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## 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, see <u>Authors & Referees</u> and the <u>Editorial Policy Checklist</u>.

Sta	atistics		
For	all statistical analys	es, confirm that the following items are present in the figure legend, table legend, main text, or Methods section.	
n/a	Confirmed		
	The exact sam	ple size $(n)$ for each experimental group/condition, given as a discrete number and unit of measurement	
	🗶 A statement o	n whether measurements were taken from distinct samples or whether the same sample was measured repeatedly	
	<b>Y</b>	test(s) used AND whether they are one- or two-sided ests should be described solely by name; describe more complex techniques in the Methods section.	
×	A description	of all covariates tested	
x	A description	of any assumptions or corrections, such as tests of normality and adjustment for multiple comparisons	
x	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 <i>P</i> values as exact values whenever suitable.		
×	For Bayesian analysis, information on the choice of priors and Markov chain Monte Carlo settings		
x	For hierarchic	al 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			
	1	Our web collection on <u>statistics for biologists</u> contains articles on many of the points above.	
So	ftware and c	ode	
Policy information about <u>availability of computer code</u>			
Data collection		The latest version of the software were used in this paper. MUSCLE/muscle 3.8.31) , MCscanX, SweeD/version 3.2.1	
D	ata analysis	MUSCLE software, MCscan software, SweeD software	
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/review We strongly encourage code deposition in a community repository (e.g. GitHub). See the Nature Research guidelines for submitting code & software for further information.			
Da	ta		
All	manuscripts must i - Accession codes, uni - A list of figures that	It <u>availability of data</u> nclude a <u>data availability statement</u> . This statement should provide the following information, where applicable: que identifiers, or web links for publicly available datasets have associated raw data restrictions on data availability	
Data that support the findings of this study are available from the corresponding author upon reasonable request.			
Fi	eld-speci	fic reporting	
Plea	se select the one b	elow that is the best fit for your research. If you are not sure, read the appropriate sections before making your selection.	
x	Life sciences	Behavioural & social sciences Ecological, evolutionary & environmental sciences	

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

## Life sciences study design

Validation

All studies must dis	close on these points even when the disclosure is negative.	
Sample size	we used 166 rice accessions to conduct population genetics study, 134 F8 RIL lines and 451 near-isogenic F2 individuals were grown for phenotype-genotype association analysis. These samples are generally considered large for population analysis.	
Data exclusions	No data exclusion	
Replication	We planted 3 replicates and each replicate have 12 individuals in two rows (6 individuals in each row), with a row spacing of 30 centimetres and a plant spacing of 20 centimetres. Each replicate contain all lines and the there replicates plant according to randomized block design.	
Randomization	Not applicable	
Blinding	Not applicable	
We require informati system or method list  Materials & exp n/a Involved in the state of the system	cell lines  cell lines  x	
Antibodies		
Antibodies used	All the antibodies used in this paper are monoclonal antibody of the fusion tag in the end of interest protein.	

Mouse anti-Myc tag and GFP- tag antibody are brought from abcam company. And used as the protocol provided.