QTL mapping of leaf angle on eight nodes in maize enable the optimize canopy by differential operating of leaf angle at different levels of plant

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Running title: Regulatory network for leaf angle on eight nodes in maize

Highlight: QTL were identified for maize Leaf angle on different nodes, which has illuminated the genetic control networks of maize LA and empowered canopy ideotype design by manipulate leaf angle at individual leaves.

Abstract

Leaf angle (LA) is one of the most important canopy architecture traits of maize (Zea mays L.). To date, there is an urgent need to characterize the genetic control of LA at multiple nodes to bridge the information gap remain in optimizing canopy architecture for maximum yield at different canopy levels. In this study, through the cross between B73 (compact plant architecture) and SICAU1212 (expanded plant architecture), 199 derived RIL families were used to perform QTL mapping for LA from eight leaves at different nodes in three environments, utilizing single-environment analysis and joint mapping. Combining the results of two mapping strategies, we identified 15 common QTL associated with LA at eight nodes. The phenotypic variation explained by the individual QTL ranged from 0.39% to 20.14% and the number of leaves controlled by a single QTL varied from 1 to 8. Among them, QTL $qLA2.1$ and $qLA5.1$ simultaneously controlled LA of all the eight nodes; however, $qLA2.2$ only affected that of 1stLA. The total phenotypic variation explained by all QTL identified for LA at eight nodes ranged from 15.69% (8thLA) to 51.73% (1stLA). The number of QTL detected for LA at each nodes ranged from 4 (7thLA) to 11 (1stLA). These results provide comprehensive insights into the molecular bases of regulatory networks in LA morphogenesis, and will benefit the molecular design breeding of ideotype and further cloning of LA QTL at different plant levels in maize.

Keywords: Leaf angle, eight nodes, canopy architecture, QTL, Regulatory networks, maize
Introduction

Maize (*Zea mays* L.) is one of the most important cereal crops worldwide, and the primary goal of maize breeding programs is to generate high-yielding varieties. During the past several decades, the increase in maize yield was largely due to the increased plant density, rather than improving the potential yield per plant (Duvick, 2005; Ma *et al*., 2014b; Mock and Pearce, 1975; Russell, 1991; Tollenaar and Wu, 1999). A suite of dramatic changes in plant architecture have been observed, which play a pivotal role in adaptation to high plant density. Some key parameters of optimal plant characteristics were identified as early as in 1970s, including upright leaves, maximum photosynthetic efficiency, and small tassel size and so on (Mock and Pearce, 1975).

Early studies suggested that LA was a critical parameter of plant architecture of impact on light interception and photosynthesis, and plant breeding practices in maize have also shown that LA was an essential agronomic trait in the development and adoption of high-yielding varieties of maize. As breeders focused on improving grain yield, LA score has decreased remarkably, which has mainly shaped plant architecture from expanded to compact plant architecture (Anderson and Denmead, 1969; de Wit, 1965; Duncan *et al*., 1967; Ku *et al*., 2010a). Erect canopies can increase in light interception efficiency at higher plant densities, and eventually, leading to increased production. Comprehensive analysis of the correlation between LA trait and grain yield revealed two interesting facts: (1) although the LA had significantly decreased over the past few decades, smaller LA does not guarantee higher yield; and (2) further increase in light interception efficiency needs to vary LA at different parts of maize plant (Duncan, 1971; Lambert and Johnson, 1978; Ma *et al*., 2014a; Mickelson *et al*., 2002; Pepper *et al*., 1977; Winter and Ohlrogge, 1973; Zhang *et al*., 2017). More recently, Mantilla *et al*. (2017) has proposed that the optimization of canopy architecture could be manipulated with varying LA at different levels for maximum production potential in cereal species (Mantilla-Perez and Salas Fernandez, 2017).

With the advent of QTL mapping strategies, linkage mapping and association analysis have been conducted in maize to dissect the genetic basis of LA, and hundreds of quantitative trait loci (QTL) for LA have been identified throughout all
ten of the maize chromosomes. Among these studies, there were widely different in
number and node position of selected leaves, statistical methods of phenotype data,
types of mapping populations and QTL mapping strategies. Detailed information of
previous studies showed that research groups chose different number and node
positions of leaves for analysis. In most circumstances, three continuous leaves
including the ear leaf, the leaves above and below ear were selected for analysis (Ding
et al., 2015; Ku et al., 2016; Ku et al., 2012; Ku et al., 2010b; Li et al., 2015;
Mickelson et al., 2002; Ming et al., 2007; Shi et al., 2017; Zhang et al., 2017), and in
some instances, that of the first leaf below the flag (Pan et al., 2017; Tian et al., 2011;
Wang et al., 2017a; Yang et al., 2015b) or two leaves near the ear (Chen et al., 2015;
Hou et al., 2015). There were two kinds of statistical methods to phenotype data. QTL
mapping was performed using average values for leaves or not/the value of individual
leaf. Furthermore, different mapping populations have been adopted, such as F_{2:3}
(Chen et al., 2015; Hou et al., 2015; Ku et al., 2012; Ku et al., 2010b; Ming et al.,
2007; Yu et al., 2006), F_4 (Chen et al., 2015), RIL (Ku et al., 2016; Li et al., 2015;
Mickelson et al., 2002; Shi et al., 2017; Wang et al., 2017a; Yang et al., 2015b; Zhang
et al., 2017), Four-Way Cross Mapping Population (Ding et al., 2015), NAM (Tian et
al., 2011) and ROAM (Pan et al., 2017). Together with QTL mapping, Tian et al.
(2011) and Pan et al. (2017) identified 203 and 10 single-nucleotide polymorphisms
(SNPs) associated with LA through GWAS studies, respectively.

Identification of actual genes responsible for LA QTL and isolation of mutants
with altered LA is the critical step to unravel the genetic and molecular mechanisms
underlying maize LA. So far, only two LA QTL, ZmTAC1 (Ku et al., 2011) and
ZmCLA4 (Zhang et al., 2014) were identified, and six LA mutants, liguleless1 (lg1)
(Moreno et al., 1997), lg2 (Walsh et al., 1998), Liguleless3-O (Lg3-O) (Muehlbauer et
al., 1999), Liguleless narrow-Reference (Lgn-R) (Moon et al., 2013), droopingleaf1
(drl1) and drl2 (Strable et al., 2017) have been cloned. Specifically, lg1, lg2 and
Lgn-R mutants exhibit a defect in ligule and auricle tissues and a decrease in leaf
angle (Harper and Freeling, 1996; Moon et al., 2013; Sylvester et al., 1990; Walsh et
Notably, \textit{LG1}, \textit{LG2} and \textit{LGN} were shown to act in a common pathway involved in ligule development (Harper and Freeling, 1996; Moon \textit{et al.}, 2013). Similarly, the \textit{Liguleless3-O (Lg3-O)} mutant also develops a decreased leaf angle, which might be due to the defect in transformation of blade-to-sheath at the midrib region in leaf (Fowler \textit{et al.}, 1996; Muehlbauer \textit{et al.}, 1997; Muehlbauer \textit{et al.}, 1999). Nevertheless, the \textit{drl} genes are required for properly development of leaf and leaf support tissues, and for restricting auricle expansion at the midrib, and the LA in the \textit{drl1} and \textit{drl2} mutants are increased (Strable \textit{et al.}, 2017).

As mentioned above, only one or a few or average values for leaves were characterized in previous studies, which could not provide valuable information for manipulate LA at canopy level. In the present study, the QTL mapping for LA at eight consecutive leaves were performed with an RIL population, providing valuable information in support of canopy ideotype design and fine mapping of QTL controlling maize LA at different canopy levels.

\textbf{Materials and Methods}

\textit{Mapping populations and Field experiment}

The recombinant inbred line population (RIL) was derived from a cross of B73 and SICAU1212, as described previously (Yang \textit{et al.}, 2016; Yang \textit{et al.}, 2015a). The parent B73 with erect leaves is widely used as elite line from the stiff stalk heterotic group and has been partly attributed to the changes in LA of maize varieties since 1970, and another parent SICAU1212 with extremely expanded leaves that have been developed from a waxy maize landrace Silunuo by continuously self-pollinating 10 times, which was cultivated at least 100 years ago (Tian \textit{et al.}, 2008). 199 of the 325 RIL families randomly subsampled from their initial population were used in the present study.

The 199 RIL families and their parent lines were grown in a complete randomized block designed with two replications in three distinct environments of China. The three environments locate at Jinghong of Yunnan province (21°57'N, 100°45'E, elevation 551 m) in 2015 (15JH), Chengdu of Sichuan province (30°43'N, 106°10'E, elevation 520 m) in 2016 (16CD), and Yangzhou of Jiangsu province (32°45'N, 119°09'E, elevation 52 m) in 2017 (17Ye).
103°52'E, elevation 500 m) in 2016 (16CD) and Guiyang of Guizhou province
(26°29'N, 106°39'E, elevation 1277 m) in 2016 (16GY), respectively. Fourteen plants
were cultivated for each single-row plot with a planting density of 52,500 plants ha⁻¹
in all environments. Row length was 3.0 m, and row spacing was 67 cm. Field
management was the same as the standard cultivation management in accordance with
growing season.

Phenotypic measurements and analysis
Five plants from the middle of each plot were used to evaluate the phenotype at 10 d
after pollen shed. The leaf angle (LA) of eight consecutive leaves below tassel from
each plant was measured as the angle between the vertical line and the base of leaf
midrib (Hou et al., 2015). LA of the first leaf was abbreviated to 1stLA, LA of the
second leaf below tassel abbreviated to 2ndLA, and so forth. The phenotypic data of
LA was determined as the average of each family from two replications in a single
environment.

Statistical analysis and Pearson’s phenotypic correlations were computed
employing IBM SPSS Statistics version 20.0 software (http://www.spss.com).

Broad-sense heritability (h²) for each LA was estimated as

\[ h² = \frac{\sigma_g^2}{(\sigma_g^2 + \sigma_{ge}^2/n + \sigma^2/nr)} \]

where \( \sigma_g^2 \) is the genetic variance, \( \sigma_{ge}^2 \) is the
interaction variance between genotype and environment, \( \sigma^2 \) is the error variance, \( n \) is
the number of environments and \( r \) is the number of replications in each environment
(Hallauer et al., 2010).

Linkage map and QTL mapping
We re-constructed the linkage map of RIL population that is described in a recent
report (Yang et al., 2016). Briefly, the current linkage map was shortened to have 106
SSRs and 154 indels, which spanned 1133.57 cM for the whole genome, and was
drawn in the MapChart software version 2.3 (Voorrips, 2002). The QTL for each LA
were detected by including composite interval mapping (ICIM) (Li et al., 2008; Li et
al., 2007) using the QTL IciMapping software, version 4.0 (http://www.isbreeding.net/)
in a single-environment. A walking speed of 1.0 cM was selected for QTL mapping,
and the probability in the stepwise regression was set to 0.001. Threshold LOD scores were computed by 1,000 permutations, and a type I error was set at 0.05. The joint mapping, epistatic interaction and QTL by environment interaction (QEI) detection were performed by a mixed-model based composite interval mapping (MCIM) (Wang et al., 1999) using QTLNetwork software version 2.1 (Yang et al., 2008). The testing window size, walk speed and filtration window size of genome scan configuration was set to 10, 1 and 10 cM, respectively. 1,000 permutations at a significance level of $P = 0.05$ was performed to calculate the threshold for declaring the presence of a significant QTL. Positive additive effect indicates that the allele resulting in increased LA is from the compact parent B73, whereas negative effect showing the allele from expanded parent SICAU1212. The QTL for LA of the leaves were considered identical only upon the confidence intervals of QTL were overlapped. The name of the QTL was assigned as ‘q’ followed by ‘LA’, ‘maize chromosome on which the corresponding QTL locates’, ‘.’, and ‘serial number of QTL’. In addition, QTL with PVE (%) $> 10\%$ were declared as major QTL. QTL were detected repeatedly in more than one environments were considered as stable QTL.

**Results**

*Phenotypic performance of LA from eight consecutive leaves*

The phenotypic values of LA were analyzed in RIL families and their parent lines cultivated in three distinct habitats (Table 1). It was obvious that each LA in B73 was significantly different from that in SICAU1212 ($P < 0.01$). All eight leaves tested in the parent line B73 display an almost vertical angle, whereas the other parent line SICAU1212 has more horizontal leaf orientations. In addition, the strong variation extent in all eight LAs can be detected in the RIL lines, although it displays a continuous distribution (Table 1). Specifically, the LA of topper leaves displays a larger variation (around 11-fold) than that of lower leaves (around 3-fold) in the RIL population. In addition, the values of all traits in RIL families exhibited obvious transgressive segregation, indicating that the LA is under polygenic quantitative genetic control.
The estimated broad-sense heritability ($h^2_B$) for LA in eight leaves ranges from 79.47 to 83.46 % (Table 2), indicating that genetic factors dominantly determine the formation of LA. The lines within the RIL population were significantly ($P < 0.001$) different for all traits (Table 2). Significant ($P < 0.001$) genotype × environment interactions for all traits were also observed. With the exception of 8thLA, the variances of replications for all traits were non-significant ($P < 0.05$). Therefore, the mean of two replications in one environment for each RIL family was used for single-environment mapping. The phenotypic correlation coefficients among distinct LAs within the RIL population are shown in Supplementary Table S1. It revealed that the LA of all leaves has a significantly positive correlation ($P < 0.001$) (Supplementary Table S1). Moreover, the LA of more adjacent leaves displays a much higher correlation coefficients (Supplementary Table S1).

**QTL analysis**

Using inclusive composite interval mapping, a total of 56 putative QTL for LA at eight nodes were identified, and 11 QTL were common between leaves or environments (Supplementary Table S2, Supplementary Fig.S1). Of these 11 common QTL, three QTL were identified on chromosome 5, two QTL on chromosome 3 and one QTL on chromosome 1, 2, 4, 6, 8 and 9. Five QTLs were detected in more than one environment. With the exception of $qLA6.1$ controlling 1stLA, all QTL had the negative additive effect in the single-environment analysis. The individual effect of QTL ranged from 5.62% to 20.14% and the number of leaves controlled by a single QTL varied from 1 to 8. The number of QTL detected for LA at each node ranged from 3 (8thLA) to 8 (1stLA). The total phenotypic variation explained by all QTL identified for LA at eight nodes ranged from 15.69% (8thLA) to 51.73% (1stLA).

Meanwhile, forty-four significant QTL for LA at eight nodes were detected in joint analysis, and 12 QTL were common between leaves (Supplementary Table S3, Supplementary Fig.S1). Of these 12 common QTL, eight were identical to those of the QTL identified by single-environment QTL analysis. The other 4 QTL were specific for the joint mapping, which are QTL $qLA2.2$ on chromosome 2, $qLA4.2$ on...
chromosome 4, \( q_{LA5.4} \) on chromosome 5 and \( q_{LA7.1} \) on chromosome 7. These QTL controlled one, five, one and four nodes, respectively; and accounted for 2.11%, 2.306%, 1.29% and 2.34% of phenotypic variance, respectively.

All QTL except \( q_{LA2.2} \) and \( q_{LA4.2} \) had a negative additive effect. Together, the results suggested that most of the alleles with a contribution on increasing LA are segregated from SICAU1212 with expanded plant architecture. Notably, Several alleles associated with increased LA were contributed by SICAU1212.

**QEIs**

Four QTL were involved in significant QTL × environment interaction (QEI) (Table 4) through joint analysis. Three of them are common QTL (\( q_{LA5.3} \)), which affected the LA of the 2nd, 3rd and 7th leaves simultaneously, and the additive × environment interactions for LA were responsible for 1.89-2.46 % of phenotypic variation. The other QTL (\( q_{LA5.2} \)) association with the LA of 6th leaf, and the effect of additive × environment interaction was 1.50 %.

**Epistatic interaction**

A total of twenty (\( P < 0.05 \)) epistatic interactions with additive-by-additive effects were identified for eight LAs, and individual variance ranged from 0.39 to 3.54% (Table 5). Three types of epistatic interactions were identified, including interactions occurred within the genetic region of the QTL identified, between significant QTL and non-significant QTL region, and in non-significant QTL regions. Moreover, the number of LA that individual epistatic interaction affected varied, ranging from 1 to 4. For example, the interaction between \( q_{LA5.3} \) and \( q_{LA7.1} \) affected 1stLA to 4thLA, the interaction between marker intervals of chr9-90756–mmc0051 and chr10-77445–umc1336 affected just 2ndLA. On the other hand, the interactions identified for the eight LAs also varied, ranged from 0 to 6. For example, five epistatic interactions were identified for 2ndLA, while no epistatic interaction was identified for 8thLA. In addition, the interaction between epistasis and the environment was not detected in this study.
Discussion

LA exhibits extensive phenotypic variation in single maize plant and population

Both allelic LA in maize population and LA between nodes of a single plant exhibits extensive phenotypic variation. For LA in maize population, among plant architecture-related traits, LA was the second highest degree of variation traits, which second only to tassel branch number (Pan et al., 2017). Early evidence suggested that the upper LA of 26 parental lines for maize NAM population ranged from approximately 30° to 80° (Tian et al., 2011). Likewise, in a recent study, the range of allelic LA in ten maize elite inbred lines was up to four fold (Pan et al., 2017). Additionally, the allelic LA of our mapping population displayed larger variation, ranging from 3-fold (1stLA in 2015JH) to 11-fold (8thLA in 2016CD) change. It's worth mentioning that transgressive phenotypes were observed in various segregating populations.

Phenotypic variation is the variability in phenotypes that exists at different nodes in a single maize plant. Taking a high-yielding maize hybrid (Pioneer 335) as an example, the average LA of three leaves above ear, three ear leaves and three leaves below ear were 22.9°, 32.8° and 41.9°, respectively, and there was ~2.0-fold change between uppermost and lower LA (Ma et al., 2014b). In present study, approximately 1.6-fold difference was observed among the eight LAs of SICAU1212 parental inbred lines, while approximately 2.6-fold that of B73 (Supplementary Table S1). In summary, these results suggest that diversified genetic mechanisms were responsible for LA morphogenesis of a single plant and population.

QTL hotspot regions and pleiotropic QTL response for morphogenesis of LA

Hundreds of QTL or SNPs associated with LA were identified from various genetic backgrounds, which provided valuable genetic information. However, the difference between research methods led to problems in comparative analysis, particularly node position of selected leaves. For example, ear position of maize was frequently used as a reference position to selection leaves for LA analysis. Nevertheless, the study of the leaves of the relative positions to ear leaf is not conducive to the comparative analysis
of the results. In spite of that, distribution characteristics of QTL controlling LA trait still could be seen to a certain degree by comparative analysis. In present study, fifteen QTL controlled LAs of eight consecutive leaves were detected, which was highly consistent with the results of previous studies. Moreover, these QTL were not randomly distributed on whole genome, but were located on a few genomic regions. There were eight hotspot regions for LA, which are distributed on chromosomes 1, 2, 4, 5, 8 and 9. Of these QTL, the number of times detected by sixteen different LA studies ranged from 5 to 11 (Table 3). Among them, *qLA2.1*, located on short arm of chromosome 2, were detected in most previous studies (Ding *et al.*, 2015; Hou *et al.*, 2015; Ku *et al.*, 2012; Mickelson *et al.*, 2002; Ming *et al.*, 2007; Pan *et al.*, 2017; Shi *et al.*, 2017; Tian *et al.*, 2011; Wang *et al.*, 2017b; Yang *et al.*, 2015b; Zhang *et al.*, 2017). In addition, the hotspot QTL *qLA1.1*, harbors the cloned LA mutant gene, *drooping leaf1 (drl1)* (Strable *et al.*, 2017); *qLA2.1* contains *liguleless1 (lg1)* (Becraft *et al.*, 1990; Becraft and Freeling, 1991; Moreno *et al.*, 1997; Sylvester *et al.*, 1990).

Pleiotropic QTL played an important role of LA at different nodes, and tended to control the consecutive LAs. Of fifteen identified QTL in this study, approximately seventy percent QTL controlled more than two LAs. The frequency of QTL that were identified in different genetic backgrounds was positively correlated with number of LA on different nodes that were controlled by QTL. More interestingly, the distribution of these LAs controlled by pleiotropic QTL exhibited spatial-specificity. Some QTL controlled all eight LAs, some QTL controlled LA of upper layers, and some QTL controlled LA of lower layers. For example, *qLA3.1* affected LA of upper layers, 1stLA to 4thLA, while *qLA9.1* controlled LA of lower layers, 5thLA to 8thLA. It is worth noting that the fine mapping of pleiotropic QTL can be carried out for a specific leaf controlled by the QTL with stable and major effects.

*The genetic architecture of the eight leaf angles exhibits extensive diversity*

Not only the result of present study, but also that of most previous studies (Ding *et al.*, 2015; Hou *et al.*, 2015; Ku *et al.*, 2010b; Pan *et al.*, 2017; Shi *et al.*, 2017; Tian *et al.*, 2011; Wang *et al.*, 2017b; Yang *et al.*, 2015b) demonstrated that genetic factors were a
major determinant of LA traits and the heritability of LA trait estimates were largely
similar between adjacent nodes. More interestingly, QTL analysis for LA at eight
nodes had indicated that there were different sets of QTL that controlled LA at
different nodes. The result revealed a rich genetic architecture of LA trait: (i) The
results showed that the LA trait was controlled by major gene plus polygenes genetic
model with difference in detail. The number and effects size of major QTL were not
the same for different nodes, and the number of major QTL ranged 1 (8thLA) to 4
(1stLA), and the highest PVE value was 20.14%. Additionally, the major QTL had
been identified in most QTL mapping studies of LA in maize (Chen et al., 2015; Ku et
al., 2010b; Pan et al., 2017), however, a few studies fail to find major QTL (Hou et al.,
2015; Tian et al., 2011). (ii) The epistatic interaction identified for LA of different leaf
nodes showed that there were difference in absence or presence with the different
number of loci. No epistatic interactions were observed for the 8thLA, while epistatic
interaction was identified for others, ranging from 1 (5thLA) to 5 (2ndLA). (iii) the
LA at different nodes have different QTL × environment interaction (QEI). Four LA,
including 2ndLA, 3rdLA, 6thLA: QEI of qLA5.2 and 7thLA, were affected by QEI at
different levels. The rest LA had not detected QEI. Taking together, the data indicated
that the genetic architecture underlying the eight LAs exhibits extensive diversity.

*Application for canopy ideotype breeding by design at different canopy levels design*
*in maize*

The result of this research bridges the crucial knowledge gap between theory and
breeding practice. For instance, node-specificity QTL could be applied directly to the
manipulation of the leaf angle, and region-specificity QTL could shape the LA of
upper, middle or lower canopy, separately, and environment-specificity QTL could be
used in ecological idotype breeding. Additionally, other controls of leaf angle operate
could be employed by combined use of the pleiotropic QTL, based on different
effect size of QTL. It is however worth mentioning that this study not only provides
the theoretical basis of regulatory network for designing LA differently but also
unlock the door for further dissecting the LA trait in cereal crop.

Furthermore, QTL mapping of LA at multiple levels, together with our previous research of QTL controlled leaf wide on eight nodes, provides a comprehensive strategy for maize “smart canopy” breeding, which was proposed by Ort et al. (2015) for improving canopy architecture and metabolic features of leaves interacting cooperatively throughout the canopy to maximize yield potential (Ort et al., 2015; Yang et al., 2016).

Supplementary data

Table S1. Phenotypic correlation coefficients between eight LAs across three environments

Table S2. Putative QTL for LA in the RIL population through single-environment QTL mapping

Table S3. QTL for eight LAs detected in joint analysis across three environments

Fig S1. Genetic linkage map of the RILs and locations of QTLs for eight LAs in single-environment QTL mapping and joint analysis

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Reveals the Genetic Architecture of Maize Plant Growth. *Plant Physiol* **173**, 1554-1564.
| Trait  | 15JH  | 16CD  | 16GY^b  |
|--------|-------|-------|---------|
|        | B73   | SICAU1212 | RIL | B73   | SICAU1212 | RIL | B73   | SICAU1212 | RIL |
| 1stLA  | 15.11 | 87.77** | 41.80 | 17.46 | 10.95-122.05 | 13.27 | 69.65** | 33.18 | 14.63 | 13.72-105.15 | 20.04 | 71.39** | 41.13 | 15.95 | 16.17-87.64 |
| 2ndLA  | 11.63 | 84.58** | 34.57 | 13.08 | 11.35-86.85 | 14.53 | 48.97** | 27.35 | 9.62 | 12.48-68.24 | 21.26 | 62.23** | 38.26 | 13.036 | 17.40-86.73 |
| 3rdLA  | 15.78 | 80.52** | 33.68 | 12.21 | 14.05-78.44 | 16.49 | 43.82** | 26.83 | 9.13 | 12.92-63.64 | 24.78 | 52.47** | 38.15 | 12.18 | 18.41-82.90 |
| 4thLA  | 18.33 | 69.02** | 34.64 | 11.97 | 13.66-103.63 | 18.50 | 46.95** | 27.67 | 9.53 | 12.26-67.42 | 28.05 | 51.76** | 38.15 | 11.49 | 18.22-81.88 |
| 5thLA  | 23.78 | 60.84** | 37.54 | 10.58 | 13.41-83.10 | 20.54 | 50.86** | 31.02 | 9.92 | 12.44-78.84 | 29.66 | 55.72** | 40.77 | 12.05 | 19.48-83.71 |
| 6thLA  | 29.41 | 58.12** | 40.82 | 9.65 | 16.07-83.97 | 27.43 | 59.25** | 35.29 | 9.75 | 15.62-77.34 | 34.70 | 63.25** | 44.32 | 13.29 | 24.00-83.81 |
| 7thLA  | 34.00 | 49.33** | 40.02 | 8.36 | 17.29-69.70 | 34.87 | 60.82** | 37.46 | 8.76 | 16.50-67.52 | 38.8 | 67.29** | 45.04 | 12.38 | 23.08-88.90 |
| 8thLA  | 30.93 | 46.31** | 39.20 | 8.04 | 19.49-73.40 | 37.56 | 59.77** | 38.22 | 8.31 | 21.42-66.96 | 42.42 | 64.52** | 44.93 | 12.60 | 21.27-81.90 |

15JH, 16CD and 16GY represent Jinghong of Yunnan province in 2015, Chengdu of Sichuan province and Guiyang of Guizhou province in 2016, respectively, respectively.

** indicates significant level at $P < 0.01$
Table 2. Analysis of variance (ANOVA) for LA for the RIL population in three environments

| Source of variation | Mean square 1stLA | Mean square 2ndLA | Mean square 3rdLA | Mean square 4thLA | Mean square 5thLA | Mean square 6thLA | Mean square 7thLA | Mean square 8thLA |
|---------------------|------------------|------------------|------------------|------------------|------------------|------------------|------------------|------------------|
| Environment (E)     | 9966.08***       | 12394.80***      | 13288.85***      | 12193.37***      | 9536.30***       | 7790.14***       | 6213.45***       | 6187.40***       |
| Genotype (G)        | 705.19***        | 357.06***        | 292.89***        | 291.92***        | 267.25***        | 269.66***        | 238.37***        | 240.52***        |
| G × E               | 349.42***        | 197.47***        | 180.40***        | 198.24***        | 168.31***        | 181.75***        | 170.71***        | 163.32***        |
| Replication         | 2.76             | 23.50            | 180.95           | 79.70            | 29.56            | 129.28           | 3.47             | 180.63*          |
| Error               | 139.70           | 64.72            | 62.40            | 55.98            | 52.13            | 50.75            | 46.40            | 39.71            |

$h_b^2$ the broad-sense heritability

*, ** and *** indicate significant level at $P < 0.05$, $P < 0.01$ and $P < 0.001$, respectively
Table 3. Main features of the QTL for eight LAs based on single-environment QTL mapping and joint analysis across three environments

| QTL   | Bin     | 1stLA | 2ndLA | 3rdLA | 4tYLA | 5tYLA | 6tYLA | 7tYLA | 8tYLA | References b |
|-------|---------|-------|-------|-------|-------|-------|-------|-------|-------|--------------|
| qLA1.1| 1.02-1.03| Y     | -     | Y     | -     | -     | Y     | -     | -     | 1,2,3,4,5,6,10,11,12,13,15 |
| qLA2.1| 2.01-2.02| Y, J  | Y, S, J| Y, J  | Y, J  | J     | Y, S, J| Y     | -     | 2,3,5,7,8,9,10,12,13,14,16 |
| qLA2.2| 2.07    | J     | -     | -     | -     | -     | -     | -     | -     | 6,12,14      |
| qLA3.1| 3.06    | Y, J  | Y, S, G; J| Y, J  | Y, S  | -     | -     | -     | -     | 9,12         |
| qLA3.2| 3.06-3.07| G     | -     | G     | G; J  | G; J  | J     | -     | -     | 12           |
| qLA4.1| 4.05    | -     | -     | -     | -     | Y     | -     | -     | -     | 12,14        |
| qLA4.2| 4.09    | -     | J     | -     | -     | J     | -     | J     | -     | 3,10,12,14,15 |
| qLA5.1| 5.03    | Y, S; J| Y, S, G; J| Y, S, G; J| S, G; J| Y, S, G; J| S, J  | Y, S, G; J| S, G; J| 8,10,12,13,16 |
| qLA5.2| 5.04    | -     | -     | -     | J     | -     | G; J  | -     | -     | 1,2,3,5,6,12 |
| qLA5.3| 5.06-5.07| G; J  | G; J  | G; J  | G; J  | -     | J     | G; J  | -     | 3,8,9,10,12  |
| qLA5.4| 5.07-5.08| -     | -     | -     | -     | J     | -     | -     | -     | 3,10,12      |
| qLA6.1| 6.01    | Y     | -     | -     | -     | -     | -     | -     | -     | 8,12         |
| qLA7.1| 7.01-7.02| J     | J     | J     | J     | -     | -     | -     | -     | 2,12,14      |
| qLA8.1| 8.06-8.07| S; J  | -     | -     | -     | -     | -     | -     | -     | 2,7,8,10,11,12,14,16 |
| qLA9.1| 9.04    | -     | -     | -     | -     | S; J  | S; J  | S     | Y, S; J| 3,7,10,12,14 |

a Bin 1stLA – 8tYLA (1st – 8th Years) in position (cm) 

b References: 1. Li et al. (2010); 2. Zhang et al. (2011); 3. Wang et al. (2012); 4. Chen et al. (2013); 5. Sun et al. (2014); 6. Li et al. (2015); 7. Zhang et al. (2016); 8. Wang et al. (2017); 9. Chen et al. (2018); 10. Sun et al. (2019); 11. Li et al. (2020); 12. Zhang et al. (2021); 13. Wang et al. (2022); 14. Chen et al. (2023)
The capital letters Y, S and G represent QTL detected at Jinghong of Yunnan province in 2015, Chengdu of Sichuan province and Guiyang of Guizhou province in 2016, respectively. J represents QTL detected by joint analysis across three environments.

The number 1 represents Chen et al. 2015, and 2 represents Ding et al. 2015, and 3 represents Hou et al. 2015, and 4 represents Ku et al. 2016, and 5 represents Ku et al. 2012, and 6 represents Ku et al. 2010, and 7 represents Li et al. 2015, and 8 represents Mickelson et al. 2002, and 9 represents Ming et al. 2007, and 10 represents Pan et al. 2017, and 11 represents Shi et al. 2017, and 12 represents Tian et al. 2011, and 13 represents Wang et al. 2017, and 14 represents Yang et al. 2015b, and 15 represents Yu et al. 2006, and 16 represents Zhang et al. 2017.
Table 4. QTLs × Environment (QE) interactions effects for LA identified in the RIL population using QTLNetwork

| Traits | QTL   | Marker interval                  | AE1(15JH) | AE2(16CD) | AE3(16GY) | H² (ae)(%) |
|--------|-------|----------------------------------|-----------|-----------|-----------|------------|
| 2ndLA  | qLA5.3| chr5-199388–umc2198              | 1.90*     | -2.79**   | 2.05      |            |
| 3rdLA  | qLA5.3| chr5-199388–umc2198              | 1.92**    | -2.32**   | 1.89      |            |
| 6thLA  | qLA5.2| chr5-139354–chr5-160457          |           | -1.29*    | 1.50      |            |
| 7thLA  | qLA5.3| chr5-199388–umc2198              |           | -2.07**   | 2.46      |            |

AE is the additive by designated environment interaction effect

H² (ae)(%) is contribution rate of additive by environment interaction

*, ** and *** indicate significant level at P < 0.05, P < 0.01 and P < 0.001, respectively
Table 5. Digenetic epistatic QTL for LA identified in the RIL population across three environments

| Trait | QTL_i/Marker interval_i | range_i (cM) | QTL_j/Marker interval_j | range_j (cM) | AA    | H^2(aa)(%) |
|-------|-------------------------|--------------|-------------------------|--------------|-------|------------|
| 1stLA | qLA3.1                  | 55.2-59.5    | qLA8                    | 73.3-80.0    | 1.93**| 1.35       |
| 1stLA | qLA5.3                  | 97.9-103.0   | qLA7-1                  | 15.2-19.3    | 2.42**| 1.96       |
| 2ndLA | qLA3.1                  | 55.2-59.5    | qLA4-2                  | 107.8-109.2  | -1.90**| 2.66       |
| 2ndLA | qLA4.2                  | 107.8-109.2  | qLA5-3                  | 88.2-97.2    | -2.82**| 2.47       |
| 2ndLA | qLA4.2                  | 107.8-109.2  | qLA7-1                  | 15.2-19.3    | -1.64**| 0.6        |
| 2ndLA | qLA5.3                  | 97.9-103.0   | qLA7-1                  | 15.2-19.3    | 2.38**| 2.42       |
| 2ndLA | chr9-90756–mm0051       | 47.3-48.1    | chr10-77445–umc1336     | 20.6-22.4    | -2.05**| 2.66       |
| 3rdLA | qLA3.1                  | 55.2-59.5    | qLA4-2                  | 107.8-109.2  | -1.55**| 2.06       |
| 3rdLA | qLA4.2                  | 107.8-109.2  | qLA5-3                  | 88.2-97.2    | -1.74**| 1.38       |
| 3rdLA | qLA5.3                  | 97.9-103.0   | qLA7-1                  | 15.2-19.3    | 1.83**| 2.37       |
| 3rdLA | chr9-23536–chr9-32338   | 44.7-46.6    | umc1380–chr10-12923     | 12.0-13.3    | -2.09**| 3.04       |
| 4thLA | qLA3.2                  | 67.9-76.6    | qLA5-3                  | 81.6-88.2    | 1.49**| 0.69       |
| 4thLA | qLA3.2                  | 67.9-76.6    | qLA7-1                  | 15.2-19.3    | 1.99**| 1.61       |
| 4thLA | qLA5.3                  | 97.9-103.0   | qLA7-1                  | 15.2-19.3    | 2.30**| 2.57       |
| 5thLA | chr4-150464–bnlg1137    | 49.8-65.8    | qLA4-2                  | 98.4-105.3   | 1.07* | 0.39       |
| 5thLA | chr8-103366–chr8-111393 | 39.4-42.6    | chr8-165985–chr8-167777 | 84.7-88.3   | -2.26**| 3.54       |
| 6thLA | qLA4.2                  | 107.8-109.2  | qLA5-3                  | 88.2-97.2    | -1.06**| 1.2        |
| 6thLA | chr8-103366–chr8-111393 | 39.4-43.6    | qLA8-1                  | 72.3-80.0    | -2.28* | 3.52       |
| 7thLA | qLA2.1                  | 4.6-14.6     | qLA5-1                  | 46.3-49.2    | 1.13* | 1.08       |
| 7thLA | phi213984–chr4-10484    | 14.6-18.6    | bnlg1759–umc1350        | 93.2-94.2    | -1.99**| 3.25       |

AA is the additive-by-additive epistatic interaction effect
$H^2_{(aa)(\%)}$ are percentage of variance explained by the additive-by-additive epistatic interaction effect.

*, ** and *** indicate significant level at $P < 0.05$, $P < 0.01$ and $P < 0.001$, respectively.
Fig. 1. The network diagram of identified QTL controlled the eight LAs. The 1.1 of Chromosome 1 (Chr1) represents $qLA1.1$, and so forth. The capital letters S and J represent the QTL detected only in single-environment and joint analysis, respectively. The arrowhead lines mean that the QTL control the corresponding LA. Different colored lines represent different QTL. The thick lines represent the PVE (%) of QTL $> 10\%$, fine lines represent the PVE (%) of QTL $\leq 10\%$. When one QTL was identified in more than one environment or both single-environment and joint analysis, we choose the highest value of PVE.