Potential Root Foraging Strategy of Wheat (*Triticum aestivum* L.) for Potassium Heterogeneity

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Potassium (K) distribution is horizontally heterogeneous under the conservation agriculture approach of no-till with strip fertilization. The root foraging strategy of wheat for K heterogeneity is poorly understood. In this study, WinRHIZO, microarray, Non-invasive Micro-test Technology (NMT) and a split-root system were performed to investigate root morphology, gene expression profiling and fluxes of K\(^+\) and O\(_2\) under K heterogeneity and homogeneity conditions. The split-root system was performed as follows: C. LK (both compartments had low K), C. NK (both compartments had normal K), Sp. LK (one compartment had low K) and Sp. NK (the other compartment had normal K). The ratio of total root length and root tips in Sp. NK was significantly higher than that in C. NK, while no significant differences were found between Sp. LK and C. LK. Differential expression genes in C. LK vs. C. NK had opposite responses in Sp. LK vs. C. LK and similar responses in Sp. NK vs. C. NK. Low-K responsive genes, such as peroxidases, mitochondrion, transcription factor activity, calcium ion binding, glutathione transferase and cellular respiration genes were found to be up-regulated in Sp. NK. However, methyltransferase activity, protein amino acid phosphorylation, potassium ion transport, and protein kinase activity genes were found to be down-regulated in Sp. LK. The up-regulated gene with function in respiration tended to increase K\(^+\) uptake through improving O\(_2\) influx on the root surface in Sp. NK, while the down-regulated genes with functions of K\(^+\) and O\(_2\) transport tended to reduce K\(^+\) uptake on the root surface in Sp. LK. To summarize, wheat roots tended to perform active-foraging strategies in Sp. NK and dormant-foraging strategies in Sp. LK through the following patterns: (1) root development in Sp. NK but not in Sp. LK; (2) low-K responsive genes, such as peroxidases, mitochondrion, transcription factor activity, calcium ion binding and respiration, were up-regulated in Sp. NK but not in Sp. LK; and (3) root K\(^+\) and O\(_2\) influxes increased in Sp. NK but not in Sp. LK. Our findings may better explain the optimal root foraging strategy for wheat grown with heterogeneous K distribution in the root zone.

**Keywords:** K heterogeneity, root foraging strategy, root morphology, root gene expression, K\(^+\) flux, O\(_2\) flux

**Abbreviations:** C. LK, a homogeneous K-deprived environment with both compartments had low K; C. NK, a homogeneous K-replete environment with both compartments had normal K; NMT, non-invasive Micro-test Technology; Sp. LK, a heterogeneous split environment with the compartment had low K; Sp. NK, a heterogeneous split environment with the compartment had normal K.
INTRODUCTION

Potassium (K), which accounts for 2–10% of the plant's total dry weight, is one of the most important macronutrients for crop growth (Leigh and Wyn Jones, 1984; Ma et al., 2012; Ruan et al., 2015). K is heterogeneously distributed in soils under minimum tillage (Jackson and Caldwell, 1993; Richard et al., 2000). As a major conservation agriculture approach, no-till has been adopted on 180 million hectares (Mha), or almost 12.5% of the global arable land area (Kassam et al., 2018). In China, no-till, which is usually combined with strip fertilization, is being increasingly advocated for wheat. Approximately 24% of the wheat crop was planted under the agriculture approach of no-till combined with strip fertilization in China by 2009 (Zhao et al., 2017). No-till combined with strip fertilization increases the horizontal spatial variation of K in soils (e.g., a higher K concentration on the fertilized sides and a lower K concentration on the unfertilized sides). This spatial variation occurs due to the wide-row spacing, the fixed crop rows and the fertilizer bands, the high crop residue concentration on the crop rows, the differences in soil properties in relation to their position in the crop-row, and limited K mobility in dry land soils (Ma et al., 2007; Farmaha et al., 2011, 2012; Fernández et al., 2011, 2015; Williams et al., 2017). In rainfed systems, such as those of wheat growing areas in Northern China, the horizontal spatial variation of K is more noteworthy since soil drying conditions generally occur at the end of the growing season when K and water are in greatest demand by wheat (Fernández et al., 2011; Zhao et al., 2017).

Under heterogeneous K distribution, root systems have foraging behaviors for K (Hodge, 2004; Li et al., 2017). In the dry season, K presents relatively low mobility when drying reduces soil water and increases the diffusion path length of K ions to the root surface (Römheld and Kirkby, 2010; Fernández et al., 2011). Root morphology characteristics (especially the root tips, root surface area and root length) are the criteria for evaluating root abilities of penetrating soil and contacting K when K presents with relatively low mobility (Hassan and Arshad, 2010). Root systems can enhance their length and surface area in the soil K-rich patches and restrict them in K-deficient patches (Fernández et al., 2011; Kellermeier et al., 2013). The underlying molecular mechanisms, which are involved in the root morphological changes in response to K heterogeneity, are mainly dependent on the regulation of genes encoding hormones. These hormone genes mainly include auxin, ethylene and jasmonic acid-related genes (Jung et al., 2009; Liu et al., 2013; Giehl et al., 2014; Gupta et al., 2017; Lee et al., 2017). Root development-related genes are also involved in the root morphological changes in response to K heterogeneity (Giehl and von Wirén, 2014). In addition, other physiological changes, such as the release of nutrient mobilizing root exudates or the expression of nutrient transporters, also provide contributions to root adaptions to K heterogeneity (Gruber et al., 2013; Giehl and von Wirén, 2014; Zhao et al., 2016).

Compared with root foraging strategies for K heterogeneity, root foraging strategies for nitrogen (N) and phosphorus (P) heterogeneity are more abundant. In addition to the regulation of root morphology, root exudates, nutrient transporters, hormone genes and root development-related genes (Shen et al., 2005; Paterson et al., 2006; Guan et al., 2014; Sun et al., 2018), other physiological and molecular changes have also been found to be involved in the root foraging strategy for N and P heterogeneity. Cytoskeleton activation, cell wall modification, arbuscular mycorrhizal fungi and so on are involved in root foraging for N and P heterogeneity (Duan et al., 2015; Trevisan et al., 2015). Various genes, such as genes involved in cell expansion and division, transcription factors, kinases, sugar transportation and utilization, NO\textsuperscript{3−} absorption and assimilation, cytokinin biosynthesis and so on, have been proven to be involved in root foraging responses to N and P heterogeneity (Liu et al., 2008; Krouk et al., 2010; Ruffel et al., 2011; Alvarez et al., 2012; Robaglia et al., 2012; Tabata et al., 2014; Shabnam and Iqbal, 2016; Zhou et al., 2017; Haling et al., 2018). However, unlike N and P, K does not convert into other organic compounds in plants. Thus, K\textsuperscript{+} can transport between tissues in plants. K heterogeneity results in the roots of the same plant being exposed to different K conditions. K\textsuperscript{+} in the root exposed to a low-K condition may be supplemented by K\textsuperscript{+} absorbed by a root exposed to a high-K condition. Therefore, root foraging strategies in the presence of K heterogeneity may be quite different from that for N or P. In most cases, the transport of K\textsuperscript{+} between tissues involves active transport, which requires a lot of energy (Szczera et al., 2009). Aerobic respiration, which provides energy, may play a key role in dealing with K heterogeneity. However, aerobic respiration is rarely considered when studying root forging strategies for K heterogeneity.

Therefore, we proposed that plants could realize the maximum utilization of energy and the optimum state of the root systems through root foraging behaviors under K heterogeneity (Mcniclue et al., 2009). In order to investigate the optimal root foraging strategy of wheat for K heterogeneity, we selected a K efficient wheat genotype “Tongzhou916” as the research object, which has a high K efficiency coefficient, great root development, strong K uptake ability and up-regulation of low-K responsive genes (Ruan et al., 2015). Because using localized fertilization in row crops is a similar phenomenon to the split-root system, we used the split-root system to mimic a heterogeneous soil environment. In this study, root parameters, gene expression profiling and root fluxes of net K\textsuperscript{+} and O\textsubscript{2} were analyzed by WinRHIZO, Affymetrix GeneChip and NMT, respectively. The results in this study will give us a basis to better understand the optimal root foraging strategy for K heterogeneity in the root zone of wheat.

MATERIALS AND METHODS

Plant Materials and Split-Root System

According to a previous study, Tongzhou916 has the highest K efficiency coefficient among 50 representative wheat varieties. In addition, its great root development, strong K uptake ability and the up-regulation of low-K responsive genes make a great contribution to the low-K tolerance of Tongzhou916 (Ruan et al., 2015). Therefore, “Tongzhou916” was selected as a low-K\textsuperscript{+} tolerant wheat genotype and used as the test object in this research. The hydroponics test of the wheat was carried out in
an artificial climate box. The specific environmental parameters are shown below: (1) the day and night time were 16 and 8 h, respectively; (2) the day and night temperature were 25 and 18°C, respectively; (3) the illumination intensity was 30000 lux in the day; (4) the atmospheric humidity was 70%.

In order to simulate heterogeneous soil K availability, we used a split-root system. Two separated physical spaces, in which different K supplies could be provided, were created in this split-root system. During the split-root experiment, we selected wheat seedlings with the same growth of roots and shoots. Then, the wheat roots were planted on two separated physical spaces uniformly. Each separated physical space contained the same root tip number and root length at the beginning of the experiment. Three different K environments were provided as follows (Ruffel et al., 2011): (1) a homogeneous K-deprived condition (C. LK: both compartments had low K). (2) a homogeneous K-replete condition (C. NK: both compartments had a normal K), and (3) a heterogeneous split condition (Sp. LK/Sp. NK: one compartment had low K, and the other had normal K). The nutrition solution was (mmol L\(^{-1}\)): NH\(_4\)NO\(_3\) 1.0, Ca(NO\(_3\))\(_2\)-2H\(_2\)O 1.0, NaH\(_2\)PO\(_4\) 0.25, MgSO\(_4\)-H\(_2\)O 1.0, CaCl\(_2\) 1.5, Fe-EDTA 0.1, ZnSO\(_4\)-7H\(_2\)O 1.0 × 10\(^{-3}\), (NH\(_4\))\(_6\)MoO\(_7\)2H\(_2\)O 5.0 × 10\(^{-5}\), MnSO\(_4\)-H\(_2\)O 1.0 × 10\(^{-3}\), CuSO\(_4\)-5H\(_2\)O 5.0 × 10\(^{-4}\), and H\(_3\)BO\(_4\) 1.0 × 10\(^{-3}\). The normal K treatment contained 1.0 mmol L\(^{-1}\) K\(_2\)SO\(_4\), while the low K treatment contained 0.005 mmol L\(^{-1}\) K\(_2\)SO\(_4\). All of the treatments had a pH of 6.5. The nutrient solution was changed once a day. The nutrient solution was supplied 12 h O\(_2\)/d by pumps.

In the first experiment, to analyze the shoot dry weight, K biological utilization and root development, seedlings were transferred to the split-root system for 4 weeks. Then, the seedlings were harvested, and we investigated the phenotypes, shoot dry weights, K contents and root parameters. In the second experiment, for the investigation of the molecular basis of the wheat root responses to heterogeneous vs. homogeneous K environments, 3-week-old seedlings, which were first planted in normal K solutions, were transplanted to the above split-root system for 5 days. Then, the wheat roots on each side were collected. The harvested roots were put into liquid N quickly transferred the balanced roots to a new Petri dish that contained a fresh testing solution. K\(^+\) precipitation and O\(_2\) content was measured after digesting the material with a mixture of H\(_2\)O\(_2\) and H\(_2\)SO\(_4\) (Mills and Jones, 1996) followed by flame photometer to measure the K\(^+\) content of the digested solution. For the plant biomass and K\(^+\) concentration measurement experiments, three biological replications were performed (each replicate included 20 individual plants). The formula of the K biological utilization is shown below:

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K\text{ biological utilization} = \text{dry weight of shoot/K concentration of shoot (1)}.
\]

**Microarray Analysis**

With the help of the Shanghai Biotechnology Corporation, the microarray analysis was performed successfully. Three biological replicates (each replicate included 20 individual plants) were carried out in this experiment. According to the instructions of the manufacturer, we extracted the total RNA by using TRIzol reagent (Cat#15596-018, Life Technologies, Carlsbad, CA, United States). With the help of Agilent Bioanalyzer 2100 (Agilent Technologies, Santa Clara, CA, United States), we checked the integrity of the RNA by determining its RIN number. We purified the total RNA by using an RNeasy micro kit (Cat#74004, QIAGEN, GmBH, Germany) and RNase-Free DNase Set (Cat#79254, QIAGEN, GmBH, Germany) (Ruan et al., 2015).

According to the instructions of the manufacturer, we amplified, labeled and purified the total RNA through the use of the GeneChip 3’IVT Express Kit (Cat#901229, Affymetrix, Santa Clara, CA, United States) in order to obtain biotin labeled cRNA (Wu et al., 2013). According to the instructions of the manufacturer, we performed array hybridization and washing by carrying out GeneChip® Hybridization, Wash and Stain Kit (Cat#900720, Affymetrix, Santa Clara, CA, United States) in a Hybridization Oven 645 (Cat#00-0331-220V, Affymetrix, Santa Clara, CA, United States) and Fluidics Station 450 (Cat#00-0079, Affymetrix, Santa Clara, CA, United States). We scanned the slides by using a GeneChip® Scanner 3000 (Cat#00-00212, Affymetrix, Santa Clara, CA, United States) and Command Console Software 3.1 (Affymetrix, Santa Clara, CA, United States) with default settings (Zeng et al., 2014; Zhu et al., 2016).

**Determination of the Flux Rates of K\(^+\)**

**and O\(_2\)** **on the Wheat Root Surfaces**

In order to determine the K\(^+\) and O\(_2\) net flux rates on the wheat root surfaces, wheat roots were cut off from the whole root system of each plant. Before detection, we transferred the wheat roots to a Petri dish that contained 10 mL of testing solution. After 10 min of incubation to allow them to adapt and reduce any instability induced by environmental changes, the wheat roots were used for measurements. During the detection stage, we transferred the balanced roots to a new Petri dish that contained a fresh testing solution. K\(^+\) flux and O\(_2\) influx were determined by utilizing NMT (Non-invasive Micro-test System, BIO-001A;
Younger USA LLC., Amherst, MA, United States). Six repeats were set for each test treatment. The net K⁺ flux rates along the root axis (from 0 to 250 mm) were measured to determine the maximal net K⁺ influx areas along the root axis (Supplementary Figure S1). In the areas of 7 cm to 10 cm from the root apex, we found that the K⁺ net flux rate reached the peak value. Therefore, the areas between 7 and 10 cm from the root apex were selected as the measurement site. The determining time for each tested root was 10 min. The measurement details of ion flux, such as the NMT system and the related applications, have been reported in previous research (Zheng et al., 2012; Ruan et al., 2016; Zhang et al., 2016).

Quantitative Real-Time PCR (qRT-PCR) Analysis

To verify the reliability of our microarray, Quantitative Real-time PCR (qRT-PCR) was performed. The testing roots for qRT-PCR were the part of the roots used for the microarray experiment. The total RNA extractions of root samples were performed by using the method for the TRIZOL reagent (Cat#15596-018, Life technologies, Carlsbad, CA, United States). The endogenous control used in this experiment was the actin gene of *Triticum aestivum* L. According to the instructions of the manufacturer, we synthesized first-strand cDNA by using the extracted total RNA with a cDNA synthesis Kit (Promega, United States). Three biological replicates (each replicate included 20 individual plants) were carried out in this experiment. A 20 µL reaction system, which contained 10, 6, 2, 1, and 1 µL of 2X-RT mix, nuclease-free water, template (0.2 µM), forward primer and reverse primer, was used to carry out the experiment of qRT-PCR. In the experiment of the qRT-PCR, the software of Primer 5 and DNAMAN were used to design the PCR primers (Supplementary Table S1). At least three technical duplications were performed for each PCR experiment. The relative expression of the chosen genes was calculated through the relative quantitative method of ΔΔCT (Revel et al., 2002; Ruan et al., 2015).

Statistical Analysis

The data of the microarray were normalized through the utilization of the software of MAS 5.0 algorithm, Gene Spring Software 11.0 (Agilent technologies, Santa Clara, CA, United States). We only selected the genes that showed significant differences ($P < 0.05$ and fold-change $> 1.5$) as further research objects. The data analysis was carried out with the help of
RESULTS

Wheat Growth State Under the Split-Root System

In order to simulate heterogeneous soil K distribution, we used a split-root system in the current research. Two separated physical spaces, in which different K supplies could be provided, were created in this split-root system. To study the potassium (K) foraging behavior of wheat roots, we designed three treatments (Figure 1A). The phenotypic differences were significant between all of the treatments (Figure 1B). Significant differences in shoot growth were observed between the treatments. The growth state of wheat roots under the split-root system is shown in Figure 2A. The functional annotation was performed with the help of AgriGO online service. For figure drawing, OriginPro 8.1 (Origin Inc., Chicago, IL, United States) was used. The significance of the group differences was calculated by using ANOVA and t-tests. Microsoft Excel (Microsoft Corporation, United States) and SPSS 18 (SPSS Inc., Chicago, IL, United States) were performed for the data statistical analysis.

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1 http://sas.shbio.com/portal/root/molnet_shbh/index.jsp
2 http://bioinfo.cau.edu.cn/agriGO
FIGURE 3 | Gene Ontology (GO) classifications and amount of DEGs in wheat root response to heterogeneous vs. homogeneous K environments. (A) Functional category distributions of main up-regulated genes in Sp. NK vs. C. NK. (B) Functional category distributions of main down-regulated genes in Sp. LK vs. C. LK. All microarray experiments were performed in three biological replicates.

FIGURE 4 | Unique differentially expressed gene and net flux rates of K$^+$ and O$_2$ in heterogeneous vs. homogeneous normal-K environments. (A) Expression pattern and GO classifications of unique differentially expressed genes in heterogeneous (Sp. NK) vs. homogeneous (C. NK) normal-K environments. The means ± SD (n = 3) are shown. * indicates differences between means at P < 0.05. All microarray experiments were performed in three biological replicates. (B) The rates of net K$^+$ fluxes in Sp. NK and C. NK. (C) The rates of net O$_2$ fluxes in Sp. NK and C. NK. Six biological replicates were established in the net flux experiments.

dry weight and K biological utilization were found among all of the treatments (Figures 1C,D). Shoot dry weight and K biological utilization were ranked as follows: C. NK > Sp. LK/Sp. NK > C. LK. Shoot dry weight and K biological utilization in the treatment of Sp. LK/Sp. NK were 64.6 and 70.8% of values for C. NK, respectively. In addition, there were significant differences in root development among any of the treatments (Figures 1E–H). The root system grew uniformly in homogeneous K environments (C. LK and C. NK) and non-uniformly in a heterogeneous split environment (Sp. LK/Sp. NK). The total root length, total root surface area and root tips in C. LK were 46.6, 38.0, and 51.8% of those in C. NK (Figures 1E–G). The total root length, total root surface area and root tips in Sp. NK were 85.9, 71.1, and 68.8% of those in C. NK (Figures 1E–G). The ratio of total root length and root tips in Sp. NK was significantly higher than that in C. NK (Figure 1H). No significant differences were found in root development between Sp. LK and C. LK (Figures 1E–H). The total root length, total root surface area and root tips in Sp. NK were more than half of those in C. NK (Figures 1E–G). The ratio of total root length and root tips in Sp. NK was significantly
higher than that in C. NK (Figure 1H). In addition, we compared two types of K heterogeneity as follows: (a) Sp0. MK/Sp0. NK: one compartment had 0.5 mmol L\(^{-1}\) K\(_2\)SO\(_4\), and the other had 1.0 mmol L\(^{-1}\) K\(_2\)SO\(_4\); (b) Sp. LK/Sp. NK: one compartment had 0.005 mmol L\(^{-1}\) K\(_2\)SO\(_4\), and the other had 1.0 mmol L\(^{-1}\) K\(_2\)SO\(_4\). The results showed that the total root length, total root surface area and root tips in Sp. NK were significantly higher than those in Sp0. NK (Supplementary Table S2). The ratio of total root length and root tips in Sp. NK was also significantly higher than that in Sp0. NK (Supplementary Table S2).

### Differentially Expressed Genes (DEGs) in Wheat Root Responses to Heterogeneous vs. Homogeneous K Environments

In this study, more than 61200 probe signals were detected from the Affymetrix GeneChip. The raw data sets (CEL) and the normalized expression data sets have been deposited in the Gene Expression Omnibus (GSE115111) at the National Center for Biotechnology Information\(^3\). For further analysis, the genes with significant differences (\(P < 0.05\) and fold-change > 1.5) were chosen to include as many candidate genes as possible.

There were 59 up-regulated genes in C. LK vs. C. NK. Among these genes, there were 8, 7, 0, and 0 shared genes in Sp. LK vs. C. LK down-regulated genes, Sp. NK vs. C. NK up-regulated genes, Sp. LK vs. C. LK up-regulated genes, and Sp. NK vs. C. NK down-regulated genes, respectively (Figure 2A). There were 33 down-regulated genes in C. LK vs. C. NK. Among these genes, there were 11, 2, 0, and 0 shared genes in Sp. LK vs. C. LK up-regulated genes, Sp. NK vs. C. NK down-regulated genes, Sp. LK vs. C. LK down-regulated genes, and Sp. NK vs. C. NK up-regulated genes, respectively (Figure 2A). There were 10 up-regulated genes shared in Sp. LK vs. C. LK and Sp. NK vs. C. NK (Figure 2A). The heat map of the above shared genes is shown in Figure 2B. Differential expression of genes in C. LK vs. C. NK had opposite responses in Sp. LK vs. C. LK and similar responses in Sp. NK vs. C. NK (Figure 2B).

### Functional Annotations of Differentially Expressed Genes in Wheat Root Responses to Heterogeneous vs. Homogeneous K Environments

To study the functions of putative key genes of wheat in response to heterogeneous vs. homogeneous K environments (Supplementary Table S3), Gene Ontology (GO) analysis was performed to analyze the functional differences between differentially expressed genes in Sp. NK vs. C. NK and Sp. LK vs. C. LK (Figure 3). There were five shared functions of up-regulated genes in Sp. NK vs. C. NK and down-regulated genes in Sp. LK vs. C. LK, including iron ion binding, mitochondria, transcription factor activity, cytoplasmic membrane-bounded vesicle and plastids (Figure 3). In addition, functional categories of up-regulated genes unique in Sp. NK vs. C. NK included calcium ion binding, glutathione transferase, cellular respiration, jasmonic acid mediated signaling pathway

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\(^3\)http://www.ncbi.nlm.nih.gov/geo/

![Figure 5](image-url)
FIGURE 6 | Validation of microarray data by the method of quantitative real-time PCR (qRT-PCR). All microarray and qRT-PCR experiments were carried out in three biological replications. Data are the log2 ratio (heterogeneous K/homogeneous K) for the selected genes. The correlation coefficient ($r^2$) is shown in the diagram.

and so on (Figure 3A). Functional categories of down-regulated genes unique in Sp. LK vs. C. LK included methyltransferase activity, protein amino acid phosphorylation, potassium ion transport, protein serine/threonine kinase activity and so on (Figure 3B).

Unique Differentially Expressed Genes and Net Flux Rates of K$^+$ and O$_2$ in Heterogeneous vs. Homogeneous K Environments

In order to investigate the key genes involved in K uptake, unique up-regulated genes in Sp. NK vs. C. NK was shown in Figure 4A. The unique genes had the following functions: respiratory gaseous exchange, mitochondrial respiratory chain and cellular respiration (Figure 4A). The up-regulated gene in Sp. NK vs. C. NK was up-regulated in C. LK vs. C. NK at the same time. In addition, the fold change in Sp. NK vs. C. NK was higher than that in C. LK vs. C. NK (Figure 4A). To compare the flux rate of K$^+$ and O$_2$ on the root surface in Sp. NK and C. NK, NMT was performed in this research. The net flux rate of K$^+$ and O$_2$ in Sp. NK and C. NK were shown in Figures 4B,C. For the flux rate of K$^+$, the curves presented relatively steady absorption states for both Sp. NK and C. NK (Figure 4B). However, the overall influx rate of K$^+$ in C. LK was higher than that in Sp. LK. For the flux rate of O$_2$, the curves presented gradual upward trends for both Sp. NK and C. NK (Figure 4C). For both Sp. NK and C. NK, the root surfaces presented the absorption state. The influx rate of O$_2$ in C. NK was higher than that in Sp. NK.

To investigate the key genes involved in K uptake, unique down-regulated genes in Sp. LK vs. C. LK were shown in Figure 5A. The unique genes had the following functions: potassium ion transport, potassium ion transmembrane transporter activity, oxygen transporter activity and oxygen transport (Figure 5A). In contrast, the down-regulated genes in Sp. LK vs. C. LK were up-regulated in C. LK vs. C. NK (Figure 5A). To compare the flux rate of K$^+$ and O$_2$ on the root surface in Sp. LK and C. LK, NMT was applied in this study. The net flux rate of K$^+$ and O$_2$ in Sp. LK and C. LK were shown in Figures 5B,C. For the flux rate of K$^+$, the curves presented relatively steady absorption states for both Sp. LK and C. LK (Figure 5B). However, the overall influx rate of K$^+$ in C. LK was higher than that in Sp. LK. For the flux rate of O$_2$, the curves presented gradual upward trends for both Sp. LK and C. LK (Figure 5C). For both Sp. LK and C. LK, the root surfaces presented the absorption state. The influx rate of O$_2$ in C. LK was higher than that in Sp. LK.

Quantitative Real-Time PCR (qRT-PCR) Analysis

To verify the reliability of our microarray results, quantitative real-time PCR (qRT-PCR) was carried out. In the test of qRT-PCR, several genes were selected randomly from the above four treatments. The specific-primers of these selected genes were designed as shown in Supplementary Table S1. The data of qRT-PCR showed that gene expression trends were significantly similar ($r^2 = 0.75$) to the results of the microarray data (Figure 6). This indicated that the results of the microarray results were credible in this study.
DISCUSSION

As shown above, coordinated physiological and molecular responses suggested that a variety of processes took part in the wheat root responses to heterogeneous vs. homogeneous K environments. Thus, we will focus on the three questions below: (1) What kind of root foraging strategy was optimal for K heterogeneity in wheat? (2) What were the major physiological processes taking part in the adaption to K heterogeneity of wheat? (3) What key genes were involved in regulating the above physiological processes?

The active-foraging strategy in Sp. NK and dormant-foraging strategy in Sp. LK was the optimal root foraging strategy for K heterogeneity in wheat. Such a root foraging strategy made the treatment of Sp. LK/Sp. NK produce more than half of the K heterogeneity in wheat. Such a root foraging strategy made the system in Sp. LK was the optimal root foraging strategy for K heterogeneity. Since low-K responsive genes made great contributions to the strong ability of K deficiency tolerance in plants (Ma et al., 2012; Ruan et al., 2015; Li et al., 2017), low-K responsive genes might improve the wheat low-K tolerance under K heterogeneity by improving the low-K tolerance of wheat roots on the normal-K supply side (Sp. NK), rather than by improving the low-K tolerance of wheat roots on the low-K supply side (Sp. LK). The low-K responsive genes screened in this study included genes previously reported, such as peroxidase, transcription factors, calcium ion binding, glutathione transferase, cellular respiration, jasmonic acid, methyltransferase, protein amino acid phosphorylation, and protein amino acid phosphorylation related genes. These genes play a key role in various processes of low-K responses, such as the production and elimination of reactive oxygen species (ROS), regulation of the high-affinity K⁺ absorption transporters, the promotion of root development, modifications of K transporters or channels, respiration and so on (Armengaud et al., 2004; Mittler et al., 2004; Shin and Schachtman, 2004; Shin et al., 2005; Hong-Hermesdorfa et al., 2006; Lee et al., 2007; Ho and Tasy, 2010; Kim et al., 2010, 2012; Held et al., 2011; Ma et al., 2012; Hong et al., 2013; Wang and Wu, 2013; Hafsi et al., 2014). Moreover, some new low-K responsive genes, such as genes encoding mitochondrial and oxygen transport genes were also screened in this study.

Mitochondria, as the main site for aerobic respiration, are structures that generate most of the ATP in cells (Stoimenova et al., 2007). Therefore, the up-regulations of genes encoding mitochondria could provide energy for K⁺ transmembrane transportation. In the present study, genes encoding mitochondria were up-regulated in Sp. NK vs. C. NK and down-regulated in Sp. LK vs. C. LK, indicating that mitochondria might tend to provide more driving forces for K transmembrane transportation in the treatment of Sp. NK. In addition, the root K⁺ uptake was markedly reduced under conditions of oxygen deprivation, while the K⁺ uptake was improved when the oxygen supply was increased (Mugnai et al., 2011; Pottosin, 2014). In this study, the down-regulation of genes encoding oxygen transport reduced the O₂ uptake ability in Sp. LK, suggesting that the energy produced by the oxygen transport-mediated aerobic respiration decreased in Sp. LK. Less energy led to a lower K⁺ influx rate in Sp. LK than in C. LK. Above all, the expression patterns of the above genes supported the active-foraging strategy in Sp. NK and a dormant-foraging strategy in Sp. LK.

The major physiological processes taking part in the adaption to K heterogeneity of wheat were the variations of root K⁺ and O₂ flux rates between Sp. NK and Sp. LK. The root K⁺ uptake was markedly reduced under conditions of oxygen deprivation, while the K⁺ uptake was improved when the oxygen supply was increased (Mugnai et al., 2011; Pottosin, 2014). Therefore, O₂ played an important role in K⁺ absorption. In the present study, both K⁺ and O₂ influx rates in Sp. NK were higher than those in C. NK, while both K⁺ and O₂ influx rates in Sp. LK were lower than those in C. LK. Root aerobic respiration could improve K transport ability by supplying more energy for plants under low-K stress (Singh and Blanke, 2000; Hafsi et al., 2014). The higher O₂ influx rates provided more energy for K transport in Sp. NK, while the lower O₂ influx rates led to a weaker K transport.
ability in Sp. LK. Therefore, the variations in root $K^+$ and $O_2$ flux rates between Sp. NK and Sp. LK supported the active-foraging strategy in Sp. NK and dormant-foraging strategy in Sp. LK. These physiological processes allow wheat the maximum utilization of energy, which could improve the adaption ability to the K heterogeneity of wheat.

The key genes involved in regulating the above physiological processes include genes with functions of respiratory gaseous exchange, mitochondrial respiratory chains and cellular respiration (Ta.28112.1.S1_at), potassium ion transport (Ta.113.1.S1_at, TaAffx.52479.1.S1_at), and oxygen transport (Ta.14486.1.S1_at). The up-regulation of the genes, which had the functions of respiratory gaseous exchange, mitochondrial respiratory chain and cellular respiration, led to increased $O_2$ influx in Sp. NK. The increased $O_2$ influx improved root aerobic respiration, which could supply more energy for $K$ transport in Sp. NK (Singb and Blanke, 2000; Hafsi et al., 2014). Indeed, the overall influx rate of $K^+$ in Sp. NK was higher than that in C. NK. On the other hand, the down-regulation of genes that have the functions of potassium ion transport and potassium ion transmembrane transporter activity, led to decreased $K^+$ influx in Sp. LK.

Various researchers have shown that potassium ion transporters, such as OsHAK1, HvHAK1, TaHKT1, and AtHAK5, make great contributions to $K^+$ uptake when the external $K$ supply is low (Bañuelos et al., 2002; García de Blas et al., 2003; Gierth et al., 2005; Fulgenzi et al., 2008; Ruan et al., 2015). However, the potassium ion transport genes are down-regulated in Sp. LK, which is different from the previous studies performed under homogeneous low-$K$ conditions. The main reason for this difference is that the root system in Sp. LK receives the control of system signals rather than local signals under the K heterogeneity condition. The system signals result in the root system performing the dormant-foraging strategy in Sp. LK. Likewise, the oxygen transport gene was down-regulated in Sp. LK, which could result in the $O_2$ influx rate decreasing in Sp. LK. A lower $O_2$ influx could make the energy supply for $K$ transport decrease, which would lead to a lower $K^+$ influx in Sp. LK. Overall, the key genes involved in regulating the above physiological processes could result in the active-foraging strategy in Sp. NK and dormant-foraging strategy in Sp. LK.

CONCLUSION

The root foraging strategy of wheat for K heterogeneity has been reported. Based on the root morphology, root gene expression profiling and root fluxes of net $K^+$ and $O_2$, it was hypothesized that wheat roots tended to perform active-foraging strategies in Sp. NK and dormant-foraging strategies in Sp. LK through the following patterns: (1) roots elongated in Sp. NK rather than in Sp. LK; (2) low- K responsive genes, such as peroxidases, mitochondria, transcription factor activity, calcium ion binding and respiration, were up-regulated in Sp. NK rather than in Sp. LK; and (3) root $K^+$ and $O_2$ influxes increased in Sp. NK rather than in Sp. LK. Our findings may help in better understanding of the optimal root foraging strategy of wheat for K heterogeneity. The physiological traits (e.g., root morphology, root flux rates of $K^+$ and $O_2$) can be used as germplasm screening indicators for high yield wheat under the conservation tillage method of no-till with strip fertilization. In addition, the four key genes, with the functions of root respiration, potassium ion transport and oxygen transport, can be used as a reference for molecular marker assisted breeding for high yield wheat under the conservation agriculture approach of no-till with strip fertilization.

AUTHOR CONTRIBUTIONS

JZ and LR designed the experiments. LR and XX conducted the measurements, data analysis, and wrote the manuscript. BZ, HC, CZ, and DM assisted with the data analysis. LC assisted with the experiments. All authors reviewed the manuscript before the submission.

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SUPPLEMENTARY MATERIAL

The Supplementary Material for this article can be found online at: https://www.frontiersin.org/articles/10.3389/fpls.2018.01755/full#supplementary-material

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CONCLUSION

The root foraging strategy of wheat for K heterogeneity has been reported. Based on the root morphology, root gene expression profiling and root fluxes of net $K^+$ and $O_2$, it was hypothesized that wheat roots tended to perform active-foraging strategies in Sp. NK and dormant-foraging strategies in Sp. LK through
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