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Proteomics of *Arabidopsis* leaves subjected to low or high temperature stress.

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Proteomic analysis of temperature stress-responsive proteins in *Arabidopsis thaliana* rosette leaves

Mariapina Rocco,^a Simona Arena,^b Giovanni Renzone,^b Stefania Scippa,^c Tonia Lomaglio,^c Francesca Verrillo,^a Andrea Scalonib and Mauro Marra*^d

Plants, as sessile organisms, are continuously exposed to temperature changes in the environment. Low and high temperature stresses have a great impact on agricultural productivity, since they significantly alter plant metabolism and physiology. Plant response to temperature stress is a quantitative character, being influenced by the degree of stress, time of exposure, as well as plant adaptation ability; it involves profound cellular changes at the proteomic level. We describe here the quantitative variations of the protein repertoire of *Arabidopsis thaliana* rosette leaves after exposing seedlings to either short-term cold or heat temperature stress. A proteomic approach, based on two-dimensional electrophoresis and MALDI-TOF peptide mass fingerprinting and/or nanoLC-ESI-LIT-MS/MS experiments, was used for this purpose. The comparison of the resulting proteomic maps highlighted proteins showing quantitative variations induced by temperature treatments. Thirty-eight protein spots exhibited significant quantitative changes under at least one stress condition. Identified, differentially-represented proteins belong to two main broad functional groups, namely energy production/carbon metabolism and response to abiotic and oxidative stress. The role of the identified proteins is discussed here in relation to plant adaptation to cold or heat stresses. Our results suggest a significant overlapping of the responses to opposite temperature extremes.

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1 Introduction

Plants, being in intimate contact with the environment, are continuously challenged by unfavourable conditions, such as excess/lack of water, light, nutrients or temperature extremes, all of which severely impair plant growth and reduce crop yields.¹ In order to cope with unfavourable environmental constraints, plants have evolved molecular mechanisms involving profound changes in gene expression to bring about metabolic adaptation improving fitness under stress conditions.² Increasing evidence from functional genomic studies suggests that information from transcriptomic data does not match necessarily the actual cellular protein complement,³ since different modifications may affect gene products, including post-transcriptional, co-translational and degradative ones.

Hence, integration of genomic data with characterization of actual protein effectors of plant stress response is necessary for a better understanding of the underlying molecular mechanisms.

Low and high temperature extremes are among the most common stresses that hamper plant growth and limit crop productivity. Response of plants to low temperatures is modulated by environmental parameters like duration and intensity of the challenge. In many plants, while a gradual exposure to moderately low temperatures for a sufficiently long time brings about an adaptation response known as acclimation,⁴ which involves an extensive genetic and metabolic reprogramming, short time exposure to a more severe cold challenge triggers a rapid response whose biochemical features are only partially overlapping with those of acclimation. Molecular mechanisms underlying cold acclimation have been studied at the gene level in *Arabidopsis*,⁵ where key regulators, such as the transcriptional cascade ICE1-CBF⁶ and HOS9 factor,⁷ have been identified. In recent years, dedicated proteomic studies have been realized on *Arabidopsis*,^{8–11} rice^{12,13} and other species,¹⁴ including the chilling-tolerant *Thellungiella halofila*,¹⁵ they identified a number of cold-responsive elements, among which some stress-responsive proteins and components involved in fundamental cellular processes, such as photosynthesis, carbon

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1 assimilation, sucrose synthesis, RNA processing, ROS detoxifi- 1
cation and protein folding.¹⁴ These studies have been per-
formed both at the whole and subproteome levels,^{8,16} and
5 were mostly related to long-lasting cold treatments and to the
acclimation response. Conversely, plant response to fast tem- 5
perature drop has much less been investigated. Accordingly,
proteomic characterization of the molecular mechanisms
underlying fast cold changes is still fragmentary, relying on a
10 limited number of papers related to various species, such as
Arabidopsis, rice or *Thellungiella*, under different experimental
conditions;^{8,11,15} for a review see ref. [15]. It is known that short
term chilling induces rapid structural and metabolic changes,
like alteration of membrane composition or synthesis of stress-
protecting metabolites and/or proteins, processes that can be
15 predominantly regulated at the post-transcriptional level.¹¹

Heat stress has a severe impact on crop productivity, and the
gradual increase in the ambient temperature observed in the
last few decades enhances its economic impact on a global
scale. Heat has an adverse effect on both vegetative growth and
20 reproduction of plants. At the molecular level, temperature rise
can have either structural effects, altering protein, membrane
and cytoskeleton stability, or metabolic ones, due to alteration
of enzyme activities, which in turn result in accumulation of
toxic metabolites, *i.e.* reactive oxygen species (ROS).¹⁷ Two types
25 of heat stress response can be distinguished in plants: (i) basal
thermotolerance, which is the response to a sharp temperature
increase (42–45 °C) for a short time (2–6 h); (ii) acquired
thermotolerance, in which plants are first exposed to moder-
ately high temperatures or to a gradual increase, allowing them
30 to acclimate to subsequent more severe heating.¹⁷ Experi-
mental evidence suggests that molecular players of the two
responses are only partially shared.¹⁸ In fact, it has been
reported that some classes of chaperones, *e.g.* sHSPs, HSP70s
or APX, accumulate preferentially in the acquired thermotoler-
ance,¹⁷ whereas the transcription factor MBF1c and catalase are
35 key effectors in thermotolerance.¹⁹ Proteomic studies concern-
ing plant response to heat stress have been conducted both on
model species, *i.e.* *Arabidopsis thaliana*,^{20,21} as well as on crop
species, *e.g.* rice,^{20,22,23} wheat and barley.²⁰ Globally, these
40 studies indicated that HSPs, as involved in protein stability,
play a major role in thermotolerance; other differentially-repre-
sented components acting in redox homeostasis, carbohydrate
metabolism and protein synthesis/degradation seem to be also
involved in the plant response.^{20–23}

45 In this work, we report on the main quantitative changes of
the protein repertoire of *Arabidopsis thaliana* rosette leaves after
exposing seedlings to short-term (6 h), low (4 °C) or high (42 °C)
temperature extremes. Proteins were resolved by two-dimen-
sional electrophoresis (2-DE) and [15] labeling proteomic patterns
50 were compared to control. Differentially represented components
were identified by means of combined MS experiments. A
significant overlap between the response to short-term cold
and heat shock was originally ascertained. Functional proper-
ties of identified protein species are here discussed according
55 to available literature data concerning response of plant to
different kinds of temperature stresses.

2 Materials and methods 1

2.1 Plant growth and stress treatments 5

Arabidopsis thaliana, ecotype Columbia (Col-0), seedlings were
grown in a growth chamber at 22 °C, 80% humidity, under a
16 h light/8 h dark cycle.²⁴ Three-week old plants were sub-
jected to cold or heat stress treatments by incubating them in
the dark, at 4 °C or 42 °C, respectively, for 6 and 12 h. Control
plants were incubated at 21 °C for 6 and 12 h, in the dark.
Leaves from stressed and control seedlings were then sampled
10 and immediately subjected to further assays, as described
below.

2.2 Relative electrolyte leakage measure 15

The relative electrolyte leakage (REL) assay was performed
under the conditions described by Yan *et al.*;²⁵ six biological
replicates for each samples were used.

2.3 Protein extraction 20

Protein mining was performed according to the phenol extrac-
tion method,²⁶ with minor modifications. Briefly, plant leaves
were finely powdered in liquid N₂ using a mortar and dried
under vacuum. One gram of dried leaves was suspended in
20 mL of ice-cold extraction buffer (700 mM sucrose, 500 mM
Tris-HCl, pH 7.5, 50 mM EDTA, 100 mM KCl, 2% w/v
β-mercaptoethanol, 1 mM PMSF, 1% w/v PVP, 0.25% w/v
25 CHAPS, 40 mM NaF, 1 μM okadaic acid). After addition of an
equal volume of phenol saturated-500 mM Tris-HCl, pH 7.5, the
mixture was stirred for 5 min in a Waring blender and then
centrifuged at 10 000 × *g* for 10 min, at 4 °C. The upper phenol
30 phase was removed and extracted once again with the extrac-
tion buffer. Proteins were precipitated from the phenol phase
by addition of 5 vol of saturated ammonium acetate in methan-
ol, at –20 °C, overnight. Precipitated proteins were centri-
fuged at 10 000 × *g*, for 30 min. Proteins were stored at –80 °C,
35 until used. Three biological replicates for each treatment were
analyzed, which were then subjected to an independent phenol
extraction and subsequent proteomic analysis.

2.4 2D electrophoresis and gel image acquisition 40

Protein pellets were dissolved in IEF buffer (9 M urea, 4% w/v
CHAPS, 0.5% v/v Triton X-100, 20 mM DTT, 1% w/v Bio-Rad
carrier ampholytes pH 3–10). Protein concentration was esti-
mated using the Bradford assay, modified according to Ramagli
and Rodriguez.²⁷ IPG strips (17 cm, pH 4–7, Bio-Rad Ready-
Strip, Bio-Rad) were rehydrated overnight with 300 μL of IEF
45 buffer containing 300 μg of total proteins. Proteins were
focused using a Protean IEF Cell (Bio-Rad) at 12 °C, by applying
the following voltages: 250 V (90 min), 500 V (90 min), 1000 V
(180 min) and 8000 V for a total of 52 KVh.²⁶ After focusing,
50 the proteins were reduced by incubating the IPG strips with 1% w/v
DTT in 10 mL of equilibration buffer (50 mM Tris-HCl, pH 8.8,
6 M urea, 30% w/v glycerol, 2% w/v SDS and a dash of
bromophenol blue) for 15 min, and then alkylated with 2.5%
55 w/v iodoacetamide in 10 mL of equilibration buffer, for 15 min.
Electrophoresis in the second dimension was carried out on

1 12% polyacrylamide gels (180 × 240 × 1 mm) using a Protean
apparatus (Bio-Rad), using electrophoresis buffer (25 mM Tris-HCl,
pH 8.3, 1.92 M glycine and 1% w/v SDS), with 120 V applied for
12 h, until the dye front reached the bottom of the gel. 2-DE gels
5 were then stained with colloidal Coomassie G250; resulting
images were acquired using a GS-800 imaging system (Bio-Rad).
For quantitative analysis, each sample was analyzed in technical
triplicate.

10 2.5 Gel image analysis

Digitized images of Coomassie-stained gels were analyzed
using the PD Quest (ver 7.4) 2-D analysis software (Bio-Rad),
which allowed spot detection, landmarks identification, align-
ing/matching of spots within gels, quantification of matched
15 spots and their analysis, according to manufacturer's instruc-
tions. Manual inspection of the spots was performed to verify
the accuracy of automatic gel matching; any errors in the
automatic procedure were manually corrected prior to final
data analysis. The spot volume was used as the analysis para-
20 meter for quantifying protein expression. The protein spot
volume was normalized to the spot volume of the entire gel
(*i.e.*, of all the protein spots). Fold-changes in protein spot
levels were calculated between spot volumes in the treated
group relative to that in the control one. Statistically significant
25 changes in protein expression were determined using two
sequential data analysis criteria. First, a protein spot has to
be present in all gels for each sample to be included in the
analysis. Next, statistically significant changes in protein
expression were determined using the distribution of fold-
30 change values in the data. Spots were determined to be statistically
significant (by Student's *t* test) if the difference between
the average intensity of a specific protein spot in the control
and treated plants (three technical replicates of three biological
35 samples) was greater than one standard deviation of the spot
intensities for both groups. An absolute two-fold change in
normalized spot densities was considered indicative of a differ-
entially modified protein; thus, values ≥ 2 or ≤ 0.5 were
associated with increased or decreased protein amounts after
40 treatment, respectively.

40 2.6 Protein digestion and MS analysis

Spots from 2-DE were manually excised from gels, triturated
and washed with water. Proteins were *in-gel* reduced,
S-alkylated and digested with trypsin, as previously reported.²⁸

45 Protein digests were subjected to a desalting/concentration step
on microZipTipC18 pipette tips (Millipore Corp., Bedford, MA,
USA) before MALDI-TOF-MS peptide mass fingerprinting (PMF)
and/or nanoLC-ESI-LIT-MS/MS analysis.

50 During MALDI-TOF PMF experiments, peptide mixtures
were loaded on the instrument target together with α -cyano-4-
hydroxycinnamic acid as a matrix, using the dried droplet
technique. Samples were analysed using a Voyager-DE PRO
mass spectrometer (Applied Biosystems, USA). Peptide mass
spectra were acquired in reflectron mode; internal mass cali-
55 bration was performed with peptides derived from trypsin
autoproteolysis. Data were elaborated using the DataExplorer

5.1 software (Applied Biosystems). PSD fragment ion spectral
analysis of the most abundant mass signal within each
MALDI-TOF-MS spectrum was performed as previously
reported.²⁹ Peptide mixtures were eventually analyzed by
5 nanoLC-ESI-LIT-MS/MS using a LTQ XL mass spectrometer
(ThermoFinnigan, USA) equipped with a Proxeon nanospray
source connected to an Easy-nanoLC (Proxeon, Denmark).³⁰
Peptide mixtures were separated on an Easy C₁₈ column (100 ×
0.075 mm, 3 μ m) (Proxeon) using a gradient of acetonitrile
10 containing 0.1% formic acid in aqueous 0.1% formic acid;
acetonitrile ramped from 5% to 35% over 15 min and from
35% to 95% over 2 min, at a flow rate of 300 nL min⁻¹. Spectra
were acquired in the range *m/z* 400–2000. Acquisition was
controlled by a data-dependent product ion scanning proce-
15 dure over the three most abundant ions, enabling dynamic
exclusion (repeat count 2 and exclusion duration 1 min). The
mass isolation window and collision energy were set to *m/z* 3
and 35%, respectively.

20 2.7 Protein identification

MASCOT software package version 2.2.06 (Matrix Science, UK)³¹
was used to identify spots unambiguously from an updated
plant non-redundant sequence database (UniProt 2009/05/03).
MALDI-TOF PMF data were searched using a mass tolerance
value of 40–80 ppm, trypsin as a proteolytic enzyme, a missed
25 cleavage maximum value of 2 and Cys carbamidomethylation
and Met oxidation as fixed and variable modification, respec-
tively. NanoLC-ESI-LIT-MS/MS data were searched using a mass
tolerance value of 2 Da for precursor ions and 0.8 Da for MS/MS
fragments, trypsin as a proteolytic enzyme, a missed cleavage
30 maximum value of 2 and Cys carbamidomethylation and Met
oxidation as fixed and variable modification, respectively.
MALDI-TOF PMF candidates with a cumulative MASCOT score
> 83, which were also confirmed by PSD data (data not shown),
or nanoLC-ESI-LIT-MS/MS candidates with at least 2 assigned
35 peptides with an individual MASCOT score > 25, both cor-
responding to *p* < 0.05 for a significant identification, were
further evaluated by the comparison with their calculated *M_r*
and *pI* values, using the experimental ones obtained from 2-DE.

3 Results and discussion

3.1 Effect of temperature stresses on cell membranes

45 Since cold and heat short-term treatments did not result in any
evident alteration of plant growth or phenotype, we evaluated
the increase of ion permeability of cell membranes, which is
considered as an indicator of damage induced by different
stresses including cold¹⁵ and heat²³ ones. In particular, we
measured the relative electrolyte leakage of *Arabidopsis* rosette
50 leaves incubated at 4 °C or at 42 °C, for 6 and 12 h. After 6 h,
both temperature treatments altered cell membrane ion perme-
ability, as witnessed by the slight increase of REL we observed
over control values, which further raised after 12 h (Fig. 1).
These results indicated that opposite temperature challenges
55 for the same periods of time effectively determined stress
conditions in *Arabidopsis* leaves, thereby eliciting cellular

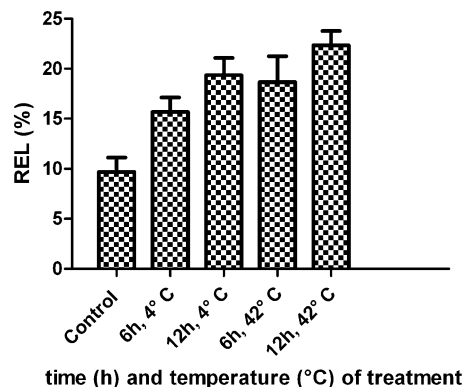


Fig. 1 Relative electrolyte leakage (REL) from rosette leaves of *Arabidopsis thaliana*. Three-week-old seedlings were treated at 4 °C or 42 °C, for 6 or 12 h, and then REL was measured. Mean values \pm SD from six replicates are reported. Control refers to plants incubated at 21 °C for 6 h, in the dark; identical data were obtained for control plants incubated at 21 °C for 12 h, in the dark.

response mechanisms suitable to be investigated by proteomic analysis. Our results were fairly in accordance with previous reports on other plant species, which indicated a relatively low level of membrane damage in response to short-term cold^{12,15} or heat challenge.^{21–23} Similarly to these studies, a higher sensitivity of *Arabidopsis* cell membranes to short-term heat than cold stress was observed; this phenomenon was associated with the capacity of heat stress to induce ROS production more rapidly than the cold one, subsequently determining a higher lipid peroxidation and final damage to membranes.²³

3.2 Proteomic analysis of *Arabidopsis* leaf proteins under cold and heat shock

A proteomic approach was then used in order to identify proteins whose abundance changed upon short term heat or cold stress of *Arabidopsis thaliana* rosette leaves. Total proteins were extracted from leaves of plants incubated at 4 °C (cold shock) or 42 °C (heat shock) for 6 h, and the corresponding control, and then resolved by 2D electrophoresis, within the *pI* range 4–7 and mass range 10–150 kDa; representative 2D gels are shown in Fig. 2. To ascertain quantitative changes in relative spot volumes for temperature-treated leaves compared to control, colloidal Coomassie-stained gels were subjected to comparative software-assisted image analysis. Average proteomic maps showed 296 ± 25 (cold stress), 302 ± 31 (heat stress) and 267 ± 28 (control) spots. Statistical evaluation of relative spot volumes allowed us to detect spots significantly varying ($p < 0.05$) in abundance in challenged leaves, as compared to control ones. In total, 38 protein spots were detected, whose abundance changed in leaves in response to cold or heat shock. Among that, 28 resulted from cold-stressed *vs.* control comparison, while 35 resulted from heat-stressed *vs.* control comparison; they represented 9% and 11% of the protein spots resolved within the leaf reference gel, respectively. All spots showed variably-represented quantitative levels among samples, except two that were present only in control. These figures

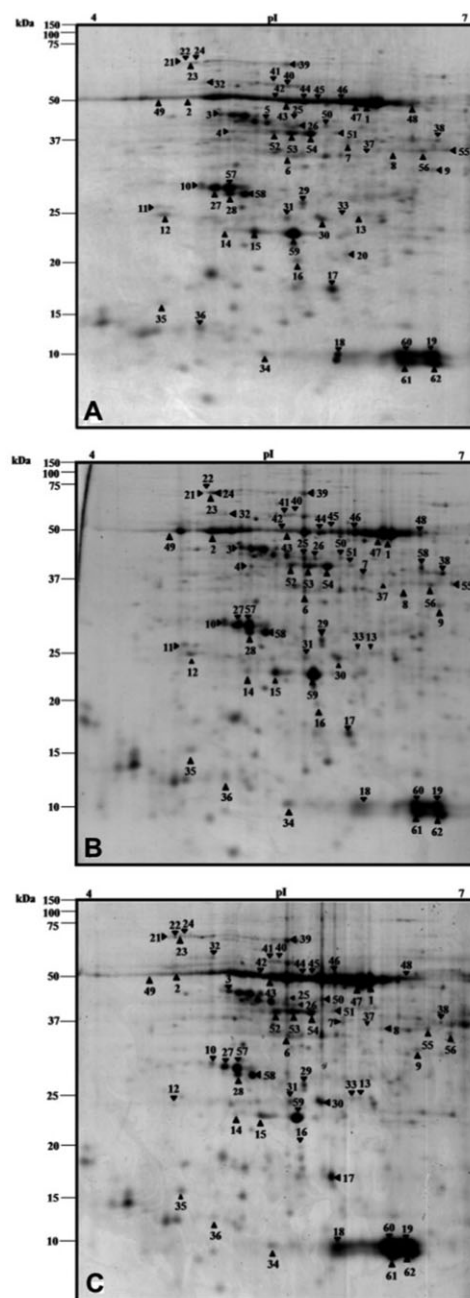


Fig. 2 Two-dimensional electrophoresis maps of total protein extracts from *Arabidopsis* rosette leaves. Panel A, plants treated at 21 °C, for 6 h (control); panel B, plants treated at 4 °C, for 6 h (cold shock); panel C, plants treated at 42 °C, for 6 h (heat shock). Differentially represented (38 in number) or invariant (24 in number) protein spots identified are numbered. Proteins were separated over the *pI* range 4–7 in the first dimension and on 12% SDS–polyacrylamide gels in the second dimension. Gels were stained with colloidal Coomassie G250.

are comparable to those of most proteomic studies concerning the effect of short and even long-term cold or heat stress in *Arabidopsis* and other species, based on 2D gel resolution of cellular protein complement (for a review see ref. 14). Hence, probably, the relatively low number of differential proteins detected is not related to the low level of cellular damage induced under short time stress, but, rather due to the intrinsic

Table 1 Proteins with changed or constant expression levels in *Arabidopsis thaliana* rosette leaves after cold or heat shock challenge. Spot number, protein name, accession number (UniProt and TAIR), identification (ID) method, sequence coverage, number of observed peptides, identification score, theoretical and experimental pI and M_r values, and fold change temperature-treated vs. control plants are listed. PMF, peptide mass fingerprinting; TMS, tandem mass spectrometry, N.d., not detectable in temperature-stressed samples. – refers to values comprised between 0.6 and 1.9 fold changes

| Spot | Protein name | UniProt accession | TAIR accession | ID method | Sequence cov% (peptides) | ID score | Theor. M_r (kDa)/ pI | Exp. M_r (kDa)/ pI | Relative fold change vs. control | |
|------|--|-------------------|----------------|-----------|--------------------------|----------|--------------------------|------------------------|----------------------------------|------|
| | | | | | | | | | Cold | Heat |
| 1 | Ribulose biphosphate carboxylase large chain | O03042 | AtCg00490 | PMF | 60 (31) | 233 | 52.7/5.88 | 51.1/5.92 | 2.0 | 0.3 |
| 2 | Ribulose biphosphate carboxylase large chain | O03042 | AtCg00490 | PMF | 60 (31) | 233 | 52.7/5.88 | 52.1/4.91 | 4.8 | 0.3 |
| 3 | Ribulose biphosphate carboxylase/oxygenase activase, chloroplastic | P10896-1 | At2g39730 | PMF | 47 (17) | 209 | 46.2/5.09 | 44.4/5.10 | 2.2 | 4.3 |
| 4 | Magnesium-chelatase subunit ChlI-1, chloroplastic | P16127 | At4g18480 | PMF | 34 (10) | 174 | 39.9/5.05 | 39.0/5.17 | 2.0 | N.d. |
| 5 | Glutamine synthetase, chloroplastic/mitochondrial | Q43127 | At5g35630 | PMF | 49 (19) | 220 | 42.4/5.28 | 42.4/5.36 | N.d. | N.d. |
| 6 | Elongation factor Tu, chloroplastic | P17745 | At4g20360 | PMF | 36 (13) | 154 | 44.7/5.31 | | | |
| 7 | Probable fructose-bisphosphate aldolase 2, chloroplastic | Q944G9 | At4g38970 | PMF | 48 (12) | 203 | 38.0/5.36 | 35.3/5.51 | 2.1 | 0.4 |
| 8 | ATP synthase gamma chain 1, chloroplastic | Q01908 | At4g04640 | TMS | 26 (9) | 409 | 35.7/6.16 | 37.2/5.81 | 2.1 | 0.4 |
| 9 | Chloroplast stem-loop binding protein of 41 kDa, chloroplastic | Q9LYA9 | At3g63140 | TMS | 21 (6) | 340 | 36.3/6.04 | 35.5/6.23 | 0.5 | 0.3 |
| 10 | Photosystem I reaction center subunit II-2, chloroplastic | Q9SA56 | At1g03130 | TMS | 49 (19) | 585 | 17.7/9.30 | 30.2/6.51 | — | 0.4 |
| 11 | Soluble inorganic pyrophosphatase 1, chloroplastic | Q9LXC9 | At5g09650 | PMF | 26 (8) | 157 | 33.4/5.72 | 26.8/4.98 | — | 0.4 |
| 12 | 29 kDa ribonucleoprotein, chloroplastic | Q43349 | At3g53460 | TMS | 40 (11) | 698 | 29.2/4.76 | 25.8/4.74 | 2.1 | N.d. |
| 13 | 29 kDa ribonucleoprotein, chloroplastic | Q43349 | At3g53460 | TMS | 40 (11) | 656 | 29.2/4.76 | 25.0/4.78 | — | 0.1 |
| 14 | Photosystem I reaction center subunit II-2, chloroplastic | Q9SA56 | At1g03130 | TMS | 44 (9) | 577 | 17.7/9.30 | 25.0/5.90 | — | 0.4 |
| 15 | Oxygen-evolving enhancer protein 2-1, chloroplastic | Q42029 | At1g06680 | PMF | 48 (7) | 129 | 20.2/5.27 | 22.0/5.13 | 3.4 | 0.5 |
| 16 | Oxygen-evolving enhancer protein 2-1, chloroplastic | Q42029 | At1g06680 | PMF | 87 (11) | 148 | 20.2/5.27 | 22.0/5.34 | — | 0.3 |
| 17 | Germin-like protein subfamily 3 member 3 | P94072 | At5g20630 | TMS | 14 (7) | 255 | 19.5/5.84 | 20.7/5.55 | 0.5 | 0.1 |
| 18 | Peptidyl-prolyl <i>cis-trans</i> isomerase CYP20-3, chloroplastic | P34791 | At3g62030 | PMF | 75 (15) | 199 | 19.9/5.47 | 19.1/5.70 | 0.5 | 0.5 |
| 19 | Ribulose biphosphate carboxylase small chain 2B, chloroplastic | P10797 | At5g38420 | PMF | 93 (20) | 179 | 14.8/5.71 | 10.5/5.75 | 2.1 | 0.5 |
| 20 | Ribulose biphosphate carboxylase small chain 2B, chloroplastic | P10797 | At5g38420 | PMF | 93 (20) | 179 | 14.8/5.71 | 10.5/6.51 | 2.3 | 0.5 |
| 21 | Outer membrane lipoprotein Blc | Q9FGT8 | At5g58070 | TMS | 23 (4) | 391 | 21.4/5.98 | 21.0/5.81 | N.d. | N.d. |
| 22 | Ribulose biphosphate carboxylase large chain (fragment) | O03042 | AtCg00490 | PMF | 60 (31) | 233 | 52.7/5.88 | | | |
| 23 | Heat shock 70 kDa protein 6, chloroplastic | Q9STW6 | At4g24280 | TMS | 51 (31) | 813 | 67.2/4.79 | 63.8/4.89 | 4.3 | 6.9 |
| 24 | Heat shock 70 kDa protein 6, chloroplastic | Q9STW6 | At4g24280 | TMS | 51 (31) | 795 | 67.2/4.79 | 64.0/4.90 | 3.2 | 2.9 |
| 25 | Heat shock 70 kDa protein 6, chloroplastic | Q9STW6 | At4g24280 | TMS | 51 (31) | 889 | 67.2/4.79 | 63.9/4.92 | 2.0 | 2.2 |
| 26 | Heat shock 70 kDa protein 6, chloroplastic | Q9STW6 | At4g24280 | TMS | 51 (31) | 609 | 67.2/4.79 | 64.0/4.93 | 4.1 | 2.8 |
| 27 | Elongation factor Tu, chloroplastic | P17745 | At4g20360 | PMF | 36 (13) | 154 | 44.7/5.31 | 42.4/5.53 | 2.5 | 2.7 |
| 28 | Actin-8 | Q96293 | At1g49240 | PMF | 30 (7) | 136 | 41.9/5.37 | 41.9/5.56 | 2.4 | 0.3 |
| 29 | Oxygen-evolving enhancer protein 1-1, chloroplastic | P23321 | At5g66570 | PMF | 43 (10) | 185 | 26.6/4.93 | 26.5/5.08 | — | 0.3 |
| 30 | Oxygen-evolving enhancer protein 1-1, chloroplastic | P23321 | At5g66570 | PMF | 43 (10) | 185 | 26.6/4.93 | 26.2/5.15 | — | 0.4 |
| 31 | Triosphosphate isomerase, cytosolic | P48491 | At3g55440 | PMF | 61(9) | 150 | 27.2/5.39 | 25.6/5.57 | — | 0.4 |
| 32 | Glutathione S-transferase F10 | P42761 | At2g30870 | TMS | 33 (7) | 456 | 24.1/5.49 | 24.1/5.65 | 2.1 | 2.2 |
| 33 | 20 kDa chaperonin, chloroplastic | O65282 | At5g20720 | TMS | 72 (10) | 623 | 21.4/5.23 | 24.2/5.51 | 2.0 | 3.5 |
| 34 | Heat shock 70 kDa protein 10, mitochondrial | Q9LDZ0 | At5g09590 | TMS | 17 (10) | 238 | 68.8/5.14 | 54.8/4.99 | — | 4.5 |
| 35 | Ribulose biphosphate carboxylase large chain (fragment) | O03042 | AtCg00490 | PMF | 60 (31) | 233 | 52.7/5.88 | 25.0/5.80 | 2.1 | — |
| 36 | Uncharacterized protein At1g13930/F16A14.27 | Q9XI93 | At1g13930 | PMF | 27 (10) | 171 | 16.2/4.82 | 10.5/5.36 | 13.6 | 8.2 |
| 37 | Translationally-controlled tumor protein homolog | P31265 | At3g16640 | TMS | 27 (4) | 403 | 18.9/4.52 | 16.3/4.75 | 2.0 | 2.7 |
| 38 | Thioredoxin M4, chloroplastic | Q9SEU6 | At3g15360 | TMS | 16 (3) | 253 | 12.5/5.42 | 13.2/4.98 | 0.3 | 0.1 |
| 39 | Annexin D1 | Q9SYT0 | At1g35720 | TMS | 19 (4) | 248 | 36.2/5.21 | 35.5/5.93 | 0.3 | 0.4 |
| 40 | Glyceraldehyde-3-phosphate dehydrogenase A, chloroplastic | P25856 | At3g26650 | PMF | 14 (5) | 91 | 36.3/6.67 | 37.5/6.58 | — | 3.2 |
| 41 | Cell division control protein 48 homolog A | P54609 | At3g09840 | TMS | 8 (5) | 188 | 89.3/5.13 | 63.9/5.52 | — | — |
| 42 | Mediator of RNA polymerase II transcription subunit 37a | Q9LKR3 | At5g28540 | PMF | 26 (17) | 149 | 70.8/5.05 | 54.7/5.52 | — | — |
| 43 | Mediator of RNA polymerase II transcription subunit 37f | Q39043 | At5g42020 | TMS | 24 (14) | 668 | 71.1/5.08 | 54.8/5.43 | — | — |

Table 1 (continued)

| Spot | Protein name | UniProt accession | TAIR accession | ID method | Sequence cov% (peptides) | ID score | Theor. M_r (kDa)/pI | Exp. M_r (kDa)/pI | Relative fold change vs. control | |
|------|---|-------------------|----------------|-----------|--------------------------|----------|-----------------------|---------------------|----------------------------------|------|
| | | | | | | | | | Cold | Heat |
| 42 | Ribulose biphosphate carboxylase large chain | O03042 | AtCg00490 | TMS | 11 (6) | 91 | 52.7/5.88 | 52.1/5.44 | — | — |
| 43 | Ribulose biphosphate carboxylase large chain | O03042 | AtCg00490 | TMS | 26 (12) | 161 | 52.7/5.88 | 52.0/5.51 | — | — |
| 44 | Ribulose biphosphate carboxylase large chain | O03042 | AtCg00490 | TMS | 20 (9) | 415 | 52.7/5.88 | 51.9/5.57 | — | — |
| 45 | ATP synthase subunit beta, chloroplastic | P19366 | AtCg00480 | TMS | 34 (12) | 91 | 53.9/5.38 | 52.0/5.63 | — | — |
| 46 | Ribulose biphosphate carboxylase large chain | O03042 | AtCg00490 | TMS | 16 (7) | 91 | 52.7/5.88 | 51.9/5.78 | — | — |
| 47 | Ribulose biphosphate carboxylase large chain | O03042 | AtCg00490 | PMF | 28 (21) | 161 | 52.7/5.88 | 51.8/5.89 | — | — |
| 48 | Ribulose biphosphate carboxylase large chain | O03042 | AtCg00490 | TMS | 23 (9) | 1441 | 52.7/5.88 | 51.7/6.30 | — | — |
| 49 | ATP synthase subunit beta, chloroplastic | P19366 | AtCg00480 | PMF | 11 (3) | 269 | 53.9/5.38 | 52.1/4.74 | — | — |
| 50 | Protein disulfide-isomerase like 2-1 | O22263 | At2g47470 | TMS | 34 (10) | 475 | 37.1/5.65 | 41.9/5.66 | — | — |
| 51 | Cysteine synthase, chloroplastic/chromoplastic | P47999 | At2g43750 | PMF | 25 (7) | 554 | 35.1/5.54 | 39.8/5.74 | — | — |
| 52 | Ribulose biphosphate carboxylase/oxygenase activase, chloroplastic | P10896-2 | At2g39730 | PMF | 30 (16) | 169 | 43.4/5.42 | 39.8/5.43 | — | — |
| 53 | Ribulose biphosphate carboxylase/oxygenase activase, chloroplastic | P10896-2 | At2g39730 | PMF | 37 (15) | 186 | 43.4/5.42 | 39.8/5.53 | — | — |
| 54 | Ribulose biphosphate carboxylase/oxygenase activase, chloroplastic | P10896-2 | At2g39730 | PMF | 32 (17) | 188 | 43.4/5.42 | 39.5/5.58 | — | — |
| 55 | Malate dehydrogenase 1, mitochondrial | Q9ZP06 | At1g53240 | PMF | 42 (9) | 741 | 33.3/6.00 | 35.5/6.70 | — | — |
| 56 | Malate dehydrogenase 1, mitochondrial | Q9ZP06 | At1g53240 | TMS | 26 (9) | 643 | 33.3/6.00 | 35.5/6.45 | — | — |
| 57 | Eukaryotic translation initiation factor 2 subunit beta | Q41969 | At5g20920 | TMS | 10 (3) | 110 | 30.7/6.79 | — | — | — |
| 58 | Ribulose biphosphate carboxylase/oxygenase activase, chloroplastic (fragment) | P10896 | At2g39730 | PMF | 16 (8) | 114 | 46.2/5.09 | 26.5/5.15 | — | — |
| 59 | Oxygen-evolving enhancer protein 1-2, chloroplastic | Q9S841 | At3g50820 | TMS | 49 (14) | 1154 | 26.6/5.02 | 26.2/5.25 | — | — |
| 60 | Carbonic anhydrase, chloroplastic | P27140 | At3g01500 | PMF | 39 (12) | 202 | 25.6/6.14 | 21.9/5.53 | — | — |
| 61 | Ribulose biphosphate carboxylase small chain 3B, chloroplastic | P10798 | At5g38410 | PMF | 47 (9) | 184 | 14.8/5.71 | 10.5/6.25 | — | — |
| 62 | Ribulose biphosphate carboxylase small chain 3B, chloroplastic | P10798 | At5g38410 | PMF | 52 (10) | 199 | 14.8/5.71 | 8.3/6.25 | — | — |
| 62 | Ribulose biphosphate carboxylase small chain 3B, chloroplastic | P10798 | At5g38410 | PMF | 43 (8) | 152 | 14.8/5.71 | 8.3/6.52 | — | — |

limitations of 2D gel-based proteomics, in which most abundant proteins and/or soluble ones are preferentially identified.

Differential spots were excised from gels, trypsinolyzed and subjected to MS analysis for further protein identification assignment. Twenty-two positive identification results derived from MALDI-TOF PMF data, whereas 18 from nanoLC-ESI-LIT-MS/MS ones; two spots were associated with multiple proteins and were not further discussed. Globally, spots assayed were associated with 28 non-redundant protein entries. The list of identified protein species, together with their quantitative variations as a result of cold and heat shock treatments, is reported in Table 1. Some proteins, such as heat shock 70 kDa protein 6, ribulose biphosphate carboxylase large and small 2B subunits, 29 kDa ribonucleoprotein (CP29A) and oxygen-evolving enhancer proteins 1-1 and 2-1, occurred as multiple spots whose structural differences were not further characterized. Ribulose biphosphate carboxylase large chain and ribulose biphosphate carboxylase/oxygenase activase were also identified among constant spots (see below). Probably, they were the result of post-translational modification or differential splicing events. Functional categorization according to Gene Ontology annotation and literature data (data not shown) showed that differentially-represented proteins grouped into two main broad classes including components involved in energy and metabolism (55%) or in stress response (34%);

the remaining proteins (11%) were related to different processes and were categorized as a miscellaneous group.

A Venn diagram representation of the differentially-represented spots indicated that a significant overlap occurred between the response to short-term cold and heat shock (Fig. 3). In fact, quantitative levels of 25 protein spots were affected by both

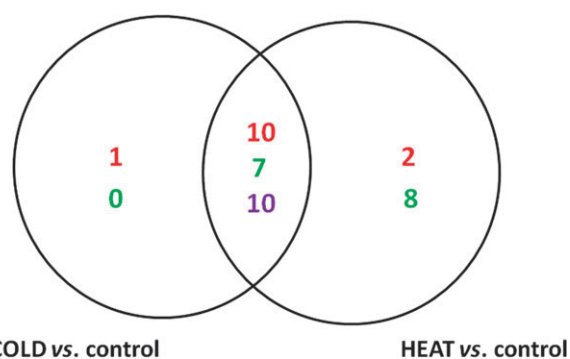


Fig. 3 Venn diagram analysis of the differentially expressed proteins during cold or heat shock treatments. Twenty-five protein spots were affected by both temperature treatments, 23 protein spots were differentially-represented following treatments, whereas 2 protein spots were detected only in control samples. Red, over-represented spots; green, down-represented spots; purple, over-represented spots in cold vs. control and down-represented in heat vs. control.

1 temperature treatments. In particular, 10 proteins, namely
ribulose biphosphate (RuBis) carboxylase/oxygenase activase
(spot 3), heat shock 70 kDa protein (HSP70) 6 (spots 21–24),
5 elongation factor Tu (EF-Tu) (spot 25), glutathione S-transferase
(GST) F10 (spot 30), 20 kDa chaperonin (spot 31), uncharacterized
protein At1g13930/F16A14.27 (spot 34) and translationally-
10 controlled tumor protein (TCTP) homolog (spot 35) were over-
represented following both cold and heat stress; five proteins,
namely chloroplast stem-loop binding protein of 41 kDa (spot 8),
germin-like protein subfamily member 3 (spot 16), peptidyl-
15 prolyl *cis-trans* isomerase CYP20-3 (CYP20-3) (spot 17), thio-
redoxin (TRX) M4 (spot 36) and annexin (ANX) D1 (spot 37),
together with those comigrating in spots 5 and 20, were down-
represented or absent following both cold and heat stress; ten
20 protein spots, namely RuBis carboxylase large chain (spots 1
and 2), magnesium chelatase subunit ChlI-1 (spot 4), fructose-
biphosphate aldolase 2 (spot 6), ATP synthase (ATPase)
gamma chain 1 (spot 7), 29 kDa ribonucleoprotein (spot 11),
oxygen-evolving enhancer (OEE) protein 2-1 (spot 14), RuBis
carboxylase small chain 2B (spots 18 and 19) and actin 8 (spot
26) were over-represented after cold shock and down-repre-
sented following heat shock.

On the other hand, a number of highly represented protein
spots were recognized as constantly present within the different
25 2-DE maps. Among that, 24 (spots 39–62) were similarly
sampled from gels, trypsinolyzed and subjected to MS analysis
for protein identification; all spots were associated with a
unique protein sequence entry, with a unique exception (spot
56) in which multiple components migrated together (Table 1).
30 Constant spots were identified as RuBis carboxylase small
chain 3B, ATPase subunit **beta**, protein disulfide-isomerase like
2-1, cysteine synthase, malate dehydrogenase 1, eukaryotic
translation initiation factor 2 subunit **beta**, OEE protein 1-2,
35 carbonic anhydrase, cell division control protein 48 homolog A,
and mediator of RNA polymerase II transcription subunits 37a
and 37f, as well as specific isoforms of RuBis carboxylase large
chain and RuBis carboxylase/oxygenase activase.

40 3.3 Proteins affected by cold treatment

Proteomic analysis suggested that short-term cold treatment of
Arabidopsis thaliana had a significant effect on specific pro-
cesses of plant photosynthesis. In fact, incubation of seedlings
at 4 °C for 6 h, brought about a quantitative increase in proteins
45 involved both in electron transport/energy production and
carbon metabolism reactions or, at least, specific isoforms of
them. As far as the first protein family, ATPase γ chain (spot 7),
and isoforms of OEE protein 2-1 (spot 14) and of 29 kDa
chloroplastic ribonucleoprotein (spot 11) were up-regulated,
50 respectively. Among Calvin cycle enzymes, levels of RuBis
carboxylase small chain 2B (spots 18 and 19) and of specific
isoforms of RuBis carboxylase large chain (spots 1, 2 and 33)
and RuBis carboxylase/activase (spot 3) were significantly
increased. On the other hand, ATPase β chain, OEE proteins
55 1-1 and 1-2, photosystem I reaction center subunit II-2, RuBis
carboxylase small chain 3B, and other isoforms of RuBis

carboxylase large chain and RuBis carboxylase/activase
remained constantly represented. 1

On the overall, the observed trends are fairly in accordance
with previous literature data. In fact, increased abundance of
specific components of the thylakoid photosynthetic apparatus,
5 including OEE proteins, has been already observed in cold-
treated *Arabidopsis*¹⁰ or in the cold-tolerant species *Thellungiella*
halophila subjected to short-term cold challenge.¹⁵ Similarly,
increased levels of certain chloroplast electron transport chain
10 components have been related to improved cold resistance.³² As
far as CP29A, our findings point to a pivotal role of this protein in
cold tolerance. In fact, it has been identified by previous proteo-
mic investigations as a cold up-regulated species both in
*Arabidopsis*¹⁰ and *Thellungiella*,¹⁵ while differences in CP29A
15 phosphorylation status have been reported between cold-tolerant
and sensitive maize cultivars, and correlated to photoinhibition
of cold-sensitive lines.³³ In *Arabidopsis*, it has recently been
shown that CP29A and CP31A are essential to maintain transcript
stability of different chloroplast mRNAs under cold stress.³⁴
20 Conversely, contrasting data have been reported for the effect
of cold stress on the abundance of Calvin cycle enzymes, depend-
ing on the species investigated and/or the type of cold treat-
ment.¹⁴ For example, increased levels of RuBis carboxylase large
subunit have been reported in rice, together with extensive
25 protein degradation,¹³ whereas a heterogeneous pattern was
observed in *Thellungiella*, depending on time of treatment and
spot multiplicity.¹⁵ The latter scenario may result from the
occurrence of post-translational modifications (*i.e.* phosphoryla-
tion, sumoylation) that can affect electron transport/energy pro-
duction and carbon metabolism enzymes under stress
30 conditions.^{35,36} Their detection needs dedicated approaches to
ascertain the presence of specific isoforms in 2-DE maps, as we
detected in this study. Under our conditions, the stress treatment
caused a significant increase in the abundance of RuBis carbox-
ylase small chain 2B (spots 18 and 19) and of specific isoforms of
35 both RuBis carboxylase large chain and RuBis carboxylase/acti-
vase; proteolytic degradation of RuBis carboxylase large chain was
also observed. Finally, cold stress has been reported to determine
an up-regulation of carbon catabolism enzymes.¹⁴ In this
context, we identified only fructose-biphosphate aldolase 2 as
40 up-regulated enzyme. This protein is involved in chloroplast
glycolysis, and its levels have been reported to increase in rice
leaves in response to long-term (48 h) cold stress.¹³

The second prominent functional group was represented by
proteins involved in the response to environmental stresses. In
45 particular, an over-representation of four spots (21, 22, 23
and 24) associated with 70 kDa class of heat shock protein 6
was evident following cold shock. Their electrophoretic pattern
suggests that, notwithstanding the occurrence of post-transla-
tional modifications affecting corresponding polypeptide
50 chain, increased amounts of all protein isoforms were detected.
Quantitative levels of 20 kDa chaperonin (spot 31) and GST F10
(spot 30) were also significantly increased, while those of
protein At1g13930/F16A14.27 (spot 34), an uncharacterized
component “involved in salt stress response”, showed the
55 largest increase among up-regulated components.

1 HSP70s belong to a conserved family of molecular chaperones that assist folding, assembly, translocation and degradation of proteins in all cellular compartments. By stabilizing protein conformation, they are essential to maintain cellular homeostasis under stressing conditions;³⁷ their increase in response to a wide range of environmental stresses (including cold) has been reported in different plant species.¹⁴ Interestingly, HSP70s have been reported in *Arabidopsis* as nuclear proteins induced after short-term cold challenge.⁸ 20 kDa chaperonin is chloroplast-located and its induction, which in this study parallels to that of many other chloroplastic polypeptides (34% of deregulated components concerned chloroplast-located species), confirms the importance of this organelle in temperature stress response. Cold stress alters cellular redox homeostasis, hence it is not surprising that accumulation of different enzymes deputed to ROS neutralization has been reported in many plants,¹⁴ including components of the ascorbate–glutathione cycle and various GST isoforms.^{14,32,38} Here, we identified a member of the phi class of the GST family, which appears to be ubiquitous in plant tissues and whose down-regulation in transgenic plants causes reduced tolerance to abiotic stresses.³⁹ Indeed, At1g13930 was previously identified by means of a functional gene-mining method to isolate salt stress tolerance genes in *Thellungiella halofila*.⁴⁰ Its function is unknown but it has been associated with salt tolerance; its strong up-regulation in our conditions suggests that it can also play a key role in cold tolerance.

On the other hand, the abundance of chloroplastic TRX M4 (spot 36), chloroplastic CYP20-3 (spot 17), ANX D1 (spot 37) and germin-like protein 3 (spot 16) was decreased after short-term cold stress. Thioredoxins m-type have been proposed to specifically regulate chloroplast glucose-6-phosphate dehydrogenase, inhibiting in the light oxidative generation of NADPH, when it is produced by the photosynthetic electron flow.⁴¹ Decreased levels under cold stress may reflect the necessity of the cell to increase reducing power with the aim of contrasting oxidative stress; this observation is in accordance with the general increase of glycolytic catabolic reactions detected during cold stress.¹⁴ CYP20-3 belongs to the large family of cyclophilins, whose primary function is to assist protein folding; they are induced in response to both abiotic and biotic stresses.²⁴ On the other hand, annexins are a conserved family of eukaryotic Ca²⁺-dependent phospholipid-binding proteins; although their expression pattern is developmentally regulated, different lines of evidence indicate that they are also involved in the protection from biotic and abiotic stresses.⁴² The rationale for the observed down-regulation under our conditions of the latter two proteins is unclear at present. Indeed, germins, which belong to the conserved cupin superfamily, have been described in barley as pathogenesis-related proteins. By eliciting an oxalate oxidase or superoxide dismutase activity, they are able to generate H₂O₂; they are very likely involved in cell wall stiffening or signalling against pathogen infection. Germin-like genes are activated also in response to abiotic stress, such as high salinity or heat.⁴³ Intriguingly, germins have already been identified in *Arabidopsis* leaves as nuclear proteins down-regulated upon short-term cold challenge.⁸

Other variably-represented proteins were identified as components whose function was related to various cellular processes; they included actin 8 (spot 26), TCTP homolog (spot 35), EF-Tu (spot 25) and magnesium-chelatase subunit ChII-1 (CHLI) (spot 4). The first two proteins are structural components involved in cytoskeleton organization and stabilization, respectively. Plasma membrane is an early target of cold injury and experimental evidence suggests that actin filaments of the cytoskeleton, as linked to the plasma membrane, are involved in the sensing of and in the response to low temperature.⁴⁴ In the moss *Psyscomitrella patens*, cellular actin content dramatically increases after long term cold stress.⁴⁵ Under our conditions, actin and TCTP homolog increased as well, even at much shorter times of cold exposure, although to a lesser extent. These findings reinforce the idea that cytoskeleton is an early target of cold damage; its rearrangement may play a pivotal role in cold tolerance. On the other hand, EF-Tu and CHLI are chloroplast-located proteins involved in the regulation of plastidial protein synthesis and chlorophyll biosynthesis, respectively; under our conditions, both proteins were up-regulated after cold stress. An EF-Tu over-representation has been already reported in cytokinin over-expressing creeping bentgrass under drought conditions;⁴⁶ it was paralleled by an increase of chloroplastic enzymes involved in photosynthesis and energy production. Our results also point to a role for this protein in sustaining protein synthesis within the chloroplast, in order to preserve the integrity of the energy production machinery under cold stress conditions. As far as magnesium-chelatase is concerned, it is worth mentioning that different subunits (CHLH, CHLI) of this enzyme have recently been shown to be involved in abscisic acid (ABA) signaling modulation.⁴⁷ ABA is a well-known environment-responsive phytohormone that plays a key role in adaptation to different stresses, including the cold one.⁴⁸

3.4 Proteins affected by heat treatment

Short-term heat treatment (42 °C, 6 h) of *Arabidopsis* seedlings also affected the abundance of leaf proteins involved in the photosynthetic electron transport and carbon metabolism. Unlikely to cold stress, a general protein down-representation was observed in this case. In fact, levels of OEE protein 2-1 (spots 14 and 15), OEE protein 1-1 (spots 27 and 28), subunit II-2 of the photosystem I reaction center (spots 9 and 13) and chloroplastic 29 kDa ribonucleoprotein (spots 11 and 12) were significantly reduced; the same trend was also observed for RuBis carboxylase small chain 2B (spots 18 and 19) and specific isoforms of RuBis carboxylase large chain (spots 1 and 2). Similarly to cold stress, a consistent increase of a specific isoform of RuBis carboxylase/activase was conversely detected (spot 3). On the other hand, RuBis carboxylase small chain 3B, and other isoforms of RuBis carboxylase large chain and RuBis carboxylase/activase remained constantly represented. Indeed, heat stress was shown to down-regulate different key enzymes of the Calvin cycle in rice, including RuBis carboxylase,⁴⁹ while an up-regulation of RuBis carboxylase/activase has already been reported in rice⁴⁹ and wheat.⁵⁰ It has been suggested that a

1 RuBis carboxylase/activase over-representation may be part of
the adaptative response, in order to maintain CO₂ fixation
under stress conditions.²² Considerations analogous to that
5 reported above for cold stress can be formulated to justify the
isoform-specific pattern of electron transport/energy produc-
tion and carbon metabolism enzymes as a result of post-
translational modifications.^{35,36}

As far as carbon metabolism and energy generation, it has
been shown that heat stress negatively affects the glycolytic
10 pathway and reduces energy production in rice, bringing about
a marked decrease in the levels of different glycolytic enzymes
or proteins involved in energy-generating reactions.²³ In agree-
ment with this scenario, we observed a down-representation of
fructose-bisphosphate aldolase 2 (spot 6), cytosolic triosepho-
15 sphate isomerase (spot 29), ATPase γ chain (spot 7) and soluble
inorganic pyrophosphatase 1 (PPase1) (spot 10), whereas an
increase in the abundance of chloroplastic glyceraldehyde
3-phosphate dehydrogenase (GAPDH) (spot 38) was detected.
By catalyzing the exergonic hydrolysis of pyrophosphate,
20 PPase1 plays a crucial role in cellular energy conservation. On
the other hand, plants contain three forms of GAPDH encoded
by distinct genes: a cytosolic form that participates in glycolysis
and two chloroplast forms involved in photosynthesis. It has
been suggested that glycolytic GAPDH may be also involved in
25 stress response, since its levels are increased by different
environmental challenges, including heat;²³ no literature data
are available on stress modulation of chloroplastic GAPDH.

Similarly to cold stress, the second functional group most
influenced by heat treatment was that of proteins involved in
30 stress response, particularly molecular chaperones. In fact, heat
stress is primarily associated with improper protein folding or
denaturation. Likewise cold stress, four spots (21, 22, 23 and
24) identified as 70 kDa class of heat shock protein 6 were
consistently up-regulated, together with another strongly
35 induced mitochondrial HSP70, *i.e.* isoform 10 (spot 32), appar-
ently specific for the heat stress condition. Finally, levels of 20
kDa chaperonin (spot 31) were more prominently up-regulated
than in cold stress, while that of CYP20-3 (spot 17) were
similarly decreased. A general deregulation of proteins with a
40 protecting role from oxidative damage was also observed in this
case, with a pattern similar to that induced by cold stress. In
fact, GST F10 (spot 30) was over-represented, while levels of
chloroplastic TRX M4 (spot 36) were decreased. Among other
proteins involved in stress response, worth noting is that
45 uncharacterized protein At1g13930/F16A14.27 (spot 34) was
the most prominently over-represented component, as in the
case of cold stress; this suggests a general role for this protein
in adaptation to fast temperature changes. On the other hand,
ANX D1 (spot 37) and germin-like protein 3 (spot 16) were both
50 down-regulated, similarly to cold stress.

As far as proteins involved in other cellular functions, the
pattern of expression following heat stress was qualitatively
reminiscent of that resulting from cold stress, concerning a
limited number of components involved in similar functions.
55 Cytoskeleton structure was apparently a target also for heat
stress injury. Microtubule stabilizer TCTP homolog (spot 35)

was up-regulated after heat shock, analogously to cold stress,
while actin 8 (spot 26) was down-regulated. Finally, EF-Tu
(spot 25) abundance was increased to the same extent as under
low temperature conditions. Besides promoting aminoacyl-
5 tRNA binding to ribosomes, it has been reported that this
protein can function as a molecular chaperone under heat
stress conditions, thus protecting chloroplast proteins from
misfolding and aggregation;⁵¹ accordingly, its synthesis is
differentially regulated in heat-sensitive and tolerant maize
10 cultivars.⁵²

4 Concluding remarks

Proteomics can significantly contribute to the understanding of
physiological modifications underlying response of plants to
15 temperatures stresses, which generally involves profound
changes in the cellular protein repertoire.¹⁴ Proteomic informa-
tion on plant temperature stress is still fragmentary, mostly
concerning the study of adaptative responses of crop species.
Since stress response is influenced by duration and intensity of
20 temperature challenge (*i.e.* shock or adaptation), corresponding
patterns of proteome variations are expected to be (very likely)
only partially coincident. Here we have reported on the com-
parative proteomic analysis of the effect of short-term cold and
heat treatment (shock) on leaves of the model species *Arabi-*
25 *dopsis thaliana*. Results allowed us to identify a number of
proteins, participating in different cellular functions, some of
which are already known to be involved in response to pro-
longed cold or heat stresses or acclimation in *Arabidopsis* and/
or other species.^{14,15,23} Novel proteins were also identified as
30 potential markers to be associated with the alteration of
cellular functions imposed by temperature stress or to the
acquisition of stress tolerance. In this context, original infor-
mation was derived for At1g13930/F16A14.27, CHL1, EF-Tu,
chloroplastic GAPDH, TCTP, EF-Tu, chloroplastic TRX M4,
35 GST F10, chloroplastic 20 kDa chaperonin and mitochondrial
HSP70 isoform 10. On the other hand, data on the remaining
differentially-represented proteins corroborate previous observa-
tions on temperature treatments under various experimental
conditions in *Arabidopsis*, *Thellungiella*, maize or rice.^{10,13,15,33,34,52}
40

Our experiments ascertained that a significant degree of
overlapping exists between metabolic alterations induced by,
and/or in response to cold and heat short-term challenge. In
fact, many differentially-represented proteins were modulated
45 by both stimuli, with some remarkable exceptions, from which
specific features of cold or heat responses can be inferred.
According to literature data, chloroplast and also plasma
membrane/cytoskeleton were confirmed as early targets for
temperature injury. Cold and heat influenced photosynthesis
50 in opposition. Over-representation of enzymes involved in
electron transport and Calvin cycle seems to be distinctive of
the cold response, aimed at protection of plant from photo-
inhibition. This response is apparently hampered by heat
stress, which probably affects thylakoidal structures more
55 severely; on the other hand, the shared proteomic signatures
we observed may be part of a common cellular strategy to

1 sustain carbon fixation under general stress conditions. In this
 2 context, 29 kDa ribonucleoprotein (specifically induced by cold
 3 stress) may deserve supplemental studies to envisage its possi-
 4 ble use as a marker for cold tolerance selection. On the other
 5 hand, CHLI, chloroplastic GAPDH and uncharacterized protein
 6 At1g13930/F16A14.27 emerged as novel intriguing chloroplast
 7 proteins, induced by cold, heat stress or both, respectively,
 8 which could also be investigated for their possible use as
 9 temperature tolerance markers. Our results also confirmed that
 10 major components of the early response to temperature stress
 11 are HSPs. Among the different chaperones whose increase has
 12 been reported in different species, HSP70 isoform 10 and 20
 13 kDa chaperonin were here preferentially induced either follow-
 14 ing heat and cold stresses, while the HSP70 isoform 10 was
 15 apparently specific to the heat shock condition. Finally, cyto-
 16 plasmic detoxification of ROS relied preferentially on the
 17 induction of GST under both cold or heat stress. In conclusion,
 18 our investigation corroborates previous observations of quanti-
 19 tative proteomic changes in other species and defines a more
 20 focused picture of the early protein changes associated with
 21 both temperature challenges, which can hopefully orient future
 22 integrated approaches to gain a deeper insight into the complex
 23 network of plant response to environmental stresses.

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