Salt Stress Encourages Proline Accumulation by Regulating Proline Biosynthesis and Degradation in Jerusalem Artichoke Plantlets

Zengrong Huang1,2, Long Zhao1, Dandan Chen1, Mingxiang Liang1*, Zhaopu Liu1*, Hongbo Shao1,2,3*, Xiaohua Long1

1 Jiangsu Key Lab of Marine Biology, College of Resources and Environmental Sciences, Nanjing Agricultural University, Nanjing, China, 2 Key Laboratory of Coastal Biology and Bioresources Utilization, Yantai Institute of Coastal Zone Research (YIC), Chinese Academy of Sciences (CAS), Yantai, China, 3 Institute for Life Sciences, Qingdao University of Science and Technology, Qingdao, China

Abstract
Proline accumulation is an important mechanism for osmotic regulation under salt stress. In this study, we evaluated proline accumulation profiles in roots, stems and leaves of Jerusalem artichoke (Helianthus tuberosus L.) plantlets under NaCl stress. We also examined HpP5CS, HtOAT and HtPDH enzyme activities and gene expression patterns of putative HpP5CS1, HpP5CS2, HtOAT, HtPDH1, and HtPDH2 genes. The objective of our study was to characterize the proline regulation mechanisms of Jerusalem artichoke, a moderately salt tolerant species, under NaCl stress. Jerusalem artichoke plantlets were observed to accumulate proline in roots, stems and leaves during salt stress. HpP5CS enzyme activities were increased under NaCl stress, while HtOAT and HtPDH activities generally repressed. Transcript levels of HpP5CS2 increased while transcript levels of HtOAT, HtPDH1 and HtPDH2 generally decreased in response to NaCl stress. Our results supports that for Jerusalem artichoke, proline synthesis under salt stress is mainly through the Glu pathway, and HpP5CS2 is predominant in this process while HtOAT plays a less important role. Both HtPDH genes may function in proline degradation.

Introduction
Salt stress is a serious abiotic stresses in agricultural production [1]. Soil salinity has been intensified more by global climate change due to the increased temperature in the future. There is the urgent need to identify and use salinity tolerant plants. Since plants can’t move, many have evolved sophisticated mechanisms to adapt to undesirable environments. Osmotic regulation is an important mechanism for plant cellular homeostasis in saline conditions. Under salt stress, plants accumulate several compatible solutes in the cytosol, such as polyols, betaine, trehalose, ectoine, proline and others [2]. As a most common osmolyte for osmoprotection, proline has been extensively researched.

Two proline biosynthetic pathways have been elucidated in higher plants: the glutamate and ornithine pathways [3,4]. In the glutamate (Glu) pathway, glutamate is first catalyzed by P5CS (Pyrroline-5-carboxylate synthetase) to generate GS (Glu) and PSC (Pyrroline-5-carboxylate). Proline is produced from P5C when it is reduced by P5CR (Pyrroline-5-carboxylate reductase). In this pathway, P5CS is identified to be the rate-limiting enzyme [5]. In the ornithine (Orn) pathway, the precursor ornithine is transaminated to P5C by 6-OAT (Orn-threonine transferase), which is considered to be the key enzyme in this pathway. In general, proline biosynthesis is considered to be mainly through Glu pathway under stress conditions while the Orn pathway is involved in seedling development [6]. Proline catabolism is mainly through proline dehydrogenase or oxidase (PDH) and P5C dehydrogenase, which produce P5G and Glu, respectively [7].

Several reports reveal that overexpression of P5CS results in an increased proline level as well as osmotolerance in transgenic plants [8–11]. However, the relationship between proline accumulation and stress tolerance is not always clear. Proline content was not consistent with salt tolerance in barley [12]. In addition, excess accumulation of proline was toxic in pdh knockout mutant plants [13]. There are also indications that AtPDH is induced by proline while repressed by osmotic stress [14]. Therefore, the relationship between proline accumulation and stress tolerance is
unclear for individual plant species. Different plants may use individual strategies to fine-tune gene regulation under stress.

Previously, we demonstrated that proline content was significantly increased under NaCl treatments in the roots of Jerusalem artichoke (*Helianthus tuberosus* L.) [15]. Jerusalem artichoke, a species of sunflower, is considered as moderately salt tolerant [16]. While proline levels in plants are determined by biosynthesis, catabolism, and transport, the proline biosynthesis and degradation pathways are not clear for Jerusalem artichoke. In this paper, we cloned several putative proline-related genes in order to further investigate proline metabolism under salinity stress in Jerusalem artichoke, a multifunctional crop in food or potential bioethanol fuel [17]. Based on proline tissue distribution, enzyme activities and gene expression profiles, the objective of our investigation was to characterize proline metabolism and regulation mechanisms under NaCl stress.

**Materials and Methods**

**Plant Material, Growth Condition and Stress Treatments**

Jerusalem artichoke (Nanyu No.1) was vegetatively propagated as previously described [15]. Briefly, tuber slices with one bud (about 2.0 g) were rinsed thoroughly with deionized water after surface sterilization. The tuber slices were covered with moist acid-washed quartz sand, supplied with 1/2 Hoagland solution and then sprouted for seven days in plastic containers [18]. Plantlets of uniform size (about 20.0 cm) with three fully-extended leaves and one expanding leaf were chosen for salinity treatments after about one month. Salinity treatments included 500 ml 1/2 Hoagland solution plus 50 mM, 100 mM or 200 mM NaCl, individually. Control plantlets received only 500 ml 1/2 Hoagland solution. The plants were grown in the greenhouse with a light intensity of 392–415 µmol m⁻² s⁻¹ during a 12-h light at 25°C and 12-h dark at 18°C. Tissue samples were harvested and immediately dipped in liquid nitrogen and stored at −80°C until use for gene expression analysis. To study relative gene expression in time-course experiments, roots, stems, and leaves were collected at 0, 1, 4 and 12 h under 0 or 100 mM NaCl treatments. For NaCl dosage experiments, samples were supplied with 500 ml 1/2 Hoagland solution including 0, 50, 100 or 200 mM NaCl and harvested 12 h after treatments. Each treatment had three biological replications.

**Proline Content Determination**

Fresh roots, stems, and leaves were collected 0, 12, 24, 48, and 72 h after NaCl treatments (0 or 100 mM). Proline content was determined by ninhydrin assay at λ350 nm according to the method of Bates et al. [19].

**P5CS, OAT and PDH Enzymes Activity Assay**

The above tissue samples for proline content measurements were also used for the following enzyme activity assays.

For the P5CS enzyme activity assay, protein was extracted according to Hayzer et al. [20] with minor modification. Briefly, 1.0 g sample was homogenized in 3.0 ml of ice-cold extraction buffer containing 0.5 M Tris-HCl (pH 7.5), 10 mM MgCl₂, 2 mM PMSF [phenylmethylsulfonyl fluoride], and 2% PVP (polyvinylpyrrolidone). The homogenate was centrifuged at 20,000 g for 20 min (4°C). The supernatant was collected as crude extract. The reaction was initiated by adding 1 ml crude extract into 3 ml reaction medium containing 50 mM Tris-HCl (pH 7.0), 20 mM MgCl₂, 10 mM ATP, 100 mM Hydroxamate-HCl, 50 mM L-glutamate. The reaction was stopped after 30 min at 37°C by adding 3 ml terminating solution containing FeCl₃ (5%), w/v) and trichloroacetic acid (12%, w/v) in 5 M HCl. The protein was removed by centrifugation and the supernatant was recorded at λ350 nm against the tube without ATP as control. The enzyme activity expressed as U/g FW. One U was defined for the enzyme content producing 1 µmol γ-gutamine in 1 min.

The enzyme activity of OAT in each sample was determined by homogenizing 1.0 g plant tissue in 3.0 ml 50 mM ice-cold phosphate buffer (pH 8.0) containing 1 mM DTT (dithiothreitol). The supernatant was collected for the enzyme assay after centrifugation at 7000 g for 20 min (4°C). The enzyme activities were determined by ninhydrin method described by Kim et al. [21]. One U of OAT activity represented 1 nmol P5C produced per hour.

PDH activity was assayed according to Lutts et al. [22]. Plant tissue (1.0 g) was homogenized in 3.0 ml ice-cold extraction buffer containing 50 mM Tris-HCl (pH 7.4), 10 mM β- mercaptoethanol, 7 mM MgCl₂, 5 mM EDTA (ethylene diamine tetraacetic acid), and 0.6 M KCl. The homogenate was centrifuged at 12,000 g for 20 min (4°C) and the supernatant was collected as crude extract. For activity determination, 0.2 ml crude extract was added into reaction mixture buffer containing 0.15 mM NaCO₃-HCl (pH 10.3), 15 mM L-proline and 1.5 mM NAD⁺. A₅40 was recorded at 25°C. The result was expressed as U/g FW. One U represented the reduction value of 0.01 A₅40 per minute.

**RNA Isolation and cDNA Synthesis**

Total RNA was extracted using E.Z.N.A. Plant RNA Kit (OMEGA, R6827-02). In order to remove residual genomic DNA, 1 μg total RNA was treated with RNase-Free DNase Set (OMEGA, E1091-02). PrimeScript RT-PCR kit (Takara, D6110A) was used for cDNA synthesis following the instructions of the manufacturer.

**Cloning cDNA for P5CS, OAT, and PDH**

Gene cloning proceeded according to Liang et al. [23]. Briefly, Jerusalem artichoke ESTs (Expressed Sequence Tags) were downloaded from NCBI website. A database of more than 40,000 retrieved ESTs was created in Bioedit software (Tom Hall, 7.0.9). P5CS, OAT and PDH cDNA sequences from Arabidopsis and rice were used to Localblast this EST database. Assembled by high similarity EST sequences, candidate genes were cloned for further bioinformatic analysis. All five Jerusalem artichoke gene sequences were deposited into the NCBI GeneBank. The accession numbers for *HtP5CS1, HtP5CS2, HtOAT, HtPDH1* and *HtPDH2* in the genebank are KC700610, KC700611, KC700612, KC700613, and KC700614, respectively.

**Bioinformatic Analysis**

The protein sequences of target genes from Genebank were used for sequence alignment analysis by the ClustalW method in the MegAlign program (DNASTAR, Inc., Madison, WI). A phylogenetic tree was generated based on the ClustalW protein alignment analysis using a Neighbor-Joining method in the MEGA 4 program. The following sequences with corresponding accession numbers were used for bioinformatics analysis: *AtP5CS1* (NM_1129539), *AtP5CS2* (NM_1154194), *OpP5CS1* (D49714.1), *OpP5CS2* (NM_001051337), *PpP5CS1* (EU340347), *PpP5CS2* (EU407263), *SbP5CS1* (GG377719), *SbP5CS2* (GQ377720), *AtOAT1* (NM_123987.3), *BoOAT* (EU375566), *GmOAT* (NM_01250221.1), *MsOAT* (AY278819), *NanoAT* (AD474377), *ApiPDH1* (NM_113981.5), *ApiPDH2* (NM_123232.2), *MsPDH* (AY556386), *NipDH1* (AY639145.1), *NipDH2* (AY639146.1).
Quantitative PCR (qPCR) Assay and Data Analysis

Quantitative PCR was performed using SYBR Premix EX Taq (Takara, DRR420A) in the ABI 7500 Real Time PCR System according to the manufacturer’s instructions. The 20 μl reaction system contained 10 μl SYBR Premix Ex Taq, 0.4 μl each of 10 μM primers, 0.4 μl ROX Reference DyeII (50×), 2.0 μl DNA template, and 6.8 μl dd H2O. The program consisted of initial denaturation at 95°C for 30 s, followed by 40 cycles of 95°C for 3 s, annealing for 34 s at 60°C. The dissociation stage includes 95°C for 15 s, 60°C for 1 min and 95°C for 15 s. The Jerusalem artichoke housekeeping gene actin was used for normalization of the amount of cDNA in each qPCR reaction. Since the deviation error of amplification efficiency between target genes and reference gene were less than 10% according to our trial experiments, data were processed using the method of 2^-ΔΔCT according to Livak et al. [24]. For gene transcript level analysis on time-course or salt dosage experiments, gene transcript level in each treatment sample was further normalized to corresponding time-point control sample to avoid circadian rhythm interference. Fold changes were used for statistical analysis.

Primers

Primers for gene cloning and qPCR are listed in Table S1 and Table S2.

Statistical Analysis

Data were analyzed by Microsoft Excel 2003 (Microsoft Corp., Redmond, WA, USA) and SPSS 16.0 (SPSS Corp., Chicago, IL, USA). Each data point shown in the figures is the mean of three biological replications. For each individual sample in real-time PCR experiments, three further technical replications were conducted. Internal significant differences among treatments were statistically evaluated by One-Way ANOVA using Duncan’s t-test. Figures were created using SigmaPlot 10.0 (Systat Software, Inc. Germany).

Results

NaCl Stress Enhanced Proline Accumulation in Roots, Stems and Leaves

To determine the relationship between salt tolerance and proline content in Jerusalem artichoke, we first examined whether salinity stress would enhance the proline levels in different tissues. Our previous experiments indicated that proline level gradually increased with salt concentrations in roots [15]. Since Jerusalem artichoke plants are mostly planted in shoreland or saline land in China, salt concentrations in the soil are usually below 150 mM. As a moderately salt-tolerant species, Jerusalem artichoke can accomplish its life cycle under 100 mM NaCl treatment [25].

Fig. 1 shows a significant increase in proline content could be observed in roots, stems and leaves treated with 100 mM NaCl solution during the initial 72 h period. Proline accumulated in roots, stems and leaves were 28.42, 38.39, and 45.10 μg/g at 72 h, respectively, which were three fold higher than each control. Proline accumulated more in leaf samples than roots or stems at the same time point.

NaCl Stress Increased HtP5CS Activities while Repressed HtOAT and HtPDH Activities

Recent evidence indicates that proline accumulation is mainly initiated by de novo synthesis [26]. To explore the reason why proline was accumulated in different tissues, enzyme activities in proline metabolism pathways were investigated. HtP5CS activity

Figure 1. Effect of NaCl stress on proline accumulation of Jerusalem artichoke in roots (A), stems (B) and leaves (C). 30-day-old Jerusalem artichoke plantlets were treated with 0 or 100 mM NaCl (see Materials and Methods) for 12, 24, 48 and 72 h, respectively. All treatments had three biological replicates. Proline contents were measured by ninhydrin assay at A520 nm. Values represent means ± SE of three independent experiments. Significant differences (P≤0.05) between treatments are indicated by different letters.

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increased in response to NaCl stress in all three tissues. A significant increase of HtP5CS activity could be found in stems of Jerusalem artichoke after 24 h NaCl exposure, while for roots and leaves, significant changes could be found after 48 h stress treatment. Leaf samples showed the highest HtP5CS enzyme activity under 72 h NaCl treatments (Fig. 2). HtP5CS enzyme activities in all tissues were consistent with proline levels which implies that proline biosynthesis mainly originates from the Glu pathway under salt stress conditions.

In contrast to HtP5CS enzyme activities, HtOAT enzyme activities showed the opposite trend. HtOAT activities were generally inhibited under NaCl stress duration from 12 to 72 h. NaCl stress decreased HtOAT activities significantly in stems and leaves of Jerusalem artichoke after 12 h stress treatment. In roots, significant decrease happened after 24 h exposure to stress (Fig. 3).

PDH enzyme plays a key role in proline catabolism. Our results demonstrated that salinity stress gradually decreased the HtPDH activities in roots and stems of Jerusalem artichoke at 24 h while activities stayed constant in leaves except at 72 h (Fig. 4). HtPDH activities were relative lower in the leaves, compared to that of roots and stems regardless of NaCl stress.

It was a little surprising that proline content is increased in all measured tissues while HtP5CS enzyme activities were almost the same in all treatments of NaCl at 12 h. We hypothesized that our enzyme activity measurement method may not be responsive enough to detect rapid changes as a result of external stimuli. To provide further information about proline metabolism in Jerusalem artichoke, several possible proline-metabolism genes were then cloned.

Bioinformatics Analysis of Five Proline-metabolism Genes

In contrast to yeast and animal system, plant genes always form a multimember gene family. Family members may play redundant roles or have function differentiations. According to published Arabidopsis and rice information, we postulated at least two P5CS and PDH isoforms in Jerusalem artichoke. Gene information of $\alpha$-OATs were reported in bacterium, while only $\delta$-OATs were found in higher plants [27], therefore we only studied $\delta$-OAT in this paper.

Using Arabidopsis and rice P5CS nucleotide sequences to Blastn the Jerusalem artichoke EST database, twelve ESTs with high similarities were retrieved, that is,125441546, 125423128, 125430151, 125443641, 125446643, 125448887, 125449956, 12550355, 125439651, 125439845 and 125442867. Assembled by SeqManTMII (DNASTAR), two contigs were formed. The first Contig was 2516 bp length and the deduced protein showed high identity to reported P5CS sequences. Primers were designed to amplify contig 1 (using primer 1, Table S1), and the results revealed a 2496 bp cDNA harboring a full-length putative P5CS (2154 bp) based on the alignment with other P5CS sequences. Contig 2 only had partial CDS sequences, so efforts were made to search against other Compositae crop's EST database, such as sunflower. It was found that 58736240 in sunflower EST database showed high similarity with the front part of contig 2 (http://compbio.dfci.harvard.edu/cgi-bin/tgi/gimain.pl?gdb=arundinacea), and 113187185 from silverleaf sunflower (Helianthus argophyllus L.) EST database hits to the back region, so these two sequences were then assembled to form contig 2 to design the primer pair. Sequenced after PCR amplification (using primer 2, Table S1), a 2257 bp sequence was obtained without full-length CDS. A 3'-end RACE PCR (Clontech, Cat.No. 634923, USA) was carried out to extend the 3'-end (using primer 3, Table S1). By using this method, contig 2 was

Figure 2. Effect of NaCl stress on P5CS activities of Jerusalem artichoke in roots (A), stems (B) and leaves (C). 30-day-old Jerusalem artichoke plantlets were treated with 0 or 100 mM NaCl (see Materials and Methods) for 12, 24, 48 and 72 h. All treatments had three biological replicates. P5CS activities were measured by hydroxylamine hydrochloride assay at A$_{535}$ nm (see Materials and Methods). Values represent means ± SE of three independent experiments. Significant differences (P≤0.05) between treatments are indicated by different letters. doi:10.1371/journal.pone.0062085.g002
Figure 3. Effect of NaCl stress on α-OAT activities of Jerusalem artichoke in roots (A), stems (B) and leaves (C). 30-day-old Jerusalem artichoke plantlets were treated with 0 or 100 mM NaCl (see Materials and Methods) for 12, 24, 48 and 72 h. All treatments had three biological replicates. OAT activities were measured by ninhydrin method (see Materials and Methods). Values represent means ± SE of three independent experiments. Significant differences (P<0.05) between treatments are indicated by different letters. doi:10.1371/journal.pone.0062085.g003

Figure 4. Effect of NaCl stress on PDH activity of Jerusalem artichoke in roots (A), stems (B) and leaves (C). 30-day-old Jerusalem artichoke plantlets were treated with 0 or 100 mM NaCl (see Materials and Methods) for 12, 24, 48 and 72 h. All treatments had three biological replicates. PDH activities were measured by spectrophotometry at A_{340} at 25°C (see Materials and Methods). Values represent means ± SE of three independent experiments. Significant differences (P<0.05) between treatments are indicated by different letters. doi:10.1371/journal.pone.0062085.g004
successfully prolonged and a full-length sequence of 2178 bp CDS was amplified (using primer 4, Table S1). According to the similarity with other P5CSs, contig 1 was named as HtP5CS1, whose deduced amino sequence showed highest similarity with PpP5CS1 (79.1%), and contig 2 was named as HtP5CS2, with 72.9% identity with PvP5CS2.

Using HtOAT and BnOAT CDS sequences to Blastn Jerusalem artichoke EST database, we found nine ESTs with high similarities, that is, 125423045, 125426820, 125427805, 125431428, 125437709, 125439430, 125440189, 125443071 and 125450077. Those ESTs were then assembled into a 1265 bp template. Using Ht cDNA as template, a putative HtOAT with a product of 1146 bp length was amplified (using primer 4, Table S1), however, without a complete CDS. The sequence of this putative HtOAT was compared with other composites plants. We retrieved 211619016 from the sunflower EST database, 113237927 from the serpenpine sunflower (Helianthus exilis L.) EST database, and 90497972 from the plains sunflower (Helianthus petiolaris L.) EST database. The retrieved sequences were used for assembling the putative partial CDS into a 1681 bp sequence. Sequenced after PCR (using primer 4, Table S1), a sequence including a full-length sequence of 1410 bp was obtained, whose deduced amino acid sequences shared 77.2% similarity with NiOAT.

Using BnPDH to Blastn against Ht EST database, we identified thirteen ESTs with high similarities and assembled these into two contigs (125424918, 125425174, 125425617, 125427731, 125428116, 125437754, 125456916, 125457059, 125422940, 125432267, 125434747, 125459104 and 125450265). Contig 1 had an 867 bp length was used for primer design. Using Ht cDNA as template, a 1618 bp product was amplified (using primer 7, Table S1), which was found to include a full-length CDS (1497 bp). Contig1 had 57.7% similarity with NtPDH1 and was named as HtPDH1. For contig 2, eight sunflower ESTs were sequenced after PCR (using primer 8, Table S1), which was found to include a full-length sequence of 1410 bp was obtained, whose deduced amino acid sequences shared 77.7% similarity with NtPDH.

Expression Profiles of Five Proline-metabolism Genes under Salt Treatments

To link proline-metabolism genes with physiological processes, transcript levels of each Jerusalem artichoke gene were studied in different tissues or at different developmental stages or stress conditions. Relative transcript levels of each gene in different tissues and at different developmental stages were quantified and compared with the level of control roots (arbitrary selection) to demonstrate the relative transcript levels of the same gene.

Gene expression of HtP5CS1 only changed significantly in roots after 4 h treatment. Transcript levels of HtP5CS1 were increased in stems after 1 h exposed to 100 mM NaCl stress and decreased after that. However, the expression level of HtP5CS1 in the leaves was almost the same as the control. Generally speaking, expression pattern of HtP5CS2 showed a similar trend with proline accumulation profiles in all three tissues examined. Transcript level of HtP5CS2 was initially induced in roots and leaves after 4 h treatment and reached the highest level at 12 h. In stems, expression levels of HtP5CS2 were only increased at 12 h (Fig. 6).

NaCl stress mildly inhibited HtOAT expression in roots within the 12 h experiment period while in leaves a significant decrease appeared only after 12 h exposed to 100 mM NaCl treatment. In stems, the expression level of HtOAT was raised after 1 h stress treatment, however, expression decreased to the control level after that (Fig. 6).

Discussion

NaCl Stress Promoted Proline Accumulation by Activating Proline Synthesis and Inhibiting Proline Catabolism Simultaneously

Previously, Newton et al. [25] reported that Jerusalem artichoke was a moderately salt tolerant crop when EC2 was 7.5 dS m−1 in soil. In the present study, we simulated a series of NaCl-induced stresses ranging from 3.48 dS m−2 (30 mM NaCl+1/2 Hoagland) to 9.82 dS m−2 (200 mM NaCl+1/2 Hoagland). Our results indicated that proline content was enhanced in Jerusalem artichoke roots, stems and leaves based on time-course experiments under 100 mM NaCl stress (EC2 = 6.26 dS m−2) (Fig. 1). The same finding was reported in canola [33] and rice callus tissue [9]. By comparing different cultivars of sunflower in response to salt stress, Shahbaz et al. [37] also found that salt tolerant cultivars exhibited greater proline accumulation. Proline content was observed higher in leaves in this study, which was same with previous study in green gram [34], mulberry [35] and canola [36]. It has been suggested that leaves accumulate more proline in order to maintain chlorophyll level and cell turgor to protect photosynthetic activity under salt stress [38]. This is consistent with our results indicating that proline accumulation was greatest in the leaves of stressed plants.

Tissue accumulations of proline under stress may result mainly from de-novo biosynthesis since proline content was increased in all three tissues compared to the control, similar to previous results [39]. To explore the mechanism behind this, we measured the rate-limiting enzyme activities in the two primary proline synthetic pathways respectively. Our results revealed that HtP5CS activities were unambiguously stimulated while HtOAT activities were
slightly repressed after the plantlets were exposed to NaCl treatments (Fig 2, Fig. 3). In general, we found that tissues that had higher HtP5CS enzyme activities accumulated more proline. In our study, HtOAT enzyme activities were lower in all three tissues under salt treatment. However, Wang et al [40] claimed that in *S. amara*, a typical halophyte, δ-OAT activity was also enhanced, contributing to proline synthesis. We inferred that proline synthesis under stress is likely species-dependent. For Jerusalem artichoke, the Glu pathway was predominant while the Orn pathway played a minor role, if any, under our stress conditions. Our findings were also consistent with Delauney et al [41].

Interestingly, PDH activity was repressed under NaCl stress (Fig. 4), which had been illustrated previously in rice [42]. Moreover, our results showed that PDH activity was higher in roots than in leaves, which was also accordance with the study in rice [43]. Higher P5CS enzyme activity and lower PDH activity may both contribute to the highest proline content in leaves. These results indicate that proline accumulates in Jerusalem artichoke plantlets under NaCl stress by activation of proline synthesis through the glu pathway, while at the same time, repression of degradation.

Figure 5. Phylogenetic analysis of HtP5CS, HtOAT and HtPDH. Protein sequences which had been reported were selected for sequence alignment. Sequence alignment analysis was performed using the ClustalW method in the Megalign program (DNASTAR, Inc., Madison, WI, USA). These genes’ accession numbers are: AtP5CS1 (NM_129539) and AtP5CS2 (NM_115419.4) in Arabidopsis; OsP5CS1 (D49714.1) and OsP5CS2 (NM_001051337) in rice (*Oryza sativa*); PvP5CS1 (EU340347), PvP5CS2 (EU407263) in common bean (*Phaseolus vulgaris*); SbP5CS1(GQ377719) and SbP5CS2(GQ377720) in sorghum (*Sorghum bicolor*); AtOAT (NM_123987.3) in Arabidopsis; BnOAT (EU375566.1) in rapeseed (*Brassica Napus*); GmOAT (NM_001250221.1) in soybean (*Glycine max*); MtOAT (AJ278819) in alfalfa (*Medicago truncatula*); NtOAT (ADM47437) in tobacco (*Nicotiana tabacum*); AtPDH1 (NM_113981.5) and AtPDH2 (NM_123232.2) in Arabidopsis; MsPDH (AY556386.1) in alfalfa (*Medicago sativa*); NtPDH1 (AY639145.1) and NtPDH2 (AY639146.1) in tobacco (*Nicotiana tabacum*).

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Sequences Analysis of Five Putative Proline-metabolism Genes in Jerusalem Artichoke Revealed High Similarity to Identified Genes

Since at least two P5CS genes, one δ-OAT and two PDH genes were already characterized in the plants, to better illustrate the proline metabolism in Jerusalem artichoke, we cloned several putative proline metabolism genes through EST alignment according to the reported nucleotide sequences. Deduced amino sequences of the five cloned genes all showed high similarities to the reported genes. According to phylogenetic analysis, the putative HtP5CS1 protein shared the highest similarity with PvP5CS1 (79.1%), and HtP5CS2 showed 73.7% similarity with PvP5CS1. HtOAT showed the highest homology with NtOAT (77.2%). Two HtPDHs were grouped with NtPDHs. The highest similarity of HtPDH1 could be found with NtPDH2 (57.9%), and HtPDH2 shared 58.4% homolog with NtPDH2. By comparisons of function-identified protein sequences, the deduced protein sequences showed that both HtP5CS1 and HtP5CS2 had a conserved AA_Kinase and Aldedh domains and an aminotran_3 was found in HtOAT (http://pfam.sanger.ac.uk/). For HtPDH1 and HtPDH2, a pro_dh domain exists. Via alignment with other species, all five genes have conserved domains, such as ATP-binding domain in P5CS genes, pyridoxal phosphate-binding in OAT gene and a proline dehydrogenase domain in the PDH gene. Sequences analysis indicated that our cloned genes could be the actual proline-metabolism genes.

Transcript Levels of HtP5CS2 were Increased while Expression Levels of HtOAT, HtPDH1 and HtPDH2 were Generally Decreased in Response to NaCl Stress

Since enzyme activity measurement methods could not promptly reflect an immediate change, we examined expression patterns in five genes in the proline metabolic pathway. Our results revealed that the induction of HtP5CS2 was initiated within 12 h by NaCl stress in different tissues. Also, the same trend was found when the plantlets were exposed to different NaCl dosages in 12-h treatments (Fig. S4). In our study, NaCl stress did not repress relative expression of HtP5CS1 (Fig. 6), however, the change of gene expression level was much less than that of HtP5CS2. In cactus pear (Opuntia streptacantha L.), Silva-Ortega et al. [38] found that the transcript level of OsP5CS (accession number:D49714.1, higher similarity to HtP5CS1) was increased with NaCl concentrations at the 9th day after treatment. Szekely et al [44] reported that knock-out Arabidopsis p5cs1 mutant plants were hypersensitive to salinity, whereas AtP5CS2 was essential for seedling development. The results presented in the study indicated that for Jerusalem artichoke, both P5CS1 and P5CS2 in Glu pathway may function synergistically. HtP5CS2 seemed to play predominant roles under salt stress based on its constantly increasing expression. Nevertheless, our results were not completely in line with Armengaud et al. [6], who indicated that...
which was consistent with the expression profile of in Fig. 4, we found that PDH activities were repressed over time, expression levels of relative expression levels of control in all salt treatments (Fig. S4). Compared to control, no significant change could be found in stems compared to the PDH1 had the greatest effect on the stress (Fig. 6 and Fig. S4). Moreover, 50 mM NaCl stress down-clarify Orn roles under salt stress. Based on the previous finding, further research should be carried out in plant developmental stage based on metabolic network to out in plant developmental stage based on metabolic network to clarify Orn roles under salt stress.

PDH is responsible for catabolizing proline into P5C. As shown in Fig. 4, we found that PDH activities were repressed over time, which was consistent with the expression profile of HtPDH2 under stress (Fig. 6 and Fig. S4). Moreover, 50 mM NaCl stress down-regulated the expression of HtPDH1 in stems. A stress of 200 mM had the greatest effect on the PDH1 transcript level in roots, while no significant change could be found in stems compared to the control in all salt treatments (Fig. S4). Compared to control, relative expression levels of HtPDH2 were significantly inhibited under different NaCl stresses in roots, stems and leaves at 12 h. Expression levels of HtPDH2 in leaves were gradually repressed with higher salt concentrations (Fig. S4). The same trend was found in Medicago sativa while Miller et al. [47] reported that OAT expression varied with growth period in Arabidopsis under salt stress, whereas the predominant role of the Glu pathway for proline synthesis was identified in adult plants. Based on the previous finding, further research should be carried out in plant developmental stage based on metabolic network to clarify Orn roles under salt stress.

Conclusions
Our experiments indicate that proline is induced in Jerusalem artichoke plantlets under salinity stress. More proline is accumulated in leaves as a possible protective metabolic adaptation to prevent leaf tissue from damage under salinity. Proline accumulation is primarily caused by activation of proline synthesis via the glu pathway, while depression of proline degradation also serves to increase proline levels. Using bioinformatic methods to analyze Jerusalem artichoke ESTs, we cloned five putative proline metabolically-related genes, whose translated amino acid sequences exhibit high similarities to published sequences. Under salinity, proline synthesis activated by HtP5CS2 may be predominant, while for proline degradation, both HtPDHs may have function.

This study supports the hypothesis that proline plays an important role in Jerusalem artichoke osmoprotection. Although extensive work regarding proline metabolism under salt stress has been done, it still remains to be investigated how these five genes in proline metabolism are regulated and how each member in these same gene families plays different roles under stress.

Supporting Information

Figure S1 Amino acid sequence alignments of HtP5CS1 and HtP5CS2. Alignment was performed using the ClustalW program. The following sequences with corresponding accession number were used for bioinformatic analysis: AIP5CS1 (NM_129539), AIP5CS2 (NM_1154194); OIP5CS1 (D49714.1), OIP5CS2 (NM_001051337); PtP5CS1 (EU340347), PtP5CS2 (EU407263); SbP5CS1 (GQ377719), SbP5CS2 (GQ377720). Asterisk, semicolon and dot represent the amino that are “identical”, “conserved substitution” and “semi-conserved substitution”, respectively. Boxed sequences showed conserved putative ATP and NAD/PH-binding sites, GK and GSA-DH domains, and putative Leu-rich regions. (TIF)

Figure S2 Amino acid sequence alignments of HtOAT. Alignment was performed using the ClustalW program. The following sequences with corresponding accession number were used for bioinformatic analysis: AtOAT (NM_123987.3); BoOAT (EU375566); GmOAT (NM_001250221.1); MtOAT (AJ278819); NtOAT (ADM47437). Asterisk, semicolon and dot represent the amino that are “identical”, “conserved substitution” and “semi-conserved substitution”, respectively. Boxed sequences showed putative pyridoxal phosphate-binding domain. (TIF)

Figure S3 Amino acid sequence alignments of HtPDH1 and HtPDH2. Alignment was performed using the ClustalW program. The following sequences with corresponding accession number were used for bioinformatic analysis: AIPDH1 (NM_113981.5), AIPDH2 (NM_123232.2); MtPDH (AY565386.1); MPDH1 (AY639145.1), MPDH2 (AY639146.1). Asterisk, semicolon and dot represent the amino that are “identical”, “conserved substitution” and “semi-conserved substitution”, respectively. Boxed sequences showed dihydrogenase domain. (TIF)

Figure S4 Expression profiles of HtP5CS1, HtP5CS2, HtOAT, HtPDH1, and HtPDH2 from roots, stems, and leaves under different NaCl dosage stresses at 12 h. 30-day-old Jerusalem artichoke plantlets were treated with 0, 50, 100 and 200 mM NaCl (see Materials and Methods) for 12 h, respectively. All treatments had three biological replicates. Total RNA was extracted from roots, stems and leaves for quantitative PCR (qPCR) analysis. Transcript levels were first normalized to the level of a control gene actin (Hkt104). The normalized transcript levels were then compared between treatments (50, 100 and 200 mM NaCl) and control (0 mM NaCl) to obtain a fold change. Values represent means ± SE of three independent experiments. Significant differences (P<0.05) between treatments are indicated by different letters. Values represent means ± SE of three independent experiments. Significant differences (P≤0.05) between treatments are indicated by different letters. (TIF)

Table S1 Primers for gene cloning of HtP5CS1, HtP5CS2, HtOAT, HtPDH1, and HtPDH2. (DOC)
Author Contributions

Revised, enriched, and submitted the manuscript: HS. Conceived and designed the experiments: ML, ZH, XL. Performed the experiments: ZH, LZ, DC. Analyzed the data: ZH, ML, HS. Contributed reagents/materials/analysis tools: ZL, XL. Wrote the paper: ZH, ML, HS.

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