



Short Communication

Discovery of a high-altitude ecotype and ancient lineage of *Arabidopsis thaliana* from Tibet

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Arabidopsis thaliana (*A. thaliana*) has long been a model species for dicotyledon study, and was the first flowering plant to get its genome completed sequenced [1]. Although most wild *A. thaliana* are collected in Europe, several studies have found a rapid *A. thaliana* west-east expansion from Central Asia [2]. The Qinghai-Tibet Plateau (QTP) is close to Central Asia and known for its high altitude, unique environments and biodiversity [3]. However, no wild-type *A. thaliana* had been either discovered or sequenced from QTP. Studies on the *A. thaliana* populations collected under 2000 m asl have shown that the adaptive variations associated with climate and altitudinal gradients [4]. Hence a high-altitude *A. thaliana*

provides a precious natural material to investigate the evolution and adaptation process.

Here, we present the genome of a new ecotype of *A. thaliana* collected in the Gongga County, Tibet (4200 m asl) (Fig. 1a), to demonstrate its evolutionary history and adaptation to high-altitude regions. The Tibetan samples were identified as *A. thaliana* by comparing the nuclear internal transcribed spacer (ITS), four chloroplast genes (*matK*, *rbcl*, *rpoB*, and *rps16*), and three chloroplast intergenic spacers (IGS, *trnL-trnF*, and *trnT-trnL*) with *A. thaliana* (Col-0) and *A. lyrata* (Supplementary Fig. 1). This is the first report that an *A. thaliana* population has been collected in the QTP over 4000 m asl and identified by molecular analysis. Moreover, the new Tibetan ecotype (herein referred to as “Tibet-0”) is diploid ($2n = 10$) according to karyotype analysis of its pollen mother cells during meiosis (Supplementary Fig. 2 online), suggesting that the ploidy of the Tibet-0 is stable and capable of further sequence analysis.

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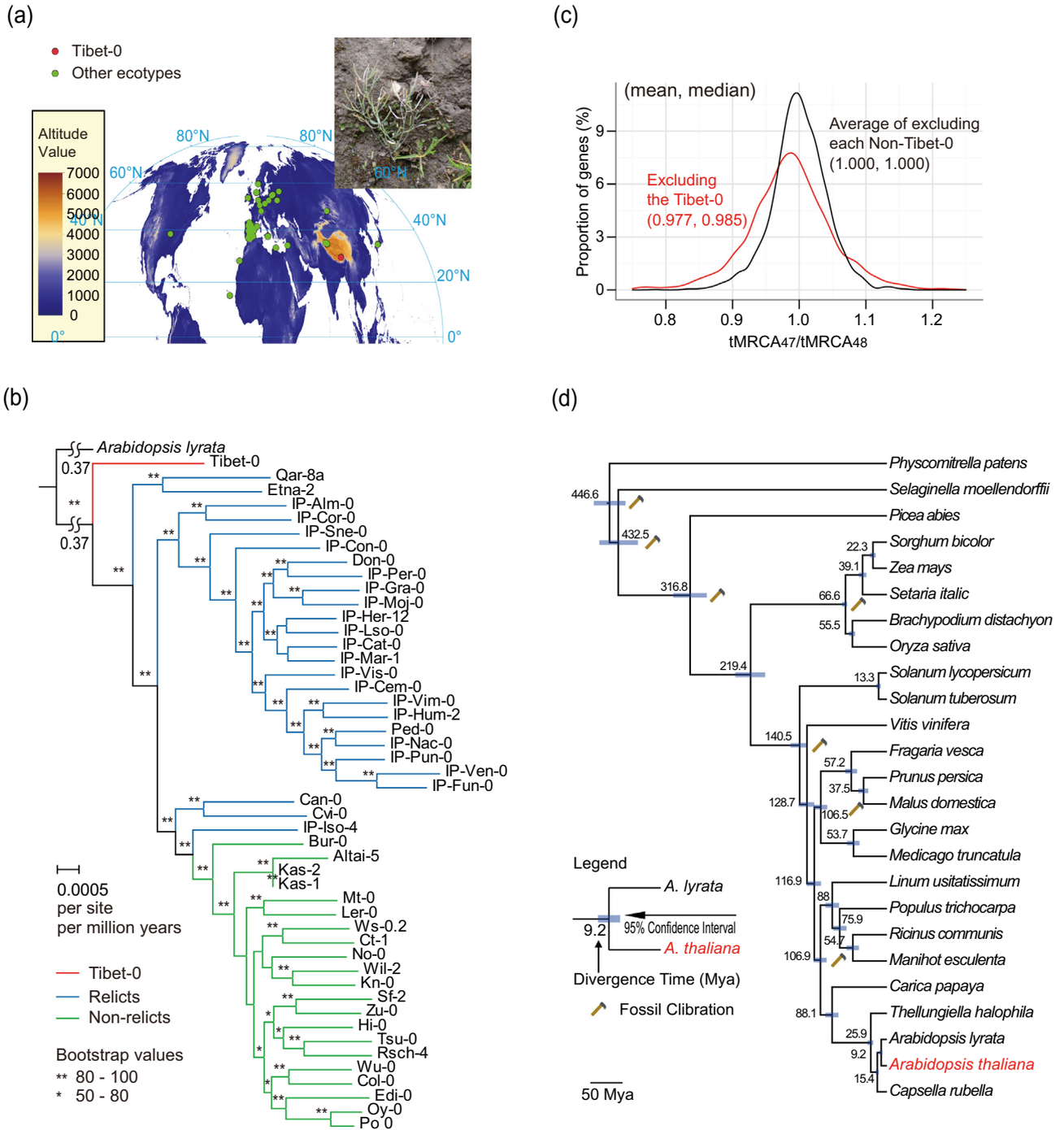


Fig. 1. Collection, phylogenetic analysis and adaptation analysis of the Tibet-0 and other 47 *A. thaliana* ecotypes. **a** Origins of Tibet-0 and other 47 *A. thaliana* ecotypes that we have analyzed in this paper (Supplementary Table 4). Elevation data were downloaded from WorldClim (<http://www.worldclim.org/>). Colors indicate altitudes, going from low to high elevation: Deep blue, blue, white, yellow, orange, and brown. **b** The maximum likelihood phylogenetic trees based on 5611 orthologues of Tibet-0, 47 *A. thaliana* ecotypes and *A. lyrata* used as outgroup. Bootstrap values based on 100 replications are listed as percentages at each node. The Tibet-0 was marked in red. **c** The time of most recent common ancestor (tMRCA) based on 2788 single-copy orthologues. The red line represents the tMRCA47/tMRCA48 distribution where the tMRCA47 values exclude Tibet-0. The black line represents the mean of tMRCA47/tMRCA48 distribution where the tMRCA47 values exclude each Non-Tibet-0. **d** Phylogenetic affinities inferred from the maximum likelihood analysis of nucleotide sequence of 334 single copy orthologues in 25 plants. The divergence time of *A. thaliana* and *A. lyrata* was about 9.2Mya (million years ago). 7 fossil calibrations used in the study were marked as the hammer symbol. Branch lengths are proportional to the number of expected nucleotide substitutions. The number on the branch is the divergence time and unit is Mya.

We then conducted genome-wide resequencing of Tibet-0 with a mean coverage of 40x of the reference genomes Col-0 and TAIR10, by using Illumina Hiseq2000 (Supplementary Tables 5, 6, online). We compared Tibet-0 with 47 other *A. thaliana* ecotypes

that have been genome-wide sequenced, and found that Tibet-0 was of high divergence, including a higher proportion of SNPs (Supplementary Tables 7–9, online). Evolutionary relationships between Tibet-0 and other ecotypes were evaluated by the

following two independent approaches based on 5611 single-copy orthologues in 47 *A. thaliana* ecotypes including 26 relicts and 21 non-relicts defined by the 1001 Genomes Consortium [5]. The first approach is the phylogenetic method. The genealogy among the individuals was inferred based on the concatenated genomic data, and Tibet-0 was placed at the root of the *A. thaliana* populations with high support value (Fig. 1b) [5,6]. It makes Tibet-0 the most ancestral lineage.

However, since this phylogenetic approach assumes that all gene loci have the same genealogy, coalescent method was also applied as a cross check [7]. In this method, 2788 single-copy orthologues were independently analyzed, and the distributions of the tMRCA (the time to the most recent common ancestor) for these genes were estimated. If Tibet-0 is the most basal lineage among the *A. thaliana* populations and Tibet-0 specific alleles has generally older histories than others, tMRCA excluding Tibet-0 will be smaller than tMRCA of all *A. thaliana* populations. Otherwise, if there is no such genetic structure and Tibet-0 specific alleles are included within the genetic diversity of other *A. thaliana* populations, tMRCA excluding Tibet-0 will be equal to the tMRCA of all *A. thaliana* populations. To examine the differences among the distributions, the tMRCA were first estimated based on 48 *A. thaliana* (tMRCA48). Subsequently, each ecotype was excluded once, and the tMRCA of 47 remaining *A. thaliana* were estimated (tMRCA47: there are 48 combinations of tMRCA47). Finally, the relative tMRCA (tMRCA47/tMRCA48) were estimated. Fig. 1C illustrates the distributions of the relative tMRCA. When Tibet-0 was excluded, the distribution of the relative tMRCA (tMRCA47_{excluding Tibet-0}/tMRCA48) significantly shifted (*t* test, $P = 9.77E-32$), while the average of tMRCA47_{excluding one ecotype other than Tibet-0}/tMRCA48 showed no significant change (Fig. 1c). These findings confirm that Tibet-0 has the most ancestral positions among *A. thaliana* populations.

To understand the correlation between the evolution of *A. thaliana* and major geological events, especially Tibetan uplifts, the divergence time between Tibet-0 and other ecotypes were estimated. Since there is no suitable fossil calibrations within *A. thaliana*, the divergence time between *A. lyrata* and *A. thaliana* was estimated based on the genomic data in the framework of whole land plant evolution with reliable fossil records, and it was estimated to be about 9 million years ago (Fig. 1d, Supplementary Fig. 3). Then, the time of the common ancestor of *A. thaliana* was estimated by multiplying the divergence time between *A. lyrata* and *A. thaliana* and the ratio of the divergence time between *A. lyrata* and *A. thaliana*. The divergence time between Tibet-0 and other ecotypes was found to be 126–149 Ka (kili annum: thousand years ago). Interestingly, the Gonghe movement, which was the last phase of Tibetan uplift, isolated the Qinghai Lake and raised the QTP to its present height began at about 150 Ka [8]. Besides, the divergence time of Tibet-0 and other ecotypes is in the middle Pleistocene late Middle Pleistocene from 781 to 126 Ka [9].

A. thaliana has been widely used in studies of plant biology. By collecting and sequencing *A. thaliana* collected from the QTP over 4200 m asl, we have found that the Tibet-0 is a new and divergent ecotype that isolated from other *A. thaliana* ecotypes since the last uplift of the QTP. After 126–149 thousands years evolution in the extreme plateau environment, Tibet-0 possesses a distinctive genome with a high proportion of SNPs compared to other ecotypes. According to the strongly negatively skewed Tajima's *D* of 5611 single-copy orthologues, a recent selective sweep or population expansion might have occurred in the *A. thaliana*, which is consistent with previous studies (Supplementary Fig. 4, online) [10].

Considering the ancestral position of Tibetan populations as well as the subsequent selective sweep or population expansion, possibly in the Last Glacial Period, suggested by the negative Tajima's *D*, we suppose that some mutations might have emerged in the ancient *A. thaliana* population located around the QTP, and then spread to most other populations. Following step is investigating phenotypic traits of Tibet-0 to study the adaptive evolution of *A. thaliana* to high altitudes. As a new model plant, the Tibet-0 from QTP would provide an invaluable material for further study.

Author contributions

F.C. and Y. Zhong conceived the project. L. Zeng, Z.G., T.Y., F.C. and Y. Zhong contributed to the design of the project and extensive discussions. M.X., N.Z., W.Z., L.Q. and T.T. collected samples from Tibet. L. Zeng and H. G. helped with sample identification. L.X., R. X., F.X., J.L., L.Z., Z.G., N.Z., Y.H., T.Y., M.H., F.Z., F.C., Y.G., L. Zhang, Y. Zhang, Z.Y., M.J.C.C. and Y. Zhong performed the common garden experiments, and sequence analyses and evolutionary analyses. L. Zeng, Z.G., Y. Zhang, M.J.C.C., N.S., F.C. and Y. Zhong wrote the manuscript. Other authors revised the manuscript.

Conflict of interest

The authors declare that they have no conflict of interest.

Availability of data and materials

The genomic DNA of Tibet-0 has been deposited in the Sequence Read Archive (SRA, <http://www.ncbi.nlm.nih.gov/sra/>) under accession number SRP052218.

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Appendix A. Supplementary data

Supplementary data associated with this article can be found, in the online version, at <https://doi.org/10.1016/j.scib.2017.10.007>.

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