration of 1 g/L. To gauge the effect of the 2mM nitrate, we had no-nitrate controls (0 mM

531 nitrate) for both CHL+/- treatments in the unperturbed pH conditions. With antifungal cyclohex-  
532 imide controls (200 ppm) for all 20 soils, we confirmed that fungal activity minimally affects nitrate  
533 utilization dynamics (Fig. S4). We also confirmed that abiotic nitrate/nitrite reduction does not oc-  
534 cur by measuring metabolic dynamics of autoclaved soil (120 °C, 99 minutes, autoclaved 5 times  
535 every 2 days) (Fig. S5). To offset the effect of increasing metabolite concentration due to evapora-  
536 tion throughout the 4-day incubation period, we used the wells with just 2mM nitrate, nitrite, and  
537 ammonium solutions to correct for evaporation in the slurry samples for every time point. The val-  
538 ues of the gravimetric water content of each soil were taken into account to correct for the dilution  
539 of 2mM nitrate due to moisture in the soil. After the incubation, the plates were stored in -80 °C  
540 for sequencing endpoint communities.

### 541 **Time-series slurry sampling, extraction, and colorimetric assays to measure** 542 **nitrate, nitrite, ammonium, WSOC, and pH**

543 To obtain the metabolic dynamics, we subsampled 60 µL of the slurry into 96-well plates 10 times  
544 throughout 4 days (0, 4, 8, 19, 25, 31, 43, 55, 67, 91 hrs), where the initial time point ( $T_0$ ) is  
545 the time of pH perturbation and the start of anaerobic incubation. To measure nitrate and nitrite  
546 dynamics, extracts were prepared from the sampled slurries by adding and vortexing 2 minutes with  
547 90 µL of 3.33 M KCl solution (final concentration of 2 M KCl) and centrifuging at 4000rpm for 5  
548 minutes. The supernatant was filtered at 0.22 µm with a vacuum manifold to remove soil particles  
549 that could interfere with colorimetric assays. Concentrations of nitrate and nitrite in the extracts  
550 were determined colorimetrically using the Griess assay [74] and vanadium (III) chloride reduction  
551 method, following the protocol outlined previously [73]. We confirmed that 95%–99% of the nitrate  
552 in the soil can be accurately retrieved and detected using this method, as verified by nitrate spike-in  
553 and extraction experiments in the soil. For a subset of 20 soils (Soil 1, 5, 12, and 17), the ammonium  
554 dynamics were measured colorimetrically using the Salicylate-hypochlorite assay from the soil  
555 extracts [75]. Chloramphenicol treatments in the samples (CHL+) led to consistent detection of 0.5  
556 mM  $\text{NH}_4^+$  due to its N-H moiety. The salicylate-hypochlorite assay is also affected by the amount

557 of base (NaOH) in the samples, resulting in slightly lower detection of chloramphenicol in the  
558 CHL+ samples (0.45mM  $\text{NH}_4^+$  in 100mM NaOH perturbations). Taking advantage of these control  
559 measurements, we used the constant  $\text{NH}_4^+$  levels in the controls without 2mM  $\text{NO}_3^-$  (No Nitrate  
560 controls) in the CHL+ samples for each soil to offset the NaOH effect in the CHL- samples and  
561 subtracted  $\text{NH}_4^+$  levels caused by chloramphenicol in CHL+ samples.

562 For water-soluble organic carbon (WSOC) measurements, we subsampled 60  $\mu\text{L}$  of the slurry  
563 into 96-well plates at  $T_0$  and endpoint (4 days). Then, soil extracts were prepared by adding,  
564 vortexing with 90  $\mu\text{L}$  Milli-Q water, centrifuging at 4000rpm for 5 minutes, and 0.22  $\mu\text{m}$  filtering  
565 the supernatant. Concentrations of the organic carbon in the supernatant were measured colori-  
566 metrically by the Walkley-Black assay, which uses dichromate in concentrated sulfuric acid for  
567 oxidative digestion [76]. We subtracted 0.4 Cmg/ml from the CHL+ samples because chloram-  
568 phenicol gave rise to a measured value of 0.4 WSOCmg/ml without additional carbon. For pH  
569 measurements, we subsampled 100  $\mu\text{L}$  of the slurry into 96-well plates at  $T_0$  and the endpoint.  
570 Then, soil extracts were prepared by adding, vortexing with 150  $\mu\text{L}$  KCl solution (final concentra-  
571 tion of 1 M KCl), centrifuging at 4000rpm for 5 minutes, and 0.22  $\mu\text{m}$  filtering the supernatant. pH  
572 of the 120  $\mu\text{L}$  supernatant was determined colorimetrically by adding 4ul of the multiple indicator  
573 dye mixture via the protocol described previously [77]. The reason we used 1 M KCl method for  
574 pH measurement (ISO 10390:2021) was that, contrary to the KCl method, the  $\text{H}_2\text{O}$  method (using  
575 water instead of 1M KCl) resulted in a highly yellow coloration of the supernatants in strong basic  
576 perturbed samples, which interfered with the wavelength of the colorimetric pH assay. For samples  
577 of pH outside the range of the assay (below pH 3 and above pH 9), we used a pH micro-electrode  
578 micro (Orion 8220BNWP, Thermo Scientific, Waltham, MA, USA).

### 579 **DNA extraction with internal standards, library preparation, and 16s rRNA** 580 **amplicon sequencing**

581 We performed 16S amplicon sequencing on half of all samples: 10 (3, 5, 6, 9, 11, 12, 14, 15, 16,  
582 17; Table S3) out of 20 soils were sequenced before perturbation and at the endpoint in both CHL+

583 and CHL- conditions, totaling 1,085 amplicon sequencing measurements. Genomic DNA was ex-  
584 tracted from 500  $\mu$ L aliquots in a combined chemical and mechanical procedure using the DNeasy  
585 96 PowerSoil Pro Kit (Qiagen, Hilden, Germany). Extraction was performed following the manu-  
586 facturer's protocol, and extracted DNA was stored at  $-20^{\circ}\text{C}$ . To estimate the absolute abundance  
587 of bacterial 16S rRNA amplicons, we added known quantities of genomic DNA (gDNA) extracted  
588 from *Escherichia coli* K-12 and *Parabacteroides* sp. TM425 (samples sourced from the Duchos-  
589 sois Family Institute Commensal Isolate Library, Chicago, IL, USA) to the slurry subsamples be-  
590 fore DNA extraction. Equal concentrations of gDNA from these two strains were added. Both  
591 strains have identical rRNA copy numbers of 7 and comparable genome sizes of approximately  
592 5000 kb. DNA Library preparation was performed using the 16S Metagenomic Sequencing Li-  
593 brary Preparation protocol with a 2-stage PCR workflow (Illumina, San Diego, CA, United States).  
594 The V3–V4 region was amplified using forward primer 341-b-S-17 (CCTACGGGNGGCWGCAG)  
595 and reverse primer 785-a-A-21 (GACTACHVGGGTATCTAATCC) [78]. We confirmed using gel  
596 electrophoresis that the negative samples containing all reagents did not show visible bands after  
597 PCR amplification. Sequences were obtained on the Illumina MiSeq platform in a  $2 \times 300$  bp  
598 paired-end run using the MiSeq Reagent Kit v3 (Illumina, San Diego, CA, United States) with  
599 25% PhiX spike-ins. A standardized 10-strain gDNA mixture (MSA-1000, ATCC, Manassas, VA,  
600 USA) was sequenced as well to serve as a positive control, which was confirmed to have relatively  
601 uniform read counts after assigning taxa.

## 602 **Model and fitting**

### 603 **Consumer-resource model**

604 Consider a consumer-resource model with one consumer variable (functional biomass  $x(t)$ , OD/biomass)  
605 and two resource variables (nitrate  $A(t)$  and carbon-nutrient  $C(t)$ , mM), which evolves in time ( $t$ ,  
606 day). The ordinary differential equations (ODEs) of the consumer-resource model can be expressed

607 as:

$$\begin{aligned}
 \dot{A}(t) &= -r_A x(t) \frac{A(t)}{A(t) + K_A}, \\
 \dot{C}(t) &= -r_C x(t) \frac{C(t)}{C(t) + K_C}, \\
 \dot{x}(t) &= \gamma x(t) \frac{A(t)}{A(t) + K_A} \frac{C(t)}{C(t) + K_C}.
 \end{aligned}
 \tag{1}$$

608 The first two equations of (1) represent the resource consumption rates, which are determined by  
 609 the functional biomass ( $x$ , biomass), the maximum consuming rates per unit biomass ( $r_A$  and  $r_C$ ,  
 610 mM/biomass/day), and the Monod functions ( $A/(A + K_A)$  and  $C/(C + K_C)$ , dimensionless).  
 611 Here assume the affinities ( $K_A$  and  $K_C$ , mM) to be fixed and small. So the Monod functions can  
 612 be deduced to 1 when  $A \gg K_A$  or  $C \gg K_C$ , and can be deduced to 0 when  $A \rightarrow 0$  or  $C \rightarrow 0$ . The  
 613 third equation represents the growth of functional biomass, which is determined by the maximum  
 614 growth rate per biomass ( $\gamma$ , 1/day) and the multiplication of two Monod terms indicating the fact  
 615 that nitrate and carbon are non-substitutable (electron acceptor and donor respectively). The growth  
 616 is exponential ( $x(t) = x(0)e^{\gamma t}$ ) when both  $A \gg K_A$  or  $C \gg K_C$ , but growth stops when either  
 617  $A \rightarrow 0$  or  $C \rightarrow 0$ . Therefore, in this model, the growth of biomass is limited by both resources, but  
 618 the consumption of one resource can continue when the other resource runs out and the biomass  
 619 growth stops. For example, we believe this happens when  $C \rightarrow 0$  in Regime II and the consumption  
 620 of  $A$  continues (Fig. 2).

### 621 **Solution for nitrate dynamics**

622 To find the solution for nitrate dynamics, we rescale the equations by combining parameters:  $\tilde{x} =$   
 623  $r_A x$ ,  $\tilde{C} = C r_A / r_C$ ,  $\tilde{K}_C = K_C r_A / r_C$ . Therefore, the equations become:

$$\begin{aligned}
 \dot{A}(t) &= -\tilde{x}(t) \frac{A(t)}{A(t) + K_A} \\
 \dot{\tilde{C}}(t) &= -\tilde{x}(t) \frac{\tilde{C}(t)}{\tilde{C}(t) + \tilde{K}_C} \\
 \dot{\tilde{x}}(t) &= \gamma \tilde{x}(t) \frac{A(t)}{A(t) + K_A} \frac{\tilde{C}(t)}{\tilde{C}(t) + \tilde{K}_C}
 \end{aligned}
 \tag{2}$$

624 In the rescaled equations (2), the parameters and variables all have units of rates (nitrate per time):  
 625  $[\tilde{x}] = mM/day$  and  $[\tilde{C}] = [\tilde{K}_C] = mM$ . Therefore, the solution of nitrate dynamics only depends  
 626 on three parameters ( $\gamma, K_A, \tilde{K}_C$ ) and three initial conditions ( $A_0, \tilde{C}(0), \tilde{x}(0)$ ). Since the affinities  
 627 are very small ( $K_A \approx 0.01mM, \tilde{K}_C \approx 0.01mM$ ), the solution of biomass approximately equals  
 628 to  $\tilde{x} = \tilde{x}(0)e^{\gamma t}$  before the time at which growth stops  $t^*$ . So the resource dynamics before  $t^*$  are  
 629 approximately  $A = A_0 - \tilde{x}(0)(e^{\gamma t} - 1)/\gamma$  and  $\tilde{C} = \tilde{C}(0) - \tilde{x}(0)(e^{\gamma t} - 1)/\gamma$ . Accordingly, the time  
 630 at which growth stops is given by  $t^* = \log(\min(A_0, \tilde{C}(0))\gamma/\tilde{x}(0) + 1)/\gamma$ . If  $\tilde{C}(0) < A_0$ , the  
 631 nitrate dynamics after  $t^*$  and before running out are given by  $A = A(t^*) - (\gamma\tilde{C}(0) + \tilde{x}(0))(t - t^*)$ .  
 632 As a result, the nitrate consumption rate after  $t^*$  is  $\gamma\tilde{C}(0)$  larger than the initial rate  $\tilde{x}(0)$ .

### 633 Least-square fitting scheme

634 To infer the model parameters from the metabolite measurement, we use the least-square fitting  
 635 scheme to find the closest dynamic curves to the time-series data. Our metabolite measurement  
 636 including the time points ( $\underline{t}^- = [t_1^-, t_2^-, \dots, t_N^-]$ ) and nitrate amount ( $\underline{a}^- = [a_1^-, a_2^-, \dots, a_N^-]$ ) for each  
 637 CHL- sample, and the measurements of  $\underline{t}^+$  and  $\underline{a}^+$  for a corresponding CHL+ sample. We set up  
 638 the loss function as the mean-squared-error (MSD):

$$L = \frac{1}{2N} \left( \sum_{k=1}^N (A(t_k^-) - a_k^-)^2 + \sum_{k=1}^N (A^c(t_k^+) - a_k^+)^2 \right). \quad (3)$$

639 Here the functions  $A(t)$  and  $A^c(t)$  are theoretical solutions of the consumer-resource model (2) for  
 640 CHL-/± conditions, respectively. Because the nitrate dynamics  $A(t)$  and  $A^c(t)$  are determined by  
 641 the parameter set  $\Theta = \{\tilde{x}(0), \tilde{C}(0), A_0, A_0^c, \gamma, K_A, \tilde{K}_C\}$ , we minimize the loss function  $L(\Theta)$  to  
 642 get the optimal model parameters  $\Theta^*$ . We note to the readers that three parameters are fixed ( $\gamma =$   
 643  $4.8day^{-1}, K_A = \tilde{K}_C = 0.01mM$ ) as justified by the sensitivity analysis in the following paragraph.  
 644 Note, that these parameters were globally fixed across all the data. We also use different initial  
 645 nitrate ( $A_0$  and  $A_0^c$ ) in the functions  $A(t)$  and  $A^c(t)$ . The optimization algorithm is the interior-  
 646 point method which is built in the MATLAB `fmincon` function. The codes and data are available at  
 647 <https://github.com/SiqiLiu-Math/xxx>. The fitting errors over all samples are shown

648 in Fig. S7, in which the root-mean-squared-error (RMSE,  $\sqrt{L(\Theta^*)}$ ) and the error per datapoint  
649 ( $|A(t_k^-) - a_k^-|$  or  $|A^c(t_k^+) - a_k^+|$ ) are normalized by the input value of nitrate (2mM).

### 650 **Sensitivity analysis on model parameters**

651  $\gamma$ ,  $K_A$ , and  $\tilde{K}_C$  were globally fixed to one value across all data. Here we justify this decision. We  
652 analyzed the sensitivity of  $\gamma$ ,  $K_A$ , and  $\tilde{K}_C$  on simulated dynamic data. To reflect the three typical  
653 dynamics (regimes) observed from the measurement, we simulated three nitrate curves by setting  
654 up the initial conditions to be  $\tilde{x}(0) = 0.01, 0.1, 0.001mM/day$  and  $\tilde{C}(0) = 0.005, 0.05, 2mM$ ,  
655 respectively. Other parameters are given by  $A_0 = A_0^c = 2mM$ ,  $K_A = \tilde{K}_C = 0.01mM$ ,  $\gamma =$   
656  $4day^{-1}$ . We then used different fixed values of parameters to fit the three examples. In the first  
657 row of Fig. S8, we used different fixed  $\gamma$  values - from  $\gamma = 2day^{-1}$  to  $\gamma = 6day^{-1}$  - to fit  
658 three simulations. We demonstrate very small mismatches (RMSE < 5%) from these variations of  
659 parameter values, which are almost invisible in Regime I and Regime II fittings. In the second and  
660 the third row of Fig. S8, we use different fixed  $K_A$  and  $\tilde{K}_C$  values - from  $10^{-4}mM$  to  $1mM$  - to  
661 fit three simulations. When  $K_A < 0.1mM$  or  $\tilde{K}_C < 0.1mM$ , the mismatches were again very  
662 small (RMSE < 1%) and invisible. These results indicate that the fixed values of  $\gamma$ ,  $K_A$  and  $\tilde{K}_C$   
663 are insensitive in large ranges.

### 664 **Determination of regime boundary with model parameters**

665 To define the regime boundaries, we examined the distributions of each parameter's value.  $\tilde{x}(0)$   
666 had a bimodal distribution (Fig. S9A). This bi-modality becomes more evident when we separately  
667 observe its distribution from the left half (perturbed pH < 4) and right half (perturbed pH > 6)  
668 of the parameter space displayed in the perturbed pH vs. native pH grid in Figure 3C (Fig. S9B).  
669 Therefore, we set the threshold for the  $\tilde{x}(0)$  boundary where these two modes are evidently sepa-  
670 rated ( $\tilde{x}(0) = 0.05$ ). The distribution of  $\gamma\tilde{C}(0)$  exhibited a significant mode around 0, prompting  
671 us to set the threshold ( $\gamma\tilde{C}(0) = 1.5$ ) at the tail region, where the  $\gamma\tilde{C}(0)$  threshold also separated  
672 the Regime III samples in the top-left quadrant of the  $\tilde{x}(0)$  vs.  $\gamma\tilde{C}(0)$  scatter plot (Fig. 3A). The



673 separation of Regime I and Regime II data points may not be clear cut in the  $\tilde{x}(0)$  vs.  $\gamma\tilde{C}(0)$  scat-  
674 ter plot (Fig. 3A). However, when we plot  $\tilde{x}(0)$  of individual soils from different native pH levels  
675 (Fig. S9E), especially in the natively acidic soils, the transition from Regime II (large  $\tilde{x}(0)$ ) to  
676 Regime I (small  $\tilde{x}(0)$ ) is evident going towards more acidic pH perturbations because of the large  
677  $\tilde{x}(0)$  levels sustained over a wide pH range in Regime II.

## 678 **Sequence data analysis**

### 679 **Sequencing data preprocessing and assigning taxonomy to ASVs with DADA2**

680 Raw Illumina sequencing reads were stripped of primers, truncated of Phred quality score below  
681 2, trimmed to length 263 for forward reads and 189 for reverse reads (ensuring a 25-nucleotide  
682 overlap for most reads), and filtered to a maximum expected error of 4 based on Phred scores;  
683 this preprocessing was performed with USEARCH ver. 11.0 [79]. The filtered reads were then  
684 processed with DADA2 ver. 1.18 following the developers' recommended pipeline [80]. Briefly,  
685 forward and reverse reads were denoised separately, then merged and filtered for chimeras. For  
686 greater sensitivity, ASV inference was performed using the DADA2 pseudo-pooling mode, pooling  
687 samples by soil. After processing, the sequencing depth of denoised samples was  $10^4$ – $10^6$  reads per  
688 sample. Low-abundance ASVs were dropped ( $\lesssim 10$  total reads across all 1085 samples), retaining  
689 34 696 ASVs for further analysis. Taxonomy was assigned by DADA2 using the SILVA database  
690 ver. 138.1, typically at the genus level, but with species-level attribution recorded in cases of a  
691 100% sequence match.

### 692 **Computing absolute abundance with internal standards of each ASV per sample**

693 As an internal control, we verified that the ASVs corresponding to the two internal standard genera  
694 *Escherichia-Shigella* and *Parabacteroides* were highly correlated with each other as expected (per-  
695 son correlation  $\rho = 0.94$ ). These ASVs were removed from the table and combined into a single  
696 reference vector of "spike-in counts". The spike-in counts constituted  $8.9 \pm 8.8\%$  of the total reads  
697 in each sample. For downstream analysis, the raw ASV counts in a sample were divided by the

698 spike-in counts of the internal standard per sample to obtain the absolute abundance of the ASV in  
699 a sample. Total biomass per sample was obtained by dividing the total raw read counts with the  
700 spike-in counts of the sample.

### 701 **Differential abundance analysis to identify enriched ASVs**

702 We conducted differential abundance analysis to statistically determine which amplicon sequence  
703 variants (ASVs) were significantly enriched in the Nutrient-limiting regime (Regime II) or the  
704 Resurgent growth regime (Regime III), respectively. To do so, we identified enriched ASVs for  
705 each perturbed pH condition in each native soil comparing endpoint chloramphenicol-untreated  
706 (CHL-) samples with endpoint chloramphenicol-treated (CHL+) samples. For each native soil,  
707 we then compiled a list of enriched ASVs by aggregating a union set of enriched ASVs across  
708 perturbed conditions that belong to Regime II (or Regime III). To remove ASVs that could be  
709 false-positive nitrate reducers, we similarly performed differential abundance analysis to identify  
710 ASVs that are enriched in no-nitrate controls (nitrate<sup>-</sup>) by comparing endpoint chloramphenicol-  
711 untreated (CHL- & nitrate<sup>-</sup>) samples with endpoint chloramphenicol-treated (CHL+ & nitrate<sup>-</sup>)  
712 samples. This filtering was done when inferring nitrate reducer biomass (Fig. S11C&D) and infer-  
713 ring the Regime III strains (Fig. S17). For each native soil, we only had nitrate<sup>-</sup> controls for the  
714 condition without acidic/basic perturbation. We assumed that these enriched ASVs in no-nitrate  
715 conditions (NNresponders) without acid/base perturbation would also be false-positive nitrate re-  
716 ducers in other acidic or basic perturbation levels. For each native soil, we filtered out these false-  
717 positive NNresponders from the aggregated list of Regime II (or Regime III) enriched ASVs.

718 To identify the ASVs enriched for each perturbed pH level, it was necessary to determine what  
719 change in recorded abundance constitutes a significant change, relative to what might be expected  
720 for purely stochastic reasons. The relevant null model would combine sampling and sequencing  
721 noise with the stochasticity of ecological dynamics over a 4-day incubation, and cannot be de-  
722 rived from first principles. However, since all measurements were performed in triplicate with

723 independent incubations, the relevant null model can be determined empirically. The deviations of  
724 replicate-replicate comparisons from 1:1 line were well-described by an effective model combining  
725 two independent contributions, a Gaussian noise of fractional magnitude  $c_{\text{frac}}$  and a constant Gaus-  
726 sian noise of magnitude  $c_0$  reads, such that repeated measurements (over biological replicates) of an  
727 ASV with mean abundance  $n$  counts are approximately Gaussian-distributed with a standard devia-  
728 tion of  $\sigma(c_0, c_{\text{frac}}) = \sqrt{(c_{\text{frac}}n)^2 + c_0^2}$  counts. In this expression,  $c_{\text{frac}}$  was estimated from moderate-  
729 abundance ASVs ( $> 50$  counts) for which the other noise term is negligible; and  $c_0$  was then de-  
730 termined as the value for which 67% of replicate-replicate comparisons are within  $\pm\sigma(c_0, c_{\text{frac}})$  of  
731 each other, as expected for 1-sigma deviations. This noise model was inferred separately for each  
732 soil and each perturbed pH level, as the corresponding samples were processed independently in  
733 different sequencing runs. For example, the parameters in Soil 11 were  $c_{\text{frac}} = 0.21 \pm 0.04$  and  
734  $c_0 = 4.5 \pm 0.7$  counts (Fig. S25).

735 The model was used to compute the z-scores for the enrichments of absolute ASV abundances  
736 in CHL- treatments against CHL+ controls (three independent z-scores from three replicate pairs;  
737 rep1-rep1, rep2-rep2, rep3-rep3). The median z-score was assigned to each ASV for each per-  
738 turbed condition. In consideration of ASVs with 0 read count in either CHL-/± samples, all raw  
739 ASV counts were augmented by a pseudocount of 0.5 and divided by the per-sample spike-in  
740 counts, yielding values that can be interpreted as the absolute biomass of each taxon (up to a factor  
741 corresponding to the copy number of the 16S operon), measured in units where 1 means as many  
742 16S fragments as the number of DNA molecules in the spike-in. Significantly enriched ASVs were  
743 identified in each perturbed condition as those with z-scores greater than  $z = \Phi^{-1}(1 - \alpha/2/n_{\text{ASV}})$ ,  
744 where  $\Phi^{-1}(x)$  is the inverse CDF of the standard normal distribution,  $\alpha = 0.05$ , and  $n_{\text{ASV}}$  as the  
745 number of nonzero ASVs in a given sample. This critical z-score ( $z = 4.2$ , when  $n_{\text{ASV}} = 2000$   
746 for enriched ASVs and  $z = 4.3$ , when  $n_{\text{ASV}} = 2500$  for filtering no-nitrate responders (NNRe-  
747 sponders)) corresponds to a two-tailed Bonferroni-corrected hypothesis test at significance level  $\alpha$   
748 under the null hypothesis that counts in the CHL- and CHL+ conditions are drawn from the same

749 distribution. These analyses were performed using custom MATLAB (Mathworks, Inc) and R  
750 scripts, which are available on the GitHub data repository for the present manuscript; for additional  
751 technical details, the reader is referred to the detailed comments in these scripts.

## 752 **Non-negative matrix factorization (NMF) analysis on phylum-level growth folds**

753 To analyze the abundance change at the phylum level, we compute the growth fold of each phylum  
754 at each condition. For each phylum, we compute the absolute abundance by aggregating the abun-  
755 dances of all ASVs within that phylum. Taking CHL+ abundance  $Abs^+$  as the reference abundance  
756 and CHL- abundance  $Abs^-$  as the endpoint abundance (where  $Abs$  denotes taxon abundance nor-  
757 malized to internal standard), the logarithm of the growth fold for phylum  $i$  and condition  $j$  is given  
758 by  $g_{ij} = \log(Abs_{ij}^- + 10^{-3}) - \log(Abs_{ij}^+ + 10^{-3})$ . Note that we use CHL+ abundance as reference in-  
759 stead of the initial abundance (at T0), to account for any effects on read counts unrelated to growth  
760 which would be common between CHL+ and CHL- conditions (e.g. direct effect of acid/base ad-  
761 dition), allowing us to focus only on growth-mediated abundance changes. We also set all negative  
762  $g_{ij}$  to 0 since we are focusing on growth. For all 130 conditions (10 soils  $\times$  13 perturbations) and  
763 40 phyla, the phylum-level growth folds  $G$  is a  $40 \times 130$  matrix. For each phylum, the row vector  $\vec{g}_i$   
764 represents how it grows under different conditions (see Fig. S15 for the growth vectors of the first  
765 10 phyla). In order to reduce the dimensionality of the growth matrix and extract the main features  
766 of the growth vectors, we use non-negative matrix factorization (NMF) to decompose the growth  
767 matrix  $G = W * H$  by factor 2. Here the feature matrix  $H$  is of size  $2 \times 130$ , and the two rows  
768  $\vec{h}_1$  and  $\vec{h}_2$  are two modes of growth vectors (shown in Fig. S15B). Therefore, the growth vector  
769 of phylum  $i$  is thus decomposed as  $\vec{g}_i \approx w_{i1}\vec{h}_1 + w_{i2}\vec{h}_2$ , while the weights  $w_{i1}$  and  $w_{i2}$  are from  
770 the  $40 \times 2$  weight matrix  $W$ . The weights of all 40 phyla are plotted in Fig. S15B, showing that  
771 Firmicutes are mostly composed by the second mode  $\vec{h}_2$  and other phyla are mostly composed by  
772 the first mode  $\vec{h}_1$ . Additionally, Bacteroidota and Proteobacteria show the most significance of the  
773 first mode. This decomposition keeps 93.36% of the original growth matrix.

## 774 **Genotyping enriched ASVs with PICRUSt2**

775 To understand what traits make Resurgent growth strains unique, we used PICRUSt2 ver. 2.5.2 [40]  
776 to infer putative genotypes of the enriched ASVs in the Resurgent growth regime (Regime III)  
777 (Fig. S18C). Using the script "place\_seqs.py" in the pipeline, we matched the representative 16S  
778 rRNA sequences of each amplicon sequence variant (ASV) to PICRUSt2's curated reference genome  
779 database (multiple sequence alignment). Then, using the "hsp.py" script and default parameters,  
780 we predicted KEGG orthologs (KO) abundance of each ASV with the matched reference genome  
781 (hidden-state prediction). To narrow down to KOs/genes related to denitrification and Dissimila-  
782 tory Nitrate Reduction to Ammonium (DNRA), we focused on nitrate reductase in denitrification  
783 (*narG*/K00370, *narH*/K00371, *narI*/K00374, *napA*/K02567, *napB*/K02567) and nitrite reductase  
784 to ammonium (*nirB*/K00362, *nirD*/K00363, *nrfA*/K03385, *nrfH*/K15876). To track which KOs  
785 were enriched at which pH in the 89 families used in the peak pH analysis (see SM for peak pH  
786 analysis), we summed the relative abundance (reads / total reads of each perturbed pH level in  
787 CHL-) of the ASVs belonging to each family that possessed at least 1 predicted gene respectively  
788 for *narGHI*, *napAB*, *nirBD*, and *nrfAH*. Then, we plotted their relative abundance values across pH  
789 for all soils, indicated by the intensity of the point's colors (Fig. S18).

## 790 **Taxonomy of growing strains in Regime III varies with soil native pH**

791 To further investigate whether the taxonomic identity of Resurgent growth (Regime III) strains  
792 varies across natural pH environments, we performed a regularized regression analysis to see if we  
793 can predict the native pH level of the source soil from the presence or absence of taxa that grow  
794 in Regime III at the ASV, Species, Genus, Family, or higher taxonomic levels. The Resurgent  
795 growth strains were determined by the differential abundance analysis as described previously.  
796 Should our findings confirm that the prediction of native soil pH is feasible based on the taxonomic  
797 variation of these strains, it means that the strains responsible for growth in Regime III *depend*  
798 *on the long-term pH of the soil*. To do so, we used the sequencing data to build a matrix where

799 the rows are samples (including three biological replicates) belonging to the Resurgent growth  
800 regime (Regime III), where each row has a corresponding native pH value of the original soil.  
801 There are 10 source soils with different native pH levels, and each soil has 3 to 6 pH perturbed  
802 samples (replicates) of which metabolite dynamics are classified as the Resurgent growth regime.  
803 The matrix's columns are different taxa belonging to the identified Resurgent growth strains, either  
804 in ASV, species, genus, family, or higher levels. Each element of the matrix is 0 if absent and  
805 1 if present in the sequencing data of the sample. Because the presence and absence of taxa can  
806 randomly depend on the random sampling depth of each sample, we test varying threshold values  
807 (0, 0.001, 0.005 relative abundance) to call the taxa present if their relative abundance is greater  
808 than the threshold (effects shown in Fig. S22F).

809 The regularized regression was performed to predict the native pH of the source soil of the  
810 samples from the presence and absence of taxa using only additive terms and LASSO regularization  
811 to avoid overfitting [81]. To estimate the regularization hyperparameter, tenfold cross-validation  
812 was performed on the samples from ten different soils with different native pH levels. All models  
813 were fit using the package glmnet in R version 4.1.4. To make predictions of the native pH, we  
814 used two strategies. First, 'in-sample' predictions used all available data points to fit the regression  
815 coefficients and predicted native pH using those coefficients. Second, to ask whether we can still  
816 predict the native pH without the model seeing the samples belonging to that specific native pH  
817 level, we implemented a 'Leave-one-soil-out' (LOSO) procedure where all the perturbed samples  
818 from one native soil were left out as a test set, and the model was trained on the remaining data  
819 to fit the regression coefficients. Then, we used the model to predict the native pH of the left-out  
820 samples (out-of-sample prediction). The observed versus predicted pH values are shown in the  
821 scatter plots (Fig. S22A). The prediction quality ( $R^2$ , coefficient of determination =  $1 - SS_E/SS_T$ ,  
822 sum of squares error over total sum of squares) was computed using the mean predicted and mean  
823 observed native pH levels for each soil (for different taxonomic levels and prediction strategies, see  
824 Fig. S22 E & F, negative  $R^2$  values indicate the predictions are worse than just predicting the pH as

825 the mean predicted pH). To ascertain that our high prediction quality was not a random artifact, we  
826 randomly permuted the native pH values of our soils 1000 times and then predicted in-sample the  
827 native pH to obtain 1000  $R^2$  values. We computed the threshold  $R^2$  value that corresponds to the  
828 p-value of 0.05 (top 50th  $R^2$  value out of 1000 instantiations) and compared it with the  $R^2$  value  
829 that we have obtained with our true native pH predictions (Fig. S22G).

### 830 **Testing the effect of different bases and salts on nutrient release**

831 To see the effects of different bases (NaOH and KOH) on nitrate reduction dynamics, we added  
832 different concentrations of NaOH and KOH (final concentration of 0, 8, 16, 24 mM in the slurry),  
833 following the same protocol previously described (without chloramphenicol), to measure the nitrate  
834 and nitrite dynamics (Fig. S12) using Soil 6 (Table S3). In addition, to test the effects of  $\text{Na}^+$ ,  $\text{K}^+$ ,  
835  $\text{Cl}^-$  separately, we added different concentrations of salts (NaCl, KCl) (without chloramphenicol  
836 and without adding any acid/base) and measured the metabolite dynamics (Fig. S12).

### 837 **Nutrient amendment experiments with slurries**

838 To experimentally determine what nutrient was limiting growth in the Nutrient-limiting regime, we  
839 conducted nutrient amendment experiments respectively with glucose, succinate, sodium acetate,  
840 ammonium chloride ( $\text{NH}_4\text{Cl}$ ), monosodium phosphate ( $\text{NaH}_2\text{PO}_4$ ), and sodium sulfate ( $\text{Na}_2\text{SO}_4$ )  
841 (for results, see Fig. 4D and Fig. S14). Among them, succinate ( $\text{pK}_a = 4.21$  and  $5.64$ ,  $25^\circ\text{C}$ ), acetate  
842 ( $\text{pK}_a = 4.76$ ,  $25^\circ\text{C}$ ), and phosphate ( $\text{pK}_a = 2.2$ ,  $7.2$ , and  $12.4$ ,  $25^\circ\text{C}$ ) were strong candidates for  
843 the limiting nutrient according to our soil nutrient release hypothesis, due to their anionic nature  
844 in mid-range pH (5-7). The incubation was conducted following the same protocol using Soil 6  
845 (Table S3) without chloramphenicol and without adding any acid/base. Concentrations were either  
846 in C mM, N mM, S mM, or P mM with final concentrations in slurry varying from 0 to 5 mM,  
847 each in triplicates. Because we have previously tested the effect of  $\text{Na}^+$  and  $\text{Cl}^-$  to be negligible  
848 in nitrate dynamics, the effect of these amendments can be attributed solely to C/N/S/P nutrients  
849 other than  $\text{Na}^+$  and  $\text{Cl}^-$ .

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889 [//github.com/KiseokKeithLee/Soil\\_pH\\_perturbation/](https://github.com/KiseokKeithLee/Soil_pH_perturbation/) upon publication.

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## 1096 **Supplementary Materials**

### 1097 **A detailed description of the three functional regimes: the Acidic death regime** 1098 **(Regime I), Nutrient-limiting regime (Regime II), and Resurgent growth regime** 1099 **(Regime III)**

1100 By quantitatively distinguishing the impact of pH on the consumer side (microbial community,  
1101  $\tilde{x}(0)$ ) and the resource side (growth-limiting nutrient,  $\tilde{C}(0)$ ), we can ask the mechanism behind  
1102 functional adaptation during different regimes. Regime II can be called the “Nutrient-limiting”  
1103 regime. Within this pH range (Fig. 3C), conditions favor the resident population of nitrate reducers;  
1104 hence it allows a large indigenous nitrate reducer population to perform nitrate reduction (large  
1105  $\tilde{x}(0)$ ). This specific range of favorable pH levels is determined by the long-term exposure to the  
1106 native pH of soils (Fig. 3C). In this regime, the increase of nitrate reduction rate is determined  
1107 by the biomass growth from the available growth-limiting nutrient. Therefore, in Regime II, the  
1108 adaptive strategy employed by the nitrate-reducing population is to utilize the pre-existing resident  
1109 species which are rather robust to pH perturbations, and at the same time incrementally increase  
1110 the resident’s biomass as the resource availability changes with pH perturbations. Going back to  
1111 the functional dynamics, that is the reason we see a relatively high slope of CHL+ conditions and  
1112 a slight increase of denitrification rate in CHL- conditions ((b) and (e) of Fig. 3B), demonstrating  
1113 that the resident nitrate reducers adapt to the new environment in a “nutrient-limiting” manner.

1114 Regime III can be called the “Resurgent growth” regime. As the perturbed pH is increased  
1115 from Regime II, there comes a critical pH of around 8 where the adaptive mechanism abruptly  
1116 transitions. When the pH is perturbed beyond the critical point, the previously large biomass of  
1117 the nitrate reducer population can no longer adapt and perform nitrate reduction ( $\tilde{x}(0) \rightarrow 0$ ). On  
1118 the resource aspect in Regime III, there is a surplus of limiting nutrients, and thus the system is  
1119 no longer limited by  $C$  but limited by nitrate  $A$ . These two effects of short-term pH change (both  
1120 on the consumer and resource aspect) set the stage for the “resurgence”. A rare population, which  
1121 we will investigate the composition later, appears to have a small biomass initially showing a flat



1122 slope in both CHL<sup>-</sup> and CHL<sup>+</sup> conditions but later grows exponentially to exponentially deplete the  
1123 nitrate in Regime III (panel c, f of Fig. 3F). This shows that in Regime III, the adaptive mechanism  
1124 of the community is to rely on the rare uprising nitrate reducer biomass to rapidly grow in the  
1125 absence of nutrient limitation.

1126 Regime I can be called the “Acidic death” regime. This regime is at the other end (acidic) of the  
1127 Nutrient-limiting regime (Regime II). As the perturbed pH is decreased from Regime II, it transi-  
1128 tions into a regime where the system fails to adapt. The boundary pH of Regime I and Regime II is  
1129 influenced by the native pH of the soil, where relatively acidic soils have a lower boundary of pH 3  
1130 or 4 and relatively neutral soils have a higher boundary of pH 5 to 6. Similar to what happens when  
1131 the community enters Regime III, the unfavorable pH diminishes the indigenous nitrate-reducing  
1132 activity of the soil, indicated by the flat CHL<sup>+</sup> dynamics in panels a, d of Fig. 3B. However, unlike  
1133 Regime III, the perturbed pH does not make the growth-limiting nutrient superfluous but makes  
1134 it unavailable, making the divergence of CHL<sup>-</sup> and CHL<sup>+</sup> dynamics nonexistent (panel a, d of  
1135 Fig. 3A). These two effects of pH perturbation make it extremely difficult for the community to  
1136 adapt to the new environment of Regime III, hence the “Acidic death” regime. Another reason we  
1137 call it that is to highlight the asymmetric effect of acidic and basic perturbations, which has been  
1138 seldom acknowledged in the literature.

### 1139 **Detailed mechanism of nutrient release in soils due to change in pH**

1140 We turned to soil literature to develop a comprehensive mechanism for the nutrient release mecha-  
1141 nism in soil [38, 39]. Soil comprises minerals, organic matter, water, and air. Minerals and organic  
1142 matter form aggregated clumps of soil particles, categorized by size into sand, silt, and clay. Clay  
1143 particles, the smallest among them, consist of layers of phyllosilicates. Each layer includes tetrahe-  
1144 dral structures of Si<sup>4+</sup> covalently bonded to four oxygens and octahedral structures of continuous  
1145 Mg<sup>2+</sup> or Al<sup>3+</sup> covalently bonded to six hydroxides [82]. Due to this chemical structure, clay par-  
1146 ticles possess numerous electrostatically charged sites, including negatively charged (oxygen atom

1147 and hydroxide) and positively charged [39] (shown as - and + sites in Fig. 4B). The clay particle's  
1148 cation exchange sites are negatively charged and form ionic bonds with cations (positively charged  
1149 ions), while the anion exchange sites are positively charged and bind to anions (negatively charged  
1150 ions). Both cations and anions can serve as potential nutrient sources. When they are bound to  
1151 the clay's exchange sites (brown section in Fig. 4B), they are protected from microbes. However,  
1152 when they are released and dissolved in the pore water (light blue or pink section in Fig. 4B), they  
1153 become available to the microbial community. To understand how NaOH or HCl impacts nutrient  
1154 availability, it's essential to track whether cations and anions are bound to clay particles or dissolved  
1155 in the pore water.

1156 The literature on nutrients and pH in soils proposes the following mechanism for nutrient re-  
1157 lease (Fig. 4B, see more detailed cartoon Fig. S13B) [38, 39]. When NaOH is added to the soil  
1158 solution, both  $\text{Na}^+$  and  $\text{OH}^-$  ions (pH-mediated) act to release the anionic nutrients (case 2 of  
1159 Fig. 4B). First,  $\text{OH}^-$  deprotonates ion exchange sites in the clay particles increasing the number  
1160 of cation exchange sites (- charge) and decreasing the number of anion exchange sites (+ charged)  
1161 reducing the capacity of the clay to hold anions. Secondly,  $\text{Na}^+$  can either bind to the clay particle  
1162 or remain in solution [38]. The ( $\text{Na}^+$ ) that remains in solution increases the stability of released an-  
1163 ions. Overall, increased  $\text{OH}^-$  increases the anionic nutrients available to the microbial community  
1164 (Fig. 4B). The converse happens for HCl perturbations (case 1 of Fig. 4B).

## 1165 **Effect of base cation on nutrient release in soils**

1166 Nutrient release is not solely driven by  $\text{OH}^-$  ions. The base cation plays an important role, and  
1167 thus whether the base cation prefers to be in the clay particle or the water solution can influence  
1168 the amount of nutrients available to microbes. It is known that the bigger the size and greater  
1169 the charge of the cation, the more selectively the cation binds to clay particles. For example, the  
1170 cation's binding specificity to the clay particle instead of staying in the solution is in the order  
1171 of  $\text{NH}_4^+ > \text{K}^+ > \text{Na}^+ > \text{Li}^+$ , divalent cations having greater binding specificity than monovalent

1172 cations (e.g.,  $\text{Ca}_2^+ > \text{K}^+$ ) [39]. Therefore, when the amount of dissolved organic carbon (DOC) was  
1173 measured after adding  $\text{Ca}(\text{OH})_2$  vs. KOH in equimolar hydroxide, adding KOH resulted in a much  
1174 higher concentration of dissolved organic carbon (DOC) [38]. Because  $\text{K}^+$  ions less specifically  
1175 bind to the clay particle and more likely remain in the solution, it would stabilize the released  
1176 anions better in the solution, presuming that the DOC is mostly anion due to many negatively  
1177 charged moieties of  $\text{O}^-$ s. To check if there was a significant difference between monovalent cations,  
1178 we compared NaOH and KOH treatments for basic perturbations. We found that there was no  
1179 significant difference in the nitrate utilization rates in the CHL- condition when we added the same  
1180 concentrations of NaOH and KOH, indicating that the amount of limiting nutrient released was  
1181 similar (Fig. S12), although the stabilized endpoint pH was different. As a sanity check, we further  
1182 tested KCl and NaCl treatments and found that  $\text{K}^+$ ,  $\text{Na}^+$ , or even  $\text{Cl}^-$  (relevant in the HCl addition)  
1183 ions themselves without  $\text{OH}^-$  did not affect nutrient release (Fig. S12), which agrees with previous  
1184 findings [38].

## 1185 **Recapitulating linear dynamics with monoculture experiments without car-** 1186 **bon**

1187 Our model and functional dynamics data suggest that the limited carbon leads to a constant rate of  
1188 nitrate reduction. However, it is difficult to understand the mechanism behind this phenomenon,  
1189 because if the organic carbon, an electron donor in the electron transport chain, is coupled to the  
1190 reduction reaction of nitrate (terminal electron acceptor), the depletion of organic carbon will likely  
1191 stop the nitrate reduction performed by the nitrate reductase enzymes. This will cause the nitrate  
1192 reduction rate to be close to 0 rather than the observed constant rate. To resolve this contradiction,  
1193 one hypothesis can be that the cells internally store carbon nutrients (electron donor) to power the  
1194 electron transport chain (consuming nitrate) without needing to import external carbon nutrients to  
1195 generate ATPs for the cell's maintenance energy. To test this hypothesis, we conducted monoculture  
1196 experiments with a *E. coli* strain and a known denitrifier *Pseudomonas sp.* strain.

## 1197 **Culturing protocol**

1198 Strains were pre-cultured in two stages under aerobic conditions before being transferred to den-  
1199 itrifying (anaerobic) conditions for phenotyping. First, wells of a sterile 24-well plate (Thermo  
1200 Scientific Nunc Non-Treated Multidishes) were loaded with 1.7 mL of R2B medium. Wells were  
1201 inoculated with *E. coli* K12 and *Pseudomonas sp.* PDM04 [73] strains from glycerol stocks stored  
1202 at  $-80^{\circ}\text{C}$ . The plates were then sealed with a gas-permeable sterile membrane (Breathe-Easier,  
1203 USA Scientific, 9126-2100). After sealing, the culture was incubated overnight at 0.5 rcf (400 RPM  
1204 in Fisherbrand Incubating Microplate Shakers 02-217-759, 3 mm orbital radius) and  $30^{\circ}\text{C}$  in aer-  
1205 obic conditions. These cultures reached saturation during this time. Second, wells of a sterile  
1206 24-well plate were loaded with 1.7 mL of defined media (15 mM ammonium, 40 mM phosphate  
1207 buffer with the final medium pH adjusted to 7.3, and trace metals and vitamins, as described in  
1208 Ref [73]) with 25 mM succinate. Wells were then inoculated with 17  $\mu\text{L}$  of the saturated R2B *E.*  
1209 *coli* K12 and *Pseudomonas sp.* PDM04. After sealing, the cultures were incubated at 0.5 rcf and  
1210  $30^{\circ}\text{C}$  in aerobic conditions overnight. These cultures reached saturation during this time. Saturated  
1211 defined media (DM) cultures were washed and normalized to a desired optical density (measured  
1212 at 600 nm) via dilution into pH 7.4 phosphate-buffered saline (8 g/L  $\text{H}_2\text{O}$ , 0.2 g/L KCl, 2.68 g/L  
1213  $\text{Na}_2\text{HPO}_4 \cdot 7\text{H}_2\text{O}$ , 0.24 g/L  $\text{KH}_2\text{PO}_4$ ).

1214 Wells of a sterile 96-deep well plate (Axygen PDW20C) were loaded with carbon-free 1.2 mL  
1215 DM supplemented with 2mM sodium nitrate which had been allowed to equilibrate in the anaerobic  
1216 glovebox. These wells were inoculated in the glovebox with 12  $\mu\text{L}$  of OD-normalized aerobic pre-  
1217 cultures, resulting in starting ODs of 0.1 and 0.01. Additional wells were left blank as no-growth  
1218 controls. Plates were sealed with a gas-permeable sterile membrane. Cultures were incubated at  
1219  $30^{\circ}\text{C}$  and shaken at 950 RPM (Fisherbrand Incubating Microplate Shakers 02-217-759 or Talboys  
1220 Professional 1000MP, 3 mm orbital radius) for 72 h. Optical densities of initial and endpoint anaer-  
1221 obic pre-cultures were measured using 300  $\mu\text{L}$  of cultures in 96-well optical plates. Nitrate and  
1222 nitrite concentrations were assayed over time via manual sampling and subsequent Griess assay

1223 and vanadium (III) chloride reduction via the protocol described in Ref. [73].

### 1224 **Linear metabolite dynamics were recapitulated with monoculture experiments**

1225 Both *E. coli* K12 and *Pseudomonas sp.* PDM04 strains were able to reduce nitrate even without  
1226 carbon in the culture media (Fig. S6A). The reduction rate was negligible for *E. coli* strain at a  
1227 starting OD600 of 0.01 (optical density at 600nm). However, for the denitrifier *Pseudomonas sp.*  
1228 PDM04 at a starting OD600 of 0.01, not only the rate of nitrate reduction was comparable to what  
1229 we observed in soils at the Nutrient-limiting regime (Regime II), but the reduction dynamics were  
1230 strikingly linear. This result directly demonstrates that nitrate reduction can proceed even when  
1231 carbon is not exogenously available. Our observations are consistent with this hypothesis that the  
1232 cells can internally store carbon and oxidize this carbon to provide electrons (NADH) to reduce  
1233 nitrate to nitrite. If the nitrate reduction rate had increased, it would have meant that the functional  
1234 biomass, or the quantity of nitrate reductase enzyme, increased. The nitrate reduction rates did  
1235 not increase throughout the experiment (top panel in Fig. S6A). This supports the idea that cells  
1236 are using nitrate to maintain biomass. Consistently, final OD600 measurements did not detect any  
1237 significant increase from the initial OD600 as expected. We can now more confidently presume  
1238 that soil microbial communities are also in the same maintenance state during the linear nitrate  
1239 dynamics. In sum, these results suggest that the functional biomass in soils can utilize nitrate at a  
1240 constant rate, even after external carbon is no longer available. Note our model assumes this to be  
1241 the case (Fig. 2).

1242 In the monoculture experiment, we observed biphasic behavior in the high initial OD600 con-  
1243 dition. This phenomenon is challenging to interpret. In the starting OD600 of 0.1, the initial slopes  
1244 of nitrate reduction dynamics are constant, then after some time, the rates decrease and remain  
1245 constant (we will call this "late slope") until the end of the experiment (bottom panel of Fig. S6A).  
1246 The linear dynamics observed in the "late slope" again still recapitulate the linear dynamics we  
1247 observed in the soils in the Nutrient-limiting regime, where the microbes could be using nitrate and

1248 internally stored carbon to generate maintenance energy. In the soil experiments, the "late slopes"  
1249 were determined by the increased functional biomass in the model. However, in these monoculture  
1250 experiments, it was difficult to understand what determines the late slope values. Their biomass had  
1251 not changed from the beginning according to the endpoint OD600 measurements, hence requiring  
1252 further investigation of the bacterial physiology. The initial slopes can be roughly explained by  
1253 their starting biomass (Fig. S6B), where the initial slope for the OD600 0.1 condition was roughly  
1254 10 times greater than that for the OD600 0.01 condition, with the fitted initial slopes showing an  
1255 increase factor ranging between 5 to 19 times.

### 1256 **Confirming the linear dependence between functional biomass and acid/base** 1257 **input**

1258 Although the linear relationship between acid/base input with the total biomass increase during  
1259 the Nutrient-limiting regime (Regime II) corroborates our proposed nutrient release mechanism,  
1260 to be more precise, we need to further show that the fold increase of the "functional" biomass is  
1261 equal to the fold increase of nitrate reduction rate from the nitrate dynamics data. However, when  
1262 we observe ASVs increasing in absolute abundance from the start to the end of the experiment  
1263 we cannot assume all ASVs are performing nitrate reduction (for example, some may be grow-  
1264 ing via fermentation). To address this we filtered out the ASVs that are likely not nitrate reducers  
1265 by removing ASVs that were enriched in no-nitrate conditions (dark grey NNresponders bar in  
1266 Fig. S11D, no pH perturbation). To detect the fractional biomass that performs nitrate reduction,  
1267 we used a differential abundance analysis to statistically determine which amplicon sequence vari-  
1268 ants (ASVs) were significantly enriched in each pH perturbed condition compared to the CHL+  
1269 counterpart serving as a baseline of no growth (see Materials and Methods for details). Then, we  
1270 summed up the absolute abundance of these ASVs that we inferred as true nitrate-reducing biomass  
1271 to obtain the functional biomass for each condition (red bar in Fig. S11D). By comparing the fold  
1272 increase of these functional biomass values (endpoint/initial functional biomass), we showed that  
1273 indeed the functional biomass increase and nitrate reduction rate increases are similar in different

1274 soils (Fig. S11C). While some soils showed very close agreement between the inferred increase in  
1275 functional biomass and increases in nitrate reduction rate (Soil 11, inset of Fig. S11C), for many  
1276 soils the relationship was not quantitative. This discrepancy likely arises from the fact that we in-  
1277 ferred the taxa that are not nitrate reducers from slurries where the pH was not perturbed. Thus  
1278 the no-nitrate responders may be distinct as pH is perturbed and this may increase errors in our  
1279 inference of changes in functional biomass.

## 1280 **Investigating the taxonomy, pH niche, and the phylogeny of the Resurgent** 1281 **growth strains**

### 1282 **Determination of peak pH for each family**

1283 To elucidate the pH niche of each family in Fig. S18A, we analyzed the relative abundance of the  
1284 chloramphenicol-untreated (CHL-) conditions of ASVs identified as being enriched in different pH  
1285 levels (see Differential abundance analysis). Due to the challenge of visualizing a large number  
1286 of ASVs, we aggregated the relative abundance of ASVs in the same family for each sample,  
1287 visualizing the data at the family level. To get the representative relative abundance of the family  
1288 in each perturbed pH level, we took the median of the relative abundance from three biological  
1289 replicates. To incorporate abundance values at each perturbed pH from all soils with different  
1290 native pH levels, we placed the perturbed pH values from all native soils from smallest to largest  
1291 and then binned neighboring 2–4 perturbed pH values (depending on the total number of relative  
1292 abundance values greater than 0) to compute the median relative abundance of each family within  
1293 each bin corresponding to its mean pH value. Now, for each family, we have a median relative  
1294 abundance value assigned to the mean pH of each bin. For each family, we ranked those median  
1295 relative abundance values across perturbed pH and found the peak pH value with the highest relative  
1296 abundance, as well as the second peak pH and its corresponding relative abundance. To reduce the  
1297 number of families to plot, we chose 89 families that have relative abundance values at the second  
1298 peak pH greater than 0.002 (= 0.2%). After aligning the 89 families with their peak pH from  
1299 smallest (top of the plot, dark blue color) to largest (bottom of the plot, yellow color), we plotted a

1300 ridge plot with the x-axis being perturbed pH level and height corresponding to the median relative  
1301 abundance of each bin (Fig. S18A). The maximum heights of the ridge for each family were set to  
1302 the same level by normalizing the maximum relative abundance for each family. Indeed, the family  
1303 with peak pH over 8 were mostly Firmicutes phylum (Bacillaceae, Clostridiaceae, Paenibacillaceae,  
1304 Caloramatoraceae, Peptostreptococcaceae, Lachnospiraceae), other than Yersiniaceae family which  
1305 was Proteobacteria.

### 1306 **Constructing a phylogenetic tree with 16S rRNA sequences**

1307 To see whether there was phylogenetic convergence among strains with similar pH niches, we  
1308 used the 16S rRNA sequences of the ASVs to construct a phylogenetic tree (Fig. S18B). To be  
1309 consistent with the previous peak pH analysis, we used the ASVs that belonged to the 89 families  
1310 in the previous analysis. We selected one ASV with the largest relative abundance from each  
1311 family to represent the family and used its 16S rRNA sequence to construct the phylogenetic tree.  
1312 The phylogenetic tree was constructed by approximating the Maximum likelihood tree with the  
1313 General time reversible model in FastTree ver. 2.1.9 [83]. The tree was plotted using the plot.phylo  
1314 function in ape package in R, each node (labeled with the classified genera or species name) colored  
1315 by its peak pH.

### 1316 **Lower-level taxonomy and traits of the Regime III strains**

1317 To identify the specific taxa accountable for the emergence of Regime III at a finer taxonomic  
1318 level, we conducted a differential abundance analysis that statistically determined which Amplicon  
1319 sequence variants (ASVs) were significantly more abundant in Regime III CHL- samples, com-  
1320 pared to CHL+ samples under same perturbed conditions (see Methods). Then, we aggregated the  
1321 relative abundance of these differential ASVs (i.e., Regime III strains) to assess their contribution to  
1322 the emergence of Regime III. Notably, their abundance began to rise between pH 7-8 (Fig. S17C),  
1323 which aligns with or slightly precedes the transition between Regime II and III (Fig. S17D). This  
1324 increase in relative abundance corresponded with the shift of the nutrient growth parameter  $\gamma\tilde{C}(0)$



1325 from zero (Fig. S17E). The analysis revealed that 10 families belonging to Firmicutes (Bacillaceae,  
1326 Paenibacillaceae, Clostridiaceae, Caloramatoraceae, Peptostreptococcaceae, etc.) and 2 families  
1327 belonging to Proteobacteria phylum (Legionellaceae and Yersiniaceae) were significantly enriched  
1328 in Regime III (Fig. S17B). At the genus level, *Bacillus*, *Clostridium*, *Paenibacillus*, and others were  
1329 identified as the primary contributors to Regime III (Fig. S17A).

1330 We lastly sought to find distinct features of the Regime III strains that differentiated them from  
1331 other strains to understand why these strains better adapt to perform nitrate reduction in high pH  
1332 and high carbon conditions. To do so, we classified families by their peak pH obtained by finding  
1333 the pH level at which its median relative abundance across soils was the highest across different  
1334 perturbed pH levels (Fig. S17A). We indeed found that the Regime III families had distinct pH  
1335 niches compared to other strains, having high relative abundance in basic pH (over 8) and in some  
1336 cases acidic pH (less than 4) but remained rare (< 0.1%) in the mid-range of pH 4-8. One can  
1337 speculate that their ability to survive and persist in extreme pH perturbations (see Fig. S20A) may be  
1338 because many taxa in the phylum Firmicutes are spore-forming bacteria species [84]. These strains  
1339 did not cluster phylogenetically and were dispersed throughout the phylogenetic tree (Fig. S17B,  
1340 see Methods).

## 1341 **pH titration curves and soil's native pH are shaped by soil's physicochemical** 1342 **properties**

1343 We've constructed pH titration curves for the 20 soils from different native pH levels (see Methods,  
1344 Fig. 6B). Because we titrated both in acid and basic directions with H<sup>+</sup> and OH<sup>-</sup> respectively, we  
1345 unified the x-axis to OH<sup>-</sup> (m mol) by shifting the curves to the right by 0.2 m mol, ensuring each  
1346 curve starts at 0 m mol OH<sup>-</sup>. We then fitted the pH titration curves with a logistic function with 4  
1347 parameters (*a*, *x<sub>mid</sub>*, *b*, *c*) as below (visualized in Fig. S23B):

$$pH = \frac{a}{1 + e^{\frac{-(x-x_{mid})}{b}}} + c \quad (4)$$

1348 Parameter  $x_{mid}$  strongly correlated with soil's native pH level ( $R^2 = 0.8$ , Fig. S23D), while parame-  
1349 ter  $c$  (y-intercept) and  $a+c$  (asymptotic y value), scaling parameter ( $b$ ) stayed mostly constant across  
1350 soils with different pH levels. This indicates that the titration curve's general shape is similar for  
1351 all soils, but the titration curve shifts horizontally depending on the soil's original pH level.

1352 Soil's native pH, which determines the horizontal shift of the titration curves, was strongly cor-  
1353 related to the cation exchange capacity (CEC, milliequivalent charge / 100g) ( $R^2 = 0.88$ , Fig. S23D).  
1354 This was expected because soils with higher CEC will have a greater number of negative charges in  
1355 the clay particles and hence more likely to adhere to protons. This will result in fewer free protons  
1356 in the soil pore water and, thus result in more basic pH levels. In the literature, CEC is reported to  
1357 be determined by soil's clay particles and its organic matter, because CEC is proportional to how  
1358 much negative charge the soil has on the aggregate's surface. However, in our dataset, the percent  
1359 clay and organic matter did not correlate strongly with CEC (Fig. S23D, see Fig. S23E for percent  
1360 clay). CEC appeared to be determined by  $Ca^{2+}$  ion concentration in the soil and not by other  
1361 cations ( $Mg^{2+}$ ,  $K^+$ ,  $Na^+$ ). Soil pH was inversely correlated with S, P, Al, and Fe concentrations,  
1362 which can either be the cause or result of the soil's pH (Fig. S23D). In sum, we can attribute the  
1363 horizontal shift of the pH titration curves to their varying native soil pH levels, which is potentially  
1364 determined by the CEC and the  $Ca^{2+}$  ion concentrations (see summarized diagram in Fig. S23C).

## 1365 **Evidence for long-term pH adaptation from phyla's differential response to** 1366 **pH perturbations**

1367 We observed that long-term pH variation (different native soil pH) shifts the pH boundaries between  
1368 functional regimes (Fig. 6). To see if those shifts of pH boundary can be explained by taxa's differ-  
1369 ential response to perturbed pH due to long-term pH adaptation, we further asked whether the pH  
1370 values where the abundances of taxa (Firmicutes, Bacteroidota, and Proteobacteria) exhibit large  
1371 changes also agree with the boundaries between regimes determined solely by nitrate utilization dy-  
1372 namics (Fig. 3). We observed the growth folds of taxa for the transition from Regime II to Regime  
1373 III and the survival folds of taxa for the transition from Regime II to Regime I (Fig. S21A). Growth

1374 folds were computed by endpoint absolute abundance ratio of  $Abs_{CHL-}/Abs_{CHL+}$  (chlorampheni-  
1375 col untreated/treated conditions) and survival folds were computed by absolute abundance ratio of  
1376  $Abs_{CHL+}/Abs_{T_0}$ , representing taxa's endpoint absolute abundance in CHL+ conditions compared  
1377 to the initial time point ( $T_0$ ) for each perturbed pH level.

1378 To understand the transition to the Acidic death regime (Regime I), we observed the survival  
1379 folds of Proteobacteria and Bacteroidota phyla across perturbed pH levels. Then, we set an identical  
1380 survival fold threshold for all soils (red lines in Fig. S21) to compute the pH at which the survival  
1381 fold goes below that threshold during acidic perturbation. We used two distinct definitions to  
1382 choose a threshold for the survival fold. The first was a definition of “dying” where the taxa's  
1383 abundance started to decline in abundance compared to  $T_0$  (survival fold threshold  $< 1$ , red solid  
1384 lines in Fig. S21A). The second was a definition of “dead” where the taxa's abundance was close  
1385 to 0 (survival fold threshold  $\rightarrow 0$ , red dashed lines in Fig. S21A). For each of these definitions,  
1386 the pH transition points were plotted (Fig. S21B with the first “dying” definition and Fig. S21C  
1387 with the second “dead” definition) and compared to the trends of functional regime boundaries  
1388 (transition from Regime II to I). Employing the ‘dying’ definition with Proteobacteria, Bacteroidota  
1389 (Fig. S21B) allowed us to recapitulate the phenomenon observed in the functional data, where the  
1390 fitted slope of Boundary I-II was less than 1, as shown in Fig. 6A. This suggests that these phyla in  
1391 the relatively neutral soil are more tolerant of larger  $\Delta$ pH change until they start to die than those in  
1392 acidic soils, possibly due to variations in soil titration curves (Fig. 6B). Because the fitted slope is  
1393 greater than 0, this also means that these phyla in relatively acidic soils can tolerate lower acidic pH  
1394 conditions than those in neutral soils, which signals long-term pH adaptation. The ‘dead’ definition  
1395 threshold resulted in a flat slope close to 0. This suggests that, despite long-term adaptation to  
1396 varying native soil pH levels, these taxa have similar pH thresholds at which complete decimation  
1397 occurs.

1398 Similarly, to understand the transition to the Resurgent growth regime (Regime III), we ob-  
1399 served the growth folds of Firmicutes phylum across perturbed pH levels. Then, we applied an

1400 identical growth fold threshold for all soils (red lines in Fig. S21A) to compute the pH at which  
1401 the growth fold goes above the threshold during basic perturbations. These pH transition points  
1402 were plotted (Fig. S21B&C) and compared to the trends of functional regime boundaries (transi-  
1403 tion from Regime II to III). Consistent with the trend of functional regime boundary II-III (Fig. 6A),  
1404 the abundance of Firmicutes began to increase at higher pH values as the native soil pH increased.  
1405 Since the NaOH amount, and consequently, the carbon nutrient level, remains constant at the pH  
1406 boundary of II-III (Fig. S24), the reason Firmicutes increases at higher pH values is not linked to  
1407 the amount of nutrients available. Therefore, this is another signal for taxa adaptation to long-term  
1408 pH variation.

## 1409 **Supplementary Tables**

Table S1: Varying experimental conditions in previous studies

Paper	Soil/source	Carbon	Chloramphenicol	Water content	Measurement (time)
Wijler & Delwiche (1954)	1 soil's pH modified by H <sub>2</sub> SO <sub>4</sub> or KOH	Dried alfalfa added	Untreated	Added water and dried	N <sub>2</sub> O/N <sub>2</sub> gas (17 days)
Nömmik (1956)	6 soils (pH 5.2-7.6), each pH modified by HCl or KOH	Untreated, wheat straw, glucose conditions	Untreated	100% water saturation	N <sub>2</sub> O/N <sub>2</sub> gas, NO <sub>3</sub> <sup>-</sup> , NO <sub>2</sub> <sup>-</sup> , NH <sub>4</sub> <sup>+</sup> (12 days)
Bremner & Shaw (1958)	8 soils (pH 3.6-8.6) and 2 of those soils (pH 3.6-4.1)'s pH modified by CaOH	Glucose	Untreated	Waterlogged (slurry)	Loss of N (30 days)
Valera & Alexander (1961)	5 pure cultures of denitrifiers tested in buffered media of different pH levels	Glucose and yeast extract	Untreated	Defined buffered medium	N <sub>2</sub> gas
Burford & Bremner (1975)	17 soils (pH 5.8-7.8)	Untreated	Untreated	Waterlogged (slurry)	Nitrous oxide/NO/N <sub>2</sub> gas, NO <sub>3</sub> <sup>-</sup> , NO <sub>2</sub> <sup>-</sup> , NH <sub>4</sub> <sup>+</sup> (7 days)
Van Cleemput et al. (1975)	1 soil's pH modified by HCl or NaOH	Dried ground plant material	Untreated	Waterlogged (slurry)	N <sub>2</sub> O/N <sub>2</sub> gas, NO <sub>3</sub> <sup>-</sup> , NO <sub>2</sub> <sup>-</sup> (100 hrs)
Smith et al. (1978)	4 soils (pH 6.4-7.6)	Untreated	Chloramphenicol and untreated	Waterlogged (slurry)	N <sub>2</sub> O while acetylene inhibition (10 hrs)
Smith & Tiedje (1979)	3 soils and pure cultures of denitrifiers	Succinate or glucose added and untreated	Chloramphenicol and untreated	Waterlogged (slurry)	N <sub>2</sub> O while acetylene inhibition (10 hrs)
Koskinen & Keeney (1982)	4 soils (pH 4.6-6.9)	Untreated	Untreated	20% water content	N <sub>2</sub> O/NO/N <sub>2</sub> gas, NO <sub>3</sub> <sup>-</sup> , NO <sub>2</sub> <sup>-</sup> , NH <sub>4</sub> <sup>+</sup> (16 days)
Waring & Gilliam (1983)	15 acidic soils (pH 3.42-5.54) and 5 of those soils limed to increase pH	Glucose and untreated	Untreated	Waterlogged (slurry)	NO <sub>3</sub> <sup>-</sup> , NO <sub>2</sub> <sup>-</sup> (20 days)

Paper	Soil/source	Carbon	Chloramphenicol	Water content	Measurement (time)
Parkin et al. (1985)	2 soils (pH 4, 6), each pH modified by H <sub>2</sub> SO <sub>4</sub> or NaOH	Glucose for short-term (DEA), untreated for long-term (DP)	Chloramphenicol (DEA) and untreated (DP)	Waterlogged (slurry) for DEA	N <sub>2</sub> O while acetylene inhibition. Denitrifying enzyme activity (DEA) (2 hrs), Denitrification potential (DP) (5 days)
Nägele & Conrad (1990)	3 soils (pH 4-7.8), each pH modified by HCl or NaOH	Glucose	Chloramphenicol and untreated	Waterlogged (slurry)	N <sub>2</sub> O/NO/N <sub>2</sub> gas, NO <sub>3</sub> <sup>-</sup> , NO <sub>2</sub> <sup>-</sup> , NH <sub>4</sub> <sup>+</sup> (20 hrs)
Drury et al. (1991)	13 soils (pH 4.88-7.79)	Glucose for denitrification potential, untreated for background denitrification rate	Untreated	Field capacity	N <sub>2</sub> O while acetylene inhibition (75 hrs)
Bandibas et al. (1994)	18 soils (pH 4-7.8)	Untreated	Untreated	Field capacity, saturation, waterlogged	N <sub>2</sub> O/N <sub>2</sub> gas, NO <sub>3</sub> <sup>-</sup> , NO <sub>2</sub> <sup>-</sup> , NH <sub>4</sub> <sup>+</sup> (20 days)
Yamulki et al. (1997)	3 soils (pH 3.9, 5.9, 7.6) and the acid soil (pH 3.9)'s pH modified by NaOH	Untreated	Untreated	Unadjusted	N <sub>2</sub> O/NO/N <sub>2</sub> gas, NO <sub>3</sub> <sup>-</sup> , NO <sub>2</sub> <sup>-</sup> (6hrs), field measurements (12 months)
Ellis et al. (1998)	4 soils (pH 3.3-6.1)	Untreated	Chloramphenicol and untreated	Waterlogged (slurry) for anaerobic condition and aerobic condition	N <sub>2</sub> O with and without acetylene inhibition, NO <sub>3</sub> <sup>-</sup> , NO <sub>2</sub> <sup>-</sup> , NH <sub>4</sub> <sup>+</sup> (48 hrs)
Šimek et al. (2000)	13 soils (pH 5.62-7.77)	Glucose	Chloramphenicol (DEA) and untreated (DP)	Waterlogged (slurry)	N <sub>2</sub> O while acetylene inhibition. Denitrifying enzyme activity (DEA) (30-60 min), Denitrification potential (DP) (48 hrs)

Paper	Soil/source	Carbon	Chloramphenicol	Water content	Measurement (time)
Simek et al. (2002)	5 soils (pH 4.9-7.9), each pH modified by H <sub>2</sub> SO <sub>4</sub> or NaOH	Glucose	Chloramphenicol (DEA) and untreated (DP)	Waterlogged (slurry)	N <sub>2</sub> O while acetylene inhibition. Denitrifying enzyme activity (DEA) (30 min-3 hrs), Denitrification potential (DP) (48 hrs)
Liu et al. (2010)	3 soils (pH 3-4), each pH modified by long-term liming	Glutamate and untreated	Untreated	Unadjusted (1st run), flooding and draining (2nd run)	N <sub>2</sub> O/N <sub>2</sub> gas (21 hrs)
Cuhel et al. (2010)	3 soils (pH 5.5-7.7), each pH modified by 10-month application of H <sub>2</sub> SO <sub>4</sub> and KOH	Glucose	Chloramphenicol for short-term (DEA), untreated for in-situ field chamber measurements	Waterlogged (slurry) for DEA	N <sub>2</sub> O/N <sub>2</sub> gas, NO <sub>3</sub> -, NO <sub>2</sub> -, NH <sub>4</sub> <sup>+</sup> . In-situ field chamber measurements (74 hrs), Denitrifying enzyme activity (DEA) (30-60 min)
Bergaust et al. (2010)	1 pure culture in 5 pH levels (pH 6.0-7.5)	Succinate	Untreated	Defined buffered medium	N <sub>2</sub> O/NO/N <sub>2</sub> gas, NO <sub>3</sub> -, NO <sub>2</sub> - (72 hrs)
Dörsch et al. (2012)	Cells extracted from 3 soils (pH 5.4-7.1) perturbed to pH 5.4 and 7.1	Glutamate	Untreated	Defined buffered medium	N <sub>2</sub> O/NO/N <sub>2</sub> gas, NO <sub>3</sub> -, NO <sub>2</sub> - (130 hrs)
Samad et al. (2016)	13 soils (pH 5.57-7.03)	Untreated	Untreated	Flooded and drained	N <sub>2</sub> O/NO/N <sub>2</sub> gas (200 hrs)

Table S2: Relevant conclusions from previous studies

Paper	Conclusions relevant to this paper
Wijler & Delwiche (1954) [85]	Total denitrification rates were quite constant above pH 6.
Nömmik (1956) [63]	Total denitrification rates are low below pH 5 and increased until 7-8. Further increase in pH suppressed denitrification. Different soils showed similar trends.
Bremner & Shaw (1958) [59]	When compared with soils without pH modification, soil with native pH of 7.5 had a higher denitrification rate. When pH was increased in soils with acidic pH levels, the denitrification rate increased. Denitrification potential correlated very well with mineralizable carbon. Easily decomposable substances (glucose, mannitol, and sucrose) have greater effects on increasing denitrification rate than difficultly decomposable materials (lignin and sawdust).
Valera & Alexander (1961) [86]	Pure cultures exhibited the highest denitrification rate between pH 7-8.
Burford & Bremner (1975) [36]	Denitrification capacity was very highly correlated with water-soluble organic carbon and mineralizable carbon, and less but significantly correlated with total organic carbon in soils.
Van Cleemput et al. (1975) [11]	Soils in very acidic conditions also show substantial denitrification. Authors suggested pH might affect the availability of readily decomposable organic matter, which correlated with the denitrification rate.
Smith et al. (1978) [55]	The denitrification rates increased from the 0hr-3hr period until a linear rate was attained. The linear rate corresponded to mineralizable carbon.
Smith & Tiedje (1979) [31]	Chloramphenicol does not affect the denitrifying rate in 0-3hrs, called linear Phase I, but prevented the rate from increasing during the transition to linear Phase II (4-8hrs) compared to the untreated samples. Even when glucose was added, the linear rate of Phase I did not change or just increased slightly, but it later prompted a logarithmic increase of denitrifying activity. Irrigating the soils before the incubation increased the linear rate of Phase I. Therefore, it can be concluded that Phase II is determined by a state where derepression of de novo enzyme synthesis is complete and therefore impacted by available electron donor, pH, and temperature, while Phase I rate is determined by indigenous enzyme activity.
Koskinen & Keeney (1982) [10]	Denitrification rate was correlated with mineralizable carbon but not consistently related to pH or total organic carbon. Therefore, the rate of organic C mineralization rather than pH controls the rate of denitrification in C-limited systems.
Waring & Gilliam (1983) [87]	Soluble organic carbon is more important in denitrification than pH. Significant denitrification can also happen in strongly acidic soils (below pH 4).



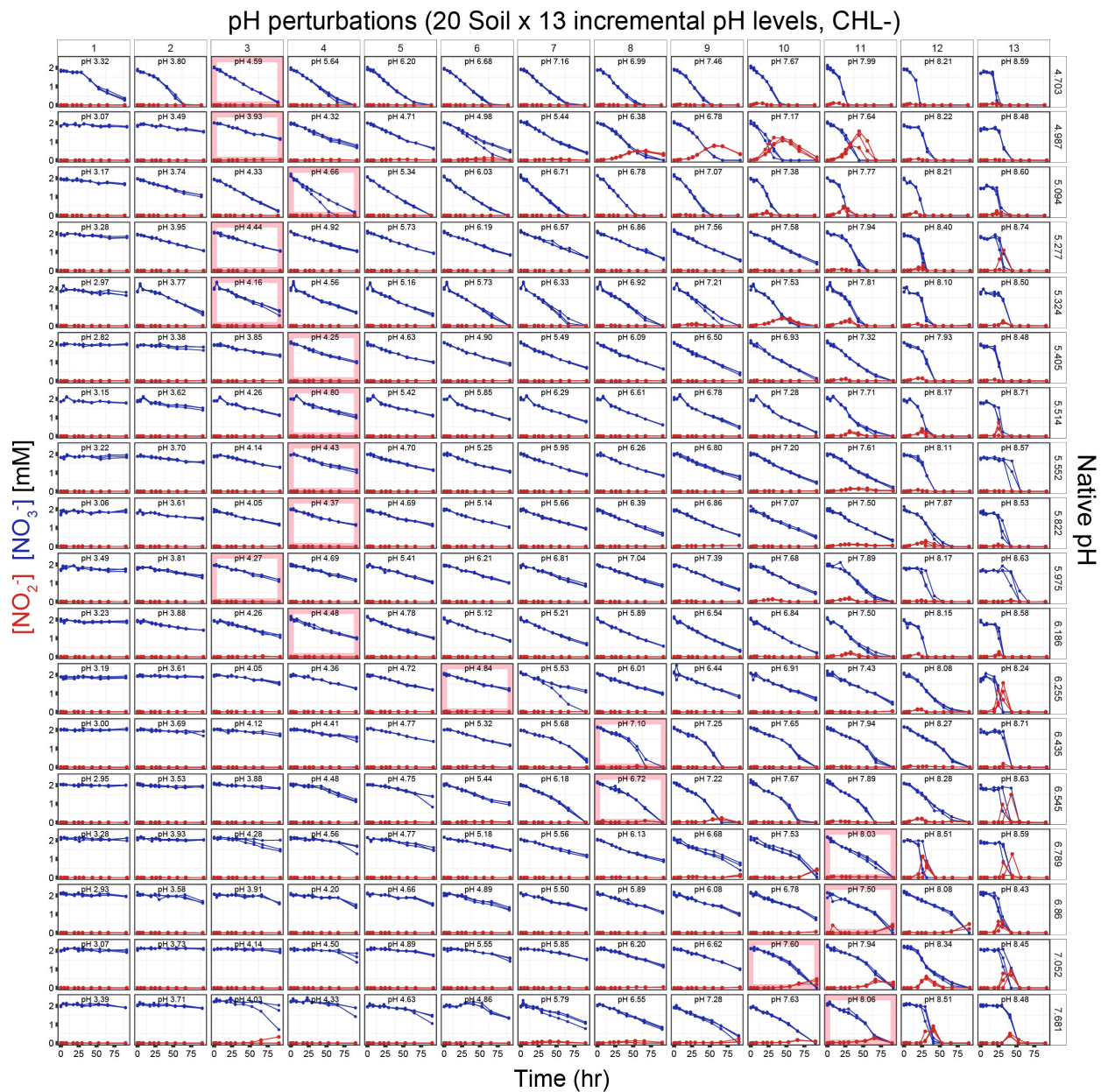
Paper	Conclusions relevant to this paper
Parkin et al. (1985) [33]	Both natural denitrification rates and denitrifying enzyme activity (DEA) were greater in neutral soil, but denitrification rate of similar magnitude occurred in acid soil, which can be due to long-term selection of acid-tolerant denitrifying populations. With DEA measurements (activity of existing enzymes), acid soil (native pH 4) had the highest denitrification rate at 3.9 and neutral soil (native pH 6) had an optimal pH at 6.3.
Nägele & Conrad (1990) [88]	Unlike Parkin et al. (1985), under chloramphenicol treatment, acid soil (native pH 4) had greater reduced nitrate when pH was adjusted to 7 compared to pH 4. On the other hand, neutral soil (native pH 7.8) had greater reduced nitrate when pH was 7 and decreased as pH was lowered.
Drury et al. (1991) [58]	For background denitrification rate (C, nitrate unamended), denitrification rates were highly correlated with biomass C, which was also highly correlated with available organic carbon. This shows that soil denitrification is limited by available organic carbon. Background and potential denitrification rates did not correlate with each other. pH nor other soil physicochemical properties did not correlate with either background and potential denitrification rate (other than the correlations stated above).
Bandibas et al. (1994) [89]	Saturated conditions produced the most N <sub>2</sub> O compared to field capacity and waterlogged moisture conditions. Soil pH was the soil characteristic that best predicted N <sub>2</sub> O emission.
Yamulki et al. (1997) [32]	Field measurements did not show a clear effect of pH on N <sub>2</sub> O emissions. The highest N <sub>2</sub> O emission rate was observed after a period of relatively high rainfall in the pH 5.9 plot. Average N <sub>2</sub> O emission rates were higher in the pH 7.6 and 5.9 plots than in the pH 3.9 plot. In lab measurements. When the soil's pH was increased in the lab from pH 3.9 to above 6, NO <sub>2</sub> emission dropped 40%.
Ellis et al. (1998) [56]	In anaerobic conditions, denitrification rates increased with higher pH levels both in CHL+ and CHL- conditions, which is contrary to Parkin et al. (1985) probably due to longer (48hr) time scale. Chloramphenicol reduced the denitrification rate in all soils with different pH levels. However, the CO <sub>2</sub> production was not significantly reduced in anaerobic conditions. Increasing pH increased the CO <sub>2</sub> production rate in both CHL+ and CHL- conditions.
Šimek et al. (2000) [60]	No significant relationship was found between short-term denitrifying enzyme activity (DEA) and longer-term denitrification potential (DP), indicating the independence of the existing metabolic activity of the denitrifier community (DEA) and its potential for rapid development when the environmental conditions change (DP). Existing enzyme activity did not correlate with pH. However, denitrification potential positively correlated with pH, available organic carbon, and biomass.

Paper	Conclusions relevant to this paper
Šimek et al. (2002) [17]	In agreement with Parkin et al. 1985, for existing denitrifying enzymes, the denitrification rate was highest close to their native pH. Optimal pH shifted to neutral pH (pH 6 - 8) as denitrification rate was measured in longer periods. It is proposed that this is either due to the development of a community of denitrifiers that can grow better at the neutral pH or due to the accommodation of the existing populations to new conditions.
Liu et al. (2010) [90]	Soil pH has little effect on denitrification rate apart from the low denitrification rates in the very acid peat soil (pH 4.0). For acidic soil, neither the gene pools ( <i>nirS</i> vs. <i>nosZ</i> ) nor their transcription rates could explain the observed effects of low pH on N <sub>2</sub> O reductase activity, which implies that this low N <sub>2</sub> O reductase activity is due to post-transcriptional level, either by interfering with translation, protein assembly or by directly affecting the activity of the functional enzyme. Denitrification in unamended soil appeared to be based on the activation of a pre-existing denitrification proteome, because constant rates of N <sub>2</sub> and N <sub>2</sub> O production were observed, and the transcription of functional genes was below the detection level, whereas glutamate-amended soils showed sharp peaks in the transcripts of <i>nirS</i> and <i>nosZ</i> , increasing the denitrification rates.
Cuhel et al. (2010) [18]	Denitrification rates were higher when soil's pH was perturbed to higher pH and were lower when perturbed to lower pH. Microbial biomass C followed a similar pattern.
Bergaust et al. (2010) [7]	Pure culture experiment of <i>Paracoccus denitrificans</i> revealed that the denitrification rate was highest at pH 7 and was slower at pH 6. This can be explained by the change in transcription of genes, where the maximum numbers of <i>norB</i> and <i>nosZ</i> transcripts were higher at pH 7 than at pH 6 and the maximum number of <i>nirS</i> transcripts was higher at pH 7 than at pH 6. However, as in Liu et al. 2010, transcription cannot explain the low N <sub>2</sub> O reduction (N <sub>2</sub> OR) rate at pH 6. They rule out the possibility of N <sub>2</sub> OR enzyme activity directly inhibited by low pH, because functional N <sub>2</sub> O reductase activity was only marginally affected by lowering pH to 6. Therefore, the loss of N <sub>2</sub> OR activity is due to unsuccessful assemblage/folding of the protein due to low pH in the periplasm.
Dorsch et al. (2012) [91]	Denitrification rates in extracted communities are higher in neutral pH 7.1 than in pH 5.4.
Samad et al. (2016) [61]	The denitrification rate (NO+N <sub>2</sub> O+N <sub>2</sub> $\mu\text{mol N/h/vial}$ ) was higher in more neutral soils. Emission kinetics across all soils revealed that the denitrification rates under anoxic conditions were significantly associated with C mineralization (CO <sub>2</sub> $\mu\text{mol/h/vial}$ ).

Table S3: Physical and chemical characteristics of soil samples

Soil No.	Soil ID	$pH_{H_2O}$	Latitude	Longitude	Sand (%)	Silt (%)	Clay (%)	Total N (%)	Total C (%)	C:N ratio	Depth	Date
Soil1	Acidic4	4.703	46.785703	-117.079245	0.00	58.15	41.85	0.13	1.61	12.79	0-10cm	9/10/22
Soil2	Acidic12	5.094	46.781117	-117.080513	0.00	63.80	36.20	0.15	1.97	13.19	0-10cm	9/12/22
Soil3	CE239	4.987	46.781153	-117.080455	0.00	61.20	38.80	0.11	1.35	12.27	10-20cm	9/8/22
Soil4	SE56b	5.277	46.778744	-117.082738	0.00	57.40	42.60	0.10	1.26	13.31	10-20cm	9/8/22
Soil5	CE201	5.324	46.781049	-117.086242	0.00	58.70	41.30	0.12	1.57	13.07	10-20cm	9/8/22
Soil6	CE73	5.405	46.779637	-117.086172	0.00	61.20	38.80	0.13	1.65	12.97	10-20cm	9/8/22
Soil7	CE153	5.514	46.7805	-117.085431	0.00	58.70	41.30	0.13	1.76	13.82	10-20cm	9/8/22
Soil8	CE56a	5.552	46.778855	-117.082968	0.00	58.70	41.30	0.10	1.35	13.64	10-20cm	9/8/22
Soil9	CE277	5.822	46.781883	-117.0835	0.00	61.20	38.80	0.09	1.19	12.65	10-20cm	9/8/22
Soil10	CE253	5.975	46.781534	-117.084192	0.00	62.50	37.50	0.08	0.98	12.00	10-20cm	9/8/22
Soil11	CE234	6.186	46.781301	-117.082533	0.00	62.50	37.50	0.13	1.88	14.04	10-20cm	9/8/22
Soil12	CE229	6.255	46.781206	-117.084623	0.00	58.80	41.20	0.09	1.20	12.73	10-20cm	9/8/22
Soil13	Neutral7	6.435	46.781523	-117.084533	0.00	60.00	40.00	0.08	0.99	12.06	10-20cm	9/11/22
Soil14	Neutral2	6.545	46.781308	-117.084696	0.00	58.80	41.20	0.11	1.28	12.06	10-20cm	9/11/22
Soil15	Neutral5	6.789	46.781416	-117.084818	0.00	63.70	36.30	0.10	1.31	13.79	10-20cm	9/11/22
Soil16	Neutral6	6.860	46.781524	-117.084694	0.00	61.20	38.80	0.09	1.07	12.30	10-20cm	9/11/22
Soil17	Neutral3	7.052	46.781194	-117.084732	0.00	60.00	40.00	0.09	1.09	12.56	10-20cm	9/11/22
Soil18	Neutral1	7.681	46.781354	-117.084812	0.00	63.70	36.30	0.10	1.21	12.37	10-20cm	9/11/22
Soil19	Neutral4	8.232	46.781222	-117.084882	0.00	62.50	37.50	0.08	1.05	13.78	10-20cm	9/11/22
Soil20	CE251	8.323	46.781492	-117.085028	0.00	67.50	32.50	0.09	1.47	16.14	10-20cm	9/8/22

## 1410 **Supplementary Figures**



**Figure S1: Flux dynamics of nitrate and nitrite of the dataset.** Time series measurements of nitrate (blue points) and nitrite (red points) across 4 days are shown. (Continued)

Figure S1: (Continued from the previous page) Each row is from the identical soil sample of a native pH level ( $\text{pH}_{H_2O}$ ), indicated at the right end of each row in the order of most acidic (top) to most basic (bottom). Each row has 13 columns which are the 13 different levels of short-term pH perturbations. The targeted perturbed pH levels were determined by constructing a soil pH titration curve before the experiment and computing how much acid (HCl) or base (NaOH) to add to the slurries. Perturbed pH levels are indicated inside each panel, which are obtained by measuring the stabilized pH values at the endpoint of the experiment (see Methods). Each line connects the point of measurements of a replicate, constituting the 3 replicates per perturbed condition. The pink-colored box for each row indicates the condition without any acid/base addition, where the pH of these conditions also changes with incubation. Soil 19 and Soil 20 are not shown due to having different numbers of perturbed pH levels (7 and 3, respectively).

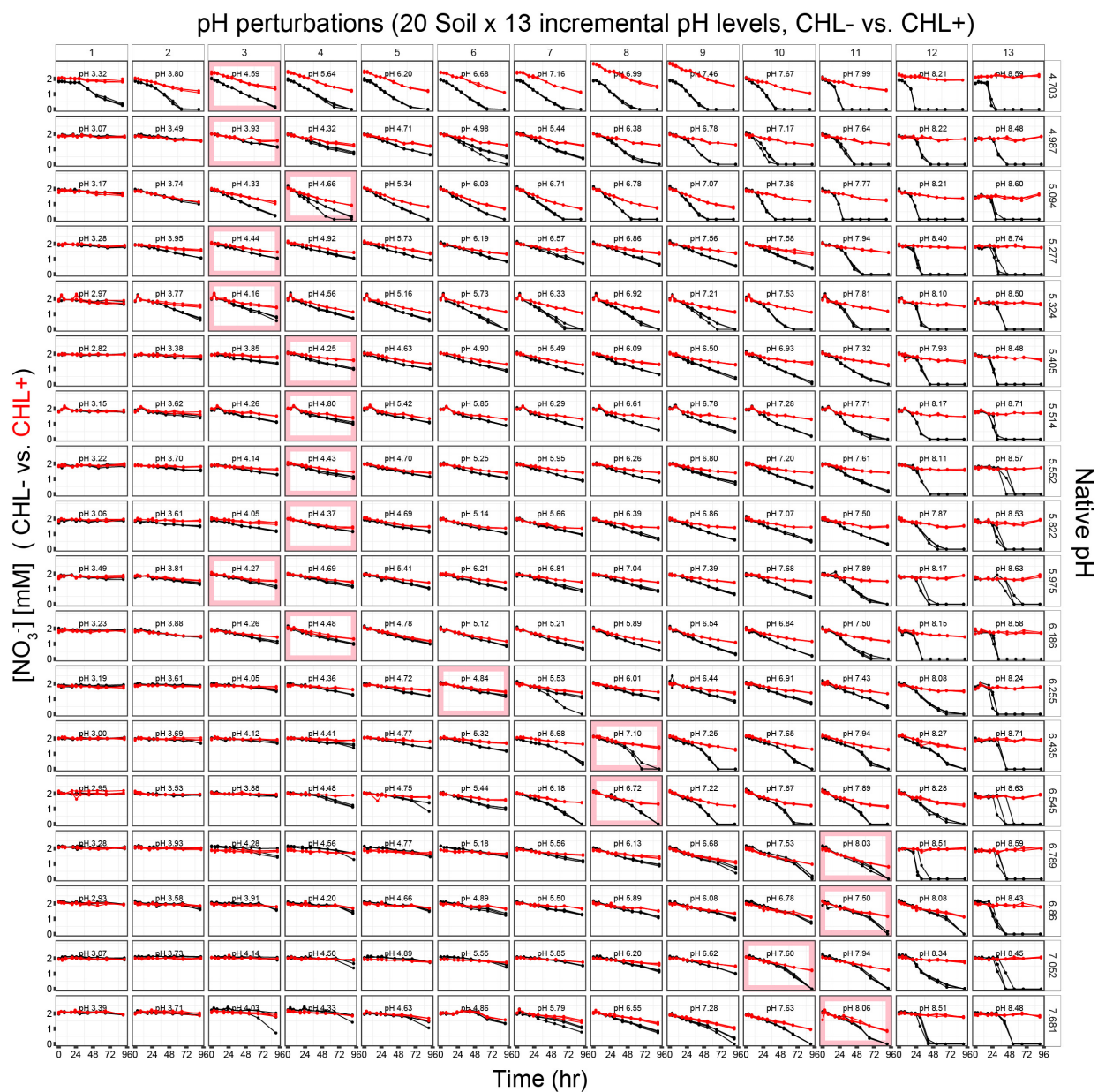
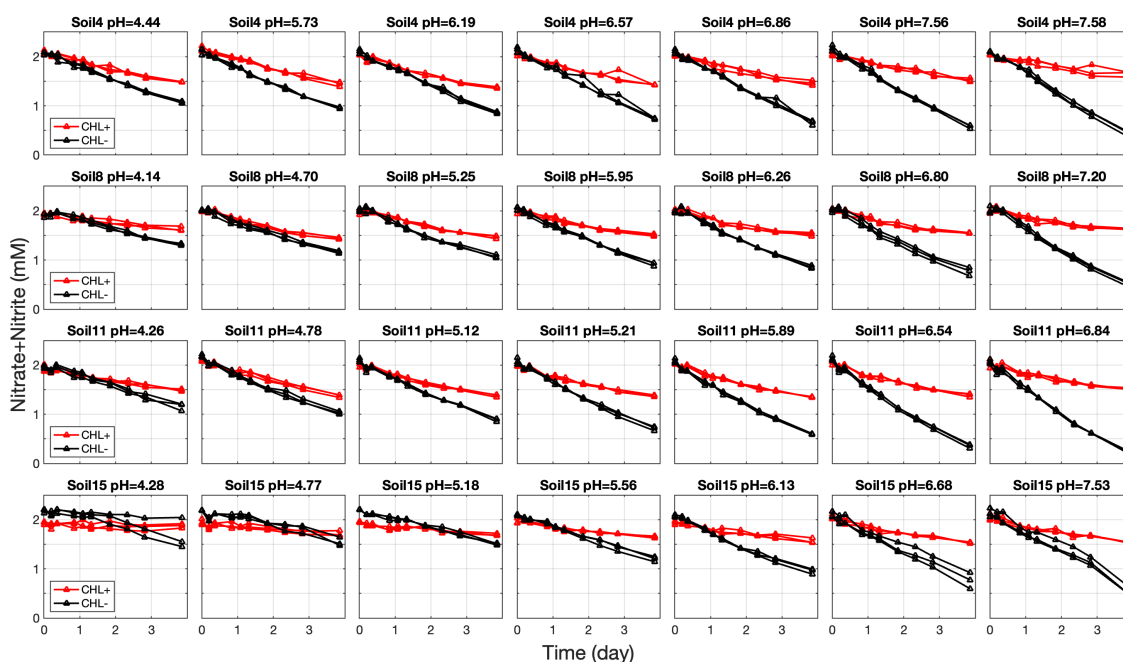


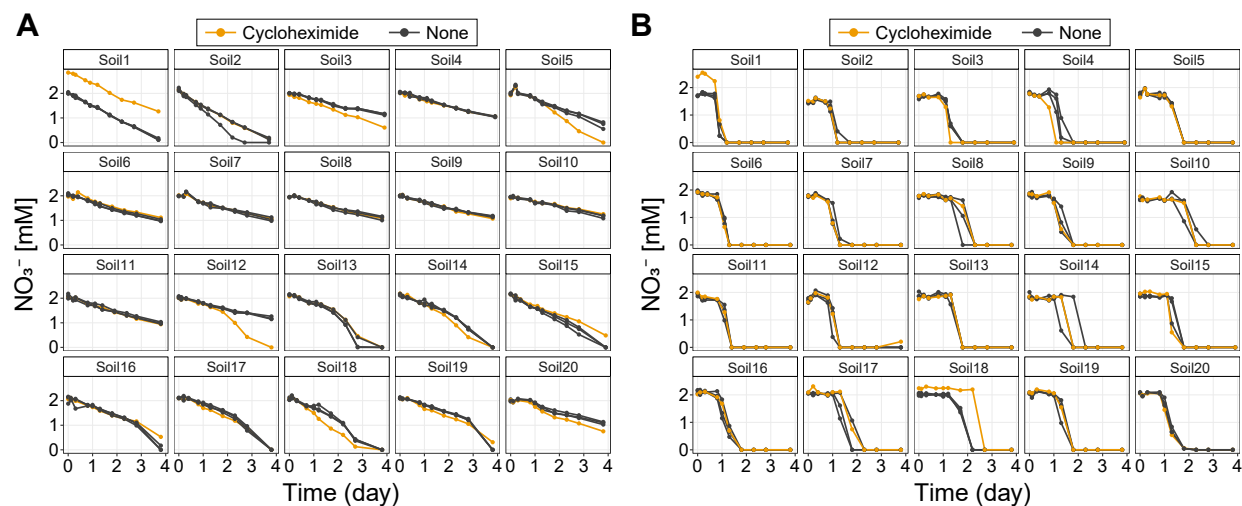
Figure S2: Nitrate dynamics of chloramphenicol untreated (CHL-) and treated (CHL+) conditions in the dataset. Time series measurements of nitrate in chloramphenicol-untreated (CHL-, black points) and treated (CHL+, red points) across 4 days are shown. (Continued)

Figure S2: (Continued from the previous page) Each row is from the identical soil sample of a native pH level ( $\text{pH}_{H_2O}$ ), indicated at the right end of each row in the order of most acidic (top) to most basic (bottom). Each row has 13 columns which are the 13 different levels of short-term pH perturbations. The targeted perturbed pH levels were determined by constructing a soil pH titration curve before the experiment and computing how much acid (HCl) or base (NaOH) to add to the slurries. Perturbed pH levels are indicated inside each panel, which are obtained by measuring the stabilized pH values at the endpoint of the experiment (see Methods). Each line connects the point of measurements of a replicate, constituting the 3 replicates per perturbed condition. The pink-colored box for each row indicates the condition without any acid/base addition, where the pH of these conditions also changes with incubation. Soil19 and Soil20 are not shown due to having different numbers of perturbed pH levels (7 and 3, respectively).





**Figure S3: Nitrate + nitrite concentration dynamics to show constant nitrite reduction rates.** The points indicate the time-series measurement of the sum of nitrate and nitrite concentrations. Concentrations from chloramphenicol-treated (CHL+) samples are in red and untreated (CHL-) samples are in black, with lines connecting each of the three biological replicates. A subset of pH perturbed conditions (each row is from the same native pH soil, with varying perturbed pH levels) is shown. Nitrate + nitrite dynamics (in CHL- conditions) are linear, indicating that the community's nitrite consumption is constant across time.



**Figure S4: Cycloheximide antifungal controls suggest a minimal role for fungi in nitrate utilization dynamics.** Nitrate dynamics across a 4-day anaerobic incubation with and without cycloheximide treatment for all 20 soils. Panel (A) shows the nitrate dynamics of pH-unperturbed samples with (orange data points, 1 replicate) and without cycloheximide (black data points, 3 biological replicates) treatment, while (B) illustrates the nitrate dynamics for basic-perturbed samples, also with (orange, 1 replicate) and without cycloheximide (black, 3 biological replicates) treatment. Most of the dynamics were not affected by the application of 200 ppm cycloheximide. Only 1 case out of 40 samples (Soil 18 in B) showed delayed nitrate reduction when the antifungal was treated. This means that fungi do not play a significant role during nitrate reduction.

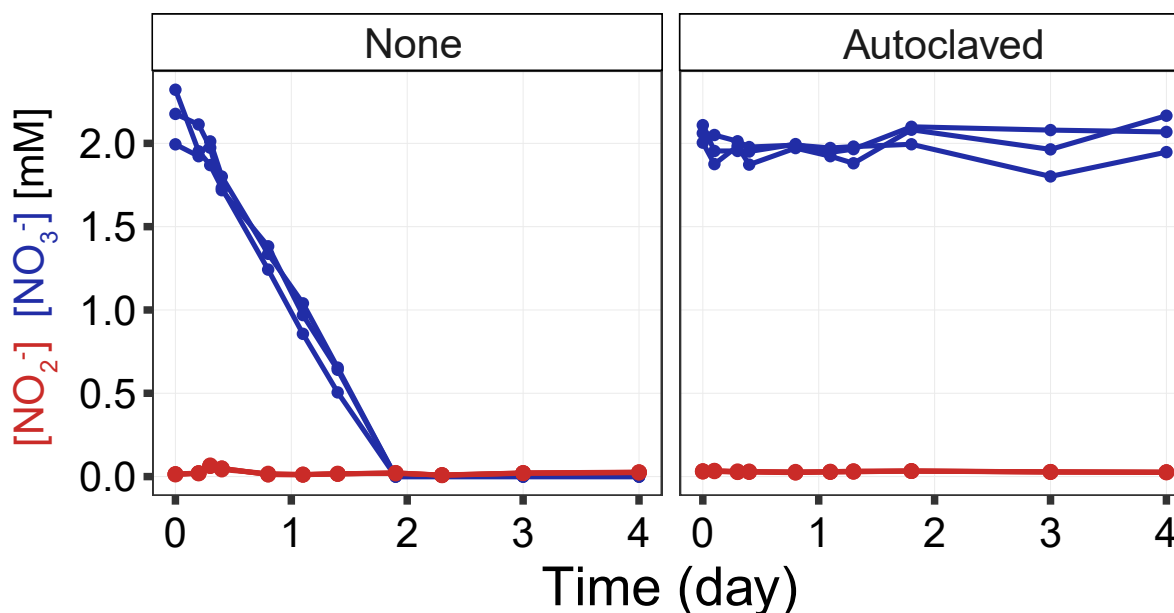
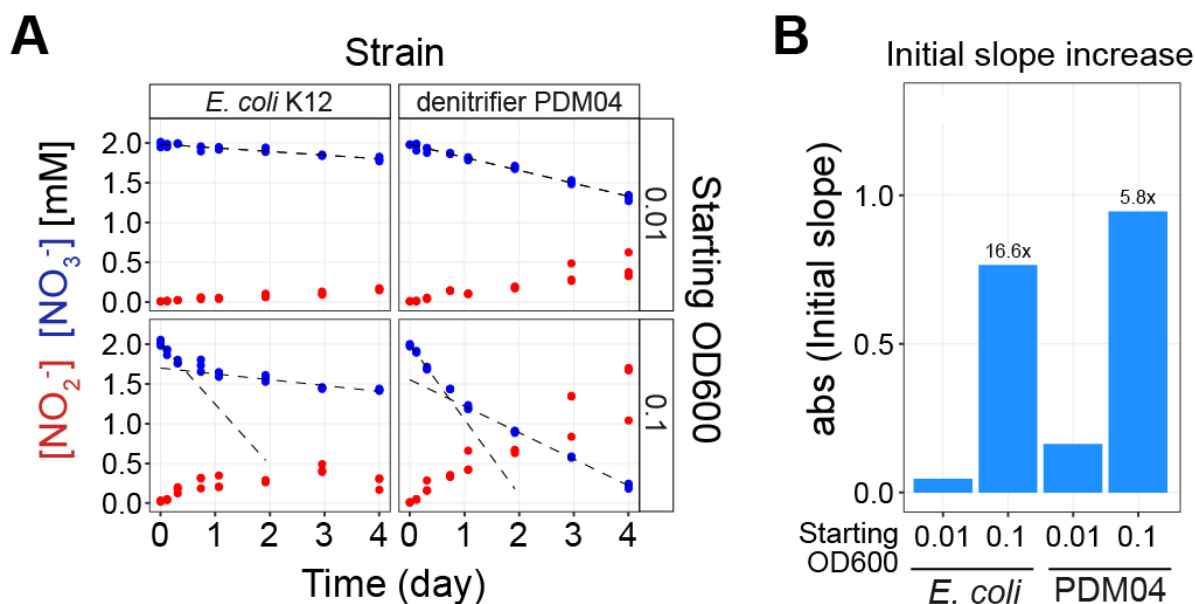
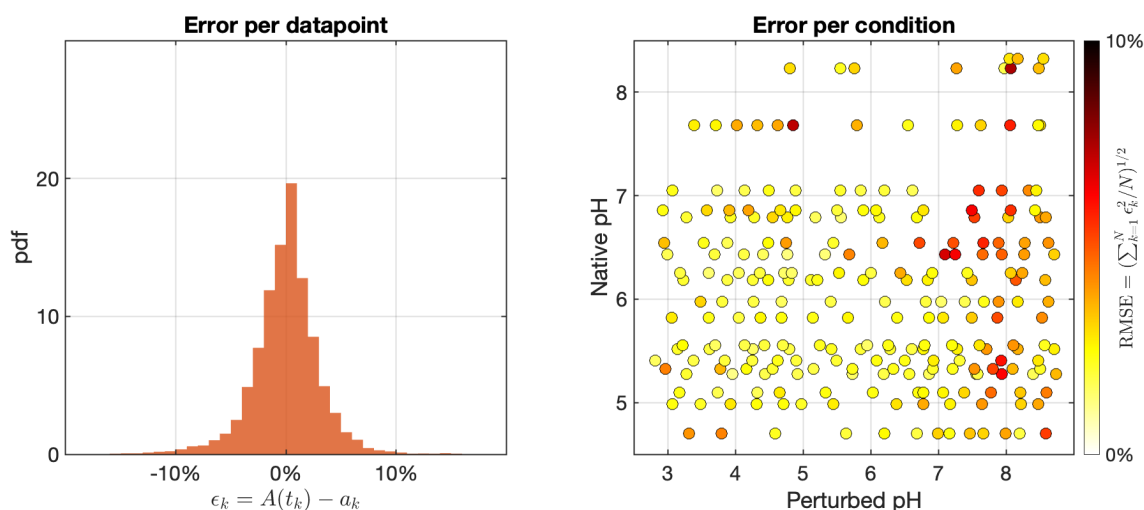


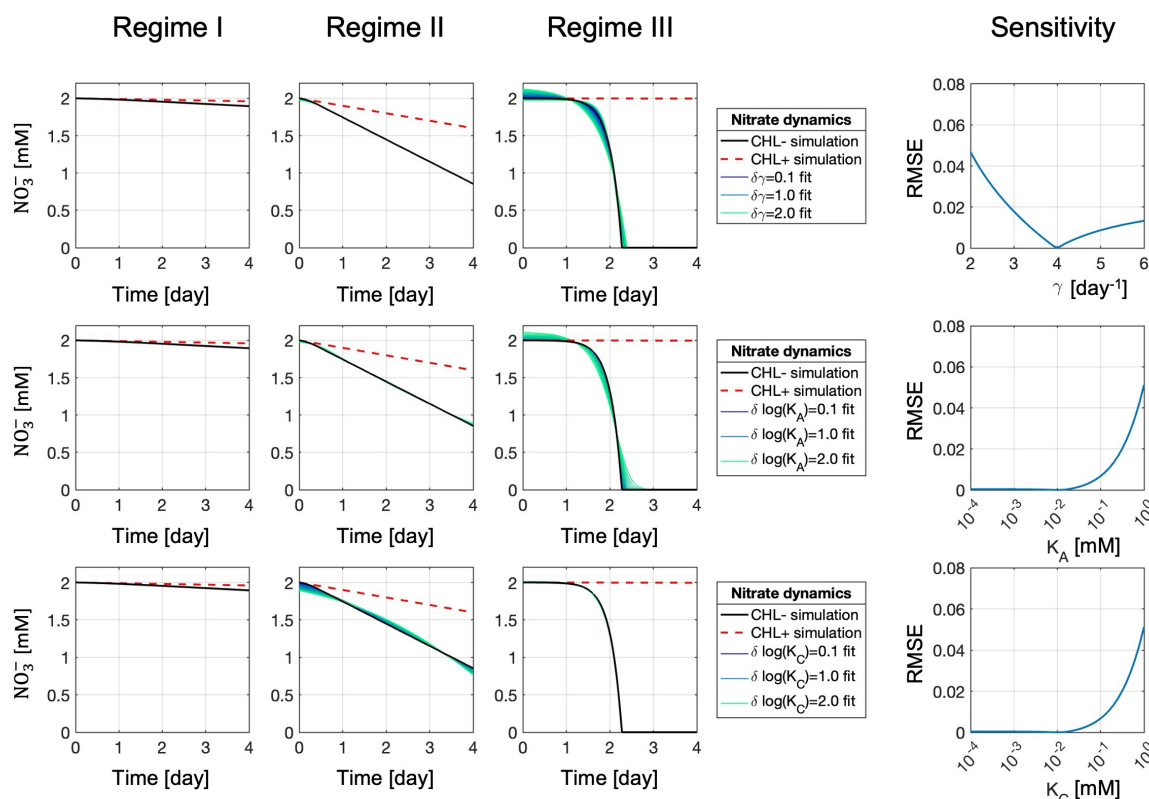
Figure S5: **The results from autoclaving soil suggest the absence of abiotic (chemical) nitrate reduction.** Nitrate (blue data points) and nitrite (red data points) dynamics of a soil sample with (right, Autoclaved) and without (left, None) autoclaving procedure. The autoclaving was performed at 120 °C for 99 minutes and repeated five times at two-day intervals. The soil used in this experiment was collected from LaBagh Woods (latitude 41.977855, longitude -87.742585), Sauganash Prairie, Chicago, IL, USA, on January 18, 2022. Contrary to the soil without the sterilization, nitrate reduction did not occur in the soil with the sterilization process (autoclaving). This rules out the possibility of abiotic (chemical) nitrate reduction occurring in soils.



**Figure S6: Linear metabolite dynamics recapitulated from monoculture experiments in the absence of external carbon sources (A)** Nitrate and nitrite dynamics of monoculture experiments using *E. coli* K12 and the denitrifier *Pseudomonas sp.* PDM04 strains over 4 days with no external carbon provided in the culture media (see SM for detailed experimental methods). The top two panels have a starting OD600 (optical density at 600nm) of 0.01 and the bottom panels have a starting OD600 (optical density at 600nm) of 0.1. The x-axis represents time in days, and the y-axis represents the concentration of nitrate (blue points) or nitrite (red points) reduced (mM), each condition having three biological replicates. The linear dynamics demonstrate that nitrate reduction can occur even in the absence of external carbon, resolving the previous contradiction about the necessity of carbon for this process. The dashed lines represent linear regression of the dynamics: for the top panels, linear regression used all data points, while in the bottom panels, initial slopes were derived from fitting the first three points, and late slopes were calculated using the last four points. **(B)** The fitted initial slope values in nitrate reduction (using initial slopes in A). The different bars indicate the initial slopes from different conditions of starting OD600 values (0.01 and 0.1) and two strains. The plot underscores the effect of starting biomass on initial nitrate reduction rates, with factor increase of initial slopes annotated on top of the bars.



**Figure S7: Quantification of error in model fitting** Error per data point (Left panel): The probability density function (pdf) represents the distribution of errors for individual data points of nitrate measurements at time point  $k$ . Errors are calculated as the difference between the model's predicted nitrate concentration  $A(t_k)$  and the observed nitrate amounts  $a_k$  for either the chloramphenicol-untreated(CHL-) or treated(CHL+) conditions, normalized by dividing by the input nitrate concentration (2mM) to be expressed as a percentage. Error per condition (right panel): Each dot represents the error for a specific experimental condition (triplicates), with the native pH of the sample on the y-axis and the perturbed pH on the x-axis. The error per condition, indicated by the color of each point, is the square root of the mean-squared error (MSD) loss function minimized during parameter optimization of both CHL-/+ conditions of triplicates, normalized by the input nitrate concentration (2mM) to be expressed as a percentage (refer to Methods for the error computation).



**Figure S8: Sensitivity analysis on model parameters  $\gamma$ ,  $K_A$ , and  $\tilde{K}_C$  to justify fixing these parameters**

To justify the fixed parameters in the fitting scheme, we analyzed the sensitivity of  $\gamma$ ,  $K_A$ , and  $\tilde{K}_C$  by simulating dynamic data. To reflect the three typical dynamics (regimes) observed from the measurement, we simulated three nitrate curves by setting up the initial conditions to be  $\tilde{x}(0) = 0.01, 0.1, 0.001 \text{ mM/day}$  and  $\tilde{C}(0) = 0.005, 0.05, 2 \text{ mM}$ , respectively. Other parameters are given by  $A_0 = A_0^C = 2 \text{ mM}$ ,  $K_A = \tilde{K}_C = 0.01 \text{ mM}$ ,  $\gamma = 4 \text{ day}^{-1}$ . Black curves indicate the simulated nitrate dynamics from the chloramphenicol-untreated (CHL-) conditions, and red dashed lines indicate the simulated nitrate dynamics from the chloramphenicol-treated (CHL+) conditions. We then used different fixed values of parameters to fit the three examples. In the first row, we used different fixed  $\gamma$  values - from  $\gamma = 2 \text{ day}^{-1}$  to  $\gamma = 6 \text{ day}^{-1}$  - to fit three simulations. The square root of the mean-squared error (RMSE) is computed by the loss function (mean-squared difference of predicted and observed nitrate concentration for both CHL-/+ conditions) minimized during parameter optimization, normalized by the input nitrate concentration (2mM) to be expressed as a percentage (refer to Methods for loss function). We demonstrate very small mismatches ( $\text{RMSE} < 5\%$ ) from these variations of parameter values, which are almost invisible in Regime I and Regime II fittings (purple lines indicate fitted results from  $\gamma = 4 \pm 0.1 \text{ day}^{-1}$ , blue lines indicate fitted results from  $\gamma = 4 \pm 1 \text{ day}^{-1}$ , green lines indicate fitted results from  $\gamma = 4 \pm 2 \text{ day}^{-1}$ ). In the second and the third row, we used different fixed  $K_A$  and  $\tilde{K}_C$  values - from  $10^{-4} \text{ mM}$  to  $1 \text{ mM}$  - to fit three simulations. When  $K_A < 0.1 \text{ mM}$  or  $\tilde{K}_C < 0.1 \text{ mM}$ , the mismatches were again very small ( $\text{RMSE} < 1\%$ ) and invisible (purple lines indicate fitted results from  $K_{A,C} = 10^{-2 \pm 0.1} \text{ mM}$ , blue lines indicate fitted results from  $K_{A,C} = 10^{-2 \pm 1} \text{ mM}$ , and green lines indicate fitted results from  $K_{A,C} = 10^{-2 \pm 2} \text{ mM}$ ). These results indicate that the fixed values of  $\gamma$ ,  $K_A$  and  $\tilde{K}_C$  are insensitive in large ranges.

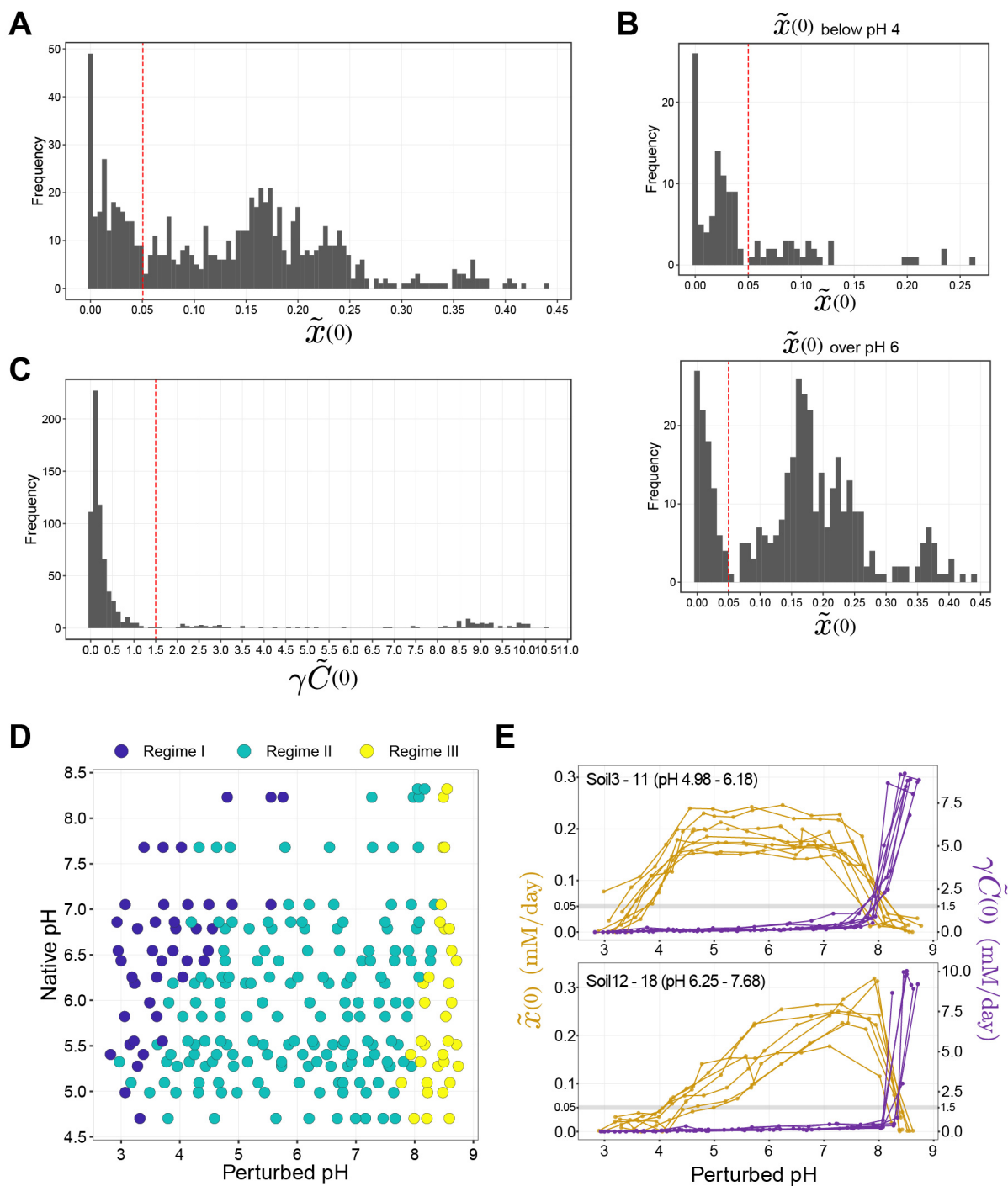
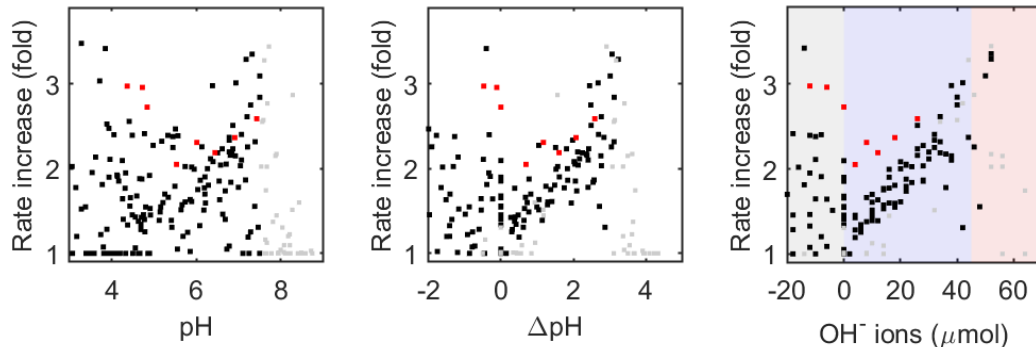


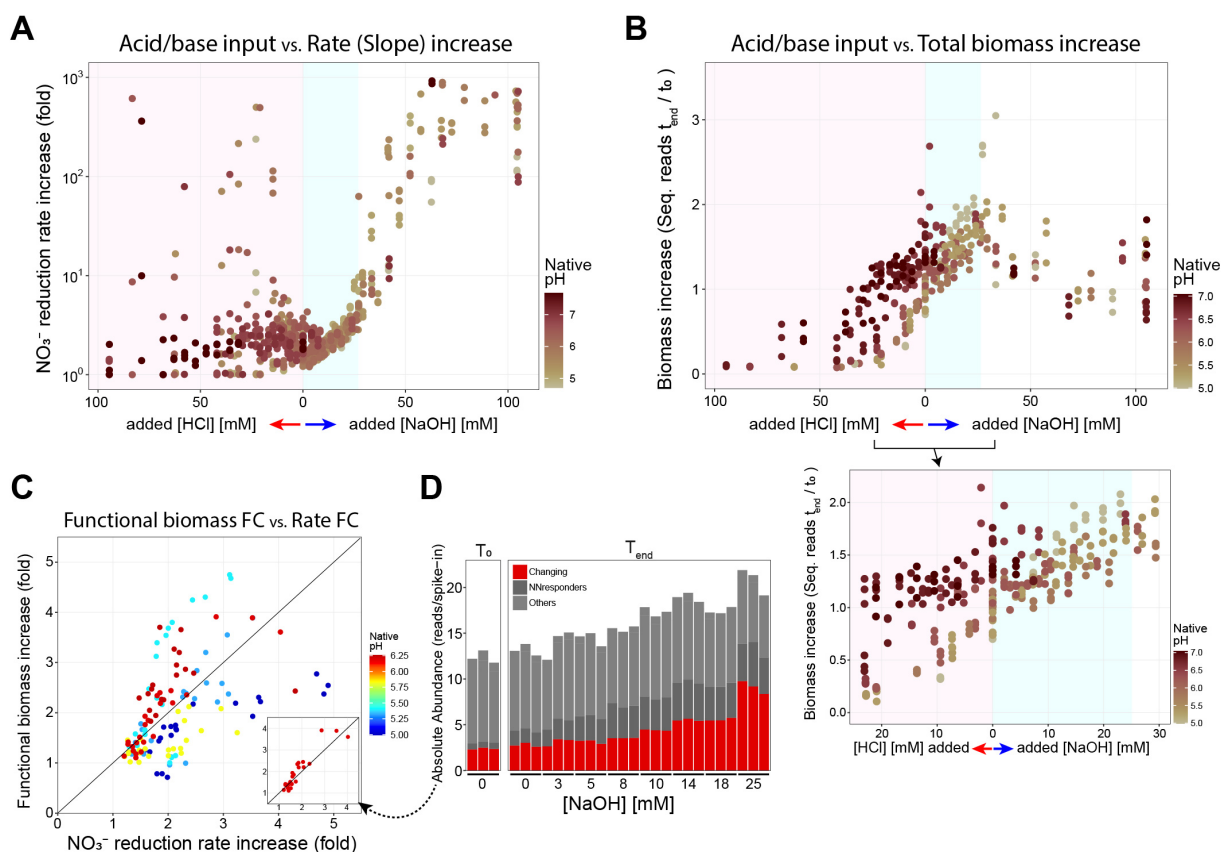
Figure S9: **Determining regime boundary thresholds with distributions of the parameters  $\tilde{x}(0)$  and  $\gamma\tilde{C}(0)$**  To determine the regime boundaries, we examined the distributions of parameters fitted to the functional data for  $\tilde{x}(0)$  and  $\gamma\tilde{C}(0)$ . (Continued)

Figure S9: (Continued from the previous page) **(A)**  $\tilde{x}(0)$  had a bimodal frequency distribution, having two peaks. **(B)** This bi-modality becomes more evident when we separately observe its distribution from the left half (perturbed pH < 4) and right half (perturbed pH > 6) of the parameter space displayed in the perturbed pH vs. native pH grid in Figure 3C. We set the threshold for the  $\tilde{x}(0)$  boundary where these two modes are separated ( $\tilde{x}(0) = 0.05$ ). **(C)**  $\gamma\tilde{C}(0)$  showed an uni-modal frequency distribution. We set the threshold ( $\gamma\tilde{C}(0) = 1.5$ ) at the tail of the distribution, where the  $\gamma\tilde{C}(0)$  threshold also separated the Regime III samples in the top-left quadrant of the  $\tilde{x}(0)$  vs.  $\gamma\tilde{C}(0)$  scatter plot (Fig. 3A). The separation of Regime I and Regime II data points may not be clear cut in the  $\tilde{x}(0)$  vs.  $\gamma\tilde{C}(0)$  scatter plot (Fig. 3A). However, when we plot  $\tilde{x}(0)$  of different soils (dark yellow colored lines) by grouping them into relatively acidic (Soil3–11 (pH 4.98-6.18), **(E)** top panel) and neutral soils (Soil12–18 (pH 6.25-7.68), **(E)** bottom panel), the transition from Regime II (large  $\tilde{x}(0)$ ) to Regime I (small  $\tilde{x}(0)$ ) is evident going towards more acidic pH perturbations, especially in the naturally acidic soils (top panel), because the large  $\tilde{x}(0)$  levels are sustained over a wide pH range in Regime II. **(D)** With these thresholds of two parameters, we can define the three different regimes of adaptive behavior across native pH and perturbed conditions (colored differently by regimes).



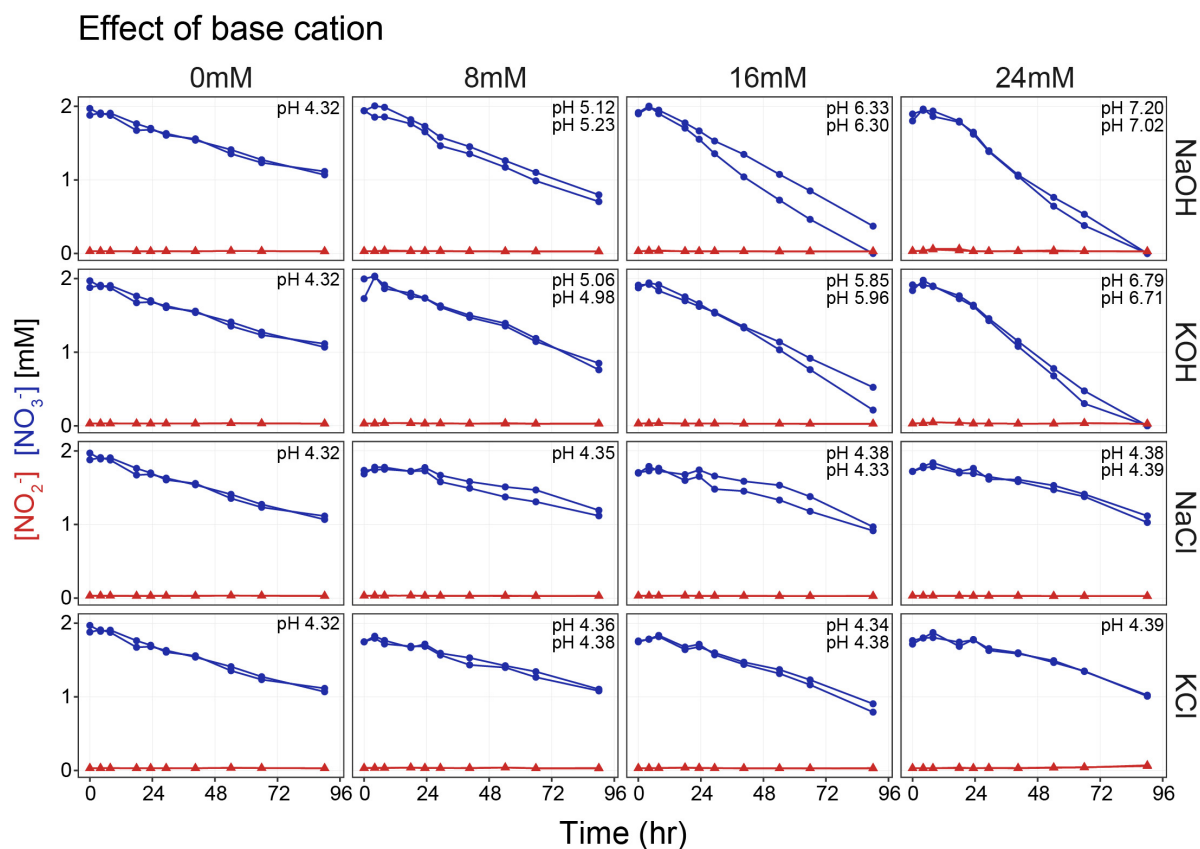


**Figure S10: NaOH input has a more consistent linear relationship with rate increase than pH or delta pH** To provide additional evidence that the  $\text{NO}_3^-$  reduction rate increase (fold) has a linear relationship with the added base, we calculated the rate increase (y-axis) independently from the model. To do so, we performed linear regression on the linear nitrate dynamics of chloramphenicol treated (CHL+) and untreated (CHL-) conditions, determining the slope ratio (CHL-/CHL+). On the leftmost plot, the rate increase is plotted against the perturbed pH (x-axis), against delta pH (= perturbed pH - native pH) in the central plot, and against the added amount of  $\text{OH}^-$  ions (in  $\mu$  moles, negative values indicate the amount of  $\text{H}^+$  ions) input on the rightmost plot. We are using perturbed samples from all soils with varying native pH levels. As we progress from left to right plots, we observe a greater collapse of data into a linear relationship with the rate increase. This confirms that NaOH is the most reliable descriptor for consistently explaining the growth due to nutrient release across soils with various native pH levels. Data points from Soil12 are colored in red owing to its slope being different from the collapsed slope of other soils (black points). Treatments with pH greater than 7.5 were colored gray, as they predominantly belong to the Resurgent growth regime (Regime III), while linearity is expected to only hold in the Nutrient-limiting regime (Regime II). The blue background in the rightmost plot is a guide for the eye, indicating the range of perturbations that typically remain within the Nutrient-limiting regime (Regime II).

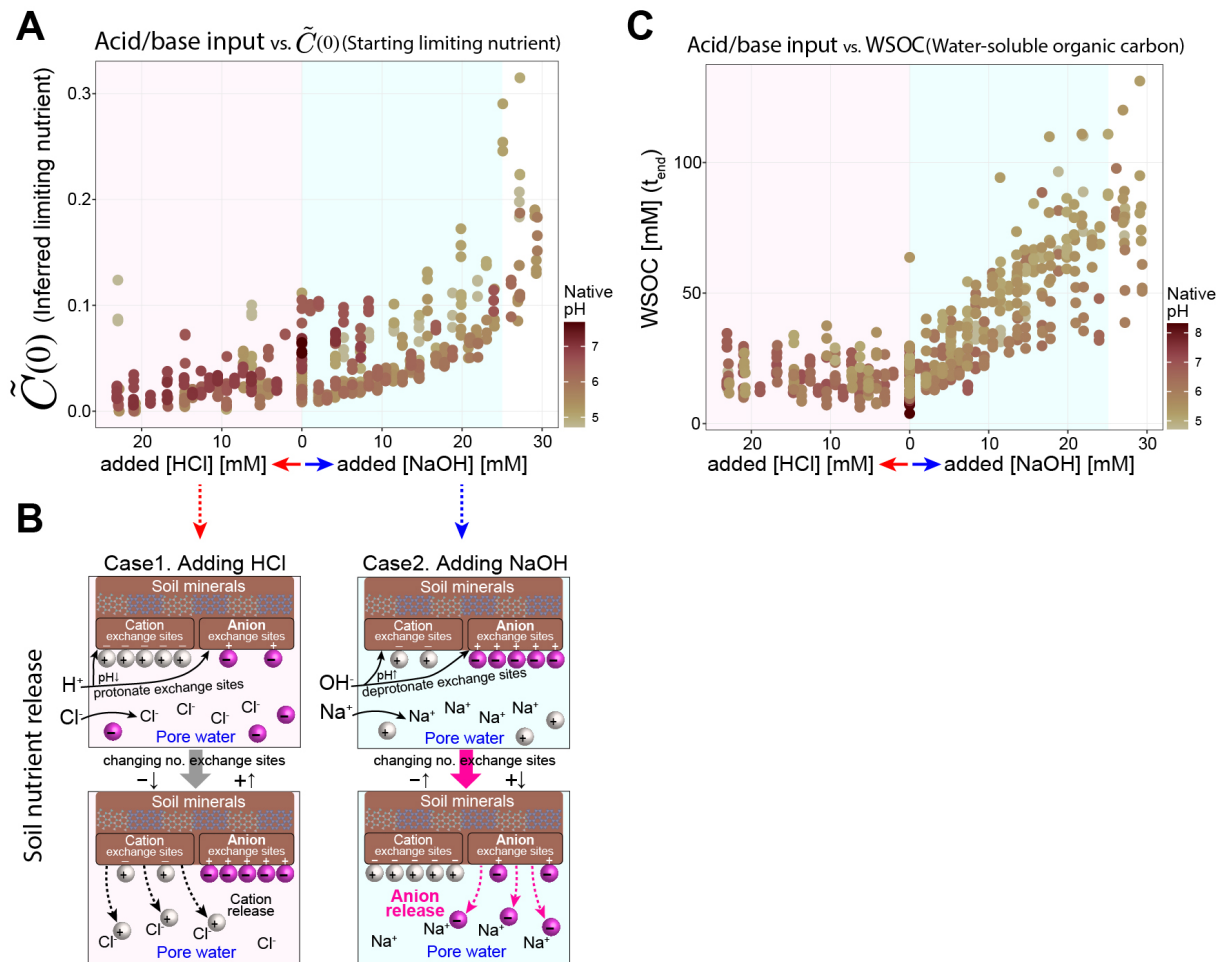


**Figure S11: Confirming the linear dependence between functional biomass and acid/base added** A more detailed analysis, accounting for individual Amplicon sequence variant) ASVs (C, D) that responded to the amendment of nitrate, further confirmed the linear dependence between biomass and acid/base added. (A) Showing the full range of acid/base input (x-axis) against the NO<sub>3</sub><sup>-</sup> reduction rate increase (fold) (y-axis) in chloramphenicol-untreated (CHL-) conditions compared to treated conditions (CHL+) for all soils from different native pH (color gradient of data points). The rate fold increase is computed from the fitted model parameters ( $1 + \gamma \tilde{C}_0 / \tilde{x}_0$ ). This linear relationship is observed within the range of NaOH addition from 0mM to 25mM, which belongs to the Nutrient-limiting regime (Regime II) (light blue background in A). This was not the case for acidic perturbations (> 0mM HCl addition) and basic perturbations beyond 25mM NaOH addition. Therefore, the fitted model parameter suggests that the addition of NaOH causes the release of limiting nutrients in the soil, increasing biomass growth. (B) Showing the full range of acid/base input against biomass growth measured by the sequencing data. Biomass increase (fold) was computed with the ratio of the total absolute abundance of initial and end time points samples ( $T_{end}/T_0$ ). We plotted an inset to highlight a zoomed-in range (< 25mM HCl, < 25mM NaOH). In this range, the amount of biomass growth evidently increases with the addition of NaOH (light blue background) and decreases with the addition of HCl (pink background) for all soils from different native pH levels (color gradient of data points). Although this linear relationship corroborates our proposed nutrient release mechanism, to be more precise, we need to prove further that the factor increase of the “functional” biomass equals the factor increase of nitrate reduction linear rate from the flux dynamics data. This is because not all biomass performs NO<sub>3</sub><sup>-</sup> reduction. (Continued)

Figure S11: (Continued from the previous page) To detect the fractional biomass that performs denitrification, we used a differential abundance analysis to statistically determine which ASVs were significantly enriched in each pH perturbed condition compared to the CHL+ counterpart serving as a baseline of no growth (see Methods). We filtered out the ASVs that could be false-positive nitrate reducers by removing ASVs that were statistically enriched in no-nitrate conditions (dark grey NNresponders bar in **(D)**). Then, we summed up the absolute abundance of these ASVs that we inferred as true nitrate reducer biomass to obtain the functional biomass for each condition (red bar in **(D)**). **(C)** By comparing the factor increase of these functional biomass values (endpoint/initial functional biomass), we showed that indeed the functional biomass increase and denitrification rate increase are aligned in different soils (color spectrum in soils with different native pH). Some soils had these two values lie very close to the 1:1 diagonal line (Soil11, inset of **(C)**), which validates our inference procedure.

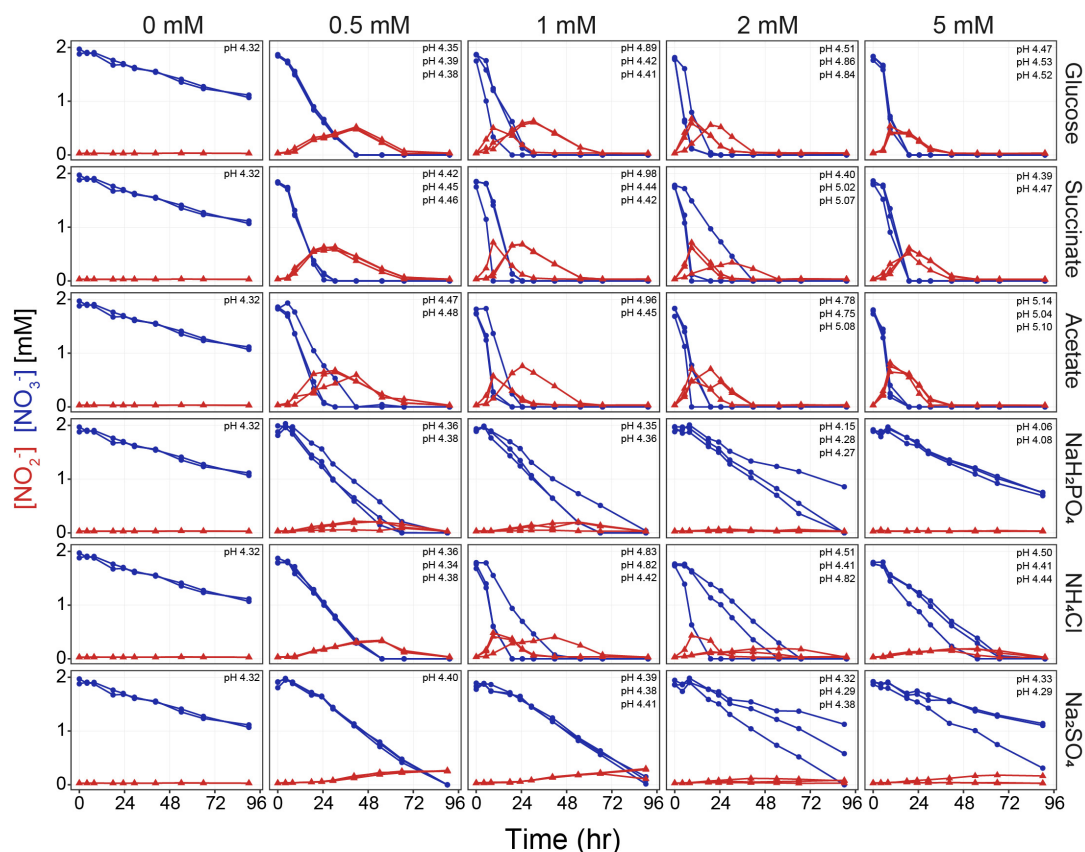


**Figure S12: Testing the effect of different bases and salts on nutrient release** To see the effects of different bases (NaOH and KOH) on nitrate reduction dynamics, we added different concentrations of NaOH and KOH (final concentration of 0, 8, 16, 24mM in the slurry), following the same protocol previously described (without chloramphenicol), to measure the nitrate and nitrite dynamics using Soil6 (Table S3). In addition, to test the effects of  $\text{Na}^+$ ,  $\text{K}^+$ , and  $\text{Cl}^-$  separately, we added different concentrations of salts (NaCl, KCl) (without chloramphenicol and without adding any acid/base) and measured the metabolite dynamics. Blue points denote nitrate measurements and red points denote nitrite measurements. The lines connect the data points of two biological replicates. Each panel displays the stabilized endpoint pH (1M KCl method). Identical pH values across biological replicates are noted once.

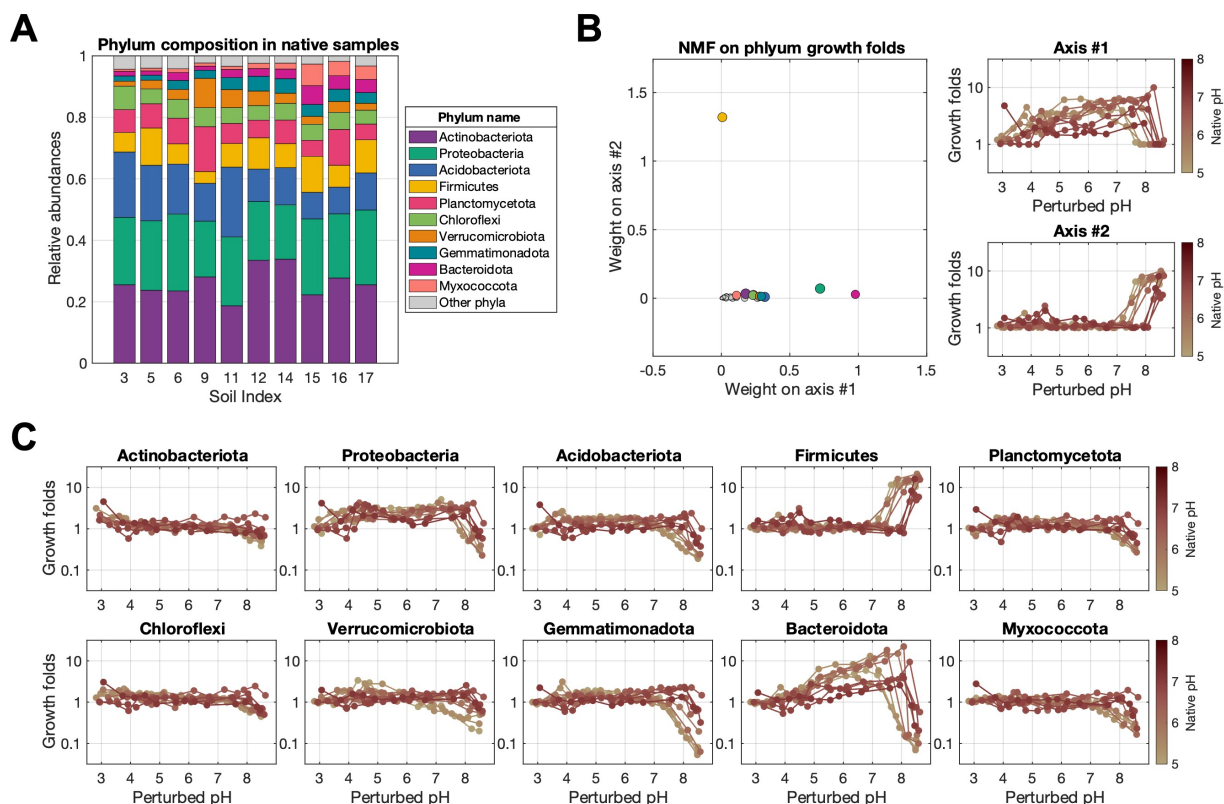


**Figure S13: Water-soluble organic carbon (WSOC) measurement aligns with the nutrient release hypothesis in the Nutrient-limiting regime (Regime II)** (B) Cartoon illustrating the soil nutrient release hypothesis; NaOH results in the release of anion nutrients from soil clay particles (brown region), while the addition of HCl releases cation nutrients and adsorbs anion nutrients. Microbes cannot access the nutrients adsorbed in the soil particles but can access the nutrients dissolved in soil pore water. Added  $\text{OH}^-$  ions deprotonate both cation and anion exchange sites, hence decreasing the number of anion exchange sites in the soil particles and increasing the number of cation exchange sites. This releases anion nutrients from the clay particles to the pore water, while cations in the pore water are adsorbed to the clay particles. In concert, added  $\text{Na}^+$  ions stabilize the released anions in the pore water facilitating the release. On the other hand, during HCl addition, Added  $\text{H}^+$  protonates both cation and anion exchange sites, hence increasing the number of anion exchange sites in the soil particles and decreasing the number of cation exchange sites. This releases cations from the clay particles to the pore water, while anion nutrients in the pore water are adsorbed to the clay particles no longer available to the microbes. In concert, added  $\text{Cl}^-$  ions stabilize the released cation in the pore water. (Continued)

Figure S13: (Continued from the previous page) **(A)** With this proposed mechanism of nutrient release by NaOH and HCl, we can further specify the type of growth-limiting nutrient by observing the change of the fitted model parameter of  $\tilde{C}(0)$  (starting limiting nutrient). In natively acidic soils, increasing NaOH concentrations linearly increased the  $\tilde{C}(0)$  (light blue region), which indicated that the limiting nutrient is negatively charged (anion nutrient). In natively neutral soils, increasing HCl concentration linearly decreased the  $\tilde{C}(0)$  (light pink region). This is congruent with our statement that the growth-limiting nutrients are anions, because when HCl is added, anions are sequestered to the clay particles becoming unavailable to the microbes (purple spheres in **B**). **(C)** Coincidentally, adding NaOH linearly increased the water-soluble organic carbon (WSOC) concentrations present in the slurry at the endpoint, while adding HCl did not. This suggests two aspects related to our nutrient release hypothesis. Firstly, it appears that most water-soluble organic carbon (WSOC) may be negatively charged (anion). Secondly, the growth-limiting nutrient might be either the WSOC itself or another nutrient that is concomitantly released with organic carbon in the form of organic matter, including all carbon (C), nitrogen (N), sulfur (S), and phosphorus (P). If the limiting nutrient were WSOC, only a fraction of WSOC would be used as nutrient, because while  $\tilde{C}(0)$  and WSOC are well correlated, the concentration of released WSOC is disproportionately higher ( $\approx 20\text{-}75$  C mM) than the amount of limiting nutrient needed to deplete all  $2\text{mM NO}_3^-$  in the system, as shown in Fig. 4D).



**Figure S14: Nitrate and nitrite dynamics of soils amended with different nutrients** To experimentally determine what nutrient was limiting growth in the Nutrient-limiting regime (Regime II), we conducted nutrient amendment experiments respectively with varying concentrations of glucose, succinate, sodium acetate, ammonium chloride ( $\text{NH}_4\text{Cl}$ ), monosodium phosphate ( $\text{NaH}_2\text{PO}_4$ ), and sodium sulfate ( $\text{Na}_2\text{SO}_4$ ). Nitrate dynamics (blue) and nitrite dynamics were measured following the same protocol with  $2\text{mM NO}_3^-$  (see Methods) using Soil6 (Table S3) without chloramphenicol and not adding any acid/base. Columns in the plot are different concentrations of C mM, N mM, S mM, or P mM in final concentrations in the slurry varying from 0 to 5 mM, each with biological replicates. 0mM amendment conditions are the same for all nutrients. Rows in the plots are different nutrients: C source (glucose, succinate, acetate), P source (phosphate), N source (ammonium), and S source (Sulfate). Each panel displays the stabilized endpoint pH (1M KCl method), and identical pH values across biological replicates were noted once. Among them, succinate ( $\text{pK}_a = 4.21$  and  $5.64$ ,  $25^\circ\text{C}$ ), acetate ( $\text{pK}_a = 4.76$ ,  $25^\circ\text{C}$ ), and phosphate ( $\text{pK}_a = 2.2$ ,  $7.2$ , and  $12.4$ ,  $25^\circ\text{C}$ ) were strong candidates for the limiting nutrient according to our soil nutrient release hypothesis, due to their anionic nature in mid-range pH (5-7). Because we have previously tested the effect of  $\text{Na}^+$  and  $\text{Cl}^-$  to be negligible in nitrate dynamics, the effect of these amendments can be attributed solely to C/N/S/P nutrients other than  $\text{Na}^+$  and  $\text{Cl}^-$ . We observed a transition from linear dynamics to exponential depletion of nitrate, when we amended the soil with a carbon source. Ammonium also made the nitrate consumption dynamics exponential in 1 mM amendment, but not in other amendment concentrations.



**Figure S15: NMF (Non-negative matrix factorization) reveals low-dimensional shifts in growth at the phylum level** (A) Initial community composition ( $T_0$ ) of native soils in the phylum level. The x-axis indicates soils with different native pH levels (Soil 3, 5, 6, 9, 11, 12, 14, 15, 16, 17, see Table S3 for their properties). The y-axis represents the relative abundance (summed to 1) of the top 10 phyla out of 40, with the cumulative abundance of the remaining phyla depicted in gray as 'Other phyla'. (C) By using the absolute abundance of each taxon in chloramphenicol-treated (CHL+) conditions as a baseline value for growth in each perturbed pH condition, we computed the fold increase of each taxon's absolute abundance in chloramphenicol-untreated (CHL-) conditions, which we call growth fold ( $Abs_{CHL-}/Abs_{CHL+}$ ). Ten different phyla showed idiosyncratic patterns of growth response along the varying perturbed pH. Soils with different native pH, indicated by the line color, showed relatively conserved growth trends in each phylum. (B) To systematically identify the underlying lower-dimensional growth response to pH, we used non-negative matrix factorization (NMF) on the growth fold values to decompose the growth response of all phyla into two modes (Axis #1 and Axis #2 in B, see Methods for details). Intriguingly, these two response patterns across pH matched the trend of functional parameters fitted with our consumer-resource model respectively for  $\tilde{C}(0)$  and  $\tilde{x}(0)$ . The growth folds of each phylum are the linear combination of two modes whose weights are plotted on the left panel of (B) (points are colored by phylum as in (A)). Firmicutes phylum is mainly composed of mode #2, while other phyla are mainly composed of mode #1. Proteobacteria and Bacteroidota have higher weight #1 than other phyla. Therefore, this enabled us to focus our analysis on these phyla to explain the transition from the Nutrient-limiting regime (Regime II) to the Resurgent growth regime (Regime III).



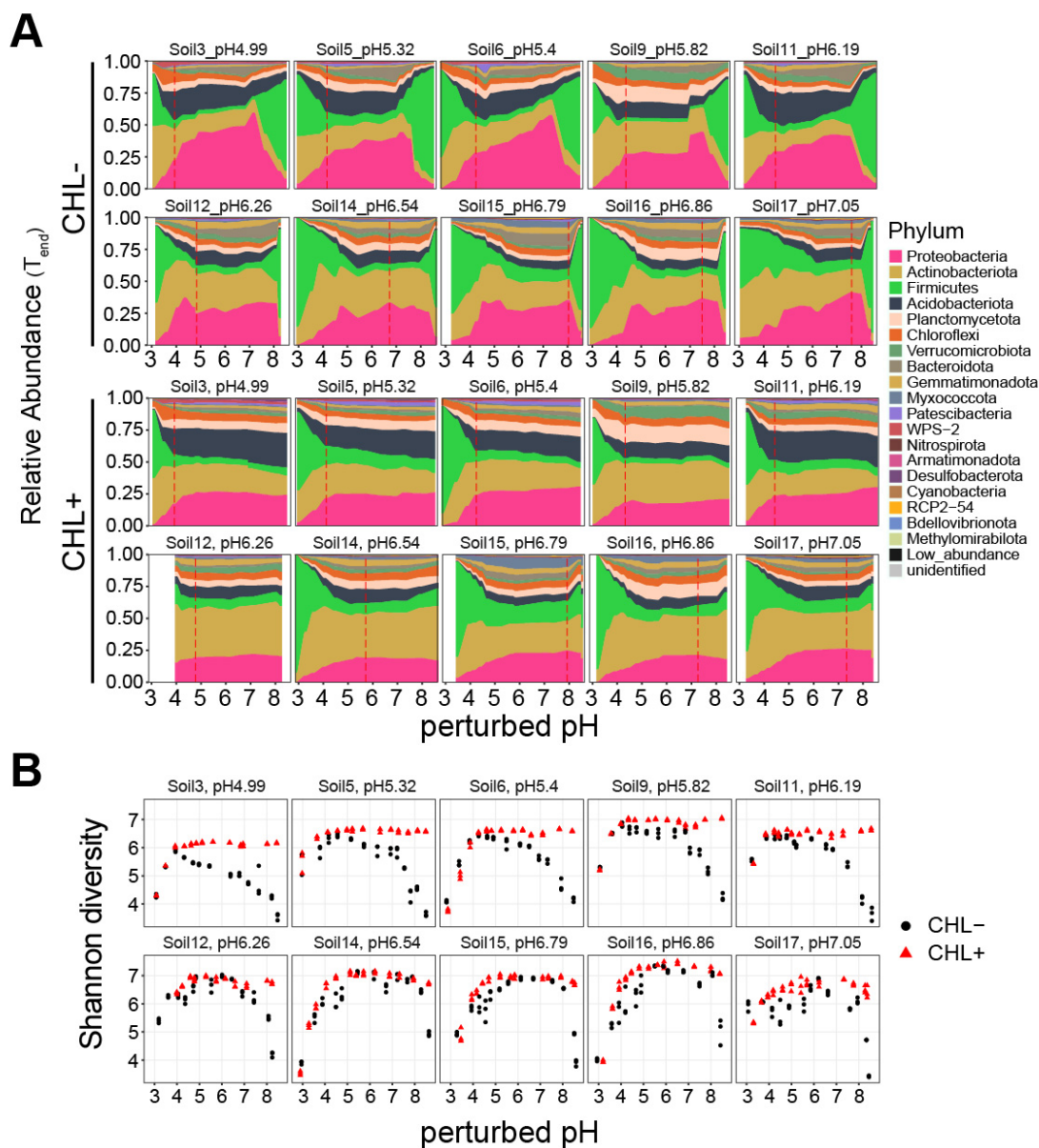
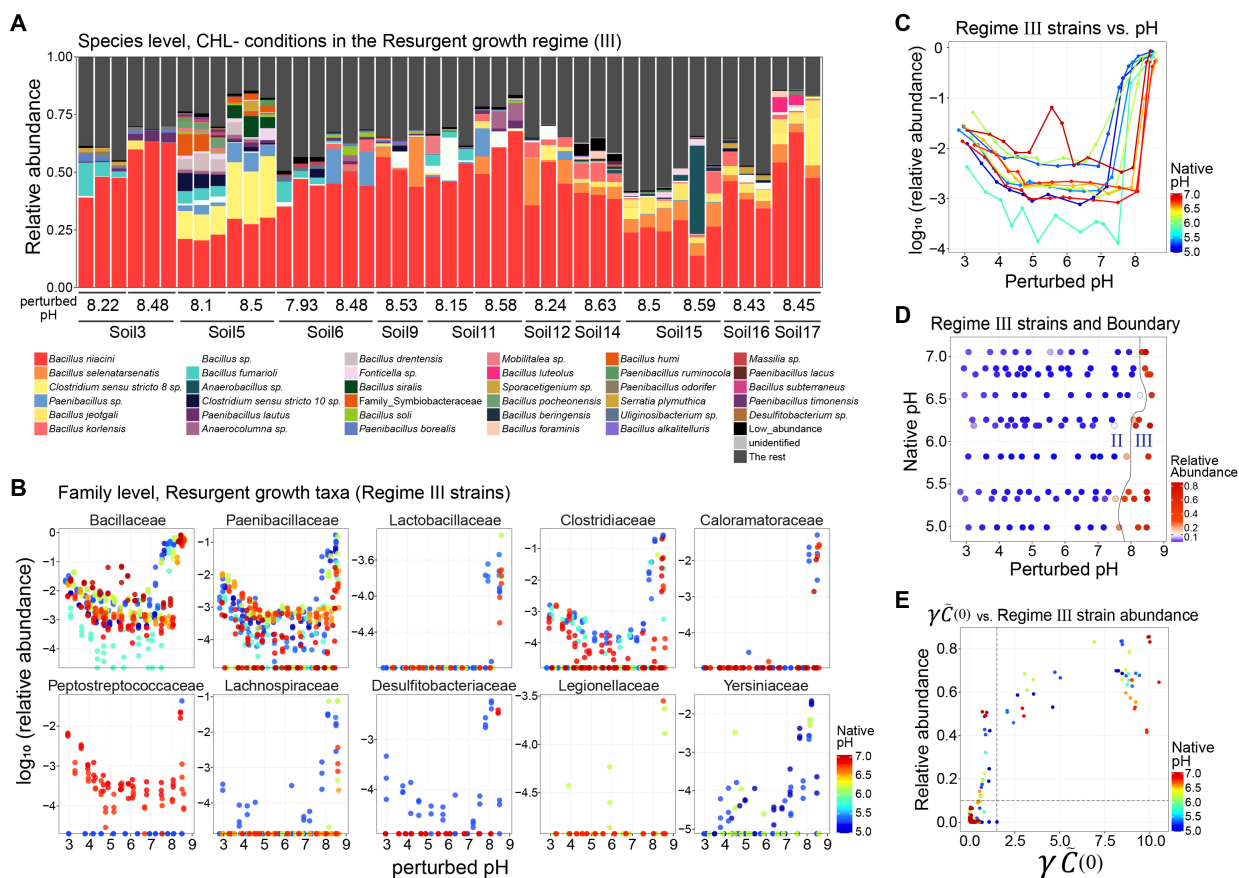


Figure S16: **Relative abundance and diversity of different soils across perturbed pH levels (A)** End-point relative abundance in the phylum level is plotted across the perturbed pH for ten different soils. CHL- indicates endpoint samples without chloramphenicol treatment. CHL+ indicates endpoint samples with chloramphenicol treatment. The alluvial plots were constructed by connecting the relative abundance values of 13 different pH perturbed levels. Red vertical dashed lines indicate the stabilized endpoint pH (1M KCl method) of the unperturbed samples. **(B)** Shannon diversity of the endpoint community is plotted across the perturbed pH for ten different soils in chloramphenicol-untreated(CHL-) and chloramphenicol-treated(CHL+) conditions.



**Figure S17: Taxonomy of the identified Resurgent growth strains (Regime III strains) and their abundance agreeing with the functional Regime III** To identify the specific taxa accountable for the emergence of Regime III at a finer taxonomic level, we conducted a differential abundance analysis that statistically determined which Amplicon sequence variants (ASVs) were significantly more abundant in the Regime III CHL- samples, compared to CHL+ samples under same perturbed pH conditions (see Methods). **(A)** The relative abundance of the ASVs in all Regime III samples is highlighted and colored by their assigned species level. The ASVs not significantly enriched in Regime III samples are colored dark gray and labeled as "The rest". At the genus level, *Bacillus*, *Clostridium*, *Paenibacillus*, and others were identified as the primary contributors to the Resurgent growth regime (Regime III) as plotted. **(B)** The analysis revealed that 10 families belonging to Firmicutes (Bacillaceae, Paenibacillaceae, Clostridiaceae, Caloramatoraceae, Peptostreptococcaceae, etc.) and 2 families belonging to Proteobacteria phylum (Legionellaceae and Yersiniaceae) significantly enriched in the Resurgent growth regime (Regime III). Their relative abundance (log<sub>10</sub> scale) increases at basic perturbed pH levels, patterns differing in soils with different native pH levels. Their relative abundance also slightly increases in Regime I, due to their high tolerance to pH perturbations. **(C-D)** Then, we aggregated the relative abundance of these differential ASVs (i.e., Regime III strains) to assess their contribution to the emergence of Regime III. Notably, their abundances (log<sub>10</sub> scale) rise between pH 7-8, which aligns with or slightly precedes the transition between Regime II and III. **(E)** This increase in relative abundance corresponded with the rise of the nutrient growth parameter  $\gamma \tilde{C}_0$  from zero.

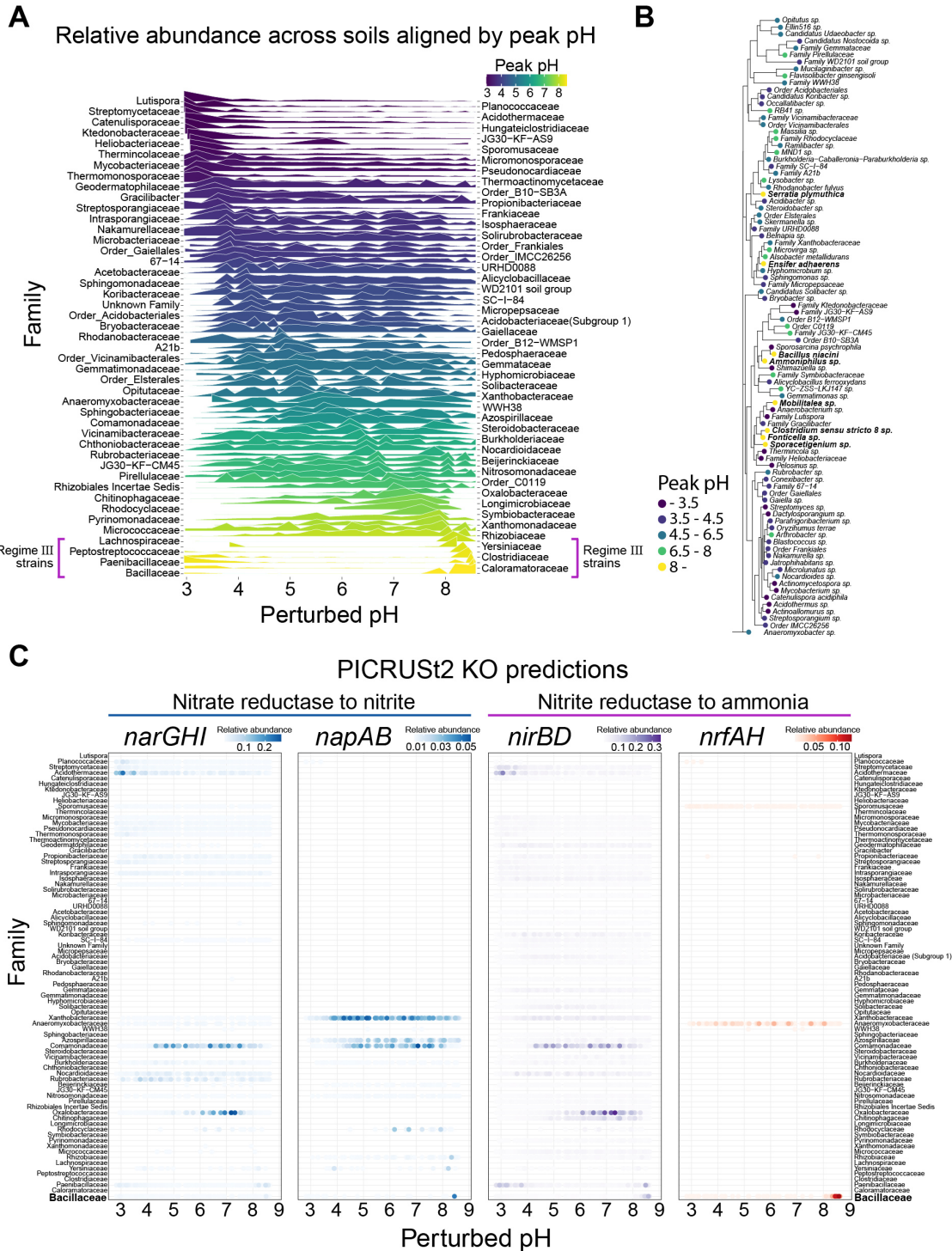
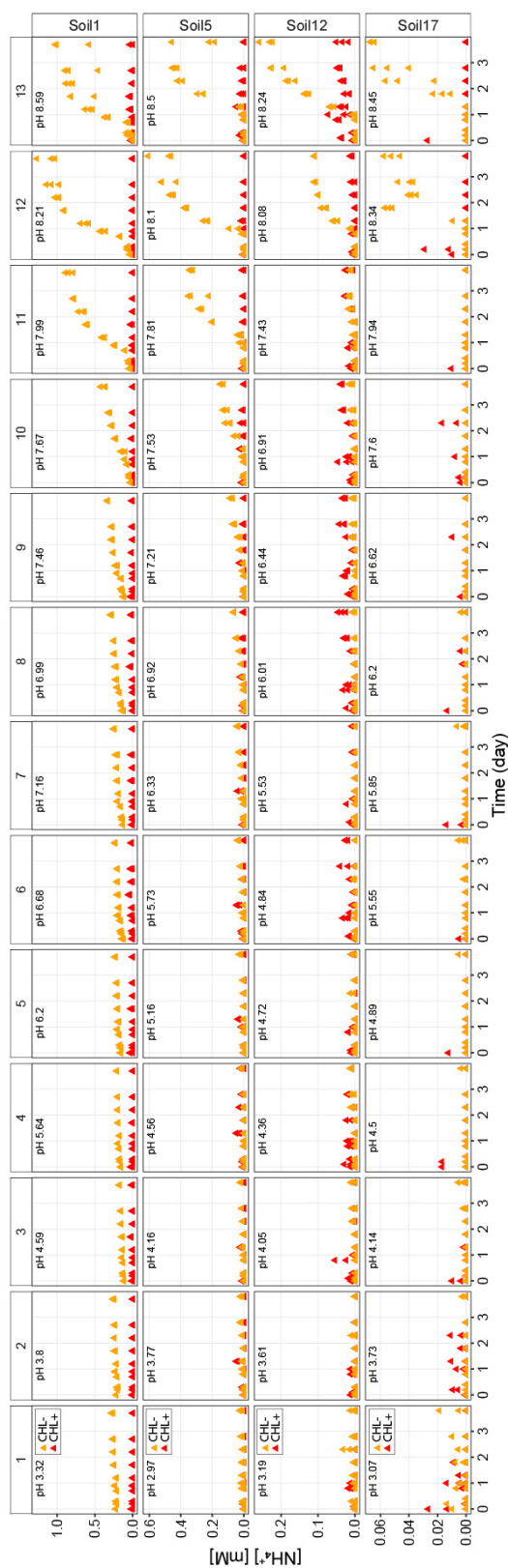
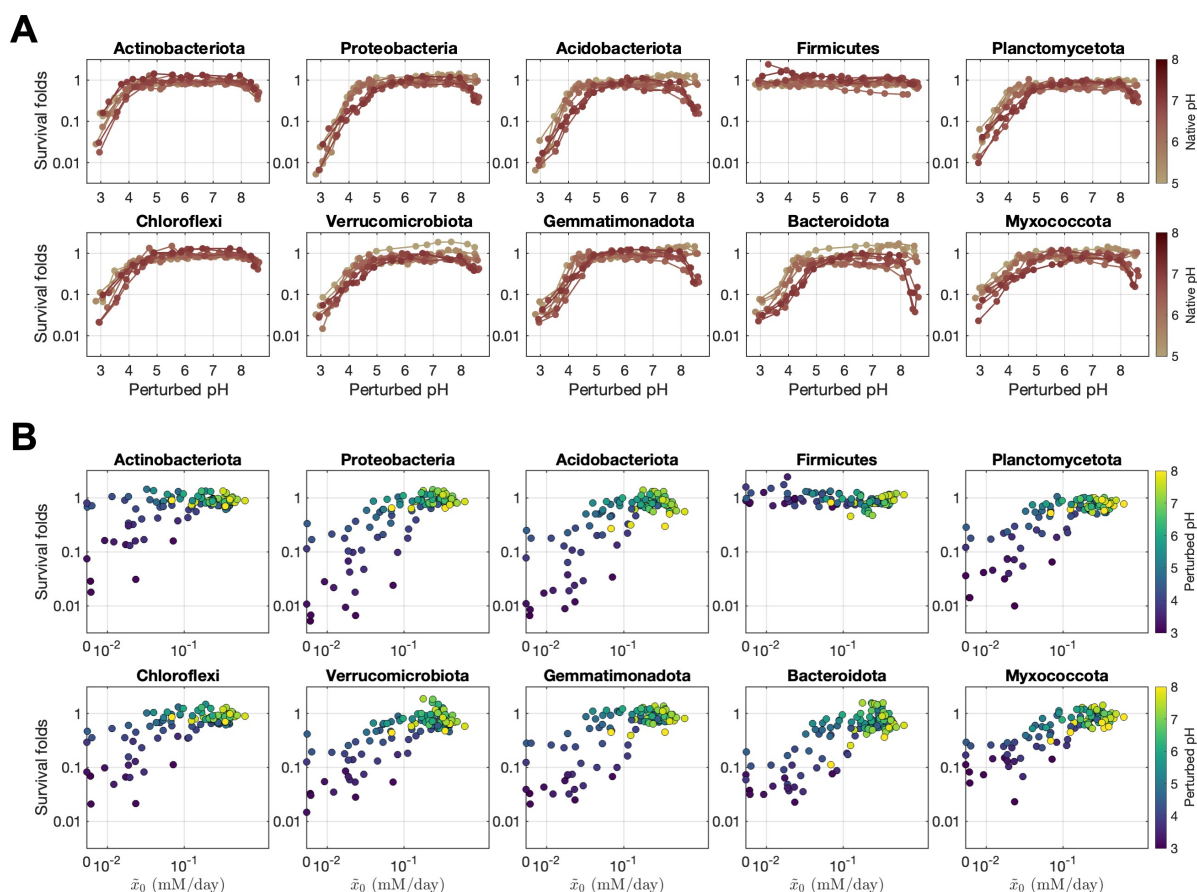


Figure S18:

**Figure S18: Traits of the Resurgent growth strains (Regime III strains) are analyzed through pH niche, phylogenetic distance, and gene predictions from PICRUSt2** (A) To elucidate the pH niche of all taxa, we analyzed the relative abundance of amplicon sequence variants (ASVs) identified as being enriched in different pH levels (see Differential abundance analysis in Methods). We aggregated the relative abundance of ASVs in the same family for each sample, and then computed the median relative abundance of samples across different soils belonging to each pH bin (see Methods). The families were ordered by their peak pH (acidic peak pH on top with dark blue color and basic peak pH on the bottom with yellow color), which was the pH corresponding to the pH bin with the highest median relative abundance for each family. Indeed, the families belonging to the Resurgent growth strains had a peak pH over 8: Bacillaceae, Clostridiaceae, Paenibacillaceae, Caloramatoraceae, Peptostreptococcaceae, Lachnospiraceae (Firmicutes phylum), and Yersiniaceae (Proteobacteria phylum). (B) To see whether there was phylogenetic convergence among strains with similar pH niches, we used the 16S rRNA sequences of the ASVs to construct a phylogenetic tree. We selected one representative ASV with the largest relative abundance from each family to represent the family and used its 16S rRNA V34 region sequence to construct the phylogenetic tree. (see Method). Each node is labeled by the genus or species name and colored by its peak pH. The Resurgent growth strains (yellow-colored) did not cluster phylogenetically and were dispersed throughout the phylogenetic tree. (C) To infer genotypes of the Resurgent growth strains (Regime III strains), we used PICRUSt2 to predict the KEGG ortholog (KO) gene abundance from the 16S rRNA sequence of each ASV (see Methods). We focused on KOs/genes related to denitrification and Dissimilatory Nitrate Reduction to Ammonium (DNRA): nitrate reductase in denitrification (*narG*, *narH*, *narI*, *napA*, *napB*) and nitrite reductase to ammonium (*nirB*, *nirD*, *nrfA*, *nrfH*). To track which KOs were enriched at which pH in the 89 families used in the peak pH analysis in A, we summed the relative abundance (reads / total reads of each perturbed pH level in CHL-samples) of the ASVs belonging to each family that possessed at least 1 predicted gene respectively for *narGHI*, *napAB*, *nirBD*, and *nrfAH*. Then, we plotted their relative abundance values across pH for all soils, indicated by the intensity of the point's colors. For the Resurgent growth strain, Bacilli family exhibited a notable enrichment in *nrfAH* genes (indicated by red points), which are DNRA-related genes producing ammonium from nitrite.

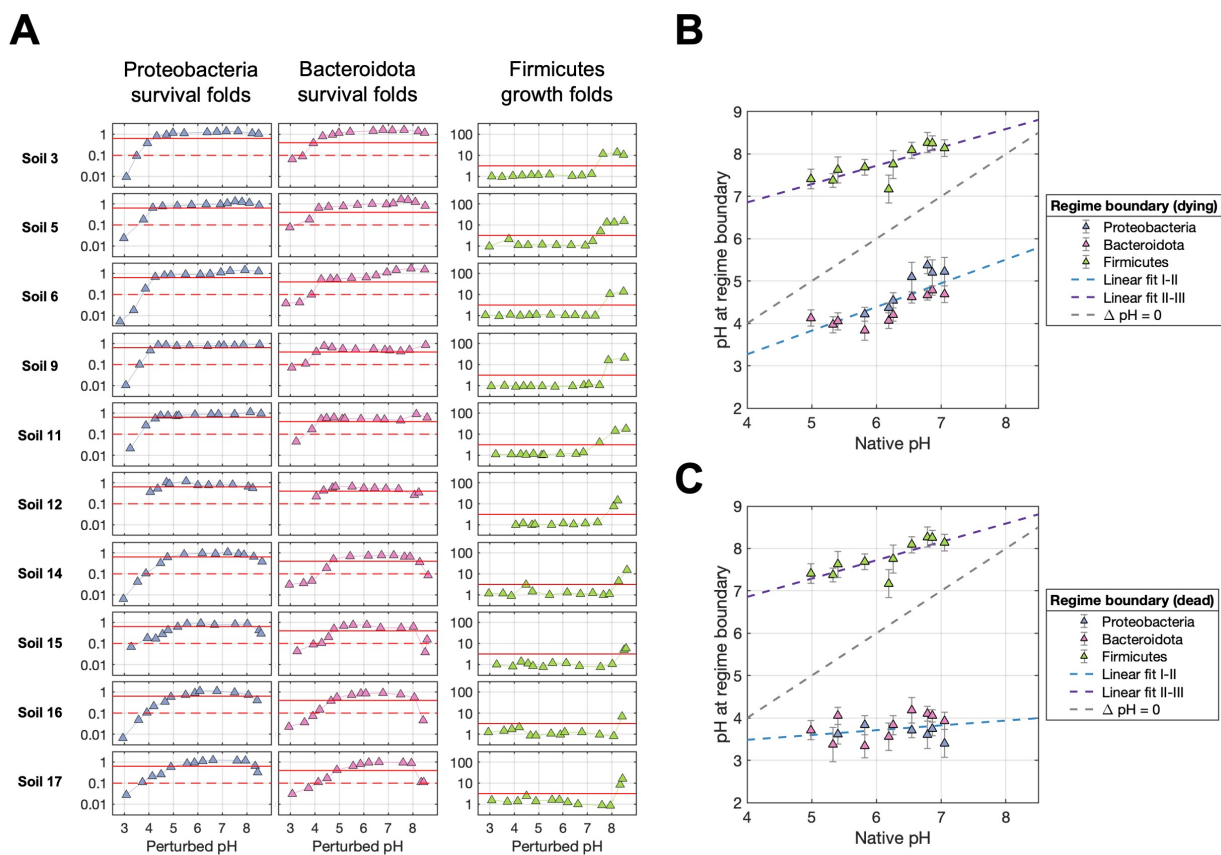


**Figure S19: Ammonium dynamics in chloramphenicol-untreated (CHL-) and treated (CHL+) samples in 4 soils with different native pH levels** For Soil 1, 5, 12, and 17 (for details of soils, refer to Table S3), the ammonium dynamics were measured colorimetrically using the Salicylate-hypochlorite assay [75]. Ammonium dynamics show that varying levels of  $\text{NH}_4^+$  (from 3% (Soil17) to 50% (Soil1) of the provided  $2\text{mM NO}_3^-$  converted to  $\text{NH}_4^+$ ) accumulated in the Resurgent growth regime (Regime III). This indicates the activation of dissimilatory nitrate to ammonia (DNRA) pathway by a subset of strains responsible for the Resurgent growth regime. Chloramphenicol treatments in the samples (CHL+) led to consistent detection of  $0.5\text{mM NH}_4^+$  due to its N-H moiety. The NaOH concentration in perturbed samples also impacted  $\text{NH}_4^+$  measurements, because the Salicylate-hypochlorite assay includes a step that  $\text{OCl}^-$  reacts with the N-H moiety resulting in N-Cl and  $\text{OH}^-$ , higher NaOH concentrations resulting in slightly lower detection of chloramphenicol in the CHL+ samples ( $0.45\text{mM NH}_4^+$  in  $100\text{mM NaOH}$  perturbations). Taking advantage of these control measurements, we used the constant  $\text{NH}_4^+$  levels in the controls without  $2\text{mM NO}_3^-$  (No-Nitrate controls) in the CHL+ conditions for each soil to offset the NaOH effect in the CHL- samples (by computing the conversion factor ratio of  $\text{NH}_4^+$  levels of No-Nitrate controls in CHL+ conditions to the initial  $\text{NH}_4^+$  levels of each condition with different NaOH additions in CHL- samples) and to subtract  $\text{NH}_4^+$  levels caused by chloramphenicol in CHL+ samples. pH indicated inside each panel is the endpoint stabilized pH (see Methods).



**Figure S20: Death of phyla during acidic pH perturbations explains Acidic death regime (Regime I)**

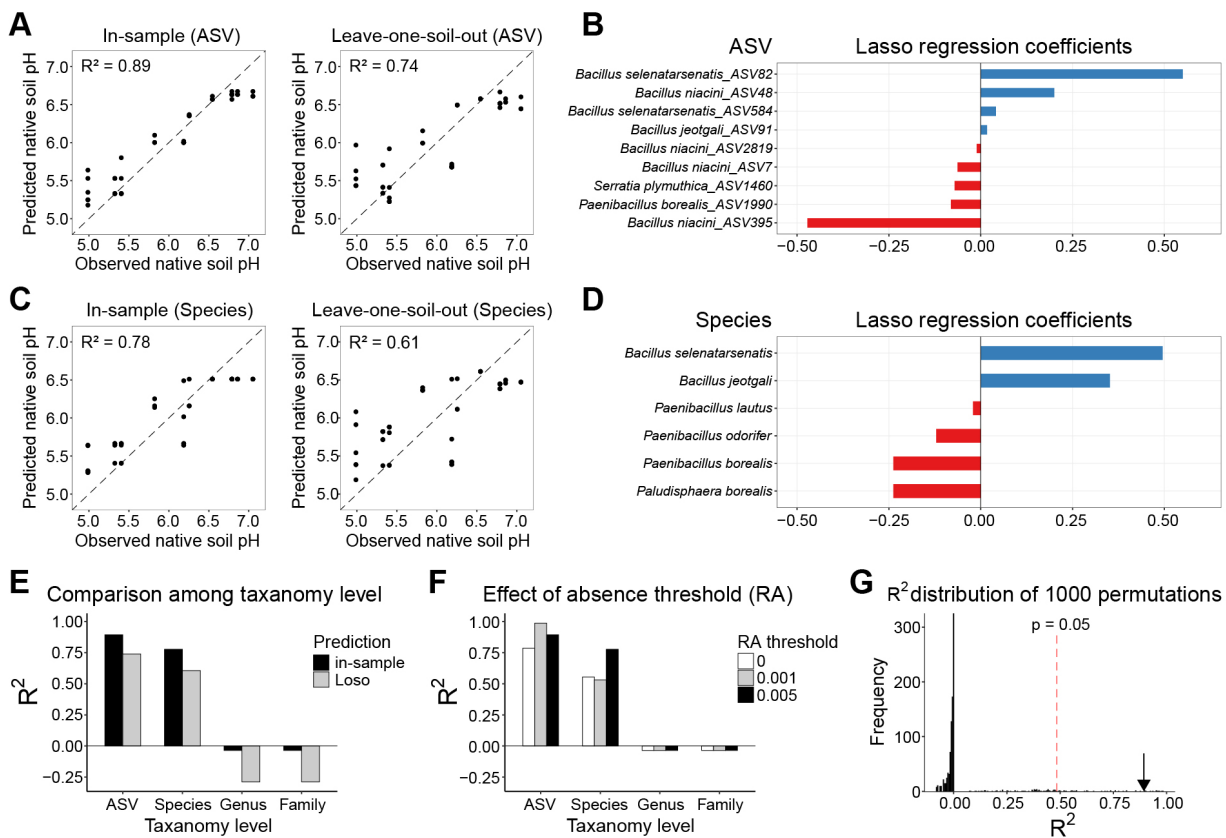
(A) To infer death, the survival folds of the top 10 phyla (relative abundance-wise) among 40 taxa were determined for each perturbed pH condition by computing the fold difference in the endpoint absolute abundance of each Phylum under CHL+ conditions, relative to their baseline levels at the initial time point ( $Abs_{CHL+}/Abs_{T_0}$ ). We used the abundance in CHL+ conditions to rule out growth and only compute the death effect of pH. A consistent drop of survival folds during acidic perturbation was observed across all phyla except for the Firmicutes phylum. (B) To check if the sequencing data supports our model parameter fitted from functional dynamics, we plotted the survival folds against the fitted  $\tilde{x}(0)$  parameter (indigenous biomass activity) for each perturbed pH level denoted by the color gradient. We removed Regime III data points to focus on the Acidic death regime (Regime I). Indeed, during acidic perturbations (dark blue points), the survival folds decreased with the indigenous biomass activity  $\tilde{x}(0)$  parameter fitted with functional data, except for only the Firmicutes. These results suggest that the impacts of acidic and basic pH perturbations on death are asymmetric rather than symmetric: while acidic conditions cause widespread death, death from basic conditions is less prominent.



**Figure S21: Inferring regime boundary with phyla abundance dynamics** To understand the transition to the Acidic death regime (Regime I) with sequencing data, we computed the survival folds of Proteobacteria and Bacteroidota phyla across perturbed pH levels, which is the absolute abundance ratio of  $Abs_{CHL+}/Abs_{T_0}$  (endpoint absolute abundance in CHL+ conditions compared to the initial time point ( $T_0$ ) for each perturbed pH level) (blue and pink data points in the left two panels). We set an identical survival fold threshold for all soils (red lines) to compute the pH at which the survival fold goes below that threshold during acidic perturbation. We used two distinct definitions to choose a threshold for the survival fold; (1) the "dying" definition where the taxa's abundance started to decline in abundance compared to  $T_0$  (survival fold threshold  $< 1$ , red solid lines), and (2) the "dead" definition where the taxa's abundance was close to 0 (survival fold threshold  $\rightarrow 0$ , red dashed lines). To understand the transition to the Resurgent growth regime (Regime III) with sequencing data, we computed the growth folds of the Firmicutes phylum (green data point in the rightmost panel) by endpoint absolute abundance ratio of  $Abs_{CHL-}/Abs_{CHL+}$  (chloramphenicol untreated/treated conditions), and similarly computed the boundary pH at which the growth folds started increasing (threshold = 3, red solid lines). **(B)** These pH transition points were plotted against the native soil pH level. For the pH transition points in Proteobacteria and Bacteroidota, we used the "dying" definition. Employing the 'dying' definition with Proteobacteria, Bacteroidota allowed us to recapitulate the phenomenon observed in the functional data, where the fitted slope of Boundary I-II was less than 1, as shown in Fig. 6A. This suggests that these phyla in the relatively neutral soil are more tolerant of larger  $\Delta$ pH change until they start to die than those in acidic soils, possibly due to variations in soil titration curves (Fig. 6B). (Continued)

Figure S21: (Continued from the previous page) Because the fitted slope is greater than 0, this also means that these phyla in relatively acidic soils can tolerate lower acidic pH conditions than those in neutral soils, which signals long-term pH adaptation. **(C)** The pH transition points were plotted against the native soil pH level using the "dead" definition for Proteobacteria and Bacteroidota. The "dead" definition threshold resulted in a flat slope close to 0. This suggests that, despite long-term adaptation to varying native soil pH levels, these taxa have similar pH thresholds at which complete decimation occurs. For **B and C**, error bars represent the pH difference between the two samples neighboring the survival or growth fold threshold. The points indicate the mid-point pH value of these boundary samples. The linear fit was determined using a least squares method (blue and purple dashed lines). The grey dashed line represents  $y = x$ , indicating hypothetical points where there is no difference between native and perturbed pH values ( $\Delta pH = pH_{perturbed} - pH_{native} = 0$ ).





**Figure S22: Strain (ASV) and species-level variation of the Resurgent growth strains arise from the soil's natural pH environment** To investigate whether the taxonomic identity of Resurgent growth (Regime III) strains is determined by long-term adaptation to or selection from different natural pH environments, we performed a regularized regression analysis to see if we can predict the native pH level of the source soil from the presence or absence of Resurgent growth strains across different taxonomic levels (Amplicon sequence variant (ASV), Species, Genus, Family, or higher taxonomic levels) (see Methods). (Continued)

Figure S22: (Continued from the previous page) **(A)** Predicted and observed native soil pH with the Lasso regression from the presence and absence of the ASVs of Resurgent growth strains, using 0.005 (out of the relative abundance of 1) as a threshold for presence (see Methods). Left is the in-sample predictions, and the right is 'Leave-one-soil-out' (Loso) predictions where we leave out samples from one soil when we fit the regression model and then use the left-out samples to make predictions of their native pH as shown in the scatter plot. The prediction quality ( $R^2$ ) was computed using the mean predicted and mean observed native pH levels for each soil. **(B)** Bar plots indicate the regression coefficients of all ASVs with non-zero coefficients from in-sample predictions in **A**. **(C)** Predicted and observed native soil pH from the Lasso regression from the presence and absence of the Resurgent growth strains in the species level. **(D)** Bar plots indicate the regression coefficients of all species with non-zero coefficients from in-sample predictions in **C**. **(E)** In-sample and Loso predictions are good only until the ASV and species level. From the genus level or higher, the predictions are worse than random (negative  $R^2$  values). A relative abundance (RA) threshold of 0.005 was used for the presence and absence. **(F)** Effect of RA threshold values for the presence and absence (0, 0.001, and 0.005 out of 1). In the case of in-sample predictions, imposing the RA threshold improved the predictions at the ASV and species level. **(G)** To ascertain our prediction quality is not an artifact, we randomly permuted the native pH values 1000 times, and then predicted in-sample the native pH, showing that the  $R^2$  value of 0.89 from in-sample predictions from **A** is greater than the top 50th  $R^2$  value (0.475) out of 1000 shuffled predictions ( $p = 0.05$ ).

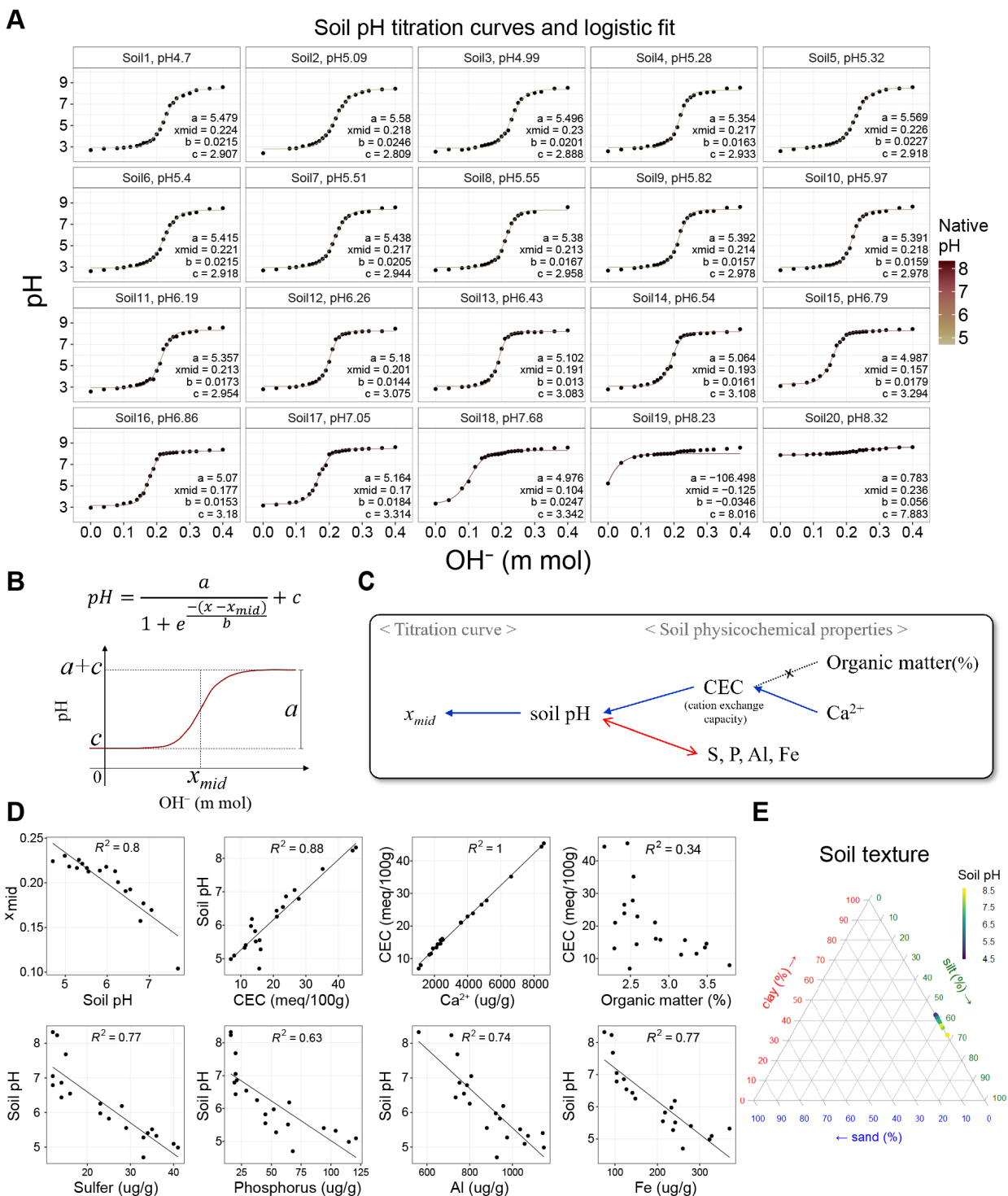


Figure S23:

Figure S23: **Relationship between pH titration curves and soil physicochemical properties** (A) Fitting logistic function to pH titration curves of the 20 soils from different native pH levels (see Methods). (B) Logistic function and parameters. (C) Summary of how soil's physicochemical properties can influence the soil pH titration curves. We can attribute the horizontal shift of the pH titration curves to their varying native soil pH levels, which are potentially determined by the Cation exchange capacity (CEC) and the  $Ca^{2+}$  ion concentrations. (D) The correlations that support the claim from the summary diagram C are shown with  $R^2$  values. (E) Soil texture (sand, silt, clay percentage composition) was mostly identical for different soils, thereby not explaining the difference in soil pH levels and the titration curves.

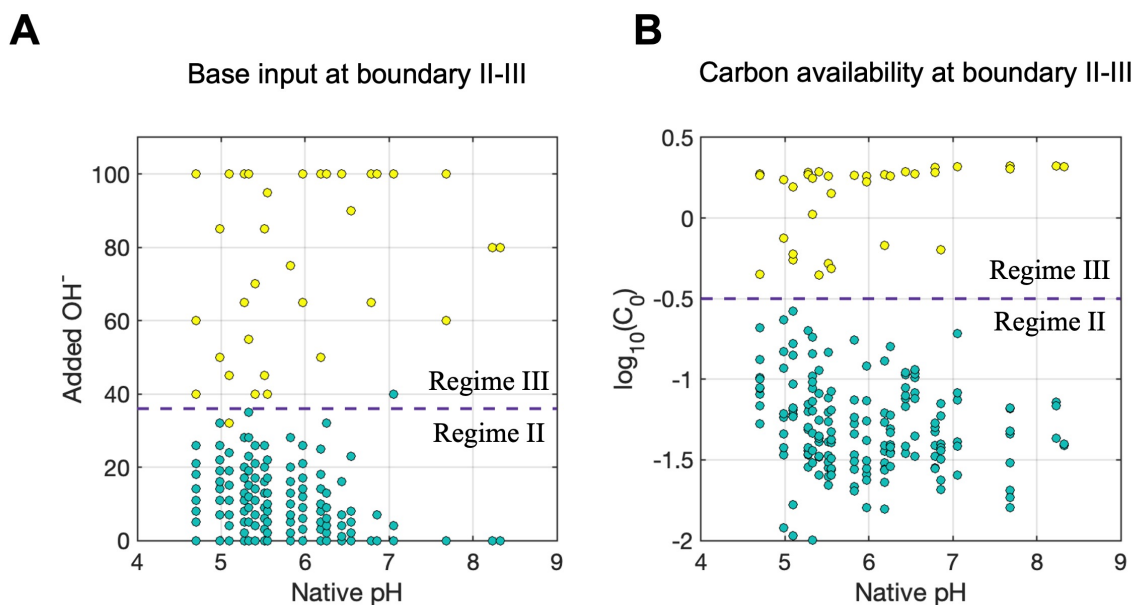
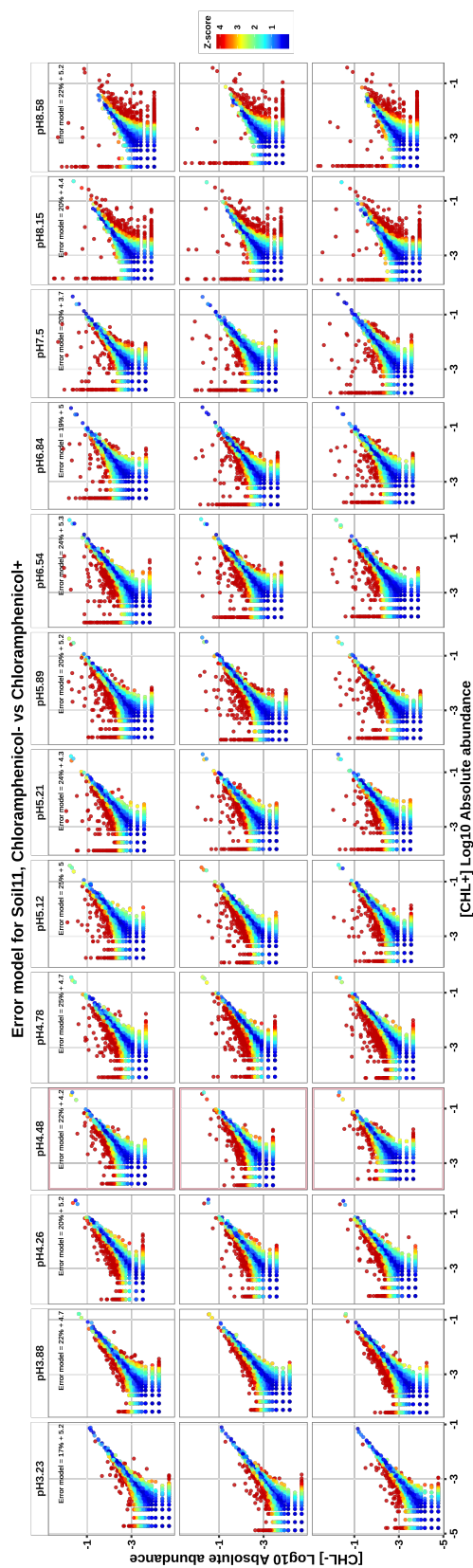


Figure S24: **Functional regime boundary II-III is dictated by the amount of NaOH and the available carbon nutrients.** (A) The relationship of the native soil pH and the amount of NaOH input (y-axis in mM) in Regime II samples (green data points) and Regime III (yellow data points). The dashed purple horizontal line indicates the NaOH input required to transition from Regime II to Regime III. The rather flat slope of pH boundary II-III vs. native pH in Fig. 6A can be explained by the fixed amount of NaOH input (dashed purple line in Fig. 6B). From our previous results, the amount of available carbon corresponds to the NaOH input. (B) Therefore, we plotted the relationship of the native soil pH and the fitted  $\tilde{C}(0)$  values (log scale). We indeed observe the consistent amount of available carbon required to transition from Regime II to Regime III (dashed purple horizontal line).



**Figure S25: Error model z score** To identify the ASVs enriched for each perturbed pH level, we empirically constructed a null model with the three biological replicates (see Methods). For each soil (Soil11 shown in the plot) and for each perturbed pH level (pH indicated in the header of each panel), we plot the log-scale absolute abundance of each ASV in chloramphenicol-treated samples (CHL+, x-axis) against the absolute abundance in chloramphenicol-untreated samples (CHL-, y-axis). Three rows of the panel indicate three biological replicate pairs. The deviations of replicate-replicate comparisons from 1:1 line are well-described by an effective model combining two independent contributions, a Gaussian noise of fractional magnitude  $c_{\text{frac}}$  and a constant Gaussian noise of magnitude  $c_0$  reads, such that repeated measurements (over biological replicates) of an ASV with mean abundance  $n$  counts are approximately Gaussian-distributed with a standard deviation of  $\sigma(c_0, c_{\text{frac}}) = \sqrt{(c_{\text{frac}}n)^2 + c_0^2}$  counts. In this expression,  $c_{\text{frac}}$  was estimated from moderate-abundance ASVs ( $> 50$  counts) for which the other noise term is negligible; and  $c_0$  was then determined as the value for which 67% of replicate-replicate comparisons are within  $\pm\sigma(c_0, c_{\text{frac}})$  of each other, as expected for 1-sigma deviations. This noise model was inferred separately for each soil and each perturbed pH level, as the corresponding samples were processed independently in different sequencing runs. The noise parameters are denoted inside the panel. The points are colored by the z-score computed from the Gaussian-distributed error model. The pink peripheral box indicates the condition without any acid/base addition.