Population transcriptomic characterization of the genetic and expression variation of a candidate progenitor of Miscanthus energy crops

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Abstract

The use of transcriptome data in the study of the population genetics of a species can capture faint signals of both genetic variation and expression variation, and can provide a broad picture of a species’ genomic response to environmental conditions. In this study, we characterized the genetic and expression diversity of *Miscanthus lutaria* by comparing more than 16,225 transcripts obtained from 78 individuals, belonging to 10 populations distributed across the species’ entire geographic range. We only observed a low level of nucleotide diversity (π = 0.000434) among the transcriptome data of these populations, which is consistent with highly conserved sequences of functional elements and protein-coding genes captured with this method. Tests of population divergence using the transcriptome data were consistent with previous microsatellite data but proved to be more sensitive, particularly if gene expression variation was considered as well. For example, the analysis of expression data showed that genes involved in photosynthetic processes and responses to temperature or reactive oxygen species stimuli were significantly enriched in certain populations. This differential gene expression was primarily observed among populations and not within populations. Interestingly, nucleotide diversity was significantly negatively correlated with expression diversity within populations, while this correlation was positive among populations. This suggests that genetic and expression variation play separate roles in adaptation and population persistence. Combining analyses of genetic and gene expression variation represents a promising approach for studying the population genetics of wild species, and may uncover both adaptive and non-adaptive processes.
Introduction

Rapid developments in sequencing technologies are beginning to offer powerful tools for studying genomic patterns in non-model organisms at the population level, and promise to provide new insights into the fundamental processes underlying genetic differentiation and adaptation of species. Starting from the late 1960s, non-coding or anonymous gene markers have been used to analyze genetic data for populations of species with the goal of understanding demographic processes such as mutation-selection balance, bottlenecks and migrations etc. (Slatkin 1987; Orr 1998; Hedrick 2009). Beyond these classical applications, the need to study process of adaptation and adaptive evolution has resulted in the development of novel method for population genetics. For example, new genomic and transcriptomic methods provide population geneticists with tools to more accurately characterize patterns of variation in natural populations and study the ecological and evolutionary processes that underlie species adaptations (Hartl & Clark 2006; Barrett & Schluter 2008; Stapley et al. 2010; Ekblom & Galindo 2011; Lamichhaney et al. 2012; Tine et al. 2014; Pan et al. 2017; Bowsher et al. 2017).

RNA-Seq is one of the most commonly used next-generation sequencing (NGS) approaches in non-model organisms (Shendure et al. 2008; Robertson et al. 2010; Grabherr et al. 2011). It accurately quantifies the relative levels of each transcript present in a sample by mapping reads to a reference sequence assembly (Morin et al. 2008; Harrison et al. 2012), yielding high-throughput gene expression data at low effort (Gahlan et al. 2012; Annadurai 2012). In addition, single nucleotide polymorphisms (SNP) identified from RNA-Seq data can potentially capture key sequence variation involved in the adaptation of a species to its...
environment. Because variation in gene expression could similarly be involved in local adaptation, and gene expression levels are at least in part affected by regulatory elements or epigenetic mechanisms, several studies have indicated that gene expression levels may be heritable and be acted upon via natural selection (Ellison et al. 2011; Guggisberg et al. 2013; Nabholz et al. 2014). Variation in genetic expression may thus be altered even before genetic variants arise in the population (West-Eberhard 2003), hence such changes could reflect the early processes that underlie adaptive divergence (e.g., in Oleksiak et al. 2002; Derome et al. 2006; Jeukens et al. 2010).

Transcriptomes represent a cost-effective method to explore the genetic mechanisms that govern ecological interactions and that underlie adaptive divergence at levels of sequences, genes or metabolic pathways in non-model organisms (Alvarez et al. 2015). For example, Derome et al. (2006) sequenced the transcriptomes of lake whitefish (Coregonus clupeaformis) from two separate lakes and found that sixteen genes related to energetic metabolism and regulation of muscle contraction showed true parallelism of transcription, i.e., parallel directional changes in expression in independently evolving populations. Harris et al. (2015) compared the transcriptomes of the white-footed mouse (Peromyscus leucopus) from urban and rural populations, and found the two populations to differ in 104,655 high-quality SNPs and 65 single sequence repeats (SSRs), as well as having 19 differentially expressed contigs related to the modification of proteins and immune function. These studies provide insights into the divergence of genetic variation and the level of gene expression that may drive the responses to different ecological environments. Therefore, variation in gene sequences and gene expression may be complementary mechanisms for a species to respond.
to its environment, and each mechanism may have unique roles in governing species adaptation (Whitehead et al. 2012). However, few studies to date have investigated the patterns and relationships between genetic variation and gene expression within and among populations across the geographic distribution of a species.

Endemic species provide an ideal opportunity to investigate the patterns and relationships of genetic variation and gene expression diversity in response to dynamic environments as they are restricted to narrowly defining geographic regions. Miscanthus lutario riparius is endemic to central China and found on seasonally flooded river banks, farm lands and ruderal land in cities. The species has a distinct distribution range from Miscanthus sacchariflorus and produces thicker, taller tillers (Chen & Renvoize 2005). M. lutario riparius plays a key role in the Yangtze River ecosystem and constitutes the major food source and habitat for many types of birds, mammals, and other animals. Miscanthus species are considered promising second-generation energy crops (Jorgensen & Schwarz 2000; Clifton-Brown et al. 2004; Somerville et al. 2010; Clark et al. 2014; 2015), and M. lutario riparius could be developed as a bioenergy resource because of its considerable biomass, presumably adapted to marginal lands, and abundant genetic variation found in natural populations (Yan et al. 2012; 2016).

Similar to other Miscanthus species, M. lutario riparius can spread clonally from rhizomes, and propagate sexually from seeds (Quinn et al. 2010; Nishiwaki et al. 2011; Yan et al. 2012). Once established, the plant develops strong rhizomes and an intense root system that enhances drought tolerance and carbon sequestration of the ecosystem (Sang & Zhu 2011; Mi et al. 2014). Despite this clonality, morphological surveys and physiological
measurements in different common gardens at different latitudes have indicated that there is considerable variation in *M. lutarioriparius*, and that there is evidence for local adaptation of plant genotypes (Yan *et al.* 2012; 2015). When *M. lutarioriparius* was transplanted from its native habitat to the Loess Plateau (Fig. 1), the low survival rate in the new environment drastically decreased genetic diversity of the transplanted population, while in the surviving plants the expression of stress-tolerance genes showed significantly increased variation (Fan *et al.* 2015; Xu *et al.* 2015; Xing *et al.* 2016; Song *et al.* 2017). However, while these studies characterized the genetic responses of *M. lutarioriparius* under extremely novel conditions, we still lack information on the performance of the species in its natural populations.

In this study, we used population RNA-Seq data from ten geographically isolated populations of *M. lutarioriparius* to examine the patterns of nucleotide and transcriptional variation across the natural range of this species, and to identify the potential mechanisms underlying the plant’s strong adaptive capacity. Additionally, we proposed to test 1) whether transcriptome data may be used as a new tool to explore population genetics via comparisons of diversity patterns obtained from SNP and expression data and 2) whether pairing nucleotide diversity with expression data from the same set of genes is a useful strategy for studying evolutionary and ecological genomics. The results of this study are expected to potentially reveal novel insights on local adaptations and evolutionary processes.
Materials and Methods

Sample collection, cDNA preparation and RNA-Seq

*M. lutarioriparius* is a riparian plant that can be established in semiarid regions, and which has a fast-growing vegetative stage and high photosynthetic activity during the flood season (from May to August) (Yan et al. 2015). Additionally, the species has a simple genetic structure as well as long-distance bidirectional gene flow with certain genetic barriers between populations, as observed using microsatellite data to estimate genetic information for populations across the native distribution range (Yan et al. 2016). To identify underlying characteristics of wild populations at the transcriptomic level, we sampled 80 individuals (8 individuals per population) from ten populations representing different habitats in the same region at noon from 18 to 28 June 2013 (Fig. 1). Plants were sampled at regular intervals of 25 m along transects in each population. This minimized the chance of sampling clonal individuals, as the largest clonal root network of *M. lutarioriparius* in these habitats was found to be 21.213 m in diameter (Yan et al. 2016). The fourth mature leaf from the apex of each plant was cut and immediately placed into liquid nitrogen for storage until RNA extraction.

Total RNA was isolated from each individual using Trizol reagent (Invitrogen, Carlsbad, California, United States), purified using the RNAeasy Mini RNA Kit (Qiagen, Schnackenburgallee, Hamburg, Germany), quantified with a NanoDrop 1000 instrument (Thermo Scientific) and stored at -80 °C. The mRNA of each individual was isolated from 5 µg of purified total RNA using one round of purification with oligo d (T) beads (Dynabeads®)
mRNA Purification Kit, Invitrogen). The cDNA libraries were prepared with the NEBNext Ultra RNA Library Prep Kit for Illumina (New England BioLabs). First-strand cDNA was synthesized using random hexamer-primed reverse transcription. After second-strand cDNA synthesis and adaptor ligation, approximately 450-bp cDNA fragments were isolated using Ampure XP beads (Beckman). The isolated cDNA fragments were amplified in 10 PCR cycles. The concentration and size of the library were assayed using the Agilent 2100 Bioanalyzer (Agilent Technologies) and Qubit® 2.0 fluorometer (Life Technologies), respectively, and 100-bp paired-end sequencing was performed on an Illumina HiSeq 2500.

**Pre-processing, quantifying gene expression, and calling SNPs**

The RNA-Seq data were filtered and trimmed to control the quality of raw reads using FASTQC (http://www.bioinformatics.bbsrc.ac.uk/projects/fastqc/) and the FASTX-Toolkit (http://hannonlab.cshl.edu/fastx_toolkit). Because of the unstable sequence content in the first 9 bases and certain sharp declines in quality in the last 11 bases, the data were trimmed for all samples, and a total of 231 Gb of sequence was generated from 2,457,041,020 raw reads (Table S1). Although no reference genome sequence was available for *M. lutaria*riparius, Xu *et al.* (2015) assembled a high-resolution reference transcriptome consisting of 18,503 unigenes based on reference genomes of related species (*Brachypodium distachyon, Oryza sativa, Sorghum bicolor*, and *Zea mays*). Therefore, the trimmed reads of each individual could be mapped to the Bowtie-build index of the reference transcriptome of *M. lutaria*riparius with an N50 of 1425 bp, and the total mapped reads were calculated using TopHat with the default settings (Langmead *et al.* 2009; Trapnell *et al.* 2009). The proportion of the reference transcriptome that was detected in each individual ranged from 55.0% to
87.4%, with an average of 74.5% for the 80 individuals of *M. lutarioriparius*. After eliminating two individuals with low coverage (<60%), the transcriptomic coverage of remaining individuals (78 samples) subjected to further analysis ranged from 65.1% to 87.4% (Fig. 2).

The expression abundance of each gene was described in terms of the expected fragments per kilobase of transcript per million fragments mapped (FPKM) using Cufflinks (v2.2.1) with the minimum number of fragments needed for new transfrags set at 2 (Trapnell *et al.* 2010). To filter out extreme values, the FPKM values per gene were added and then log₂ transformed (Xu *et al.* 2013). The transformed values were then subjected to a quartile analysis to filter out values greater than 1.5 times the inter-quartile range. After a quartile analysis of the filtered values, 16,225 transcripts were obtained with at least 40 log₂-transformed FPKMs larger than 0.

Hidden paralogous genes might lead to spurious SNPs and an excess of heterozygosity; therefore, the READS2SNP software was used to exclude spurious SNPs (Gayral *et al.* 2013). First, raw data were mapped onto the reference transcriptome using the BWA software (v0.7.12) with mismatch penalty set at 2, gap open penalty set at 6, and gap extension penalty set at 4 (Li & Durbin 2009), and then the aligned reads were sorted and indexed using SAMtools (v1.3.1) with default settings (Li *et al.* 2009). The BAM files for each individual were used to screen putative paralogous loci with a likelihood ratio test (Gayral *et al.* 2013). Candidate SNPs were identified with READS2SNP (Gayral *et al.* 2013) and SAMtools (Li *et al.* 2009). To ensure the accuracy of SNPs identification, only SNPs identified by both methods were kept, and SNPs with a quality score ≤ 10, missing data, or a minor allele This article is protected by copyright. All rights reserved.
frequency (MAF) ≤ 0.05 were deleted. All subsequent analyses were carried out on the remaining high-quality SNPs.

Genetic variation and population structure based on SNP data

Based on the SNP data, the nucleotide diversity per site (π) (Nei 1983), observed heterozygosity (H₀) (Brookfield 1996), Nei’s genetic diversity (Hₑ) (Nei 1983) and genetic differentiation (Fₛₜ) (Nei 1973) of the transcripts in each population were calculated using the formulas below: π = \frac{\sum_{i=1}^{n} (\sum_{j=1}^{x_i} x_j \pi_{ij})}{L}, where π_{ij} is the proportion of different nucleotides between the ith and jth sequences, x_i and x_j are the frequencies of these sequences, respectively, and L is the length of the gene sequences; H₀ = \frac{\sum_{i=1}^{m} (1 if a_{i1} ≠ a_{i2})}{n}, where m is the number of individuals in the population and a_{i1} and a_{i2} are the alleles of individual i at the target locus; Hₑ = 1 - \sum_{i=1}^{n} p_i^2, where n is the number of alleles at the target locus and p_i is the allele frequency of the i allele at the target locus; Fₛₜ = \frac{H_T - H_S}{H_T}, where H_S is based on the expected heterozygosity in subpopulations and H_T is based on the expected heterozygosity for the overall total population (Nei 1973); H_S = \frac{H_{E1} \cdot N_1 + H_{E2} \cdot N_2 + \cdots + H_{E_n} \cdot N_n}{N_n}, where H_{E_n} is the expected heterozygosity of the population n and N_n is the number of population n; and H_T = 1 - \sum(p^2 + q^2), where p and q represent the allele frequency of the total population.

The genetic structure of the species based on SNP data was investigated by Principal component analysis (PCA) and Bayesian clustering. The PCA was performed for 78 individuals using the program SMARTPCA of the EIGENSOFT software package (v4.2) (Patterson et al. 2006). Bayesian clustering was performed with a series of predefined populations (K = 2-10) using STRUCTURE v2.3 (Pritchard et al. 2000). Overall, 500,000
Markov chain Monte Carlo (MCMC) iterations were employed after a burn-in period of 100,000 iterations. Each $K$ value was repeated five times to assign individuals to clusters with the maximum likelihood. Structure Harvester was used to detect the optimal value of $K$ based on the recommended $\Delta K$ (Earl & Vonholdt 2012).

Mantel tests were performed in IBD 2.0 (Bohonak 2002) with 10,000 permutations between the pair-wise genetic distance [$F_{ST}/(1-F_{ST})$] and geographic distance (km) in all populations. We used the nonparametric Analysis of Molecular Variance (AMOVA) to describe the variability within and among populations and loci. A hierarchical analysis of variance and locus-by-locus analysis was performed with ARLEQUIN 3.5.1.2 (Excoffier & Lischer 2010), and significance tests were performed for 1,000 permutations.

**Variation in gene expression and population differentiation based on FPKM**

Xu et al.’s (2015) method was used to evaluate variation in gene expression in the populations. The population gene expression ($E_p$) was calculated as the average FPKM of the individuals sampled from the population using $E_p = \frac{\sum_{i=1}^{n} E_i}{n}$, where $n$ represents the number of individuals sampled from the population and $E_i$ represents the FPKM of a given gene of the $i$th individual in the population. The variation of gene expression in the population, which is denoted as the expression diversity ($E_d$), was calculated using $E_d = \frac{\sum_{i=1}^{n} |E_i - E_p|}{(n-1)E_p}$.

PCAs and Pearson correlation analyses were performed between individuals to characterize the relationships among the 78 samples based on the normalized FPKMs of each gene. The PCAs were performed in the R 3.2.1 missMDA package (Josse & Husson 2016) using the normalized FPKMs of each gene. The Pearson correlation coefficients among...
individuals were calculated in the R 3.2.1 Hmisc package using the normalized FPKMs of
each gene. Then, to estimate the gene expression relationship among populations ($E_p$
similarity), we calculated Pearson’s correlation coefficients based on the average correlation
genomes of individuals and used the method proposed by Wolf et al. (2010) for
standardization. Mantel tests were performed in IBD 2.0 (Bohonak 2002) with 10,000
permutations between $E_p$ similarity and geographic distance (km) in all populations.

The Kolmogorov-Smirnov test ($K-S$ test) as implemented in R 3.2.1 was applied to
measure the differentially expressed genes (DEGs) among the populations. To further
identify genes that were differentially expressed among the 10 populations, one-way
ANOVAs were performed for 16,225 genes (transcripts) of the $M. lutaria riparius$
transcriptome obtained from the 78 individuals of 10 populations, and the Bonferroni
correction was used to adjust each $P$ value with a multiplier of 16,225. During the one-way
ANOVA, the requirements of normality and homogeneity of variance were fulfilled.
Adjusted $P$-values of 0.05 were used to identify genes that were significantly differentially
expressed among populations.

Additionally, we also identified differentially expressed genes among the ten
populations using the DESeq package (v1.26.0) because DESeq is for pairwise comparisons,
whereas ANOVA is for any number of populations. First, filtered and trimmed reads were
aligned to the reference transcriptome using Bowtie2 (Langmead et al. 2009) and TopHat
(Trapnell et al. 2009). Then, Cufflinks, cuffmerge (Trapnell et al. 2012), and HTSeq (Anders
et al. 2015) were used to calculate the read counts. DEGs among populations were then
identified according to a false discovery rate (FDR) of 5% (Anders & Huber 2010). The
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significant DEGs for each pair-wise comparison were analyzed using the functional
annotation tools of clusterProfiler in R (Yu et al. 2012) at FDR < 0.05.

Detecting the relationship between expression diversity and nucleotide diversity among
populations

To detect if the gene expression diversity was consistent with the nucleotide diversity for the
16,225 genes within and among populations, we calculated the correlation coefficients ($r$) of
the $E_d$ and $\pi$ for these genes in each population. Finally, a Pearson’s correlation analysis was
performed between the median $E_d$ and $\pi$ of each population to explain the relationship
between genetic variation and gene expression diversity. R 3.2.1 was used for the analysis
and $P$-values were estimated from 1000 permutations.

Results

Genetic variation and population structure based on SNP data

Overall, 12,285 SNPs from 5,920 transcripts were identified across the 78 individuals. The
average nucleotide diversity ($\pi$) of the populations was 0.000434 (0.000418
(LU24)-0.000444 (LU5)) (Table 1). The majority of transcripts harbored a low level of
genetic variation (Fig. 3A). The $\pi$ quantiles in each population were similar (Fig. S1A), and
the $\pi$ distributions were similar among the following seven populations according to the $K$-$S$
test: LU10, LU17, LU5, LU7, LU14, LU19, and LU24 (Table S2). The $\pi$ distributions of
LU4 and LU9 were both significantly different from these seven populations, and in addition

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the distribution of $\pi$ in LU23 was significantly different from the populations LU7, LU14, and LU19 (Table S2). The genetic diversity ($H_O$ and $H_E$) of the 10 populations varied from 0.4636 (LU10) to 0.5144 (LU4) and from 0.3596 (LU10) to 0.3704 (LU9), respectively (Table 1). The pair-wise $F_{ST}$ values among the 10 populations ranged from 0.0274 to 0.0578 (Table S3).

The STRUCTURE analysis revealed a consistent increasing trend in the mean lnP($K$) value from $K = 2$ to $K = 10$, and low variance was observed across replicate runs. $K = 3$ was found to be the most appropriate $K$ value ($\Delta K$), which suggested that a model with three gene pools captured a major split (Fig. S1B). The PCA showed that the populations could cluster in three groups that corresponded to populations in the western, central, and eastern portions of the Yangtze River (Fig. 3B). Moreover, a significant association was observed between the level of genetic differentiation and geographic distance among the 10 populations by IBD analysis ($r = 0.318, P = 0.033$) (Fig. 3C).

A locus-by-locus AMOVA of the SNP data (12,285 loci) showed that only 0.83% of the total genetic variation partitioned in the 10 $M$. lutaria riparius populations was attributed to differences among populations ($d.f. = 9; P < 0.0001$), while 99.17% was attributed to the differences within populations ($d.f. = 150; P < 0.0001$) (Fig. 3D).

**Variation in gene expression within and among populations based on FPKM**

For the distribution of expression density, the $E_p$ shapes showed a right-skewed distribution and peak at the expression level intervals of 10-30 FPKM (Fig. 4A), and the $K$-S test showed that 43 pairs (excluding the pairs LU19-LU5 and LU19-LU4) were significantly different.
Based on the quantiles of the seven populations, the $\log_2 E_p$ quantile shifted down in LU23 and LU24 and up in LU9 (Fig. S2A). The $E_d$ of these populations also showed a right-skewed distribution and peak at 0.2 - 0.3 (Fig. 4B). The average $E_d$ values of 10 populations ranged from 0.304 (LU24) to 0.426 (LU19) (Fig. S2B). Each pair-wise comparison of the $E_d$ among the 10 populations was significantly different (Table S5).

The PCA analysis of the expression level of the 16,225 genes from all individuals showed that these individuals were distributed to some degree based on their geographic locations (Fig. 4C). Moreover, a significantly negative correlation was observed between the population similarity and geographic distance among the 10 populations ($r = -0.437, P = 0.003$) (Fig. 4D). According to the one-way ANOVA, most of variation in the gene expression occurred within populations (69.64%), although the variation range was wider among populations ($0.80e^{-4}$-32.59) than within populations (0.02-26.47). 1,570 genes showed significantly differentiated expression, with the majority of differences occurring among populations (FDR < 0.05) (Fig. 4E). All significant enriched terms for each pair-wise comparison are listed in Table S6.

**Relationship between gene expression variations and genetic variations within and among populations**

A comparison of variation in nucleotide diversity and gene expression revealed a contrasting pattern within and among populations. At the genes level, the relationship between the $E_d$ and $\pi$ of 16,225 genes in each population was significantly negatively correlated ($r = -0.03 \sim -0.10; P < 0.0001 \sim 0.019$) (Fig. 5). At the population level, the relationship between average

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$E_d$ and $\pi$ of each population was significantly positively correlated ($r = 0.054; P = 0.046$) (Fig. 6A). Among populations, the expression similarity matrix (Table S7) was significantly negatively correlated with the genetic distance matrix ($r = -0.576; P = 0.001$) (Fig. 6B).

**Discussion**

Transcriptome analysis performed across the natural range of a species can reveal a broad and detailed picture of the species’ genomic response to environmental differences (Gibson 2008; Whitehead et al. 2012). Our results for *M. lutarioriparius* provide several notable insights. First, variation among transcriptomes revealed population structures consistent with those previously reported from nuclear microsatellites data. Specifically, low levels of genetic differentiation among populations were found by both transcriptome and microsatellite data, suggesting high levels of gene flow or recent historical connections among certain populations (McGlashan et al. 2001). Second, differentiation among populations was greater for gene expression data than for nucleotide diversity. Variation in gene expression is affected by the environment but has at least some heritable component (Ellison et al. 2011; Guggisberg et al. 2013; Nabholz et al. 2014). Thus, gene expression data is a compelling complement to SNP markers and has the potential to use as an additional tool to explore population genetics. Moreover, gene expression diversity and nucleotide diversity are significantly negatively correlated within all populations, while they are positively correlated among populations. These differences could provide insights into driving mechanisms underlying plant adaptations to natural habitats.

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Based on the population transcriptomes, *M. lutarioriparius* had a lower nucleotide diversity (\(\pi = 0.000434\)) than other crop grasses, such as *Z. mays* (Wright *et al*. 2005; Hufford *et al*. 2012) or *Pennisetum glaucum* (Clotault *et al*. 2012), though similarly low nucleotide diversity has been reported in *Oryza barthii* (Nabholz *et al*. 2014). The alternate measure of genetic diversity \(H_E\) was also lower than the diversity estimated by microsatellite analyses across the entire distribution range (\(H_E\) Transcriptome: 0.3596-0.3704 vs. \(H_E\) Microsatellites: 0.682-0.786) (Yan *et al*., 2016). The low nucleotide variation and genetic diversity observed in the study could have several explanations. First, we used highly conservative criteria of SNP calling, and SNPs with a quality score \(\leq 10\), having missing data, or a minor allele frequency (MAF) \(\leq 0.05\) were all eliminated, which decreases the number of SNPs from transcripts in the study. Second, RNA-Seq analysis attempts to reconstruct the transcribed sequence of a given sample from short transcript fragments, but a bias towards highly expressed transcripts and current technological limitations often mean that the transcriptome cannot be retrieved in full (Novembre & Stephens 2008). Third, transcribed regions of genes targeted by RNA-Seq are likely subject to stronger negative selection than introns or intergenic regions (Gossmann *et al*. 2010). Although SNPs from RNA-seq data might therefore underestimate genetic variation, they have several advantages for population genetics, including higher genomic density, data quality, reproducibility, and genotyping efficiency (Schunter *et al*. 2014).

The genetic differentiation of *M. lutarioriparius*, as determined by SNPs was much lower than that of the close relative *Miscanthus sinensis* (Clark *et al*. 2015). This can be explained by distinct differences in the species’ ecologies, such as the narrow distribution. 

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along riparian corridors and the frequent human activity in the native habitats of *M. lutarioriparius*, which has been shown to increase gene flow among populations (Yan *et al.* 2016). From the evolutionary and ecological perspectives, the lower genetic differentiation may have been caused by geographic variations in the Yangtze River region, which indicates that the species was exposed to stressors or environmental factors that produced local genetic differentiation and genetic homogeneity. Critically, natural selection that favored adaptations to local environmental conditions also affected the genetic differentiation of the populations (Slatkin 1987).

We could demonstrate that SNPs from RNA-Seq data revealed local adaptations by plants to different environments. Specifically, population LU23 had a different distribution of \( \pi \) compared with the other populations (Table S2), and the results further demonstrated the genetic disconnection of LU23 from adjacent populations (Yan *et al.* 2016). Moreover, the \( \pi \) distributions showed that several populations, such as LU4 and LU9, were considerably different from the other populations, which suggests that different environments could cause the divergence of populations under non-controlled conditions (Cheviron *et al.* 2008). LU4 is located close to Dongting Lake and endures flood stress throughout the growing season; LU9 is located at the junction of Dongting Lake and the Yangtze River; and LU23 is located along a dam, which is the source of intensive anthropogenic activities. Thus, the study of transcriptomic variation could provide new insights into the adaptive strategies of *M. lutarioriparius* in distinct geographical regions, which will help in the designation of conservation units based on the degree of variation among populations.

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Variation in gene expression is produced by a combination of genetic and environmental factors, and can be very sensitive to environmental differences; therefore, it has the potential to reveal early phases of species divergence (Gibson 2008; Wolf et al. 2010). Indeed, all $E_p$ and $E_d$ pairs of the 10 populations were significantly different, whereas the levels of nucleotide diversity in the populations were almost indistinguishable. Studying expression patterns using transcriptome techniques therefore represents an important complementary approach for studying population divergence, even though the relative contribution of such gene expression divergence to overall species differentiation remains unclear, and should be addressed in future studies.

The PCA of gene expression revealed strong differentiation of the population structure, for example, LU5, LU9, LU14, and LU19 were clearly distinguished with the longitude and latitude of the populations. In addition, we found a significant pattern of isolation by distance (IBD) at the kilometers scale, and increasingly strong correlations between genetic distance/expression similarity and geographic distance were observed for microsatellite data ($r = 0.295$) (Yan et al. 2016), SNP data ($r = 0.318$), and gene expression data ($r = -0.437$). The ANOVA of gene expression showed the same pattern as the AMOVA of genetic variation, with the majority of variation occurring within populations. According to the functional annotations for biological processes (differential expression analysis), we found that LU19 exhibits greater differentiation than the other populations, while LU4 and LU5 did not show significantly enriched terms compared with those of the other populations. All of the above analyses suggest that variation in gene expression is more sensitive to detect weak differentiation than genetic diversity (Microsatellite and SNP).

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Intriguingly, gene expression related to photosynthetic processes was significantly enriched between LU4 and LU14, LU7 and LU23, LU9 and LU23, and LU14 and LU23. In addition, the genes responding to temperature stimuli were enriched between LU9 and LU14 as well as LU14 and LU19. Between LU10 and LU19, the responses to temperature stimulus, hydrogen peroxide and reactive oxygen species were enriched. Specifically, the differential gene expression was primarily observed among populations and not within populations, and this pattern appears to have been the result of natural selection (Whitehead & Crawford 2006a; 2006b). As such, analyses of gene expression in population genetics research could be used to more precisely characterize population differentiation among populations and identify genes influenced by natural selection.

Variation in gene expression among populations might result from genetic, environmental, developmental or random biological effects, which are important for adaptive evolution (Oleksiak et al. 2005). Understanding the patterns of gene expression and genetic variation in populations from different habitats could shed light on the plant responses to novel environments via variations in allelic properties and/or gene expression diversity. We found a significantly negative correlation between nucleotide diversity and gene expression diversity at the gene level in all populations (Fig. 5), which suggests that the expression and nucleotide diversity of genes have a reciprocal relationship when a gene adapts to an organism’s surroundings. The pattern might be attributed to Miscanthus suffering from the chromosomal duplication in its evolution history (Hodkinson et al. 2015). Indeed, duplicated genes usually increase gene expression diversity, but with slightly weaker genetic diversity than single-copy genes (Gu et al. 2004; Jordan et al. 2004). Additionally, divergence in gene
expression may well be an important contributor to adaptation in different environments and could thereby maintain the stability of phenotype in a population via genetic canalization (Wagner et al. 1997; Wilkins 1997).

Plants respond to stressors either via plasticity (i.e., phenotypic changes that do not depend on immediate heritable genetic change) within the lifetimes of individuals or via evolutionary adaptations over multiple generations (Harrisson et al. 2013; Lempe et al. 2013). Regardless of the pathways involved, the response depends on altering intrinsic materials (e.g., gene expression or allelic frequencies) responsible for adaptations that enhance the fitness and survival of individuals. For example, variation in DNA sequence (genetic variation) likely causes changes in gene expression, and in turn variation in gene expression reflects underlying genetic variations. In the study, we found a significantly positive correlation between nucleotide diversity and gene expression diversity (Fig. 6A), and a significantly negative correlation between similarity of expression diversity and genetic distance among populations (Fig. 6B), which indicate gene expression variation represents an important component in maintaining a stable population (Roger & Bernatchez 2005; 2007; Scott et al. 2009; Fu et al. 2012; Ackermann et al. 2013; Leder et al. 2015). Thus, inter-individual variations in gene expression are likely heritable and provide genomic evidence of the maintenance of genetic differentiation via natural selection as the natural counterpart of genetic variation.

In conclusion, gene expression data can be used as a powerful tool to explore population genetics by comparing the diversity patterns obtained from SNPs and expression data. The results obtained using SNPs from the transcriptome were consistent with results from...
microsatellites in the nuclear genome, but the gene expression data revealed additional levels of population structure not captured with conventional methods. The reciprocal relationship between genetic variation and gene expression is an important pattern that reflects the responses of a species to its environment, and both are likely involved in local adaptation. In short, RNA-Seq is a useful tool for assessing the variations in gene expression that underlies phenotypic differences among wild populations, and the SNP library and extensive transcriptome sequences developed in this study will facilitate future functional analyses of molecular signatures capable of indicating adaptations in populations of *M. lutarioriparius*.

**Acknowledgements**

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Data accessibility

Author contributions:
Sang T and Li JQ funded project and designed the study. Yan J & Song ZH conducted analyses of data and wrote the manuscript. Kang LF & Xing SL carry out the experiments and data analysis. Züst T, Greimler J, Liu W, Xu Q and Zhu CY contributed to the manuscript revisions.

Figure legends

Figure 1 Ten locations of the studied Miscanthus lutariaoriparius populations across the species distribution range. These populations were sampled because of their special geographic position, genetic variation or adaptability (Yan et al. 2012; 2015; 2016). The population identities correspond to the codes used in Yan et al. 2016. In the upper left corner, the gray colors in the map correspond to locations where M. lutariaoriparius has been transplanted from warm and wet sites to semiarid regions and was able to adapt to the environment (Yan et al. 2012). The degree of suitability for growing M. lutariaoriparius in those environments is indicated by darker shades of gray.

Figure 2 Coverage of the reference transcriptome of the 78 M. lutariaoriparius samples. The dots represent an individual with a certain number of reads after rigorous quality testing. The regression line is based on the best least squares fit to the function at $r^2 = 0.269$.

Figure 3 Genetic analysis based on SNPs in M. lutariaoriparius: (A) The distribution of nucleotide diversity; (B) PCA of SNP data in each individual; (C) Relationship between genetic distance and geographic distance between each two populations; and (D) Distribution of genetic variation within populations and among populations based on SNP data.

Figure 4 Expression analysis based on FPKM in M. lutariaoriparius: (A) The distribution of population expression ($E_p$); (B) Expression diversity ($E_d$) relationship between genetic distance and geographic distance; (C) PCA of gene expression (FPKM); (D) Relationship between population expression similarity and geographic distance; and (E) Distribution of expression variation within populations and among populations based on FPKM. Genes with differential expression are indicated by red dots.

Figure 5 Correlation between the nuclear diversity and expression diversity of all genes in each population.

Figure 6 Transcriptomic patterns of M. lutariaoriparius across the distribution range: (A) relationship between nucleotide diversity and expression diversity; and (B) relationship between genetic distance and population expression similarity.
Supporting information

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

Table S1 The RNAseq data after being filtered and trimmed.

Table S2 K-S test of nucleotide diversity (π) between populations.

Table S3 The pair-wise $F_{ST}$ values among the 10 populations based on SNP data.

Table S4 K-S test of population expression ($E_p$) between populations.

Table S5 K-S test of expression diversity ($E_d$) between populations.

Table S6 Significantly enriched GO terms in biological processes of DEGs between populations.

Table S7 Similarity of population expression ($E_p$) among populations. Pearson correlation coefficients based on scaled $E_p$ of 10 populations.

Fig. S1.Summary of genetic variation and gene structure in *M. lutarioriparius* (A) nucleotide diversity, (B) population genetic structure based on STRUCTURE.

Fig. S2.Summary of gene expression in *M. lutarioriparius*: (A) population expression ($E_p$), (B) expression diversity ($E_d$).
Table 1 Measures of genetic diversity for each population of *M. lutarioriparius* based on population transcriptome data.

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