

#### SUPPLEMENTARY INFORMATION

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# Shared strategies for $\beta$ -lactam catabolism in the soil microbiome

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#### Supplementary table 1. Description of previously selected antibiotic-catabolizing soil isolates

Isolate	Prior	Organism	Initial	β-lactam catabolism	
name <sup>a</sup>	designation		selection	Penicillin	Carbenicillin
ABC02	CA-S3F-1	Paraburkholderia	Carbenicillin	+	+
		caledonica CA-			
		S3F-1			
ABC07	PE-S1G-1	Pseudomonas sp.	Penicillin	+	-
		PE-S1G-1			
ABC08	PE-S2R-1	Pandoraea sp.	Penicillin	+	-
		PE-S2R-1			
ABC10	PE-S2T-3	Pandoraea sp.	Penicillin	+	-
		PE-S2T-3			

<sup>&</sup>lt;sup>a</sup>Data, isolate selection and growth described in Dantas et al (2008)<sup>15</sup>

Supplementary table 2. Total  $\beta$ -lactamase and  $\beta$ -lactamase-like genes identified in ABC strain genomes

Ambler class	ABC02	ABC07	ABC08	ABC10	Total
Class A	1	0	0	0	1
Class B	0	0	0	0	0
Class C	5	2	4	4	15
Class D	0	0	1	1	2
Incomplete annotation*	3	8	3	3	17
Total	9	10	8	8	35

<sup>\*</sup>Includes β-lactamases without Ambler class identification, and putative β-lactamases annotated as serine proteases or, predominantly, MBL fold metallo-hydrolases

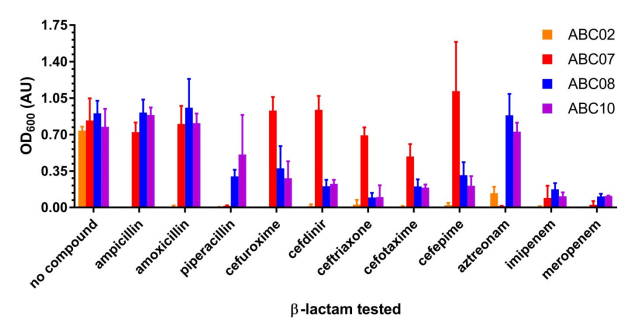
## Supplementary table 3. $\beta\text{-lactamases}$ up-regulated by growth on penicillin in ABC strain genomes

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Ambler class	ABC02	ABC07	ABC08	ABC10	Total
Class A	1	0	0	0	1
Class B	0	0	0	0	0
Class C	1	2	1	1	5
Class D	0	0	1	1	2
Total	2	2	2	2	8

**Supplementary table 4. Primer sequences** 

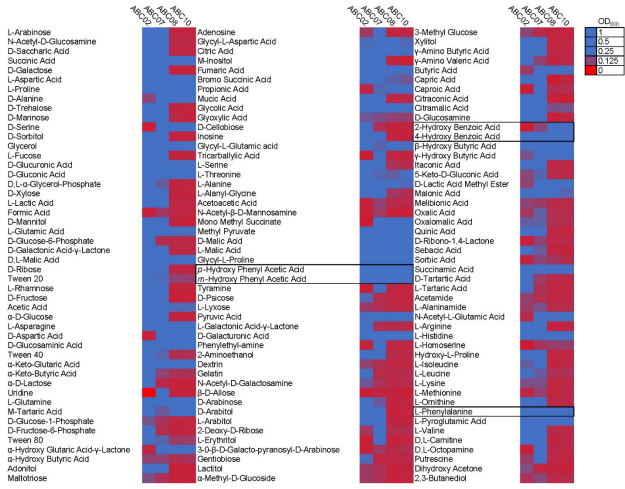
	ntary table 4. Primer sequences	T
Primer	~	_
name	Sequence	Purpose
	GTCGAC TCTAGA	Amplification of <i>put1</i> gene for
5093BW	ATGAGTATTGTTGAACTGGATGC	cloning into pET28B vector
	CCAGTGAA CTCGAG	Amplification of <i>put1</i> gene for
5094BW	TCAACCGGTCGCCGAGAGGC	cloning into pET28B vector
	GTCGAC TCTAGA	Amplification of <i>put2</i> gene for
5095BW	ATGAGCCCACACCAGCACATCC	cloning into pET28B vector
	CCAGTGAA CTCGAG	Amplification of <i>put2</i> gene for
5096BW	TCAGGCGGCTTGAGCCA	cloning into pET28B vector
		Diagnostic PCR for cloning of
54TSC	CGAGATCTCGATCCCGCG	inserts into pET28B vector
		Diagnostic PCR for cloning of
55TSC	CTAGTTATTGCTCAGCGGTGG	inserts into pET28B vector
		Amplification of <i>E. coli</i>
	ATGGGTACCATGAAAAATAGAAAT	penicillin G amidase gene for
5777TSC	CGTATGATCGTG	cloning into pZA11 vector
		Amplification of <i>E. coli</i>
	ATGCTGCAGTTATCTCTGAACGTGC	penicillin G amidase gene for
5779TSC	AACACTTC	cloning into pZA11 vector
		Amplification of pZA11 vector
5780TSC	TTCCTGCAGCCCGG	by inverse PCR
		Amplification of pZA11 vector
5781TSC	CCCGGTACCTTTCTCC	by inverse PCR
		Diagnostic PCR for cloning of
22TSC	GATACTGAGCACATCAGCAGGA	inserts into pZA11 vector
		Diagnostic PCR for cloning of
5723TSC	GGCATCTTCCAGGAAATCTCC	inserts into pZA11 vector
		Diagnostic PCR for gentamicin
6057TSC	atgttacgcagcagcaacg	resistance cassette
		Diagnostic PCR for gentamicin
6058TSC	ttaggtggcggtacttggg	resistance cassette
6059TSC	atgtccaccaacttatcagtgataaag	Diagnostic PCR for <i>tetA</i>
6060TSC	tcagcgatcggctcg	Diagnostic PCR for tetA
	gtcgactctagaggatccccGCTGTGGAGAGG	Amplification of ~1,000 bp
6032TSC	CCGGCT	upstream of paaF
	agttggtggacatGAGTTCACTCGAATTAT	Amplification of ~1,000 bp
6033TSC	TTTTGTGTATCAGCCCAC	upstream of paaF
	gagtgaactcATGTCCACCAACTTATCA	Amplification of <i>tetA</i> for <i>paaF</i>
6034TSC	GTGATAAAG	allelic replacement
		Amplification of <i>tetA</i> for <i>paaF</i>
6035TSC	ggtgcagcagTCAGCGATCGGCTCGTTG	allelic replacement
	cgatcgctgaCTGCTGCACCTTTTTGGCC	Amplification of ~1,000 bp
6036TSC		downstream of <i>paaF</i>
6037TSC	C ccgaattcgagctcgagcccTTGCGCCTGGGT	Amplification of ~1,000 bp

	GCCCTG	downstream of paaF
		Amplification of ~1,000 bp
	tgtaaagcaagcttctgcaggtcgactctagaggatcccc	upstream of penicillin
6087TSC	CCTCCAGCGCCCCGGCC	catabolon
		Amplification of ~1,000 bp
	actgataagttggtggacatGGTGATTCTCCTC	upstream of penicillin
6088TSC	GCAGGCGGTG	catabolon
		Amplification of <i>tetA</i> for
	cgcctgcgaggagaatcaccATGTCCACCAAC	penicillin catabolon allelic
6089TSC	TTATCAGTGATAAAG	replacement
		Amplification of <i>tetA</i> for
	agatacaggcccgtcgcggcTCAGCGATCGG	penicillin catabolon allelic
6090TSC	CTCGTTG	replacement
		Amplification of ~1,000 bp
	ggcaacgagccgatcgctgaGCCGCGACGGG	downstream of penicillin
6091TSC	CCTGTAT	catabolon
		Amplification of ~1,000 bp
	aattaaggtaccgaattcgagctcgagcccGACATA	downstream of penicillin
6092TSC	AAGACTCTTGGGGCAAAAAACCTG	catabolon
		Amplify pEXG2 plasmid for
		Gibson assembly to make
6093TSC	GGGCTCGAGCTCGAATTCGG	ABC07 KOs
		Amplify pEXG2 plasmid for
		Gibson assembly to make
6031TSC	GGGGATCCTCTAGAGTCG	ABC07 KOs
	ATGAACATGCCTATTCCAAACGCCG	
6098TSC	TG	Diagnostic PCR for paaF
6099TSC	TCATGCCTTAGGGCGACGATCC	Diagnostic PCR for paaF



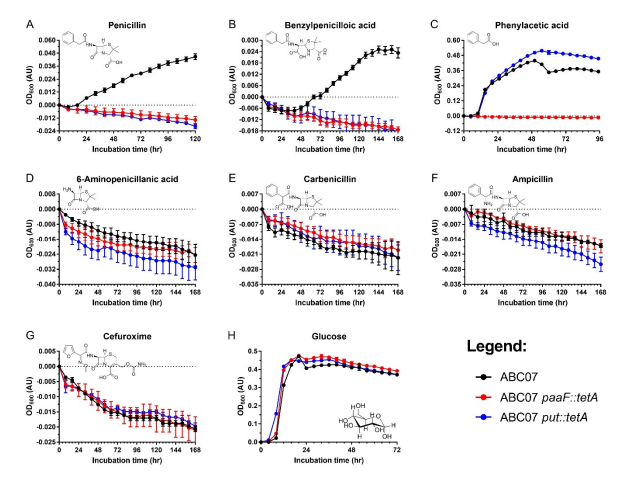
Supplementary figure 1. ABC strains exhibit multi-class, cross-generational resistance to  $\beta$ -lactam antibiotics

ABC strains were grown in rich media with 1 g/l  $\beta$ -lactam antibiotics including  $3^{rd}$  and  $4^{th}$  generation penicillins (ampicillin/amoxicillin and piperacillin respectively),  $2^{nd}$ ,  $3^{rd}$ , and  $4^{th}$  generation cephalosporins (cefuroxime, cefdinir/ceftriaxone/cefotaxime, and cefepime respectively), a monobactam (aztreonam) and two carbapenems (imipenem and meropenem). Average culture density measured by  $OD_{600}$  of three cultures with standard deviation is shown.



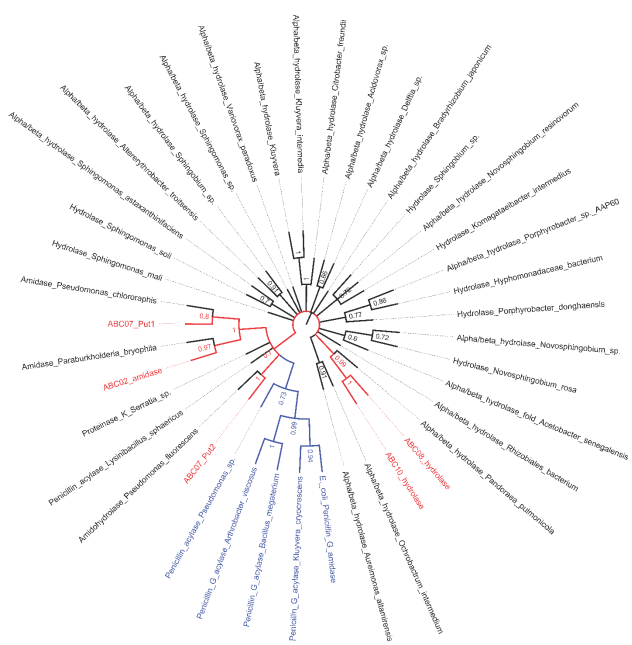
### Supplementary figure 2. ABC strains catabolize diverse substrates including aromatic compounds

Growth of four ABC strains were assayed on 190 different carbon sources. Only carbon sources for which growth exceeded  $OD_{600}$  0.05 for at least one strain are displayed. Aromatic carbon sources are boxed for emphasis.

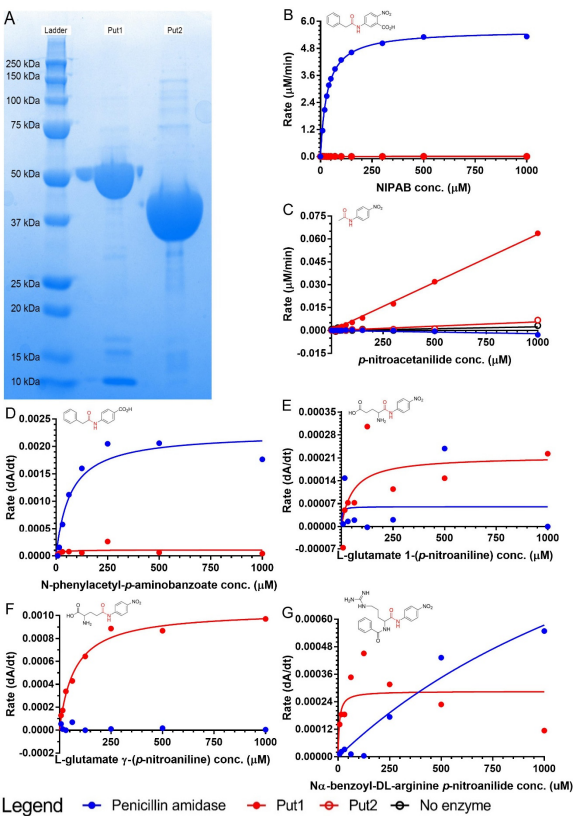


Supplementary figure 3. Growth of ABC07, paaF, and put knock-outs on penicillin degradation products and other β-lactam antibiotics

Growth curves of wild-type ABC07 (black), and *paaF::tetA* (red) and *put::tetA* (blue) strains in minimal media containing (a) penicillin, (b) benzylpenicilloic acid, (c) phenylacetic acid, (d) 6-aminopenicillanic acid, (e) carbenicillin, (f) ampicillin, (g) cefuroxime, or (h) glucose as sole carbon source. Measurements are the average of triplicate cultures with standard error of the mean displayed.



Supplementary figure 4. Phylogenetic analysis of penicillin-responsive amidases, amidohydrolases, and hydrolases from ABC strains compared to penicillin amidases Maximum Likelihood tree of penicillin amidase amino acid sequences (shown in blue), including *E. coli* penicillin amidase and penicillin-upregulated amidase, amidohydrolase, and hydrolase amino acid sequences from ABC strains (shown in red). Node support is given by bootstrap fraction.



Supplementary figure 5. *In vitro* characterization of ABC07 Put1 and Put2 from *E. coli* overexpression

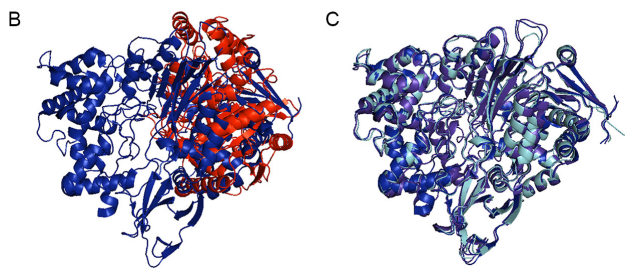
(a) SDS-PAGE gel of his-tagged Put1 and Put2 following over expression in E. coli and purification by IMAC resin. Expected protein masses are approximately 54.4 kDa and 44.0 kDa for Put1 and Put2 respectively. (b through g) Enzyme kinetic analysis of Put1, Put2 (panels b and c only), commercial E. coli penicillin amidase, or no enzyme with (b) 6-nitro-3-(phenylacetamido)benzoic acid (NIPAB) (penicillin amidase K<sub>M</sub> 70.12±16.13 μM), (c) pnitroacetanilide, (d) N-phenylacetyl-p-aminobenzoate (penicillin amidase K<sub>M</sub> 68.97±24.45 μM), (e) L-glutamate 1-(p-nitroaniline), (f) L-glutamate  $\gamma$ -(p-nitroaniline) (Put1 K<sub>M</sub> 73.33±10.53  $\mu$ M), or (g) Nα-benzoyl-DL-arginine p-nitroaniline (penicillin amidase K<sub>M</sub> 1.99±1.93 mM) as substrates.

Α 132 OTYNSVFGTTGNAY------VMSLCESALKDFAAL 292 ABC07 Put1

280 ANGLAGYPTTSNMWVIGKSKA...NPOSGYIANWNNSPOKDYPAS 540 E. coli K. cryocrescens 280 ANGLAGYPTTSNMWVIGKNKA...NPQSGYIANWNNSPQKDYPAS 539 256 EELGLPLKIGSNAAIVGSEKS...NPKNGYVVNWNNKPSKEWVN- 519 B. megaterium

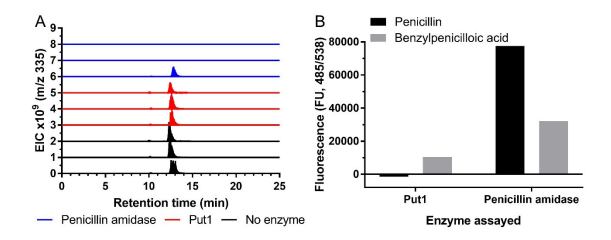
A. viscosus 256 EELGLPLKIGSNAAIVGSEKS...NPKNGYVVNWNNKPSKEWVN- 519

Pseudomonas sp. 230 GDASDAAGGGSNNWAVAPGRT...DPPGGLIVTANNRVVADDHP- 490



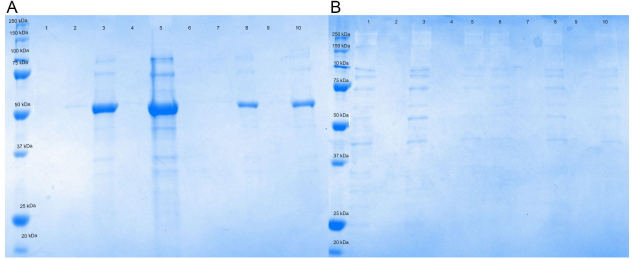
#### Supplementary figure 6. Sequence and structural comparison of Put1 with penicillin amidase enzyme family

(a) Multiple sequence alignment of Put1 enzyme alongside penicillin G amidase homologs. Amino acid residues conserved among penicillin amidases and N-terminal nucleophile hydrolases, the catalytic serine (\*) and the oxyanion hole (#), are highlighted. Overlays and fit (RMSD/aligned residues) of E. coli penicillin amidase with (b) the Phyre predicted Put1 three dimensional structure (6.79 Å/88 residues) and (b) penicillin G amidases homologs from P. rettgeri (1.71 Å/744 residues), and K. citrophilia (1.39 Å/552 residues).



## Supplementary figure 7. Put1 hydrolyzes the amide bond of benzylpenicilloic acid but shows little degradative activity with penicillin

(a) LCMS analysis of penicillin degradation by  $E.\ coli$  penicillin amidase or Put1 by extracted ion count monitoring at 335 m/z corresponding to singly protonated penicillin. Within each condition traces represent (bottom to top) incubation for 30 min, 300 min, and 22.5 hr. (b) Put1 and  $E.\ coli$  penicillin amidase were incubated with penicillin or benzylpenicilloic acid as substrates overnight. Hydrolysis of an amide bond (the phenylacetamide bond in penicillin and benzylpenicilloic acid or the  $\beta$ -lactam ring in penicillin) was detected by derivatization of newly revealed amino groups by the reagent NBD-Cl followed by fluorescent quantification.



Supplementary figure 8. Test conditions for the expression and purification of Put1 from *E. coli* 

SDS-PAGE gels of (a) insoluble and (b) soluble protein fractions from cell-free extracts of *E. coli* BL21(DE3) cells expressing *put1* on pET28B vector under the following conditions: TB brand MB induced with (1) no IPTG, (2) 100 µM IPTG at inoculation, (3) 100 µM IPTG at density, (4) 500 µM IPTG at inoculation, (5) 500 µM IPTG at density, and TB brand FB induced with (6) no IPTG, (7) 100 µM IPTG at inoculation, (8) 100 µM IPTG at density, (9) 500 µM IPTG at inoculation, and (10) 500 µM IPTG at density. For all conditions the majority of Put1 protein (MW 54.4 kDa) is found in the insoluble fraction.