

Relevance of the Unfolded Protein Response to Spaceflight-Induced Transcriptional Reprogramming in *Arabidopsis*

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Abstract

Plants are primary producers of food and oxygen on Earth and will likewise be indispensable to the establishment of large-scale sustainable ecosystems and human survival in space. To contribute to the understanding of how plants respond to spaceflight stress, we examined the significance of the unfolded protein response (UPR), a conserved signaling cascade that responds to a number of unfavorable environmental stresses, in the model plant *Arabidopsis thaliana*. To do so, we performed a large-scale comparative transcriptome profiling in wild type and various UPR-defective mutants during the SpaceX-CRS12 mission to the International Space Station. We established that orbital culture substantially alters the expression of hundreds of stress-related genes compared with ground control conditions. Although expression of those genes varied in the UPR mutants on the ground, it was largely similar across the genotypes in the spaceflight condition. Our results have yielded new information on how plants respond to growth in orbit and support the hypothesis that spaceflight induces the activation of signaling pathways that compensate for the loss of UPR regulators in the control of downstream transcriptional regulatory networks. Key Words: Spaceflight—Unfolded protein response—*Arabidopsis*—Microgravity. *Astrobiology* 21, 367–380.

1. Introduction

EXTRATERRESTRIAL HABITATION AND prolonged space travel require successful plant growth to recreate livable environments for humans (Ferl *et al.*, 2002; Massa *et al.*, 2016; Zhou *et al.*, 2019). Studies over the past 70 years have sought to develop a better understanding of how plants are affected by and adapt to the significant stresses imposed by spaceflight (*e.g.*, microgravity, radiation, vibration, limited exchange of gases), which can affect plant development and yield (Paul *et al.*, 2013). Recent iterations of the sophisticated chamber hardware for plant growth housed on the International Space Station (ISS) have allowed for multigenerational plant growth in space and analyses of plant responses to this environment (Massa *et al.*, 2016). However, these facilities are insufficient for large-scale plant growth on extraterrestrial environments due to their size and resource cost. Therefore, generation of germplasm adapted to stresses experienced during growth in extraterrestrial environments is a critical contribution to the realization of sustainable plant cultivation in space.

The unfolded protein response (UPR) is a signaling cascade that responds to a number of unfavorable environmental and cellular stresses. The UPR is generally activated by a buildup of unfolded proteins in the endoplasmic reticulum (ER), a condition known as ER stress (Ron and Walter, 2007). The ER stress sensors conserved across metazoans and plants include the ER-associated protein kinase and ribonuclease inositol-requiring enzyme 1 (IRE1) and ER membrane-tethered transcription factors (TFs) (metazoan activating transcription factor 6 [ATF6] and plant basic leucine zipper 17 [bZIP17] and bZIP28). Activation of IRE1 leads to unconventional splicing of an intron from the mRNA of an IRE1-downstream bZIP-TF (metazoan X-box binding protein 1 and plant bZIP60). The UPR TFs are translocated to the nucleus to control expression of UPR target genes and restore ER homeostasis (Koizumi *et al.*, 2001; Chen and Brandizzi, 2012; Halbleib *et al.*, 2017; Kim *et al.*, 2018; Ruberti *et al.*, 2018; Tam *et al.*, 2018; Mishiba *et al.*, 2019; Pastor-Cantizano *et al.*, 2019; Pu *et al.*, 2019). Insufficient UPR leads to the actuation of cell death (Ron and Walter, 2007; Walter and Ron, 2011).

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In terrestrially grown plants, the UPR is a key mediator of responses to a variety of stresses, including heat, pathogen, and high light/singlet oxygen (Deng *et al.*, 2011; Moreno *et al.*, 2012; Guillemette *et al.*, 2014; Zhang *et al.*, 2015, 2017; Pastor-Cantizano *et al.*, 2019; Beaugelin *et al.*, 2020). Additionally, analyses of higher order UPR mutants have demonstrated that the UPR regulators are necessary for post-embryonic growth and reproductive development in *Arabidopsis* under unstressed conditions as well (Chen and Brandizzi, 2012; Kim *et al.*, 2018; Mishiba *et al.*, 2019; Pu *et al.*, 2019). Therefore, a better understanding of the UPR can enable efforts to potentiate plant stress responses and improve plant yield.

Given the broad responsiveness of the UPR to environmental stresses, we hypothesized that the UPR effectors could coordinate gene expression reprogramming in spaceflight stress conditions. To test this hypothesis, we analyzed global gene expression changes in wild type (WT) *Arabidopsis* as well as loss-of-function mutants of IRE1 (*ire1a ire1b*, herein dubbed *ire1*), bZIP28, and bZIP60 (single and double mutants: *bzip28*, *bzip60*, *bzip28 bzip60*), cultivated in orbit during the SpaceX CRS12 mission to the ISS. We used these genetic backgrounds to identify genes controlled jointly or specifically by the UPR sensors and UPR TFs and define the extent to which the known signaling pathways of the UPR functionally interact in a whole organism under microgravity-associated conditions. We established that, in space and on ground, gene expression undergoes a substantial reprogramming on a genome-scale. Growth in orbit substantially altered the expression of thousands of genes associated with significant biological traits compared with ground controls. However, while many of these spaceflight-responsive genes were regulated uniquely in certain UPR mutants compared with WT in the ground control, such a genotype-specific regulation was not observed in the spaceflight condition. These observations not only provide new insight into how plants respond to spaceflight, but also establish that spaceflight-induced transcriptional responses mitigate the need for the gene-regulatory networks controlled by the UPR sensors.

2. Materials and Methods

2.1. Launch hardware and experimental timeline

This flight experiment utilized 4 Biological Research In a Canister (BRIC) containing a total of 22 Petri Dish Fixation Unit (PDFU) hardware (Wells *et al.*, 2001) to cultivate sterile dark-grown seedlings germinated aboard the ISS for a 14 day period. PDFU actuation chambers were loaded with a tissue fixative (RNAlater; Invitrogen) to preserve samples at the conclusion of the flight experiment. An identical set of samples was prepared and grown on Earth with a 2-day offset at Kennedy Space Center (KSC) ISS Environmental Simulator to allow for data transmission and reproduction of incubation conditions experienced by flight samples in orbit. HOBO data loggers equipped with temperature sensors were integrated into two of the four BRICs to record temperatures experienced by samples during the experiment for *post hoc* analysis. Launch samples (*i.e.*, *Arabidopsis* seeds) were integrated into BRIC flight hardware in a sterile hood ~48 h before the August 14, 2017, launch of SpaceX CRS12 spacecraft. Integrated science/hardware was kept at 4°C to

maintain seed dormancy before packing in cold storage bags while being loaded onto Dragon capsule and during launch. After docking, samples were removed from cold storage bags by ISS astronauts, warmed to ambient ISS temperature, allowing seed germination and experiment initiation in the BRIC. After 14 days, ISS astronauts actuated PDFUs, which were then incubated at room temperature for a further 3 h before being transferred to the ISS –80 MELFI freezer. Samples were kept at approximately –80°C until BRICs were conditioned to –32°C in double cold bag storage (Hutchison and Campana, 2009) and stowed in the Dragon capsule before undocking and atmospheric reentry on September 16. After returning to KSC, samples were stored at –80°C until de-integration of the flight and ground samples. De-integration occurred on November 1, 2017.

2.2. Germplasm and culture conditions

Arabidopsis thaliana seeds of the following genotypes were used for flight and ground controls: WT (Col-0 ecotype), *atire1* (Nagashima *et al.*, 2011; Chen and Brandizzi, 2012), *bzip60* (Moreno *et al.*, 2012), *bzip28* (Gao *et al.*, 2008), and *bzip28 bzip60* (Deng *et al.*, 2013). Petri dishes (60 mm) were prepared with 6.7 mL of sterile ½ Murashige and Skoog media supplemented with Gamborg's B5 Vitamins (PhytoTechnology Laboratories), 0.5% sucrose (Sigma-Aldrich), 0.4% Phytigel (Sigma-Aldrich), pH adjusted to 5.7. In a sterile hood, seeds of WT and UPR-mutant genotypes were surface sterilized with one wash of 70% ethanol, one wash of 50% bleach containing 0.5% Tween 20, and then nine additional washes with sterile H₂O distilled twice. After the final wash was removed, seeds were resuspended in 1.5 mL sterile water for wet plating by using a 1 mL pipette equipped with sterile filter tip. For each experimental unit (flight and ground control), five plate replicates of WT and *bzip28/bzip60* genotypes and four plate replicates of *atire1*, *bzip28*, and *bzip60* genotypes were prepared. Each plate replicate contained 70–80 seeds evenly spaced in a grid pattern on the plate surface. The Petri dishes were sealed with Parafilm (Heathrow Scientific), and individual plates were then wrapped twice with sterile aluminum foil. Individually wrapped plates were grouped by BRIC configuration and wrapped together with two more layers of sterile aluminum foil before sample removal from the sterile hood. Plates were placed at 4°C until the integration of samples into science hardware the following morning.

2.3. Sample processing and experimental material assessment

Samples of flight and ground control experiments were preserved in RNAlater *in situ* and kept at –80°C (see the Results section for experimental timeline). The Petri dishes were removed from packaging and thawed in groups of three to prevent excess exposure to room temperature during sample collection. After removing most of the RNAlater from the plates, sterile forceps were used to transfer seedlings from the plates to microcentrifuge tubes containing two glass beads. Seedlings were then frozen in liquid nitrogen. This procedure was carried out quickly to maximize RNA recovery. Accordingly, only some pictures were taken of plates before extraction for example purposes. Most pictures were taken after the bulk of the sample was

removed, with the remaining seedlings also imaged for *post hoc* inspection. All plates were free from any visible evidence of bacterial or fungal contamination. Frozen samples were ground to a powder with a Retsch Mixer Mill (Retsch, Haan, Germany). RNA was extracted from tissues by using a NucleoSpin RNA Plant Kit (Macherey-Nagel) according to the manufacturer's instructions including DNase digestion. The overall quality and RNA integrity number (RIN) of RNA samples were assessed with Agilent Bioanalyzer 2100 (Agilent, Santa Clara, CA).

2.4. Library preparation, sequencing, and bioinformatics analysis

RNA-sequencing (RNA-seq) libraries were constructed by using the Illumina TruSeq Stranded mRNA Library (Illumina, San Diego, CA) and sequenced in single-end mode on the Illumina HiSeq 4000 platform (50-nt) at Research Technology Support Facility Genomics Core at Michigan State University. For each library, read quality was assessed with the FastQC (version 0.11.3) software (<https://www.bioinformatics.babraham.ac.uk/projects/fastqc>). Reads were cleaned for quality and adapter sequences with Cutadapt (version 1.8.1) using a minimum base quality 20 retaining reads with a minimum length of 30 nucleotides after trimming (Martin, 2011). Quality-filtered reads were aligned to the Col-0 reference genome (TAIR10) with Bowtie (version 2.3.1) and TopHat (version 2.1.1) with a 10 bp minimum intron length and 15,000 bp maximum intron length (Langmead and Salzberg, 2012; Kim *et al.*, 2013). Fragments per kilobase exon model per million mapped reads (FPKM) were measured by using TAIR10 gene model annotation with Cufflinks (version 1.3.0) (Trapnell *et al.*, 2010). The log₂ transformed and normalized gene expression levels (FPKM +1) were used for correlation analysis (Spearman's rank correlation coefficient) between biological replicates and principal component analysis (PCA). Per-gene read counts were identified with HTSeq (version 0.6.1p1) in the union mode with a minimum mapping quality of 20 with stranded reverse counting (Anders *et al.*, 2015). Differential gene expression analysis was performed in four biological replicates (for WT and *bzip28 bzip60*, selected based on the correlation with other biological replicates) with DESeq2 (version 1.16.1) within R (version 3.4.0) based on a comparison of spaceflight to ground with adjusted *p*-value <0.01 and absolute log₂-transformed fold change >1.5 (Love *et al.*, 2014). Genes of which the total count across all samples is <100 were not included in the analysis. Gene Ontology (GO) overrepresentation was performed with PANTHER (Fisher's exact type with false discovery rate correction) (www.pantherdb.org) (Mi *et al.*, 2019a, 2019b). K-means clustering analysis on average FPKM values from biological replicates was performed using the Morpheus tool (Morpheus, <https://software.broadinstitute.org/morpheus>). The optimal number of K-means clusters was determined with the factoextra package in R.

3. Results

3.1. Spaceflight alters the growth of seedlings independent of an intact UPR signaling

When we inspected the WT and UPR-mutant seedlings (*atire1*, *bzip60*, *bzip28*, and *bzip28 bzip60*) of the ground control and flight samples at the completion of the mission,

we found that in the ground control samples, etiolated hypocotyls (*i.e.*, pale and elongated due to the lack of light) were above the surface of the solidified media while roots had penetrated the growth medium perpendicular to the surface (Fig. 1A). However, in flight samples, we found that etiolated hypocotyls as well as roots had generally penetrated the growth medium regardless of genotype tested (Fig. 1A). Interestingly, we also observed that cotyledon petioles of flight sample seedlings were elongated (Fig. 1B) compared with ground control seedlings (Fig. 1A). Overall, these observations are consistent with plant growth in the darkness and space, conditions leading to elongated hypocotyls and petioles, and a lack of directional growth, respectively (Paul *et al.*, 2017). These observations also suggest that the UPR unlikely exerts a noticeable role in growth direction in response to altered gravity levels.

3.2. Spaceflight results in an increase of total RNA

RNA degradation has previously been observed in independent *Arabidopsis* spaceflight experiments performed in BRIC-PDFUs (Paul *et al.*, 2012; Johnson *et al.*, 2017). To test the RNA quality of our samples, we measured RIN as an indicator of overall RNA quality (RIN, 1=low quality; 10=high quality) of each sample and compared size peaks of 25S and 18S ribosomal RNA (rRNA) among samples (Mueller *et al.*, 2004). Note that these measurements include smaller plastid rRNA peaks, which lower the maximum RIN value to around 8, independent of RNA quality (Babu and Gassmann, 2016). We established that all flight samples had RIN values between 7.5 and 8.0, indicating that RNA was of high quality. For ground samples, RIN values were found to be between 4.0 and 7.7, which would ordinarily indicate mild degradation of some samples. However, closer analyses revealed that, in nearly all ground samples (77%), the RIN algorithm failed to identify the correct peaks. In these samples, the algorithm identified the 18S rRNA peak as the 25S rRNA peak, and a putative organelle rRNA peak (Babu and Gassmann, 2016) as the 18S rRNA (Supplementary Fig. S1). Therefore, to compare RNA quality between treatments and genotypes from the Bioanalyzer data outputs, we used the ratio of 25S/18S peak heights as a substitute measure. Because the 25S peak height is reduced more quickly than the 18S peak in RNA-degrading conditions (*e.g.*, elevated temperature, exogenous RNases, endogenous apoptotic RNase activity) (Mueller *et al.*, 2004; Babu and Gassmann, 2016), a decreased 25S/18S ratio would indicate RNA degradation. We observed no significant differences of the 25S/18S ratio across all genotypes (Fig. 2A); therefore, both flight and ground samples had no significant RNA degradation.

Interestingly, by comparing the 25S peak height with one of the two other peaks near the 18S peak [*i.e.*, a putative organelle rRNA peak, designated 18S (-3); Supplementary Fig. S1] in each sample, it was clear that the relative ratio of 25S to 18S (-3) was significantly lower in the ground samples compared with flight samples (Fig. 2B and Supplementary Fig. S1). This low ratio indicates that the ground samples were depleted of nuclear-encoded rRNAs (*i.e.*, 25S and 18S rRNAs) compared with other RNA species. This observation is also consistent with our findings that our ground samples contained significantly less total

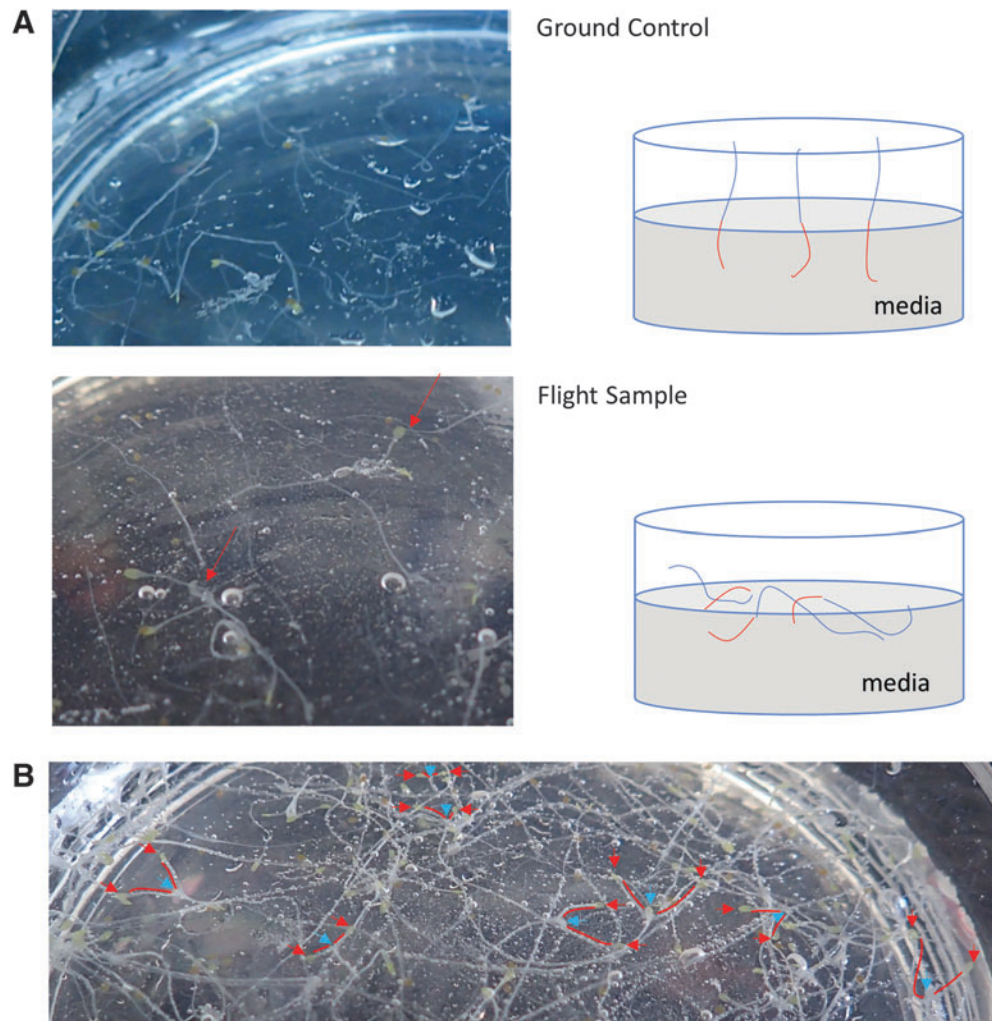


FIG. 1. (A) Representative ground and flight sample plates were imaged after the bulk of etiolated hypocotyls, and RNAlater was removed to ensure maximal RNA integrity. Remaining seedlings were used for *post hoc* analysis of morphological/growth differences. (B) Left: Example plate of a WT flight sample, which was imaged after thawing, but before seedling removal. Individual seedlings that could be distinguished from the bulk were marked with a blue arrow to mark the shoot meristem. The cotyledons of elongated petioles were marked with a red arrow and a red line used to connect the cotyledon to the meristem of the same seedling. Petioles were elongated compared with ground sample petioles [representative morphology of ground control presented in (A)]. WT, wild type.

RNA [of which rRNA is a significant fraction (Lodish, 2000)] than flight samples (Fig. 2C). Because we extracted RNA from a similar number of seedlings from flight samples and ground samples, our results indicate that flight samples contained larger amount of total RNA compared with ground samples, which is likely to be caused by elevated levels of nuclear encoded rRNA in flight samples, and unlikely to be due to RNA degradation of the ground samples.

3.3. Global transcriptomic analyses indicate that gene expression reprogramming in response to spaceflight depends partially on intact UPR signaling

Having established that the total RNA from ground and flight samples was of acceptable quality, we next proceeded to RNA-seq to investigate the impact of spaceflight on global gene expression changes in the UPR mutants. In

RNA-seq library preparation, mRNA was enriched by purification to efficiently remove rRNA (Zhao *et al.*, 2018) and mitigate potential sequencing bias due to higher rRNA levels in flight samples. We obtained an average of ~ 32 million reads per sample, of which 95–99% were successfully mapped to the *Arabidopsis* reference genome (Supplementary Fig. S2). Spearman's rank correlation coefficients calculated between biological replicates showed a high reproducibility of our RNA-seq data set (Fig. 3A). Furthermore, PCA exhibited a strong separation of ground samples from flight samples (Fig. 3B) and indicated that ground samples located more closely to each other than the flight samples. Overall, these analyses further supported a statistical robustness of the RNA-seq and justified further investigation.

To investigate gene expression changes in response to spaceflight, we identified differentially expressed genes (DEGs) in each genotype by comparing gene expression values in ground and flight samples (Supplementary Table S1). A total

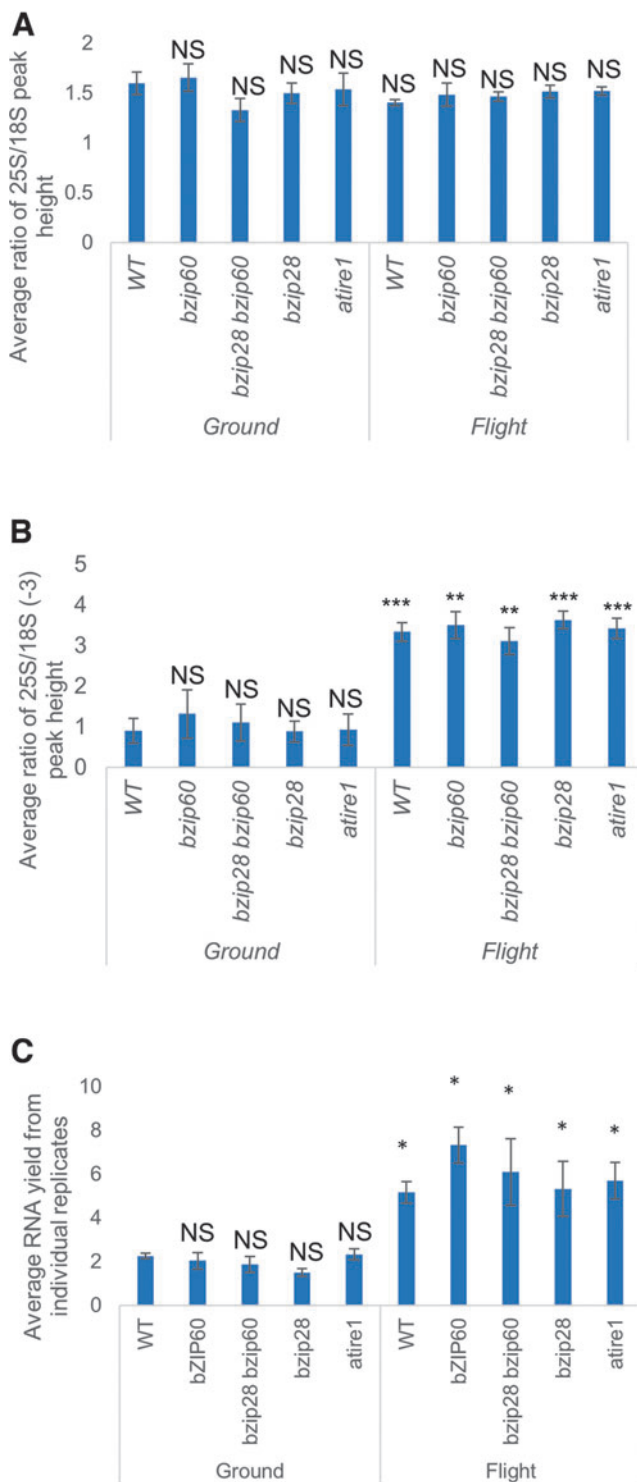


FIG. 2. (A) Average ratio of 25S/18S peak heights as determined from Bioanalyzer traces from each sample was used as a secondary measure of RNA quality due to the RIN algorithms incorrect identification of the appropriate peaks. (B) The relative content of rRNA found in each genotype in both conditions was determined by comparing the height of the 25S peak with an organellar rRNA peak [18S (-3)], which was found in each sample. (C) Average RNA yields from each genotype from flight and ground samples. Statistical significance determined by Welch's *t*-test, *p*-value represented by not significant (NS) = >0.05; * = <0.05; ** = <0.005; *** = <0.0005. RIN, RNA integrity number.

of 3465 genes were classified as DEGs in at least one of the tested genotypes. WT had the largest number of DEGs (up-regulated DEGs in flight compared with ground, $n=1675$; downregulated DEGs, $n=831$) among the genotypes tested. The *bzip28 bzip60* mutant had the smallest number of DEGs (upregulated DEGs, $n=1293$; downregulated DEGs, $n=562$) (Fig. 4A, B). In all genotypes, the number of upregulated DEGs was higher than that of downregulated DEGs (WT, 2.02-fold; *atire1*, 1.98-fold; *bzip28*, 2.28-fold; *bzip60*, 1.80-fold; *bzip28 bzip60*, 2.30-fold), indicating a higher impact of spaceflight on inducing gene expression rather than suppressing it. While the identity of 34.8% (783/2249) of upregulated DEGs and 27.5% (335/1217) of downregulated DEGs overlapped across all genotypes, relatively smaller numbers of DEGs were found to be genotype-specific, ranging from 30 (downregulated exclusively in *bzip28 bzip60*) to 207 (upregulated in WT).

In summary, based on the verified number of upregulated and downregulated DEGs in flight samples compared with ground samples across genotypes, the UPR mutants had consistently fewer overall DEGs than WT, indicating that the UPR could play at least a partial role in regulating the transcriptional reprogramming in space compared with ground control.

3.4. Biological pathways connected to the DEGs between flight and ground

To gain insights in the biological pathways altered in spaceflight in our experimental setup, we performed separate GO analyses on upregulated and downregulated DEGs in WT (Supplementary Table S2), generating a list of parental-GO terms (more general, represented by a larger number of genes in the reference gene set) and cognate child-GO terms (more specific, smaller numbers of genes in the reference gene set) (Fig. 5). Intriguingly, we found that stress-responsive genes (*e.g.*, “response to abscisic acid,” “response to hypoxia,” “response to water deprivation,” and “response to oxidative stress”) as well as genes involved in physiological responses often associated with stress response adaptation were enriched in downregulated DEGs in WT. This result is consistent with previous studies that reported downregulation of water stress-related genes when using *Arabidopsis* BRIC-PDFU microarray transcriptomes (Johnson *et al.*, 2017) and that found abscisic acid response and water stress response overrepresentation in misregulated DEGs in Col-0 WT with RNA-seq (Choi *et al.*, 2019). In addition, we observed that metabolic processes associated with stress adaptation (Hildebrandt Tatjana *et al.*, 2015; Batista-Silva *et al.*, 2019), including amino acid catabolism and sucrose starvation response, were overrepresented in flight-downregulated DEGs. Ribosome biogenesis, translation, and gene expression processes were highly underrepresented in this category, that is, they were more likely to be upregulated by flight or remain unchanged. By further analyzing the normalized gene expression values (FPKM), we also found that ribosome biogenesis and rRNA processing GO terms appeared significantly overrepresented in genes upregulated by >2-fold changes (flight/ground). These, however, were not considered as DEGs based on the strict statistical criteria applied in our analyses (see the Materials and Methods section). The lower FPKMs of

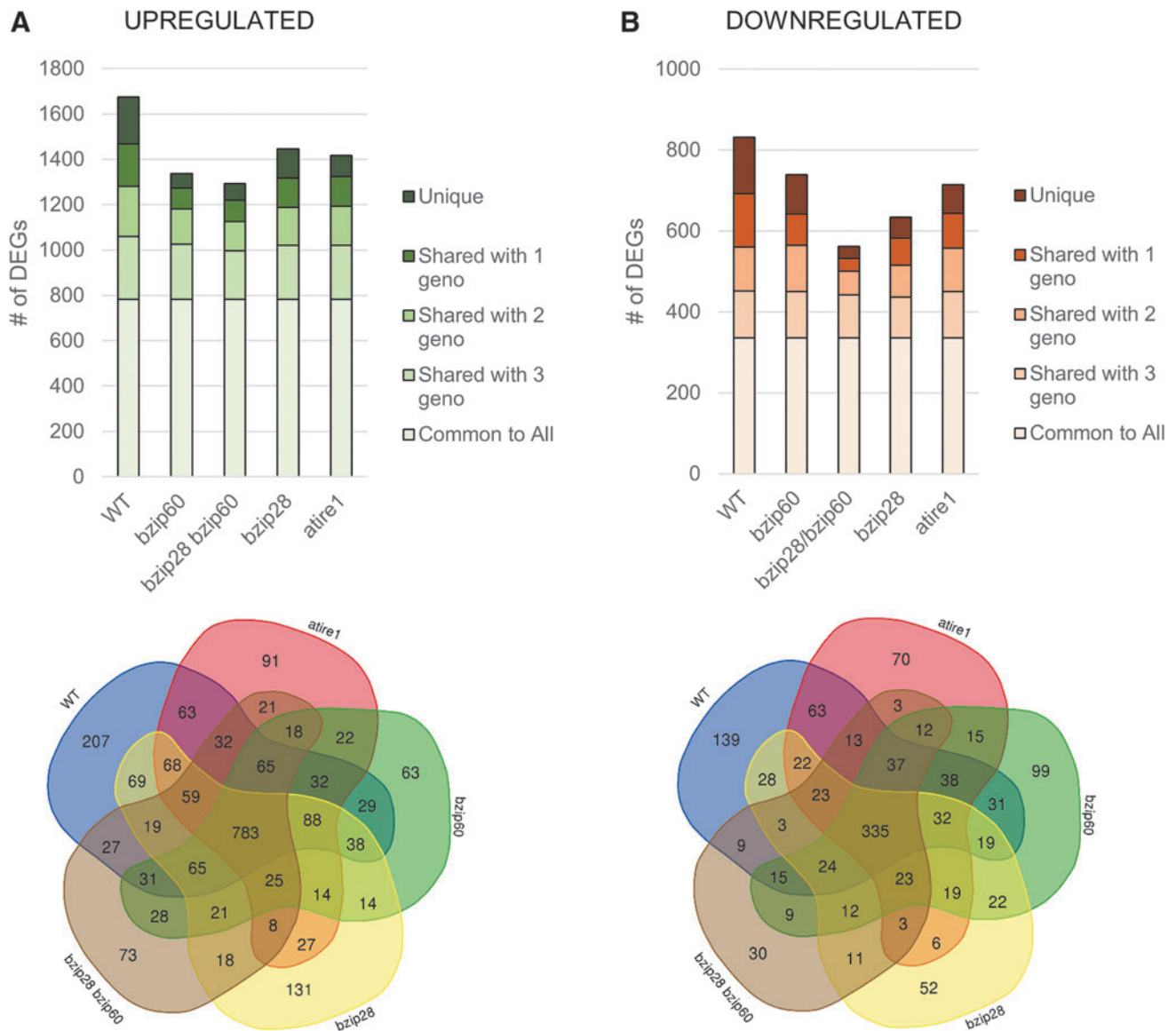


FIG. 4. Differential expression analysis using the four biological replicates with the highest correlation was performed via HTseq v0.6.1pl and DESeq v2. For each genotype and for genes, DEGs in flight samples relative to ground samples were determined using a strict criterion: adjusted *p*-value <0.01; |log₂FC| >1.5. Total number of upregulated (**A**) and down-regulated (**B**) DEGs in each background was analyzed to determine what proportion was shared between the different genotypes. DEG, differentially expressed gene.

In Cluster 2 (DEGs; *n*=546), expression of DEGs was induced exclusively in ground *bzip60* compared with other ground genotypes while highly suppressed to similar levels of expression in flight samples of all genotypes. As such, the proportion of DEGs in *bzip60*, whose expression was significantly different from WT, was much higher (17%) than other genotypes (*bzip28 bzip60*, 0.9%; *bzip28*, 1.0%; *atire1*, 1.2%) (Supplementary Fig. S4). We reasoned that bZIP28 and bZIP60 could have a negative feedback relationship in which the absence of bZIP28 suppressed the effect of *bzip60* mutation exclusively in the ground condition. The over-represented GO terms in Cluster 2 were largely similar to the GO terms enriched in the downregulated DEGs genes in the WT genotype: “response to abscisic acid,” “response to water deprivation,” and “response to hypoxia” (Fig. 5). In

addition, Cluster 2 DEGs showed significant enrichment of GO terms associated with biotic stress responses that were not found in analyses of the WT genotype (Fig. 5) or found to be strongly enriched in the any of the other clusters (Fig. 6 and Supplementary Table S4). These results indicate that bZIP60 may have repressive roles in regulating genes involved in both abiotic and biotic stress responses.

The gene expression pattern of Cluster 3 (DEGs; *n*=274) was also characterized largely by genes with lower expression values in spaceflight compared with ground control across genotypes, except for *bzip28*. However, contrasting with Cluster 2, the FPKM values in the *bzip60* genotype was not different from WT FPKM values; only 3.2% of DEGs showed significant different FPKM compared with WT (Supplementary Fig. S4). Instead, the absence of bZIP28

DNA replication, DNA repair	# in Ref.	# in DEGs	Expected	Fold Enrichment	FDR
mitotic DNA replication initiation	4	4	0.24	16.81	3.08E-02
cell cycle	486	76	28.91	2.63	4.06E-09
double-strand break repair via break-induced replication	12	8	0.71	11.21	7.31E-04
double-strand break repair via homologous recombination	81	14	4.82	2.91	4.05E-02
DNA replication	124	31	7.38	4.2	5.54E-07
Far Red/Red Light Response	# in Ref.	# in DEGs	Expected	Fold Enrichment	FDR
response to far red light	51	15	3.03	4.94	3.99E-04
response to red or far red light	197	26	11.72	2.22	2.17E-02
response to light stimulus	680	74	40.45	1.83	5.04E-04
response to radiation	705	75	41.94	1.79	6.65E-04
Photosynthesis	# in Ref.	# in DEGs	Expected	Fold Enrichment	FDR
photosynthetic electron transport in photosystem I	17	8	1.01	7.91	3.76E-03
photosynthesis, light reaction	122	31	7.26	4.27	5.93E-07
generation of precursor metabolites and energy	338	42	20.11	2.09	3.54E-03
photosynthesis, light harvesting in photosystem I	23	9	1.37	6.58	3.86E-03
photosynthesis, light harvesting	44	15	2.62	5.73	1.54E-04
Secondary Metabolite Biosynthesis	# in Ref.	# in DEGs	Expected	Fold Enrichment	FDR
glucosinolate metabolic process	112	22	6.66	3.3	6.59E-04
glucosinolate biosynthetic process	39	16	2.32	6.9	1.42E-05
secondary metabolite biosynthetic process	147	22	8.75	2.52	1.65E-02

Abscisic Acid, Response to Stress	# in Ref.	# in DEGs	Expected	Fold Enrichment	FDR
response to abscisic acid	524	61	15.16	4.02	5.26E-16
response to water deprivation	340	48	9.83	4.88	2.58E-15
cellular response to hypoxia	234	46	6.77	6.8	2.91E-19
response to oxidative stress	392	41	11.34	3.62	3.04E-09
response to stress	3079	200	89.06	2.25	4.75E-24
Amino Acid Catabolism, Sucrose Starvation	# in Ref.	# in DEGs	Expected	Fold Enrichment	FDR
tyrosine catabolic process	5	3	0.14	20.74	4.01E-02
alpha-amino acid catabolic process	65	11	1.88	5.85	7.03E-04
leucine catabolic process	7	4	0.2	19.76	9.01E-03
branched-chain amino acid catabolic process	18	7	0.52	13.44	3.87E-04
cellular amino acid catabolic process	73	15	2.11	7.1	3.02E-06
isoprenoid catabolic process	23	5	0.67	7.52	3.71E-02
lipid catabolic process	108	11	3.12	3.52	2.33E-02
cellular response to sucrose starvation	4	4	0.12	34.57	2.55E-03
Absence of Light	# in Ref.	# in DEGs	Expected	Fold Enrichment	FDR
response to absence of light	44	13	1.27	10.21	7.39E-07
response to light intensity	142	19	4.11	4.63	1.86E-05
response to abiotic stimulus	2062	158	59.64	2.65	9.37E-25
Ribosome Biogenesis, Translation	# in Ref.	# in DEGs	Expected	Fold Enrichment	FDR
ncRNA metabolic process	407	1	11.77	0.08	1.08E-02
ribosome biogenesis	351	1	10.15	0.1	3.57E-02
ribonucleoprotein complex biogenesis	422	2	12.21	0.16	3.57E-02
translation	431	1	12.47	0.08	5.79E-03
RNA metabolic process	1227	6	35.49	0.17	2.94E-07
gene expression	1403	6	40.58	0.15	4.86E-09
cellular protein metabolic process	2699	43	78.07	0.55	8.55E-04

FIG. 5. Representative biological processes gene ontologies (GO terms) over- or underrepresented by upregulated or downregulated DEGs in the WT background. GO, Gene Ontology.

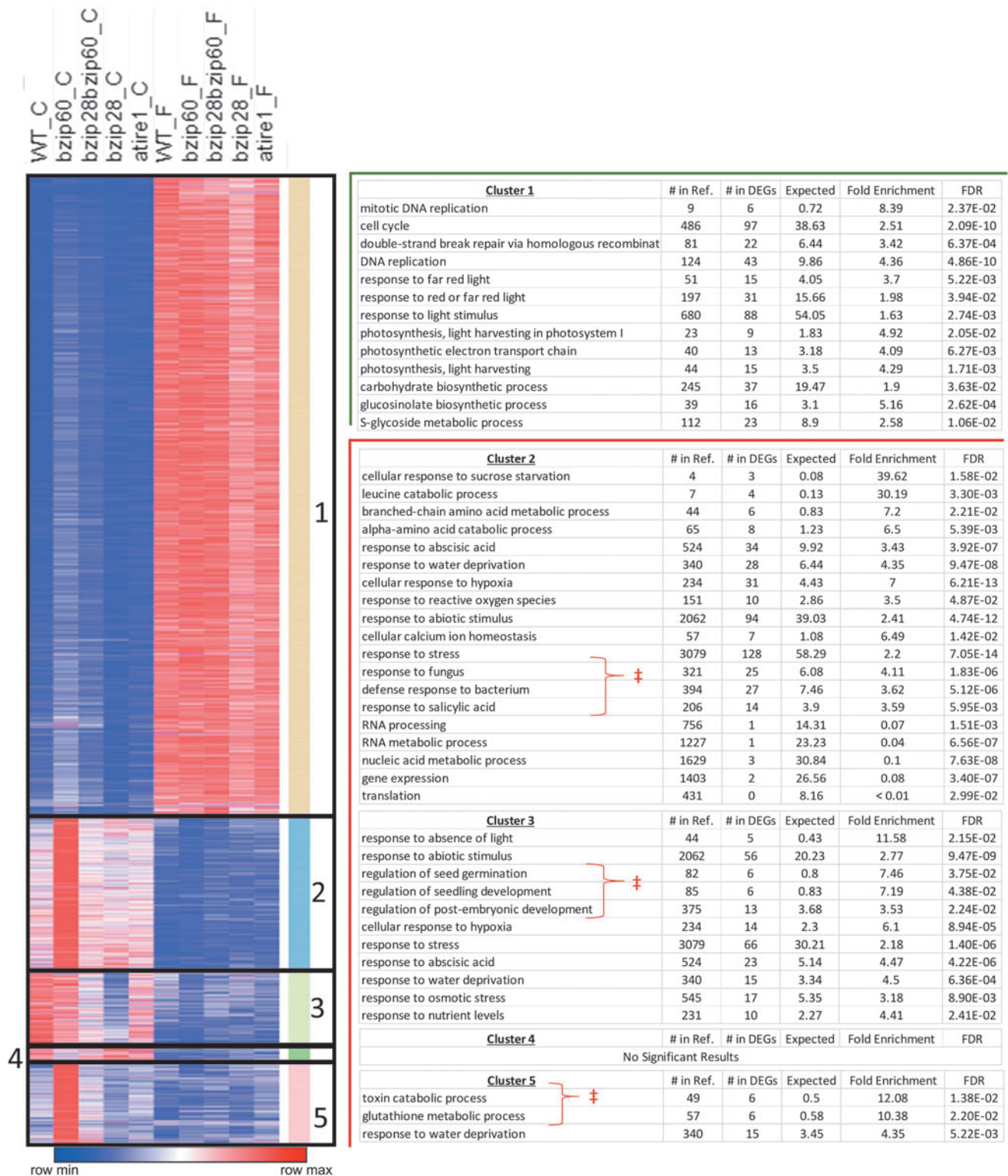


FIG. 6. K-means clustering analysis of all 3465 DEGs in at least 1 background. Each row represents fragments per kilobase exon model per million mapped reads values of an individual gene averaged between biological replicates. For each row, blue represents the minimum relative expression value, red the maximum expression value, and white is the middle value. For each cluster, GO biological process analysis of DEGs demonstrates a role for UPR regulation of ground control stress responses, which are largely repressed by flight. UPR, unfolded protein response.

(i.e., in the *bzip28* mutant) had a higher impact on gene expression in the ground condition compared with other mutants (Supplementary Fig. S4). Interestingly, these Cluster 3 DEGs had intermediate FPKM values in the *bzip28 bzip60* genotype in ground control compared with the extremes of the *bzip60* and *bzip28* single mutants, indicating an antagonistic regulation of bZIP28 and bZIP60 on these genes in the ground condition, which was largely compensated for in the *bzip28 bzip60* genotype.

Similar to Clusters 2 and 3, Cluster 4 (DEGs; $n=55$) contained genes, whose expression exhibited overall lower FPKMs in spaceflight compared with ground across genotypes. However, Cluster 4 showed a unique pattern: the expression of the DEGs in this cluster was significantly lower in *bzip60* compared with WT and the other genotypes in the ground condition (Supplementary Fig. S4) and showed no prominent differences across genotypes in the spaceflight condition. Although Cluster 4 was not significantly represented by any biological process GO terms, a closer analysis revealed that 22% of all DEGs in this cluster were encoded on the mitochondrial genome; mitochondria-encoded genes comprise only 0.5% of all protein-coding genes in the *Arabidopsis* genome annotation (<https://www.arabidopsis.org>) and 0.3% of DEGs called in this study.

Cluster 5 (DEGs; $n=291$) showed a gene expression pattern similar to Cluster 2 with significantly higher levels in the *bzip60* genotype compared with the other genotypes in the ground control samples (Supplementary Fig. S4). In this Cluster, the DEGs were more affected by the absence of bZIP28 (*bzip28 bzip60* and *bzip28*) in the ground condition compared with Cluster 2. The GO term “response to water deprivation,” which was found to be enriched in Cluster 2, was significantly enriched in Cluster 5. In addition, relatively narrow child GO terms “toxin catabolic process” and “glutathione metabolic process,” which were not enriched in other clusters, were enriched in Cluster 5.

Overall, by comparing the *bzip60* and *bzip28* single mutants with the *bzip28 bzip60* double mutant, our transcriptomic profiling provides evidence for a highly complex, unconventional, regulatory relationship between bZIP60 and bZIP28 under the conditions experienced by ground control seedlings. Furthermore, a small number of significant differences between WT and the UPR mutants in spaceflight were found, supporting a small but significant role of the UPR in gene expression in spaceflight.

4. Discussion

In this study, we examined the transcriptional responses of WT and seedlings with a compromised UPR to spaceflight to set the foundations to manipulate a critical growth and stress signaling pathway for improving plant adaptation to extraterrestrial environments. We utilized the BRIC-PDFU sterile plant culture hardware during the SpaceX CRS12 mission to compare the transcriptional responses with the spaceflight between *A. thaliana* WT and mutants defective in one or more components of the UPR, namely the TFs bZIP60, bZIP28, and the ER-resident kinase/ribonuclease IRE1.

The BRIC-PDFU hardware has been employed in a number of dark-grown *Arabidopsis* transcriptome experi-

ments toward different aims (Paul *et al.*, 2012; Kwon *et al.*, 2015; Johnson *et al.*, 2017; Choi *et al.*, 2019). Variability in technical experimental details and limited overlap between spaceflight/ground DEGs has been verified even between the same WT control genotype in simultaneous experiments (Johnson *et al.*, 2017). However, some broad biological pathways have been found to be induced or repressed in response to spaceflight, including cell wall modification, response to light/high light, oxidative stress, osmotic stress response, heat shock, and biotic defense/secondary metabolite synthesis (Paul *et al.*, 2012; Johnson *et al.*, 2017; Choi *et al.*, 2019). Many of these responses were also noted in our study, including the downregulation of water stress response in space, which has been identified in four separate experiments (Johnson *et al.*, 2017; Choi *et al.*, 2019). However, we also observed correlative differences in seedling growth and an overall gene expression landscape not noted in previous studies. Our ground control seedlings had grown in a predictable manner, consistent with the expected morphology of terrestrially grown *Arabidopsis* etiolated hypocotyls (Fig. 1). Flight sample growth was also largely in line with expectations for seedlings grown in microgravity, including the observed petiole elongation, which was shared in all flight samples. Indeed, a petiole elongation of flight samples was also present in the images published of dark-grown *Arabidopsis* experiments in the Col-0 background in previous experiments (Johnson *et al.*, 2015; Paul *et al.*, 2017).

This growth phenotype is consistent with low R/FR ratio and shade avoidance syndrome mediated by phytochrome signaling (Franklin, 2008). The correlative responses observed in the transcriptome analysis (Fig. 4) support that differentially regulated growth phenotypes and the large transcriptional rearrangements in flight during our experiment might have been mediated by phytochrome-related signaling, which is also known to constitutively repress abscisic acid signaling (Yang *et al.*, 2016), and were found to be repressed in our flight samples. However, these differences in the response to light and the increased expression of photosynthetic components between ground and flight samples are anomalous when considering the spaceflight culture methods used in this study. BRIC-PDFUs are autoclavable, black polymer containers, which are sealed with metal lids during science integration, and allow injection of the chemical preservative without opening the unit. After being sealed on August 12, the seeds and seedlings germinated from these seeds in the BRICs were not exposed to light during the launch and the 14-day growth period on the ISS. Therefore, the dichotomous occurrence of the “absence of light” overrepresentation in downregulated and of “response to R/FR light response” in upregulated DEGs is unlikely to be the result of actual differential exposure to light.

One likely explanation is related to the hypocotyl-tissue media contact that occurred in our flight samples, which lacked a clear growth vector in microgravity, but had not occurred in our ground samples where roots grew perpendicular into the media. In previous experiments by Johnson *et al.* (2015) and Paul *et al.* (2017), the dark-grown ground control plates were oriented vertically and both sample sets displayed petiole elongation, although the precise differences in length or extent of petiole elongation between flight and ground were not quantified. How media contact could

induce R/FR light signaling in the dark is not immediately obvious; however, earlier studies showed that media containing sucrose modulated the R/FR signaling mediated by phytochrome A, promoting a red light response (Dijkwel *et al.*, 1997). Coincidentally, the higher rRNA and RNA content observed in flight samples compared with ground control samples in our study (Fig. 2) is also consistent with an increased exposure of cells to sucrose or glucose, which is known to induce RNA accumulation, rRNA transcription, and ribosome biogenesis in plants (Kojima *et al.*, 2007; Ishida *et al.*, 2016), yeast (Kunkel *et al.*, 2019), and mammals (Hannan *et al.*, 2003). Nonetheless, we cannot rule out the remote possibility that our observations may be influenced by possible interactions between ionizing radiation and phytochrome R/FR signaling. For example, low-dose gamma (γ) irradiation of lettuce seeds was found to mimic the effects of FR deactivation of red light-activated phytochromes (Hsiao and Vidaver, 1974). Additionally, a structural study of the bacterial phytochrome

from the radiation-resistant bacteria *Deinococcus radiodurans* established that X-ray radiation induced deprotonation of chromophore in the inactive phytochrome, a biochemical step thought to be involved in light-induced activation of this protein (Li *et al.*, 2015). Although the dose required to deprotonate 50% of the phytochrome (Li *et al.*, 2015) was orders of magnitude larger than that expected to be experienced during our experimental period on the ISS, differences between prokaryotic and eukaryotic phytochromes could affect relevant properties of a hypothetical phytochrome–radiation interaction.

Interpreting a role for the UPR in the transcriptional response to spaceflight is complicated. We observed clear differences in the number of spaceflight DEGs in the UPR-mutant backgrounds compared with WT (Fig. 4); and only 1118 of the 3465 DEGs were common to all genotypes. This would normally suggest that the transcriptional readjustments that occurred in response to spaceflight were at least partially the result of UPR-dependent processes. However,

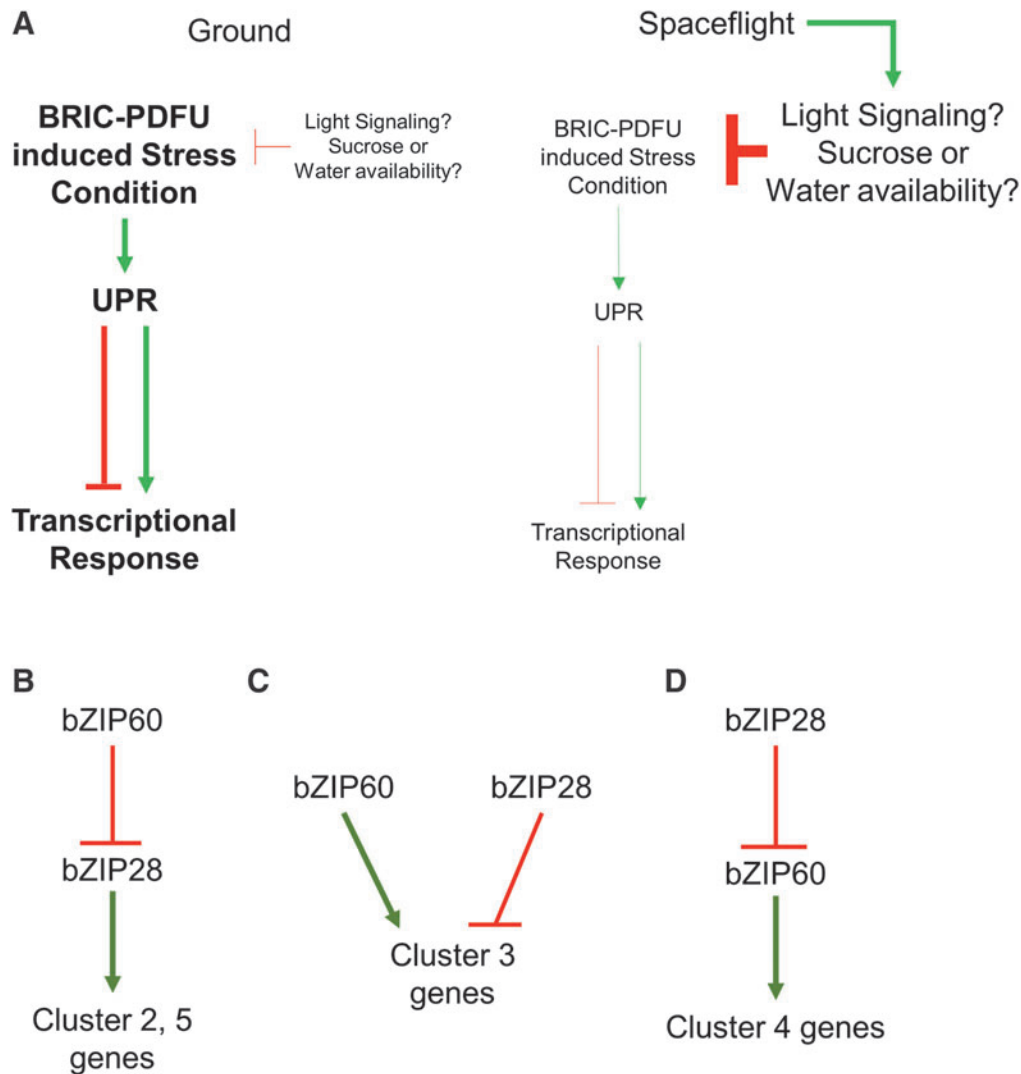


FIG. 7. (A) Hypothesized regulatory framework for stress-responsive DEGs found to be regulated by the UPR on the ground but not in flight conditions. Possible interactions between bZIP60 and bZIP28 in the regulation of DEGs were found in (B) Clusters 2 and 5, (C) Cluster 3, and (D) Cluster 4.

upon closer analysis of the underlying FPKM values, it became clear that the differences in fold change values (flight/ground) across genotypes were more heavily influenced by the differential regulation of expression in the ground samples by the UPR regulators (Fig. 6). The number of DEGs with FPKM values significantly different from WT FPKM values in at least one UPR-mutant genotype was four times greater in the ground samples compared with flight samples (Supplementary Fig. S3). The heat map visualization of these values (Fig. 6) further suggests that the variations in ground samples expression levels were largely muted by spaceflight, as the endpoint transcript levels in flight samples were nearly uniform in the different genotypes. Overall, this would suggest that the UPR does not have a broad involvement in the response to spaceflight. One explanation for this observation may be related to the concerted downregulation of many stress-responsive processes in the flight samples compared with the ground samples (Fig. 6). In spaceflight conditions, it seems likely that alternative signaling pathways are actuated, which repress the observed stress responses regulated by the UPR. Given the prevalence of starvation responses in ground samples, it is possible that microgravity-induced changes in growth habit provide better nutrient availability (Fig. 7). As such, plants in flight may be able to better handle the stresses imposed by culture conditions, without requiring UPR regulator involvement.

Nonetheless, the observations related to an interaction between the UPR regulators bZIP60 and bZIP28 and the stresses imposed on ground control seedlings have yielded important information, which should be explored in the future. In the canonical ER stress response induced chemically or via environmental stress, bZIP60 and bZIP28 TFs interact in the nucleus and direct the actions of the COMPASS DNA methylation complex to increase transcription of target genes (Song *et al.*, 2015). Furthermore, these TFs can also bind independently to gene promoters to activate downstream UPR genes, as evidenced by the weaker activation of ER chaperones in the *bzip28 bzip60* double mutant compared with either of the *bzip28* or *bzip60* single mutants (Ruberti *et al.*, 2018; this work). Although it has been suggested that bZIP60 and bZIP28 may also have unique target genes (Pastor-Cantizano *et al.*, 2019), in our ground control samples, the transcriptomic data suggest that bZIP60 and bZIP28 may have a more complex antagonistic relationship in the control of genes related to the response to abscisic acid, hypoxia, water deprivation, and to oxidative stress (Fig. 6). In Clusters 2 and 5, and Cluster 4, we observed that the expression levels of the DEGs were higher or lower than WT in the *bzip60* genotype, respectively (Fig. 6 and Supplementary Fig. S3). However, these differences were not observed in the *bzip28 bzip60* double mutant as would be expected. Conversely, in Cluster 3, we observed in ground samples that the expression levels of the DEGs were sharply lower in the *bzip28* genotype compared with WT. In the *bzip28 bzip60* genotype, the genes in Cluster 3 had expression levels that were higher than the *bzip28* genotype but were also lower than those of these genes in the WT or *bzip60* genotypes. In response to chemically induced UPR conditions, bZIP60 and bZIP28 cooperatively upregulate several UPR genes (Song *et al.*, 2015; Ruberti *et al.*, 2018). In our ground controls, the stress-responsive genes, largely

represented by “response to abscisic acid,” “response to water deprivation,” and “response to hypoxia” abiotic stress responses, were regulated by bZIP28 and bZIP60 in a way that suggests that these TFs have antagonistic regulatory effects on these processes (Fig. 7B–D).

The exact nature of the stress experienced by the ground control seedlings would need to be elucidated to better understand the impact of this information on agronomic and/or spaceflight applications. Studies on the effect of plant growth in BRIC-PDFUs have already established that significant stress may be imposed on the seedlings grown in these conditions (Johnson *et al.*, 2015; Basu *et al.*, 2017). Consistent with our results, other BRIC-PDFU transcriptomes have found a downregulation of genes related to water stress responses (Johnson *et al.*, 2017). However, as evaporative water loss from the BRIC-PDFUs is unlikely because they are sealed containers, the strongly represented GO terms related to “response to water deprivation” and “response to abscisic acid” are unlikely to be a direct response to actual water loss from the plates. Instead, it may be possible that the ground seedlings with etiolated hypocotyls that are not in contact with the media or have less overall contact with the media are water stressed compared with flight seedlings that are in direct contact or have penetrated the media. Independent of the underlying stimulus for the observed differences, the possibility that bZIP60 and bZIP28 may have antagonistic interactions related to control of abscisic acid signaling or water deprivation responses should be investigated at the molecular level in the future to improve plant growth and stress responses in space and on the ground.

Data Accessibility

Raw sequencing data have been deposited at the NCBI Gene Expression Omnibus under the accession number GSE148914. Raw sequencing data have also been deposited at the NASA GeneLab data repository (Ray *et al.*, 2018) for spaceflight experiments under the accession number GLDS-321.

Authors' Contributions

F.B. designed the experiments. S.Z.-D. and E.A. performed preliminary launch preparations. E.A. executed the experiment. D.K.K. performed bioinformatics analysis. E.A., D.K.K., S.Z.-D, and F.B. analyzed data and wrote the article.

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Author Disclosure Statement

No competing financial interests exist.

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Supplementary Material

Supplementary Table S1
 Supplementary Table S2
 Supplementary Table S3
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Abbreviations Used

ATF6 = activating transcription factor 6
 BRIC = Biological Research In a Canister
 bZIP17 = basic leucine zipper 17
 bZIP28 = basic leucine zipper 28
 bZIP60 = basic leucine zipper 60
 DEG = differentially expressed gene
 ER = endoplasmic reticulum
 FPKM = fragments per kilobase exon model per million mapped reads
 GO = Gene Ontology
 IRE1 = inositol-requiring enzyme 1
 ISS = International Space Station
 KSC = Kennedy Space Center
 PCA = principal component analysis
 PDFU = Petri Dish Fixation Unit
 R/FR = red/far red
 RIN = RNA integrity number
 RNA-seq = RNA-sequencing
 rRNA = ribosomal RNA
 TF = transcription factor
 UPR = unfolded protein response
 WT = wild type