Methods for On-Orbit Germination of Arabidopsis thaliana for Proteomic Analysis

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ABSTRACT

Large-scale omics approaches make excellent choices for research aboard the International Space Station (ISS) because they provide large amounts of data that can be continually mined even after the original research has been completed. A proteomic approach can provide information about which proteins are produced, degraded, or post-translationally modified, potentially shedding light on cellular strategies that cannot be discerned from transcriptomic data. To collect sufficient tissue from a Biological Research In Canisters (BRIC)-grown experiment on the ISS for proteomic analysis, several modifications were made to existing protocols. Approximately 800-1000 seeds were housed in each Petri Dish Fixation Units (PDFU). These were germinated up to 120 h after planting by transferring the BRIC from cold stasis to room temperature. Growth continued for only 72 h after germination to allow sufficient tissue for extraction, and to minimize the impact of ethylene and crowding stress. Seedlings were then exposed to RNAlater®. Results indicate that RNAlater®-treated Arabidopsis seedlings yield an equal amount of protein to those flash-frozen in liquid nitrogen.

INTRODUCTION

The use of proteomics to track relative protein expression is becoming a common approach to understand how an organism responds on a molecular level to a variety of environmental conditions or genetic mutations. Historically, a transcriptomic approach has been used to understand changes in plant gene expression during spaceflight (Beaulieu et al., 2013; Paul et al., 2005; Paul et al., 2012; Paul et al., 2013; Salmi and Roux 2008; Salmi et al., 2011a; Salmi et al., 2011b). This is an attractive approach because vast quantities of data can be obtained from very little tissue. However, changes in gene expression do not necessarily reflect direct changes in protein quantity (Griffin et al., 2002). Considering that the presence of active protein is the final determinant of a cellular response to a given environmental stimulus, a proteomic approach can provide more reliable data. In addition to protein production, proteomics can also identify protein degradation, as well as post translational modifications, such as phosphorylation, which can be critical for activating or inactivating function.

Key words: Arabidopsis; Proteomics; Transcriptomics; BRIC; ISS; RNAlater®

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One of the main drawbacks to using proteomics as a tool to analyze the microgravity environment is the amount of tissue required to obtain sufficient material for analysis. Experiments that use the popular and accessible Biological Research In Canisters (BRIC) hardware are typically limited to a handful of seedlings growing on a few 60 mm Petri dishes, often in the dark (Kuznetsov et al., 2001; Millar et al., 2011). Under these conditions, lifespan is short, plants are limited to carbon sources from the media when producing biomass, and stress from etiolated growth can confound analysis of the stress induced by microgravity by alternatively regulating, and therefore masking, those genes that would normally appear in a microgravity transcriptomic data set.

Improved isobaric tags and mass spectrometry technology have begun to make proteomics more accessible for research with limited tissue availability (Reviewed by Vanderschuren et al., 2013). To date, discovery proteomics has been used to analyze a variety of bacteria, mammalian cells, *Saccharomyces cerevisiae*, rice leaf tissue, Arabidopsis callus, and de-etiolated 12-d-old Arabidopsis seedlings (Ma et al., 2007; Crabbe et al., 2011; Grimm et al., 2011; Ferl et al., 2015). Previous plant samples have utilized lighted environments that allowed photosynthesis, carbon fixation, and increased plant mass. Here we focus on refining methods for performing proteomic analysis on Arabidopsis samples grown in the BRIC hardware, a dark environment with limited growth space. Specifically, planting density to obtain sufficient tissue, coordinated on-orbit germination in darkness, growth time, and preservation of sample protein content must all be refined to collect a useable sample for analysis (Figure 1). By utilizing high-density planting, a 72 h etiolated growth plan, and RNAlater® for protein stability, the BRIC hardware can be successfully used to generate proteomic data from etiolated Arabidopsis plants germinated and grown on-orbit aboard the ISS.

**Figure 1. Protocol flow chart for proteomic analysis of BRIC-grown material.** Blue ovals indicate aspects of the approach that are addressed in this work.
MATERIALS AND METHODS

Plant Growth

Arabidopsis seeds in the Col-0 background were surface sterilized in 30% bleach/0.01% Tween 20 solution. Sterile seeds were placed on individual filter paper discs to dry, prior to random planting across a 60 mm Petri dish containing 0.8% agar, 0.5x Murashige & Skoog media (Caisson Laboratories, Smithfield, UT), and 1% sucrose. Unless otherwise indicated, Petri dishes were sealed with parafilm, wrapped in foil, and stratified for 48 h to 72 h prior to receiving 2 h white light to induce germination. Plants were then wrapped in foil and placed inside a modified coffee mug to approximate the BRIC environment.

Figure 2. Comparison of mean protein yield by starting seed weight. Seed sets weighing between 0.026 g (~1300 seeds) to 0.006 g (~300 seeds) were grown on plates for 96 h prior to protein extraction. * indicates p<0.03 using a Student’s T-test. Bars represent an average of three separate extractions, each using approximately 1300 seeds. Error bars represent SEM.

Cold Treatment to Delay Germination

Modifications to the standard germination and growth timeline were made to accommodate the requirements of a flight schedule. After seeds were sown, plates were placed at 4°C in the dark for 18 h, followed by a 2 h light treatment at room temperature. After light treatment, plates were returned to the cold. A subset of individually wrapped plates was removed every 48 h to score germination, preventing light contamination of the remaining plates. This germination and growth protocol was used to analyze cold storage prior to germination, as well as optimal growth time after transfer to room temperature.

RNAlater® Treatment

After growth for 96 h at room temperature, Petri dishes were flooded with 10 mL RNAlater® and resealed for 12 h. Petri dishes with RNAlater® were then placed into the -80°C freezer for 12 h to 72 h, prior to protein isolation. Total soluble protein was extracted from whole seedlings as described previously (Basu et al., 2015). Protein yield was quantified using the Bradford Assay (Fisher Scientific). Absorption at 595 nm was compared to a bovine serum albumin (BSA) standard curve.

RESULTS

High-Density Seed Plating to Improve Protein Yield

Unlike transcriptomic approaches, proteomics requires a minimum of 50 μg of protein per replicate (Murad et al., 2011). One option for obtaining this material, while limiting the number of Petri Dish Fixation Units (PDFUs) necessary, is the use of high-density planting. To determine the optimal number of seeds per plate, between 0.006 g and 0.016 g of Arabidopsis seed was sown on a 60 mm Petri dish and grown for 96 h prior to protein extraction. As seedlings from 3 plates were combined to form one replicate, 15-20 μg protein per plate would be sufficient for the amount of protein required for one replicate. However, increasing the number of seeds above 0.016 g did not further improve results. Therefore, we determined that under these growth conditions 0.016 g was the optimal amount for planting.

Cold Storage Can Delay Germination

Research utilizing the ISS often requires a several-day stasis between integration of the biological sample into the flight hardware and experiment activation. It is critical that seedlings germinate in near-unison to prevent life-stage fluctuations in the data. However, seedlings must also not germinate early, exposing them to the
accelerated gravity of launch and potential problems created by growth while in cold stowage. To mitigate this problem, seeds were sown on Petri dishes and allowed to imbibe at 4°C for approximately 18 h, exposed to white light for 2 h, and then returned to darkness at 4°C to determine how long germination could be delayed. Plants left at 4°C after the light treatment did not show visible indications of germination up to 168 h (7 d), when a small percentage did begin to germinate. However, plants placed at 23°C immediately after light treatment began to show germination after 72 h, with full germination apparent after 120 h (Figure 3). After 144 h (6 d) in the cold, return to room temperature led to visible germination of 95% of seeds after 44 h (Figure 4), indicating that this is an effective method to temporarily inhibit, yet uniformly induce germination.

Figure 3. Cold storage of planted seed delays germination for up to 120 h. Approximately 100 seeds were planted on ½ MS agar +1% sucrose plates. Plates were cold treated for 18 h, followed by 2 h white light. Plates were then returned to cold, with a subset of plates removed and imaged every 24 h. Representative plates are shown. Arrows indicate seedlings emerging from the seed.
Figure 4. Germination efficiency after return to room temperature. Plates were cold treated for 18 h, exposed to 2 h white light, followed by an additional 6 days at 4°C. After return to room temperature, germination percentage was recorded every 6 h for 44 h. * (p<0.03) and # (p<0.01) indicate significant result from a Student’s T-test. n=3 replicates of approximately 1000 seeds each. Bars represent standard error.

72 Hours of Growth Maximized Protein Yield

Uniform germination is induced by shifting the BRICs from cold storage at 4°C to ambient temperature aboard the ISS. To determine how long plants should be allowed to grow prior to fixation, a time series was performed analyzing protein yield after three, four, and five days post-transfer to room temperature. Prior to three days, growth was not sufficient to perform extractions. Increasing growth time from three to four days did not appear to significantly impact protein yield (Figure 5). Increasing the time to five days did improve the output. However, due to the high density planting, three days was chosen as the preferred growth time to minimize the impact of crowding stress and ethylene build-up within the PDFUs. Furthermore, we reasoned that most growth after three days could be attributed to vacuole expansion rather than a marked increase in cell division (Gendreau et al., 1997).

RNAlater® Successfully Preserves Protein for Proteomic Analysis

After three days of growth, samples within the BRICs must be preserved for analysis, potentially months later. Direct transfer to the onboard Minus-Eighty Degree Laboratory Freezer for ISS (MELFI) may not be the best choice because slow freezing may cause changes in the transcriptome or proteome. RNAlater® has been the fixative of choice for previous transcriptomic experiments on the ISS (Beaulieu et al., 2013; Ferl et al., 2011; Kittang et al., 2010; Paul et al., 2005b; Paul and Ferl 2011; Paul et al., 2013), and has been used to preserve protein in Saccharomyces cerevisiae, Arabidopsis callus, and 12 d-old de-etiolated Arabidopsis seedlings (Ferl et al., 2015; Van Eijden et al., 2013; Zhang et al., 2015). To

Figure 5. Impact of growth time on mean protein yield. Plates containing 0.016 g of seed were grown for 1, 2, or 3 d prior to protein extraction and quantification of total protein. * indicates p<0.03 using a Student’s T-test. n=6 plates of approximately 1,300 seedlings. Error bars represent SEM.

Figure 6. RNAlater™ stabilizes protein yield. Plates containing 0.016 g of seed were grown in darkness for 96 h. Seedlings were either soaked in RNAlater™ for 12 h prior to freezing at -80°C, or subject to immediate protein extraction. Results indicate no significant difference in total protein yield between the two approaches. n=3 plates of approximately 1300 seedlings. Bars represent SEM.
determine if RNAlater® treatment was also able
to preserve protein content, plates were flooded
with RNAlater® and incubated at 23°C for 12 h,
frozen at -80°C for 24 h, and thawed prior to
protein extraction. Compared to samples extracted
immediately after growth, RNAlater®/-80°C
treatment had no significant impact on protein
yield (Figure 6). This suggests that the approach
prevents major protein degradation during
fixation.

DISCUSSION

Large-scale omics approaches to under-
standing space biology are appealing because they
allow collection of a vast amount of data but are
relatively light on space and astronaut-time
requirements. For years, microarray, and later
RNA-seq, has been used to gather transcriptomic
data for plants grown on orbit (Beaulieu et al.,
2013; Ferl et al., 2015; Paul et al., 2005a; Paul et
al., 2012; Paul et al., 2013; Salmi and Roux, 2008;
Salmi et al., 2011a; Salmi et al., 2011b). Unlike
physiological data, analyses of these results can
continue well beyond the spaceflight, and are
often utilized by many researchers to understand
specific problems unrelated to the original, but
relevant to space biology.

To truly understand the physiological impact
of spaceflight, however, gene expression data is
insufficient. Protein levels, and often the
reversible, post-translational modifications of
proteins, are responsible for physiological
outcomes on both a cellular and whole-plant level.
Therefore, a proteomic approach is more
appropriate for determining precisely how plants
respond to growth in microgravity. Using
proteomics, researchers can isolate specific
cellular responses, including protein synthesis,
degradation, and activity-altering modifications
such as phosphorylation.

High-Density Planting Allows Sufficient Tissue
for Proteomic Analysis

There are technical challenges associated with
proteomics experiments aboard the ISS, especially
when coupled with BRIC hardware. Proteomics
requires much more plant material than
transcriptomic approaches. Successful experi-
ments necessitate at least 50-100 g of protein per
replicate (Murad et al., 2011). To generate

enough tissue within the confines of the PDFUs,
we used a dense planting approach, resulting in
over 800 seedlings per 60 mm Petri dish. The
primary concern with this approach was the
buildup of ethylene altering growth, development,
and the proteomic results. To mitigate this
possibility, seedlings were allowed to grow only
72 h after germination induction. This timeline
had minimal impact on the overall protein yield,
but reduced the time the seedlings spent in
crowded conditions. Controls were also run in
crowded conditions. Like transcriptomic
approaches, proteomics relies on comparison of
conditions. Even if ethylene-responsive genes
were upregulated in these conditions, the same
genes would also be activated in the control
samples, negating the impacts. Overall, ethylene-
responsive genes and proteins were not detected
in the transcriptomic or proteomic data (data not
shown), indicating that they were not activated, or
that they were similarly expressed in both
samples.

RNAlater® Fixation Protects Protein Content

Terrestrial experiments normally utilize rapid
freezing in liquid nitrogen to preserve protein
samples, followed by immediate homogenization
or long-term storage at -80°C. However, aboard
the ISS, liquid nitrogen is not an option. Direct
transfer to the -80°C freezer is not desirable
because slow freezing could induce gene and
protein expression in response to temperature
change and ice crystal induced cell lysis.

Traditional fixatives, such as glutaraldehyde, pose
health hazards and are not allowed on board the
ISS.

The preservative RNAlater® has long been
used as a fixative appropriate for on-orbit
transcriptomic experiments, as it works well to
preserve nucleic acids (Drakulovski et al., 2013;
Saito et al., 2011). Other recent work has shown
RNAlater® to also preserve protein in a wide
variety of biological samples (Abbaraju et al.,
2011; Gorokhova et al., 2005; Saito et al., 2011).
RNAlater® contains high levels of quaternary
ammonium sulfates and cesium sulfates, leading
to protein denaturation within samples (Bennike et
al., 2016). This activity prevents sample
degradation by DNase, RNase, and protease, as
well as the activity of protein modifying enzymes.
Results here indicate that overall the quantity and
quality of protein are consistent between RNAlater®-treated and flash-frozen samples. However, this work does not address the efficacy of RNAlater® to limit changes to the proteome.

CONCLUSION

The approach outlined here has been executed successfully to obtain sufficient protein from BRIC samples on the ISS. Based on these results, we recommend allocating three PDFUs per replicate. Through managing seed number and growth time, these methods allow proteomic analysis of Arabidopsis tissue using this specific hardware. Furthermore, manipulation of germination through temperature change, as well as RNAlater® fixation, drastically limit crew time, making this an attractive approach for future proteomic analysis of Arabidopsis mutants requiring only short, etiolated growth.

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REFERENCES

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