## natureresearch

Corresponding author(s):	Aurélie Ernst
Last updated by author(s):	Dec 4, 2019

## Reporting Summary

Nature Research 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 Research policies, seeAuthors & Referees and theEditorial Policy Checklist.

For all statistical analyses, confirm that the following items are present in the figure legend, table legend, main text, or Methods section.  n/a Confirmed				
n/a Confirmed				
Tya Commined				
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.				
For Bayesian analysis, information on the choice of priors and Markov chain Monte Carlo settings				
For hierarchical and complex designs, identification of the appropriate level for tests and full reporting of outcomes				
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 <u>statistics for biologists</u> contains articles on many of the points above.				
Software and code				
Policy information about <u>availability of computer code</u>				
Data collection All resources used are described in the Methods section. Publicly available software included the human reference genome.				
Data analysis  All software used are described in the Methods section and all codes are publically available.				
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/reviewers. We strongly encourage code deposition in a community repository (e.g. GitHub). See the Nature Research <u>guidelines for submitting code &amp; software</u> for further information.				
Data				
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 list of figures that have associated raw data  - A description of any restrictions on data availability				
Provide your data availability statement here.				
Field-specific reporting				
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.				

Ecological, evolutionary & environmental sciences

Behavioural & social sciences

For a reference copy of the document with all sections, see  $\underline{\text{nature.com/documents/nr-reporting-summary-flat.pdf}}$ 

	•			- 1	•	
Lita	CCION	$\sim$ $\sim$ $\sim$	tiidv	$^{\prime}$	$\cap$ CI	$\tau$ n
$\Box$	sciend	$\mathcal{L} $	LUUV	- ( ) (	- 215	211
	001011		c	<u> </u>	,	$\neg \cdot \cdot$

Outcomes

	1 0			
all studies must discl	ose on these points even when the disclosure is negative.			
Sample size	Sample size was determined by the number of available tumor and control samples.			
Data exclusions	No data were excluded.			
	Whole-genome sequencing (WGS) and whole-exome sequencing (WES) were performed once per sample (typical convention in cancer genome sequencing and personalized oncology).			
Randomization [	No randomization was performed			
Blinding	The investigators were blind when performing the chromothripsis scoring.			
We require information ystem or method listed Materials & expenda Involved in the Antibodies Eukaryotic column Palaeontolog Animals and	ChIP-seq  Flow cytometry			
Human resea	rch participants			
	out <u>studies involving human research participants</u>			
Population charact	we performed WES, WGS and RNAseq of tumor tissue (and matched blood for WGS and WES) from 634 patients with advanced cancer and who were enrolled in the MASTER (Molecularly Aided Stratification for Tumor Eradication Research) program, a registry trial for younger adults with advanced cancer across all histologies and patients with rare tumors. All patients had progressive disease prior to molecular analysis. Detailed information on the diagnoses is provided in Supplementary Table 1.			
Recruitment	For WES and WGS, fresh-frozen tumor specimens and matched normal control samples were collected from adult patients who had been diagnosed with chordoma according to World Health Organization criteria at four German cancer centers (NCT Heidelberg and Heidelberg University Hospital, Heidelberg; West German Cancer Center, Essen; University Hospital Carl Gustav Carus, Dresden; Frankfurt University Hospital, Frankfurt). Samples were pseudonymized, and tumor histology and cellularity were assessed at the Institute of Pathology, Heidelberg University Hospital, prior to further processing.			
Ethics oversight	All patients provided written informed consent under a protocol approved by the Ethics Committee of Heidelberg University, and the study was conducted in accordance with the Declaration of Helsinki.			
lote that full informati	on on the approval of the study protocol must also be provided in the manuscript.			
ماداد ادماد				
Clinical data				
olicy information ab Il manuscripts should	comply with the ICMJE guidelines for publication of clinical research and a completed CONSORT checklist must be included with all submissions.			
Clinical trial registr	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.			

Describe how you pre-defined primary and secondary outcome measures and how you assessed these measures.