

Genetic dissection of the shoot and root ionomes of Brassica napus grown with contrasting phosphate supplies

Article

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1	Original Article
2	Genetic dissection of the shoot and root ionomes of Brassica napus grown with
3	contrasting phosphate supplies
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Background and Aims Mineral elements have many essential and beneficial functions
in plants. Phosphorus (P) deficiency can result in changes in the ionomes of plant organs.
The aims of this study were to characterize the effects of P supply on the ionomes of
shoots and roots, and to identify chromosomal quantitative trait loci (QTLs) for shoot
and root ionomic traits, as well as those affecting the partitioning of mineral elements
between shoot and root in *Brassica napus* grown with contrasting P supplies.

Methods Shoot and root concentrations of eleven mineral elements (B, Ca, Cu, Fe, K,
Mg, Mn, Na, P, S and Zn) were investigated by ICP-OES in a *Brassica napus* double
haploid population grown at an optimal (OP) and a low phosphorus supply (LP) in an
agar system. Shoot, root and plant contents, and the partitioning of mineral elements
between shoot and root were calculated.

Key Results The tissue concentrations of B, Ca, Cu, K, Mg, Mn, Na, P and Zn were reduced by P starvation, while the concentration of Fe was increased by P starvation in the *Bna*TNDH population. A total of 133 and 123 QTLs for shoot and root ionomic traits were identified at OP and LP, respectively. A major QTL cluster on chromosome C07 had a significant effect on shoot Mg and S concentrations at LP and was narrowed down to a 2.1-Mb region using an advanced backcross population.

Conclusions The tissue concentration and partitioning of each mineral element was affected differently by phosphorus starvation. There was a significant difference in mineral element composition between shoots and roots. Identification of the genes underlying these QTLs will enhance our understanding of processes affecting the uptake and partitioning of mineral elements in *Brassica napus*.

2	Key	words:	Mineral	element,	concentration,	content,	phosphorus,	Brassica	napus,
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1 INTRODUCTION

Mineral elements have many essential and beneficial functions in plants (Grusak et al., 2 3 2016). They serve in structural components, such as cell walls and membranes, in energy transduction, in proteins and metabolites, in nucleotides, in the osmotic and 4 electrochemical balance of cellular compartments, in detoxification of the cytoplasm, 5 and in the regulation of biological activities (Baxter, 2009; Grusak et al., 2016). The 6 elemental composition of an organism, or a constituent part, is referred to as its ionome 7 (Lahner et al., 2003; Neugebauer et al., 2018). The ionome of plant tissues often varies 8 9 depending on both genetic and non-genetic factors, such as environmental conditions and plant development, and on the interactions between these (Ghandilyan et al., 2009a; 10 Neugebauer et al., 2018). A lack or excess of a mineral element will limit plant growth, 11 12 and tissue concentrations of plants must be maintained within an appropriate range (Grusak et al., 2016). 13

Phosphorus (P) is an essential macronutrient for plant growth and development 14 (Hawkesford et al., 2012). It is present in membranes, as a component of phospholipids, 15 participates in photosynthesis, energy transduction, and primary and secondary 16 metabolism as a component of metabolites, is essential for gene replication and 17 expression as a component of nucleotides, and serves in intracellular signal transduction 18 via phosphorylation reactions (Hawkesford et al., 2012; Grusak et al., 2016). Lack of 19 available P in soils restricts plant growth, delays development, and reduces crop yields 20 21 (Hawkesford et al., 2012).



1	processes, including mobilization from the soil, uptake by roots, translocation from root
2	to shoot in the xylem and recirculation within the plant via the phloem (White and
3	Broadley, 2009; White, 2012a, b). When the supply of an essential mineral element
4	(nutrient) is compromised this can affect the bioavailability, uptake, transport and
5	utilization of other mineral elements (Watanabe et al., 2015; Neugebauer et al., 2018).
6	Hence, P deficiency can result in changes in the ionomes of plant organs. For example,
7	P deficiency in Arabidopsis results in the inhibition of primary root growth and affects
8	Fe homeostasis through modulation of LPR1 and LPR2 ferroxidases, leading to the
9	apoplastic accumulation of Fe ₃₊ (Ward et al., 2008; Müller et al., 2015; Balzergue et
10	al., 2017; Gutiérrez-Alanís et al., 2018). In Arabidopsis shoots, P deficiency results in
11	increased As, B, Fe, and Zn concentrations and decreased Co and Cu concentrations
12	(Baxter et al., 2008). In Brassica napus, P deficiency results in reduced concentrations
13	of Ca, Fe, Mg, Mn, and Zn in seeds (Ding et al., 2010). An appreciation of the
14	interactions between P nutrition and the accumulation of other mineral elements in
15	plants may help researchers understand the physiological responses of plants to P
16	deficiency better. However, very little is known about the molecular determinants of
17	alterations in the ionome caused by fluctuations in P supply.
18	The uptake and accumulation of mineral elements in roots and shoots are most likely
19	affected by many genetic factors (Huang and Salt, 2016). Quantitative trait locus (QTL)

analysis is a powerful technique to identify chromosomal regions containing genetic
factors linked to variation in complex traits (Paran and Zamir, 2003). A large number

of QTLs have been detected that affect the acquisition and accumulation of mineral

1	elements in plant tissues (e.g. Bentsink et al., 2003; Loudet et al., 2003, 2007; Payne et
2	al., 2004; Vreugdenhil et al., 2004; Harada and Leigh, 2006; Waters and Grusak, 2008;
3	Ghandilyan et al., 2009a, b; Sánchez-Bermejo et al., 2014; Wang et al., 2019) and a
4	number of genes impacting the plant ionome have been cloned using forward genetics
5	in Arabidopsis (reviewed in Huang and Salt, 2016). In Brassica rapa, several QTLs
6	have been discovered that affect seed and leaf phosphate concentrations (Zhao et al.,
7	2008) and leaf Al, Fe, Mg, Mn, Na, P, Sr and Zn concentrations (Wu et al., 2008). In
8	Brassica oleracea, QTLs have been identified for shoot Ca and Mg concentrations
9	(Broadley et al., 2008), shoot P concentration (Hammond et al., 2009), and shoot K and
10	Na concentrations (White et al., 2010). In B. napus, QTLs influencing shoot B, Ca, Cu,
11	Fe, Mg, P and Zn concentrations (Liu et al., 2009), seed Ca, Cu, Fe, Mg, Mn, P and Zn
12	concentrations (Ding et al., 2010) and seed S concentration (Körber et al., 2016), have
13	been identified.
14	In addition to traditional linkage mapping using biparental recombinant populations,
15	genome-wide association analysis (GWAS) can be used to uncover the genetic basis of
16	complex traits, including ionomic traits (e.g. Atwell et al., 2010; Yang et al., 2018).
17	More QTLs with a narrower mapping interval can be detected by GWAS using natural
18	populations that contain extensive genetic diversity and have undergone numerous
19	recombination events (Xiao et al., 2017). Bus et al. (2014) identified several significant
20	single nucleotide polymorphisms (SNPs) associated with shoot Ca, Cu, Mg, Mn, Na, S
21	and Zn concentrations in <i>B. napus</i> through GWAS. Additionally, a number of SNPs and

22 gene expression markers were discovered for leaf nitrate, phosphate and sulfate

concentrations (Koprivova *et al.*, 2014), leaf Ca and Mg concentrations (Alcock *et al.*,
 2017), and leaf nitrate, P and K concentrations (Alcock *et al.*, 2018) through associative
 transcriptomics. Although the identity of the underlying genes remains unknown, these
 studies have demonstrated the presence of allelic variation affecting the accumulation
 of mineral elements in Brassicaceae crops.

Brassica napus (AnAnCnCn, ~1130 Mb, 2n=4x=38) is an allopolyploid crop derived 6 from interspecific crosses between the diploid progenitors, *B. rapa* (ArAr, 2n=2x=20) 7 and B. oleracea (C_0C_0 , 2n=2x=18) (Chalhoub et al., 2014). It is the third largest source 8 9 of vegetable oil globally. However, it is highly susceptible to P deficiency (Duan et al., 2009). To date, few studies have been performed to detect QTLs associated with shoot 10 and root ionomes under the same conditions in any Brassicaceae crop. In the present 11 12 study, a *B. napus* double haploid (*Bna*TNDH) population derived from a cross between cultivars Tapidor and Ningyou 7 was employed (Qiu et al., 2006). The shoot and root 13 ionomic profiles (B, Ca, Cu, Fe, K, Mg, Mn, Na, P, S and Zn) of 182 genotypes from 14 the BnaTNDH population grown in an agar system with an optimal (OP) or a low P 15 supply (LP) were examined. The objectives of the study were (i) to characterize the 16 effects of P supply on the uptake of mineral elements and the ionomes of shoot and root 17 tissues of B. napus, (ii) to identify QTLs and QTL clusters associated with shoot and 18 19 root ionomic traits, as well as those affecting the partitioning of mineral elements between shoot and root tissues, at OP and LP, and (iii) to elucidate differences in the 20 21 genetic control of shoot and root ionomic traits when plants are grown with contrasting P supplies. 22

1 MATERIALS AND METHODS

2 Plant materials and growing conditions

The *Bna*TNDH mapping population employed consisted of 182 lines generated through anther culture of the F1 generation of a cross between a European winter-type cultivar Tapidor containing a low glucosinolate concentration (Sharpe and Lydiate, 2003), and Ningyou 7, a Chinese semiwinter-type cultivar with high glucosinolate concentration (Qiu *et al.*, 2006).

To fine map candidate QTLs, an advanced backcross population consisting of 860 8 9 BC4F1 lines, which were generated with cultivar Tapidor as the recurrent parent and cultivar Ningyou 7 as the donor parent (Zeng, 2011), were employed to confirm and 10 resolve the QTLs identified in the BnaTNDH population. Each BC4F1 line was self-11 12 pollinated to produce the BC4F2 population. Genomic DNA of six BC4F2 individuals selected randomly from each BC₄F₁ line was bulked in an equal ratio to generate bulk 13 DNA, and these six individuals were further self-pollinated to construct the BC4F2:3 14 15 population. These DNA bulks and the genomic DNA of the two parental cultivars were 16 subjected to specific-locus amplified fragment sequencing using an Illumina HiSeq 2500 sequencer with a paired-end pattern. 17

The root traits and biomass traits of the *Bna*TNDH population and its parents had been screened previously in an agar system both at a phosphate (Pi) concentration of 0.625 mM (an optimal phosphorus supply, OP) and at a Pi concentration of 0 mM (a low phosphorus supply, LP) (Shi *et al.*, 2013). In this study, five BC4F2:3 lines (1757-3, 1856-3, 1856-4, 2292-3 and 2303-4) were screened in the agar system at LP, and 1 Tapidor and Ningyou 7 at both OP and LP.

Briefly, surface sterilized seeds were sown into vented polystyrene trays (QTray; 240 2 \times 240 \times 20 mm; Molecular Devices, Hampshire, UK) containing 300 mL 0.8% (w/v) 3 agar and a modified basal salt mix (Murashige and Skoog, 1962) with either K and Pi 4 added as KH2PO4 for OP or with 0.625 mM KCl added to provide K for LP. Seeds were 5 sown 3 cm from the top edge of a tray, with four seeds per line and two lines per tray. 6 Trays were sealed with Nescofilm and placed 10° from vertical in a growth room under 7 a 16-h photoperiod at a constant temperature of 24 °C. Illumination was provided by a 8 9 bank of 84 100-W cool fluorescent tubes (Philips, Eindhoven, Netherlands), giving a photon flux density between 400 and 700 nm of 80–100 µmol photons m-2 s-1 at plant 10 height. For each line, 16 seeds were sown across four independent replicates, at both 11 12 OP and LP. Trays were placed randomly within the growth room.

13

14 *Phenotypic analyses*

Shoots and roots were harvested separately and dried at 80 °C 12 d after sowing. Shoot 15 dry weight (SDW) and root dry weight (RDW) were determined, and total plant dry 16 weight (TDW) was calculated as the sum of SDW and RDW. Prior to analysis, TDW, 17 SDW and RDW were natural logarithm (ln)-transformed to improve the normality and 18 19 variance of the data. To acquire adjusted line means, the REML (residual maximum likelihood) procedure in GenStat (15th Edition, VSN International Ltd, Hemel 20 Hempstead, UK) was performed using the $[(P]_{ext} \times Line)]$ term as a fixed factor and 21 [(Replicate/Run/Plate/Position)] as a random factor. 22

1	The mineral element concentrations in shoot and root dry matter were measured
2	using inductively coupled plasma optical emission spectrometry (ICP-OES) (JY Ultima
3	2; Jobin Yvon Ltd, Stanmore, Middlesex, UK). All the replicate samples for each line
4	were mixed together, oven dried at 80 °C for at least 48 hours and then milled.
5	Approximately 0.1 g dried ground powder was put into a PTFE digestion tube together
6	with 1 ml concentrated nitric acid, closed tightly and processed in a closed vessel acid
7	digestion microwave (MARSXpress; CEM Corporation, Matthews, NC, USA). After
8	cooling, each digest was diluted to a final volume of 10 ml with deionized water. These
9	dilutions were used to measure mineral element concentration by ICP-OES. For each
10	mineral element, shoot content was calculated as the product of shoot concentration and
11	SDW, and root content as the product of root concentration and RDW. Plant mineral
12	element contents were calculated as the sum of shoot mineral element content and root
13	mineral element content. The partitioning of a mineral element to the shoot was
14	calculated as the quotient of shoot mineral element content divided by plant mineral
15	element content (Wu et al., 2015).

17 *QTL mapping and integration of the QTL clusters*

The *Bna*TNDH linkage map used in this study was constructed as previously described (Zhang *et al.*, 2016). This map spanned 2077.9 cM in length and contained 1698 SNP markers and 343 original markers on 19 chromosomes, with an average distance of 1.02 cM between adjacent markers. The additive and epistatic QTLs for different traits at both OP and LP were determined using the QTL IciMapping v4.1 (Meng *et al.*, 2015)

using single environment phenotypic values. Briefly, for the additive QTL, the ICIM-1 ADD mapping method was exploited in the software. The walk speed was 1 cM, and 2 3 the P values for entering variables (PIN) and removing variables (POUT) were set at 0.001 and 0.002, respectively. The epistatic QTLs were identified by the ICIM-EPI 4 mapping method. The walk speed was 5 cM, and PIN and POUT were set at 0.0001 and 5 0.0002, respectively. The LOD thresholds for the additive QTL and epistatic QTL were 6 set to 2.5 and 5.0 as the default manual input value, respectively. The phenotypic 7 variation explained by each additive QTL or epistatic QTL and the corresponding 8 9 additive effects were also estimated using the same software.

A QTL cluster was defined as two or more significant QTLs with overlapping 10 confidence intervals. For a QTL cluster, the coincidence of QTLs for two or more traits 11 12 was considered to be positive if the alleles increasing trait values were from the same parent, while it was considered to be negative if the alleles increasing trait values were 13 from different parents (Coque et al., 2008). QTL meta-analysis was performed using 14 BioMercator v4.2 (Arcade et al., 2004). Meta-analysis computing was based on the 15 position of each input QTL, and on the variance of this position, assessed through 16 confidence interval values. The algorithm developed by Goffinet and Gerber (2000) 17 was employed to perform the analysis. 18

19

20 Confirmation and resolution of the QTL cluster Cl17.1 and prediction of candidate
21 genes

22 The 860 BC₄F₁ substitution lines of *B. napus*, in which the segments of cultivar Ningyou

1	7 were introgressed into the genetic background of cultivar Tapidor, were genotyped
2	with 17116 genome-wide markers (InDels and SNPs) produced from SLAF-seq (data
3	not shown). Nine BC4F1 lines were screened based on the two flanking markers of the
4	QTL cluster Cl17.1. These lines showed more than 90% genetic similarity to the
5	recurrent parent (Tapidor) and did not harbour any other QTLs containing Ningyou 7
6	alleles affecting shoot K, Mg and S concentrations at LP. The BC4F2:3 lines derived from
7	these nine BC4F1 lines were further genotyped using the five InDel markers located in
8	the region of the QTL cluster Cl17.1. Finally, five BC4F2:3 lines (1757-3, 1856-3, 1856-
9	4, 2292-3 and 2303-4) were identified with whole or part homozygous donor segments
10	of the QTL cluster Cl17.1. The five BC4F2:3 lines were grown at LP in the agar system
11	for 12 d, and the shoot K, Mg and S concentrations of these five lines were measured
12	by ICP-OES. The QTL cluster Cl17.1 was confirmed and resolved from the phenotypes
13	and genotypes of the five BC4F2:3 lines. The resolved QTL cluster Cl17.1 was mapped
14	to the reference genome (cultivar Darmor-bzh) based on the physical position of the
15	two flanking markers. The available reference genome of B. napus (Chalhoub et al.,
16	2014) and the functional annotation of the Arabidopsis genome
17	(https://www.arabidopsis.org/) were employed for the prediction of putative candidate
18	genes.
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1 **RESULTS**

2 Variation and correlation of shoot concentrations of eleven mineral elements among

3 *the* Bna*TNDH population at OP and LP*

To establish variation in the ionome, a meta-analysis of shoot and root ionomes in the parental lines Tapidor and Ningyou 7 was conducted. Ionome data from shoots and roots of plants grown with a range of external P concentrations in an agar system obtained from this and previous work (Shi *et al.*, 2013) and of plants grown with low and high B concentrations in a hydroponic system (Liu *et al.*, 2009) were analysed. Across these multiple environments, Tapidor had higher shoot B, Cu and P concentrations, but lower shoot K and S concentrations, than Ningyou 7 (Fig. 1A).

Shoot concentrations of Cu, K, Mg and P were reduced by P starvation in both Tapidor and Ningyou 7, while shoot concentrations of Ca and Mn were only reduced by P starvation in Ningyou 7 (Fig. 2A; Supplementary Data Table S1). By contrast, shoot Fe and S concentrations of both cultivars and the shoot Na concentration of Tapidor were increased by P starvation.

The mean shoot B, Fe, Na, S and Zn concentrations of the *Bna*TNDH population were greater at LP than at OP, but the mean shoot Ca, Cu, K, Mg and P concentrations were lower at LP than at OP (Fig. 3; Supplementary Data Table S1). Shoot Ca and Mn concentrations of the *Bna*TNDH population had the highest correlation coefficients among the eleven mineral elements studied at both OP (r = 0.75) and at LP (r = 0.84; Table 1). Among the genotypes of the *Bna*TNDH population there were significant positive correlations in shoot concentrations between OP and LP for all eleven mineral

- 1 elements studied, except for Zn (Table 1).
- 2

3 *Variation and correlation of root concentrations of eleven mineral elements among the*

4 BnaTNDH population at OP and LP

5 Tapidor had significantly higher root K and Mg concentrations, but a lower root Na 6 concentration than Ningyou 7 across the multiple growing environments (Fig. 1B). The 7 root P concentration in both Tapidor and Ningyou 7 was significantly reduced by P 8 starvation (Fig. 2B; Supplementary Data Table S1). Root B, Fe, K, Mn and Na 9 concentrations were reduced only in cultivar Tapidor and root Mg concentration was 10 reduced only in cultivar Ningyou 7 by P starvation. By contrast, root Cu and S 11 concentrations of both cultivars were increased by P starvation.

12 The mean root Cu and S concentrations of the BnaTNDH population were greater at LP than at OP (Fig. 3; Supplementary Data Table S1). By contrast, root B, K, Mg, Mn, 13 Na and P concentrations were lower at LP than at OP. Root Ca and Mn concentrations 14 15 had the highest correlation coefficients among the eleven mineral elements at both OP (r = 0.70) and LP (r = 0.76; Table 1). There were significant positive correlations in root 16 concentrations between OP and LP for all eleven mineral elements studied in the 17 BnaTNDH population, except for Cu, P and Zn (Table 1). Generally, stronger 18 correlations were detected between shoot concentrations of mineral elements and 19 between root concentrations of mineral elements at LP than at OP (Table 1). 20

21

22 Difference in concentrations of eleven mineral elements between shoot and root among

1 *the* Bna*TNDH* population at OP and LP

Shoot mineral element concentrations of the BnaTNDH population clustered separately 2 from their root mineral element concentrations in a PCA (principal component analysis) 3 biplot at both OP and LP (Fig. 4). The first component accounted for 58.3% and 60.3% 4 of the total variation at OP and LP, respectively, and mainly represented the contrast 5 6 between B+Ca+Mg+Mn+Na+P and Fe at both OP and LP. Interestingly, Cu was loaded positively onto PC1 at OP, but negatively onto PC1 at LP. The second component, 7 accounting for 10.6% and 14.7% of the total variation at OP and LP, respectively, 8 9 mainly represented K and Zn at both OP and LP. In addition, S was loaded in nearly equal proportion between PC1 and PC2 at both OP and LP. 10

11

Variation and correlation of shoot and root contents of eleven mineral elements among
the BnaTNDH population at OP and LP

14 Tapidor had lower shoot contents than Ningyou 7 of all eleven mineral elements studied

at both OP and LP, except for Cu at LP and Fe at OP (Supplementary Data Table S2).

16 The mean shoot Fe content of the *Bna*TNDH population was higher at LP than at OP

17 (Fig. 3; Supplementary Data Table S2). By contrast, the mean shoot Ca, Cu, K, Mg, Mn

and P contents of the *Bna*TNDH population were lower at LP than at OP.

Tapidor had lower root contents of all the eleven mineral elements studied than Ningyou 7 at both OP and LP, except for Cu at LP (Supplementary Data Table S2). The mean root Cu content in the *Bna*TNDH population was higher at LP than at OP, while the mean root B, Ca, Fe, K, Mg, Mn, Na, P and Zn contents were lower at LP than at OP (Fig. 3; Supplementary Data Table S2). Significant correlations were observed

1	among shoot and root contents of all the eleven mineral elements in the BnaTNDH
2	population at both OP and LP (Supplementary Data Table S3). Significant positive
3	correlations were also observed in shoot and root contents of all the eleven mineral
4	elements between OP and LP (Supplementary Data Table S3).
5	
6	Variation and correlations of whole plant mineral element contents among the
7	BnaTNDH population at OP and LP
8	Tapidor had lower plant contents of all mineral elements than Ningyou 7 at both OP
9	and LP, except for Cu at LP (Supplementary Data Table S4). The mean plant Fe content
10	of the <i>Bna</i> TNDH population was greater at LP than at OP, while the mean plant B, Ca,
11	Cu, K, Mg, Mn, Na, P and Zn contents were lower at LP than at OP (Fig. 3;
12	Supplementary Data Table S4).
13	Significant positive correlations among the plant contents of all eleven mineral
14	elements were observed at both OP and LP in the BnaTNDH population, with the lowest
15	significant correlation coefficient ($r = 0.28$) between plant Fe and P content at LP and
16	the highest correlation coefficient ($r = 0.95$) between plant Ca and Mn content at both
17	OP and LP (Table 2). The plant contents of all the eleven mineral elements studied were
18	significantly positively correlated between OP and LP in the BnaTNDH population

(Table 2).

21 Variation and correlations in partitioning of eleven mineral elements to the shoot

among the Bna*TNDH population at OP and LP*

1	Tapidor and Ningyou 7 differed less than 5% in their partitioning of individual mineral
2	elements to the shoot at OP and LP, except for Fe (11.1% at OP and 23.6% at LP)
3	(Supplementary Data Table S4). The mean partitioning to the shoot of all eleven mineral
4	elements studied in the <i>Bna</i> TNDH population was greater than 80% at both OP and LP,
5	with the exception of Cu at LP (66.7%) and Fe at OP (19.9%) and LP (36.6%). The
6	partitioning of B, Fe, Na and Zn to the shoot in the BnaTNDH population was greater
7	at LP than at OP (Fig. 3; Supplementary Data Table S4). In contrast, partitioning of Ca,
8	Cu, K and P to the shoot was less at LP than at OP.
9	There were significant positive correlations in the partitioning of all the eleven
10	mineral elements studied at both OP and LP, except between Cu and Ca, Fe, K, Mg, Mn
11	and Na at OP (Table 2). The partitioning of Ca to the shoot was highly correlated with
12	the partitioning of Mn to the shoot in the <i>Bna</i> TNDH population at both OP ($r = 0.86$)
13	and LP ($r = 0.87$), suggesting that the distribution of these two mineral elements within
14	the plant might be controlled by similar transport processes. It is noteworthy that, for
15	example, neither Ca nor Mn are readily mobile in the phloem (White, 2012b; Grusak et
16	al., 2016). The partitioning to the shoot of all eleven mineral elements studied in the
17	BnaTNDH population were significantly positively correlated between OP and LP
18	(Table 2). However, Cu and Fe partitioning to the shoot at OP had a relatively weak
19	correlation with their partitioning at LP, suggesting that there might be a difference in
20	the control of the partitioning of these two mineral elements to the shoot between OP
21	and LP.

1 QTLs and epistatic interactions for mineral element concentrations, contents, and

2 partitioning to the shoot at OP and LP

Approximately normal distributions and transgressive segregations were observed for 3 all the ionomic traits studied in the BnaTNDH population at both OP and LP 4 (Supplementary Data Tables S1, S2 and S4; Supplementary Data Figs. S1-S6), 5 indicating a quantitative inheritance pattern suitable for QTL identification. A total of 6 41 QTLs distributed across twelve chromosomes were associated with shoot 7 concentrations of the eleven mineral elements at OP and LP, explaining 1.4-19.3% of 8 9 the phenotypic variation (Table 3). Among these QTLs, CaconcLPS-A07 for shoot Ca concentration at LP, KconcOPS-C08a for shoot K concentration at OP, and SconcOPS-10 A09 and SconcOPS-C07 for shoot S concentration at OP accounted for 18.0%, 17.1%, 11 12 19.3% and 19.3% of the phenotypic variation for these traits, respectively. For both shoot Ca and Mn concentrations, the associated QTLs identified at OP and LP were 13 closely linked on chromosome A07. 14

A total of 34 QTLs for root concentrations of the eleven mineral elements at OP and LP explained 4.6–23.1% of the phenotypic variation, and were located on 15 of the 19 chromosomes of *B. napus* (Table 3). Among these QTLs, SconcOPR-A04 and SconcOPR-C01 for root S concentration at OP and SconcLPR-A04a for root S concentration at LP accounted for 19.2%, 20.5% and 23.1% of the phenotypic variation, respectively. In addition, the QTLs for the shoot and root S concentrations were colocated on chromosome A09 at OP.



1	eleven mineral elements at OP and LP, which accounted for 0.7-23.0% of the
2	phenotypic variation (Table 3). Among these QTLs, CucontOPS-A03 for shoot Cu
3	contents and PcontOPS-A02 for shoot P contents at OP explained 16.6% and 23.0% of
4	the phenotypic variation, respectively. There was a close linkage relationship between
5	the QTLs for the shoot contents of five mineral elements (B, Ca, Mn, Na and Zn)
6	identified at OP and at LP on chromosome A03, which may be due to the close linkage
7	relationship between the QTLs identified for SDW at OP and at LP (Table 3). A close
8	linkage relationship was also observed between the QTLs detected for the shoot
9	contents of four mineral elements (Ca, K, Mg and S) at OP and LP on chromosome A04.
10	In addition, a stable QTL affecting shoot K content was detected on chromosome A03
11	at OP and LP.
12	Twenty-nine QTLs across ten chromosomes for root contents of the eleven mineral
13	elements at OP and LP explained 4.9-14.5% of the phenotypic variation (Table 3).
14	Moreover, the QTLs for the shoot and root contents of five mineral elements (Ca, K,
15	Mn, Na and Zn) were co-located on chromosome A03 at LP, and QTLs for the shoot
16	and root S contents were co-located on chromosome C04 at OP.
17	A total of 46 QTLs distributed across eight chromosomes were associated with plant
18	contents of the eleven mineral elements at OP and LP, explaining 5.1-16.0% of the
19	phenotypic variation (Table 3). One QTL, CucontOPP-A03 for plant Cu content at OP,
20	accounted for 16.0% of the phenotypic variation. The QTLs for the plant contents of B,
21	Ca, K, Mn, Na and S identified at OP and at LP had close linkage relationships.
22	A total of 54 significant QTLs, including 33 at OP and 21 at LP, were detected for

partitioning of the eleven mineral elements to the shoot (Table 3). These QTLs were
located on 15 chromosomes, explaining 4.0–24.9% of the phenotypic variation. The
QTLs for the partitioning of Mg to the shoot at OP and LP and those for the partitioning
of S to the shoot at OP and LP were closely linked on chromosomes C04 and A04,
respectively. Moreover, a QTL for the partitioning of Ca to the shoot on chromosome
C09 was identified at both OP and LP (Table 3).

A total of 54 epistatic interactions, including 31 at OP and 23 at LP, were detected 7 for shoot and root concentrations, shoot, root and plant contents, and partitioning to the 8 9 shoot of the eleven mineral elements in the BnaTNDH population (Fig. 5; Table 4). There were 25 epistatic interactions in the A genome (A01–A10), 13 in the C genome 10 (C01-C09), and 16 between the two genomes of B. napus (Table 4). The individual 11 12 phenotypic contributions of these epistatic interactions for different traits ranged from 6.8% to 24.4%, and three pairs of them explained more than 20.0% of the phenotypic 13 variation. None of these epistatic interactions involved any additive QTL except for one 14 pair showing a QTL/non-QTL interaction for root Mn concentration at OP 15 (Supplementary Data Table S5). 16

A pleiotropic epistatic interaction affecting root B, K and Mn contents at OP was identified on chromosome A10, and another one affecting root B, K, Mn and Zn contents at OP was found on chromosome C09 (Fig. 5A). A pleiotropic epistatic interaction for shoot K content and plant K and Na contents at LP was detected between chromosome A08 and C05, and another one for root B content and plant Mn content at LP was discovered between chromosome A03 and A08 (Fig. 5B). Although an epistatic interaction was detected for TDW at both OP and LP, these two pairs did not overlap
with any of the previously identified 54 epistatic interactions (Fig. 5), implying that
these 54 epistatic interactions may affect the uptake and transport of different mineral
elements. All these epistatic interactions for the same trait were not detected
consistently across OP and LP (Table 4).

The number of additive QTLs and epistatic interactions varied from zero to five and 6 from zero to four, respectively (Supplementary Data Table S5). The additive QTLs and 7 epistatic interactions accounted for 0-51.9% and 0-48.7% of the total phenotypic 8 9 variation, respectively. The total phenotypic variation explained by the additive QTL was more than 50% for shoot and root S concentrations at OP, which could provide 10 targets for *B. napus* breeding programmes. The presence of additive QTLs and epistatic 11 12 interactions with positive and negative effects could provide the genetic basis for the transgressive segregation of the traits studied. 13

14

QTL clusters for shoot and root concentrations, shoot, root and plant contents, and partitioning to the shoot of eleven mineral elements at OP and LP

A total of 49 QTL clusters were identified across twelve chromosomes in a metaanalysis (Fig. 6; Supplementary Data Table S6). There were 17, 14 and 18 QTL clusters detected at OP, LP, and both OP and LP, respectively. Six QTL clusters (Cl2.3, Cl3.3, Cl3.4, Cl4.2, Cl4.3 and Cl19.2) were associated with more than three traits impacting mineral element composition (Supplementary Data Table S6). Among these QTL clusters, four QTL clusters (Cl2.3, Cl3.3, Cl3.4 and Cl4.2) overlapped with QTLs for

1	biomass traits, while two QTL clusters Cl4.3 and Cl19.2 were not associated with
2	biomass traits. Cl4.3 was associated with shoot B, Ca, K, Mg, Mn, Na, S and Zn
3	contents, plant Ca, K, Mg, Mn, Na and S contents, and the partitioning of B to the shoot
4	at LP. The alleles with positive effects in this QTL cluster were contributed by Ningyou
5	7. Cl19.2 was associated with root Zn content and plant Ca, Mn and Na contents at OP.
6	The alleles with positive effects in this QTL cluster were contributed by Tapidor.

8 Confirmation and refinement of the QTL cluster Cl17.1 associated with shoot K, Mg,
9 and S concentrations at LP

Given the consistent difference in the shoot K and S concentrations between Tapidor 10 and Ningyou 7 across multiple environments (Fig. 1A), the QTL cluster Cl17.1 may be 11 12 a robust locus. The presence of the QTL cluster Cl17.1 was confirmed and resolved further using substitution lines. Shoot K, Mg and S concentrations of the five BC4F2:3 13 lines (1757-3, 1856-3, 1856-4, 2292-3 and 2303-4) and the two parental lines were 14 15 determined at LP in the agar system (Fig. 7). There was no significant difference in shoot K concentration between Tapidor and any of the five BC4F2:3 lines, although an 16 obvious difference in this trait was observed between the cultivars Tapidor and Ningyou 17 7 (Fig. 7A). This suggests that the QTL cluster Cl17.1 had only a minor effect on shoot 18 K concentration, in line with the small fraction of phenotypic variation (3.6%) 19 explained by the QTL KconcLPS-C07 (Table 3). In contrast, all five BC4F2:3 lines had 20 significantly higher shoot Mg concentrations than Tapidor, although there was no 21 significant difference in this trait between the cultivars Tapidor and Ningyou 7 (Fig. 22

7B). In addition, four of the five BC4F2:3 lines had significantly higher shoot S 1 concentrations than Tapidor, the exception being line 2303-4 (Fig. 7C). The allele from 2 Ningyou 7 within the QTL cluster Cl17.1 had a positive effect on both shoot Mg and S 3 concentrations in all five BC4F2:3 lines. Thus, it was consistent with the positive 4 contribution of the Ningyou 7 allele to the trait value in the BnaTNDH population 5 (Table 3). The QTL cluster Cl17.1 was narrowed down to 28.5-30.6 Mb on 6 chromosome C07 using knowledge of the introgression regions of these five lines (Fig. 7 7D). 8

9

10 **DISCUSSION**

The uptake and partitioning to the shoot of eleven mineral elements and their QTLs
were differentially influenced by P starvation in B. napus

The application of high-throughput elemental analysis has facilitated the quantitative 13 and simultaneous measurement of the elemental composition of living organisms (Salt 14 et al., 2008). In this study, the biomass and concentrations of eleven mineral elements 15 (B, Ca, Cu, Fe, K, Mg, Mn, Na, P, S and Zn) in the shoot and root were investigated in 16 the *Bna*TNDH population grown at OP and LP (Figs. 1–3). Direct interactions between 17 cations and anions in their uptake are rare since they occur through different 18 transporters, but the uptake of one mineral element can affect the uptake of another 19 indirectly through effects on the membrane potential, the proton electrochemical 20 21 gradient or via feedback regulation through plant growth, metabolism or cellular homeostasis (White, 2012a). Thus, the decrease in the plant content of Ca, Cu, K, Mg, 22

1	Mn, Na and Zn in P-deficient B. napus (Fig. 3; Supplementary Data Table S4) might be
2	a consequence of reduced growth, as was reported previously in B. napus grown in a
3	hydroponics system (Maillard et al., 2016). Traits affecting root morphology and
4	anatomy play a key role in the acquisition of mineral elements by plants (White et al.,
5	2013) and a significant positive correlation between leaf Ca (and Zn) concentrations
6	and lateral root density (LRD) was observed in field trials with <i>B. napus</i> (Thomas <i>et al.</i> ,
7	2016a, b). In the experiments reported here, the concentrations of mineral elements in
8	the shoot had no correlations, or significant negative correlations, with LRD, but shoot
9	B, Ca, Cu, Fe, K, Mg, Mn, Na, P and Zn concentrations had significant positive
10	correlations with PRL and/or LRL at OP and LP (Supplementary Data Table S7),
11	suggesting that greater root length might improve nutrient acquisition in an agar system
12	with a homogeneous nutrient availability. By contrast, the shoot Fe concentration was
13	greater in plants at LP than at OP (Fig. 3; Supplementary Data Table S4), possibly
14	because P and Fe can precipitate together when Pi concentrations are high, which results
15	in reduced Fe availability (Dalton et al., 1983; Ward et al., 2008).

In addition to the uptake of mineral elements by roots, the translocation of mineral elements from the root to the shoot in the xylem and their recirculation within the plant via the phloem affect the accumulation of mineral elements in the shoot. The mean partitioning of all eleven mineral elements to the shoot was more than 80% in the *Bna*TNDH population grown at OP, except for Fe (Supplementary Data Table S4), suggesting that mineral nutrients were preferentially partitioned to the shoot to maintain plant growth and development. The rather low partitioning of Fe to the shoot (19.9%)

1	led to a relatively low mean shoot Fe concentration (92.7 μ g g-1 DW) and a relatively
2	high mean root Fe concentration (2013 μ g g-1 DW) in the <i>Bna</i> TNDH population at OP
3	(Supplementary Data Table S1). Similarly, mean rosette Fe concentration (39 μ g g-1
4	DW) was much lower than mean root Fe concentration (1680 μg g-1 DW) in an
5	Arabidopsis RIL population (Ghandilyan et al., 2009b), and root Fe concentrations
6	were greater than shoot Fe concentrations in B. oleracea genotypes, which was
7	exacerbated by greater P supply (Pongrac et al., 2020). It was also found that Fe
8	concentrations in the roots of many Brazilian tree species were greater than their shoot
9	Fe concentrations when grown hydroponically in a complete nutrient solution
10	(Neugebauer et al., 2019). It may be argued that the large amounts of Fe stored in roots
11	are necessary to sustain the normal growth of roots, serve as a Fe reserve for periods of
12	reduced Fe availability, or protect the shoot from Fe toxicity.
13	The partitioning to the shoot of each of the eleven mineral elements was affected
14	differently by P starvation in B. napus (Fig. 3; Supplementary Data Table S4). The
15	partitioning of Cu and Fe to the shoot were most strongly influenced by P starvation.
16	The partitioning of Cu to the shoot decreased by 23% and the partitioning of Fe to the
17	shoot increased by 84% at LP (Supplementary Data Table S4). Iron is transported
18	mainly in the form of Fe ₃₊ citrate in the xylem (Welch, 1995; von Wirén et al., 1999).
19	In Arabidopsis, AtFRD3, a member of the multidrug and toxin efflux (MATE)

20 transporter family, is expressed in the root pericycle and appears to be involved in

loading citrate into the xylem (Durrett *et al.*, 2007; Puig *et al.*, 2007). *BnaA05g29700D*

and *BnaC05g44030D* are the homologous genes of *AtFRD3* in *B. napus*. The expression

1	of both these genes was significantly induced in roots of P-deficient plants (Li et al.,
2	2019), which might result in greater Fe translocation from roots to shoots and account
3	for an increased Fe partitioning to the shoot in <i>B. napus</i> at LP. The Arabidopsis P-type
4	ATPase HMA5 (AtHMA5) is involved in loading Cu into the xylem for root-to-shoot
5	translocation and/or Cu detoxification in the root (Andrés-Colás et al., 2006; Kobayashi
6	et al., 2008). The expression of BnaA10g06240D (a homologous gene of AtHMA5 in B.
7	napus) was reduced in roots of B. napus plants at LP (Li et al., 2019), which might
8	account for a significantly reduced Cu partitioning to the shoot at LP.
9	A total of 133 and 123 QTLs for shoot and root ionomic traits were identified at OP
10	and LP, respectively. For each ionomic trait, most of the QTLs identified at OP differed
11	from the QTLs detected at LP in the present study. Similar observations were made by
12	Ding et al. (2010) that the QTLs affecting seed mineral element concentrations in B.
13	napus in P-deficient plants differed from those affecting seed mineral element
14	concentrations in P-replete plants. Moreover, different QTLs affected shoot mineral
15	element concentrations in B-deficient and B-replete B. napus plants (Liu et al., 2009).
16	Given the large number of solute-specific and non-specific transport proteins in plants
17	(Mäser et al., 2001) and the sophisticated regulation of their activities in response to
18	plant nutritional status, it is perhaps unsurprising that the ionomes of plant organs and
19	QTLs affecting the uptake and partitioning of mineral elements between organs should
20	differ at OP and LP.

Several QTLs for shoot Ca, K and Mg concentrations detected at OP in this study
were mapped to the same chromosome of *B. napus* on which a number of SNPs

1	associated with shoot Ca, K and Mg concentrations were also identified by associative
2	transcriptomics (Alcock et al., 2017, 2018). However, it is difficult to determine
3	whether these QTLs for the same trait were located on the same chromosomal regions
4	because the physical positions of the SNPs detected by Alcock et al (2017, 2018) are
5	not available. Further studies should be performed to confirm the stability of these
6	QTLs across different populations and/or environments.

8 Significant differences in the concentrations of mineral element in shoots and roots and
9 their genetic control

Mean shoot B, Ca, Mg, Mn, Na and P concentrations in the *Bna*TNDH population were 10 greater than those in the root regardless of P supply (Fig. 4; Supplementary Data Table 11 12 S1). By contrast, root Fe, S and Zn concentrations were greater than those in the shoot at both OP and LP. These observations indicate that mineral element composition is 13 organ specific in plants. In Arabidopsis, higher shoot B, Mg, Mn and Na concentrations, 14 15 but lower shoot Fe, S and Zn concentrations were also observed at both OP and LP (Ghandilyan et al., 2009b; Gruber et al., 2013). The variation in concentrations of 16 mineral elements among different organs might be associated with the specific 17 biological functions of these organs. For example, relatively high Mg and Mn 18 concentrations in the shoot might be important for photosynthetic efficiency (Black et 19 al., 2006; Kering et al., 2009). 20

In general, the concentrations of mineral elements in shoots had relatively weak correlations with those in the root (Supplementary Data Table S8), as was observed previously in *B. napus* (Thomas *et al.*, 2016*a*). QTLs associated with shoot and root
concentrations of most mineral elements did not overlap although co-located QTLs for
shoot and root S concentrations were identified on chromosome A09 at OP. Similarly,
a striking difference in QTLs associated with the concentrations of mineral elements in
different plant organs (root, rosette and seed) was found in Arabidopsis (Ghandilyan *et al.*, 2009*b*).

7

8 *QTLs for ionomic traits for different mineral elements mapped to the same locus*

9 Significant positive correlations were observed among many ionomic traits, such as shoot and root concentrations, shoot, root and plant contents, and partitioning to the 10 shoot, of the eleven mineral elements in the *Bna*TNDH population across P treatments 11 12 (Table 1; Table 2; Supplementary Data Table S3). Correlations among traits for mineral elements could be the result of element-element interactions, but could also result from 13 genetic linkage of the QTLs controlling these traits (Fig. 6; Supplementary Data Table 14 S6). For example, shoot and root concentrations, shoot, root and plant contents, and 15 partitioning of Ca to the shoot were highly correlated with those traits for Mn at both 16 OP and LP (Table 1; Table 2; Supplementary Data Table S3), and QTLs associated with 17 most of these traits for these two mineral elements were co-located (Supplementary 18 Data Table S6). Co-localization of QTLs associated with ionomic traits of different 19 mineral elements were reported for shoot Ca/Mg in B. oleracea (Broadley et al., 2008), 20 21 shoot Mg/Sr in B. rapa (Wu et al., 2008), rosette K/Mg/Zn in Arabidopsis (Ghandilyan et al., 2009a), shoot B/Cu, B/P and Ca/Mg (Liu et al., 2009), seed Ca/Mg and 22

Cu/Fe/Mn/Zn (Ding et al., 2010), and shoot Cu/Mn/Zn (Bus et al., 2014) in B. napus. 1 Together, these findings reflect the observation that some mineral elements share 2 3 common uptake and transport pathways (White et al., 2012a, b), especially those mineral elements with chemical similarities, such as Ca and Mg, or Ca and Mn observed 4 in this study. Mei et al. (2007) found that expression of an activated Arabidopsis 5 Ca2+/H+ antiporter CAX1 variant that increased Ca accumulation also increased 6 7 concentrations of other mineral elements, such as Mg and Mn, in the root of tobacco. It is anticipated that studies of the genetic basis of shoot mineral element composition 8 9 will contribute to improvements in the nutritional content of leafy vegetables, such as B. rapa and B. oleracea (White and Broadley, 2009). In this study, ten QTL clusters 10 were identified that affected shoot concentrations and/or contents of more than two 11 12 mineral elements but did not affect SDW (Fig. 6; Supplementary Data Table S6). Among these QTL clusters, the QTL cluster Cl17.1 on chromosome C07 had a 13 significant effect on both shoot Mg and S concentrations at LP (Fig. 7). Glucosinolates 14 are a group of sulfur-rich secondary metabolites that are abundant in Brassicaceae. An 15 obvious phenotypic segregation is observed for total glucosinolate concentration and 16 the majority of the individual glucosinolates in seeds and leaves in the BnaTNDH 17 population (Feng et al., 2012). Sulfur concentration is tightly positive correlated with 18 glucosinolate concentrations in seed (Körber et al., 2016) and the QTL cluster Cl17.1 19 was co-located with QTLs affecting seed concentrations of three different 20 21 glucosinolates (4-methylsulfinylbutyl glucosinolate, 2-hydroxy-4-pentenyl glucosinolate and 3-indolyl-methyl glucosinolate) previously identified in the 22

BnaTNDH population (Feng et al., 2012). This major QTL cluster was further narrowed
down to a 28.5–30.6 Mb region in which 236 annotated genes were located
(Supplementary Data Table S9). There was a promising candidate gene,
BnaC07g22430D (homologous to At2G03620, magnesium transporter 3), for Mg
transport in this region but no obvious candidate genes for S transport. Further research
should be conducted to investigate if the pleiotropic effect of this locus is conferred by
one gene or two closely linked genes.

8

9 Epistatic interactions for the ionomic traits

A total of 54 epistatic interactions were identified for various ionomic traits in the 10 BnaTNDH population at OP and LP (Table 4), but most of these epistatic interactions 11 12 did not involve any additive QTL (Supplementary Data Table S5). A number of epistatic interactions were also found for different ionomic traits in Arabidopsis (Ghandilyan et 13 al., 2009b) and B. napus (Liu et al., 2009). The large number of epistatic interactions 14 15 discovered in this study suggest a complex genetic network controlling the *B. napus* ionome at both OP and LP. The epistatic interactions could account for 0-48.7% of the 16 phenotypic variation for different ionomic traits, implying that epistasis is a major 17 genetic component for some ionomic traits. The additive QTLs could explain between 18 19 0-51.9% of the phenotypic variation for different ionomic traits, suggesting that it might be feasible to improve ionomic traits genetically. 20

21

1 CONCLUSIONS

The reductions in plant Ca, Cu, K, Mg, Mn, Na and Zn contents in P-deficient B. napus are likely to be a consequence of reduced growth. The Fe concentration was higher in plants at LP than at OP, possibly because P and Fe can precipitate together when Pi concentrations are high, which results in reduced Fe availability. Significant positive correlations were observed among many ionomic traits across P treatments, which could be the result of element-element interactions, but also could result from the genetic linkage of QTLs controlling traits for different elements. Six QTL clusters were associated with more than three traits impacting mineral element composition, suggesting that some mineral elements share common uptake and transport pathways. Near-isogenic lines should be developed to allow finer mapping of the quantitative genes underpinning the major QTLs identified in this study. This will contribute to a greater understanding of processes affecting the uptake and partitioning of mineral elements in *B. napus*.

1 SUPPLEMENTARY DATA

Figure S1: frequency distribution of shoot concentrations of eleven mineral elements in 2 the BnaTNDH mapping population grown at an optimal (OP) and a low P supply (LP). 3 Figure S2: frequency distribution of root concentrations of eleven mineral elements in 4 the BnaTNDH mapping population grown at an optimal (OP) and a low P supply (LP). 5 Figure S3: frequency distribution of shoot contents of eleven mineral elements in the 6 BnaTNDH mapping population grown at an optimal (OP) and a low P supply (LP). 7 Figure S4: frequency distribution of root contents of eleven mineral elements in the 8 9 BnaTNDH mapping population grown at an optimal (OP) and a low P supply (LP). Figure S5: frequency distribution of plant contents of eleven mineral elements in the 10 BnaTNDH mapping population grown at an optimal (OP) and a low P supply (LP). 11 Figure S6: frequency distribution of partitioning to the shoot of eleven mineral elements 12 in the BnaTNDH mapping population grown at an optimal (OP) and a low P supply 13 (LP). Table S1: shoot and root concentrations of eleven mineral elements in the 14 15 BnaTNDH lines and their parents at an optimal (OP) and a low P supply (LP). Table S2: shoot and root contents (ug plant-1) of eleven mineral elements in the *Bna*TNDH 16 17 lines and their parents at an optimal (OP) and a low P supply (LP). Table S3: Pearson's correlation coefficients among shoot contents and among root contents of eleven 18 mineral elements in the BnaTNDH mapping population at an optimal (upper right 19 triangle) and a low P supply (lower left triangle). Table S4: plant contents (µg plant-1) 20 and partitioning to the shoot (%) of eleven mineral elements in the BnaTNDH lines and 21 their parents at an optimal (OP) and a low P supply (LP). Table S5: the number and 22

explained phenotypic variation of the additive QTL and epistatic QTL for shoot and 1 root concentrations, shoot, root and plant contents, and partitioning to the shoot of 2 eleven mineral elements detected in the BnaTNDH population at an optimal (OP) and 3 a low P supply (LP). Table S6: meta-analysis of QTL clusters for shoot and root 4 concentrations, shoot, root and plant contents, and partitioning to the shoot of eleven 5 mineral elements, shoot dry weight, root dry weight and total dry weight in the 6 BnaTNDH population at an optimal and a low P supply. Table S7: Pearson's correlation 7 coefficients between root traits and shoot concentrations of eleven mineral elements in 8 9 the BnaTNDH mapping population at an optimal (OP) and a low P supply (LP). Table S8: Pearson's correlation coefficients between shoot and root concentrations of eleven 10 mineral elements in the *Bna*TNDH mapping population at an optimal (OP) and a low P 11 12 supply (LP). Table S9: annotated genes underlying the QTL cluster Cl17.1 in Brassica napus. 13

14

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22

LITERATURE CITED

2	Alcock TD, Havlickova L, He Z, et al. 2017. Identification of candidate genes for
3	calcium and magnesium accumulation in Brassica napus L. by association
4	genetics. Frontiers in Plant Science 8: 1968.
5	Alcock TD, Havlickova L, He Z, et al. 2018. Species-wide variation in shoot nitrate
6	concentration, and genetic loci controlling nitrate, phosphorus and potassium
7	accumulation in Brassica napus L. Frontiers in Plant Science 9: 1487.
8	Andrés-Colás N, Sancenón V, Rodríguez-Navarro S, et al. 2006. The Arabidopsis
9	heavy metal P-type ATPase HMA5 interacts with metallochaperones and functions
10	in copper detoxification of roots. The Plant Journal 45: 225-236.
11	Arcade A, Labourdette A, Falque M, et al. 2004. BioMercator: integrating genetic
12	maps and QTL towards discovery of candidate genes. Bioinformatics 20: 2324-
13	2326.
14	Atwell S, Huang YS, Vilhjálmsson BJ, et al. 2010. Genome-wide association study
15	of 107 phenotypes in Arabidopsis thaliana inbred lines. Nature 465: 627-631.
16	Balzergue C, Dartevelle T, Godon C, et al. 2017. Low phosphate activates STOP1-
17	ALMT1 to rapidly inhibit root cell elongation. Nature Communications 8: 15300.
18	Baxter I. 2009. Ionomics: studying the social network of mineral nutrients. Current
19	Opinion in Plant Biology 12: 381–386.
20	Baxter IR, Vitek O, Lahner B, et al. 2008. The leaf ionome as a multivariable system
21	to detect a plant's physiological status. Proceedings of the National Academy of
22	Sciences, USA 105: 12081–12086.
1	Bentsink L, Yuan K, Koornneef M, Vreugdenhil D. 2003. The genetics of phytate
----	---
2	and phosphate accumulation in seeds and leaves of Arabidopsis thaliana, using
3	natural variation. Theoretical and Applied Genetics 106: 1234–1243.
4	Black JR, Yin Q, Casey WH. 2006. An experimental study of magnesium-isotope
5	fractionation in chlorophyll-a photosynthesis. Geochimica et Cosmochimica Acta
6	70: 4072–4079.
7	Broadley MR, Hammond JP, King GJ, et al. 2008. Shoot calcium and magnesium
8	concentrations differ between subtaxa, are highly heritable, and associate with
9	potentially pleiotropic loci in Brassica oleracea. Plant Physiology 146: 1707-
10	1720.
11	Bus A, Körber N, Parkin IA, et al. 2014. Species- and genome-wide dissection of the
12	shoot ionome in Brassica napus and its relationship to seedling
13	development. Frontiers in Plant Science 5: 485.
14	Chalhoub B, Denoeud F, Liu S, et al. 2014. Early allopolyploid evolution in the post-
15	Neolithic Brassica napus oilseed genome. Science 345: 950–953.
16	Coque M, Martin A, Veyrieras JB, Hirel B, Gallais A. 2008. Genetic variation for N-
17	remobilization and postsilking N-uptake in a set of maize recombinant inbred lines.
18	3. QTL detection and coincidences. Theoretical and Applied Genetics 117: 729-
19	747.
20	Dalton CC, Iqbal K, Turner DA. 1983. Iron phosphate precipitation in Murashige and
21	Skoog media. Physiologia Plantarum 57: 472–476.
22	Ding G, Yang M, Hu Y, et al. 2010. Quantitative trait loci affecting seed mineral

1	concentrations in Brassica napus grown with contrasting phosphorus							
2	supplies. Annals of Botany 105: 1221–1234.							
3	Duan HY, Shi L, Ye XS, Wang YH, Xu FS. 2009. Identification of phosphorous							
4	efficient germplasm in oilseed rape. Journal of Plant Nutrition 32: 1148–1163.							
5	Durrett TP, Gassmann W, Rogers EE. 2007. The FRD3-mediated efflux of citrate							
6	into the root vasculature is necessary for efficient iron translocation. Plant							
7	<i>Physiology</i> 144: 197–205.							
8	Feng J, Long Y, Shi L, Shi J, Barker G, Meng J. 2012. Characterization of							
9	metabolite quantitative trait loci and metabolic networks that control glucosinolate							
10	concentration in the seeds and leaves of Brassica napus. New Phytologist 193: 96-							
11	108.							
12	Ghandilyan A, Barboza L, Tisné S, et al. 2009a. Genetic analysis identifies							
12 13	Ghandilyan A, Barboza L, Tisné S, et al. 2009a. Genetic analysis identifies quantitative trait loci controlling rosette mineral concentrations in Arabidopsis							
12 13 14	Ghandilyan A, Barboza L, Tisné S, et al. 2009a. Genetic analysis identifies quantitative trait loci controlling rosette mineral concentrations in Arabidopsis thaliana under drought. New Phytologist 184: 180–192.							
12 13 14 15	 Ghandilyan A, Barboza L, Tisné S, et al. 2009a. Genetic analysis identifies quantitative trait loci controlling rosette mineral concentrations in Arabidopsis thaliana under drought. New Phytologist 184: 180–192. Ghandilyan A, Ilk N, Hanhart C, et al. 2009b. A strong effect of growth medium and 							
12 13 14 15 16	 Ghandilyan A, Barboza L, Tisné S, et al. 2009a. Genetic analysis identifies quantitative trait loci controlling rosette mineral concentrations in Arabidopsis thaliana under drought. New Phytologist 184: 180–192. Ghandilyan A, Ilk N, Hanhart C, et al. 2009b. A strong effect of growth medium and organ type on the identification of QTLs for phytate and mineral concentrations in 							
12 13 14 15 16 17	 Ghandilyan A, Barboza L, Tisné S, et al. 2009a. Genetic analysis identifies quantitative trait loci controlling rosette mineral concentrations in Arabidopsis thaliana under drought. New Phytologist 184: 180–192. Ghandilyan A, Ilk N, Hanhart C, et al. 2009b. A strong effect of growth medium and organ type on the identification of QTLs for phytate and mineral concentrations in three Arabidopsis thaliana RIL populations. Journal of Experimental Botany 60: 							
12 13 14 15 16 17 18	 Ghandilyan A, Barboza L, Tisné S, et al. 2009a. Genetic analysis identifies quantitative trait loci controlling rosette mineral concentrations in Arabidopsis thaliana under drought. New Phytologist 184: 180–192. Ghandilyan A, Ilk N, Hanhart C, et al. 2009b. A strong effect of growth medium and organ type on the identification of QTLs for phytate and mineral concentrations in three Arabidopsis thaliana RIL populations. Journal of Experimental Botany 60: 1409–1425. 							
12 13 14 15 16 17 18 19	 Ghandilyan A, Barboza L, Tisné S, et al. 2009a. Genetic analysis identifies quantitative trait loci controlling rosette mineral concentrations in Arabidopsis thaliana under drought. New Phytologist 184: 180–192. Ghandilyan A, Ilk N, Hanhart C, et al. 2009b. A strong effect of growth medium and organ type on the identification of QTLs for phytate and mineral concentrations in three Arabidopsis thaliana RIL populations. Journal of Experimental Botany 60: 1409–1425. Goffinet B, Gerber S. 2000. Quantitative trait loci: a meta-analysis. Genetics 155: 							
12 13 14 15 16 17 18 19 20	 Ghandilyan A, Barboza L, Tisné S, et al. 2009a. Genetic analysis identifies quantitative trait loci controlling rosette mineral concentrations in Arabidopsis thaliana under drought. New Phytologist 184: 180–192. Ghandilyan A, Ilk N, Hanhart C, et al. 2009b. A strong effect of growth medium and organ type on the identification of QTLs for phytate and mineral concentrations in three Arabidopsis thaliana RIL populations. Journal of Experimental Botany 60: 1409–1425. Goffinet B, Gerber S. 2000. Quantitative trait loci: a meta-analysis. Genetics 155: 463–473. 							
12 13 14 15 16 17 18 19 20 21	 Ghandilyan A, Barboza L, Tisné S, et al. 2009a. Genetic analysis identifies quantitative trait loci controlling rosette mineral concentrations in Arabidopsis thaliana under drought. New Phytologist 184: 180–192. Ghandilyan A, Ilk N, Hanhart C, et al. 2009b. A strong effect of growth medium and organ type on the identification of QTLs for phytate and mineral concentrations in three Arabidopsis thaliana RIL populations. Journal of Experimental Botany 60: 1409–1425. Goffinet B, Gerber S. 2000. Quantitative trait loci: a meta-analysis. Genetics 155: 463–473. Gruber BD, Giehl RF, Friedel S, von Wirén N. 2013. Plasticity of the Arabidopsis 							

1	Grusak MA, Broadley MR, White PJ. 2016. Plant macro- and micronutrient minerals
2	(Version 2.0). In: eLS, John Wiley & Sons, Chichester UK. doi:
3	10.1002/9780470015902.a0001306.pub2
4	Gutiérrez-Alanís D, Ojeda-Rivera JO, Yong-Villalobos L, Cárdenas-Torres
5	L, Herrera-Estrella L. 2018. Adaptation to phosphate scarcity: tips from
6	Arabidopsis roots. Trends in Plant Science 23: 721–730.
7	Hammond JP, Broadley MR, White PJ, et al. 2009. Shoot yield drives phosphorus
8	use efficiency in Brassica oleracea and correlates with root architecture
9	traits. Journal of Experimental Botany 60: 1953–1968.
10	Harada H, Leigh RA. 2006. Genetic mapping of natural variation in potassium
11	concentrations in shoots of Arabidopsis thaliana. Journal of Experimental Botany
12	57: 953–960.
13	Hawkesford M, Horst W, Kichey T, et al. 2012. Functions of macronutrients. In:
14	Marschner P, ed. Marschner's Mineral Nutrition of Higher Plants. London:
15	Academic Press, 135–189.
16	Huang XY, Salt DE. 2016. Plant ionomics: from elemental profiling to environmental
17	adaptation. Molecular Plant 9: 787–797.
18	Kering MK, Lukaszewska K, Blevins DG. 2009. Manganese requirement for
19	optimum photosynthesis and growth in NAD-malic enzyme C-4 species. Plant and
20	<i>Soil</i> 316: 217–226.
21	Kobayashi Y, Kuroda K, Kimura K, et al. 2008. Amino acid polymorphisms in
22	strictly conserved domains of a P-type ATPase HMA5 are involved in the
	37

1	mechanism of copper tolerance variation in Arabidopsis. Plant Physiology 148:						
2	969–980.						
3	Koprivova A, Harper AL, Trick M, Bancroft I, Kopriva S. 2014. Dissection of the						
4	control of anion homeostasis by associative transcriptomics in Brassica						
5	napus. Plant Physiology 166: 442–450.						
6	Körber N, Bus A, Li J, et al. 2016. Agronomic and seed quality traits dissected by						
7	genome-wide association mapping in Brassica napus. Frontiers in Plant Science						
8	7: 386.						
9	Lahner B, Gong J, Mahmoudian M, et al. 2003. Genomic scale profiling of nutrient						
10	and trace elements in Arabidopsis thaliana. Nature Biotechnology 21: 1215–1221.						
11	Li Y, Wang X, Zhang H, et al. 2019. Molecular identification of the phosphate tr						
12	ansporter family 1 (PHT1) genes and their expression profiles in response to ph						
13	osphorus deprivation and other abiotic stresses in Brassica napus. PLoS One 1						
14	4: e0220374.						
15	Liu J, Yang J, Li R, et al. 2009. Analysis of genetic factors that control shoot mineral						
16	concentrations in rapeseed (Brassica napus) in different boron						
17	environments. Plant and Soil 320: 255-266.						
18	Loudet O, Chaillou S, Merigout P, Talbotec J, Daniel-Vedele F. 2003. Quantitative						
19	trait loci analysis of nitrogen use efficiency in Arabidopsis. Plant Physiology 131:						
20	345–358.						
21	Loudet O, Saliba-Colombani V, Camilleri C, et al. 2007. Natural variation for sulfate						
22	content in Arabidopsis thaliana is highly controlled by APR2. Nature Genetics 39:						

1 896–900.

2	Maillard A, Etienne P, Diquélou S, et al. 2016. Nutrient deficiencies modify the
3	ionomic composition of plant tissues: a focus on cross-talk between molybdenum
4	and other nutrients in Brassica napus. Journal of Experimental Botany 67: 5631-
5	5641.
6	Mäser P, Thomine S, Schroeder JI, et al. 2001. Phylogenetic relationships within
7	cation transporter families of Arabidopsis. Plant Physiology 126: 1646–1667.
8	Mei H, Zhao J, Pittman JK, Lachmansingh J, Park S, Hirschi KD. 2007. In planta
9	regulation of the Arabidopsis Ca2+/H+ antiporter CAX1. Journal of Experimental
10	<i>Botany</i> 58: 3419–3427.
11	Meng L, Li H, Zhang L, Wang J. 2015. QTL IciMapping: integrated software for
12	genetic linkage map construction and quantitative trait locus mapping in bi-
13	parental populations. The Crop Journal 3: 265–279.
14	Müller J, Toev T, Heisters M, et al. 2015. Iron-dependent callose deposition adjusts
15	root meristem maintenance to phosphate availability. Developmental Cell 33: 216-
16	230.
17	Murashige T, Skoog F. 1962. A revised medium for rapid growth and bioassays with
18	tobacco tissue cultures. Physiologia Plantarum 15: 473–497.
19	Neugebauer K, Broadley MR, El-Serehy HA, et al. 2018. Variation in the angiosperm
20	ionome. Physiologia Plantarum 163: 306–322.
21	Neugebauer K, El-Serehy HA, George TS, et al. 2019. The influence of phylogeny
22	and ecology on root, shoot and plant ionomes of fourteen native Brazilian species.

1	Physiologia Plantarum, in press. doi: 10.1111/ppl.13018.
2	Paran I, Zamir D. 2003. Quantitative traits in plants: beyond the QTL. Trends in
3	<i>Genetics</i> 19: 303–306.
4	Payne KA, Bowen HC, Hammond JP, et al. 2004. Natural genetic variation in
5	caesium (Cs) accumulation by Arabidopsis thaliana. New Phytologist 162: 535-
6	548.
7	Pongrac P, Fischer S, Thompson JA, Wright G, White PJ. 2020. Early responses of
8	Brassica oleracea roots to zinc supply under sufficient and sub-optimal
9	phosphorus supply. Frontiers in Plant Science 10: 1645.
10	Puig S, Andrés-Colás N, García-Molina A, Peñarrubia L. 2007. Copper and iron
11	homeostasis in Arabidopsis: responses to metal deficiencies, interactions and
12	biotechnological applications. Plant, Cell & Environment 30: 271–290.
13	Qiu D, Morgan C, Shi J, et al. 2006. A comparative linkage map of oilseed rape and
14	its use for QTL analysis of seed oil and erucic acid content. Theoretical and
15	Applied Genetics 114: 67–80.
16	Salt DE, Baxter I, Lahner B. 2008. Ionomics and the study of the plant ionome.
17	Annual Review of Plant Biology 59: 709–733.
18	Sánchez-Bermejo E, Castrillo G, del Llano B, et al. 2014. Natural variation in
19	arsenate tolerance identifies an arsenate reductase in Arabidopsis thaliana. Nature
20	Communications 5: 4617.
21	Sharpe AG, Lydiate DJ. 2003. Mapping the mosaic of ancestral genotypes in a cultivar
22	of oilseed rape (Brassica napus) selected via pedigree breeding. Genome 46: 461-

468.

2	Shi L, Shi T, Broadley MR, et al. 2013. High-throughput root phenotyping screens
3	identify genetic loci associated with root architectural traits in Brassica napus
4	under contrasting phosphate availabilities. Annals of Botany 112: 381-389.
5	Thomas CL, Alcock TD, Graham NS, et al. 2016a. Root morphology and seed and
6	leaf ionomic traits in a Brassica napus L. diversity panel show wide phenotypic
7	variation and are characteristic of crop habit. BMC Plant Biology 16: 214.
8	Thomas CL, Graham NS, Hayden R, et al. 2016b. High-throughput phenotyping
9	(HTP) identifies seedling root traits linked to variation in seed yield and nutrient
10	capture in field-grown oilseed rape (Brassica napus L.). Annals of Botany 118:
11	655–665.
12	von Wirén N, Klair S, Bansal S, et al. 1999. Nicotianamine chelates both Fem and Fem.
13	Implications for metal transport in plants. Plant Physiology 119: 1107–1114.
14	Vreugdenhil D, Aarts MGM, Koornneef M, Nelissen H, Ernst WHO. 2004. Natural
15	variation and QTL analysis for cationic mineral content in seeds of Arabidopsis
16	thaliana. Plant, Cell & Environment 27: 828–839.
17	Wang W, Ding GD, White PJ, et al. 2019. Mapping and cloning of quantitative trait
18	loci for phosphorus efficiency in crops: opportunities and challenges. Plant and
19	<i>Soil</i> 439: 91–112.
20	Ward JT, Lahner B, Yakubova E, Salt DE, Raghothama KG. 2008. The effect of
21	iron on the primary root elongation of Arabidopsis during phosphate deficiency.
22	<i>Plant Physiology</i> 147: 1181–1191.

1	Watanabe T, Urayama M, Shinano T, Okada R, Osaki M. 2015. Application of
2	ionomics to plant and soil in fields under long-term fertilizer trials. SpringerPlus 4:
3	781.
4	Waters BM, Grusak MA. 2008. Quantitative trait locus mapping for seed mineral
5	concentrations in two Arabidopsis thaliana recombinant inbred populations. New
6	Phytologist 179: 1033–1047.
7	Welch RM. 1995. Micronutrient nutrition of plants. Critical Reviews in Plant Sciences
8	14: 49–82.
9	White PJ, Broadley MR. 2009. Biofortification of crops with seven mineral elements
10	often lacking in human diets – iron, zinc, copper, calcium, magnesium, selenium
11	and iodine. New Phytologist 182: 49-84.
12	White PJ, George TS, Gregory PJ, Bengough AG, Hallett PD, McKenzie BM. 2013.
13	Matching roots to their environment. Annals of Botany 112: 207-222.
14	White PJ, Hammond JP, King GJ, et al. 2010. Genetic analysis of potassium use
15	efficiency in Brassica oleracea. Annals of Botany 105: 1199-1210.
16	White PJ. 2012a. Ion uptake mechanisms of individual cells and roots: short-distance
17	transport. In: Marschner P, ed. Marschner's Mineral Nutrition of Higher Plants.
18	London: Academic Press, 7–47.
19	White PJ. 2012b. Long-distance transport in the xylem and phloem. In: Marschner P,
20	ed. Marschner's Mineral Nutrition of Higher Plants. London: Academic Press,
21	49–70.

22 Wu D, Sato K, Ma JF. 2015. Genome-wide association mapping of cadmium

1	accumulation in different organs of barley. New Phytologist 208: 817-829.
2	Wu J, Yuan YX, Zhang XW, et al. 2008. Mapping QTLs for mineral accumulation
3	and shoot dry biomass under different Zn nutritional conditions in Chinese
4	cabbage (Brassica rapa L. ssp. pekinensis). Plant and Soil 310: 25-40.
5	Xiao Y, Liu H, Wu L, Warburton M, Yan J. 2017. Genome-wide association studies
6	in maize: praise and stargaze. Molecular Plant 10: 359–374.
7	Yang M, Lu K, Zhao FJ et al. 2018. Genome-wide association studies reveal the
8	genetic basis of ionomic variation in rice. The Plant Cell 30: 2720–2740.
9	Zeng LP. 2011. Construction of reciprocal introgression lines and evaluation in
10	Brassica napus L. Master Thesis, Huazhong Agricultural University, China.
11	Zhang Y, Thomas CL, Xiang J, et al. 2016. QTL meta-analysis of root traits in
12	Brassica napus under contrasting phosphorus supply in two growth systems.
13	Scientific Reports 6: 33113.
14	Zhao J, Jamar DC, Lou P, et al. 2008. Quantitative trait loci analysis of phytate and
15	phosphate concentrations in seeds and leaves of Brassica rapa. Plant, Cell &
16	<i>Environment</i> 31: 887–900.

1 Figure legends

Fig. 1. Shoot (A) and root concentrations (B) of eleven mineral elements in cultivars 2 Tapidor and Ningyou 7 grown in various environments. Shoot concentrations of B, Ca, 3 Cu, Fe, Mg, P and Zn were determined in three studies (study 1, 2 and 3), comprising 4 eleven growth environments: two environments were from study 1 (0.25 and 50 μ M B 5 in a hydroponic system, Liu *et al.*, 2009), five from study 2-1 (0, 6, 312.5, 625 and 1250 6 μM P in an agar system, Shi et al., 2013), two from study 2-2 (0 and 625 μM P in an 7 agar system, Shi et al., 2013), and two from study 3 (0 and 625 µM P in an agar system 8 9 in the present paper); Shoot concentrations of K and Mn were from nine environments, of which five were from study 2-1, two were from study 2-2, and two were from study 10 3; Shoot concentrations of Na and S were from seven environments, of which five were 11 12 from study 2-1 and two were from study 2-2; Root concentrations of B, Ca, Cu, Fe, K, Mg, Mn, P and Zn were from nine environments, of which five were from study 2-1, 13 two were from study 2-2, and two were from study 3; Root concentrations of Na and S 14 were from seven environments, of which five were from study 2-1 and two were from 15 study 2-2. The two walls of the box correspond to first and third quartiles. Whiskers are 16 separated from the box by a 1.5 interquartile range (3rd quartile minus 1st quartile). 17 Circles represent individual measures outside the whiskers. The central black line in the 18 box is a median. A significant difference for each mineral element tested by one-sample 19 t test between the mean and 1 is indicated by an asterisk (*P < 0.05, **P < 0.01). 20 21 Figure 1.

22



Fig. 2. The effect of P deficiency on shoot (A) and root concentrations (B) of eleven 1 mineral elements in Tapidor and Ningyou 7. The data for Tapidor and Ningyou 7 at a 2 3 low P supply (LP) were scaled to that of cultivars Tapidor and Ningyou 7 at an optimal P supply (OP). The black dotted line indicates shoot and root concentrations of the 4 eleven mineral elements of Tapidor and Ningyou 7 at OP (Supplementary Data Table 5 S1). The black and gray lines indicate shoot and root concentrations of the eleven 6 mineral elements in Tapidor and Ningyou 7 at LP, respectively. Traits that show a 7 significant difference between the two P treatments are labelled by a black asterisk for 8 Tapidor and a gray asterisk for Ningyou 7 (*P < 0.05, Student's *t*-test). Each trait 9 contains eight to eleven biological replicates. 10





1	Fig. 3. Shoot and root concentrations, shoot, root and plant contents, and partitioning
2	to the shoot of eleven mineral elements in the BnaTNDH population grown at an
3	optimal (OP) and a low P supply (LP). Boxes represent the mid two quartiles with the
4	median drawn; whiskers are the 95% confidence limits, and extreme values are plotted
5	individually. For each trait, a significant difference between two P treatments is labelled
6	by an asterisk (* $P < 0.05$, ** $P < 0.01$, Student's <i>t</i> -test).



$ \begin{array}{c} Total contract in the second secon$	** 12 ** 10 ** 8 1 0 8 1 0 20 1 20 1 1 0 25 1 0 1 </th <th>400 350 300 250 250 200 150 100 4000 1 400 400</th> <th>5000 5000 5000 5000 5000 5000 5000 5000 5000 5000 5000 5000 5000 1 1 1 5000 1 1 1 1 1 1 1 1 1 1 1 1 1</th> <th>5000 </th> <th>400 350 300 250 260 240 200 180 160 2.5 1.5 1.0 1.5 1.0 1.5 1.5 1.0 1.5 1.5 1.5 1.0 1.5 1.5 1.5 1.5 1.5 1.5 1.5 1.5</th> <th>4000 3500 2500 2500 2500 2500 2000 1500 25 25 20 15 10 5 </th> <th>18000 1** 16000 1 14000 1 12000 1 10000 1 8000 1 6000 1 4000 1 10000 1 10000 1 10000 1 10000 1 10000 1 1400 1 140 1 120 1 140 1 120 1 140 1 120 1 140 1 120 1 140 1 120 1 140 1 120 1 140 1 120 1 140 1 120 1 140 1 140 1 140 1 140 1 140 1 140 1 140</th> <th>18000 *** 16000 *** 14000 1 12000 1 18000 *** 18000 *** 18000 *** 18000 *** 18000 *** 16000 1 12000 1 12000 1 12000 1 100 1 100 1 100 1 100 1 100 1 100 1 100 1 100 1 100 1 100 1 100 1 100 1 100 1 100 1 100 1 100 1 110 1 120 1 140 1 100 1 100 1 100 1 100 1</th> <th>240 200 180 100 140 200 300 200 100 1.4 120 200 300 200 100 1.4 1.2 1.0 0.8 0.4 0.5 0.4 0.5 0.4 0.5 0.4 0.5 0.4 0.5 0.5 0.4 0.5 0.5 0.4 0.5 0.5 0.4 0.5 0.5 0.5 0.5 0.5 0.5 0.5 0.5 0.5 0.5</th>	400 350 300 250 250 200 150 100 4000 1 400 400	5000 5000 5000 5000 5000 5000 5000 5000 5000 5000 5000 5000 5000 1 1 1 5000 1 1 1 1 1 1 1 1 1 1 1 1 1	5000	400 350 300 250 260 240 200 180 160 2.5 1.5 1.0 1.5 1.0 1.5 1.5 1.0 1.5 1.5 1.5 1.0 1.5 1.5 1.5 1.5 1.5 1.5 1.5 1.5	4000 3500 2500 2500 2500 2500 2000 1500 25 25 20 15 10 5 	18000 1** 16000 1 14000 1 12000 1 10000 1 8000 1 6000 1 4000 1 10000 1 10000 1 10000 1 10000 1 10000 1 1400 1 140 1 120 1 140 1 120 1 140 1 120 1 140 1 120 1 140 1 120 1 140 1 120 1 140 1 120 1 140 1 120 1 140 1 140 1 140 1 140 1 140 1 140 1 140	18000 *** 16000 *** 14000 1 12000 1 18000 *** 18000 *** 18000 *** 18000 *** 18000 *** 16000 1 12000 1 12000 1 12000 1 100 1 100 1 100 1 100 1 100 1 100 1 100 1 100 1 100 1 100 1 100 1 100 1 100 1 100 1 100 1 100 1 110 1 120 1 140 1 100 1 100 1 100 1 100 1	240 200 180 100 140 200 300 200 100 1.4 120 200 300 200 100 1.4 1.2 1.0 0.8 0.4 0.5 0.4 0.5 0.4 0.5 0.4 0.5 0.4 0.5 0.5 0.4 0.5 0.5 0.4 0.5 0.5 0.4 0.5 0.5 0.5 0.5 0.5 0.5 0.5 0.5 0.5 0.5
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1 Figure 3

Fig. 4. Principal component (PC) plot of the first two PC scores of shoot and root
 concentrations of eleven mineral elements in the *Bna*TNDH population at an optimal
 (A) and a low P supply (B). Colours indicate organs (black = shoot, gray = root).



Fig. 5. The epistatic QTLs for shoot and root concentrations, shoot, root and plant 1 contents, and partitioning to the shoot of eleven mineral elements as well as the total 2 dry weight (TDW) at an optimal (A) and a low P supply (B). The circle indicates the 3 19 linkage groups and the black and gray lines denote the epistatic interactions between 4 each two loci for different traits. Two pleiotropic epistatic QTLs are presented at both 5 optimal and low P supplies. Each ionomic trait was denominated as "S (abbreviation of 6 shoot) or R (abbreviation of root) or P (abbreviation of plant) + [mineral element] + 7 conc (abbreviation of concentration) or cont (abbreviation of content) or part 8 (abbreviation of partitioning)". 9



1 Figure 5

Fig. 6. Location of the QTL clusters for shoot and root concentrations, shoot, root and plant contents, and partitioning to the shoot of eleven mineral elements in the *Bna*TNDH population grown at an optimal (OP) and a low P supply (LP) identified by meta-QTL analysis. The vertical columns represented linkage groups of the *Bna*TNDH population. The black blocks inside each vertical column represent the QTL clusters. The name of each QTL cluster is on the right of the linkage group. The number in brackets indicates the number of mineral elements controlled by different QTL clusters.





1	Fig. 7. Shoot K (A), Mg (B) and S (C) concentrations of cultivars Tapidor, Ningyou 7
2	and five BC4F2:3 lines generated with cultivar Tapidor as the recurrent parent and
3	cultivar Ningyou 7 as the donor parent grown at LP in the agar system. Introgressed
4	regions of cultivar Ningyou 7 in the QTL cluster Cl17.1 in these five lines are indicated
5	in white in (D), and the physical positions of the five InDel markers are shown. All lines
6	have five replicates except for the line 2292-3, which has three replicates. A significant
7	difference between data for cultivar Tapidor and other genotypes is indicated by an
8	asterisk (* $P < 0.05$, ** $P < 0.01$) according to Student's <i>t</i> -test.





	Correlation coefficients											
Shoot concentration		В	Ca	Cu	Fe	Κ	Mg	Mn	Na	Р	S	Zn
	В	0.45**	0.11	0.33**	0.16*	0.39**	0.13	0.22**	0.25**	0.35**	-0.02	0.00
	Ca	0.36**	0.53**	0.14	-0.04	0.36**	0.37**	0.75**	0.49**	0.14	0.16*	0.45**
	Cu	-0.02	0.18*	0.18*	0.25**	0.09	0.11	0.08	0.11	0.14	0.02	0.12
	Fe	0.39**	0.42**	0.05	0.20*	-0.07	0.07	-0.10	0.11	-0.08	0.01	0.01
	Κ	0.47**	0.48**	0.15	0.25**	0.50**	-0.01	0.40**	0.26**	0.16*	0.05	0.18*
	Mg	0.11	0.40**	0.24**	0.02	0.44**	0.52**	0.27**	0.04	0.20**	0.28**	0.08
	Mn	0.45**	0.84**	0.07	0.37**	0.43**	0.27**	0.44**	0.19*	0.10	0.04	0.34**
	Na	0.58**	0.71**	0.19*	0.53**	0.57**	0.34**	0.56**	0.45**	0.17*	0.07	0.28**
	Р	0.09	0.15	0.53**	-0.03	0.30**	0.54**	0.00	0.17*	0.16*	0.09	0.10
	S	0.14	0.13	0.15	-0.11	0.33**	0.28**	0.18*	0.12	0.15	0.69**	0.18*
	Zn	0.44**	0.62**	0.16*	0.66**	0.27**	0.05	0.62**	0.53*	0.05	0.11	-0.11
Root concentration		В	Ca	Cu	Fe	Κ	Mg	Mn	Na	Р	S	Zn
	В	0.21**	0.38**	0.11	-0.02	0.10	0.07	0.46**	0.16*	0.02	-0.02	0.15
	Ca	0.44**	0.38**	-0.06	-0.14	0.17*	0.40**	0.70**	0.34**	0.08	-0.02	-0.01
	Cu	0.06	0.31**	0.05	0.14	-0.04	-0.06	-0.05	0.01	0.10	0.05	0.05
	Fe	-0.02	0.12	0.45**	0.32**	0.13	-0.14	0.06	0.06	0.18*	0.22**	0.04
	Κ	0.05	0.09	0.08	-0.02	0.56**	0.13	0.12	0.30**	0.08	0.17*	-0.09
	Mg	-0.08	0.17*	0.04	-0.03	0.22**	0.73**	0.22**	0.28**	0.33**	0.25**	0.17*
	Mn	0.43**	0.76**	0.14	0.19*	0.02	-0.02	0.34**	0.32**	0.01	0.04	0.06
	Na	0.29**	0.42**	0.11	0.11	0.31**	0.19*	0.35**	0.50**	-0.05	0.27**	0.11
	Р	-0.06	-0.02	0.08	0.21**	0.28**	0.21**	0.03	-0.05	0.09	0.14	0.04
	S	0.01	0.10	0.06	0.13	0.42**	0.38**	-0.06	0.24**	0.09	0.58**	0.23**
	Zn	0.17*	0.46**	0.24**	0.14	0.15	-0.09	0.52**	0.17*	0.06	0.11	0.03

Table 1. Pearson's correlation coefficients among shoot concentrations and among root concentrations of eleven mineral elements in the *Bna*TNDH mapping population at an optimal (upper right triangle) and a low P supply (lower left triangle)

Grey values on the diagonal indicate the correlations between the two P treatments for each trait, **P*<0.05, ***P*<0.01.

						Correlat	ion coeffici	ents				
Plant content		В	Ca	Cu	Fe	Κ	Mg	Mn	Na	Р	S	Zn
	В	0.67**	0.85**	0.68**	0.65**	0.94**	0.88**	0.88**	0.88**	0.89**	0.80**	0.72**
	Ca	0.84**	0.70**	0.66**	0.56**	0.90**	0.89**	0.95**	0.90**	0.84**	0.79**	0.81**
	Cu	0.47**	0.51**	0.27**	0.57**	0.66**	0.62**	0.62**	0.65**	0.62**	0.57**	0.59**
	Fe	0.54**	0.53**	0.40**	0.51**	0.64**	0.54**	0.59**	0.59**	0.57**	0.54**	0.58**
	Κ	0.93**	0.86**	0.53**	0.52**	0.66**	0.89**	0.93**	0.90**	0.88**	0.82**	0.77**
	Mg	0.81**	0.82**	0.50**	0.41**	0.90**	0.75**	0.87**	0.88**	0.88**	0.85**	0.73**
	Mn	0.89**	0.95**	0.49**	0.54**	0.90**	0.82**	0.69**	0.87**	0.85**	0.79**	0.81**
	Na	0.90**	0.91**	0.50**	0.59**	0.90**	0.81**	0.88**	0.70**	0.84**	0.80**	0.79**
	Р	0.60**	0.57**	0.61**	0.28**	0.70**	0.75**	0.55**	0.59**	0.41**	0.78**	0.72**
	S	0.81**	0.74**	0.46**	0.38**	0.85**	0.80**	0.79**	0.75**	0.57**	0.76**	0.70**
	Zn	0.86**	0.87**	0.54**	0.62**	0.81**	0.68**	0.90**	0.84**	0.50**	0.71**	0.50**
Partitioning		В	Ca	Cu	Fe	Κ	Mg	Mn	Na	Р	S	Zn
	В	0.56**	0.54**	0.30**	0.28**	0.52**	0.58**	0.60**	0.55**	0.40**	0.47**	0.61**
	Ca	0.58**	0.56**	0.06	0.23**	0.58**	0.58**	0.86**	0.59**	0.26**	0.48**	0.33**
	Cu	0.21**	0.32**	0.26**	0.15	0.05	0.13	0.04	0.10	0.26**	0.27**	0.20*
	Fe	0.32**	0.39**	0.33**	0.32**	0.19*	0.30**	0.24**	0.37**	0.28**	0.34**	0.37**
	Κ	0.51**	0.48**	0.16*	0.23**	0.69**	0.51**	0.57**	0.60**	0.31**	0.59**	0.36**
	Mg	0.47**	0.52**	0.21**	0.31**	0.58**	0.76**	0.50**	0.54**	0.48**	0.64**	0.56**
	Mn	0.62**	0.87**	0.25**	0.41**	0.49**	0.50**	0.52**	0.53**	0.25**	0.45**	0.38**
	Na	0.56**	0.71**	0.20**	0.32**	0.54**	0.57**	0.68**	0.65**	0.21**	0.50**	0.38**
	Р	0.58**	0.43**	0.21**	0.39**	0.55**	0.54**	0.44**	0.41**	0.44**	0.54**	0.46**
	S	0.53**	0.46**	0.21**	0.35**	0.69**	0.64**	0.50**	0.46**	0.63**	0.78**	0.57**
	Zn	0.56**	0.69**	0.26**	0.37**	0.64**	0.59**	0.68**	0.59**	0.60**	0.64**	0.46**

Table 2. Pearson's correlation coefficients among plant contents and among partitioning to the shoot of eleven mineral elements in the *Bna*TNDH mapping population at an optimal (upper right triangle) and a low P supply (lower left triangle)

Grey values on the diagonal indicate the correlations between the two P treatments for each trait, **P*<0.05, ***P*<0.01.

Table 3. Significant QTLs associated with shoot and root concentrations, shoot, root and plant contents, and partitioning to the shoot of eleven mineral	
elements, shoot dry weight (SDW), root dry weight (RDW) and total dry weight (TDW) in the BnaTNDH population at an optimal (OP) and a low P	
supply (LP)	

Trait	Mineral element	P treatment	QTL name	Chromoso me	Position (cM)	LOD score	Confidence interval (cM)	Additive effect	R2 (%)
Shoot concentration	В	ОР	BconcOPS-A01	A01	48	4.73	47.5–49.5	1.617	13.6
		LP	BconcLPS-A03	A03	49	2.52	48.5–49.5	-1.332	6.8
	Ca	OP	CaconcOPS- A03	A03	45	2.63	44.5–45.5	-215.2	6.1
			CaconcOPS- A07	A07	89	6.28	88.5–89.5	311.9	15.0
		LP	CaconcLPS- A03	A03	38	4.16	35.5–42.5	-249.1	6.8
			CaconcLPS- A07	A07	88	10.39	87.5–88.5	378.9	18.0
			CaconcLPS- A09	A09	53	6.64	52.5–54.5	295.1	10.9
			CaconcLPS- C04	C04	63	2.82	61.5–68.5	-187.7	4.4
	Fe	LP	FeconcLPS- A02	A02	52	2.73	51.5–52.5	10.94	6.3
			FeconcLPS- A09	A09	137	3.68	136.5–139.0	12.59	8.6
			FeconcLPS- C06	C06	9	4.15	6.5–10.5	13.83	10.4
	Κ	OP	KconcOPS-A03	A03	45	8.14	44.5–45.5	-2201	3.6

		KconcOPS-A04	A04	37	3.54	36.5-37.5	1258	1.4
		KconcOPS-A09	A09	68	6.10	67.5–68.5	1679	2.4
		KconcOPS- C08a	C08	32	29.92	31.5–32.5	-4455	17.1
		KconcOPS- C08b	C08	34	21.69	33.5–36.5	3555	10.9
	LP	KconcLPS-A02	A02	73	3.98	72.5–73.5	-1536	5.2
		KconcLPS-A03	A03	47	3.91	46.5-47.5	-1490	5.1
		KconcLPS-A09	A09	116	3.43	115.5–116.5	1356	4.9
		KconcLPS-C06	C06	83	8.40	80.5-84.5	-2227	12.2
		KconcLPS-C07	C07	34	2.70	32.5-35.5	-1174	3.6
Mg	OP	MgconcOPS- A09	A09	81	2.61	78.5–82.5	77.90	6.3
		MgconcOPS- C07	C07	30	4.87	29.5–31.5	-109.7	11.9
	LP	MgconcLPS- A09	A09	92	4.91	91.5–93.5	130.3	9.8
		MgconcLPS- C07	C07	35	3.40	34.5–35.5	-109.6	6.6
Mn	OP	MnconcOPS- A04	A04	12	3.39	10.5–13.5	7.743	6.9
		MnconcOPS- A07	A07	89	3.13	88.5–89.5	7.712	6.9
		MnconcOPS- A08	A08	77	2.85	74.5–79.5	7.106	5.8
		MnconcOPS- C07	C07	89	4.12	88.5–90.5	-8.629	8.3
	LP	MnconcLPS- A07	A07	92	4.39	89.5–101.5	9.852	11.1

			MnconcLPS- A09	A09	53	4.23	52.5–54.5	9.538	10.4
	Na	LP	NaconcLPS- A09	A09	136	3.10	135.5–137.5	91.76	10.7
	Р	OP	PconcOPS-A04	A04	36	3.76	31.5-36.5	437.6	8.5
			PconcOPS-C04	C04	39	4.34	36.5-39.5	-472.2	9.9
	S	OP	SconcOPS-A09	A09	11	12.76	8.5-16.5	-634.1	19.3
			SconcOPS-C02	C02	5	3.29	3.5–6.5	-300.2	4.6
			SconcOPS-C06	C06	16	6.59	15.5-18.5	-414.0	8.7
			SconcOPS-C07	C07	39	12.91	36.5-40.5	-627.2	19.3
		LP	SconcLPS-A09	A09	6	6.12	3.5–7.5	-583.3	12.7
			SconcLPS-C02	C02	46	3.17	44.5-47.5	-394.6	6.2
			SconcLPS-C07	C07	34	6.36	33.5–35.5	-588.7	13.3
Root concentration	В	OP	BconcOPR-A05	A05	12	4.83	11.5–14.5	-1.278	13.3
		LP	BconcLPR- A09a	A09	32	3.51	29.5–34.5	1.330	7.8
			BconcLPR- A09b	A09	128	3.36	124.5–128.5	1.164	6.3
	Cu	LP	CuconcLPR- A06	A06	85	2.71	84.5-85.5	0.962	7.3
	Fe	OP	FeconcOPR- C01	C01	47	3.14	45.5–48.5	126.4	11.4
	K	OP	KconcOPR- A06	A06	59	3.15	58.5–59.5	-1253	10.9
		LP	KconcLPR-A03	A03	92	3.63	91.5-92.5	1670	9.9
	Mg	OP	MgconcOPR- A07	A07	71	7.44	70.5–74.5	115.6	13.6

		MgconcOPR- C09	C09	46	3.10	45.5–47.5	-74.75	5.5
	LP	MgconcLPR- A03	A03	101	4.95	99.5–102.5	-91.12	9.2
		MgconcLPR- A04	A04	31	4.10	30.5–31.5	80.92	7.5
		MgconcLPR- A10	A10	49	3.54	48.5–49.5	-83.15	6.8
		MgconcLPR- C06	C06	76	2.52	75.5–76.5	-67.11	4.6
		MgconcLPR- C07	C07	67	3.79	65.5–67.5	79.20	6.9
Mn	OP	MnconcOPR- A08	A08	1	2.54	0–4.5	-4.041	6.6
Na	OP	NaconcOPR- A04	A04	31	2.55	30.5–31.5	61.66	4.8
		NaconcOPR- A09	A09	80	5.43	77.5–81.5	-95.59	11.6
		NaconcOPR- C03	C03	6	3.37	5.5–6.5	92.26	6.4
	LP	NaconcLPR- C03	C03	12	2.63	10.5–12.5	81.56	7.1
Р	OP	PconcOPR-A09	A09	124	4.85	123.5-124.5	353.3	12.1
		PconcOPR-C04	C04	56	3.24	55.5–56.5	-282.5	7.8
		PconcOPR-C07	C07	2	2.55	0–2.5	249.1	6.1
S	OP	SconcOPR-A04	A04	37	10.98	36.5-37.5	476.9	19.2
		SconcOPR-A09	A09	9	4.22	8.5-15.5	-291.8	6.8
		SconcOPR-C01	C01	41	11.52	40.5-41.5	496.5	20.5
		SconcOPR-C09	C09	88	2.93	87.5-89.5	-233.1	4.6

		LP	SconcLPR- A04a	A04	27	14.09	26.5–27.5	700.3	23.1
			SconcLPR- A04b	A04	53	4.78	51.5–57.5	376.9	6.7
			SconcLPR-C01	C01	8	5.72	7.5-8.5	416.0	8.1
			SconcLPR-C05	C05	98	6.24	90.5-105.5	-483.6	11.1
	Zn	OP	ZnconcOPR- A03	A03	69	4.02	68.5–69.5	-18.83	9.2
			ZnconcOPR- A04a	A04	15	3.23	13.5–15.5	15.91	7.2
			ZnconcOPR- A04b	A04	65	2.73	64.5–67.5	14.66	6.1
			ZnconcOPR- C04	C04	3	2.93	2.5–3.5	-15.17	6.5
Shoot content	В	OP	BcontOPS-A03	A03	51	4.60	50.5-51.5	-0.026	12.5
		LP	BcontLPS-A03	A03	52	5.69	51.5-52.5	-0.025	14.8
			BcontLPS-A04	A04	5	3.95	4.5–5.5	-0.019	9.9
	Ca	OP	CacontOPS- A02	A02	75	3.99	74.5–75.5	-3.181	6.8
			CacontOPS- A03	A03	51	8.09	50.5–51.5	-4.456	14.5
			CacontOPS- A04	A04	3	2.57	2.5-4.5	-2.268	4.2
			CacontOPS- A07	A07	75	2.75	74.5–75.5	2.281	4.4
		LP	CacontLPS- A02	A02	112	4.22	99.5–120.0	-2.469	8.9
			CacontLPS- A03	A03	52	6.85	51.5–52.5	-3.066	11.7

		CacontLPS- A04	A04	5	5.71	4.5–5.5	-2.611	9.8
		CacontLPS- A07	A07	69	4.47	68.5–70.5	2.189	7.1
		CacontLPS- A09	A09	67	2.98	65.5–68.5	1.810	4.8
Cu	OP	CucontOPS- A03	A03	51	6.33	50.5–51.5	-0.004	16.6
Fe	LP	FecontLPS-A09	A09	136	4.95	135.5-136.5	0.108	12.8
Κ	OP	KcontOPS-A02	A02	75	5.86	74.5-75.5	-35.47	5.2
		KcontOPS-A03	A03	52	8.38	51.5-52.5	-41.48	7.6
		KcontOPS-A04	A04	3	3.20	2.5-4.5	-22.79	2.7
	KcontOPS- A09a	A09	68	14.67	66.5–68.5	51.86	14.2	
		KcontOPS- A09b	A09	75	7.87	73.5–75.5	-36.06	6.9
	LP	KcontLPS-A02	A02	72	2.93	71.5-72.5	-19.88	6.4
		KcontLPS-A03	A03	52	4.78	51.5-52.5	-26.14	11.3
		KcontLPS-A04	A04	5	3.32	4.5–5.5	-19.93	7.5
Mg	OP	MgcontOPS- A02	A02	73	4.39	72.5–73.5	-1.556	8.1
		MgcontOPS- A03	A03	51	6.07	50.5–51.5	-1.810	11.6
		MgcontOPS- A04	A04	3	4.56	2.5–4.5	-1.443	8.3
	LP	MgcontLPS- A03	A03	59	5.12	58.5–59.5	-1.389	9.2
		MgcontLPS- A04	A04	5	6.06	4.5–5.5	-1.489	11.6

		MgcontLPS- A09	A09	74	2.91	73.5–74.5	1.028	5.7
Mn	OP	MncontOPS- A02	A02	75	2.97	74.5–75.5	-0.114	7.2
		MncontOPS- A03	A03	51	5.04	50.5–51.5	-0.142	12.2
	LP	MncontLPS- A03	A03	52	4.50	51.5–52.5	-0.117	10.3
		MncontLPS- A04	A04	5	3.91	4.5–5.5	-0.101	8.9
		MncontLPS- A07	A07	69	2.52	68.5–70.5	0.078	5.4
Na	OP	NacontOPS- A03	A03	51	3.79	50.5–51.5	-1.223	10.5
	LP	NacontLPS- A03	A03	52	4.80	51.5–52.5	-1.170	13.0
		NacontLPS- A04	A04	5	3.78	4.5–5.5	-0.963	10.2
		NacontLPS- A09	A09	132	2.68	130.5–132.5	0.775	6.8
Р	OP	PcontOPS-A02	A02	79	46.12	78.5–79.5	-29.55	23.0
		PcontOPS-A03	A03	51	4.44	50.5-51.5	-6.583	1.1
		PcontOPS-A04	A04	2	3.85	1.5–2.5	-5.667	1.0
		PcontOPS-C04	C04	22	2.93	19.5–25.5	-4.843	0.7
	LP	PcontLPS-C09	C09	123	2.64	117.5–134.5	-2.403	7.1
S	OP	ScontOPS-A02	A02	73	3.37	72.5-73.5	-4.470	6.3
		ScontOPS-A03	A03	51	4.01	50.5-51.5	-4.815	7.8
		ScontOPS-A04	A04	3	5.42	2.5-4.5	-5.288	10.5
	LP	ScontLPS-A02	A02	54	3.33	53.5-54.5	-4.130	5.8

			ScontLPS-A03	A03	57	3.74	56.5-57.5	-4.567	6.5
			ScontLPS-A04	A04	5	7.63	4.5–5.5	-6.519	14.8
	Zn	OP	ZncontOPS- A03	A03	51	4.81	50.5–51.5	-0.084	13.3
		LP	ZncontLPS- A02	A02	106	2.66	92.5–120.0	-0.062	9.1
			ZncontLPS- A03	A03	52	3.57	51.5–52.5	-0.063	8.0
			ZncontLPS- A04	A04	5	3.51	4.5–5.5	-0.058	7.9
Root content	В	LP	BcontLPR-A03	A03	50	3.47	49.5–50.5	-0.003	13.7
	Ca	OP	CacontOPR- A03	A03	69	2.65	68.5–69.5	-0.333	7.8
		LP	CacontLPR- A03	A03	52	4.51	51.5–52.5	-0.398	11.7
	Cu	LP	CucontLPR- A05	A05	14	4.28	12.5–18.5	-0.001	10.8
			CucontLPR- A09	A09	71	2.81	70.5–71.5	-0.001	6.8
	Κ	LP	KcontLPR-A03	A03	52	2.79	51.5-52.5	-3.286	8.1
	Mg	OP	MgcontOPR- A03	A03	105	2.70	104.5-107.5	-0.154	8.0
		LP	MgcontLPR- A03	A03	85	4.79	84.5-85.5	-0.159	9.9
			MgcontLPR- A07	A07	53	4.23	52.5–53.5	0.145	8.2
			MgcontLPR- C04	C04	60	3.22	58.5-60.5	-0.127	6.4

Mn	OP	MncontOPR- A03	A03	69	2.54	68.5–69.5	-0.016	7.4
	LP	MncontLPR- A03	A03	52	4.03	51.5–52.5	-0.018	10.9
Na	LP	NacontLPR- A02	A02	107	4.12	96.5–120.0	-0.200	11.8
		NacontLPR- A03	A03	52	4.35	51.5–52.5	-0.178	8.1
Р	OP	PcontOPR-A03	A03	78	4.85	76.5-81.5	-0.749	9.6
		PcontOPR-C04	C04	23	3.75	19.5–25.5	-0.652	7.4
		PcontOPR-C08	C08	69	3.77	66.5–69.5	-0.674	7.9
		PcontOPR-C09	C09	14	3.87	9.5–16.5	0.671	7.7
S	OP	ScontOPR-A03	A03	72	3.12	69.5-73.5	-0.987	8.7
	LP	ScontLPR-A03	A03	50	2.71	49.5-50.5	-0.828	8.7
		ScontLPR-A04	A04	38	5.45	37.5–38.5	0.994	14.5
		ScontLPR-C07	C07	84	2.52	83.5-84.5	-0.688	6.5
Zn	OP	ZncontOPR- A03	A03	69	6.81	68.5–69.5	-0.037	13.4
		ZncontOPR- A04	A04	65	3.49	64.5–67.5	0.025	6.5
		ZncontOPR- C04a	C04	6	2.86	3.5–9.5	-0.022	5.3
		ZncontOPR- C04b	C04	91	3.39	90.5–91.5	-0.024	6.2
		ZncontOPR- C09	C09	49	2.67	47.5–50.5	0.022	4.9
	LP	ZncontLPR- A03	A03	52	3.97	51.5-52.5	-0.018	10.4

			ZncontLPR-	C04	25	2.63	22.5–25.5	-0.013	6.6
Plant content	В	OP	BcontOPP-A03	A03	51	4.01	50.5-51.5	-0.027	11.8
		LP	BcontLPP-A03	A03	52	5.24	51.5-52.5	-0.027	13.8
			BcontLPP-A04	A04	9	3.83	8.5–9.5	-0.021	9.6
	Ca	OP	CacontOPP- A02	A02	73	4.06	72.5–73.5	-3.368	7.5
			CacontOPP- A03	A03	51	6.85	50.5–51.5	-4.380	13.6
			CacontOPP- C09	C09	49	3.11	47.5–50.5	2.724	5.7
		LP	CacontLPP- A02	A02	114	3.67	101.5-120.0	-2.603	8.3
			CacontLPP- A03	A03	52	6.67	51.5–52.5	-3.421	12.4
			CacontLPP- A04	A04	5	4.55	4.5–5.5	-2.647	8.6
			CacontLPP- A07	A07	69	4.40	68.5–70.5	2.469	7.6
	Cu	OP	CucontOPP- A03	A03	51	5.75	50.5–51.5	-0.005	16.0
		LP	CucontLPP- C05	C05	0	2.88	0–0.5	-0.002	8.4
	Κ	OP	KcontOPP-A02	A02	75	3.18	74.5-75.5	-32.69	8.8
			KcontOPP-A03	A03	51	5.17	50.5-51.5	-38.96	13.8
		LP	KcontLPP-A02	A02	72	2.77	71.5–72.5	-21.42	6.3
			KcontLPP-A03	A03	52	4.88	51.5-52.5	-29.28	12.0
			KcontLPP-A04	A04	5	2.73	4.5-5.5	-19.99	6.5

Mg	OP	MgcontOPP- A02	A02	73	4.09	72.5–73.5	-1.656	9.1
		MgcontOPP- A03	A03	51	4.63	50.5–51.5	-1.720	10.5
		MgcontOPP- A04	A04	9	4.21	8.5–9.5	-1.514	9.2
	LP	MgcontLPP- A04	A04	5	3.16	4.5–5.5	-1.291	12.4
Mn	OP	MncontOPP- A02	A02	75	3.15	74.5–75.5	-0.127	7.2
		MncontOPP- A03	A03	51	5.51	50.5–51.5	-0.160	12.7
		MncontOPP- C09	C09	48	3.12	47.5–49.5	0.113	6.8
	LP	MncontLPP- A03	A03	52	3.61	51.5–52.5	-0.120	11.3
		MncontLPP- A04	A04	5	2.58	4.5–5.5	-0.094	8.0
Na	OP	NacontOPP- A03	A03	51	6.31	50.5–51.5	-1.585	11.9
		NacontOPP- A04	A04	2	3.31	1.5–2.5	-1.063	5.9
		NacontOPP- C09	C09	48	2.93	47.5–49.5	1.000	5.1
	LP	NacontLPP- A02	A02	110	3.34	97.5–120.0	-1.188	10.8
		NacontLPP- A03	A03	52	5.19	51.5–52.5	-1.339	11.8

			NacontLPP-	A04	5	3.40	4.5–5.5	-1.015	7.8
			A04 NacontLPP-	A09	139	2.54	135.5–139.0	0.830	5.3
	Р	OP	PcontOPP-A03	A03	51	2.75	50.5-51.5	-6.400	8.4
		LP	PcontLPP-C09	C09	123	2.66	117.5–134.5	-2.567	7.2
	S	OP	ScontOPP-A02	A02	73	4.95	72.5–73.5	-6.198	11.2
			ScontOPP-A03	A03	56	3.24	55.5-56.5	-4.845	7.4
			ScontOPP-A04	A04	8	3.71	7.5-8.5	-4.869	8.3
			ScontOPP-C07	C07	50	2.72	48.5-50.5	-4.318	6.1
		LP	ScontLPP-A02	A02	54	3.31	53.5-54.5	-4.601	6.4
			ScontLPP-A03	A03	57	3.75	56.5-57.5	-5.110	7.3
			ScontLPP-A04	A04	5	5.58	4.5-5.5	-6.186	11.9
			ScontLPP-C07	C07	40	2.90	38.5-41.5	-4.381	5.8
	Zn	OP	ZncontOPP- A03	A03	69	6.35	68.5–69.5	-0.121	14.9
		LP	ZncontLPP- A02	A02	107	2.59	93.5–120.0	-0.073	8.6
			ZncontLPP- A03	A03	50	3.23	49.5–50.5	-0.072	7.0
Partitioning	В	OP	BpartOP-C04	C04	91	2.55	90.5–91.5	0.447	11.1
		LP	BpartLP-A04	A04	5	4.15	4.5-5.5	-0.532	8.9
			BpartLP-C04	C04	60	3.21	58.5-60.5	0.475	7.2
			BpartLP-C07	C07	93	3.71	92.5–93.0	0.506	8.1
	Ca	OP	CapartOP-A05	A05	89	3.07	88.5-89.5	0.383	7.0
			CapartOP-C02	C02	3	3.85	1.5-5.5	0.420	8.5
			CapartOP-C03	C03	120	5.89	118.5-120.5	0.521	12.9
			CapartOP-C05	C05	83	2.70	81.5-88.5	-0.349	5.8
			CapartOP-C09	C09	117	3.14	116.5-117.5	-0.373	6.7

	LP	CapartLP-A07	A07	88	3.97	87.5-88.5	0.576	9.6	
		CapartLP-A09	A09	61	4.49	60.5-61.5	0.617	11.0	
		CapartLP-C09	C09	117	3.36	116.5-117.5	-0.531	8.1	
Cu	LP	CupartLP-A05	A05	16	2.82	12.5-18.5	2.145	7.0	
		CupartLP-A09	A09	91	2.74	90.5–92.5	2.124	6.8	
Fe	OP	FepartOP-A01	A01	84	4.18	82.5-84.5	1.121	9.7	
		FepartOP-A04	A04	14	2.60	13.5-15.5	-0.873	6.1	
		FepartOP-C03	C03	85	4.77	83.5-85.5	1.184	11.2	
		FepartOP-C09	C09	117	2.75	116.5-117.5	-0.889	6.3	
Κ	OP	KpartOP-A02	A02	81	5.19	80.5-81.5	-0.744	7.5	
		KpartOP-A05	A05	40	5.88	37.5-40.5	-0.730	8.4	
		KpartOP-A09	A09	40	2.89	38.5-42.5	0.538	4.5	
		KpartOP-C03	C03	137	6.99	133.5-142.0	0.818	10.6	
	LP	KpartLP-A04	A04	8	3.86	7.5-8.5	-0.809	11.4	
Mg	OP	MgpartOP-A07	A07	69	3.20	68.5-70.5	-0.468	6.4	
		MgpartOP-C04	C04	54	4.35	50.5-54.5	0.551	8.9	
	LP	MgpartLP-A07	A07	53	3.58	52.5-53.5	-0.551	6.5	
		MgpartLP-C04	C04	55	5.12	54.5-55.5	0.665	9.6	
Mn	OP	MnpartOP-A09	A09	79	4.50	77.5-81.5	0.515	9.7	
		MnpartOP-C02	C02	16	2.98	15.5–19.5	0.411	6.2	
		MnpartOP-C03	C03	120	3.72	118.5-120.5	0.463	7.7	
		MnpartOP-C05	C05	91	4.87	90.5-102.5	-0.537	10.4	
Na	OP	NapartOP-A04	A04	58	5.31	54.5-58.5	-0.832	14.5	
		NapartOP-A09	A09	54	4.39	52.5-54.5	0.750	11.7	
	LP	NapartLP-A04	A04	31	7.29	30.5-31.5	-0.833	10.9	
		NapartLP-A05	A05	12	2.85	10.5-12.5	0.506	4.0	
		NapartLP-A09	A09	121	3.72	120.5-122.5	0.584	5.3	
		NapartLP-C03	C03	12	4.16	10.5-12.5	-0.710	6.1	
Р	OP	PpartOP-A03	A03	76	2.92	75.5-76.5	0.341	7.6	
			PpartOP-C04	C04	91	3.99	90.5–91.5	0.393	10.3
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	S	OP	SpartOP-A04	A04	8	11.12	7.5-8.5	-1.253	20.8
			SpartOP-A09	A09	45	3.46	44.5-45.5	0.650	5.7
			SpartOP-A10	A10	74	5.02	72.5-75.0	0.893	8.3
			SpartOP-C02	C02	45	3.16	43.5-46.5	-0.628	5.3
		LP	SpartLP-A04	A04	7	11.63	6.5-7.5	-1.464	24.9
			SpartLP-A05	A05	112	2.75	110.5-112.0	0.658	5.1
			SpartLP-C09	C09	14	4.45	10.5-17.5	-0.852	8.5
	Zn	OP	ZnpartOP-A03	A03	69	4.83	68.5–69.5	1.286	8.7
			ZnpartOP-A04a	A04	8	6.19	7.5-8.5	-1.424	11.6
			ZnpartOP-A04b	A04	65	6.33	64.5-66.5	-1.423	11.7
			ZnpartOP-C04a	C04	5	4.75	3.5-7.5	1.256	9.1
			ZnpartOP-C04b	C04	91	3.58	90.5-91.5	1.050	6.3
		LP	ZnpartLP-A04	A04	13	8.69	11.5–13.5	-1.134	18.7
			ZnpartLP-C08	C08	31	3.41	27.5-31.5	-0.705	7.2
			ZnpartLP-C09	C09	127	3.14	125.5-141.0	-0.668	6.5
Shoot dry weight		OP	SDWOP-A02	A02	75	2.98	74.5-75.5	-0.374	6.7
			SDWOP-A03	A03	51	4.81	50.5-51.5	-0.451	10.7
			SDWOP-A04	A04	3	4.06	2.5-3.5	-0.389	8.9
		LP	SDWLP-A02	A02	103	2.63	86.5-117.5	-0.328	8.3
			SDWLP-A03	A03	52	3.31	51.5-52.5	-0.324	6.8
			SDWLP-A04	A04	9	4.73	8.5–9.5	-0.354	9.7
Root dry weight		LP	RDWLP-A03	A03	52	3.01	51.5-52.5	-0.070	8.5
Total dry weight		OP	TDWOP-A02	A02	73	2.89	72.5-73.5	-0.420	6.0
			TDWOP-A03	A03	51	4.79	50.5-51.5	-0.536	10.3
			TDWOP-A04	A04	3	3.45	2.5–4.5	-0.425	7.3
		LP	TDWLP-A02	A02	72	2.53	71.5-72.5	-0.324	4.8
			TDWLP-A03	A03	52	4.94	51.5-52.5	-0.465	10.2
			TDWLP-A04	A04	9	4.25	8.5-9.5	-0.388	8.4

Note: each QTL for shoot and root concentrations of eleven mineral elements was denominated as "mineral element + conc (abbreviation of concentration) + P treatment + S (abbreviation of shoot) or R (abbreviation of root) + chromosome + the serial letter". Each QTL for shoot, root and plant contents of eleven mineral elements was denominated as "mineral element + cont (abbreviation of content) + P treatment + S (abbreviation of shoot) or R (abbreviation of plant) + chromosome + the serial letter". Each QTL for partitioning of eleven mineral elements was denominated as "mineral element + chromosome + the serial letter". Each QTL for partitioning of eleven mineral elements was denominated as "mineral element + part (abbreviation of partitioning) + P treatment + chromosome + the serial letter". Each QTL for biomass traits was denominated as "trait + P treatment + chromosome". A positive additive effect indicates a positive contribution of the Tapidor allele to the trait value, and a negative additive effect indicates a positive contribution of the Ningyou 7 allele to the trait value. R₂, the explained phenotypic variation.

Trait	Mineral element	P treatm ent	Chromo some1	Position 1 (cM)	LeftM arker1	RightM arker1	Chromo some2	Position 2 (cM)	LeftM arker2	RightM arker2	LOD score	R2 (%)	Ad d1	Ad d2	Add by Add
Shoot concentrati on	Cu	OP	A10	35	C10M 37	C10M3 8	C04	40	C14M 26	C14M2 7	5.03	13. 6	0.1 04	- 0.2 01	-0.456
			A09	45	C9M2 4	C9M25	C09	70	C19M 43	C19M4 4	5.26	15. 0	- 0.1 85	0.0 88	-0.452
	Fe	OP	A02	25	C2M1 8	C2M19	A05	30	C5M2 1	C5M22	5.32	11. 3	0.2 84	0.3 36	5.171
			A06	20	C6M1 5	C6M16	A06	35	C6M3 1	C6M32	5.26	14. 5	7.0 53	- 5.8 32	-7.992
			A05	35	C5M2 7	C5M28	A07	70	C7M9 4	C7M95	5.12	11. 0	0.3 87	- 1.3 63	-5.010
			C04	40	C14M 26	C14M2 7	C07	0	C17M 1	C17M2	5.54	11. 9	0.8 05	1.2 04	5.262
		LP	A02	70	C2M8 0	C2M81	A02	85	C2M1 04	C2M10 5	5.13	17. 9	3.3 84	- 8.3 49	-19.18
	K	LP	A01	15	C1M6	C1M7	A09	30	C9M1 7	C9M18	5.37	10. 5	- 305 .9	- 98. 64	-1588

Table 4. Epistatic loci for shoot and root concentrations, shoot, root and plant contents, and partitioning to the shoot of eleven mineral elements, and total dry weight in the *Bna*TNDH population at an optimal (OP) and a low P supply (LP)

	Р	OP	A08	30	C8M2 2	C8M23	A09	75	C9M7 1	C9M72	6.26	16. 1	- 77. 59	100 .9	-529.0
Root concentrati on	S	LP	A06	30	C6M2 4	C6M25	C06	95	C16M 74	C16M7 5	5.00	8.0	- 27. 88	47. 10	-494.7
	Zn	LP	A02	70	C2M8 0	C2M81	A02	95	C2M1 06	C2M10 7	7.39	17. 0	11. 02	- 11. 09	-14.18
	В	LP	A03	130	C3M1 55	C3M15 6	A06	15	C6M6	C6M7	5.25	13. 1	- 0.3 10	- 0.1 19	1.428
	Mg	LP	A02	80	C2M1 00	C2M10 1	C07	0	C17M 1	C17M2	6.03	11. 2	10. 85	- 69. 45	-96.72
			A05	0	C5M1	C5M2	C09	60	C19M 31	C19M3 2	5.51	10. 7	7.8 97	- 27. 76	90.33
	Mn	OP	A04	5	C4M7	C4M8	A08	0	C8M1	C8M2	5.31	24. 4	0.4 56	- 4.3 38	5.191
	Zn	OP	A03	60	C3M5 4	C3M55	A10	45	C10M 54	C10M5 5	5.01	23. 0	11. 13	- 4.2 30	21.96
		LP	A06	115	C6M1 37	C6M13 8	A06	120	C6M1 37	C6M13 8	7.32	6.8	- 22. 19	20. 13	-22.93

Shoot content	Cu	LP	A05	70	C5M7 2	C5M73	C05	50	C15M 25	C15M2 6	5.13	20. 7	- 0.0 01	0.0 02	0.003
	K	LP	A08	20	C8M1 0	C8M11	C05	70	C15M 44	C15M4 5	5.09	15. 8	- 9.7 59	- 2.0 21	-23.88
	Mg	LP	C06	70	C16M 54	C16M5 5	C07	90	C17M 77	C17M7 8	5.54	16. 2	- 0.0 94	0.3 16	1.348
	Mn	OP	A06	65	C6M8 1	C6M82	A10	10	C10M 5	C10M6	5.12	12. 3	0.0 16	0.0 10	-0.126
		LP	A01	40	C1M7 0	C1M71	C05	110	C15M 54	C15M5 5	6.32	17. 4	0.0 03	- 0.0 06	0.124
	S	OP	A05	25	C5M1 6	C5M17	C04	25	C14M 13	C14M1 4	5.35	18. 6	- 2.4 48	- 0.1 75	-4.998
			A09	120	C9M1 32	C9M13 3	C08	50	C18M 37	C18M3 8	5.37	15. 7	- 1.2 88	- 0.7 71	4.806
	Zn	LP	C05	40	C15M 19	C15M2 0	C05	70	C15M 44	C15M4 5	5.31	17. 4	- 0.0 45	0.0 23	-0.080
Root content	В	OP	A10	40	C10M 46	C10M4 7	A10	45	C10M 54	C10M5 5	5.59	12. 0	- 0.0 12	0.0 13	-0.012
			C09	10	C19M 3	C19M4	C09	15	C19M 4	C19M5	5.45	13. 4	- 0.0 11	0.0 13	-0.013

	LP	A03	0	C3M1	C3M2	A08	5	C8M2	C8M3	5.05	14. 4	0.0 01	$\begin{array}{c} 0.0\\ 00 \end{array}$	0.004
Cu	OP	A03	40	C3M1 2	C3M13	A03	45	C3M1 5	C3M16	5.22	13. 6	0.0 03	- 0.0 03	-0.003
K	OP	A10	40	C10M 46	C10M4 7	A10	45	C10M 54	C10M5 5	5.94	7.6	- 17. 14	16. 64	-15.09
		C03	75	C13M 73	C13M7 4	C03	80	C13M 74	C13M7 5	5.06	8.1	- 13. 40	14. 50	-18.26
		C05	35	C15M 18	C15M1 9	C05	40	C15M 19	C15M2 0	5.36	8.8	- 12. 53	9.9 06	-19.37
		C09	10	C19M 3	C19M4	C09	15	C19M 4	C19M5	6.44	8.8	- 13. 73	16. 80	-16.69
Mn	OP	A10	40	C10M 46	C10M4 7	A10	45	C10M 54	C10M5 5	5.32	11. 1	- 0.0 56	0.0 55	-0.053
		C09	10	C19M 3	C19M4	C09	15	C19M 4	C19M5	5.57	13. 2	- 0.0 45	0.0 55	-0.058
Na	OP	C07	50	C17M 37	C17M3 8	C08	50	C18M 37	C18M3 8	5.05	16. 4	- 0.1 22	- 0.1 43	0.247
S	OP	A02	30	C2M2 0	C2M21	A02	45	C2M2 6	C2M27	5.14	11. 7	- 0.8 57	1.1 39	-1.945

	Zn	OP	A05	35	C5M2 7	C5M28	A05	40	C5M3 2	C5M33	6.54	7.4	0.0 86	0.0 88	-0.106
			C09	10	C19M 3	C19M4	C09	15	C19M 4	C19M5	5.67	8.1	- 0.0 82	0.0 85	-0.110
Plant content	В	OP	A01	25	C1M2 3	C1M24	A05	110	C5M1 03	C5M10 4	5.08	19. 0	0.0 08	- 0.0 01	0.027
	Fe	LP	C03	140	C13M 126	C13M1 27	C06	75	C16M 59	C16M6 0	5.26	15. 6	- 0.1 70	0.0 71	-0.351
	K	LP	A03	0	C3M1	C3M2	A08	20	C8M1 0	C8M11	5.10	16. 5	- 1.9 78	- 8.1 71	26.65
			A08	20	C8M1 0	C8M11	C05	70	C15M 44	C15M4 5	5.14	17. 4	- 10. 22	- 2.1 31	-26.73
	Mn	LP	A03	0	C3M1	C3M2	A08	5	C8M2	C8M3	5.76	15. 8	0.0 04	- 0.0 22	0.134
			A01	40	C1M7 0	C1M71	C05	115	C15M 55	C15M5 6	5.04	13. 1	- 0.0 10	0.0 11	0.123
	Na	LP	A08	20	C8M1 0	C8M11	C05	70	C15M 44	C15M4 5	5.74	17. 0	- 0.3 20	- 0.1 36	-1.261

	Zn	LP	C05	35	C15M 18	C15M1 9	C05	70	C15M 44	C15M4 5	5.38	15. 2	0.0 55	0.0 24	-0.092
Partitionin g	Mg	OP	C08	55	C18M 46	C18M4 7	C08	100	C18M 99	C18M1 00	5.29	11. 3	0.2 99	0.0 30	0.618
	Mn	LP	A06	100	C6M1 28	C6M12 9	C05	40	C15M 19	C15M2 0	5.41	13. 8	0.0 39	0.1 98	-0.685
	Na	OP	A01	0	C1M1	C1M2	A05	10	C5M3	C5M4	6.05	11. 7	- 0.0 13	0.1 11	-0.814
			A06	95	C6M1 24	C6M12 5	A06	120	C6M1 37	C6M13 8	5.16	11. 8	0.1 28	- 0.5 48	0.930
	Zn	ОР	A03	70	C3M7 2	C3M73	A10	55	C10M 67	C10M6 8	5.25	18. 1	- 0.8 27	0.8 84	-1.483
Total dry weight		OP	A06	25	C6M1 9	C6M20	C03	90	C13M 80	C13M8 1	5.10	7.8	- 0.0 63	0.1 15	-0.479
		LP	A08	30	C8M2 2	C8M23	C03	5	C13M 4	C13M5	5.43	11. 0	- 0.4 76	- 0.2 37	0.518

Note: a positive value of Add by Add indicates that two loci genotypes being the same as those in parent Tapidor (or Ningyou 7) take the positive effects, while the two-loci recombinants take the negative effects. The case of negative values is just the opposite. R₂, the explained phenotypic variation.