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## Reporting Checklist

This checklist is used to ensure good reporting standards and to improve the reproducibility of published results. For more information, please read the journal's Guide to Authors.

- Check here to confirm that the following information is available in the Material & Methods section:
  - the exact sample size (n) for each experimental group/condition, given as a number, not a range;
  - a description of the sample collection allowing the reader to understand whether the samples represent technical or biological replicates (including how many animals, litters, culture, etc.);
  - a statement of how many times the experiment shown was replicated in the laboratory;
  - **definitions of statistical methods and measures**: (For small sample sizes (n<5) descriptive statistics are not appropriate, instead plot individual data points)
    - very common tests, such as t-test, simple  $\chi^2$  tests, Wilcoxon and Mann-Whitney tests, can be unambiguously identified by name only, but more complex techniques should be described in the methods section;
    - o are tests one-sided or two-sided?
    - o are there adjustments for multiple comparisons?
    - o statistical test results, e.g., P values;
    - o definition of 'center values' as median or mean;
    - o definition of error bars as s.d. or s.e.m. or c.i.

Please ensure that the answers to the following questions are reported **in the manuscript itself.** We encourage you to include a specific subsection in the methods section for statistics, reagents and animal models. Below, provide the page number or section and paragraph number.

### Statistics and general methods

 How was the sample size chosen to ensure adequate power to detect a pre-specified effect size? (Give section/paragraph or page #)

For animal studies, include a statement about sample size estimate even if no statistical methods were used.

- Describe inclusion/exclusion criteria if samples or animals were excluded from the analysis. Were the criteria pre-established? (Give section/paragraph or page #)
- 3. If a method of randomization was used to determine how samples/animals were allocated to experimental groups and processed, describe it. (Give section/paragraph or page #)

For animal studies, include a statement about randomization even if no randomization was used.

## Reported in section/paragraph or page #

Page 7, and Supplemental methods page 3

Page 7, and Supplemental methods page 3

N/A

Supplemental methods page 3

N/A

4.	If the investigator was blinded to the group allocation during the experiment and/or when assessing the outcome, state the extent of blinding. (Give section/paragraph or page #)	Page 7, Supplemental methods page 2		
For	animal studies, include a statement about blinding even if no blinding was done.	N/A		
5.	For every figure, are statistical tests justified as appropriate?	Page 3		
Do the data meet the assumptions of the tests (e.g., normal distribution)?		Page 3		
	here an estimate of variation within each group of data?	Page 3		
	he variance similar between the groups that are being statistically compared? (Give section/paragraph or page #)	Page 3		
Reagents		Reported in section/paragraph or page #		
6.	Report the source of antibodies (vendor and catalog number)	Page 6		
7.	Identify the source of cell lines and report if they were recently authenticated (e.g., by STR			
	profiling) and tested for mycoplasma contamination	Page 5		
An	profiling) and tested for mycoplasma	Page 5  Reported in section/paragraph or page #		
An 8.	profiling) and tested for mycoplasma contamination			

10. We recommend consulting the ARRIVE guidelines (<u>PLoS Biol. 8(6)</u>, e1000412,2010) to ensure that other relevant aspects of animal studies are adequately reported.

#### **Human subjects**

# 11. Identify the committee(s) approving the study protocol.

- 12. Include a statement confirming that informed consent was obtained from all subjects.
- 13. For publication of patient photos, include a statement confirming that consent to publish was obtained.
- 14. Report the clinical trial registration number (at <u>ClinicalTrials.gov</u> or equivalent).

## Reported in section/paragraph or page #

Page 4	
Page 4	
N/A	
N/A	

- 15. For phase II and III randomized controlled trials, please refer to the <u>CONSORT statement</u> and submit the CONSORT checklist with your submission.
- 16. For tumor marker prognostic studies, we recommend that you follow the REMARK reporting guidelines.

### Data deposition

- 17. Provide accession codes for deposited data.

  Data deposition in a public repository is mandatory for:
  - a. Protein, DNA and RNA sequences
  - b. Macromolecular structures
  - c. Crystallographic data for small molecules
  - d. Microarray data

### Reported in section/paragraph or page #

N/A			

Deposition is strongly recommended for many other datasets for which structured public repositories exist; more details on our data policy are available in the Guide to Authors. We encourage the provision of other source data in supplementary information or in unstructured repositories such as <u>Figshare</u> and <u>Dryad</u>. We encourage publication of Data Descriptors (see <u>Scientific Data</u>) to maximize data reuse.

18. If computer code was used to generate results that are central to the paper's conclusions, include a statement in the Methods section under "Code availability" to indicate whether and how the code can be accessed. Include version information as necessary and any restrictions on availability.

V/A			