High-quality ice plant reference genome analysis provides insights into genome evolution and allows exploration of genes involved in the transition from C3 to CAM pathways

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Received 24 February 2022; revised 19 May 2022; accepted 10 July 2022.
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Keywords: ice plant, genome assembly, functional annotation, genome evolution, crassulacean acid metabolism, expression pattern.

Introduction

Mesembryanthemum crystallinum (ice plant, 2n = 2x = 18) belongs to the Aizoaceae family, which contains approximately 160 genera and 2500 species. Ice plants are native to southern and eastern Africa and are now grown worldwide. Ice plants are vegetables with high nutritional value and health functions. They are rich in pinitol, which lowers blood sugar and has potential antidiabetic effects (Diria et al., 2016; Zhang et al., 2019a). They also contain myo-inositol, which prevents fatty liver disease (Diria et al., 2016). Ice plants are inherently more resistant to diseases, insect pests and toxicity compared with most other vegetables. Therefore, these plants are capable of healthy growth without the need of pesticides but only a small amount of fertilizer (Amari et al., 2020; Sliwa-Cebula et al., 2020). Ice plants can be eaten fresh as a high-quality green vegetable.

Soil salinization is a global ecological problem that seriously affects the growth and production of most crops (Palansooriya et al., 2019). Clarifying the salt tolerance mechanisms of halophytes is important for preventing soil salinization and increasing crop yields (Mishra and Tanna, 2017). Ice plants are halophytes that convert from C3 photosynthesis to crassulacean acid metabolism (CAM) under high-salinity stress (Adams et al., 1998). CAM is a special form of photosynthetic carbon assimilation that occurs in an estimated 7% of vascular plants (Crayn et al., 2004; Cushman et al., 2008; Silvera et al., 2005). CAM is characterized by the absorption and assimilation of atmospheric carbon dioxide (CO2) catalysed by phosphoenolpyruvate carboxylase (PEPC) at night, leading to the accumulation of C4 acids (Gilman and Edwards, 2020). By absorbing large amounts of CO2 at night, when the evapotranspiration rate is low, the water use efficiency of CAM plants is three to six times higher than that of C4 and C3 plants (Cushman et al., 2008). The genes related to salt tolerance identified in ice plants are strong candidates for genetic engineering of C3 crops to improve salt tolerance (Tsukagoshi et al., 2015).

Several studies have revealed omics level changes in ice plants in response to salt stress. The first microarray experiment used...
5-week-old and 14-day salt-stressed plants (Cushman et al., 2008). Using second-generation sequencing, 53 516 cDNAs were detected in the roots of ice plants (Tsukagoshi et al., 2015). A reference transcriptome was constructed that contained 37 341 transcripts from control and salt-treated ice plant epidermal bladder cells (EBCs) (Oh et al., 2015). One hundred thirty-five conserved microRNAs (miRNAs) have been identified in the roots of 3-day-old ice plant seedlings (Chiu-Pin et al., 2016). The proteomes and metabolomes of ice plant EBCs have also been studied (Barkia et al., 2016; Barkla and Vera-Estrella, 2015; Li et al., 2021). Recently, the critical transition time for ice plants from C3 to CAM has been determined by measuring several key attributes, including gas exchange, stomatal aperture, titratable acidity, CAM enzyme activity and CAM gene expression (Kong et al., 2020).

Related transcriptome studies have analysed the changes in gene expression of ice plants after salt stress treatment (Kong et al., 2020; Oh et al., 2015; Tsukagoshi et al., 2015). However, the genome of ice plants has not yet been resolved, which has hindered studies of molecular regulation mechanisms related to salt tolerance in ice plants. In the present study, PacBio HiFi, Illumina sequencing and Hi-C technologies were used to obtain high-quality ice plant genome sequences. Several key genes for salt tolerance in ice plants were identified by combined genomic and transcriptomic analyses. These data provide an important reference for research on salt-tolerant genomics and molecular biology of ice plants.

Results
Ice plant genome sequencing, assembly, and assessment

De novo ice plant genome sequencing was performed using PacBio HiFi, Illumina and Hi-C technologies (Figure 1a, Table 1). First, the ice plant genome was estimated by the K-mer method using 90.43 Gb of data from Illumina sequencing (Tables 1 and S1). The estimated size of the ice plant genome was 394.89 Mb, and the heterozygosity rate was 0.10% (Figure S1, Table S1). The PacBio HiFi sequencer was adopted to generate 18.77 Gb of data with an average of 47.53× coverage depth (Tables 1 and S3). In total, 109.20 Gb (276.53×) of ice plant DNA sequences generated from the Illumina and PacBio platforms were used to perform the preliminary assembly. The results indicated that the cumulative length of the contig was 377.96 Mb, and that of contig N50 was 6.18 Mb (Tables S4–S6).

Hi-C technology was used to improve the ice plant genome assembly. High-quality sequences (61.80 Gb, 156.50×) were obtained by Illumina sequencing (Table 1). A Hi-C contact map was used to divide the distinct regions of each chromosome (Figure 1b). Finally, the assembled genome size was approximately 377.97 Mb, with contig N50 and scaffold N50 reaching 6.18 Mb and 40.45 Mb, respectively (Table 2). In total, 370.61 Mb sequences were anchored to nine chromosomes in the ice plant, accounting for 98.05% of the assembled genome (Figure 1c, Table S7).

The reads mapping rate exceeded 99.11%, indicating the assembled ice plant genome was relatively complete (Table S8). The core eukaryotic gene mapping approach (CEGMA) and benchmarking universal single-copy orthologs (BUSCO) methods were used to assess the quality of assembled genomes and annotations. The CEGMA results showed that 96.77% (241) of core eukaryotic genes were detected in the assembled genome (Table S9). The BUSCO results showed that 98.0% of 1641 genes were found in the ice plant genome (Table S10).

Genome annotation

Repetitive sequences accounted for 48.04% of the estimated ice plant genome (Figure 1c, Table S11). Most repetitive sequences were long-terminal repeats (LTRs; 121.34 Mb), accounting for 32.11% of the genome (Figure S2, Table S11). Long interspersed nuclear elements (LINE) and DNA transposons only accounted for 6.40% and 7.42% of the ice plant genome, respectively. Most genes were located in the terminal regions of each chromosome, displaying a similar trend to that of DNA transposable elements. However, Copia and Gypsy retrotransposons were almost inversely distributed on each chromosome compared with the genes (Figure 1c).

Among the 24 234 annotated ice plant genes (Figures S3, S4, Tables S12, S13), InterPro, Swiss-Prot, Kyoto encyclopedia of genes and genomes (KEGG) and non-redundant protein databases showed functional evidence for 24 064 (99.30%) genes, with 15 512 genes annotated in all databases (Figure S5, Table S14). Concerning RNA, 4269 rRNAs, 2446 miRNAs, 1054 tRNAs and 889 snRNAs were found, in total accounting for 0.91% of the ice plant genome (Figure S6, Table S15).

Gene family expansion analysis and divergence time estimation

We detected gene families in ice plant, nine Caryophyllales species, two other eudicots (Arabidopsis thaliana and Vitis vinifera), and Oryza sativa (Figure 2a, Table S16). In total, 28 868 gene families were identified in the ice plant and the other 12 examined species (Figure 2b, Table S17). There were 14 330 gene families in the ice plant, which was lower than that in Atriplex hortensis (15950), Chenopodium quinoa (15886), Spinacia oleracea (15020), and Chenopodium pallidicaule (14729) (Figure 2b, Table S17). However, only 12 835 and 12 894 gene families were detected in A. thaliana and Beta vulgaris, respectively. Common and specific gene families analyses among the 13 species was performed using a Venn diagram (Figure 2b). A total of 252 single-copy and 6108 common gene families were found among the 13 species (Figure 2b, Table S17). The 468 ice plant species-specific gene families exceeded the numbers for C. pallidicaule (103), B. vulgaris (258) and Hylcoreus undatus (447) but was less than that of the other nine species (Figure 2b).

Gene family contraction and expansion were explored in ice plant and the 12 other representative species (Figure 2c). In the ice plant, we detected 106 gene family expansions, which was more than that of the closely related species, H. undatus (92) (Figure 2c). The largest number of gene family expansions was found in C. quinoa (888), followed by A. hortensis (453), and S. oleracea (306). Only 29 gene family contractions were found in ice plants, which was more than in Fagopyrum tataricum (6) and S. chinensis (14), but fewer than that seen in other Caryophyllales species (Figure 2c). We performed a phylogenetic analysis and divergence time estimation using single-copy gene families among the 13 species (Figure 2c). The ice plant had the closest relationship with H. undatus among all examined species. It diverged from H. undatus 30.6–47.4 million years ago (Mya) (Figure 2c). The divergence time between Caryophyllales and the other species was 84.0–99.9 Mya (Figure 2c).
Evolution and polyploidization of the ice plant genome

We explored the genomic evolution of ice plants using the rate of synonymous nucleotide substitution (Ks) within syntenic blocks among ice plants, grapes, and nine other Caryophyllales species (Figure 2d-e). In the Ks density plot, only one peak was detected in the ice plant genome. This indicates that only one polyploidization event occurred in the ice plant, which was an ancient whole-genome triplication (WGT) event shared with grapes and most eudicots (Jaillon et al., 2007). Therefore, there were no recent whole-genome duplication (WGD) or WGT events in ice plants.

Surprisingly, two peaks were detected in the Ks density plot of the S. chinensis genome, indicating that two polyploidization events had occurred in this species (Figure 2e). However, only one polyploidization event was found in a previous report, indicating that one recent WGT event in S. chinensis was overlooked. We also verified that S. chinensis experienced two rounds of WGT events by combining dot plot and syntenic analysis data (Figure 3a-d). Based on the ancient WGT event times that occurred in most eudicots, the recent WGT event in S. chinensis...
occurred between 50.60-57.42 Mya (Ks = 0.91). This was later than the time of the recent WGD event in *F. tataricum* (53.03–59.95 Mya; Ks = 0.95) but earlier than the times of the recent WGD events in *H. undatus* (46.33–52.38 Mya; Ks = 0.83) and *A. cruentus* (29.59–33.44 Mya; Ks = 0.53) (Figure 2e, Table S18).

### Syntenic and gene retention analysis of the ice plant genome

We identified 100 intra-genomic collinear blocks in an ice plant, containing 2079 collinear genes (Table S19). We then mapped the ice plant gene sequences onto nine other Caryophyllales genomes to infer their inter-genomic collinearity (Figures 3b, S7–S10). Among the Caryophyllales species, the most collinear blocks were between the ice plant and *C. quinoa* (305). This might reflect the genomic fusion that occurred in *C. quinoa* according to a previous report (Zou et al., 2017) (Figure S7). We also found that the number of collinear blocks between ice plants and species that underwent recent WGD or WGT events was greater than that of the other species (Figures 3b, S7–S10). For example, there were more collinear blocks between the ice plant and the four species that underwent recent WGD or WGT events, including *Amaranthus cruentus* (223 collinear blocks), *F. tataricum* (217), *H. undatus* (192) and *S. chinensis* (189) (Figures 3b, S8, and S9). However, there were fewer collinear blocks between

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**Figure 2** Gene family, phylogenetic and divergence time analyses. (a) The gene numbers of each category in the ice plant and 12 representative species. (b) Common and specific-specific gene families in ice plant and the other 12 species. (c) Expansion/contraction of gene family and divergence time estimation. The green and purple pies depict the ratios of the numbers of expansion and contraction gene families, respectively. The numbers on the nodes represent the species divergence time, with the confidence range list in brackets. The grey squares, hexagons and circles represent whole-genome duplication (WGD), whole-genome triplication (WGT) and genome fusion, respectively. (d) The density of synonymous nucleotide substitutions per synonymous site (Ks) among collinear genes between ice plant (Mcr) and other related species. (e) The density plot of Ks among collinear genes within ice plant (Mcr) and each other related species.
ice plant and the other four species that had not undergone recent WGD or WGT events, including *A. hortensis* (181), *S. oleracea* (166), *B. vulgaris* (142) and *C. pallidicaule* (131) (Figures S7, S9, S10). Furthermore, the largest number of large collinear blocks (gene pairs >200) were between the ice plant and *H. undatus* (44), followed by *C. quinoa* (40) and *S. oleracea* (32). Syntenic analysis indicated that most species underwent chromosome rearrangement after divergence (Figures 3b, S7–S10). To a certain extent, the syntenic results reflect the genetic relationship between ice plants and other Caryophyllales species.

The ratio of the collinear regions between the ice plant and grape was 1:1 because no recent genome duplication has occurred in the ice plant. However, the ratio between ice plants and the three species (*H. undatus*, *A. cruentus*, and *F. tataricum*) was 1:2 because of the recent WGD event detected in these three species (Figures 3b, 4a, Table S20). The ratio of ice plants to *S. chinensis* was 1:3 because of the recent WGT event detected in this study. For example, the end of chromosome 5 in the ice plant was collinear with *S. chinensis* chromosomes 8, 14 and 18, and it was also collinear with *H. undatus* chromosomes 3 and 11 (Figure 3d). Microsynteny analysis was also consistent with the global syntenic analysis, showing a similar ratio between ice plants and other species. For example, the 4.19–4.46 Mb of chromosome 4 in the ice plant was perfectly collinear with grape, *H. undatus* and *S. chinensis* (Figure 3d).

We conducted gene retention analysis of the ice plant genome in homologous regions by comparing it with other species. Different regions on the chromosome showed divergent retention levels (Figures 4a, b, S11–S19, Table S21). Grossly, the highest retention rate of collinear ice plant genes was 51.85%
using *H. undatus* as a reference, followed by *C. palilidae* (46.60%) and *C. quinoa* (43.66%) (Figure 4c, Table S21). However, the average retention rate of different chromosomes was only 9.66% when *F. tataricum* was the reference. We counted the number of syntenic gene pairs between the ice plant and the other nine Caryophyllales genomes. The most syntenic gene pairs were found between the ice plant and *A. hortensis* (5989 gene pairs), followed by *C. quinoa* (5735), *S. chinensis* (5656), *B. vulgaris* (5527), *S. oleracea* (5350), *H. undatus* (5177), *C. pallidicaule* (4975) and *A. cruentus* (4783). However, only 2426 syntenic gene pairs were identified between the ice plant and *F. tataricum*. The collective results revealed large-scale genome fractionation and instability of the ice plant genome after its split from these plants.

Exploring key genes in the CAM pathway

The CAM pathway is highly plastic under salt stress in ice plants, which makes it a good representative model for studying the transition mechanism from C3 to CAM (Cushman et al., 2008; Kong et al., 2020). Here, we attempted to identify the genes involved in regulating the C3 to CAM pathway by combining genomic and transcriptomic analyses (Table S22). Twenty genes encoding six main enzymes involved in the CAM pathway were identified in the ice plant (Figure 5a). Most nodes in the pathway had more gene copies in ice plants, including seven malate dehydrogenase (MDH), five malic enzyme (ME), three phosphoenolpyruvate carboxylase (PEPC) and three carbonic anhydrase (CA) genes.

To further explore the expression pattern of 20 CAM-related genes in the ice plant, we used three RNA-seq data sets from different tissues (E1) and applied salt treatment for different times (E2) or at different concentrations (E3) (Figure 5b, Tables S23–25). The E1 data set was generated in this study, and the E2 and E3 data sets were collected from previous studies (Kong et al., 2020; Tsukagoshi et al., 2015). Among the 20 differentially expressed genes (DEGs), ten, seven, and one were detected in E1, E2 and E3 experiments, respectively. Under salt treatment (E2 and E3), seven DEGs were identified, including four MDH and three PEPC genes. Furthermore, the expression level of the PEPC gene (*Mc08G01316*) was higher in the salt-treated plant tissue than in the control, regardless of day or night during days 5 to 7 (Figure 5c). The expression trend of the *Mc08G01316* gene was similar to that of the PEPC gene (*Contig20312*) identified in a previous study (Kong et al., 2020). The expression patterns of the

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**Figure 4** Global alignments and gene retention analyses of the ice plant genome. (a) Global alignment of homologous regions in ice plant and 10 other plant genomes with the grape as a reference. Each genome of *H. undatus*, *A. cruentus*, *F. tataricum* and *S. chinensis* were further divided into two or three sub-genomes due to additional WGD or WGT events that occurred in their genomes after divergence from the grape. Collinear genes between each sub-genome of species and grape were shown in each circle. The curved lines of the inner circle were composed of 19 grape chromosomes corresponding to the seven chromosomes of the ancient core-eudicot hexaploid ancestor (+ event). (b) The retention of duplicated genes residing in the ice plant genome along with each chromosome of the *H. undatus*. (c) The summary of gene retention in the ice plant genome compared with other species.
other two PEPC genes (Mc07G01375, Mc06G00895) were the opposite of Mc08G01316. However, these two genes were not detected in a previous study (Kong et al., 2020). Similarly, the expression levels of the two MDH genes (Mc03G00793, Mc07G01398) were lower with salt treatment than in the control, regardless of timing (i.e., day or night).

Figure 5 Overview of crassulacean acid metabolism (CAM) pathway in ice plant. (a) The CAM pathway map in the ice plant. Blue colours showed the key enzymes involved in the CAM pathway. The numbers in parenthesis are the number of genes encoding the corresponding enzymes in the ice plant. CA, carbonic anhydrase; PEPC, phosphoenolpyruvate carboxylase; PEPCK, PEPC kinase; MDH, malate dehydrogenase; ME, malic enzyme; PPDK, pyruvate phosphate kinase. (b) Expression profile heatmap of CAM-related genes in ice plant. The expression values (FPKM) were transformed by log2. The RNA-seq data set of the experiment (E1) was obtained in this study, and the data sets of E2 and E3 were obtained according to the previous reports. The red circle indicates the DEGs in corresponding experiments (fold-change > 2, q-value < 0.01). (c) Expression level of PEPC and MDH genes under salt treatment from E2 data set in the ice plant. (d) The interaction network between CAM-related genes and DEGs identified from E2 and E3 data sets in ice plant. (e) The edge number of each CAM-related gene in the network. (f) Functional enrichment analysis of DEGs involved in the network using KEGG (q-value < 0.05).

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Interaction network construction for key genes in the ice plant CAM pathway

Based on the identified CAM-related genes and their expression patterns, we constructed an interaction network for these genes and the genes they regulate in ice plants (Figure 5d). A total of 805 gene pairs formed the network according to the Pearson correlation coefficients (PCC > 0.95) (Table S26). Among these connections, only 10 represent regulatory interactions between CAM-related genes. The other 795 connections represent interactions between CAM-related genes and other DEGs in the ice plant genome (Table S26). This network contained nine CAM-related genes (five DEGs) and 285 other DEGs (Figure 5d, Table S27). Among these genes, the PEPC gene Mc07G01375 had the most connections (163) with other genes, followed by the MDH gene Mc04G00899 (149 connections) and the MDH gene Mc01G01317 (131 connections) (Figure 5d, e, Table S27). These results suggest that genes with more connections might play a core role in the CAM pathway in ice plants.

To explore the functions of the genes involved in the network constructed in the ice plant, we performed enrichment analysis. We identified 10 significantly enriched terms (q-value < 0.05). The most significantly enriched term was pyruvate metabolism (q-value = 2.08 × 10⁻¹¹), followed by carbon metabolism, carbon fixation in photosynthetic organisms and citrate cycle (TCA cycle) (Figure 5f, Table S28). Most enriched terms were related to the CAM pathway in plants.

Morphological detection and verification of CAM pathway genes in the ice plant

Ice plants reportedly survive high-salinity conditions because their EBCs sequester up to 1 M sodium to adjust to osmotic stress (Cushman et al., 1989). The seedlings showed obvious wilt symptoms after 1 day of salt treatment. This phenotype became more pronounced in the cotyledons with increasing treatment time, while the mature leaves examined on days 3 and 6 displayed minor changes, and the whole plants seemed to be more energetic compared to day 1 (Figure 6a). The fresh weight to dry weight ratio after 1, 3 and 6 days of salt stress showed a downward trend compared with that on day 0 (Figure 6b). CAM plants assimilate carbon dioxide at night, accumulate malic acid and transfer it to vacuoles for storage, resulting in an increase in the hydrogen ion concentration in vacuoles. Therefore, the change in titratable acidity in leaves at the start (8:00 pm) and end (8:00 am) was determined as a vital indicator of the presence or absence of CAM activity. On day 0, titratable acidity was very low and did not fluctuate at the end of the day and night (Figure 6c). On day 1 of salt treatment, with obvious phenotypic differences, the nocturnal acid content at dawn was significantly higher than that on day 0. Consistent with this result, on day 0, the stomatal aperture was larger in the day and smaller at night, whereas the plant showed irreversible stomatal movement on days 1, 3 and 6 after stress (Figure 6d). Therefore, we inferred that the plant quickly shifted from C₃ to CAM photosynthesis after 1 day of salt treatment in our study.

In ice plants, genes with CAM-related functions exhibit stress-induced/putative circadian expression patterns following salt stress (Cushman et al., 2008). The aforementioned analysis showed that the expression of four MDH and three PEPC genes was largely altered during the transition of ice plants from C₃ photosynthesis to CAM (Figure 5b). Among these, the expression levels of two MDH genes decreased during the transition phase compared with the control, and two PEPCs showed opposite expression trends: one was time dependently induced, and the other was time dependently repressed with salinity stress (Figure 5c). Several PEPC and MDH genes were selected for qRT-PCR to validate the expression patterns of the individual genes. Among the four genes encoding MDH, the expression level of Mc01G01317 was relatively low and repressed by salinity stress at midday in all salt treatments (Figure 6e). Mc07G01398 was rapidly induced more than twice, at midday and midnight on day 1. However, compared with day 0, the change in the later stages was not significant. The transcript abundance of the other two MDH genes, Mc07G01448 and Mc03G00793, increased on day 1 both at midday and midnight; the genes maintained a higher expression state on days 3 and 6 after salt treatment (Figure 6e), which correlated with the nocturnal accumulation of acidity (Figure 6c). Remarkably, Mc03G00793 showed a putative circadian expression pattern that peaked at midnight and was more pronounced on day 6 of salinity, suggesting that this gene may be specialized in CAM function. Mc08G01316 and Mc07G01375 encode two PEPC genes (the key enzymes responsible for assimilating CO₂ during the night in CAM plants). In our study, two PEPC genes were immediately induced by salt treatment on day 1 and relatively abundant transcripts were maintained on subsequent days (Figure 6f). In addition, they both exhibited an apparent circadian fluctuation in expression level, but in opposite directions. Mc08G01316 had peak expression in the middle of the day, while the expression level of Mc07G01375 peaked at subjective midnight (Figure 6f).

In addition, we detected CAM-related genes in 22 other species (11 C₃, 7 C₄ and 4 CAM species) by comparative analysis (Figure 6g). Interestingly, we detected 111 genes in the C₄ plant Saccharum spontaneum, which was far more than in any other species (Figure 6g, Table S29). Only eight genes were found in another C₄ plant, Amaranthus hypochondriacus. Among the C₃ plants examined, the number of CAM genes ranged from 18 to 49, C₄ plant genes ranged from 8 to 111 and CAM plant genes with 1000 bootstrap repeats in ice plant and other 22 related species.

Figure 6 Morphological detection and CAM pathway genes verification in ice plant. (a) Representative images of plants during the course of 0, 1, 3 and 6 days of salt treatments. (b) Fresh weight to dry weight ratio. (c) Levels of titratable acidity of leaves measured at the start (8:00 am) and end (8:00 pm) of the photoperiod. Each value is mean ± standard deviation. The asterisks above the bars indicate Student’s t test significance in comparison with day 0 (*P < 0.05; **P < 0.01). (d) The representative images of stomata under different salt treatment time courses. Expression profiles of representative MDH and PEPC genes determined at midday and midnight in leaves of M. crystallinum under different stages of salt stress. (e) The relative expression level of four MDH genes under different stages of salt stress by qRT-PCR. (f) The relative expression level of two PEPC genes under different stages of salt stress by qRT-PCR. (g) The heat map of the CAM pathway genes (PPDK, PEPC, MDH, CA, PEPC and ME) number in ice plant and other 22 related species, including C₃, C₄ and CAM plants. The gene number in each species was transformed by log2. (h) Maximum-likelihood trees of PEPC genes that were constructed using the amino acid sequences with 1000 bootstrap repeats in ice plant and other 22 related species.
Deciphering of the first ice plant genome
ranged from 21 to 46 (Figures S20–S25, Table S29). This phenomenon indicates that the number of genes involved in the CAM pathway varies widely among different species, despite belonging to the same metabolic type. Interestingly, the number of phosphoenolpyruvate carboxylase kinase (PEPCK) genes in the basal angiosperm plant Amborella trichopoda was much higher than that of the other examined species. This result provides abundant data resources and ideas for the evolution of the PEPCK gene family. Cluster analysis performed according to the number of CAM-related genes revealed that ice plants clustered with C3 and CAM plants and were most similar to the CAM plant Ananas comosus (pineapple) (Figure 6g). In addition, the phylogenetic trees of genes encoding the six main enzymes indicated that most genes in the ice plant had a close relationship with CAM or C3 plants. In particular, three PEPC genes in ice plants were distributed in different branches in the phylogenetic tree, indicating their probable function in the C3 and CAM pathways (Figure 6h). To some extent, these results support the position that ice plants are C3 and CAM-facultative plants.

Discussion

The ice plant is a member of the Aizoaceae family, which belongs to the order Caryophyllales. The genomes of several species in this order have been sequenced and released, including 14 species from Amaranthaceae, four species from Drosoraceae, four species from Polygonaceae, two species from Cactaceae and one species each from Simmondsiaceae, Caryophyllaceae and Phytolaccaceae. To date, no species in Aizoaceae has been sequenced and reported. The ice plant genome data in this study will provide a rich resource for conducting comparative and functional genomic studies in Aizoaceae and other families of the order Caryophyllales.

The CAM pathway is highly plastic in ice plants (Cushman, 2001; Cushman et al., 2008; Cushman and Bohnert, 1999). Under non-stressed conditions, the ice plant shows C3 photosynthesis and can complete its entire life cycle in the C3 mode without exhibiting net CO₂ absorption at night (Winter and Holtum, 2007). However, it exhibits all the physiological characteristics of a CAM plant when grown under various stress conditions such as high salinity, water deficit or high light (Matsuoka et al., 2018; Wakamatsu et al., 2021; Winter and Holtum, 2005). The indubious identity of the CAM and the biochemical properties of C3 and CAM in the same cells of ice plants make it an excellent model for studying the transition mechanism from C3 to CAM (Bohnert and Cushman, 2000; Kong et al., 2020; Kore-eda et al., 2013; Winter and Holtum, 2014).

The annual succulent ice plant is a typical model halophyte used to explore the basic biochemical, physiological and molecular mechanisms of salt and water stress responses in plants (Barkla et al., 2009; Bohnert and Cushman, 2000). Ice plants can adjust their metabolic pathway from C3 to CAM in response to various stresses (Adams et al., 1998; Oh et al., 2015; Winter and Holtum, 2007).

In vascular plants, PEPC belongs to a multigene family, with each member encoding a specific functional enzyme (O’Leary et al., 2011). In CAM species, at least one form of the gene specializes in CAM function, which catalyses nocturnal CO₂ assimilation into C₄-dicarboxylic acids (Gehrig et al., 1998; Lepinec et al., 1993). The results of phylogenetic analyses of PEPC genes imply a single origin before the divergence of bacterial and plant lineages (Itzu et al., 2004; Westhoff and Gowik, 2004). CAM-specific PEPC genes were thought to have evolved first because of a lack of water supply from the non-photosynthetic role of PEPC by local gene duplications, which could then allow the subsequent functional specialization of genes divided into different clades (Gehrig et al., 2005; Taybi et al., 2004). In the present study, three PEPC genes were assigned to different clades, indicating functional divergence.

MDH is a ubiquitous enzyme in plants with different genes, with roles in a series of metabolic processes according to its subcellular location (Gietl, 1992). In particular, cytosolic MDH converts oxaloacetate to l-malate in CAM plants (Holtum and Winter, 1982). In our study, the expression levels of four MDH genes were determined during the transition from C3 to CAM, indicating different expression patterns of the genes. Among these, Mc03G00792 presented a putative circadian fluctuation in mRNA abundance that peaked at subjective midnight, implying its specific function in CAM photosynthesis.

Conclusions

Here, we report a high-quality and chromosomal-level ice plant genome. This is the first released genome of an Aizoaceae family member. The total length of the genome is 377.97 Mb comprising 24 234 genes. Although no recent WGD or WGT events occurred in ice plants, we detected a novel WGT event that was overlooked in a previous report on S. chinensis. Several key genes involved in the CAM pathway were identified and a comprehensive network was constructed for CAM-related genes in ice plants. The ice plant genome sequences, together with the comparative genomic analysis data, will provide rich resources for studies of gene functions and genome evolution in ice plants and other Aizoaceae plants.

Materials and methods

DNA sequencing and genome size estimation

Genomic DNA was extracted from ice plant leaves using a QIAGEN kit, according to the standard procedure. DNA purity was determined using a NanoDrop™ One spectrophotometer. DNA quantification was performed using a Qubit® 3.0 fluorometer. Sequencing libraries were sequenced using the Illumina (USA) and PacBio (Pacific Biosciences, USA) platforms according to previous reports (Song et al., 2021b; Song et al., 2022). Three sequencing strategies were used. In the first, two paired-end libraries were constructed with fragments of 350 bp and sequenced on the Illumina platform. In the second strategy, third-generation sequencing libraries were constructed and sequenced using the PacBio HiFi platform according to the manufacturer’s protocol. In the third strategy, Hi-C technology combined with Illumina sequencing was used to assist genome assembly. The ice plant genome size was estimated according to the 17 nt k-mers using Illumina sequencing data (Marcais and Kingsford, 2011).

Data quality control and de novo genome assembly

Third-generation data quality control was performed using SMRT Link (v11.0) software (https://www.pacbio.com/support/software-downloads/). The original data were polymerase reads of dumbbell-shaped structural sequences containing adapters at both ends. Subreads were obtained after the sequences were interrupted by the adapters and the adapter sequences were filtered out. Subreads were filtered using the minimum length
when the number of low-quality (< value) database (Bao et al., 2021). HiFiasm was developed based on the characteristics of PacBio HiFi reads. Therefore, it is more prominently used than other software in the assembly of HiFi data. HiFiasm assembly is divided into three steps. The first step is error correction, in which HiFiasm uses all HiFi reads for all-vs-all comparisons and error correction. The second is construction of the assembly graph. After correction, a phased string graph was constructed according to the overlap between the sequences. The third is, the generation of assembly sequence, in which HiFiasm selects one side of the bubble to build the primary assembly.

Hi-C data-assisted assembly and genome annotation

Hi-C data quality control includes alignment control and HiCUP quality control (Wingett et al., 2015). A Perl script was used for alignment control for three reasons. First, reads were removed using adapters. Second, reads with a ratio of N (N means that the base could not be determined) > 10% were removed. Third, when the number of low-quality (<5) bases contained in the single-end sequencing read exceeded 20% of the read length, the paired reads were removed. HiCUP quality control analysis was performed in three steps: (i) reads were aligned to the reference genome, (ii) the hicup_filter in the HiCUP software was used to filter the sequences in the alignment and the hicup_deduplicator in HiCUP was used to filter duplicate contacts and (iii) the ratio of the number of valid and unique read pairs after deduplication to the number of read pairs after quality control was calculated.

Based on Hi-C technology, the ALLHiC program was used to assist in ice plant genome assembly (Zhang et al., 2019b). Clustered bam files and genomes from ALLHiC were visualized using the Juicebox program (Durand et al., 2016). Manual correction was performed according to the strength of chromosome interaction. The genome was obtained at the chromosomal level. Finally, the assembled genome was assessed using BUSCO (embryophytaodb10) and CEGMA software (Manni et al., 2021; Parra et al., 2007). Genomes and second-generation data were aligned using BWA software (Li and Durbin, 2009). The alignment rate, genome coverage and depth of reads were determined to assess the integrity of the assembly and uniformity of sequencing.

Genome annotation

Repeated sequences were detected using de novo prediction and homologous alignment. First, a repeat sequence database was built using RepeatModeler, LTR_FINDER (Xu and Wang, 2007), RepeatScout (Price et al., 2005) and Piler (Edgar and Myers, 2005) for de novo estimation. Second, repeated sequences were predicted using Repeatmasker. The repeatproteinmask and Repeatmasker programs were used to conduct homologous sequence alignment by searching the RepBase database (Bao et al., 2015; Tarailo-Graovac and Chen, 2009). Tandem repeat sequences were detected using TRF software (Benson, 1999). sSnRNAs and miRNAs were identified using INFERNAL (Nawrocki and Eddy, 2013). tRNA and rRNA genes were detected using tRNAscan-SE and BLAST (E-value <1e-5), respectively (Chan and Lowe, 2019). Simple sequence repeat (SSR) was identified according to the previous reports (Song et al., 2021a; Song et al., 2021c).

Protein-coding gene prediction and functional annotation

De novo prediction was performed using three software programs: GlimmerHMM (Stanke and Morgenstern, 2005), SNAP (Korf, 2004) and Augustus (http://bioinf.uni-greifswald.de/augustus). Homologous prediction was conducted using GeneWise and BLAST (E-value <1e-5) (Birney et al., 2004; Camacho et al., 2009). The predicted results were integrated using the IntegrationModeler (EVM) pipeline (Haas et al., 2008). Finally, the results of EVM gene prediction were corrected by combining them with transcriptomic data using the PASA software (Haas et al., 2003). Several protein databases, including TrEMBL, SwissProt, InterPro and KEGG, were used to conduct gene annotation with an E-value cut-off of 1e-5. The distribution of genes, repeat sequences and non-coding genes on each chromosome was illustrated using TBtools (Chen et al., 2020).

Detection of gene families and expansion analysis

Gene families were detected using OrthoFinder (Emms and Kelly, 2019). First, alternative splicing was filtered for each species. Only the longest transcript was retained for gene family analysis. Second, genes with amino acid lengths of <50 were removed. Third, all-vs-all BLAST was performed using the protein sequences to obtain the similarity relationships of all examined species (E-value <1e-5). Finally, single-copy and multi-copy gene families were detected by conducting cluster analysis based on the MCL graph clustering algorithm (Inflation = 1.5). Gene family contraction and amplification were performed using CAFE software (–p 0.05 -t 4 -r 10 000; De Bie et al., 2006).

Phylogeny and divergence time analysis

Genes from single-copy gene families were used to perform multiple sequence alignments using MUSCLE (Edgar, 2004). The RAxML program (–m PROTGAMMAAUTO -p 12345 -x 12 345 -# 100 -f ad -t 20) was used to construct a phylogenetic tree of 13 species based on the maximum-likelihood (ML) model (Stamatakis, 2014). Single-copy gene families, combined with the species trees, were used to estimate the divergence time using the Mmctree method of the PAML program (burn-in = 50 000; sample frequency = 50; sample number = 10 000) (Yang, 2007). Time correction points were obtained from the TimeTree database (Kumar et al., 2017).

RNA-seq and public data set collection

RNA was extracted from the roots, stems and leaves of ice plants. The purity of the RNA(OD260/280) was determined using a NanoDrop spectrophotometer. RNA concentration was quantified using Qubit, and RNA integrity was detected using an Agilent 2100 device. RNA-seq libraries were constructed using an AMPure XP kit (Beckman, China), according to the manufacturer's instructions. RNA sequencing was performed on an Illumina HiSeq 4000 with 150-bp paired-end reads.

Two public RNA-seq data sets were downloaded from NCBI and DDBJ databases to comprehensively explore the expression patterns of genes under salt stress. The raw reads of the second data set from guard cells of control and salt-treated ice plants were collected at 12 am and 12 pm from 5 to 7 days after salt treatment (SRX3878746) (Kong et al., 2020). Raw reads of third
data sets were from ice plants under salt treatments with 0, 140, 250 and 500 mM NaCl (ADRP002316) (Tsukagoshi et al., 2015).

**RNA-seq analysis**

The quality of all raw reads, including our RNA-seq data and public data sets, was assessed using the FastQC program (https://github.com/}"w-]andrews/FastQC). The trim-galore program was used to filter bad quality reads and remove adaptors (https://anaconda.org/biobolna/trim-galore). Finally, the clean reads were mapped to the ice plant genome using HISAT2 (~ p 12 – x Index) (Kim et al., 2015). The expression value of each gene was normalized to fragments per kilobase of exon per million mapped fragments (Trapnell et al., 2010). Analysis of DEGs was performed using DESeq with parameters set as [log2(fold-change)] >1 and P-adj. < 0.05 (Anders and Huber, 2010; Wu et al., 2021).

**Genome collinearity and visualization**

Genome collinearity was assessed using the WGD indexing program, which integrates an improved ColinearScan version (-icl model) (Sun et al., 2021; Wang et al., 2006). First, homologous genes within or between two species genomes were identified using the Blastp program (E-value <1e-5). Then, the -icl model was adopted to run the WGD for collinearity detection. The maximal gap length of collinearity was set to 50, and over 30 gene families were deleted before running -icl. A dot plot of collinear genes was generated using WGDi (Sun et al., 2021).

Using the grape genome as a reference, we built collinear alignments for each species. Theoretically, each grape genome has two additional collinear genes because of a WGT event (Jaillon et al., 2007). In grapes, the cell of the column was filled with a gene name if a collinear gene was detected. The cell was marked with a dot if a collinear gene was absent. We also assigned the corresponding number of columns according to the situation of the WGD or WGT events for ice plants and other species. Finally, collinear alignment was visualized using a Circos plot, which was created using the –ci module in WGDi (Sun et al., 2021). Synteny and microsynteny among different species were visualized using MCScan in Python (Tang et al., 2008). The duplicate_gene_classifier program MCScanX was used to predict duplicated gene types (Wang et al., 2012).

**Ks calculation and distribution fitting**

The MUSCLE program was used to perform alignment using homologous amino acid sequences (~maxiters 1 –diags –sv –distance1 kbit20-3) (Edgar, 2004). The PAL2NAL program was used to convert the protein alignment into a codon alignment according to the CDS sequence (Suyama et al., 2006). Finally, Ka and Ks were calculated using the yn00 program of PAML with the Nei-Gojobori method according to the previous reports (Pei et al., 2021a; Pei et al., 2021b; Yang, 2007). In collinear blocks, the median values of Ks between homologous genes were used to classify blocks caused by duplication events. Ks was illustrated on a collinear block of different colours using the WGDi (Sun et al., 2021). The distribution of Ks density was determined using three modules of WGDi: Kspeaks (~kp), PeaksFit (~pf), and KsFigures (~kF). The curve of the Ks density distribution was drawn using Kspeaks. Multipeak fitting was then performed using PeaksFit. Finally, KsFigures were used to convert multiple fitted density curves into one graph.

**CAM pathway genes analyses**

Based on the Pfam annotation, CAM-related genes were identified using accession numbers with E-values <1e-5 (Table S22). The chromosomal distribution of CAM-related genes was determined using TBtools (Chen et al., 2020). The protein sequences of CAM-related genes were aligned using the Mafft program (~maxiterate 1000 –localpairs) (Nakamura et al., 2018). An ML tree was then constructed using FastTree using the JTT model with a bootstrap set of 1000 (Price et al., 2009; Yu et al., 2022). PCCs between CAM-related genes and DEGs were calculated based on gene expression using Perl scripts. Based on |PCC| >0.95, an interaction network was constructed using Gephi software with the algorithm ForceAtlas2 (https://gephi.org) (Jacomy et al., 2014). KEGG functional enrichment analysis was performed using the OmicShare platform (q-value <0.05) (https://www.omic-share.com/tools).

** Morphological detection and CAM pathway gene verification**

**Plant growth and salt stress**

Ice plants were grown in a growth chamber under 200 μmol/m²/s white light with a 12 h (26 °C) day/12 h (18 °C) night cycle. After 1 week, seedlings were transferred to 32-ounce containers and nourished using 0.5× Hoagland’s solution every 4 days. To induce CAM, salt treatment was conducted on day 28 after sowing by irrigation with 0.5 mM NaCl in 0.5× Hoagland’s solution once daily. Before salt treatment (day 0, control) and at 1, 3 and 6 days of stress, the second pair of mature leaves was harvested at midday and midnight for three replicate plants.

**Stomatal aperture assay**

For stomatal movement observation, a part of the abaxial epidermis was peeled with tweezers at 4:00 am and 4:00 pm at each treatment time point. The peeled epidermal strips were immediately placed onto a microscope slide and observed under a Leica DM6000 B microscope.

**Titratable acidity measurement**

Mature leaves were collected from control and salt-treated plants at the end of the light and dark periods. After measuring fresh weight, the samples were frozen in liquid nitrogen and ground to a fine powder using a mortar and pestle. For organic acid extraction, the powder was homogenized in 20% ethanol, boiled until volume was reduced by half, suspended in water to the original volume and boiled to half original volume. Finally, the sample was returned to its original volume using distilled water and cooled to room temperature. Titratable acidity was determined by titration with 5 mM NaOH, using phenolphthalein as an indicator. The volume of NaOH was recorded, which could be switched to leaf titratable acidity in the form of μmol H⁺ g⁻¹ fresh weight.

**RNA extraction and RT-qPCR**

Total RNA was isolated using RNAsio Plus (TaKaRa Bio, Japan). Integrity and quality were assessed by electrophoresis and spectrophotometry. First-strand cDNA was synthesized using the PrimeScript RT Reagent Kit, according to the manufacturer’s instructions. The resulting cDNA was diluted 1:5 in double-distilled water before RT-qPCR analysis using an Agilent MX3005P.
qPCR System Cycler. Each sample was run in triplicate and normalized by comparison with \textit{FNR1}. Primers for the \textit{PEPC} and \textit{MDH} genes are listed in Table S30.

Acknowledgements

This work was supported by the Natural Science Foundation for Distinguished Young Scholar of Hebei Province (C2022209010), the National Natural Science Foundation of China (32172583), the Natural Science Foundation of Hebei (C2021209005), and the China Postdoctoral Science Foundation (2020M673188, 2021T140097). The genome sequencing and Hi-C were conducted in the Novogene Corporation.

Conflicts of interest

The authors declare no competing interests.

Author contributions

X.S. conceived the project and were responsible for the project initiation. X.S. and S.S. supervised and managed the project and research. Experiments and analyses were designed by X.S., N.L. and S.S. Data generation and bioinformatic analyses were led by X.S., S.S., T.Y., Z.L., Y.W., R.Z., P.S., Z.W., X.T., C.Z., S.F. and Y.Z. The manuscript was organized, written and revised by X.S., N.L., H.L. and W.C. All authors read and revised the manuscript.

Data availability statement

The genome sequence and RNA-seq data sets of ice plant reported in this paper have been deposited in the Genome Sequence Archive (Wang \textit{et al.}, 2017) in Big Data Center (Members, 2019), Beijing Institute of Genomics (BIG), Chinese Academy of Sciences, under accession numbers CRA005754 and CRA005756 that are publicly accessible at http://bigd.big.ac.cn/gsa. All materials and related data in this study are available upon request.

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Supporting information

Additional supporting information may be found online in the Supporting information section at the end of the article.

Figure S1 K-mer distribution of the ice plant genome with the K-mer = 17.

Figure S2 The length and frequency of the main types of repetitive sequences in ice plant genome, including DNA, LINE, LTR, and SINE repeats.

Figure S3 Comparative analysis of CDS length, exon length, exon number, gene length, and intron length in ice plant and other representative species.

Figure S4 The Venn diagram of gene sets evidence in ice plant genome.

Figure S5 The Venn diagram of gene function annotations in ice plant by four databases.

Figure S6 The chromosomal distribution of ice plant ncRNAs, including tRNA, rRNA, miRNA, and snRNA.

Figure S7 Syntenic comparison between ice plant (mcr) and A. hortensis (acr) or V. vinifera (vi).

Figure S8 Syntenic comparison between ice plant (mcr) and F. tataricum (fta) or V. vinifera (vi).

Figure S9 Syntenic comparison between ice plant (mcr) and A. cruentus (acru) or B. vulgaris (bvu).

Figure S10 Syntenic comparison between ice plant (mcr) and C. pallidicaule (cpa) or S. oleracea (sol).

Figure S11 Gene retention analysis of ice plant genome comparing with grape.

Figure S12 Gene retention analysis of ice plant genome comparing with A. cruentus.

Figure S13 Gene retention analysis of ice plant genome comparing with A. hortensis.

Figure S14 Gene retention analysis of ice plant genome comparing with B. vulgaris.

Figure S15 Gene retention analysis of ice plant genome comparing with C. pallidicaule.

Figure S16 Gene retention analysis of ice plant genome comparing with C. quinoa.

Figure S17 Gene retention analysis of ice plant genome comparing with F. tataricum.

Figure S18 Gene retention analysis of ice plant genome comparing with S. chinensis.

Figure S19 Gene retention analysis of ice plant genome comparing with S. oleracea.

Figure S20 Maximum-likelihood trees of CA genes that were constructed using the amino acid sequences with 1000 bootstrap repeats in ice plant and other 22 related species.

Figure S21 Maximum-likelihood trees of MDH genes that were constructed using the amino acid sequences with 1000 bootstrap repeats in ice plant and other 22 related species.

Figure S22 Maximum-likelihood trees of ME genes that were constructed using the amino acid sequences with 1000 bootstrap repeats in ice plant and other 22 related species.

Figure S23 Maximum-likelihood trees of PPDK genes that were constructed using the amino acid sequences with 1000 bootstrap repeats in ice plant and other 22 related species.

Figure S24 Maximum-likelihood trees of PEPC genes that were constructed using the amino acid sequences with 1000 bootstrap repeats in ice plant and other 22 related species.

Figure S25 Maximum-likelihood trees of PPDK genes that were constructed using the amino acid sequences with 1000 bootstrap repeats in ice plant and other 22 related species.

Table S1 Statistics of sequencing data obtained by Illumina HiSeq platform for ice plant genome survey.

Table S2 K-mer statistics of the genomic characteristics of ice plant genome obtained by genome survey.

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