

VERSION 1

MAR 01, 2023

OPEN ACCESS

DOI:

dx.doi.org/10.17504/protocol s.io.6qpvr4qdogmk/v1

Protocol Citation: Ryoma Sato, Yuri Kondo, Sakae Agarie 2023. Methods in "The first released available genome of the common ice plant (Mesembryanthemum crystallinum L.) extended the research region on salt tolerance, C3-CAM photosynthetic conversion, and halophism". protocols.io https://dx.doi.org/10.17504/protocols.io.6qpvr4qdogmk/v1

License: This is an open access protocol distributed under the terms of the Creative Commons Attribution License, which permits unrestricted use, distribution, and reproduction in any medium, provided the original author and source are credited

Protocol status: Working We use this protocol and it's working

Created: Feb 28, 2023

Last Modified: Mar 01, 2023

Methods in "The first released available genome of the common ice plant (Mesembryanthemum crystallinum L.) extended the research region on salt tolerance, C3-CAM photosynthetic conversion, and halophism" V.1

Ryoma Sato¹, Yuri Kondo¹, Sakae Agarie²

¹Graduate school of Bioresource and Bioenvironmental Sciences, Kyushu University;

²Faculty of Agriculture, Kyushu university

Sakae Agarie: Corresponding author;



Ryoma Sato

ABSTRACT

The wild-type seeds of the common ice plant were sowed on a germination medium. The seedlings were grown in particular soil and treated with a solution that included salt and nutrients for two weeks in a greenhouse. Genomic DNA was extracted from frozen leaves and made into a library using special kits. Obtained NGS data were trimmed and assembled by *Musket*, *ALGA*, and *Redundans*. The completeness of the genome was checked using BUSCO and BLASTN.

In this protocol, five types of analysis methods were introduced, including the establishment of a phylogenetic tree based on 18S rDNA via *NGPhylogeny.fr*, detection of repetitive regions with *RepeatModeler2*, *TEclass*, and *RepeatMasker*, search for genomic sequences coding tRNA and miRNA by *tRNAscan-SE2.0* and *Infernal*, gene prediction using *BRAKER2* and *DIAMOND*, and protein domain searches based on Pfam database using HMMER.

PROTOCOL integer ID: 77834

Keywords: De novo shotgungenome assembly, the common ice plant, halophytes

MATERIALS

Seeds of the common ice plant (Mesembryanthemum crystallinum) were personally provided by Dr. John C. Cushman from the University of Nevada and stored under coolness and darkness until use. Originally, wild-type seeds were collected from the plants identified by Dr. Klaus Winter, an expert on the common ice plant, on a coastal cliff at the Mediterranean Sea shore close to Caesarea in Israel (around 32° 29' 43.4"N, 34° 53' 22.8"E) in 1978 (Winter et al. 1978). Three voucher specimens of *M. crystallinum* have been deposited in the Herbarium at the Royal Botanic Gardens Kew (55793.000, K000296094, and K000267571). In this study, our biological materials were recognized as the same plants as those specimens.

Experiments, including collecting samples for this study, were conducted in compliance with relevant institutional, national, and international guidelines and laws.

The seeds were aseptically sown on a medium for germination containing 4.6 g MS salt (mixed salts for Murashige-Skoog medium) ∆ 30 g sucrose △ 1 mL B5 vitamin (Gamborg et al., 1968; △ 1 g nicotinic acid △ 1 g pyridoxine hydrochloride , △ 10 g thiamine hydrochloride , and \perp 0.8 % (w/w) agarose , and \perp 100 g myo-inositol per \perp 1 L of B5 vitamin), ©H 5.7 per 🛕 1 L . The raising of seedlings was performed according to the methods published by Agarie et al. (2009). The two-week-old seedlings grown in a growth chamber under 12:00:00 of light and (5) 12:00:00 of darkness at 4 25 °C were transferred to plastic pots filled with the growth medium soils composed of 4 50 % peat moss, △ 30 % cocopeat , and △ 20 % perlite , specified for the ice plants (Japan Agricultural Cooperatives Ito-Shima, Fukuoka, Japan) and irrigated with a nutrient solution of 🚨 1.5 g per one litter OAT House No. 1 and 🗸 1.0 g per one litetr OAT House No. 2 (OAT Agrio Co., Ltd., Tokyo, Japan) in a greenhouse at Kyushu University 33°35'35.1"N 130°12'53.2"E for five weeks. The plants were treated with the liquid solution including 4 0.3 % (w/w) NaCl for two weeks. Approximately \(\Lambda \) 0.6 g of tissue from each leaf \(\text{was collected, quickly} \) frozen in liquid nitrogen, and stored at -80°C

Table of contents

- 1 ① DNA extraction, library construction, and sequencing
 - 2 Clean read preparation and genome size estimation
 - ③ De novo genome assembly and quality evaluation
 - 4 Phylogenetic tree creation among multiple plant species using 18S ribosomal DNA sequences
 - ⑤ Detection of repetitive regions

- © Search for genomic sequences coding transfer RNA (tRNA) and micro-RNA (miRNA)
- ② Gene prediction
- ® Protein domain searches

1 DNA extraction, library construction, and sequencing

Total genomic DNA was extracted from the leaf tissue and purified using MagExtractor™-Plant Genome Nucleic Acid Purification Kits (Toyobo Co., Ltd., Shiga, Japan), according to the manufacturer's instructions. The DNA samples were fragmented by sonication and used to construct short insert paired-end libraries construction using NEBNext® Ultra™DNA Library Prep Kits for Illumina (New England Biolabs Ltd., Ipswich, MA, USA). Briefly, in the end-repair step, fragmented DNA was phosphorylated at the 5' end and adenylated at the 3' end. During the ligation step, full-length circulated adaptor sequences were ligated to the fragments. After adaptor cleavage, purification, and size selection were performed. The indexed PCR products were taken to obtain the final sequencing libraries. The mean insert size for paired-end libraries was 300 bp. The paired-end (2×150 bp) sequencing was conducted on an Illumina NovaSeq 6000 platform (Illumina Inc., San Diego, CA, USA).

2 Clean read preparation and genome size estimation

The mean insert size was calculated using REAPR (v1.0.18)(<u>Hunt et al. 2013</u>), and raw pairedend sequences were filtered based on the frequency of 21-mer sequences using the program Musket (v1.1)(<u>Liu et al. 2013</u>).

The key parameter values were as follows: musket -omulti output -inorder pair1.fastq pair2.fastq.

```
#download of musket
#download from https://sourceforge.net/projects/musket/ and moved
it to the DDBJ NIG sUPER COMPUTER server using SFTP command
cd $HOME
tar xvzf musket-1.1.tar.gz
cd musket-1.1/
make
./musket #check
cd $
~/musket-1.1/musket -omulti output -inorder pair1.fastq pair2.fastq
-p 12
```

Sequence reads that appeared rarely or abnormally frequently were removed to obtain clean read data. In the corrected reads, unique and duplicate read numbers in the corrected reads were measured using fastqc (v0.11.9)(Simon 2010). The clean data were used for an estimate of genome size as follows.

```
#quality check (fastqc, multiqc)
fastqc XX.fq.gz -o XX
multiqc ./
```

K-mers were counted and exported to histogram files using jellyfish (v2.3)(<u>Marçais and Kingsford 2011</u>) [key parameter: jellyfish histo reads.jf].

GenomeScope2.0($\underline{Ranallo-Benavidez}$ et al. 2020) corresponding key parameters were applied to calculate the genome sizes using k-mers lengths of 21 and 25.

```
#How to estimate genome size using GenomeScope2.0
#REFERENCES:
#GenomeScope 2.0 for estimating genome size and heterozygosity of
ploidy genomes from WGS reads
#Original article ↓ Article
#Ranallo-Benavidez, T. R., Jaron, K. S., & Schatz, M. C. (2020).
#GenomeScope 2.0 and Smudgeplot for reference-free profiling of
polyploid genomes. nature communications, 11(1), 1432.
https://doi.org/10.1038/s41467-020-14998-3
git clone <a href="https://github.com/tbenavi1/genomescope2.0.git">https://github.com/tbenavi1/genomescope2.0.git</a>
cd genomescope2.0/
mkdir ~/R libs
echo "R LIBS=~/R libs/" >> ~/.Renviron
#Rscript install.
Move raw data (fastg format) to the specified directory
First, analyze the fastq file.
jellyfish count -C -m 21 -s 1000000000 -t 12 *fg -o reads k21.jf
jellyfish count -C -m 25 -s 1000000000 -t 12 *fq -o reads k25.jf
#-m Length of mer
#-s Initial hash size
#-t Number of threads (1)
Output a histogram file.
jellyfish histo -t 12 reads.jf > reads.histogram
Output a graph of k-mer spectrum. k-mer max recommended value is
1000.
~/Important Software/genomescope2.0/genomescope.R -i reads.histo -o
```

```
output_dir2 -k 21 -p 2

-k kmer length used to calculate kmer spectra [default 21]
-i input histogram file
-o output directory name
-p ploidy (1, 2, 3, 4, 5, or 6) for model to use [default 2]

When running GenomeScope2, if you make a mistake in specifying the number of ploidy, the estimated value will change.
```

If you are not sure about the ploidy and want to estimate the number of ploidy and whether it is heteroploidy or homoploidy, use smudgeplot first.

http://kazumaxneo.hatenablog.com/entry/2019/04/18/073000

3 De novo genome assembly and quality evaluation

The reads were assembled using ALGA (v1.0.3; <u>Swat et al. 2021</u>) with the default parameter – error-rate = 0.02. long DNA fragments 1 to 10 kb in length were combined, and gaps between them were filled with unknown bases (Ns) using Redundant (v0.14a; <u>Pryszcz and Gabaldón 2016</u>), a software program for scaffolding, with default parameter values.

```
#Using ALGA
#How to install #ALGA
(https://kazumaxneo.hatenablog.com/entry/2021/01/22/121538)
#From git.hub
#Depends on.
#CMake VERSION 2.8.7 or higher
#C++ 17 or higher
#Install the latest version of make
#First, check the version of make
make --version
GNU Make 3.82
#Built for x86 64-redhat-linux-gnu.
#Copyright (C) 2010 Free Software Foundation, Inc.
#License GPLv3+: GNU GPL version 3 or later
<http://gnu.org/licenses/gpl.html>
#This is free software: you are free to modify it and redistribute
it freely.
No #warranty to the fullest extent permitted by law.
If your #make version is 4 or lower, update to 4 or higher.
conda install make
make --version
```

```
Collect package metadata (current repodata.json): done
Solution environment: done
## Package plan ##
## Environment location /home/iceplant4561/anaconda3/envs/gappadder
# Add/update specifications.
# - make
# the following new packages will be installed.
# libgcc mutex conda-forge/linux-64:: libgcc mutex-0.1-conda forge
# openmp mutex conda-forge/linux-64:: openmp mutex-4.5-1 gnu
# libgcc-ng conda-forge/linux-64::libgcc-ng-11.2.0-h1d223b6 11
# libgomp conda-forge/linux-64::libgomp-11.2.0-h1d223b6 11
# make conda-forge/linux-64::make-4.3-hd18ef5c 1
\#proceed([y]/n)?y
#prepare transaction: done
#transaction validation: done
#execute transaction: done
make --version
#GNU Make 4.3
#Built for x86 64-conda-linux-gnu.
#Copyright (C) 1988-2020 Free Software Foundation, Inc.
#License GPLv3+: GNU GPL version 3 or later
<http://gnu.org/licenses/qpl.html>
#This is free software: you are free to modify it and redistribute
it.
No #warranty to the extent permitted by law.
#Update cmake
wget
https://github.com/Kitware/CMake/releases/download/v3.22.1/cmake-
3.22.1.tar.qz
tar zxvf cmake-3.22.1.tar.qz
#Build
cd cmake-3.22.1/
. /bootstrap
build
#pass through the path
echo 'export PATH=$HOME/cmake-3.22.1/bin/:$PATH' >> ~/.bashrc
Source ~/.bashrc
```

```
#Check cmake version
cmake --version
#Check cmake version.
#CMake suite is maintained and supported by Kitware
(kitware.com/cmake).
#How to install and build alga using c++ version 17 (see .2022 Jan
11 email from Mr. Ashizawa, National Institute of Genetics).
qlogin
Module load gcc/9.2.0
wget
https://github.com/swacisko/ALGA/archive/refs/tags/1.0.3.tar.gz
tar zxvf 1.0.3.tar.gz
cd ALGA-1.0.3/
# or
#git clone https://github.com/swacisko/ALGA.git
#cd ALGA/
#either is fine
mkdir build
cd build
cmake -DCMAKE CXX COMPILER=/opt/pkg/gcc/9.2.0/bin/c++ \?
-DCMAKE C COMPILER=/opt/pkg/gcc/9.2.0/bin/gcc ...
make -j 4
#ALGA CMakeCache.txt CMakeFiles cmake install.cmake Makefile
#ALGA build is now complete.
cd $HOME/WGS/iceplant draft contig
~/ALGA/ALGA --file1=output.0.fastq --file2=output.1.fastq --
threads=10 --output=Mc draft genome.fasta --error-rate=0.02
conda activate Redundans
 ~/New redundans/redundans.py -v \
 -i /home/iceplant4561/WGS/iceplant draft contig/output 1.fastg \
 /home/iceplant4561/WGS/iceplant draft contig/output 2.fastq \
 - f
/home/iceplant4561/WGS/iceplant draft contig/Mc draft genome.fasta
-o more scaffolding
```

The genome coverage of reads was estimated using the Mosdepth program (Pedersen and

Quinlan 2018).

```
#Genome Coverage Calculations Using mosdepth
#Reference:
https://kazumaxneo.hatenablog.com/entry/2018/06/06/112849
#http://kazumaxneo.hatenablog.com/entry/2018/04/04/175133
#Mapping fastg data to thegenome using minimap2
nohup singularity exec /usr/local/biotools/m/minimap2:2.9--1 \.
minimap2 -t 10 -a -x sr \setminus.
/home/iceplant4561/Agarie group/Iceplant shotgun genome assembly/mo
re scaffolding/Mc 2nd scaffold.filled.fa \?
/home/iceplant4561/Agarie group/Iceplant shotgun genome assembly/tr
immed/Mc musket 1.fastq \.
/home/iceplant4561/Agarie_group/Iceplant_shotgun_genome_assembly/tr
immed/Mc_musket 2.fastq \ \?
> Mc Genome.sam &.
#Conversion to bam file => sort
samtools view -@ 40 -bS Mc Genome.sam > Mc Genome.bam
samtools sort -@ 40 -o Mc Genome sort.bam Mc Genome.bam
samtools index Mc Genome sort.bam #index place
singularity exec /usr/local/biotools/m/m/mosdepth -t 40 -n
Mc Genome sort.bam
```

The completeness of the assembled genome was evaluated based on the content of orthologs in higher plants, using the benchmarking universal single-copy orthologs (BUSCO) program (v5.0; Manni *et al.* 2021). The lineage dataset was embryophyta_odb10 (creation date: 2020-09-10, number of BUSCOs: 1614).

```
#List creation
singularity exec /usr/local/biotools/b/busco\:5.4.3--pyhdfd78af 0
busco --list-datasets
#genome
singularity exec /usr/local/biotools/b/busco\:5.4.3---pyhdfd78af 0
busco -m geno -i Complete iceplant genome.fasta -o out dir -l
embryophyta odb10 -c 30
#Merge multiple busco data
generate plot.py -wd BUSCO summaries/
*It is necessary to store the short summary* file under
BUSCO summaries beforehand.
In short summary.specific.embryophyta odb10.*.txt, the * part is
the species name.
If you use
short summary.specific.embryophyta odb10.M.crystallinum.txt, it
will be separated by M. Use M crystallinum.
*Maybe you can do it by tinkering with python scripts (230208)
```

We also searched for core genes in the genome sequences of nine other plant species: *Kewa caespitosa, Pharnaceum exiguum, Macarthuria australis, Solanum chaucha, Populus trichocarpa, Arabidopsis thaliana*, and *Oryza sativa* using BUSCO. The first three species belong to the same order, Caryophyllales, to which the ice plants belong. Genome information was obtained from the NCBI (see Supplementary Note 1 "Address to genome information"; <u>Supplementary Information: Sato et al., 2022a</u>). The number of bases, sequences, sequences in several base number ranges, and the maximum base length of the final draft genome sequences was calculated using <u>gVolante</u> (v2.0.0)(<u>Nishimura et al. 2017</u>). BLASTN (v2.2.31+; <u>McGinnis and Madden 2004</u>) was used to investigate the number of cDNA sequences identified by transcriptome (<u>Lim et al. 2019</u>), and registered DNA sequences (retrieved from <u>NCBI</u>, last accessed February 2022) were aligned to the final assembled genome sequence.

Phylogenetic tree creation among multiple plant species .

The 18S ribosomal genes were extracted using barrnap (v0.9; <u>Seemann 2018</u>) from the obtained genome sequences of the ice plant.

```
barrnap --kingdom euk --threads 12 genome.fasta
#Extract 18 S rDNA sequences from result fasta files.
```

As comparative objectives, 25 kinds of 18S ribosomal genes from general crops (Japanese radish [*Raphanus sativus*], Soybean [*Glycine max*], Japanese trefoil [*Lotus japonicus*], Barrelclover [*Medicago truncatula*], Adzuki bean [*Vigna angularis*], Banana [*Musa acuminata*], Barley [*Hordeum vulgare*], Sorghum [*Sorghum bicolor*], Bread wheat [*Triticum aestivum*], Maize [*Zea*

mays], Apple [Malus domestica], Peach [Prunus persica], Coffee tree (Arabica var.) [Coffea arabica], Coffee tree (Robusta var.) [C. canephora], Clementine [Citrus clementina], Orange [C. sinensis], Poplar, Tobacco [Nicotiana tabacum], Tomato [Solanum lycopersicum], Eggplant [S. melongena], Potato [S. tuberosum] and Grape [Vitis vinifera]) were selected using the SILVA database (Release. 2020-08; Pruesse et al. 2007). After joining all ribosomal DNA sequences into one file, a molecular phylogenetic tree was created using implemented in NGPhylogeny.fr (Lemoine et al. 2019) (Released in 2019). SH-aLRT (Shimodaira-Hasegawa-approximate likelihood ratio test) (Shimodaira and Hasegawa 1999) was used to determine the molecular phylogenetic tree.

5 Detection of repetitive regions

Repetitive sequences were detected, and custom repeat libraries involving transposable elements and long terminal repeat-retro transposons were generated using RepeatModeler2 (v2.0.2; Flynn et al. 2020) and TEclass (v2.1.3)(Abrusán et al. 2009). Known repeat sequences were detected and classified in the assembled genome sequence with reference to the Repbase library (Bao et al. 2015) and the custom repeat libraries, using RepeatMasker(v4.1.2-p1; Smit et al. 2013-2015). The capital letters in the genome sequences were replaced with small characters as soft masking.

```
1) Creation of a repetitive array custom repeat library using
RepeatModeler2
conda create -n repeatmodeler RepeatModeler==2.0.3
conda activate repeatmodeler
BuildDatabase -name Mc
~/Important Software/Agarie group/ice plant genome/data/iceplant ge
nome.fasta
gsub -V -cwd -l medium -l s rt=120:00:00 -l d rt=120:00:00 -l
s vmem=30G -l mem req=30G -pe def slot 30 -b y -e .
/error log repeatmodeler -N RepeatModeler \
RepeatModeler \ -database Mc PacB
-database Mc PacBio -pa 29 -genomeSampleSizeMax 370000000 \
-repeatmasker dir ~/Important Software/RepeatMasker/RepeatMasker \
-abblast dir ~/Important Software/ab-blast-20200317-linux-x64/
*Note! It will probably take two days time. Sleep at home.
Mc-families.fa and Mc-families.stk (Stockholm format) will appear
*Name specified with -name during BuilDatabase
2)TEclass is used to classify TEs classified by RepeatModeler in
detail.
singularity exec ~/Important Software/teclass.sif TEclassTest.pl -c
TEclass-2.1.3c/classifiers -o ./ Mc-families.fa
cd *
cp Mc-families.fa.lib ./
~/Agarie group/ice plant genome/Repeat/RepeatMasker
cd ~/Agarie group/ice plant genome/RepeatmaskerLib/
3)RepeatMasker
makeblastdb -in Mc-families.fa.lib -dbtype nucl -blastdb version 4
*If you don't do it first, an error will occur.
RepeatMasker -pa 20 -html -gff -xsmall -lib Mc-families.fa.lib
Mc scaffold.filled.fa
```

© Search for genomic sequences coding transfer RNA (tRNA.

The tRNA genes were identified in the draft common ice plant genome using tRNAscan-SE2.0 (v2.0.9)(Chan et al. 2021).

```
#Identification of tRNAs using tRNAscanSE-2.0
#Reference:https://kazumaxneo.hatenablog.com/entry/2019/05/07/07300
0
singularity exec /usr/local/biotools/t/trnascan-se\:2.0.9--
pl5321hec16e2b_3 \
tRNAscan-SE \
~/Agarie_group/ice_plant_genome/data/iceplant_genome.fasta \
-E -o Mc_tRNA_output -f tRNA_structure -s isotype -m statistics -b
bedfiles -j gff -a fastafile -l worklog --detail --thread 30
```

The tRNA data of other nine plant species—*Arabidopsis*, rice, tomato, poplar, horseradish, potato, grape, soybean, and coffee tree (robusta species)—were obtained from the PlantRNA database (Cognat *et al.* 2013). The percentages of arbitrary tRNAs against the total tRNAs in the genome were calculated and compared to the ice plants' values with those of the other species. Smirnov-Grubbs' outlier tests were performed to select tRNAs more significantly involved. The test statistic T was calculated using the following equation:

```
T=\frac{(Percentage~of~arbitrary~tRNAs~in~the~ice~plant)-
(Sample~mean~for~all~nine~species)}{\sqrt{Sample~variance}}
```

The miRNA loci in the genome sequence were identified using the cmscan command in infernal (v1.1.4; Nawrocki and Eddy 2013) using Rfam.

#Identification of small RNA using Infernal
#Reference: http://eddylab.org/infernal/Userguide.pdf
#http://http.ebi.ac.uk/pub/databases/Rfam/

#In this case, we will use "Searching the Rfam CM database with a query sequence" on p. 29 #of the reference.

#Infernal uses the Singularity image file from the Institute of Genetics.

#Use Covariant Model (CM). I'm not sure, but I'll try to do it as written.

- (1) Get the latest covariance model from Rfam
 wget http://http.ebi.ac.uk/pub/databases/Rfam/14.8/Rfam.cm.gz
 gunzip Rfam.cm.gz
- (2) Get Rfam's clan information (like a family)
 wget http://http.ebi.ac.uk/pub/databases/Rfam/14.8/Rfam.clanin
 mv Rfam.clanin Rfam.14.8.clanin
- (3) Make data available in cmpress
 singularity exec /usr/local/biotools/i/infernal\:1.1.4-pl5321hec16e2b_1 \c
 cmpress Rfam.cm
- (4) Search with cmscan. Do as written.
 singularity exec /usr/local/biotools/i/infernal\:1.1.4-pl5321hec16e2b_1 \cmpress
 cmscan --rfam --cut_ga --nohmmonly --tblout Mc_genome.tblout --fmt
 2 --clanin Rfam.14.8.clanin --cpu 30 \
 Rfam.cm ~/Agarie_group/ice_plant_genome/data/iceplant_genome.fasta
 > Mc_genome.cmscan

7 Gene prediction

The BRAKER2 pipeline (v2.1.5; <u>Brůna et al. 2021</u>) was used for the prediction of genes in the common ice plant genome. Amino acid sequences were translated from the transcriptome profile reported by <u>Lim et al. (2019)</u> and used as additional reference data for the prediction of genes. BRAKER2 was used with the default parameters (–softmasking).

#Annotation of genomes using BRAKER2 At this point, change the header of the fasta file of the masked

genome. In the bam2hints process of BRAKER2, if there is a whitespace (" " ← this) in the fasta header, the error message "The hints file is empty. The hints file is empty. Maybe the genome and the RNA-seg file do not belong together" error occurs. Reference: https://github.com/Gaius-Augustus/BRAKER#common-problems How to change Create a new line cat Iceplant-genome fasta full softmask.fasta | awk '/^>/ { print n $0; n = "" }! /^>/ { printf "%s", $0; n = "\n" } END{ printf "%s",$ n }' > A.fasta mv A.fasta Iceplant-genome fasta full softmask.fasta Preparation for BRAKER below Reference: https://qiita.com/drk0311/items/a3ac648f2780cfee57b1 Obtaining GeneMark-ES/ET/EP http://exon.gatech.edu/GeneMark/license download.cgi Here, add your name and affiliation, and GeneMark-ES/ET/EP ver 4.69_lic LINUX 64 kernel 2.6 - 3 and click on I agree to the terms of this license agreement to go to the next page. Right-click here and click Download Download the key as well Transfer to linux via sftp sftp iceplant4561@gw2.ddbj.nig.ac.jp cd /home/iceplant4561/Important Software put gmes linux 64.tar.gz put gm key 64.gz Back to linux cd /home/iceplant4561/Important Software Extract each tar xzvf gmes linux 64.tar.gz

gunzip gm key 64.gz

#The license is valid for 200 days, so after 200 days, go back to the above site, re-enter your registration information, agree to the license, and then click on the "Download key 32 bit or 64 bit" button. #Please download the license key (gm key 64.gz or gm key 32.gz) from the link "Please download key 32 bit or 64 bit". #If you are using 64 bit now, the majority of users will probably use 64 bit. Unzip the license key with gunzip, rename it to .gm key and save it in your home directory. #Now, you can save the program anywhere you want, but I keep my tools that cannot be managed by Anaconda in a directory named "local" under my home directory (~ or /home/account name) and put them there. #local directory under your home directory (~ or /home/account name) for tools that cannot be managed by Anaconda. The following is a case of dropping the program files into the downloads folder on Windows. cp gm key 64 ~/.gm key /home/iceplant4561/Agarie group/ice plant genome from GSA/BRAKER/gm es linux 64 . /check install.bash Checking GeneMark-ES installation export GENEMARK PATH=/home/iceplant4561/Agarie group/ice plant genome from GSA/BRAKER/gmes linux 64 source ~/.bashrc Create a Docker image container for BRAKER (v2.1.5) cd ~/Important Software/ singularity build braker.sif docker://hamiltonjp/braker2:a765b80 cd ~/Agarie group/ice plant genome/BRAKER/ species="M.crystallinum" species dir="\${PWD}/\${species}" singularity exec /home/iceplant4561/Important Software/braker.sif \ braker.pl --genome=./Iceplant-genome fasta full softmask.fasta \ --species=\${species} braker2 \ --workingdir=./braker2 out \

--prot_seq=Proteins_from_iceplants.fasta \

--softmasking \

--gff3 \
--epmode \

```
--cores 45 \
--
GENEMARK_PATH=~/Agarie_group/ice_plant_genome_from_GSA/BRAKER/gmes_linux_64 \
--
AUGUSTUS_CONFIG_PATH=~/Agarie_group/ice_plant_genome_from_GSA/BRAKER/gmes_linux_64 \
--
--
--
---
AUGUSTUS_CONFIG_PATH=~/Agarie_group/ice_plant_genome_from_GSA/BRAKER/Augustus/config/ \
--useexisting
```

The total sequences, total bases, total amino acids, and N50 were computed based on the resulting fasta-format files containing information about the genes, coding sequences, and amino acids using seqkit (v2.0.0; Shen *et al.* 2016) [key parameter: seqkit stats]. Protein BLAST searches (*E*-value < 1e-5) were conducted using DIAMOND (v2.0.13.151; Buchfink *et al.* 2021) against the NCBI-non-redundant protein sequences (retrieved from NCBI in March 2022), Uniprot-swissprot(retrieved in March 18), Ensemble TAIR10 (retrieved in March 2022), and NCBI poplar amino acid sequence databases(retrieved from NCBI in March 2022).

```
#Output statistics for FASTA files using seqkit stats
seqkit stats -a *.fa

#BLASTP using DIAMOND(ex. NCBI)
singularity exec /usr/local/biotools/d/diamond:2.0.9--hdcc8f71_0
diamond makedb --in nr --db nr

singularity exec /usr/local/biotools/d/diamond:2.0.9--hdcc8f71_0
diamond blastp --query protein_output.fa \
--db ~/blast/database_for_blast/nr.dmnd --max-target-seqs 1 \
--evalue 1e-5 --outfmt 6 --out blast_vs_ncbi.txt -b12 -c1 --threads
30
```

® Protein domain searches

The protein domains in the genome were identified using the Pfam (v33.1) database(<u>Mistry et al. 2021</u>) with *E*-value < 1e-3, using HMMER (v3.1b2; <u>Potter et al. 2018</u>).

```
hmmpress ~/Pfam_db/Pfam.hmm
hmmscan --domtblout Pfam_result.out -E 1e-3 --cpu 20 \
~/Pfam_db/Pfam.hmm Protein_braker.fasta

#hmmscan can't be used because Pfam.hmm files are big data.
#https://www.biostars.org/p/438243/
```

The protein databases of rice, maize, and poplar from the <u>NCBI</u> (last accessed February 2022) were used in the domain for a detailed classification of the PKinase family, the iTAK (v18.12) web tool (<u>Zheng et al. 2016</u>; last accessed February 2022) was utilized. The ratio of families with a high ratio of genes to total genes in the ice plant was compared with that of the same families in the other plants. For statistical analysis, we used Smirnov-Grubbs' outlier tests. The following equation was used to obtain the test statistic T:

T=\frac{(Percentage~of~arbitrary~protein~families~in~the~ice~plant)-(Sample~mean~for~all~nine~species)}{\sqrt{Sample~variance}}

Finally, BLASTP was used to compare proteins generated from the ice plant genome and those from *Arabidopsis*, rice, maize, and poplar and renamed TAIR10 ID. These IDs were subjected to gene ontology (GO) enrichment analysis using DAVID (updated in 2022; accessed on March 24; Sherman et al. 2022) based on a modified Fisher exact probability test with *E*-value < 0.05.