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Heat Stress Affects Pi-related Genes Expression and Inorganic Phosphate Deposition/Accumulation in Barley

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Phosphorus (P) in plants is taken from soil as an inorganic phosphate (Pi) and is one of the most important macroelements in growth and development. Plants actively react to Pi starvation by the induced expression of Pi transporters, MIR399, MIR827, and miR399 molecular sponge – IPS1 genes and by the decreased expression of the ubiquitin-conjugating enzyme E2 (PHOSPHATE2 – PHO2) and Pi sensing and transport SPX-MFS genes. The PHO2 protein is involved in the degradation of Pi transporters PHT1;1 (from soil to roots) and PHO1 (from roots to shoots). The decreased expression of PHO2 leads to Pi accumulation in shoots. In contrast, the pho1 mutant shows a decreased level of Pi concentration in shoots. Finally, Pi starvation leads to decreased Pi concentration in all plant tissues. Little is known about plant Pi homeostasis in other abiotic stress conditions. We found that, during the first hour of heat stress, Pi accumulated in barley shoots but not in the roots, and transcriptomic data analysis as well as RT-qPCR led us to propose an explanation for this phenomenon. Pi transport inhibition from soil to roots is balanced by lower Pi efflux from roots to shoots directed by the PHO1 transporter. In shoots, the PHO2 mRNA level is decreased, leading to an increased Pi level. We concluded that Pi homeostasis in barley during heat stress is maintained by dynamic changes in Pi-related genes expression.

Keywords: barley, Pi transporters, PHO2, phosphate, high temperature, abiotic stress

INTRODUCTION

Phosphorus (P) is an essential macronutrient for plant growth and development. It is a component of vital molecules such as ATP, DNA, RNA, phospholipids, and phosphorylated sugars, making it crucial to cellular metabolism (Huang et al., 2008), energy conversion, and cell structure. Plants take up P solely as inorganic phosphate ions (Pi) from low concentrations in the soil solution. Most soil P is adsorbed, immobilized in organic matter, or precipitated as minerals (Poirier and Bucher, 2002).

Plants have a range of strategies to cope with a limiting Pi supply, including (i) favoring root growth over shoot growth, (ii) increasing the activity of high-affinity Pi transporters, (iii) exuding protons and organic anions to liberate Pi in the rhizosphere, and (iv) secreting acid phosphatases and ribonucleases to release Pi from organic compounds (Nilsson et al., 2007). Phosphate-deficient...
plants mobilize Pi from different subcellular compartments and organs and increase their Pi use efficiency by promoting metabolic bypasses that favor Pi-releasing reactions (Nilsson et al., 2007; Huang et al., 2008).

The uptake and redistribution of Pi is mediated by many different Pi transport proteins. These transporters have been classified into four families (dependent on their cellular location): PHT1 (plasma membrane), PHT2 (plastid inner envelope), PHT3 (mitochondrial inner membrane), and PHT4 (plastid inner envelope) (Karandashov and Bucher, 2005; Guo et al., 2008). The main role in Pi transportation from soil to plant roots is played by proteins belonging to the PHT1 Pi transporter family. Ten PHT1 genes have been identified in barley, and some of them have almost identical sequences; i.e., HvPHT1;1, and its paralogs PHT1;2, PHT1;9, and PHT1;10. These paralogs are expressed only in roots and are highly responsive to Pi limitation (Huang et al., 2011). HvPht1;6 is expressed in roots and in the older leaves of shoots (Preuss et al., 2010). The HvPHT1;8 gene is activated in the presence of arbuscular mycorrhizal fungi (Christophersen et al., 2009). In Arabidopsis PHT1;1 and PHT1;4, members of the PHT1 Pi transporter family play a central role in Pi acquisition (Shin et al., 2004). Another Pi transporter gene (PHO1) is primarily expressed in the root stelar cells and is involved in Pi loading into the xylem (Liu et al., 2014; Wege et al., 2016), and the pho1 mutant shows a low Pi level in shoots (Liu et al., 2014). Rice PHO1;2 is involved in the long-distance transport of Pi from roots to shoots (Wang et al., 2015). SPX-MFS proteins are, in turn, involved in Pi sensing and transport. In rice, OsSPX-MFS1 and OsSPX-MFS2 are targeted by Pi-responsive miR827 (Wang et al., 2012). This microRNA in Arabidopsis targets NLA (Nitrogen Limitation Adaptation) mRNAs that encode E3 ubiquitin ligase. Recently, it was shown that NLA protein works together with PHO2 (PHOSPHATE2, ubiquitin-conjugating enzyme E2), and they are both responsible for PHT1;4 degradation via the 26S proteasome (Park et al., 2014). PHO2 is involved in defining cellular Pi homeostasis and is responsible for the ubiquitin transfer to the protein substrate via the E3 enzyme (ubiquitin-protein ligase) (Smalle and Vierstra, 2004). The PHO2 protein itself is involved in the degradation of the PHT1;1, PHT1;2, PHT1;3, PHT1;4, PHO1, and PHF1 (PHOSPHATE TRANSPORTER TRAFFIC FACILITATOR1) (Liu et al., 2012; Huang et al., 2013). Thus, it is a general negative regulator of Pi uptake. The response of plants to Pi deprivation requires specific sensing mechanisms to monitor Pi status as well as signaling mechanisms leading to transcriptional and post-transcriptional regulation. PHR1 from Arabidopsis thaliana is a MYB-related transcription factor (TF) that regulates the expression of Pi-starvation induced genes through binding to the PiBS (PHR1-binding sequence) sequence, a DNA motif in the promoters of many Pi-related genes (Bari et al., 2006; Sobkowiak et al., 2012). There are three PiBS-like elements within the barley PHT1;1 promoter that are potentially recognized by PHR1 (Schunmann et al., 2004a). Moreover, in the Arabidopsis IPS1 promoter, there are two PiBS motifs (Bustos et al., 2010). In rice, PHR1 and PHR2 (an ortholog of AtPHR1) are involved in Pi-related gene regulation (Wang et al., 2014). Although AtPHR1 and OsPHR2 regulate the expression of Pi-related genes, they are rather stably expressed and do not belong to the group of Pi-stress responsive genes (Wang et al., 2014). The Pi homeostasis regulatory system also comprises the actions of microRNA399 and microRNA827 (Bari et al., 2006; Hackenberg et al., 2013). miR399 guides the RISC complex to target PHO2 mRNA. Furthermore, a riboregulator IPS1 can bind and quench miR399 activity because it is not cleaved by the miRNA due to the base pair mismatches, which prevents PHO2 mRNA against complete degradation (Frances-Zorrilla et al., 2007). Also, genes such as Arabidopsis SIZ1 (SIZ/PIAS-type SUMO) (Small Ubiquitin-related Modifier) E3 ligase) are involved in Pi response (Miura et al., 2005). Rice mutant OsSIZ1 shows an increase of Pi concentration in both root and shoot tissue (Wang et al., 2015). In Arabidopsis, SIZ1 is responsible for the SUMOylation of PHR1 (Miura et al., 2005). Although OsSIZ1 itself is not very responsive to Pi starvation, it regulates the expression of other Pi-related genes, like MIR399a, PHO2, PHF2, or Pi transporters (Wang et al., 2015).

Heat stress has a big impact on plant growth and development. High temperature affects flowering time, reduces number, and leads to a reduction of size as well as an increase of deformity in floral organs (Zinn et al., 2010). Heat stress is also responsible for changing gene expression, including microRNAs and (as a consequence) their target mRNA levels. Barley microRNAs like miR160a, 166a, 167b, and 5175a are upregulated in heat stress, and consequently, the appropriate target genes are downregulated (Kruszka et al., 2014).

Transcriptomic data analysis derived from 1-h heat-stressed plants shows dramatic changes in Pi-related gene expression in roots as compared to unstressed controls. Based on their expression changes, a network of particular gene expression regulation and its influence on Pi homeostasis can be proposed. The expression of the HvPHT1;1, HvPHT1;4, and HvPHT1;6 genes is downregulated in roots, which therefore should have decreased Pi concentrations. However, this is not the case; possibly, because the expression of the PHO1 gene is concurrently decreased in the root. A lower PHO1 level can lead to lower Pi efflux from roots to shoots. Pi concentration in roots (in control and in heat treated plants) was always on the same level with one exception at 4 h time-point. The observed increased Pi concentration was correlated with significant PHT1;4 up-regulation. Moreover, transcriptomic data reveals that PHO2 and SPX-MFS are downregulated in roots, but not PHR1, PHR2, nor SIZ1. In heat treated shoots at two time-points: 1 h and 12 h we observed increased Pi concentration. It was correlated with PHO2 and PHT1;6 genes expression profile changes. At these time-points, the barley shoots showed increased Pi concentration; this agrees with previous data showing that PHO2 silencing leads to an increase of Pi concentration in barley shoots (Pack et al., 2010). Since we detected neither mature miR399 nor significant changes in the expression of mature miR827 in barley shoots, we concluded that the changes observed resulted from transcriptional regulation. The transcriptomic data was mostly confirmed by the RT-qPCR results. The changes in Pi concentration in barley shoots at 1 and 12 h heat time-points are abolished in other hours of heat stress, leading
to Pi-concentration stability. Summarizing, the changes in expression of Pi-related genes after heat stress are not chaotic but strictly controlled, regarding what is necessary for the plant to maintain Pi homeostasis. During Pi re-supply, in barley roots we observed significant downregulation of the PHT1;1, PHT1;4, PHT1;6, PHO1, PHO2 but not PHR1, SPX-MFS. In contrast to heat stress condition, at 1 h after Pi re-supply we observed increased expression of PHO1 in Pi-resupplied plants compared to the control ones. We concluded that fine-tuning changes of PHO1 expression in barley roots prevents Pi over-accumulation.

MATERIALS AND METHODS

Plant Growth Conditions

In the heat experiment, barley plants cultivar Rolap were grown under conditions as previously described (Kruszka et al., 2013), and the heat stress conditions were established also as previously described (Kruszka et al., 2014) – except that the soil water content was kept at 70% to prevent drought stress. The soil/send mixture (7:2) was amended with 35 ml of a medium containing: 28 mM NH₄NO₃, 20 mM KH₂PO₄, 4 mM K₂SO₄, 16 mM MgSO₄•7H₂O, 53 μM H₂BO₃, 8 μM CuSO₄, 4 μM MnSO₄•H₂O, and 120 μM FeCl₃•6H₂O. Two-week-old barley plants exposed 48 hours to heat stress were harvested at 1, 3, 4, 5, 6, 7, 8, 12, 24, and 48 h after the initiation of 35.5°C heat stress. Unstressed control plants were also grown (at 22°C). In the Pi stress experiment, barley plants cultivars Rolap and Black Hulless were grown at different Pi regimes in a 7:2 (cv. Rolap) and 1:1 (cv. Black Hulless) (w/w) soil-sand mixture containing 8 mg Pi/kg soil (extraction by 0.5 M NaHCO₃ (Olsen et al., 1954)). The level of Pi in plant root and shoot samples was monitored sequentially as follows: P sufficiency (control), P limitation, P replenishment, 20 das. Each pot contained three barley plants.

Phosphate Measurement

The level of Pi in plant root and shoot samples was monitored by an inorganic P assay utilizing malachite-green/ammonium molybdate in 4 M HCl and 34% natrium citrate•2H₂O solutions (Lanzetta et al., 1979). Fifty milligram of liquid N₂ frozen plant tissue was homogenized in a glass homogenizer in 1 ml of ice-cold 1 M HCl. After centrifugation, the supernatants were used for a micro-plate assay at room temperature. The standard curve was constructed from a P-stock solution of 1 nmol/μl KH₂PO₄ in two replicates. The samples were measured in two technical and three biological replicates. The OD₆0₀ was read (Müller et al., 2004) using either a SpectraMax M5 Microplate Reader (Molecular Devices, LLC, USA) or Infinite F200 Pro (TECAN, Switzerland).

RNA Isolation

For the barley heat stress experiment, RNA isolation procedure was described previously (Kruszka et al., 2013, 2014). In brief, the RNA isolation method utilized a mixture containing a 38% v/v phenol solution saturated with 0.1 M sodium acetate (Roti Aqua Phenol, Roth, Karlsruhe, Germany), supplemented with 0.8 M guanidine thiocyanate, 0.4 M ammonium thiocyanate, 0.1 M sodium acetate, 5% v/v glycerol, 0.5% sodium lauroylsarcosine, and 5 mM EDTA. To remove polysaccharides, the Ambion Plant RNA Isolation Aid (Thermo Fisher Scientific, Lithuania) was used. RNA was isolated using TRIzol reagent (Thermo Fisher Scientific, Lithuania) according to the manufacturer’s instructions. The RNA yield was determined using the Nanodrop ND-1000 UV spectrophotometer (Thermo Fisher Scientific, Lithuania). The quality of RNA was checked using the Agilent 2100 Bioanalyzer (Agilent Technologies, Santa Clara, CA, USA) and the RNA integrity number (RIN) ranged from 8.0 to 10.0. The samples were stored at -80°C until use in the experiments.

**TABLE 1 | Sequences of used primers.**

| Name     | Sequence 5’ to 3’ | Task       |
|----------|------------------|------------|
| APO175   | GGGATAACACTGCTGTTGATGTC | 3’ RACE, PHR1 |
| APO176   | CATCCCTGTTGAGTGACCTGAC | 3’ RACE, PHR1 |
| APO177   | ATGAGTGAGCTGCTGTTGAC | 5’ RACE, PHR1 |
| APO178   | GAATTGGAAGCTGAACTGAGC | 5’ RACE, PHR1 |
| APO273   | CACATGACATTTGACAACACCTTA | 3’ RACE, PHO2 |
| APO274   | CACATCAGATCATGAGTGCTGCAAGAG | 3’ RACE, PHO2 |
| APOY11   | CTGCTGCAGTAAAGGAAGGAGCA | 5’ RACE, PHO2 |
| APOY22   | CTCCTCAAGAAGGAGGAGCA | 5’ RACE, PHO2 |
| APOY92   | TCTAATCAACGCGACAGAA | PCR, ADP |
| APO410   | CTGCAAGCCTGCAGAAG | PCR, ADP |
| APO623   | AAGCTGCAAGGCAAGGAGGAA | RT-qPCR, pri-miR399c |
| APO624   | CTGCCAAATAAAAGGAGGACCC | RT-qPCR, pri-miR399c |
| APO627   | CAGGCCAACACTCTCCTTGGCA | PCR, miR399c |
| APO697   | TGTTTGTTGACATGTACCTAA | Probe for miR827 |
| APO698   | GCAACCTCTTGCTTGCTGTT | RT-qPCR, Pri17/4 |
| APO699   | TTGCTGCTTGCTTGCTGTTGT | RT-qPCR, Pri17/4 |
| APO700   | CAAAGGCCAAGCAGAGTGA | RT-qPCR, PHR1 |
| APO701   | ATGACCTGAGGAGGAGGAGGA | RT-qPCR, PHR1 |
| APO702   | TTGAGAGCAAGGAGGAGGAGGA | RT-qPCR, NLA |
| APO703   | ACTCGCTCATCTGCACTTCT | RT-qPCR, NLA |
| APO704   | ATGAGTCTGACCTGACATCTG | RT-qPCR, SPX-MFS |
| APO705   | ACCAATGCTGAGGAGGAGCA | RT-qPCR, SPX-MFS |
| APO706   | Described by (Huang et al., 2011) | RT-qPCR, Pri17/6 |
| APO707   | Described by (Huang et al., 2011) | RT-qPCR, Pri17/6 |
| APO708   | Described by (Huang et al., 2011) | RT-qPCR, Pri17/1 |
| APO709   | Described by (Huang et al., 2011) | RT-qPCR, Pri17/1 |
| APO739   | ATGGGTGCAATCTCTGCTGAT | RT-qPCR, PHO1 |
| APO740   | CTGAGAAGAATCTGCTGACAG | RT-qPCR, PHO1 |
| APO741   | AACTCATGAGGCTCTTGTTG | RT-qPCR, PHR2 |
| APO742   | TGGGCAATCTGCTGATCTT | RT-qPCR, PHR2 |
| APO743   | AGATGCTCTGCTGCTTGTT | RT-qPCR, S21 |
| APO744   | CAGGGCTCTCCTTCTCTCT | RT-qPCR, S21 |
added, with the resulting supernatant being purified using a Direct-zol™ RNA MiniPrep kit according to the manufacturer’s protocol (ZYMO RESEARCH, The Epigenetics Company, USA).

**cDNA Synthesis and PCR Reactions**
cDNAs used in RT-PCR and RT-qPCR analysis were synthesized using an oligo(dT)$_{18}$ primer, SuperScript III Reverse Transcriptase (Invitrogen, USA) and 3 µg of Turbo DNAse (Ambion, Lithuania) treated RNA as a template. cDNAs were diluted four times, and 1 µl was used in RT-qPCR and diluted five times, with 2 µl used for RT–PCR amplification. PCR amplification of either an ubiquitin or ADP cDNA fragment were used as a positive control and control of the sample loading. The purity of cDNA samples containing no genomic DNA was controlled by PCR amplification of a barley HvPHT1;1 GenBank accession no. AF543197.1 promoter fragment (Schunmann et al., 2004b; Kruszka et al., 2014).

**PCR Amplification**
Primers for RT-qPCR amplification of HvPHT1;1/HvPHT1;9, PHR1, PHO2, IPS1, and ubiquitin were previously described (Pacak et al., 2010). The rest of the used primers are summarized in Table 1.

For RT-PCR, “Touchdown” PCR was performed (Tolia and Joshua-Tor, 2006). RT-qPCR was performed as previously described (Kruszka et al., 2013).

RACE and RLM-RACE were performed using a SMARTer™ RACE cDNA Amplification Kit (Clontech, Mountain View, CA, USA) and GeneRacer kit (Invitrogen), respectively (according to the manufacturers’ protocol).

For RT-qPCR analysis, an Applied Biosystems 7900HT Fast Real-Time PCR System and SYBR Green Master Mix (Applied Biosystems) were used. Data was normalized to ADP levels. The results were presented as either log$_{10}$($-\Delta$Ct) + 7 or as fold changes as previously described (Kruszka et al., 2013). The R$^2$ values of analyzed data (≥0.997) were calculated with LinRegPCR software (Ramakers et al., 2003). Three biological replications were analyzed.

**Northern Blot**
Experiments were performed using either 50 µg RNA enriched in small RNA isolated from barley shoots (Figure 6), according to (Szaryznska et al., 2009; Kruszka et al., 2013) or 7.5 µg RNA enriched in small RNA isolated from barley roots (Figure 10), according to (Pall and Hamilton, 2008).

**Transcriptome Analysis**
Total RNA was isolated using the procedure described above. Isolated RNA samples were Turbo DNase (Ambion) treated and then phenol/chloroform purified. Three biological replications of barley shoots and two biological replications of barley roots harvested after 1 h of heat treatment were analyzed (as well as the control samples). Transcriptome libraries (strand-specific) were constructed and then sequenced by BGI Tech Solutions (Hong Kong) Co., Ltd. Clean reads were mapped to the reference barley genome1 (Ensembl Plants, version: 82214v1) using TopHat2 software (Kim et al., 2013). Statistical analysis was performed to identify gene-expression differences between the samples studied (TEAM, 2010; Pinheiro et al., 2014). Numbers of reads for each gene mapped to the reference genome were counted using HTSeq software (Anders et al., 2015). The script was run for each fastq file for every biological replicate from every condition. Afterward, all of the results were gathered in tables – one table per one comparison (e.g., a table with two biological replicates from roots treated with high temperature and two replicates from control roots). These tables were then fed in a DeSeq – R package suitable for analysis of differential gene expression based on the negative binomial distribution (Anders and Huber, 2010). The data used in this study have been deposited under NCBI GEO accession GSE82134.

**Gene Identification**
Transcriptomic analysis met some difficulties caused by sequence similarities between gene paralogs. For instance, the MLOC_28370 gene encodes a 369AA protein identical

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**Table 2 | Analyzed Pi-related genes with their Ensembl Plants database numbers and characteristics.**

| Gene     | Ensembl Plants | Function                                      | Notes                                      |
|----------|----------------|----------------------------------------------|--------------------------------------------|
| PHT1;1   | MLOC_28370.1   | Pi transporter                               | four paralogs in barley                   |
| PHT1;4   | MLOC_6187      | Pi transporter                               |                                            |
| PHT1;6   | MLOC_80912.2   | Pi transporter                               |                                            |
| PHR1     | MLOC_5585      | TF, ortholog of OsPHR1                       |                                            |
| PHR2     | MLOC_60198.1   | TF, ortholog of OsPHR2                       |                                            |
| PHO1     | MLOC_12153.1   | Pi transporter, homologous to rice PHO1;2    |                                            |
| PHO2     | MLOC_53410.2   | ubiquitin-conjugating enzyme E2, catalytic (UBC9) domain | targeted by miR3999                        |
| IPS1     | not identified in Ensembl Plants | binds miR3999                         | position chr 4, 510585405-510586016       |
| NLA      | MLOC_52462.2   | E3 ubiquitin-protein ligase, SPX_BAH1-like, RING domains | no miR827 binding site in barley           |
| SPX-MFS  | MLOC_57566.4   | Pi sensing and transport SPX, MFS domains    | targeted by miR827                        |
| SIZ1     | MLOC_38182.4   | PHR1 sumoylation                             |                                            |
| Pre-miR399c | not identified in Ensembl Plants | miR399c targets PHO2                         | position chr 2, 558571927-558572025       |

Barley genes PHR1, PHR2, PHO1, NLA, SPX-MFS, SIZ1, and MIR399c were identified based on their similarity to known orthologs and then allocated to a particular MLOC number.

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1http://plants.ensembl.org/Hordeum_vulgare/Info/Index
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to HvPHT1;1 (GenBank AAN37900.1; AF543197), HvPHT1;2 (AAO72434.1, AY187020), and HvPHT1;9 (GenBank CAP17759; AM904733), both on nucleotide and amino-acid levels. Thus, transcriptomic reads reflect expression of these three Pi transporters. To distinguish the expression of the PHT1;1 gene from their paralogs particular Pi transporter, two sets of RT-qPCR primers were used for analysis. We analyzed both PHT1;1/PHT1;9, and concrete PHT1;1 gene expression. We identified the putative barley PHR1 gene by comparison with the well-known Psr1 (Phosphate starvation response 1) and PHR genes from Chlamydomonas reinhardtii, rice, and Arabidopsis. We found several barley ESTs (BU993345, BU970600 and BF621611). In further steps, these ESTs were clustered into one mRNA fragment. Based on this mRNA fragment sequence, a pair

![Figure 1](image-url)

**FIGURE 1** | (A) Neighbor-Joining phylogram of the PHR1/PHR2 transcription factor (TF) genes translated into amino acid sequences. Locus At5g06800 (protein – NP_196298.2); locus At5g29000 (PHR1-LIKE1 – PHL1, protein AAN86177.1); locus At2g20400 (protein – AAP04104.1); AtPHR1 – locus At4G28610 (protein – NP_194590); locus At1g79430 (protein – BAH194448); locus At3g13040 (protein – NP_974298); At3g04450 (protein – NP_187095.2); OsPHR1 – locus Os03g0329900 (protein – NP_001050006.1); OsPHR2 – locus Os07g0438800 (protein – BAG94425.1); CrPsr1 (nucleotide – XM_001700501, protein – XP_001700553); HvPHR1 (nucleotide – GQ337895, MLOC_5585, protein – ACT34981, MLOC_5585); HvPHR2 (MLOC_60198.1). Bootstrap value was calculated based on 100 replications; (B) Barley PHO2 gene structure. Exons, introns, CDS and UTRs are depicted by blue boxes, black lines, orange boxes, and dotted lines, respectively. Within exon 2, six miR399 binding sites are depicted; the red vertical line represents binding site no 6; (C) Alignment of six miR399 binding sites in HvPHO2 5′UTR (according to AK249253) and one binding site in HvIPS1 (GQ301528) recognized by miR399. Bold, black letters denote Hv-miR399c sequence. Sites 1, 2, 3, 4, 5, and 6 are numbered starting from the 5′ end of the barley PHO2 mRNA. Yellow arrows show cleavage position (Hackenberg et al., 2013). Site 6 (5′ CUUGGCACUUCUCUUGGCA 3′) has three mismatches to hvu-miR399d-3p (Hackenberg et al., 2013).
FIGURE 2 | Pi-related protein structures with depicted unique domains. PHT1 proteins possess MFS (Major Facilitator Superfamily) domain characteristic for the secondary membrane transporter family (Wang et al., 2012); PHR1 contains LHEQLE motif, myb-like DNA-binding, and SHAQKYF class domains. PHO1 protein possesses SPX and EXS domains (Erst1-Xpr1-Syg1), PHO2 contains UBCc domain (ubiquitin conjugating enzyme); NLA protein possesses additional SPX domain of the E3 ubiquitin-protein ligase BAH1/NLA and RING-finger (Really Interested New Gene) domain; SPX-MFS has SPX and MFS domains; SIZ1 has following domains: SAP (SAF-A/B, Acinus and PIAS), plant homeodomain (PHD) finger, and MIZ/SP-RING (Msx – Interacting – Zinc finger) domain. Protein structures were constructed based on the appropriate protein sequences, whose accession numbers are deposited in Table 2—with the exception of PHO1 (protein – BAJ96107) and SIZ1 (protein – BAJ97548). The black bar represents 100 amino-acids.

of primers was designed. The amplified PCR product (923 bp in length) – i.e., fragment of HvPHR1 – was sequenced as previously described (Pacak et al., 2010). To determine full-length HvPHR1, 5’ and 3’ RACE experiments were conducted. The coding sequence had 1356 bp and can encode a 451AA sequence that contains 99% identities to MLOC_5585. Barley PHO1 – MLOC_12153.1 contains an amino acid sequence with 298/477 (62%) and 432/475 (91%) identity (blastp analysis) to the Arabidopsis PHO1 protein (GenBank NP_188985.2) and Brachypodium distachyon phosphate transporter PHO1-2 (GenBank XP_003570364.1), respectively. We found contig TCONS_00089457 that contains the mature barley miR399c sequence. The pre-miR399c sequence was identified in the transcriptomic data derived from heat-treated shoot tissue – TCONS_00089457. This sequence was mapped to morex_contig_188578 CAJW010188578 carma = 2HL (ipk-gatersleben database). The longest derived sequence that formed a stem-loop structure was used for construction of the pre-miR399c. Using 5’ RLM-RACE, two TSS of the barley PHO2 gene were identified at positions 447916090 (+) and 447916332 (+) on the chromosome 1. Promoter region (1422 bp) upstream of the 2nd TSS (heterogenic) was used for promoter elements analysis. Later, a PHO2 genomic sequence was establish based on 5’ and 3’ RACE analysis and morex_contig_38340 CAJW010038340 carma = 1H DNA sequences deposited at the ipk-gatersleben database.

Software

The alignment and trees were constructed using CLC Main Workbench 7 software with the following parameters: (Kimura-2 parameters) and the consensus trees (neighbor-joining) with bootstrap value = 100 replications. Stem-loop structures were constructed using Folder Version 1.11 software with a RNAfold algorithm2. Significance of the results (p-value) was analyzed using student t-test. Statistical RT-qPCR analyses were performed using log10(2(−ΔΔCt)) + 7 values.

Database Searching

We searched the following databases: http://plants.ensembl.org/index.html, www.mirbase.org, and GenBank and IPK Gatersleben3. For promoter regulatory element analysis, a New PLACE (A Database of Plant cis-acting Regulatory DNA Elements) database was used (Higo et al., 1999).

Accession Numbers

Studied genes: Ensembl Plants database – Heat shock protein 90.1 – MLOC_5618, other accession numbers are deposited at Table 2. GenBank database: HvPHT1;1 – AF543197, AAN37900.1; HvPHT1;2 – AAO72434.1, AY187020; HvPHT1;4 – AY187024.1, AAO72437.1; HvPHT1;6 – FM866444.1, CAS02288; HvPHT1;9 – AM904733, CAP17759; HvPHR1 – GQ337895, ACT34981; HvPHR2 – AK363485.1, BAJ94688.1; HvPHO1 – AK364904.1, BAJ96107; HvPHO2 – AK249253 (cv. Haruna Nijo); HvPHO2 – GQ861514.1, ACV7226 (cv. Black Hulless); OsPHO2 – LOC_Os05g48390 (AU032431); AtpPHO2 – AT2G33770.1; HvNLA – AK354779, BAJ85998.1; HvIPS1 – GQ301528; HvSIZ1 – AK366345.1, BAJ97548; ADP – ADP-ribosylation factor 1-like protein gene – AJ508228.2. The use of the Hsp17 gene as a “heat” stress gene marker (GenBank: AK252765) was described previously (Kruszka et al., 2014).

RESULTS

Identification of Barley Pi-related Genes

GenBank and Ensembl Plants databases were searched to identify barley Pi-related genes. Several Pi-related barley genes like

2http://www.ncbi.nlm.nih.gov/rnafolder/rnafolder.php

3http://webblast.ipk-gatersleben.de/barley/
Pi transporters (PHT1;1, PHT1;4, PHT1;6) and PHR1, PHO2 were described previously (Schunmann et al., 2004a,b; Pacak et al., 2010; Preuss et al., 2010). Table 2 shows all barley genes identified by us as well as other Pi-related barley genes.

A phylogram was created to show the similarities of the identified barley PHR1/PHR2 TFs to the well-known PHR proteins from rice and Arabidopsis (Figure 1A) (Wang et al., 2014, 2015). Additionally, we have included the Arabidopsis

**FIGURE 3** | Barley pre-miRmiR399 structure. (A) Barley pre-miR399 stem-loop structure based on MI0017933 sequence – miRBase; (B) Alignment of the barley PHO2 5’ UTR and reverse complement miR399 related sequences. MI0017933 – miRBase, barley pre-miR399, MIMAT0020542 – miRBase barley microRNA399, fragment of TCONS_00089457 – Rolap, transcriptome identified pri-miR399c; (C) pre-miR399c stem-loop structure based on the TCONS_00089457 sequence (AGTCCAGTTTCAGGGCTCCTCTTTATTGGCAGGGAGCGTGTGAGGCCATGTAGCCTCATTCAGCGCTCTGCCAAAGGAGAGTTGCCCTGTAACTGGAAA).
PHR1-like protein (PHL1) and other MYB-CC proteins (Bustos et al., 2010).

Two splice variants of PHO2 barley mRNA encoding 847AA and 544AA long proteins are deposited in Ensembl Plants. We compared PHO2 CDS derived from MLOC_53410.2, with PHO2 CDS derived from cv. Haruna Nijo, cv. Rolap, and cv. Black Hulless, and found that CDS from Haruna Nijo, and Rolap were identical, differing from Black Hulless in seven nucleotide substitutions (three synonymous and four non-synonymous mutations). The rice PHO2 ortholog contains 11 exons and encodes an 876AA protein (Hu et al., 2011). In Arabidopsis, the PHO2 gene has nine exons and encodes 907AA. The barley PHO2 gene is located on chromosome 1, is 7320 bp in length, and contains ten exons (Figure 1B). The 5' UTR of barley PHO2 mRNA contains six potential miR399 recognition sites (Figure 1C). Only two of them were found to be cleaved in barley (Hackenberg et al., 2013). The schemes of the analyzed Pi-related proteins with depicted unique domains are presented in Figure 2.

Ten MIR399 genes and one MIR827 gene have been identified in the barley genome (Hackenberg et al., 2013). Both are induced by Pi deficiency (Hackenberg et al., 2013). In miRBase, there is only one pre-miR399 precursor available (M10017933) from which mature miR399 could be diced out. This proposed pre-miR399 stem-loop structure does not contain the typical features of plant pre-miRNA (Figure 3A) (Bologna et al., 2009). Since it is possible that the miR399-3p sequence is unusually processed in barley shoots (Hackenberg et al., 2013), we decided to analyze it in more depth. We found that this sequence is, in fact, the reverse complement fragment of the PHO2 5' UTR containing miR399 binding site no 2 (Figure 3B). We found a locus encoding the miR399c sequence and were able to fold it into a classical stem-loop structure with free energy $\Delta G = -69.10$ kcal/mol (Figure 3C).

Our bioinformatics search allowed us to identify all known important players involved in Pi homeostasis maintenance in barley plants.

**Heat Stress Affects the Expression of Pi-related Genes**

To test the heat-stress influence on Pi-related gene expression and Pi concentration, 2-week-old barley plants were stressed by exposure to a 35.5°C temperature for two days. Root and shoot samples were collected after 1, 3, 4, 5, 6, 7, 8, 12, 24, and 48 h.

Transcriptome deep sequencing was applied to RNA isolated from samples exposed to one hour of heat stress and from...
FIGURE 5 | Pi concentration. (A) in barley roots and (B) in shoots after heat stress as compared to control plants. Blue and red bars represent control and heat-treated samples. *p < 0.05, **p < 0.005 (t-student test). Pi concentration n = 3, two technical replications.
un-stressed control plants. Subsequently, our analysis was focused on the expression of Pi homeostasis maintenance-related genes. After one hour of heat stress, the expression of PHT1;1, PHT1;4, PHT1;6, PHO1, PHO2, and SPX-MFS genes was decreased in roots. Only the NLA gene showed an upregulated expression in the barley roots. The expression pattern of the PHR1, PHR2, and SIZ1 genes was not significantly influenced. In contrast, the expression of most analyzed genes in shoots remained unchanged after one hour of heat stress. The only downregulated Pi-related gene was PHO2 (Figure 4). The changes revealed in the transcriptomic data were confirmed by RT-qPCR (Supplementary Figures S1B–F,H–O,Q,R). The downregulated expression of genes in roots appeared to be permanent, as the SPX-MFS and PHO2 genes remained downregulated after 48 h of heat stress. Since PHT1;1 and PHT1;9 CDSs have identical sequences, RT-qPCR designed for their 3′ UTR fragment amplification was performed, which allowed us to discriminate between the expression of these two genes. The data shows that both PHT1;1/PHT1;9 (and also the PHT1;6, PHO1, PHR2, and SIZ1 genes) stay at the lower expression level as compared to control plants. The PHT1;4 gene shows a more-complex expression pattern; it was downregulated at 1 and 3 h, but up-regulation between 4 and 12 h of heat duration was also observed (Supplementary Figure S1C). In roots at 4 h time-point we observed the increased Pi concentration (Figure 5A). This correlated with PHT1;4 significant up-regulation at this time-point. Since at 6 h time-point also PHT1;4 gene expression was higher in heat stressed plants compared to the control ones, other factors could contribute to a mechanism of keeping the Pi concentration on the same level. The NLA gene was always upregulated, with the highest expression level at 24 h (Supplementary Figure S1H). Surprisingly, IPS1 was also upregulated after 6 h of heat stress, with the highest expression level at 48 h. The expression of PHO2 expression in roots and shoots was always downregulated, with the highest decrease at 12 h and 24 hour in roots and shoots. The PHT1;6 expression in shoots shows the highest decrease in expression at the 12 h time point (Supplementary Figure S1M). In the case of other shoot-expressed Pi-related genes, the expression remains almost at the same level. The only exception is present at the 48 h time point for NLA, SPX-MFS, and PHR1 (also 12 h) genes, which are downregulated. IPS1 expression is almost at the same level during the first eight hours but later decreased. Transcriptomic data revealed that in shoots there were no PHT1;1, PHT1;4 genes expression; PHO1 expression was also very low compared to the root expression. That is why even subtle changes in PHO2, PHT1;6 genes expression could influence Pi concentration. At 1 and 12 h time-points we observed that both these genes had been affected by heat stress. At 1 h time-point transcriptomic data showed significant decreased expression of PHO2. RT-qPCR confirmed the transcriptomic data but probably the fold change was too low to be significant in RT-qPCR analysis. Interestingly RT-qPCR showed significant increased PHT1;6 gene expression and it could be coupled with an increased Pi level (Figure 5B). At 12 h time-point we observed significant decreased expression of PHO2 and PHT1;6. Previously we noted that decreased level of PHO2 gene in barley led to an increase of Pi concentration (Pacak et al., 2010). Further studies on protein level as well as identification of the PHO2 "partner proteins" in shoots are necessary to reveal the role of PHT1;6 in Pi concentration homeostasis in shoots. We were not able to detect mature miR399 in barley shoots grown in heat stress and control conditions (Figure 6A). The expression of the miR827 fluctuates in control conditions, and these fluctuations are also observed in heat-treated plants (Figure 6B).
draw the conclusion that the changes in PHO2 and SPX-MFS gene expression in barley shoots (their mRNAs are targeted by miR399 and miR827, respectively) result from transcriptional rather than post-transcriptional regulation. Surprisingly, the observed gene-expression changes in roots do not affect the final root Pi concentration. We noted a significant Pi concentration increase at the 1- and 12-h time-points of heat stress in barley shoots, but not at other time-points (Figures 5A,B). This data shows that barley plants exposed to heat stress modulate the expression level of Pi-related genes to stabilize Pi concentration. Although the Pi concentration at the end of 48 hours of heat stress was similar to that of the control plants, heat stress inhibited barley growth and leaves no 4 development (Figures 7A and 8). The leaves of barley plants are shorter one week after the end of the heat stress period (leaf number four), and we observed faster leaf senescence (leaf number one, Figure 8). Growth impairment and premature senescence may result from the macronutrient homeostasis disturbances resulting from heat-stress-affected gene-expression regulation.

Dynamic Changes in Expression Pattern of Barley Pi-Related Genes upon Phosphate Starvation and Pi Re-supply

Analysis of Pi-related barley gene expression under heat stress revealed fast and dramatic changes at the transcriptional level, which finally led to the stabilization of Pi concentration in the plant tissues. To learn more about the regulation and pattern of expression of Pi-related genes in barley, we studied the same set of genes for their expression pattern in roots under control conditions, Pi starvation, and Pi re-supply. Plants were grown at P-limiting conditions and then exposed to P sufficiency. Time-course changes in gene expression were compared to the changes
in parallel sets of plants maintained at P-limiting or P-sufficient conditions (control conditions).

In 2-week old barley cv. Rolap PHT1;1, PHT1;4, and PHT1;6 showed increased expression under low Pi conditions compared to the control conditions. Pi-resupply down-regulated Pi transporters expression what was observed after 3 h of the Pi re-supply (Figure 9). Especially PHT1;6 gene was sensitive to the Pi re-supply. Its expression after 48 h was even lower in Pi re-supplied plants compared to the control ones (Figure 9C). After 48 h of the Pi-resupply PHT1;1 gene expression was not distinguishable from plants grown under low Pi. During heat stress we observed that PHT1;1 gene was the most affected PHT1 transporter gene by heat stress. It showed plant flexibility in gene expression regulation upon different stresses in order to control Pi homeostasis. Among two genes which mRNA levels are controlled by microRNA, SPX-MFS showed higher sensitivity to two abiotic stresses applied. It was constantly downregulated during heat stress and up-regulated after Pi-resupply at 3 h and 48 h time-points (Figure 9G). PHO2 expression was even lower in Pi re-supplied plants compared to the control ones (Figure 9F). It showed that in the experimental conditions Pi did not influence PHR1. Compared to heat stress we observed different PHO1 expression change (Figure 9E). During heat stress at 1 h time point PHO1 expression was down-regulated but after Pi re-supply its expression was even increased compared to the control plants. It could be explained by the fact that plants produced more PHO1 protein to transport available Pi from roots to shoots. After 12 h PHO1 expression in Pi re-supplied plants reached the control level. The weight of barley plants from low Pi conditions was 62% of the control plants weight. The phenotypes of the Pi control, Pi limitation, Pi re-supplied plants are presented in Figure 7B.

In our previous work we silenced PHO2 gene using VIGS approach and observed Pi concentration increase in barley shoots and the PHO2 downregulation in barley roots (Pacak et al., 2010). For this purpose we used Black Hulless barley cultivar. We used the same cultivar to analyze Pi-related gene expression under different Pi regime. Moreover we kept plants longer (23 days) in low Pi condition to observe more remarkable differences in Pi-related genes expression. Root material was collected at 10, 15, 20 das +1 h after Pi re-supply, 21, and 23 das. RNA was isolated from root tissue, and expression was analyzed for a set of Pi-related genes. The greatest differences in gene expression was observed between plants grown under P-sufficient and P-limiting conditions for IPS1, PHT1;1, and pri-miR399c. At control conditions, the highest decrease ranged from 1349-, 447-, and 347-fold reduction, respectively, as compared to...
Pi-starvation condition (Supplementary Figures S2A,K,L). After Pi re-supply, we observed a weaker reduction of IPS1, PHT1;1, and pri-miR399c (which was 26-, 35-, and 44-fold, respectively). These reduction levels were similar to those observed for PHO1 (33-fold, 24 h) and PHT1;1 (20-fold, 12 h) during the heat stress (Supplementary Figure S2I). Interestingly, IPS1 expression was downregulated after Pi re-supply but upregulated 11-fold (48 h) after heat stress (Supplementary Figures S1G and S2L). Expression of the PHO2 and SPX-MFS decreased in Pi-limited plants at the early steps of development; but later, their expression reached the same levels as in the P-fed controls (Supplementary Figures S2G,I). The expression of PHO1 is not changed significantly in different Pi-regime conditions in barley roots. During heat stress both PHT1;1 and PHO1
genes expression is downregulated, but Pi concentration affects only the PHT1;1 expression. No reduction of PHO1 expression may prevent Pi-over accumulation after Pi re-supply treatment. The higher expression of pri-miR399c at the early stages of development correlated with the lower expression of the PHO2 in Pi limited (compared to control plants). At later growth stages, the expression of PHO2 approached the level in the P-sufficient controls plants, although the abundance of pri-miR399c and mature miR399 was still on a high level in Pi starvation condition (Supplementary Figure S2K and Figure 10A). This suggests that the role of miR399 in PHO2 gene-expression regulation is not crucial at the latter developmental stages. Accordingly, Pi re-supply did not change the level of PHO2 expression significantly, although the expression of pri-miR399 and, consequently, mature miR399 was strongly decreased (Supplementary Figure S2K and Figure 10A). Like the mature miR399, miR827 also showed a reduced level after Pi re-supply, corresponding to a reduction to 57% at 3 days after Pi re-supply compared to the P-limiting plants (Figure 10B). Pi re-supply resulted in strongly increased Pi concentrations in barley roots (Supplementary Figure S2M), but the reached level was similar to the Pi concentration level in the control plants. Harvested shoot weights were suppressed by Pi limitation already by 15 days, representing only 40% of the weight of the P-sufficient control plants at 23 days (Table 3). The re-supply of P-to-P limited plants at 20 days produced a marked growth response over the subsequent three days. Symptoms of early senescence were observed for the older leaves of P-limited plants (Figure 11).

**DISCUSSION**

Studies of the expression of phosphate-related genes in crops are crucial to understand the complexity of adaptive reactions in plants grown under phosphate depletion (Baker et al., 2015). This is especially important, due to the fact that phosphate rock for fertilizer production is a limited resource that requires careful management, since it is estimated that all cheap forms of phosphorus will soon be exhausted (Cordell and White, 2014). It is known that heat stress has a big impact on plant growth. The warming climate is one of the biggest challenges for agriculture, as one degree increase in temperature may decrease crop productivity by as much as 10% (Zinn et al., 2010). Heat stress inhibits plant functions in many ways. In barley, it reduces tillering and decreases plant height and spike length (Abou-Elwafa and Amein, 2016). This work shows that a heat-stress period influenced barley morphology such that, after one week, specific leaves were shorter than the corresponding leaves of control plants, and the oldest leaf showed more severe senescence (Figure 8). Heat stress affects the expression of many genes, including those involved in calcium and sugar signaling in wheat, TFs (Hsf, NAC, AP2/ERF, WRKY, MYB, and C2H2) in rice panicles, or six different peroxidases in switchgrass (Qin et al., 2008; Zhang et al., 2012; Li et al., 2013; Banerjee and Roychoudhury, 2015). The predicted increasing exposure of crop plants to heat stress requires an understanding of how they regulate macronutrient homeostasis under such circumstances. We have shown that heat stress affects the expression of Pi-related genes. This is
especially visible in root tissues. Already after one hour of heat stress, we observed decreased levels of PHT1;1, PHT1;4, and HvPHT1;6, which should result in a decreased level of Pi concentration in roots. However, Pi concentration did not change. This could be explained by the reduced expression levels of PHO1, PHO2, and SPX-MSF genes. The pho1 mutant in Arabidopsis is defective in Pi transport from root to shoot, resulting in a strong Pi deficiency in shoot tissue (Stefanovic et al., 2011). On the other side, PHO1-gene over-expression in Arabidopsis leaves led to the increased export of Pi into the xylem vessels (Stefanovic et al., 2011). Barley PHO1 gene expression is weakly responsive to Pi starvation (similar to Arabidopsis). The heat-stress impact on PHO1 gene expression shown here reveals that this gene represents an important Pi homeostasis regulatory element (Hamburger et al., 2002). This data may explain why Pi concentration in barley roots is non-sensitive to heat stress. We assume that Pi transport to the shoots is inhibited during heat stress. The observed decrease of the PHO2 transcript level in roots may be responsible for the lower degradation of Pi transporters. Interestingly, we observed that PHT1;4 gene expression increased during heat stress, which could compensate for the decrease of PHT1;1 expression.

After one hour of heat stress, all analyzed Pi-related genes in the shoots have stable expression compared to the roots, with only one exception: PHO2 expression was downregulated significantly after one hour of heat treatment. Previously, we observed that the silencing of the PHO2 gene resulted in an increased Pi concentration (Pacak et al., 2010). Interestingly, PHR1 and PHR2 TFs expression was not changed in either barley roots or shoots. Since PHR1 expression is stable, other TFs can be responsible for either the downregulation or upregulation of Pi-related genes during heat stress. In roots PHR2 expression is downregulated in the further time-points.

In further experiments, we analyzed plants grown in soil with a different Pi supply. HvIPS1, HvPHT1;1, HvPHT1;4, HvPHT1;6, and MIR399c showed a clear response to Pi starvation. After Pi replenishment, we observed a reduction of HvPHT1;1, HvPHT1;4, HvPHT1;6, HvIPS1, and pri-miR399c expression levels. In contrast to the heat stress Pi re-supply treatment did not downregulate PHO1 gene expression (with one exception 12 h time-point, barley cv. Rolap). We conclude that barley plants utilized two strategies to regulate Pi concentration. Upon heat stress both PHT1;1 and PHO1 genes are downregulated and Pi concentration is stable. Pi re-supply treatment downregulated HvPHT1;1 expression but not PHO1. Its expression is even higher at +1 h time-point (Figure 9E). This allows for Pi concentration increments but prevent Pi over-accumulation. In Black Hulless 10- and 15-day-old plants, we observed higher HvPHO2 expression in plants grown in a high Pi supply (compared to low Pi conditions). The data showed that HvPHO2 expression was influenced by the Pi content in soil and correlated with an increase in pri-microRNA399c expression. PHO2 expression was especially affected by the concentration of Pi in the first two weeks of plant growth. MiR399 and miR827 are the only microRNAs that bind to the 5’ UTR of mRNAs in Arabidopsis (Ding et al., 2012). Their levels increased highly during Pi starvation and also decreased after Pi supply. The early senescence of leaf no. 1 of barley plants grown in low Pi

![Figure 11](image-url)
FIGURE 12 | Model of heat stress impact on Pi-related genes expression and their Pi-homeostasis-related activity. The solid and dotted lines represent regulation on the transcriptional and protein levels, respectively. Red dots represent Pi ions.
suggests the remobilization of Pi from this leaf to the other plant parts.

It was previously shown that, at an ambient temperature (23°C), miR399a, b, and d are more abundant by 1.30–1.31-fold than at 16°C (Lee et al., 2010). During heat stress at a higher temperature (35.5°C), we observed the expression of PHO2 to be downregulated in both roots and shoots. Our Northern analysis of the shoot samples did not show any miR399 signals. The low abundance of miR399 and miR827 in Pi-sufficient conditions and after heat treatment was not surprising. The abundance of these microRNAs is elevated during Pi starvation, and other stresses (like drought) in *Brachypodium distachyon* are not sufficient to induce their expression (Bertolini et al., 2013). We concluded that, apart from the post-transcriptional action of miR399 on *PHO2* expression, downregulation may be caused on the transcriptional level. We performed *PHO2*-promoter analysis and found binding sites for several TFs. Some of them are presented in Table 4.

Our data suggests that heat stress affects the expression of barley Pi-related genes with an impact on plant Pi concentration during the first period of stress. Surprisingly, remarkable changes in the gene expression after 24 and 48 h of heat stress did not result in changed Pi concentrations in the barley plants. The model of gene action during heat stress is presented in Figure 12. Generally, we observed two strategies applied by barley plants to cope with heat stress and Pi-supply treatment. During heat stress, the expression of *PHT1* transporters (especially *PHT1;1*) significantly decreased, with a simultaneous reduction of *PHO1* expression, thus blocking Pi efflux from roots to shoots. This stabilizes the Pi concentration in barley roots. During the Pi re-supply treatment, *PHT1;1* expression is also reduced (but not *PHO1*). This reduction is in response to Pi accessibility, but it did not hamper the increase of Pi concentration in the root. One could expect that the increased Pi concentration in roots should be accompanied by the incremental expression of *PHO1*. Later, after three days of Pi re-supplementation, the expression of the *PHO1* gene remains stable.

Further studies on barley Pi-homeostasis maintenance during heat stress are necessary, mainly at the protein and RNA levels. For example, it was recently shown that cis- *NATPHO1;2* long non-coding RNA promotes *PHO1;2* translation (Secco et al., 2010; Jabnoune et al., 2013). Previously it was reported that *NLA* gene involved in adaptive responses to low nitrogen conditions in *Arabidopsis* is regulated by Pi-starvation induced miR827 (Kant et al., 2011). There are more connections between nitrogen and phosphate metabolism pathways. *Arabidopsis* mutants *phf1* and *pht1;1* were identified as suppressors of the *nla* mutant. These suppressors restore the *nla* mutant phenotype to WT (Kant et al., 2011). Thus the observed decreased level of the *PHT1;1* expression during heat stress could be connected to the *NLA* gene expression up-regulation.

Studies presented in this work shed a new light on the mechanisms of Pi-homeostasis regulation and can help to improve crop plants in their response to unfavorable changes in the environment.

**AUTHOR CONTRIBUTIONS**

AP designed, performed, analyzed all experiments, prepared manuscript, tables, and figures, wrote manuscript, MB-P performed Pi concentration analysis and analyzed experiments, AS-B designed, performed heat stress experiment, KK designed, performed heat stress experiment, PS performed Pi concentration analysis, KM analyzed transcriptome data, IJ designed Pi re-supply experiment and assisted in manuscript preparation, AJ assisted in manuscript writing, editing, ZS-K designed heat experiment, wrote manuscript.

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**SUPPLEMENTARY MATERIAL**

The Supplementary Material for this article can be found online at: [http://journal.frontiersin.org/article/10.3389/fpls.2016.00926](http://journal.frontiersin.org/article/10.3389/fpls.2016.00926)
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