nature portfolio

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Reporting Summary

Nature Portfolio wishes to improve the reproducibility of the work that we publish. This form provides structure for consistency and transparency in reporting. For further information on Nature Portfolio policies, see our <u>Editorial Policies</u> and the <u>Editorial Policy Checklist</u>.

For all statistical analyses, confirm that the following items are present in the figure legend, table legend, main text, or Methods section.

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n/a	Confirmed
	\square The exact sample size (n) for each experimental group/condition, given as a discrete number and unit of measurement
	🔀 A statement on whether measurements were taken from distinct samples or whether the same sample was measured repeatedly
	The statistical test(s) used AND whether they are one- or two-sided Only common tests should be described solely by name; describe more complex techniques in the Methods section.
\boxtimes	A description of all covariates tested
	A description of any assumptions or corrections, such as tests of normality and adjustment for multiple comparisons
	A full description of the statistical parameters including central tendency (e.g. means) or other basic estimates (e.g. regression coefficient) AND variation (e.g. standard deviation) or associated estimates of uncertainty (e.g. confidence intervals)
	For null hypothesis testing, the test statistic (e.g. <i>F</i> , <i>t</i> , <i>r</i>) with confidence intervals, effect sizes, degrees of freedom and <i>P</i> value noted <i>Give P values as exact values whenever suitable.</i>
\boxtimes	For Bayesian analysis, information on the choice of priors and Markov chain Monte Carlo settings
\boxtimes	For hierarchical and complex designs, identification of the appropriate level for tests and full reporting of outcomes
\boxtimes	Estimates of effect sizes (e.g. Cohen's <i>d</i> , Pearson's <i>r</i>), indicating how they were calculated

Our web collection on $\underline{statistics\ for\ biologists}$ contains articles on many of the points above.

Software and code

Policy information about <u>availability of computer code</u>

Data collection

No software was used for data collection.

Data analysis

All data analysis was performed using open source packages, including bayestestR, accessed from RStudio (Version 1.3.1093). No custom algorithms were used. The Agilent MassHunter TOF Quantitative Analysis (Version 10.1) or Agilent MassHunter QQQ Quantitative Analysis software (Version 10.1) was used to quantify the xenobiotic compounds in each sample. TEM image analysis was performed using FIJI and CellProfiler 4.2.6. Proteomics data was processed using Proteome Discoverer v3.0. TPP data analysis was performed using R packages TPP and TPP2D.

For manuscripts utilizing custom algorithms or software that are central to the research but not yet described in published literature, software must be made available to editors and reviewers. We strongly encourage code deposition in a community repository (e.g. GitHub). See the Nature Portfolio guidelines for submitting code & software for further information.

Data

Blinding

Policy information about availability of data

All manuscripts must include a data availability statement. This statement should provide the following information, where applicable:

- Accession codes, unique identifiers, or web links for publicly available datasets
- A description of any restrictions on data availability
- For clinical datasets or third party data, please ensure that the statement adheres to our policy

All data is included in the supplementary information. All electron microscopy images are available at EBI Bioimage archive (TMP_1715003294892). The raw mass-spectrometry data are available at EBI MetaboLights repository (MTBLS9756, MTBLS9745, MTBLS10603, MTBLS10622, MTBLS10636). The raw proteomics and TPP data have been deposited to the ProteomeXchange Consortium via the PRIDE79 partner repository with the dataset identifier PXD050999, and PXD062206. Raw sequencing reads of the evolved B. uniformis and 16S rRNA genotyping were uploaded to the European Nucleotide Archive (ID PRJEB72794, PRJEB75767).

Research involving human participants, their data, or biological material

Policy information ab and sexual orientatio		rith <u>human participants or human data</u> . See also policy information about <u>sex, gender (identity/presentation),</u> thnicity and racism.	
Reporting on sex and gender		N/A	
Reporting on race, ethnicity, or other socially relevant groupings		N/A	
Population characteristics		N/A	
Recruitment		N/A	
Ethics oversight		N/A	
Note that full information	on on the appro	oval of the study protocol must also be provided in the manuscript.	
Field-spec	cific re	porting	
Please select the one	below that is	the best fit for your research. If you are not sure, read the appropriate sections before making your selection.	
Life sciences	В	ehavioural & social sciences	
For a reference copy of the	e document with a	all sections, see <u>nature.com/documents/nr-reporting-summary-flat.pdf</u>	
Life sciend	ces stu	ıdy design	
All studies must discl	ose on these	points even when the disclosure is negative.	
r a v a	No statistical methods were used to pre-determine sample sizes. Sample sizes were chosen on previous experience and according to those reported in previous publications 36,56. For animal studies, to obtain approval from the ethics committee in Germany, we performed a power analysis based on a generalized (i.e. not specific for PFAS) pilot study. Drawing from our 10 years of experience working with gnotobiotic mice, we showed that group sizes of 5 to 9 animals provide sufficient power to detect chemical-microbiome interactions. Based on this, we received approval for group sizes of 9 animals per group. No animals or data points were excluded from the analysis. All relevant and possible data points were used for statistical comparisons when applicable.		
Data exclusions	Data exclusions No animals or data points were excluded from the analysis.		
, N	Screening experiments were performed in three independent batches; all attempts at replication were successful and no data was excluded. Microbiological experiments were performed independently of each other starting with different inoculates. The number of replicates for each experiments are reported in the main text (e.g. in figure legends). All attempts at replication were successful and no data was excluded from the analysis.		
Randomization No randomization was used in microbiological experiments. Randomization was not relevant since experiments were conduced under controlled conditions and using genetically defined material. In animal experiments, female and male mice were housed in separate cages, and subsequently, cages were randomly assigned to experimental groups. Each experimental group included mice from both sexes.			

No blinding was included. Blinding was considered not critical as the overall aim of the study was observational/discovery oriented. Seeveral

experiments included involvement of different labs and thus providing robustness to the conclusions.

Behavioural & social sciences study design

All studies must disclose on these points even when the disclosure is negative.

Study description

Briefly describe the study type including whether data are quantitative, qualitative, or mixed-methods (e.g. qualitative cross-sectional, quantitative experimental, mixed-methods case study).

Research sample

State the research sample (e.g. Harvard university undergraduates, villagers in rural India) and provide relevant demographic information (e.g. age, sex) and indicate whether the sample is representative. Provide a rationale for the study sample chosen. For studies involving existing datasets, please describe the dataset and source.

Sampling strategy

Describe the sampling procedure (e.g. random, snowball, stratified, convenience). Describe the statistical methods that were used to predetermine sample size OR if no sample-size calculation was performed, describe how sample sizes were chosen and provide a rationale for why these sample sizes are sufficient. For qualitative data, please indicate whether data saturation was considered, and what criteria were used to decide that no further sampling was needed.

Data collection

Provide details about the data collection procedure, including the instruments or devices used to record the data (e.g. pen and paper, computer, eye tracker, video or audio equipment) whether anyone was present besides the participant(s) and the researcher, and whether the researcher was blind to experimental condition and/or the study hypothesis during data collection.

Timing

Indicate the start and stop dates of data collection. If there is a gap between collection periods, state the dates for each sample cohort.

Data exclusions

If no data were excluded from the analyses, state so OR if data were excluded, provide the exact number of exclusions and the rationale behind them, indicating whether exclusion criteria were pre-established.

Non-participation

State how many participants dropped out/declined participation and the reason(s) given OR provide response rate OR state that no participants dropped out/declined participation.

Randomization

If participants were not allocated into experimental groups, state so OR describe how participants were allocated to groups, and if allocation was not random, describe how covariates were controlled.

Ecological, evolutionary & environmental sciences study design

All studies must disclose on these points even when the disclosure is negative.

Study description

Briefly describe the study. For quantitative data include treatment factors and interactions, design structure (e.g. factorial, nested, hierarchical), nature and number of experimental units and replicates.

Research sample

Describe the research sample (e.g. a group of tagged Passer domesticus, all Stenocereus thurberi within Organ Pipe Cactus National Monument), and provide a rationale for the sample choice. When relevant, describe the organism taxa, source, sex, age range and any manipulations. State what population the sample is meant to represent when applicable. For studies involving existing datasets, describe the data and its source.

Sampling strategy

Note the sampling procedure. Describe the statistical methods that were used to predetermine sample size OR if no sample-size calculation was performed, describe how sample sizes were chosen and provide a rationale for why these sample sizes are sufficient.

Data collection

Describe the data collection procedure, including who recorded the data and how.

Timing and spatial scale

Indicate the start and stop dates of data collection, noting the frequency and periodicity of sampling and providing a rationale for these choices. If there is a gap between collection periods, state the dates for each sample cohort. Specify the spatial scale from which the data are taken

Data exclusions

If no data were excluded from the analyses, state so OR if data were excluded, describe the exclusions and the rationale behind them, indicating whether exclusion criteria were pre-established.

Reproducibility

Describe the measures taken to verify the reproducibility of experimental findings. For each experiment, note whether any attempts to repeat the experiment failed OR state that all attempts to repeat the experiment were successful.

Randomization

Describe how samples/organisms/participants were allocated into groups. If allocation was not random, describe how covariates were controlled. If this is not relevant to your study, explain why.

Blinding

Describe the extent of blinding used during data acquisition and analysis. If blinding was not possible, describe why OR explain why blinding was not relevant to your study.

Did the study involve field work?

Yes

Field work, collection and transport

Field conditions	Describe the study conditions for field work, providing relevant parameters (e.g. temperature, rainfall).
Location	State the location of the sampling or experiment, providing relevant parameters (e.g. latitude and longitude, elevation, water depth).
Access & import/export	Describe the efforts you have made to access habitats and to collect and import/export your samples in a responsible manner and in compliance with local, national and international laws, noting any permits that were obtained (give the name of the issuing authority, the date of issue, and any identifying information).
Disturbance	Describe any disturbance caused by the study and how it was minimized.

Reporting for specific materials, systems and methods

We require information from authors about some types of materials, experimental systems and methods used in many studies. Here, indicate whether each material, system or method listed is relevant to your study. If you are not sure if a list item applies to your research, read the appropriate section before selecting a response.

Materials & experime n/a Involved in the study Antibodies Eukaryotic cell lines Palaeontology and a Animals and other o Clinical data Dual use research of	n/a Involved in the study ChIP-seq Flow cytometry MRI-based neuroimaging ganisms
Plants	
Antibodies	
Antibodies used Describe all antibodies used in the study; as applicable, provide supplier name, catalog number, clone name, and lot number.	
Validation	Describe the validation of each primary antibody for the species and application, noting any validation statements on the manufacturer's website, relevant citations, antibody profiles in online databases, or data provided in the manuscript.
Eukaryotic cell line	es
Policy information about <u>ce</u>	Il lines and Sex and Gender in Research
Cell line source(s)	State the source of each cell line used and the sex of all primary cell lines and cells derived from human participants or vertebrate models.
Authentication	Describe the authentication procedures for each cell line used OR declare that none of the cell lines used were authenticated.
Mycoplasma contamination	Confirm that all cell lines tested negative for mycoplasma contamination OR describe the results of the testing for mycoplasma contamination OR declare that the cell lines were not tested for mycoplasma contamination.
Commonly misidentified I (See <u>ICLAC</u> register)	Name any commonly misidentified cell lines used in the study and provide a rationale for their use.

Palaeontology and Archaeology

Specimen provenance

Provide provenance information for specimens and describe permits that were obtained for the work (including the name of the issuing authority, the date of issue, and any identifying information). Permits should encompass collection and, where applicable, export.

Specimen deposition | Indicate where the specimens have been deposited to permit free access by other researchers.

Dating methods	If new dates are provided, describe how they were obtained (e.g. collection, storage, sample pretreatment and measurement), where they were obtained (i.e. lab name), the calibration program and the protocol for quality assurance OR state that no new dates are provided.		
Tick this box to confir	m that the raw and calibrated dates are available in the paper or in Supplementary Information.		
Ethics oversight Identify the organization(s) that approved or provided guidance on the study protocol, OR state that no ethical approval or given was required and explain why not.			
Note that full information on t	the approval of the study protocol must also be provided in the manuscript.		
Animals and othe	r research organisms		
Policy information about <u>st</u> <u>Research</u>	<u>cudies involving animals;</u> <u>ARRIVE guidelines</u> recommended for reporting animal research, and <u>Sex and Gender in</u>		
Laboratory animals	Germfree C57BL/6J mice were bred in house (Gnotobiotic Mouse Facility, Tübingen). The housing conditions were 12:12-h light-dark cycles; Temperature: 22 oC +/- 2 oC; Humidity: 50-56%. Female and male mice between 5 – 6 weeks old were randomly assigned to experimental groups. Mice were kept in groups of 3 mice per cage during the experiment. All animals were scored daily for their health status.		
Wild animals	No wild animals were used in the study		
Reporting on sex	Female and male (number of animals reported in the text) mice between 5 – 6 weeks were used and animals were randomly assigned to experimental groups.		
Field-collected samples	No field collected samples were used in the study		
Ethics oversight	Animal experiments were approved by the local authorities in Germany (Regierungspräsidium Tübingen, H 02/20 G).		
Note that full information on t	the approval of the study protocol must also be provided in the manuscript.		
Clinical data			
Policy information about <u>cl</u> All manuscripts should comply	inical studies with the ICMJE guidelines for publication of clinical research and a completed <u>CONSORT checklist</u> must be included with all submissions.		
Clinical trial registration	Provide the trial registration number from ClinicalTrials.gov or an equivalent agency.		
Study protocol	Note where the full trial protocol can be accessed OR if not available, explain why.		
Data collection	Describe the settings and locales of data collection, noting the time periods of recruitment and data collection.		
Outcomes	Describe how you pre-defined primary and secondary outcome measures and how you assessed these measures.		
Dual use research	n of concern		
Policy information about <u>d</u>	ual use research of concern		
Hazards			
	iberate or reckless misuse of agents or technologies generated in the work, or the application of information presented		
in the manuscript, pose a	threat to:		
No Yes			
Public health			
National security Crops and/or lives	tock		
Ecosystems	Crops and/or livestock		
Any other significant area			

Experiments of concern

Doe	the work involve any of these experiments of concern:
No	Yes
\boxtimes	Demonstrate how to render a vaccine ineffective
\boxtimes	Confer resistance to therapeutically useful antibiotics or antiviral agent
\boxtimes	Enhance the virulence of a pathogen or render a nonpathogen virulent
\boxtimes	Increase transmissibility of a pathogen
\boxtimes	Alter the host range of a pathogen
\boxtimes	Enable evasion of diagnostic/detection modalities
\boxtimes	Enable the weaponization of a biological agent or toxin
∇	Any other notentially harmful combination of experiments and agents

Plants

Seed stocks

Report on the source of all seed stocks or other plant material used. If applicable, state the seed stock centre and catalogue number. If plant specimens were collected from the field, describe the collection location, date and sampling procedures.

Novel plant genotypes

Describe the methods by which all novel plant genotypes were produced. This includes those generated by transgenic approaches, gene editing, chemical/radiation-based mutagenesis and hybridization. For transgenic lines, describe the transformation method, the number of independent lines analyzed and the generation upon which experiments were performed. For gene-edited lines, describe the editor used, the endogenous sequence targeted for editing, the targeting guide RNA sequence (if applicable) and how the editor was applied.

Authentication

Describe any authentication procedures for each seed stock used or novel genotype generated. Describe any experiments used to assess the effect of a mutation and, where applicable, how potential secondary effects (e.g. second site T-DNA insertions, mosiacism, off-target gene editing) were examined.

ChIP-seq

Data deposition

Confirm that both raw and final processed data have been deposited in a public database such as <u>GEO</u>.

Confirm that you have deposited or provided access to graph files (e.g. BED files) for the called peaks.

Data access links

May remain private before publication.

For "Initial submission" or "Revised version" documents, provide reviewer access links. For your "Final submission" document, provide a link to the deposited data.

Files in database submission

Provide a list of all files available in the database submission.

Genome browser session (e.g. <u>UCSC</u>)

Provide a link to an anonymized genome browser session for "Initial submission" and "Revised version" documents only, to enable peer review. Write "no longer applicable" for "Final submission" documents.

Methodology

Replicates

Describe the experimental replicates, specifying number, type and replicate agreement.

Sequencing depth

Describe the sequencing depth for each experiment, providing the total number of reads, uniquely mapped reads, length of reads and whether they were paired- or single-end.

Antibodies

Describe the antibodies used for the ChIP-seq experiments; as applicable, provide supplier name, catalog number, clone name, and lot number.

Peak calling parameters

Specify the command line program and parameters used for read mapping and peak calling, including the ChIP, control and index files

Data quality

Describe the methods used to ensure data quality in full detail, including how many peaks are at FDR 5% and above 5-fold enrichment.

Software

Describe the software used to collect and analyze the ChIP-seq data. For custom code that has been deposited into a community repository, provide accession details.

Flow Cytometry

Plots				
Confirm that:				
The axis labels state the marker and fluorochrome used (e.g. CD4-FITC).				
	ble. Include numbers along axes only for bottom left plot of group (a 'group' is an analysis of identical markers).			
	h outliers or pseudocolor plots.			
A numerical value for number	r of cells or percentage (with statistics) is provided.			
Methodology				
Sample preparation	Describe the sample preparation, detailing the biological source of the cells and any tissue processing steps used.			
Instrument	Identify the instrument used for data collection, specifying make and model number.			
Software	Describe the software used to collect and analyze the flow cytometry data. For custom code that has been deposited into a community repository, provide accession details.			
	Describe the abundance of the relevant cell populations within post-sort fractions, providing details on the purity of the samples and how it was determined.			
	Describe the gating strategy used for all relevant experiments, specifying the preliminary FSC/SSC gates of the starting cell population, indicating where boundaries between "positive" and "negative" staining cell populations are defined.			
Tick this box to confirm that a	a figure exemplifying the gating strategy is provided in the Supplementary Information.			
Magnetic resonance in	naging			
Experimental design				
Design type	Indicate task or resting state; event-related or block design.			
Design specifications	Specify the number of blocks, trials or experimental units per session and/or subject, and specify the length of each trial or block (if trials are blocked) and interval between trials.			
Behavioral performance measure	State number and/or type of variables recorded (e.g. correct button press, response time) and what statistics were used to establish that the subjects were performing the task as expected (e.g. mean, range, and/or standard deviation across subjects).			
Acquisition				
Imaging type(s)	Specify: functional, structural, diffusion, perfusion.			
Field strength	Specify in Tesla			
Sequence & imaging parameters	Specify the pulse sequence type (gradient echo, spin echo, etc.), imaging type (EPI, spiral, etc.), field of view, matrix size, slice thickness, orientation and TE/TR/flip angle.			
Area of acquisition	State whether a whole brain scan was used OR define the area of acquisition, describing how the region was determined.			
Diffusion MRI Used	☐ Not used			
Preprocessing				
Preprocessing software	Provide detail on software version and revision number and on specific parameters (model/functions, brain extraction, segmentation, smoothing kernel size, etc.).			
Normalization	If data were normalized/standardized, describe the approach(es): specify linear or non-linear and define image types used for transformation OR indicate that data were not normalized and explain rationale for lack of normalization.			
•	Describe the template used for normalization/transformation, specifying subject space or group standardized space (e.g. original Talairach, MNI305, ICBM152) OR indicate that the data were not normalized.			
Noise and artifact removal	Describe your procedure(s) for artifact and structured noise removal, specifying motion parameters, tissue signals and			

	censor	

Define your software and/or method and criteria for volume censoring, and state the extent of such censoring.

Statistical	modeling	8	infer	ence
Statistical	modeling	5	HILL	CITCO

Statistical modeling & inference			
	Specify type (mass univariate, multivariate, RSA, predictive, etc.) and describe essential details of the model at the first and second levels (e.g. fixed, random or mixed effects; drift or auto-correlation).		
	Define precise effect in terms of the task or stimulus conditions instead of psychological concepts and indicate whether ANOVA or factorial designs were used.		
Specify type of analysis:	in ROI-based Both		
Statistic type for inference Specify v	oxel-wise or cluster-wise and report all relevant parameters for cluster-wise methods.		
(See Eklund et al. 2016)			
Correction Describe the type of correction and how it is obtained for multiple comparisons (e.g. FWE, FDR, permutation or Monte			
Models & analysis n/a Involved in the study			
Functional and/or effective connectivity	Report the measures of dependence used and the model details (e.g. Pearson correlation, partial correlation, mutual information).		
Graph analysis	Report the dependent variable and connectivity measure, specifying weighted graph or binarized graph, subject- or group-level, and the global and/or node summaries used (e.g. clustering coefficient, efficiency, etc.).		

Multivariate modeling and predictive analysis

Specify independent variables, features extraction and dimension reduction, model, training and evaluation metrics.