1	Natural variation in a molybdate transporter controls grain molybdenum
2	concentration in rice
3	
4	Xin-Yuan Huang ^{1,*} , Huan Liu ¹ , Yu-Fei Zhu ¹ , Shannon R M Pinson ² , Hong-Xuan Lin ³ ,
5	Mary Lou Guerinot ⁴ , Fang-Jie Zhao ¹ , David E Salt ^{5,*}
6	
7	¹ State Key Laboratory of Crop Genetics and Germplasm Enhancement, College of
8	Resources and Environmental Sciences, Nanjing Agricultural University, Nanjing 210095,
9	China
10	² USDA–ARS Dale Bumpers National Rice Research Center, Stuttgart, AR 72160 USA
11	³ National Key Laboratory of Plant Molecular Genetics, CAS Centre for Excellence in
12	Molecular Plant Sciences and Collaborative Innovation Center of Genetics & Development,
13	Shanghai Institute of Plant Physiology & Ecology, Shanghai Institute for Biological
14	Sciences, Chinese Academic of Sciences, Shanghai 200032, China
15	⁴ Department of Biological Sciences, Dartmouth College, Hanover, New Hampshire 03755,
16	USA
17	⁵ School of Biosciences, University of Nottingham, Sutton Bonington Campus,
18	Loughborough, Leicestershire LE12 5RD, UK
19	
20	*Authors for Correspondence:
21	Xin-Yuan Huang
22	Tel: +86 2584399562
23	Email: <u>xinyuan.huang@njau.edu.cn</u>
24	
25	David E Salt
26	Tel: +44 1159516339
27	Email: <u>david.salt@nottingham.ac.uk</u>
28	

Total word count (excluding summary, references and legends):	6764	No. of figures:	8 (Figs 1- 8 in color)
Summary:	198	No. of Tables:	0
Introduction:	859	No of Supporting Information files:	11 (Fig. S1-S8; Table S1-S2; Methods S1)
Materials and Methods:	1504		
Results:	2846		
Discussion:	1409		
Acknowledgements:	107		

1 Summary

- Molybdenum (Mo) is an essential micronutrient for most living organisms,
 including humans. Cereals such as rice are the major dietary source of Mo.
 However, little is known about the genetic basis of the variation in Mo content in
 rice (*Oryza sativa* L.) grain.
- We mapped a quantitative trait locus (QTL) *qGMo8* that controls Mo accumulation
 in rice grain by using a recombinant inbred line population and a backcross
 introgression line population.
- We identified a molybdate transporter, *OsMOT1*;1, as the causal gene for this QTL. 9 • 10 OsMOT1:1 exhibits transport activity for molybdate, but not sulfate, when heterogeneously expressed in yeast cells. OsMOT1;1 is mainly expressed in roots 11 12 and is involved in the uptake and translocation of molybdate under molybdate limited condition. Knock-down of OsMOT1;1 results in less Mo being translocated 13 14 to shoots, lower Mo concentration in grains and higher sensitivity to Mo deficiency. We reveal that the natural variation of Mo concentration in rice grains is attributed 15 16 to the variable expression of OsMOT1;1 due to sequence variation in its promoter. Identification of natural allelic variation in OsMOT1;1 may facilitate the 17 • development of rice varieties with Mo enriched grain for dietary needs and improve 18
- 19 Mo nutrition of rice on Mo-deficient soils.
- 20

21 Key words: molybdenum, mineral nutrient, natural variation, QTL, rice

1 Introduction

2 As one of the most important staple crops, rice not only provides more than one fifth of daily calories for half of the world's human population but is also a major source of mineral 3 nutrients such as Mo, an essential micronutrient for almost all living organisms. In humans, 4 Mo is required for the biosynthesis of molybdenum cofactor (Moco), which forms the 5 active site of molybdenum-requiring enzymes (molybdoenzymes) including aldehyde 6 oxidase, xanthine dehydrogenase, sulfite oxidase and amidoxime reducing component 7 (Schwarz & Mendel, 2006). These enzymes participate in crucial processes such as purine 8 9 metabolism and sulfite detoxification and play a vital role in maintaining human health (Schwarz & Mendel, 2006). Deficiency of Moco biosynthesis in humans results in the 10 11 decrease of molybdoenzyme activity, which leads to inheritable progressive neurological damage and even early childhood death (Johnson et al., 1980; Schwarz, 2005). Although 12 Mo deficiency in humans is extremely rare, food crops grown on soils that are freely 13 drained, acidic and rich in iron oxides are known to have lower Mo concentrations 14 15 (Marschner & Rengel, 2012).

16 Mo is also an essential micronutrient for plants. Plants take up Mo mainly as molybdate. Molybdate itself is biologically inactive and must be incorporated into tricyclic pterin to 17 18 form Moco. Moco serves as electron donors and/or acceptors in molybdoenzymes and plays key roles in the assimilation and biogeochemical cycles of carbon, nitrogen, and 19 sulfur (Schwarz & Mendel, 2006; Bittner, 2014). Although Mo deficiency in humans is 20 rare, Mo deficiency in crops is becoming an agricultural problem, especially for crops 21 22 grown on acid soils. In acid soils, the high levels of reactive iron oxides/hydroxides are known to have a strong adsorption of molybdate which deceases the bioavailability of Mo 23 (Marschner & Rengel, 2012). It is estimated that up to 70% of the world's arable is 24 25 characterized acidic, Mo deficiency is thus a widespread agricultural concern (von Uexküll & Mutert, 1995). The deficiency of Mo in soils has been shown to inhibit plant growth and 26 agricultural productivity (Kaiser et al., 2005). Plants suffering from Mo deficiency develop 27 28 the typical "whiptail" phenotypes, which includes mottled lesions on the leaves, rolling of leaves and wilting of leaf edges (Arnon & Stout, 1939). 29

1 Living organisms that require Mo to synthesize molybdoenzymes take up Mo from the 2 environment in an energy dependent process. In *Escherichia coli*, Mo is taken up by a high-3 affinity ABC-type transport system. This system is encoded by modABC genes and composed of a periplasmic molybdate-binding protein (ModA), a membrane channel 4 protein (ModB), and an energy-transducing ATPase protein (ModC) (Grunden & 5 Shanmugam, 1997; Hollenstein et al., 2007). A large number of ABC transporter genes are 6 present in the genome of eukaryotes; however, none of the ABC-type Mo-specific 7 transporters have been identified in eukaryotes (Kaiser et al., 2005). The first eukaryotic 8 9 high-affinity molybdate transporter encoded by MoT1 (CrMOT1) was identified in the green alga Chlamydomonas reinhardtii (Tejada-Jimenez et al., 2007). Knock-down of 10 *CrMOT1* by an antisense RNA strategy inhibited the molybdate transport activity and the 11 12 activity of the Mo-containing enzyme nitrate reductase, indicating a function of CrMOT1 in molybdate transport (Tejada-Jimenez et al., 2007). A high-affinity molybdate 13 14 transporter AtMOT1;1 (also named as MOT1) that shows sequence similarity to CrMOT1 was also identified in Arabidopsis thaliana (Tomatsu et al., 2007; Baxter et al., 2008). 15 16 AtMOT1;1 belongs to group V of the sulfate transporter superfamily and was previously named as Sultr5;2 (Tejada-Jimenez et al., 2013). Knock-out of AtMOT1;1 led to decreased 17 18 accumulation of Mo in both roots and shoots and the atmot1;1 mutant showed Mo deficiency symptoms when grown under limited Mo supply conditions, suggesting an 19 20 essential role of AtMOT1; 1 in uptake of Mo from soil in A. thaliana (Tomatsu et al., 2007; Baxter et al., 2008). Another member of group V of the sulfate transporter superfamily, 21 22 AtMOT1;2 (Sultr5;1; previously named MOT2), was also identified in A. thaliana, which localizes to the tonoplast and functioned in vacuolar molybdate export (Gasber *et al.*, 2011). 23 24 Recently, LjMOT1 was isolated from Lotus japonicus, which is essential for L. japonicus 25 to take up Mo from the soil (Gao et al., 2016; Duan et al., 2017). Very recently, MtMOT1.3 was identified in the model legume *Medicago truncatula*, which is required for nitrogenase 26 activity in root nodules (Tejada-Jimenez et al., 2017). In C. reinhardtii, there is another 27 molybdate transporter CrMOT2, which shows no sequence similarity to CrMOT1 and thus 28 29 is not related to the MOT1 family (Tejada-Jimenez et al., 2011). CrMOT2 is also a highaffinity molybdate transporter and mainly functions under the molybdate deficient 30

conditions (Tejada-Jimenez *et al.*, 2011). To date, molybdate transporters that control Mo
 concentration have not been characterised in staple food crops.

In this study, we identify a quantitative trait locus (QTL) *qGMo8* that controls the variation 3 4 in grain Mo concentration in rice. We show that qGMo8 encodes a molybdate transporter OsMOT1;1. OsMOT1;1 is mainly expressed in roots and exhibits molybdate transport 5 activity when heterogeneously expressed in yeast cells. Loss-of-function of OsMOT1;1 6 results in decreased Mo translocation from roots to shoots, lower Mo concentration in 7 8 grains and higher sensitivity to Mo deficiency. We further reveal that the natural variation 9 of Mo concentration in rice grains is attributed to the variable expression of *OsMOT1*; 1 in 10 roots.

11

Materials and Methods

13 Plant materials and growth conditions

The rice (Oryza sativa L.) recombinant inbred lines derived from a cross between Lemont 14 (LM, *japonica*) and TeQing (TQ, *indica*) (LT-RILs), and TeQing-into-Lemont backcross 15 16 introgression lines (TILs) were described previously (Tabien et al., 2000; Pinson et al., 17 2012; Huang et al., 2016b). The heterogeneous inbred families (HIFs) of OsMOT1; 1 locus 18 were generated as previously described (Tuinstra et al., 1997; Loudet et al., 2005). A PCR 19 marker was developed based on the 222 bp deletion in the promoter of OsMOT1;1 in TQ, 20 and was used for genotyping the TILs. The line TIL669.4 was determined to be heterozygous at the OsMOT1;1 locus. Plants fixed with TQ allele (HIF669.4-TQ) and 21 22 plants fixed with Lemont allele (HIF669.4-LM) were identified in the next generation of 23 TIL669.4, resulting in HIFs for comparing phenotypic effects of the OsMOT1;1 alleles 24 from TQ and LM. The T-DNA insertion mutant for OsMOT1;1 which is in Zhonghua 11 25 background was obtained from Huazhong Agricultural University, China (http://rmd.ncpgr.cn). The segregated plants without T-DNA insertion in OsMOT1;1 gene 26 were used as a WT control. The WT and T-DNA seeds were planted into soil in a 27 28 greenhouse at the University of Aberdeen, U.K., or hydroponically grown in Nanjing 29 Agricultural University, China. The A. thaliana T-DNA insertion mutant for AtMOT1;1

(SALK_118311) was obtained from the Arabidopsis Biological Resource Center (ABRC,
 http://www.arabidopsis.org/abrc/).

The growth of LT-RIL and TIL populations in the field in Texas, U.S.A., and the growth 3 4 of the TIL population in a greenhouse in Purdue University were described previously (Zhang et al., 2014; Huang et al., 2016b). For analysis of the grains and different tissues 5 of the WT and *osmot1;1*, plants were grown in soil in a greenhouse at the University of 6 7 Aberdeen, U.K. as described previously (Huang et al., 2016b). For the hydroponic 8 experiment, WT and osmot1;1 plants were grown in 96-well plates with the bottom 9 removed. The plates were put in tip boxes containing half-strength Kimura B solution with different concentrations of Mo. The growth condition was described previously (Huang et 10 al., 2016b). A. thaliana transgenic plants were grown on MGRL agar media in a growth 11 12 chamber at the University of Aberdeen, U.K. as described (Huang et al., 2016a).

13 QTL analysis and fine mapping of *qGMo8*

The QTL mapping has been performed previously by using both multiple interval mapping 14 (MIM) and Bayesian information criterion (BIC) methods based on the least squares (LS) 15 means of the 5 years-replications for the grain Mo of LT-RIL and three replications of TIL 16 (Zhang et al., 2014). We reperformed QTL analyses based on the individual year data under 17 flooded or unflooded field conditions by using Windows QTL cartographer version 2.5 18 19 (http://statgen.ncsu.edu/qtlcart/WQTLCart.htm) with a composite interval mapping (CIM) method 20 according to previous studies (Huang et al., 2016b). Seven markers were developed in the 21 qGMo8 mapping interval and used to genotype the entire population of 123 TILs. Three representative TILs with different genotypes in the qGMo8 mapping interval and two 22 control TILs were chosen for further replicated phenotypic analyses and fine mapping. By 23 24 integration of the genotypes and grain Mo concentrations of these lines, the qGMo8 was 25 fine mapped to a 522 kb region on the top of chromosome 8. The primer sequences are 26 listed in Table S2.

27 Tissue elemental analysis

The elemental concentrations in grains of LT-RILs, TILs, WT and *osmot1;1* were determined using an inductively couple plasma mass spectrometer (ICP-MS, Elan DRCe, PerkinElmer; or NexION 300D, PerkinElmer) according to previous studies (Zhang *et al.*, 1 2014; Huang et al., 2016b). For determination of elemental concentrations in different 2 organs of WT and *osmot1*; 1 grown in soil, tissues were excised and washed with Milli-Q 3 water and dried at 88°C overnight. For analysis of roots of WT and osmot1;1 grown hydroponically, roots were excised and washed with 0.5 mM CaCl₂ solution three times 4 and rinsed with Milli-Q water once and then dried at 88°C overnight. Samples were 5 6 digested with concentrated HNO₃ at 118°C for 4 h. The elemental concentrations in the digested samples were determined by ICP-MS (Huang et al., 2016b). The Mo 7 8 concentrations in shoots and roots of transgenic A. thaliana plants were determined 9 according to previous studies (Huang et al., 2016a).

10 Genetic and transgenic complementation test

For genetic complementation, *osmot1;1* as well as WT T-DNA progeny were crossed with 11 12 HIF669.4-TQ and HIF669.4-LM, respectively. The concentrations of Mo in grains of F1 plants from each cross were then determined. For transgenic complementation, the full-13 length coding sequence of OsMOT1;1 was PCR amplified using the cDNA of TQ or LM 14 as templates. The correct PCR fragments confirmed by sequencing were ligated into the 15 16 SalI-SpeI site of p1301GFP vector (Huang et al., 2009) to generate the 17 35S:OsMOT1;1(TQ)-GFP and 35S:OsMOT1;1(LM)-GFP constructs. The resulting plasmids were transformed into Agrobacterium tumefaciens strain HA105 and introduced 18 19 into rice *japonica* cv. Zhonghua 11 as described previously (Hiei *et al.*, 1994). *osmot1*;1 20 was crossed with two transgenic OsMOT1;1 overexpression lines, 35S:OsMOT1;1(TQ)-21 GFP and 35S:OsMOT1;1(LM)-GFP. Two independent transgenic lines of 35S:OsMOT1;1(TQ)-GFP and 35S:OsMOT1;1(LM)-GFP were used for crossing. In 22 subsequent F_2 populations, plants containing the transgene in homozygous osmot1;1 23 mutant background were identified by genotyping. Plants without the transgene in WT or 24 25 homozygous osmot1;1 mutant background were used as controls. The grain Mo concentrations in grains from these F2 plants were determined by ICP-MS. The primer 26 27 sequences using for genotyping are listed in Table S2.

28 Expressing OsMOT1;1 in an A. thaliana atmot1;1 mutant

OsMOT1;1 was expressed in an A. thaliana atmot1;1 T-DNA mutant using the 35S
promoter or AtMOT1;1 native promoter. The construction of 35S:OsMOT1;1(TQ)-GFP

1 and 35S:OsMOT1;1(LM)-GFP vectors was as described above. To generate the AtMOT1;1 2 promoter driven OsMOT1:1 expression vectors, the 1,777-bp promoter sequence of 3 AtMOT1;1 was PCR amplified from the genomic DNA of Col-0 using the primers listed in Table S2. The PCR fragment confirmed by sequencing was used to substitute the 35S 4 promoters in the 35S:OsMOT1;1(TQ)-GFP and 35S:OsMOT1;1(LM)-GFP vectors to 5 generate MOT1pro:OsMOT1;1(TQ)-GFP and MOT1pro:OsMOT1;1(LM)-GFP vectors, 6 respectively. These plasmids were also transformed into Agrobacterium tumefaciens strain 7 GV3101 and introduced in A. thaliana mot1 mutant (SALK_118311) using the floral dip 8 9 method (Clough & Bent, 1998). To assay low pH sensitivity, homozygous T3 transgenic 10 plants were grown on MGRL medium with Mo omitted. The low pH media was prepared as described above by adding 35 µl 6 M HCl into 100 ml media after autoclaving. Plants 11 were grown on plates horizontally in a climate-controlled room with temperature of 19-12 22°C, photoperiod of 10 h light $(100 \pm 10 \text{ } \mu\text{mol } \text{m}^{-2} \text{ } \text{s}^{-1})/14$ h dark and humidity of 60%. 13 14 The shoots were weighed after growing for 20 d.

15 Functional analysis of OsMOT1;1 in yeast

16 The molybdate transporting activity of OsMOT1;1 in yeast was determined according to 17 previous studies with modifications [12]. To generate the plasmids for expression of OsMOT1;1 in yeast (Saccharomyces cerevisiae), the full-length coding sequence of 18 OsMOT1;1 was PCR amplified from the cDNA of TQ or LM using the primers listed in 19 20 Table S2. The correct PCR fragments confirmed by sequencing were subcloned into the 21 EcoRI-XhoI site of the yeast expression vector pYX222x (Tomatsu et al., 2007). The expression of OsMOT1;1 in this plasmid was driven by a constitutive triose phosphate 22 isomerase promoter. The resultant plasmids and empty vector were transformed into yeast 23 strain BY4741 (MATa his $3\Delta 1$ leu $2\Delta 0$ met $15\Delta 0$ ura $3\Delta 0$) using a Frozen-EZ Yeast 24 25 Transformation II Kit (ZYMO Research). For molybdate transporting assay, the BY4741 strains transformed with the empty vector pYX222x, pYX222x-OsMOT1; I(TQ) or 26 pYX222x-OsMOT1;1(LM) were inoculated at 30°C overnight in 3 mL of Mo-free SD/-His 27 media (6.7 g L^{-1} yeast nitrogen base without amino acids and without molybdenum 28 (FORMEDIUM), 1.92 g L⁻¹ dropout mix without histidine, 2% (w/v) glucose). A 100 μ L 29 sample of overnight yeast cells was then transferred to 10 ml Mo-free SD/-His media and 30 31 incubated at 30°C until the optical density at 600 nm (O.D.600) reached to around 1.

1 Hexaammonium heptamolybdate was then added to the media to a final concentration of 2 0.5μ M. After shaking at 30°C for 30 min, yeast cells were harvested by centrifugation and 3 then washed three times with 1mM EDTA-Na and once with Milli-Q water. Cells were 4 dried at 80°C overnight, then digested with concentrated HNO₃. The Mo concentration in 5 digested samples were determined by ICP-MS. The amount of yeast was converted by 1ml 6 of 1 O.D.600 culture containing 3×10^7 cells (Baxter *et al.*, 2008).

- 7 To test the sulfate transporting activity of OsMOT1;1, the empty vector pYX222x, 8 *pYX222x-OsMOT1;1(TQ)* and *pYX222x-OsMOT1;1(LM)* were transformed into a yeast mutant CP154-7B (MATa, his3, leu2, ura3, ade2, trp1, sul1::LEU2, sul2::URA3) as above 9 (Tomatsu *et al.*, 2007). An *A. thaliana* high-affinity sulfate transporter gene *SULTR1*;2 was 10 used as positive control. The overnight yeast cultures were collected by centrifugation and 11 12 washed once with Milli-Q water. After adjusting the O.D.600 of yeast cultures to 0.5, 10 μ L yeast suspensions were spotted on synthetic medium (0.5 mM sulfate, 20 g L⁻¹ glucose, 13 10 g L⁻¹ agarose, 20 mg L⁻¹ Ade, 30 mg L⁻¹ L Leu, 20 mg L⁻¹ Ura, and 20 mg of L⁻¹ Trp) 14 with or without 20 mg L^{-1} methionine. The plates were incubated at 30°C for 6 days. 15
- 16 Other methods could be found in Methods S1.
- 17
- 18

19 **Results**

20 Map-based cloning of *qGMo8*

21 We previously used two synthetic rice mapping populations composed of recombinant inbred lines derived from a cross between Lemont (LM, *japonica*) and TeQing (TQ, *indica*) 22 (LT-RILs), and TeQing-into-Lemont backcross introgression lines (TILs), to identify 23 24 QTLs that control the variation in concentration of 16 elements in unmilled rice grain (Zhang et al., 2014). 134 QTLs were identified in either one or both mapping populations 25 26 grown under flooded and/or unflooded field conditions (Zhang et al., 2014). Among 8 QTLs that control the variation in grain Mo concentration, one was detected on the top of 27 chromosome 8 (designated as qGMo8) in both mapping populations in field trials over 28 multiple years and in both flooded and unflooded fields. This QTL explains up to 35% of 29

the variation in grain Mo (Fig. 1a and 1b). The *qGMo8* first observed in grain from field
grown materials was also detected in both grain and leaf tissues from greenhouse grown
TILs, which allows us to fine map the QTL using plants cultivated in the greenhouse (Fig.
1b).

To narrow down the qGMo8 mapping interval, we developed additional markers to 5 genotype the TIL population. Integration of grain Mo concentration and genotype data of 6 7 three TILs narrowed the QTL interval down to a 522 kb region on the top of chromosome 8 8 (Fig. 1c). In this region, a gene (LOC_Os08g01120) annotated as sulfate transporter 9 encodes a protein that shows 57.8% sequence similarity to A. thaliana high-affinity molybdate transporter AtMOT1:1 (Tomatsu et al., 2007; Baxter et al., 2008), which makes 10 it a strong candidate gene for *qGMo8* (Fig. S1a). We thus designated LOC_Os08g01120 11 12 as OsMOT1;1. Sequence analysis revealed a 9-basepair (bp) deletion (Del⁹) and a single 13 nucleotide polymorphism (SNP) in the coding sequence of OsMOT1;1 in TQ (Fig. 1d, Fig. S1b). The 9-bp deletion alters the amino acid sequence from four glutamines in LM to a 14 15 glutamic acid in TQ (Fig. 1d, Fig. S1b). The SNP in the coding sequence also alters the amino acid sequence, with an alanine in LM and a serine in TQ at the position of 125 16 (S125A) (Fig. 1d, Fig. S1b). A 222-bp deletion (Del²²²) and a 16-bp deletion (Del¹⁶) were 17 found in the promoter sequence of OsMOT1;1 in TQ. Meanwhile, there are 19 SNPs in the 18 19 promoter sequence of OsMOT1;1 in TQ. Comparison of OsMOT1;1 sequences of TQ and LM with Nipponbare reference sequence identified a 12-bp deletion (Del¹²) in the coding 20 region of OsMOT1;1 in both TQ and LM, and only a SNP in the promoter sequence of 21 OsMOT1;1 between LM and Nipponbare (Fig. 1d). Sequence alignment showed that none 22 of the variable amino acids between TQ and LM are conserved in the MOT1 protein family 23 (Fig. S1a), suggesting that these amino acids might not alter the function of OsMOT1;1. 24

The existence of residual heterozygosity in TILs is useful for developing appropriate nearly isogenic lines (NIL) by generating heterogeneous inbred families (HIFs) (Tuinstra *et al.*, 1997; Loudet *et al.*, 2005; Huang *et al.*, 2016b). We identified TIL669.4 which is heterozygous at the *OsMOT1;1* locus, and isolated HIF669.4-TQ and HIF669.4-LM in the next generation (Fig. S2). These two HIFs have similar genomic background and only differ in a small genomic region containing homozygous *OsMOT1;1* alleles from TQ or LM, respectively. Both grain and leaf Mo concentrations of HIF669.4-TQ are significant lower than that of HIF669.4-LM, suggesting that the TQ allele is a weak allele (Fig. 1e).
 Quantitative RT-PCR analysis showed that the expression levels of *OsMOT1;1* in both
 shoots and roots of HIF669.4-TQ were significantly lower than that of HIF669.4-LM,
 suggesting that the low Mo in HIF669.4-TQ might be due to the lower expression level of
 OsMOT1;1 (Fig. 1f).

6 OsMOT1;1 is the causal gene for qGMo8

7 To investigate the function of OsMOT1;1, we isolated a T-DNA insertion mutant of 8 OsMOT1;1. The T-DNA insertion at the promoter of OsMOT1;1 significantly decreases its expression (Fig. S3a- S3c). The *osmot1;1* mutant has no obvious phenotype difference 9 10 from the wild-type (WT) when grown in soil in a greenhouse (Fig. 2a). Elemental profile analysis showed that the Mo concentration in the grain of *osmot1*;1 is approximately 58% 11 12 lower than that of the WT (P < 0.001, Student's *t*-test, n = 10) (Fig. 2b and 2c). The low Mo phenotype was also observed in the blades and sheaths of *osmot1;1* (Fig. 2b). Of the 13 14 22 elements determined. Mo is the only element that is significantly changed in the grain, leaves and sheaths of *osmot1;1*, suggesting a specific effect of *OsMOT1;1* mutation on Mo 15 16 concentrations (Fig. 2c). To test whether OsMOT1; 1 is the causal gene for qGMo8, we 17 performed an allelic complementation by crossing *osmot1;1* as well as WT with HIF669.4-TQ and HIF669.4-LM, respectively. The expression level of OsMOT1;1 in the roots of 18 19 both HIF669.4-TQ and HIF669.4-LM was higher than in that of *osmot1;1* (Fig. S3e). The 20 Mo concentrations in both leaves and grain of $osmot1; 1 \times HIF669.4$ -TQ and $osmot1; 1 \times$ 21 HIF669.4-LM F1 plants grown in soil in a greenhouse were significantly higher than that of osmot1;1 (Fig. 2d and 2e), similar to the levels in HIF669.4-TQ and HIF669.4-LM, 22 respectively, suggesting the complementation of *OsMOT1*; 1 from both TQ and LM to the 23 knockout allele. However, significant differences of both leaf and grain Mo concentrations 24 25 between $osmot1; 1 \times HIF669.4$ -TQ F1 and $osmot1; 1 \times HIF669.4$ -LM F1 were observed (Fig. 2d and 2e), indicating differential functional activity of OsMOT1;1 between TQ and 26 LM. To further confirm OsMOT1;1 as the causal gene, we crossed osmot1;1 with 27 OsMOT1;1 overexpression lines in which OsMOT1;1 from either TQ or LM was expressed 28 from a cauliflower mosaic virus (CaMV) 35S promoter in the cv. Zhonghua 11 background. 29 30 When grown in soil in a greenhouse, the Mo concentrations in grains of F2 plants containing the transgene in homozygous *osmot1;1* mutant background was significantly 31

1 higher than those homozygous *osmot1;1* plants without the transgene (Fig. 2e). These

2 results demonstrate that overexpression of *OsMOT1;1* is able to complement the low Mo

3 phenotype of the *osmot1;1* mutant.

4 Low Mo Phenotype of *osmot1;1* knockout mutant

5 At grain maturity stage, the *osmot1;1* mutant grown in soil in the greenhouse has a lower concentration of Mo than WT not only in the grain, but also in the blade and sheath of the 6 7 flag leaf, and most of the nodes and internodes of the main tiller (Fig. 3a). Analysis of the 8 seedlings grown hydroponically in the nutrient solution containing 1 nM Mo showed that osmot1;1 accumulated significant lower Mo in both roots and shoots (Fig. 3b and 3c). 9 10 However, the difference in the root Mo concentration between osmot1;1 and WT 11 disappeared when plants were grown in the nutrient solution containing 10 nM or higher 12 concentration of Mo (Fig. 3b). Similarly, significant differences in the shoot Mo concentration were observed only at the low levels of Mo supply (1 and 10 nM), but not at 13 14 the high levels of Mo supply (100 nm or 1 μ M) (Fig. 3c). These results suggest that OsMOT1;1 might function mainly at low Mo concentration. Consistent with lower Mo 15 16 level in shoots, the Mo concentration in the xylem sap of *osmot1;1* was significantly 17 decreased compared to the WT (Fig. 3d), suggesting that Mo translocation from roots to shoots was affected in *osmot1*;1. Further analysis showed that the Mo concentrations in all 18 tissues except the leaf sheaths of the 5th and 6th leaves of *osmot1;1* were significantly lower 19 20 than that of WT (Fig. 3e).

21 The *osmot1;1* mutant is sensitive to limited Mo supply

Given that osmot1;1 accumulates less Mo in shoots, we investigated the sensitivity of 22 23 osmot1;1 to Mo deficiency. The bioavailability of Mo is strongly dependent on the soil pH with Mo becoming much less bioavailable in acid soils (Marschner & Rengel, 2012). When 24 25 grown in acidified solid media without supplementation of Mo, most of the seeds of 26 osmot1;1 did not germinate, and the growth of the plants from seeds that did germinate was strongly inhibited (Fig. 4a and 4b). In contrast, the WT plants were able to germinate 27 and grow, even though the root growth was also inhibited by low pH (Fig. 4a and 4b). 28 29 These results establish that, without Mo added into the media, *osmot1;1* is more sensitive 30 to low pH compared to the WT. Supplementation of 1 µM Mo to the acidified solid media was able to restore the growth of *osmot1;1* to the level of WT (Fig. 4a and 4b), indicating
that the sensitivity of *osmot1;1* to low pH is due to the deficiency of Mo in the media. We
further showed that the HIF669.4-TQ, which accumulated less Mo (Fig. 1e), was more
sensitive to Mo deficiency at low pH condition compared to the HIF669.4-LM (Fig. S4a
and S4b).

6 Expression pattern and subcellular localization of OsMOT1;1

7 OsMOT1:1 was expressed in most of the plant tissues throughout the growth period of LM 8 except the panicles at the reproductive stage (Fig. 5a). However, the expression of 9 OsMOT1;1 was much stronger in the roots than in the other tissues. In roots of 2-week-old 10 seedlings, OsMOT1;1 was strongly expressed in the lateral roots as investigated by 11 OsMOT1;1 promoter-GUS transgenic rice plants (Fig. 5b). Low levels of GUS signals 12 were detected in other tissues. To test whether the expression of OsMOT1;1 was affected by Mo supply, LM plants were grown in nutrient solution containing 1 µM Mo for one 13 14 week and transferred to nutrient solution with Mo omitted for a further week. The expression of OsMOT1;1 in roots was strongly suppressed by Mo depletion as determined 15 16 by qRT-PCR. However, such suppression was not found in shoots (Fig. 5c).

To investigate the subcellular localization of OsMOT1;1, OsMOT1;1 from TQ or LM was 17 fused with GFP to the C-terminal and expressed under the control of 35S promoter in a 18 japonica variety Zhonghua 11 (Fig. S3d). The GFP fluorescence was co-localized with the 19 signal of a mitochondrial dye Mitotacker, suggesting OsMOT1;1 localizes to the 20 mitochondria (Fig. 5d). We observed that OsMOT1;1 from both TQ and LM were localized 21 to the mitochondria, indicating that the amino acid variation of OsMOT1;1 between TQ 22 and LM has no effect on the subcellular localization (Fig. 5d). Furthermore, the Mo 23 24 concentrations in roots and shoots of transgenic lines were higher than the non-transgenic control line, suggesting the mitochondria-localized OsMOT1;1 is functional (Fig. S5). 25

26 Functional analysis of OsMOT1;1 in A. thaliana

OsMOT1;1 shares 57.8% sequence similarity to AtMOT1;1 (Fig. S1a), a high-affinity
molybdate transporter in *A. thaliana*. The *atmot1;1* mutant accumulates lower levels of Mo
in leaves than WT (Tomatsu *et al.*, 2007; Baxter *et al.*, 2008). To investigate whether
expression of *OsMOT1;1* in *atmot1;1* could suppress its low Mo phenotype, we

1 heterologously expressed OsMOT1;1-GFP in atmot1;1 using 35S promoter (Fig. S6). In 2 the root cells of the transgenic plants, the GFP signals were observed to co-localize with 3 the mitochondria specific dye MitoTracker (Fig. S7), further confirming the mitochondria 4 localization of OsMOT1;1. The Mo concentrations in both roots and shoots of transgenic plants expressing OsMOT1;1 from either TQ or LM were significantly higher than that of 5 atmot1;1, even higher than the WT Col-0 (Fig. 6a). These results suggest that expression 6 of OsMOT1;1 from either TQ or LM could enhance Mo accumulation in atmot1;1. To rule 7 out the ectopic effect of overexpression of OsMOT1;1 driven by 35S promoter, we also 8 9 expressed OsMOT1;1 in atmot1;1 using AtMOT1;1 native promoter (Fig. S6). The Mo concentrations in both roots and shoots of transgenic plants were significantly higher than 10 that of *atmot1;1*, similar to the level in WT Col-0 (Fig. 6b), suggesting that expression of 11 12 OsMOT1;1 using AtMOT1;1 native promoter was able to complement the AtMOT1;1 knockout mutant. Notably, there was no significant difference in either root or shoot Mo 13 14 concentration between transgenic plants expressing *OsMOT1;1* from TQ or LM (Fig. 6b), indicating no functional difference between the *OsMOT1*; 1 allele of TQ and LM. Previous 15 16 studies have shown that the *atmot1;1* mutant is sensitive to low Mo stress (Tomatsu *et al.*, 2007) and shows defective growth in acidic soil in which the bioavailability of Mo is low 17 18 (Poormohammad Kiani et al., 2012). Similar to the sensitivity to acidic soil, atmot1;1 also showed growth inhibition on agar media at low pH without added Mo (Fig. 6c to 6e). The 19 20 transgenic plants expressing OsMOT1;1 from TQ or LM using 35S promoter or MOT1 native promoter completely restore the growth of *atmot1;1* on agar media at low pH (Fig. 21 22 6c to 6e). The results further confirm that *OsMOT1;1* is able to complement the mutation of *AtMOT1*; *1* in *A. thaliana*. 23

24 Molybdate transport activity of OsMOT1;1

The molybdate transport activity of OsMOT1;1 was tested by heterologous expression of *OsMOT1*;1 in yeast (*Saccharomyces cerevisiae*) strain BY4741. Yeast strain BY4741 transformed with empty vector or *OsMOT1;1* from TQ or LM were cultured in Mo-free media to the mid-log phase and then transferred to the media containing 0.5 μ M Mo and incubated for 30 min. The Mo concentration in yeast cells transformed with *OsMOT1;1* was significantly higher than the control strain transformed with an empty vector (Fig. 7a). These results support the conclusion that OsMOT1;1 is able to transport molybdate. Further comparison revealed no difference of Mo concentrations in the strains transformed with
 OsMOT1;1 from TQ or LM (Fig. 7a), indicating that OsMOT1;1 from TQ and LM
 exhibited similar molybdate transporting activity.

4 OsMOT1;1 shares sequence similarity to sulfate transporter genes in rice and thus was previously annotated as a member of the group five sulfate transporter family (Kumar et 5 al., 2011). To determine whether OsMOT1;1 exhibits a sulfate transport activity, we 6 performed complementation analysis of a yeast mutant CP154-7B, which is defective in 7 8 two high-affinity sulfate transporters and is unable to grow on the media containing lower 9 than 1 mM sulfate as the sole sulfur source (Shibagaki et al., 2002; Yoshimoto et al., 2002; Tomatsu et al., 2007). The yeast mutant transformed with SULTR1;2, an A. thaliana high-10 affinity sulfate transporter, was able to grow on the – Met media. However, expression of 11 12 OsMOT1;1 from either TQ or LM was unable to complement the growth defect of the 13 mutant strain on –Met media (Fig. 7b). These results suggest that OsMOT1;1 likely does 14 not exhibit sulfate transport activity.

15 Analysis of natural variation of OsMOT1;1

To investigate the contribution of natural variation at *OsMOT1*;1 to the variation of grain 16 Mo in rice, we analyzed the coding sequences of 1,479 rice accessions which have been 17 18 resequenced (Zhao et al., 2015). Five non-synonymous polymorphisms in the coding sequence of *OsMOT1*; *I* were identified, including the two polymorphic sites between TQ 19 and LM (S125A and Del⁹), and the 12 bp deletion (Del¹²) in both TQ and LM (Table S1 20 and Fig. 1d). To determine the contribution of these three polymorphic sites on the 21 variation of grain Mo, we genotyped 1,097 accessions of USDA rice core collection for 22 which we have previously reported the grain ionomic profile (Pinson et al., 2015). We 23 24 found no significant difference of grain Mo between the two alleles on any of these three polymorphic sites (Fig. 8a, Fig. S8a). We further compared the grain Mo concentrations of 25 26 eight haplotypes derived from the combination of these three polymorphic sites. We observed no significant differences of grain Mo among the eight haplotypes (Fig. 8b and 27 28 Fig. S8b). These results suggested that the variation in the coding region of OsMOT1;1 might not contribute to the variation of grain Mo in the rice population. 29

There are 19 SNPs and two sequence deletions (Del²²² and Del¹⁶) in the promoter sequence 1 of OsMOT1;1 in TQ (Fig. 1d). We determined the contribution of the two major sequence 2 variances, Del²²² and Del¹⁶, on the variation of grain Mo. We genotyped the 1,097 3 accessions of USDA core collection and compared the grain Mo concentrations of 4 accessions with or without the deletions. We found that there was no significant difference 5 between the accessions with or without the Del²²² or Del¹⁶ (Fig. 8c and Fig. S8c). 6 7 Furthermore, the grain Mo concentrations among the accessions with combination of Del²²² and Del¹⁶ were fairly similar (Fig. 8d and Fig. S8d). These results suggested that 8 Del²²² and Del¹⁶ in the promoter of *OsMOT1*;1 might not contribute to the variation of 9 grain Mo. Thus, the different promoter activity of *OsMOT1*;1 between TQ and LM was 10 likely due to the 19 SNPs, which might cause different expression level of OsMOT1;1 11 between TQ and LM. 12

13 Correlation of grain Mo with the expression of OsMOT1;1

14 To determine whether the variation of grain Mo was due to the variation in the OsMOT1;1 expression level, we selected 35 rice accessions from the USDA core collection with grain 15 Mo concentrations ranging from 0.063 to 1.52 μ g g⁻¹ under flooded growth condition and 16 0.084 to 1.23 μ g g⁻¹ under unflooded condition. The expression levels of *OsMOT1;1* were 17 determined in the roots of three-week-old plants grown hydroponically with 1 µM Mo. We 18 found highly significant correlations between the expression level of *OsMOT1*;1 in roots 19 20 from hydroponically grown plants with the Mo concentrations in grains from plants grown in the field under both flooded (Pearson's $R^2 = 0.7368$, p < 0.001) (Fig. 8e) and unflooded 21 (Pearson's $R^2 = 0.6207$, p < 0.001) conditions (Fig. S8e). These results suggest the natural 22 variation of Mo concentration in rice grains is attributed to the variable expression of 23 OsMOT1:1 in roots. 24

25

26 Discussion

As one of the essential mineral nutrients required by plants, Mo plays important roles in nitrate assimilation, abscisic acid biosynthesis, purine degradation and sulfite detoxification (Schwarz & Mendel, 2006; Bittner, 2014). However, the mechanisms of Mo uptake and transport and the regulation these processes are largely unknown in plants,

1 especially in crops. In this study, we identified a QTL qGMo8 that controls the variation 2 of Mo concentration in rice shoots and grains. We determined the causal gene for this QTL 3 to be OsMOT1;1 by genetic and transgenic complementation (Fig. 2d and 2e). OsMOT1;1 was previously annotated as a member of the group V sulfate transporter superfamily 4 (Tejada-Jimenez et al., 2013). We provided evidence that OsMOT1;1 is a molybdate 5 transporter, including the fact that OsMOT1;1 enhances molybdate but not sulfate uptake 6 when heterogeneously expressed in yeast (Fig. 7a and 2b). We found that the difference in 7 grain Mo concentrations between rice cultivars TQ and LM was not due to altered 8 molybdate transporting activity but most likely to the allelic variation at the gene 9 expression level of *OsMOT1*;1. This conclusion is supported by several lines of evidence. 10 First, the molybdate transporting activity of OsMOT1;1 from TQ and LM was similar when 11 12 heterogeneously expressed in yeast (Fig. 7a). Second, the OsMOT1;1 from either TQ or LM was able to complement the low Mo phenotype of *atmot1*; 1 to a similar level when 13 expressed by the *AtMOT1*; *1* native promoter (Fig. 6b, 6c and 6e). Third, the heterogeneous 14 inbred families line HIF669.4-LM with higher expression level of *OsMOT1*; 1 accumulates 15 16 more Mo in the leaves and grains than that of HIF669.4-TQ. Knockout of OsMOT1;1 resulted in decreased Mo concentration in the shoots and grains (Fig. 2b and 2c). We thus 17 18 conclude that OsMOT1;1 is a molybdate transporter in rice and the natural variation of Mo concentration in rice grains is attributed to the allelic variation at the gene expression level 19 20 of OsMOT1;1.

Several molybdate transporters have been identified, including CrMOT1 and CrMOT2 21 from C. reinhardtii (Tejada-Jimenez et al., 2007; Tejada-Jimenez et al., 2011), AtMOT1;1 22 and AtMOT1;2 from A. thaliana (Tomatsu et al., 2007; Baxter et al., 2008; Gasber et al., 23 2011), LjMOT1 from Lotus japonicus (Gao et al., 2016; Duan et al., 2017) and MtMOT1.3 24 from Medicago truncatula (Tejada-Jimenez et al., 2017). AtMOT1;1 has been shown to 25 control the natural variation in leaf Mo concentration in A. thaliana (Baxter et al., 2008). 26 A 53-bp deletion in the promoter of *AtMOT1*; *1* which is located 13 bp upstream from the 27 transcription start-site of AtMOT1;1 was identified as the functional polymorphism 28 29 contributing to decreased leaf Mo concentration (Tomatsu et al., 2007; Baxter et al., 2008). Further analysis revealed a total of six non-coding structural polymorphisms in the 30 AtMOT1;1 promoter, including the 53-bp deletion originally identified in Ler-0 and a 31

duplicated 330-bp insertion that has undergone a 4-bp deletion (Forsberg *et al.*, 2015). The
53-bp deletion is associated with decreased leaf Mo, while the duplicated 330-bp insertion
is associated with elevated leaf Mo. These two polymorphisms control Mo concentration
in leaves by either decreasing or increasing, respectively, the expression level of *AtMOT1;1*.

In this study, we identified 19 SNPs and two deletions, a 222-bp deletion (Del²²²) and a 16-5 bp deletion (Del¹⁶), in the promoter of OsMOT1;1 between TQ and LM (Fig. 1d). We 6 showed that the difference in grain Mo concentration between TQ and LM was due to the 7 8 different expression level of OsMOT1;1 (Fig. 1e and 1f). Further analysis of 35 rice 9 accessions revealed a significant positive correlation between the expression level of OsMOT1;1 in roots and Mo concentration in grains (Fig. 8e and Fig. S8e). However, the 10 variation in grain Mo appears unrelated to the two main non-coding structural 11 polymorphisms, Del²²² and Del¹⁶, as grain Mo concentrations of rice accessions with or 12 without these two deletions were similar (Fig. 8c and 8d, Fig. S8c and S8d). Therefore, the 13 14 difference in grain Mo concentration between TQ and LM may be due to the SNPs in the 15 promoter of *OsMOT1*; 1 which lead to different expression level. At least two SNPs which are significantly associated with the variation of leaf Mo were also identified on the 16 AtMOT1;1 locus, even though the effect of these SNPs on its expression is not clear 17 (Forsberg et al., 2015). Thus, the variation of Mo in both rice and A. thaliana is attributed 18 19 to the variable expression of the *MOT1*; 1 gene. Such conserved genetic variation structure 20 across plant species suggests a role of *MOT1*; *1* gene in the adaptation to the environment. Indeed, A. thaliana accessions with the weak allele of AtMOT1;1 from West Asia appear 21 to adapt to their native habitats where the water extractable Mo content is high in soils 22 (Poormohammad Kiani et al., 2012). We found that the osmot1;1 mutant is sensitive to 23 limited Mo supply condition (Fig. 4a and 4b), similar to the sensitivity of *atmot1*;1 to Mo 24 deficiency (Tomatsu et al., 2007) and to acidic soils in which the bioavailability of Mo is 25 low (Poormohammad Kiani et al., 2012). Therefore, the MOT1;1 gene may play an 26 important role in adaptation to variable molybdate availability in soils caused by 27 environmental changes. 28

The polymorphisms in the coding region of *AtMOT1;1* also affect its function. A single amino acid variation on AtMOT1;1 in L*er*-0 accession alters its molybdate transporting activity (Tomatsu *et al.*, 2007). The hypofunction of AtMOT1;1 in Sha accession was also

1 proved to be caused by a single amino acid change (Poormohammad Kiani et al., 2012). 2 In the present study, we found two amino acid polymorphisms in OsMOT1;1 between TQ 3 and LM (Fig. 1d). However, none of them appears to alter the function of OsMOT1;1 as these two version proteins showed similar molybdate transporting activity and was able to 4 complement the low Mo phenotype of the *atmot1*;1 mutant to a similar level (Fig. 6b, 6c 5 and 6e; Fig. 7a). A 12-bp deletion (Del¹²) in the coding region of OsMOT1;1 of both TQ 6 and LM, compared with Nipponbare, seems not to change its function because the 7 presence/absence of Del¹² is not associated with the variation of grain Mo (Fig. 8a and 8b, 8 Fig. S8a and S8b). Thus, the large sequence diversity of the *MOT1*;1 locus explains well 9 the identification of AtMOT1;1 in controlling the variation of leaf Mo by genome-wide 10 association (GWA) analysis based on either the mean or variance of leaf Mo (Shen et al., 11 2012; Forsberg et al., 2015), and OsMOT1;1 as the potential locus responsible for the 12 variation of rice grain Mo concentration (Norton et al., 2014). 13

14 The expression of *OsMOT1*; *I* is stronger in roots than in shoots (Fig. 5a), similar to the 15 expression pattern of AtMOT1; 1 in A. thaliana (Tomatsu et al., 2007), which is consistent with the fact that AtMOT1;1 mainly functions in roots (Baxter et al., 2008). Under Mo 16 limited condition, OsMOT1;1 is downregulated in roots but not in shoots (Fig. 5a). 17 However, such downregulation was not observed for AtMOT1;1 in roots but rather in 18 19 shoots (Tomatsu *et al.*, 2007), suggesting the different behaviour of *MOT1*; 1 genes in rice 20 and A. thaliana under Mo scarce condition. OsMOT1;1 may mainly function under low Mo condition, as we only observed the difference of Mo concentrations between WT and 21 osmot1;1 in the hydroponic growth system containing 1 or 10 nM Mo but not at relative 22 higher Mo condition (100 nM or 1 μ M) (Fig. 3b and 3c). The Mo concentration in the 23 xylem sap of *osmot1;1* is lower than that of WT (Fig. 3d), suggestion *OsMOT1;1* is also 24 involved in the translocation of Mo from roots to shoots. Meanwhile, the Mo 25 concentrations in the grain, blade and sheath of osmot1;1 only decrease by 58% - 82% 26 compared to the WT (Fig. 2d), suggesting the existence of additional transporters that 27 control the accumulation of Mo in rice. Further studies are required to elucidate the detailed 28 29 function of OsMOT1;1 and to identify other transporters in controlling the Mo homeostasis 30 in rice.

In summary, we have identified *OsMOT1;1* as the causal gene underlying the QTL for Mo accumulation in rice shoots and grains. OsMOT1;1 exhibits molybdate transport activity. The identification of *OsMOT1;1* provides an important insight into the regulation of Mo homeostasis in rice and a useful gene to breed rice varieties resistant to Mo deficiency in soils. Given the importance of cereals as a source of Mo in the human diet, the identification of natural variation at the *OsMOT1;1* locus provides an efficient way to breed rice varieties with Mo enrichment in the grain, which could improve the nutrient quality of grains.

8

9 Acknowledgments

10 We thank B. Lahner, E. Yakubova and John Danku for ICP-MS analysis, Long-Gang Cui 11 and Min Shi for rice transformation, Takehiro Kamiya and Toru Fujiwara for providing pYX222x and pYX222x-SULTR1;2 vectors. We also thank the Rice Mutant Database at 12 13 Huazhong Agricultural University for providing the T-DNA insertion line. This research was supported by the US National Science Foundation, Plant Genome Research Program 14 15 (Grant #IOS 0701119 to D.E.S, M.L.G and S.R.M.P.), the Fundamental Research Funds for the Central Universities (KYZ201714 to X.-Y.H), the Natural Science Foundation of 16 China (31520103914 to F.-J.Z. and D.E.S.; 31772382 to X.-Y.H) and Jiangsu Natural 17 Science Foundation for Distinguished Young Scholars (KB20180023 to X.-Y.H). 18

19 Author contribution

- 20 X.-Y.H. and D.E.S. designed the research; X.-Y.H., H.L., Y.-F.Z., S.R.M.P., H.-X.L., and
- 21 M.L.G. performed the experiments. X.-Y.H. and D.E.S. analysed the data. X.-Y.H., F.-J.Z.,
- and D.E.S. wrote the paper with contributions of S.R.M.P. and M.L.G.
- 23
- 24
- 25

26 **References**

Arnon DI, Stout PR. 1939. Molybdenum as an essential element for higher plants. *Plant Physiology* 14: 599-602.

1	Baxter I, Muthukumar B, Park HC, Buchner P, Lahner B, Danku J, Zhao K, Lee J,
2	Hawkesford MJ, Guerinot ML, Salt DE. 2008. Variation in molybdenum content across
3	broadly distributed populations of Arabidopsis thaliana is controlled by a mitochondrial
4	molybdenum transporter (MOT1). PLoS Genetics 4: e1000004.
5	Bittner F. 2014. Molybdenum metabolism in plants and crosstalk to iron. Front Plant Sci 5: 28.
6	Clough SJ, Bent AF. 1998. Floral dip: a simplified method for Agrobacterium-mediated
7	transformation of Arabidopsis thaliana. Plant Journal 16: 735-743.
8	Duan G, Hakoyama T, Kamiya T, Miwa H, Lombardo F, Sato S, Tabata S, Chen Z,
9	Watanabe T, Shinano T, Fujiwara T. 2017. LjMOT1, a high-affinity molybdate
10	transporter from Lotus japonicus, is essential for molybdate uptake, but not for the delivery
11	to nodules. <i>Plant Journal</i> 90 : 1108-1119.
12	Forsberg SK, Andreatta ME, Huang XY, Danku J, Salt DE, Carlborg O. 2015. The multi-
13	allelic genetic architecture of a variance-heterogeneity locus for molybdenum
14	concentration in leaves acts as a source of unexplained additive genetic variance. PLoS
15	<i>Genetics</i> 11 : e1005648.
16	Gao JS, Wu FF, Shen ZL, Meng Y, Cai YP, Lin Y. 2016. A putative molybdate transporter
17	LjMOT1 is required for molybdenum transport in Lotus japonicus. Physiologia Plantarum
18	158 : 331-340.
19	Gasber A, Klaumann S, Trentmann O, Trampczynska A, Clemens S, Schneider S, Sauer N,
20	Feifer I, Bittner F, Mendel RR, Neuhaus HE. 2011. Identification of an Arabidopsis
21	solute carrier critical for intracellular transport and inter-organ allocation of molybdate.
22	Plant Biology (Stuttgart, Germany) 13: 710-718.
23	Grunden AM, Shanmugam KT. 1997. Molybdate transport and regulation in bacteria. Archives
24	of Microbiology 168: 345-354.
25	Hiei Y, Ohta S, Komari T, Kumashiro T. 1994. Efficient transformation of rice (Oryza sativa L.)
26	mediated by Agrobacterium and sequence analysis of the boundaries of the T-DNA. Plant
27	Journal 6: 271-282.
28	Hollenstein K, Frei DC, Locher KP. 2007. Structure of an ABC transporter in complex with its
29	binding protein. Nature 446: 213-216.
30	Huang XY, Chao DY, Gao JP, Zhu MZ, Shi M, Lin HX. 2009. A previously unknown zinc
31	finger protein, DST, regulates drought and salt tolerance in rice via stomatal aperture
32	control. Genes & Development 23: 1805-1817.
33	Huang XY, Chao DY, Koprivova A, Danku J, Wirtz M, Muller S, Sandoval FJ, Bauwe H,
34	Roje S, Dilkes B, Hell R, Kopriva S, Salt DE. 2016a. Nuclear localised MORE

1 SULPHUR ACCUMULATION1 epigenetically regulates sulphur homeostasis in 2 Arabidopsis thaliana. *PLoS Genetics* **12**: e1006298. 3 Huang XY, Deng F, Yamaji N, Pinson SR, Fujii-Kashino M, Danku J, Douglas A, Guerinot 4 ML, Salt DE, Ma JF. 2016b. A heavy metal P-type ATPase OsHMA4 prevents copper 5 accumulation in rice grain. Nature Communications 7: 12138. Johnson JL, Waud WR, Rajagopalan KV, Duran M, Beemer FA, Wadman SK. 1980. Inborn 6 7 errors of molybdenum metabolism: combined deficiencies of sulfite oxidase and xanthine 8 dehydrogenase in a patient lacking the molybdenum cofactor. Proceedings of the National 9 Academy of Sciences of the United States of America 77: 3715-3719. Kaiser BN, Gridley KL, Ngaire Brady J, Phillips T, Tyerman SD. 2005. The role of 10 molybdenum in agricultural plant production. Annals of Botany 96: 745-754. 11 12 Kumar S, Asif MH, Chakrabarty D, Tripathi RD, Trivedi PK. 2011. Differential expression 13 and alternative splicing of rice sulphate transporter family members regulate sulphur status 14 during plant growth, development and stress conditions. Functional and Integrative Genomics 11: 259-273. 15 Loudet O, Gaudon V, Trubuil A, Daniel-Vedele F. 2005. Quantitative trait loci controlling root 16 17 growth and architecture in Arabidopsis thaliana confirmed by heterogeneous inbred family. 18 Theoretical and Applied Genetics 110: 742-753. 19 Marschner P, Rengel Z 2012. Nutrient availability in soils. *Mineral Nutrition of Higher Plants* 20 (Third Edition). San Diego: Academic Press, 315-330. 21 Norton GJ, Douglas A, Lahner B, Yakubova E, Guerinot ML, Pinson SR, Tarpley L, Eizenga 22 GC, McGrath SP, Zhao FJ, Islam MR, Islam S, Duan G, Zhu Y, Salt DE, Meharg AA, 23 Price AH. 2014. Genome wide association mapping of grain arsenic, copper, molybdenum 24 and zinc in rice (Oryza sativa L.) grown at four international field sites. PloS One 9: e89685. 25 Pinson SRM, Liu GJ, Jia MH, Jia YL, Fjellstrom RG, Sharma A, Wang YG, Tabien RE, Li 26 **ZK. 2012.** Registration of a rice gene-mapping population consisting of 'TeQing'-into-27 'Lemont' backcross introgression lines. Journal of Plant Registrations 6: 128-135. Pinson SRM, Tarpley L, Yan WG, Yeater K, Lahner B, Yakubova E, Huang XY, Zhang M, 28 29 Guerinot ML, Salt DE. 2015. Worldwide genetic diversity for mineral element 30 concentrations in rice grain. Crop Science 55: 294-311. Poormohammad Kiani S, Trontin C, Andreatta M, Simon M, Robert T, Salt DE, Loudet O. 31 32 2012. Allelic heterogeneity and trade-off shape natural variation for response to soil 33 micronutrient. PLoS Genetics 8: e1002814.

1	Schwarz G. 2005. Molybdenum cofactor biosynthesis and deficiency. Cellular and Molecular Life
2	<i>Sciences</i> 62 : 2792-2810.
3	Schwarz G, Mendel RR. 2006. Molybdenum cofactor biosynthesis and molybdenum enzymes.
4	Annual Review of Plant Biology 57: 623-647.
5	Shen X, Pettersson M, Ronnegard L, Carlborg O. 2012. Inheritance beyond plain heritability:
6	variance-controlling genes in Arabidopsis thaliana. PLoS Genetics 8: e1002839.
7	Shibagaki N, Rose A, McDermott JP, Fujiwara T, Hayashi H, Yoneyama T, Davies JP. 2002.
8	Selenate-resistant mutants of Arabidopsis thaliana identify Sultr1;2, a sulfate transporter
9	required for efficient transport of sulfate into roots. Plant Journal 29: 475-486.
10	Tabien RE, Li Z, Paterson AH, Marchetti MA, Stansel JW, Pinson SRM. 2000. Mapping of
11	four major rice blast resistance genes from 'Lemont' and 'Teqing' and evaluation of their
12	combinatorial effect for field resistance. Theoretical and Applied Genetics 101: 1215-1225.
13	Tejada-Jimenez M, Chamizo-Ampudia A, Galvan A, Fernandez E, Llamas A. 2013.
14	Molybdenum metabolism in plants. <i>Metallomics</i> 5: 1191-1203.
15	Tejada-Jimenez M, Galvan A, Fernandez E. 2011. Algae and humans share a molybdate
16	transporter. Proceedings of the National Academy of Sciences of the United States of
17	America 108: 6420-6425.
18	Tejada-Jimenez M, Gil-Diez P, Leon-Mediavilla J, Wen J, Mysore KS, Imperial J, Gonzalez-
19	Guerrero M. 2017. Medicago truncatula Molybdate Transporter type 1 (MtMOT1.3) is a
20	plasma membrane molybdenum transporter required for nitrogenase activity in root
21	nodules under molybdenum deficiency. New Phytologist 216: 1223-1235.
22	Tejada-Jimenez M, Llamas A, Sanz-Luque E, Galvan A, Fernandez E. 2007. A high-affinity
23	molybdate transporter in eukaryotes. Proceedings of the National Academy of Sciences of
24	the United States of America 104: 20126-20130.
25	Tomatsu H, Takano J, Takahashi H, Watanabe-Takahashi A, Shibagaki N, Fujiwara T. 2007.
26	An Arabidopsis thaliana high-affinity molybdate transporter required for efficient uptake
27	of molybdate from soil. Proceedings of the National Academy of Sciences of the United
28	<i>States of America</i> 104 : 18807-18812.
29	Tuinstra MR, Ejeta G, Goldsbrough PB. 1997. Heterogeneous inbred family (HIF) analysis: a
30	method for developing near-isogenic lines that differ at quantitative trait loci. Theoretical
31	and Applied Genetics 95: 1005-1011.
32	von Uexküll HR, Mutert E. 1995. Global extent, development and economic impact of acid soils.
33	<i>Plant and Soil</i> 171 : 1-15.

Yoshimoto N, Takahashi H, Smith FW, Yamaya T, Saito K. 2002. Two distinct high-affinity
 sulfate transporters with different inducibilities mediate uptake of sulfate in Arabidopsis
 roots. *Plant Journal* 29: 465-473.
 Zhang M, Pinson SR, Tarpley L, Huang XY, Lahner B, Yakubova E, Baxter I, Guerinot ML,
 Salt DE. 2014. Mapping and validation of quantitative trait loci associated with

- 6 concentrations of 16 elements in unmilled rice grain. *Theoretical and Applied Genetics* 127:
 7 137-165.
- 8 Zhao H, Yao W, Ouyang Y, Yang W, Wang G, Lian X, Xing Y, Chen L, Xie W. 2015.
 9 RiceVarMap: a comprehensive database of rice genomic variations. *Nucleic Acids*10 *Research* 43: D1018-1022.
- 11 12

13 Figure legends

14 Fig. 1. QTL analysis and cloning of *qGMo8*.

15 (a and b) The LOD profiling of *qGMo8* on chromosome 8 in the LT-RIL population (a) and TIL population (b) grown in multiple years under different conditions. F, flooded; U, 16 unflooded. G, grains of TILs grown in greenhouse; L, leaves of greenhouse grown TILs. 17 (c) Grain Mo concentration and genotype at QTL interval of selected TILs. Data are 18 presented as mean \pm SD (n = 3). Columns with different letters indicate significant 19 difference at $P \le 0.01$ (Tukey's Honestly Significant Difference (HSD) test). DW, dry 20 21 weight. (d) Gene structure and sequence variation of *OsMOT1;1* among Nipponbare (Nipp), TQ and LM. Blue bars and white bars represent exons and untranslated regions, 22 respectively. Vertical lines represent SNPs. (e) The Mo concentration in the grain and leaf 23 of HIF669.4-TQ and HIF669.4-LM. (f) Expression level of OsMOT1;1 in shoots and roots 24 25 of HIF669.4-TQ and HIF669.4-LM. Data in (e) and (f) are presented as mean \pm SD with n = 6 in (e) and 3 in (f). ** indicates significant difference at $P \le 0.01$ (Student's t test). 26

27 Fig. 2. Characterization of the *osmot1;1* mutant and complementation test.

(a) Greenhouse grown WT and *osmot1;1* plants grown at harvesting stage. (b) Mo concentrations in the grain, blade and sheath of WT and *osmot1;1*. Data are presented as means \pm SD (n = 8). ** indicates significant difference between WT and *osmot1;1* at $P \leq$

1 0.01 (Student's t test). (c) Percentage difference of 22 elements in the grain, blade and 2 sheath of *osmot1;1* compared with the WT. Data are visualized in the radar chart. (d) 3 Genetic complementation of *osmot1;1* by crossing with HIF669.4-TQ or HIF669.4-LM. The Mo concentrations in the grain of F1 plants were determined. (e) Transgenic 4 complementation of *osmot1;1* by crossing with *OsMOT1;1* overexpression lines. The grain 5 Mo concentrations of F2 plants in a homozygous mutant background containing 6 35S:OsMOT1;1(TQ)-GFP or 35S:OsMOT1;1(LM)-GFP constructs were determined. Two 7 independent complemented lines were used for crossing. Data in (d, e) are presented as 8 boxplots (n = 6). Boxes with different letters indicate significant difference at $P \le 0.01$ 9 (Tukey's HSD test). DW, dry weight. 10

11 Fig. 3. Low Mo phenotype of the *osmot1;1* mutant.

12 (a) The Mo concentrations in different tissues of greenhouse grown WT and osmot1;1 plants at harvesting stage. (b and c) The Mo concentrations in roots (b) and shoots (c) of 13 14 WT and *osmot1;1*. Plants were hydroponically grown in nutrient solution containing various concentrations of Mo for two weeks. (d) The Mo concentration in the xylem sap 15 16 of WT and *osmot1*;1. Plants were hydroponically grown in Mo-free nutrient solution for 17 one week and treated with various concentrations of Mo for another week. (e) The Mo concentrations in different tissues of WT and osmot1;1 seedlings. Plants were 18 19 hydroponically grown in nutrient solution containing 10 nM of Mo for two weeks. Data (a) 20 are presented as boxplots (n = 11 for WT and n = 10 for *osmot1*;1). Data in (b-e) are presented as means \pm SD with n = 4 in (b, c), 8 in (d, e). * and ** indicate significant 21 difference between WT and *osmot1;1* at $P \le 0.05$ and 0.01 (Student's *t* test), respectively. 22 DW, dry weight. 23

Fig. 4. The *osmot1;1* mutant is sensitive to Mo deficiency at low pH condition.

(a) The phenotype of WT and *osmot1;1* grown under normal or low pH media with or without 1 μ M Mo added. Low pH media was made by adding 50 μ L 6 M HCl to 100 mL agar media after autoclaving. Plants were grown for 20 days. Bar, 3 cm. (b) The plant height of WT and *osmot1;1* grown as in (a). Data are presented as means \pm SD with n = 6. *** indicated significant difference between WT and *osmot1;1* at $P \le 0.001$ (Student's *t* test).

1 Fig. 5. Expression pattern of OsMOT1;1.

2 (a) Expression level of *OsMOT1*; *1* in different organs of Lemont at different growth stages. 3 (b) Histochemical GUS staining of roots of transgenic rice plants transformed with 4 OsMOT1;1 promoter driven GUS constructs. The mature zone (left), elongation zone (middle) and root tip (right) of roots of 2-week-old plants are showed. Scale bars from left 5 to right are 2, 0.5 and 2 mm. (c) Expression of OsMOT1;1 was suppressed in roots under 6 7 Mo deficiency. Lemont (LM) plants were grown hydroponically with 1 μ M Mo for one 8 week and then treated with 1 μ M Mo (+Mo) or without Mo (-Mo) for another week. 9 Relative expression level of was determined by qRT-PCR with three biological replicates. ** indicates significant difference at $P \le 0.01$ (Student's *t* test). (d) Subcellular localization 10 of OsMOT1;1 in stable transgenic rice plants. OsMOT1;1 from TeQing (TQ) or LM were 11 12 fused with GFP at N-terminal and overexpressed under the control of CaMV 35S promoter. 13 Mitochondria were stained with the specific dye MitoTracker. Scale bar, 10 µm.

14 Fig. 6. Functional analysis of OsMOT1;1 in Arabidopsis atmot1;1 mutant.

(a and b) Mo concentrations in roots and shoots of the *mot1* mutant transformed with 15 OsMOT1;1-GFP from TeQing (TQ) or Lemont (LM) driven by CaMV 35S promoter (a) 16 or AtMOT1;1 native promoter (b). Plants were grown on MGRL media containing 24 nM 17 18 Mo for two weeks. Three independent transgenic lines are shown. (c) Phenotype of OsMOT1;1-GFP transgenic lines in mot1 background. Plants were grown on MGRL media 19 without added Mo at control or low pH condition for 20 days. Bar, 1 cm. (d and e) Fresh 20 weight of *mot1* transformed with OsMOT1;1-GFP from TeQing (TQ) or Lemont (LM) 21 driven by CaMV 35S promoter (d) or AtMOT1;1 native promoter (e). Plants were grown 22 as in (c). Three independent transgenic lines are shown. Data in (a) and (b) are shown as 23 24 means \pm SD with three biological replicates. Six plants were combined in each replicate. Data in (d) and (e) are shown as boxplots with n = 7 to 12. Columns or boxplots with 25 26 different letters in (a) to (e) indicate significant difference at $P \le 0.01$ (Tukey's HSD test). DW, dry weight. 27

Fig. 7. Functional analysis of *OsMOT1;1* in yeast.

(a) Molybdate transport activity of OsMOT1;1. Yeast strains transformed with empty
vector, or *OsMOT1;1* from TeQing (TQ) or Lemont (LM) were incubated in media

1 containing 0.5 μ M Mo for 30 min. Mo concentrations in yeast cells were determined. Data 2 are shown as boxplot with n = 3. Boxplots with different letters indicate significant 3 difference at $P \le 0.01$ (Tukey's HSD test). (b) Complementation analysis of a yeast mutant 4 defective in sulfate uptake. The yeast mutant strains transformed with empty vector, 5 *SULTR1;2*, or *OsMOT1;1* from TQ or LM were incubated on media containing 0.5 mM 6 sulfate with or without added Met for four days.

7 Fig. 8. Contribution of allelic variation of *OsMOT1;1* on the variation of grain Mo in

8 USDA rice core collection grown in flooded condition.

9 (a) The grain Mo in rice accessions with different alleles at three polymorphic sites in the 10 coding region of *OsMOT1;1*. (b) The grain Mo in rice accessions with different haplotypes derived from the combination of three polymorphic sites in the coding region of OsMOT1;1. 11 12 (c) The grain Mo in rice accessions with or without the deletions in the promoter region of OsMOT1;1. (d) The grain Mo in rice accessions with different promoter haplotypes derived 13 14 from the combination of two deletions in the coding region of OsMOT1;1. Data in (a - d) are shown as boxplots. Numbers under the boxes are the accession number; numbers above 15 16 the boxes are the P values; the same letters above the boxes indicate no significant 17 difference at $P \le 0.01$ (Tukey's HSD test). + and – represent the presence and absence of the deletion, respectively. The dash color boxes in (b) and (d) mark the haplotypes in TQ, 18 19 LM and Nipp. (e) The correlation of the relative expression of *OsMOT1*; *1* in roots with the 20 grain Mo concentration in 35 rice accessions grown in flooded condition. The expression 21 of OsMOT1;1 in the roots of 3-week-old plants were determined by qRT-PCR. The relative 22 expression level of OsMOT1;1 was normalized the rice actin gene, and presented as the mean of 2⁽-deltaCt) with three biological replicates. 23

24

25 Supporting Information

Additional supporting information may be found in the online version of this article.

Fig. S1 Sequence alignment of MOT1 proteins.

Fig. S2 Schematic diagram of development of heterogeneous inbred families (HIF).

29 Fig. S3 Molecular characterization of OsMOT1;1 T-DNA insertion and OsMOT1;1

30 overexpression lines.

- 1 Fig. S4 The sensitivity of HIF669.4-TQ and HIF669.4-LM to Mo deficiency at low pH
- 2 condition.
- Fig. S5 Overexpression of *OsMOT1;1* increases Mo concentrations in both roots and
 shoots.
- 5 **Fig. S6** Expression level of OsMOT1;1 in Arabidopsis transgenic lines.
- 6 **Fig. S7** Subcellular localization of OsMOT1;1 in Arabidopsis.
- 7 Fig. S8 Contribution of allelic variation of OsMOT1;1 on the variation of grain Mo in
- 8 USDA rice core collection grown in unflooded condition.
- 9 **Table S1**. Sequence variation and allele frequency of OsMOT1;1.
- 10 **Table S2**. The primers used in this study.
- 11 Methods S1 Supporting information for Materials and Methods.
- 12





(a and b) The LOD profiling of *qGMo8* on chromosome 8 in the LT-RIL population (a) and TIL population (b) grown in multiple years under different conditions. F, flooded; U, unflooded. G, grains of TILs grown in greenhouse; L, leaves of greenhouse grown TILs. (c) Grain Mo concentration and genotype at QTL interval of selected TILs. Data are presented as mean \pm SD (n = 3). Columns with different letters indicate significant difference at $P \le 0.01$ (Tukey's Honestly Significant Difference (HSD) test). DW, dry weight. (d) Gene structure and sequence variation of *OsMOT1;1* among Nipponbare (Nipp), TQ and LM. Blue bars and white bars represent exons and untranslated regions, respectively. Vertical lines represent SNPs. (e) The Mo concentration in the grain and leaf of HIF669.4-TQ and HIF669.4-LM. (f) Expression level of *OsMOT1;1* in shoots and roots of HIF669.4-TQ and HIF669.4-LM. Data in (E) and (f) are presented as mean \pm SD with n = 6 in (e) and 3 in (f). ** indicates significant difference at $P \le 0.01$ (Student's *t* test).



Fig 2. Characterization of the osmot1;1 mutant and complementation test.

(a) Greenhouse grown WT and osmot1;1 plants grown at harvesting stage. (b) Mo concentrations in the grain, blade and sheath of WT and osmot1;1. Data are presented as means \pm SD (n = 8). ** indicates significant difference between WT and osmot1;1 at $P \le 0.01$ (Student's *t* test). (c) Percentage difference of 22 elements in the grain, blade and sheath of osmot1;1 compared with the WT. Data are visualized in the radar chart. (d) Genetic complementation of osmot1;1 by crossing with HIF669.4-TQ or HIF669.4-LM. The Mo concentrations in the grain of F1 plants were determined. (e) Transgenic complementation of osmot1;1 by crossing with OsMOT1;1overexpression lines. The grain Mo concentrations of F2 plants in a homozygous mutant background containing 35S:OsMOT1;1(TQ)-GFP or 35S:OsMOT1;1(LM)-GFP constructs were determined. Two independent complemented lines were used for crossing. Data in (d, e) are presented as boxplots (n = 6). Boxes with different letters indicate significant difference at $P \le$ 0.01 (Tukey's HSD test). DW, dry weight.



Fig 3. Low Mo phenotype of the *osmot1;1* mutant.

(a) The Mo concentrations in different tissues of greenhouse grown WT and osmot1;1 plants at harvesting stage. (b and c) The Mo concentrations in roots (b) and shoots (c) of WT and osmot1;1. Plants were hydroponically grown in nutrient solution containing various concentrations of Mo for two weeks. (d) The Mo concentration in the xylem sap of WT and osmot1;1. Plants were hydroponically grown in Mo-free nutrient solution for one week and treated with various concentrations of Mo for another week. (e) The Mo concentrations in different tissues of WT and osmot1;1 seedlings. Plants were hydroponically grown in nutrient solution containing 10 nM of Mo for two weeks. Data (a) are presented as boxplots (n = 11 for WT and n = 10 for osmot1;1). Data in (b-e) are presented as means \pm SD with n = 4 in (b and c), 8 in (d, e). * and ** indicate significant difference between WT and osmot1;1 at $P \leq 0.05$ and 0.01 (Student's t test), respectively. DW, dry weight.





(a) The phenotype of WT and osmot1;1 grown under normal or low pH media with or without 1 µM Mo added. Low pH media was made by adding 50 µL 6 M HCl to 100 mL agar media after autoclaving. Plants were grown for 20 days. Bar, 3 cm. (b) The plant height of WT and osmot1;1 grown as in (A). Data are presented as means ± SD with n = 6. *** indicated significant difference between WT and osmot1;1 at $P \le 0.001$ (Student's *t* test).



Fig 5. Expression pattern and subcellular localization of OsMOT1;1.

(a) Expression level of *OsMOT1;1* in different organs of Lemont at different growth stages. (b) Histochemical GUS staining of roots of transgenic rice plants transformed with *OsMOT1;1* promoter driven GUS constructs. The mature zone (left), elongation zone (middle) and root tip (right) of roots of 2-week-old plants are showed. Scale bars from left to right are 2, 0.5 and 2 mm. (c) Expression of *OsMOT1;1* was suppressed in roots under Mo deficiency. Lemont (LM) plants were grown hydroponically with 1 μ M Mo for one week and then treated with 1 μ M Mo (+Mo) or without Mo (-Mo) for another week. Relative expression level of was determined by qRT-PCR with three biological replicates. ** indicates significant difference at *P* ≤ 0.01 (Student's *t* test). (d) Subcellular localization of OsMOT1;1 in stable transgenic rice plants. OsMOT1;1 from TeQing (TQ) or LM were fused with GFP at N-terminal and overexpressed under the control of CaMV 35S promoter. Mitochondria were stained with the specific dye MitoTracker. Scale bar, 10 µm.



Fig 6 continued

Fig 6. Functional analysis of OsMOT1;1 in Arabidopsis atmot1;1 mutant.

(a and b) Mo concentrations in roots and shoots of the *mot1* mutant transformed with *OsMOT1;1-GFP* from TeQing (TQ) or Lemont (LM) driven by CaMV 35S promoter (a) or *AtMOT1;1* native promoter (b). Plants were grown on MGRL media containing 24 nM Mo for two weeks. Three independent transgenic lines are shown. (c) Phenotype of *OsMOT1;1-GFP* transgenic lines in *mot1* background. Plants were grown on MGRL media without added Mo at control or low pH condition for 20 days. Bar, 1 cm. (d and e) Fresh weight of *mot1* transformed with *OsMOT1;1-GFP* from TeQing (TQ) or Lemont (LM) driven by CaMV 35S promoter (d) or *AtMOT1;1* native promoter (e). Plants were grown as in (c). Three independent transgenic lines are shown. Data in (a) and (b) are shown as means \pm SD with three biological replicates. Six plants were combined in each replicate. Data in (d) and (e) are shown as boxplots with *n* = 7 to 12. Columns or boxplots with different letters in (a) to (e) indicate significant difference at *P* ≤ 0.01 (Tukey's HSD test). DW, dry weight.



Fig 7. Functional analysis of OsMOT1;1 in yeast.

(a) Molybdate transport activity of OsMOT1;1. Yeast strains transformed with empty vector, or *OsMOT1;1* from TeQing (TQ) or Lemont (LM) were incubated in media containing 0.5 μ M Mo for 30 min. Mo concentrations in yeast cells were determined. Data are shown as boxplot with n = 3. Boxplots with different letters indicate significant difference at $P \le 0.01$ (Tukey's HSD test). (b) Complementation analysis of a yeast mutant defective in sulfate uptake. The yeast mutant strains transformed with empty vector, *SULTR1;2*, or *OsMOT1;1* from TQ or LM were incubated on media containing 0.5 mM sulfate with or without added Met for four days.



Fig 8. Contribution of allelic variation of *OsMOT1;1* on the variation of grain Mo in USDA rice core collection grown in flooded condition.

(a) The grain Mo in rice accessions with different alleles at three polymorphic sites in the coding region of OsMOT1;1. (b) The grain Mo in rice accessions with different haplotypes derived from the combination of three polymorphic sites in the coding region of OsMOT1;1. (c) The grain Mo in rice accessions with or without the deletions in the promoter region of OsMOT1;1. (d) The grain Mo in rice accessions with different promoter haplotypes derived from the combination of two deletions in the coding region of OsMOT1;1. Data in (a - d) are shown as boxplots. Numbers under the boxes are the accession number; numbers above the boxes are the P values; the same letters above the boxes indicate no significant difference at $P \le 0.01$ (Tukey's HSD test). + and – represent the presence and absence of the deletion, respectively. The dash color boxes in (b) and (d) mark the haplotypes in TQ, LM and Nipp. (e) The correlation of the relative expression of OsMOT1;1 in roots with the grain Mo concentration in 35 rice accessions grown in flooded condition. The expression of OsMOT1;1 in the roots of 3-week-old plants were determined by qRT-PCR. The relative expression level of OsMOT1;1 was normalized the rice actin gene, and presented as the mean of $2^{(-}$ deltaCt) with three biological replicates.