Diversity arrays technology (DArT) for studying the genetic polymorphism of flue-cured tobacco (*Nicotiana tabacum*)

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Abstract: Diversity arrays technology (DArT) is a microarray-based marker system that achieves high throughput by reducing the complexity of the genome. A DArT chip has recently been developed for tobacco. In this study, we genotyped 267 flue-cured cultivars/landraces, including 121 Chinese accessions over five decades from widespread geographic regions in China, 103 from the Americas and 43 other foreign cultivars, using the newly developed chip. Three hundred thirty three polymorphic DArT makers were selected and used for a phylogenetic analysis, which suggested that the 267 accessions could be classified into two subgroups, which could each be further divided into 2-4 sections. Eight elite cultivars, that account for 83% of the area of Chinese tobacco production, are all found in one subgroup. Two high-quality cultivars, HHDJY and Cuibi1, were grouped together in one section, while six other high-yield cultivars were grouped into another section. The 330 DArT marker clones were sequenced and close to 95% of them are within non-repetitive regions. Finally, the implications of this study for Chinese flue-cured tobacco breeding and production programs are discussed.

Key words: *Nicotiana Tabacum*, DArT, Genetic Diversity, Flue-cured Tobacco

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1 Introduction

Genetic resources play a key role in crop breeding and a wide genetic variability within the germplasm pools of crops is critical for good breeding practices. To estimate the genetic diversity of a crop and to understand the population structure or genetic background of crops, several investigations have been carried out in many crops at the country level and worldwide. Traditional investigations of genetic diversity have considered phenotypic traits, isoenzymes, etc., which usually present with relatively low levels of polymorphism, and the emergence of DNA markers in the 1980s (Botstein et al. 1980) made these investigations more efficient. Early molecular marker-based studies in crops mainly focused on the use of AFLP (amplified fragment length polymorphism) or RAPD (randomly amplified polymorphic DNA). Along with large-scale sequencing efforts in crops, sequence-based markers, such as simple sequence repeat (SSR) and single nucleotide polymorphism (SNP), have been developed and widely used in studies of crop diversity (Agarwal et al. 2008).

Tobacco (*Nicotiana tabacum*) is one of the most economically important nonfood crops. Tobacco breeding programs face challenges with regard to quality characteristics, such as low toxin content, disease resistance, drought tolerance etc., and thus, there have recently been several molecular marker-based studies on genetic diversity. Initially, as
with other crops, AFLP (Ren and Timko 2001) and then SSR (Yang et al., 2007; Moon et al., 2009a; 2009b; Fricano et al., 2012) were used. Molecular diversity in a worldwide collection of tobacco germplasm revealed the structured populations, basically by different tobacco types (Fricano et al., 2012). Flue-cured tobacco accounts for approximately 75% of world tobacco production (Moon et al., 2009a) and over 90% of that in China (Liu, 2011). However, flue-cured tobacco contains only a small portion of \textit{N. tabacum} germplasm (Moon et al., 2009a). However, its genetic diversity has been steadily decreasing over the past few decades due to strict breeding selection. (Moon et al., 2009b) investigated a population of 117 U.S. flue-cured cultivars with 71 microsatellite primer pairs and demonstrated that over 50% of the allelic diversity was lost from varieties released between the 1930s and 2005. Although there have been several investigations based on limited SSR or AFLP markers and cultivars (Zhang et al., 2006; Yang et al., 2007), little is currently known about the genetic variation within Chinese flue-cured tobacco pools and the impact of breeding practices on genetic diversity within Chinese elite cultivars.

Diversity arrays technology (DArT) is a microarray-based marker system that achieves high throughput by reducing the complexity of a DNA sample to obtain a ‘representation’ of that sample (Jaccoud et al., 2001). It has been developed and used in many plants, including wheat and barley, which have complex genomes, while their genome sequences are not yet available for genetic mapping or studies of genetic diversity etc. (Wenzl et al. 2004; Akbari et al. 2006; Tinker et al., 2009; Alsop et al., 2011). Very recently, we developed a tobacco DArT chip that includes 7680 representative sequence tags and which has been successfully used to construct tobacco genetic maps (Lu et al., 2012).

In this study, we profiled a germplasm set of 267 flue-cured tobacco cultivars including 121 elite materials over five decades of cultivar development and landraces, or farmer varieties, from widespread geographic regions in China and 103 typical flue-cured tobacco accessions from the Americas and 43 from other countries were also included for the phylogenetic analysis. DNA was extracted from fresh leaf tissue (~200mg) of these 267 tobacco accessions using a cetyltrimethylammonium bromide (CTAB) protocol (Sambrook and Russell, 2001).

2. Materials and methods

2.1 Plant materials and DNA extraction

A total of 121 Chinese flue-cured tobacco accessions, including elite materials over five decades of cultivar development and landraces or farmer varieties, from widespread geographic regions were collected and used in this study (Table 1; Supplementary Table 1). To elucidate the evolutionary origin of the Chinese accessions, 103 typical flue-cured tobacco accessions from the Americas and 43 from other countries were also included for the phylogenetic analysis. DNA was extracted from fresh leaf tissue (~200mg) of these 267 tobacco accessions using a cetyltrimethylammonium bromide (CTAB) protocol (Sambrook and Russell, 2001).

2.2 DArT marker detection

A DArT marker chip for tobacco (Lu et al., 2012) was used in this study. Briefly, a genome representation of a mixture of 5 cultivars (HHDJY, Hicks Broad Leaf, Florida 301, Burley 21 and Turkey Basma) was produced after PstI/TaqI digestion and spotted on microarray slides, and the individual genotypes were screened for polymorphism based on fluorescence signals. The 267 tobacco accessions were screened for fingerprinting.

<table>
<thead>
<tr>
<th>Geographic region</th>
<th>Country</th>
<th>Number</th>
</tr>
</thead>
<tbody>
<tr>
<td>China</td>
<td></td>
<td>121</td>
</tr>
<tr>
<td>Asia</td>
<td>Japan, Taiwan, Vietnam, N. Korea, S. Korea and The Philippines</td>
<td>18</td>
</tr>
<tr>
<td>North America</td>
<td>USA</td>
<td>82</td>
</tr>
<tr>
<td>South America</td>
<td>Canada</td>
<td>8</td>
</tr>
<tr>
<td>America</td>
<td>Brazil, Cuba and Argentina</td>
<td>13</td>
</tr>
<tr>
<td>Africa</td>
<td>Zimbabwe, Somalia, Tanzania and Zambia</td>
<td>18</td>
</tr>
<tr>
<td>Europe</td>
<td>Poland, Yugoslavia and England</td>
<td>4</td>
</tr>
<tr>
<td>Oceania</td>
<td>Australia</td>
<td>3</td>
</tr>
<tr>
<td>Total</td>
<td></td>
<td>267</td>
</tr>
</tbody>
</table>
The *Escherichia coli* clones containing the polymorphic DArT markers identified using the discovery arrays were re-arrayed into 384-deep-well microliter plates and grown at 37°C for 20 hours. Plasmid DNA, isolated using the Eppendorf PerfectPrep Plasmid 384 procedure, was sequenced in both directions using the M13R (5′-GGAAACAGCTATGACCATG-3′) and T7-ZL (5′-TAATACGACTCACTATAGGG-3′) primers. Following an ethanol precipitation cleanup step, the reactions were run on an ABI 3730xl capillary electrophoresis instrument. All sequence reads were assembled and merged to provide one high-quality read per clone where possible. Vector sequences and PstI sites were trimmed so as to not introduce biased similarity among DArT clones in current or future analyses.

### 2.3 Data analysis

The genetic relationship of the 267 tobacco varieties was investigated using the program NTSYSpc 2.11e (www.exetersoftware.com) with the following settings: Qualitative data, Dice coefficient, Sequential Agglomerative Hierarchical and Nested clustering method (SAHN), Unweighted Pair-Group Method, Arithmetic average (UPGMA).

The population structure among all 267 tobacco accessions was evaluated with STRUCTURE 2.3.3 (Hubisz *et al*. 2009) using an admixture model with no linkage. Three hundred thirty three DArT markers were used and considered haploid for this analysis. All analyses had a burn-in length of 50,000 iterations and a run length of 100,000 iterations. Three replicates were carried out at each value of K (population number, from 2 to 10). Simulations were run with uncorrelated allele frequencies.

DArT clones were collected and sequenced by the Sanger method in one direction. Repeat sequences were identified and masked by searching against Repbase16.06 (http://www.girinst.org) with RepeatMasker (http://www.repeatmasker.org/). The non-repetitive DArT sequences were further compared with the tobacco unigene set TobEA (Edwards *et al*. 2010) and GenBank nr database by BLASTX, and the sequences with significant hits (e-value < 1e-7) were annotated. GO classification was carried out for those hits (e-value < e-7) using Blast2GO (Conesa *et al*. 2005) and the GO histogram was generated by WEGO (Ye *et al*. 2006).

Two summary statistics parameters were used for diversity measure with a Perl script. The two parameters estimate the population mutation rate per locus based on the number of segregating sites (Watterson 1975) and the mean value of pairwise divergence per locus (π) (Tajima 1983), respectively.

### 3 Results

#### 3.1 Characteristics of DArT markers

A total of 330 high-quality DArT markers were identified from discovery arrays and used in the following diversity analysis. The polymorphism information content (PIC) values of the 330 identified polymorphic markers ranged from 0.07-0.50, with an average value of 0.34 (78.8% > 0.2) (Table 2).

<table>
<thead>
<tr>
<th>PIC value</th>
<th>DArTs</th>
<th>% DArTs</th>
</tr>
</thead>
<tbody>
<tr>
<td>0.5-0.4</td>
<td>152</td>
<td>55.2</td>
</tr>
<tr>
<td>0.4-0.3</td>
<td>48</td>
<td>14.5</td>
</tr>
<tr>
<td>0.3-0.2</td>
<td>30</td>
<td>9.1</td>
</tr>
<tr>
<td>0.2-0.1</td>
<td>27</td>
<td>8.2</td>
</tr>
<tr>
<td>&lt;0.1</td>
<td>43</td>
<td>13.0</td>
</tr>
</tbody>
</table>

To further characterize the 330 DArT markers, their clones were collected and sequenced by the Sanger method in one direction. Repeat sequences were identified and masked by searching against Repbase16.06 (http://www.girinst.org) with RepeatMasker (http://www.repeatmasker.org/). The non-repetitive DArT sequences were further compared with the tobacco unigene set TobEA (Edwards *et al*. 2010) and GenBank nr database by BLASTX, and the sequences with significant hits (e-value < 1e-7) were as genic regions. GO classification was carried out for those hits (e-value < e-7) using Blast2GO (Conesa *et al*. 2005) and the GO histogram was generated by WEGO (Ye *et al*. 2006).
Based on the genotyping results of 330 DArT markers, the phylogenetic tree and population structure of 267 flue-cured tobacco accessions from China and other countries (Table 1) were investigated, respectively. In general, both clustering results (Fig. 2 and Fig. 3; detailed information were provided as supplementary Table 1) fit the pedigree of the cultivars used in this study. For example, 581 were bred by selection from Dajingyuan and they initially clustered together. Yunyan85 and Yunyan87 have the same parental line (K326) and initially they clustered together. The 267 accessions appeared to be comprised of two subgroups according to the phylogenetic analysis (Fig. 2). Each subgroup can be further classified into 2-4 sections according to a phylogenetic tree (Table 3 and Fig. 2) or population structure analysis (Fig. 3). In subgroup 1, two sections (section A and B or #1 and #2) could be observed by both analyses. In contrast, subgroup 2 is relatively complex, and at least 2-4 sections can be ascertained (Supplementary Table 1). There are two main sections (C and D) in the phylogenetic tree or three sections (#3-#5) by population structure analysis in this subgroup and section C includes particularly diverse accessions (144 of 267 accessions). In general, section A, B and D by phylogenetic analysis included most accessions of section #2, #1 and #4 of population structure analysis respectively, while section C covered #3 and #5 (detailed corresponding section pairs are provided as supplementary Table 1).

Interestingly, the main cultivars in China are all contained in subgroup 1. For example, eight cultivars (K326, Yunyan85, Yunyan87, Yunyan97, Zhongyan100, Longjiang911, HHDJY and Cuibi1) from this subgroup accounted for 83% of the total area of tobacco production in 2010 (Liu, 2011). Further, among the eight elite cultivars, two with high quality (HHDJY and Cuibi1) were grouped into a section (section A by phylogenetics) that included the U.S. cultivars K346, NC89, etc., while the other six with high yield were grouped into another independent section (section B by phylogenetics) that included the U.S. cultivars K326, NC71, etc. (Table 3). To compare genetic diversity with the Chinese tobacco cultivars (released from 1950 to now) and landraces, the eight cultivar set has the smallest values of the statistic parameter (π) and number of segregating sites in the 330 DArT marker sites (Table 4; Supplementary Table 2), suggesting that they only included part of the genetic diversity of Chinese tobacco accessions. These results indicate that current Chinese tobacco breeding programs and tobacco production reflect a narrow genetic diversity or background.

Our cluster analysis appears to show that (1) the 267 cultivars were not first clustered by their geographical origins, i.e., cultivars from the same region or country were not usually grouped together. The results suggested that germplasm introduction and cross-breeding programs have given flue-cured tobacco cultivars a mixed genetic background in different countries. On the other hand, the elite cultivars that are bred and used in different countries show a similar genetic background; (2) farmer varieties show a wide genetic diversity (e.g. 315 segregating sites; Table 4) and cover each section, including even nodes that are further downstream in the phylogenetic tree. These varieties, together with foreign cultivars/germplasm (detailed information for diversity measures were provided as supplementary Table 2), provide a valuable genetic pool to meet the challenge that faces Chinese flue-cured tobacco breeding programs. (3) Elite cultivars that were widely planted at different times in the U.S. and China were usually grouped together in the same section (Table 3). In general, the trend of reduction in the number of segregating sites and π value over time was observed.
Fig. 2 DArT-based UPGMA dendrogram of 267 flue-cured tobacco cultivars. Four sections and phylogenetic codes (1-267 from top) of 267 accessions are shown. Detailed information of the four sections and their accessions were provided in Table 3 and supplementary Table 1.
in the Chinese tobacco breeding program (Table 4). The results suggests that over the past five decades, the Chinese tobacco breeding program has followed that of the U.S. (Moon et al., 2009b). However, two new cultivars that were just released by YATAS (Yunyan98 and Yunyan100) are located in an independent section (section D in the phylogenetic tree) of subgroup 2 and present a unique genetic background to other elites

### Table 4 Genetic diversity of Chinese flue-cured tobacco accessions in various breeding periods

<table>
<thead>
<tr>
<th>Group</th>
<th>Number of accessions</th>
<th>Number of segregating sites</th>
<th>Diversity value (π*)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Cultivars (1950-now)</td>
<td>72</td>
<td>302</td>
<td>0.412</td>
</tr>
<tr>
<td>Eight elites set#(now)</td>
<td>8</td>
<td>140</td>
<td>0.233</td>
</tr>
<tr>
<td>Landraces</td>
<td>49</td>
<td>315</td>
<td>0.385</td>
</tr>
<tr>
<td>1950-70s</td>
<td>36</td>
<td>294</td>
<td>0.395</td>
</tr>
<tr>
<td>1980-90s</td>
<td>24</td>
<td>282</td>
<td>0.412</td>
</tr>
<tr>
<td>2000s</td>
<td>10</td>
<td>237</td>
<td>0.325</td>
</tr>
</tbody>
</table>

*π*, the average pairwise nucleotide diversity (Tajima, 1983).

4 Discussion

In this study, we investigated the genetic diversity of 267 flue-cured accessions, including 121 Chinese cultivars/landraces, 103 cultivars from the Americas and 43 other foreign cultivars as controls,
by using 330 DArT markers. To our knowledge, this is the first attempt to investigate the genetic diversity of Chinese tobacco with the largest cultivar sample size and number of molecular markers. Our results are generally consistent with those of other studies of diversity based on SSR markers in U.S. and Chinese accessions. Moon et al. (2009a; 2009b) reported similar cluster results and limited genetic diversity for the U.S. flue-cured tobacco cultivar used in this study. Their investigation (Moon et al. 2009b) suggested that the allelic diversity in U.S. flue-cured tobacco cultivars has been declining since the 1930s. A narrow genetic diversity has also been reported in Chinese flue-cured tobacco based on AFLP and SST (Zhang et al., 2006; Yang et al., 2007).

As mentioned above, a narrow genetic diversity, which usually makes the development of molecular markers more difficult, was observed in tobacco. In addition to AFLP, RAPD and SSR markers, DArT provides an alternative for tobacco, as shown here. The DArT technology combines a reduction of genome complexity with high throughput and cost-effective hybridization-based polymorphism detection. DArT is not sequence-based, and is thus particularly helpful for species such as tobacco for which the genome sequence is not available or sequence data are limited. Many DArT markers are within genes or gene-rich regions, as shown in this study and others (e.g., Tinker et al., 2009), which is also particularly important for QTL mapping and molecular-assisted breeding of species with complex genomes (big, repetitive and polyploid), such as wheat, barley, oat and tobacco.

Our results offer several implications for Chinese flue-cured tobacco breeding programs and production. First, in Chinese tobacco production, cultivars with different genetic backgrounds could be planted in reasonable proportions or distributions. For example, an increase in the distribution of two new cultivars (Yunyan98 and Yunyan100) should improve the current situation in which only a limited number of elite cultivars with an extremely narrow genetic background are being planted. As mentioned above, the two cultivars are located in an independent section of subgroup 2 and present a unique genetic background to other elites. If they eventually become elite cultivars like K326 and HHDJY in China, they will be the first cultivars with a novel genetic background or Chinese genetic characteristics. Meanwhile, even for cultivars from subgroup 1, an increase in the distribution of cultivars with high quality (HHDJY and Cuibi1) should also be helpful for easing the current situation in China. Second, farmer varieties should be given more attention in Chinese tobacco breeding programs. They provide the same wide genetic novelty as foreign germplasm. In subgroup 2, several landraces are particularly diverse with respect to others (Fig. 2; Supplementary Table 1), and therefore provide tremendous promise for genetic improvement in current tobacco breeding programs in China.

References


