# nature portfolio

Corresponding author(s):	Georg Damm, J. Gray Camp, Barbara Treutlei
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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>.

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For a	Il statistical analyses, confirm that the following items are present in the figure legend, table legend, main text, or Methods section.
n/a	Confirmed
$\boxtimes$	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.
	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 Give P values as exact values whenever suitable.
$\boxtimes$	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 $d$ , Pearson's $r$ ), indicating how they were calculated
'	Our web collection on <u>statistics for biologists</u> contains articles on many of the points above.
Sof	tware and code

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

Data collection

Provide a description of all commercial, open source and custom code used to collect the data in this study, specifying the version used OR state that no software was used.

Data analysis

See Methods and "Code availability"

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

Policy information about availability of data

All manuscripts must include a <u>data availability statement</u>. 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

Raw and processed scRNA-seq and snRNA-seq data generated and used in this study were deposited in Mendeley under accession numbers (http://dx.doi.org/10.17632/yp3txzw64c.1) and EMBL-EBI ArrayExpress, respectively.

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Research	involving	human	narticir	nants	their	data	$\circ$ r	hin	logical	l mat	eria
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Policy information abou and sexual orientation a		ith <u>human participants or human data</u> . See also policy information about <u>sex, gender (identity/presentation),</u> :hnicity and racism.				
Reporting on sex and	Geender was not taken into consideration during sample collection and analysis. Sex of human donors has been reported for each obtained sample, and controlled for in data analysis.					
Reporting on race, etl other socially relevan groupings	• • •	No selection, distinction, or analytical accommodation was considered for race, ethnicity, or other socially relevant groupings, as sample acquisition was subject to sporadic availability for exploratory analysis.				
Population characteri	ristics See Methods, in the section "Human liver tissue samples"					
Recruitment		See Methods, in the section "Human liver tissue samples"				
Ethics oversight		Informed consent of the patients for the use of tissue for research purposes was obtained according to the ethical guidelines of Leipzig University Hospital (006/17-ek, 21 March 2017, revised and renewed 12 February 2019).				
Note that full information	on the appro	oval of the study protocol must also be provided in the manuscript.				
Field eneci	fic ro	n outing				
Field-speci						
	_	the best fit for your research. If you are not sure, read the appropriate sections before making your selection.				
X Life sciences		ehavioural & social sciences				
For a reference copy of the do	ocument with a	all sections, see <u>nature.com/documents/nr-reporting-summary-flat.pdf</u>				
_ife science	es stu	ıdy design				
All studies must disclose	e on these	points even when the disclosure is negative.				
Sample size Sam	Sample size was not determined since samples were collected from available donors based on clinical opportunity.					
Data exclusions Dat	Data was excluded if it did not pass quality control metrics as specified in the manuscript methods and supplementary tables.					
reg	enerating an	of cells were sequenced from 3 healthy fresh tissue liver samples; from 12 post-PVE fresh tissue liver samples, including 6 d 6 embolized samples. Transcriptomes of nuclei were sequenced from 3 healthy frozen tissue liver samples. All experimental mparable between technical and biological replicates.				
Randomization No	No randomization was performed since samples were collected from available donors based on clinical opportunity.					
Blinding	Blinding was not preformed due to the exploratory nature of data analysis in this manuscript.					
Reporting	for cr	pecific materials, systems and methods				
<u> </u>	<u> </u>	about some types of materials, experimental systems and methods used in many studies. Here, indicate whether each material,				
		your study. If you are not sure if a list item applies to your research, read the appropriate section before selecting a response.				
Materials & experi	mental s	ystems Methods				
n/a Involved in the stu		n/a Involved in the study				
Antibodies X		ChIP-seq				
Eukaryotic cell lines		Flow cytometry				
	Palaeontology and archaeology MRI-based neuroimaging					
Animals and oth	ner organism	S .				
Clinical data	-lf					
Dual use researd	cn of concer	1				
Plants						

### **Antibodies**

Antibodies used

See Methods, in the section "H/E staining and Immunohistochemistry"

Validation

Antibodies used are commercially available (See Methods, in the section "H/E staining and Immunohistochemistry")

#### **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.