UC Riverside

UC Riverside Previously Published Works

Title

High resolution RNA-seq profiling of genes encoding ribosomal proteins across different organs and developmental stages in *Arabidopsis thaliana*.

Permalink

https://escholarship.org/uc/item/2m58j6rw

Journal

Plant direct, 5(5)

ISSN

2475-4455

Authors

Xiong, Wei Zhang, Jiancong Lan, Ting et al.

Publication Date

2021-05-01

DOI

10.1002/pld3.320

Peer reviewed

ORIGINAL RESEARCH





High resolution RNA-seq profiling of genes encoding ribosomal proteins across different organs and developmental stages in *Arabidopsis thaliana*

¹Guangdong Provincial Key Laboratory for Plant Epigenetics, Longhua Bioindustry and Innovation Research Institute, College of Life Sciences and Oceanography, Shenzhen University, Shenzhen, China

²Key Laboratory of Optoelectronic Devices and Systems of Ministry of Education and Guangdong Province, College of Optoelectronic Engineering, Shenzhen University, Shenzhen, China

³Department of Botany and Plant Sciences, Institute of Integrative Genome Biology, University of California, Riverside, CA, USA

Correspondence

Beixin Mo, Guangdong Provincial Key Laboratory for Plant Epigenetics, Longhua Bioindustry and Innovation Research Institute, College of Life Sciences and Oceanography, Shenzhen University, Shenzhen, Guangdong 518060, China. Email: bmo@szu.edu.cn

Funding information

Guangdong Innovation Team Project, Grant/ Award Number: 2014ZT05S078; National Natural Science Foundation of China, Grant/ Award Number: 31870287; National Key R&D Program of China Grant, Grant/Award Number: 2019YFA0903902; Shenzhen Science & Technology Program, Grant/ Award Number: JCYJ20190808115005598

Abstract

In Arabidopsis thaliana, each ribosomal protein (RP) is encoded by a small gene family consisting of two or more highly homologous paralogues, which results in ribosome heterogeneity. It is largely unknown that how genes from multiple member containing RP families are regulated at transcriptional level to accommodate the needs of different plant organs and developmental stages. In this study, we investigated the transcript accumulation profiles of RP genes and found that the expression levels of RP genes are varied dramatically in different organs and developmental stages. Although most RP genes are found to be ubiquitously transcribed, some are obviously transcribed with spatiotemporal specificity. The hierarchical clustering trees of transcript accumulation intensity of RP genes revealed that different organs and developmental stages have different population of RP gene transcripts. By interrogating of the expression fluctuation trend of RP genes, we found that in spite of the fact that most groups of paralogous RP genes are transcribed in concerted manners, some RPs gene have contrasting expression patterns. When transcripts of paralogous RP genes from the same family are considered together, the expression level of most RP genes are well-matched but some are obviously higher or lower, therefore we speculate that some superfluous RPs may act outside the ribosome and a portion of ribosomes may lack one or even more RP(s). Altogether, our analysis results suggested that functional divergence may exist among heterogeneous ribosomes that resulted from different combination of RP paralogues, and substoichiometry of several RP gene families may lead to another layer of heterogeneous ribosomes which also have divergent functions in plants.

KEYWORDS

Arabidopsis thaliana, functional specialization, gene duplication, paralogue, ribosomal protein, ribosome heterogeneity, transcript profiling

Wei Xiong and Jiancong Zhang contributed equally to this work.

This is an open access article under the terms of the Creative Commons Attribution License, which permits use, distribution and reproduction in any medium, provided the original work is properly cited.

© 2021 The Authors. Plant Direct published by American Society of Plant Biologists, Society for Experimental Biology and John Wiley & Sons Ltd.



1 | INTRODUCTION

Ribosome is a ribonucleoprotein complex comprising a large and a small subunit, and is essential for catalyzing the peptidyl transferase reaction during polypeptide synthesis in all living cells. In plants, the large ribosomal subunit is composed of 48 RPL (Ribosomal Protein of Large subunit) proteins in conjunction with three rRNAs (25S, 5.8S, and 5S), whereas the small subunit is composed of 33 RPS (Ribosomal Protein of Small subunit) proteins in conjunction with the 18S rRNA (Chang et al., 2005; Savada & Bonham-Smith, 2014). Biogenesis of cytoplasmic ribosome is a highly orchestrated process involving the coordinated production and transport of four rRNAs and 81 RPs (Saez-Vasquez & Delseny, 2019).

In Arabidopsis thaliana, each RP is encoded by a small gene family containing two or more highly homologous family members (Barakat et al., 2001). The presence of multiple gene paralogues for RPs in plants, which leads to the production of heterogeneous ribosomes (Giavalisco et al., 2005), might be a consequence of high frequency of ancestral polyploidy events and could reflect a need to maintain adequate ribosome dose or to maintain some degree of ribosome heterogeneity (Blanc & Wolfe, 2004; Horiguchi et al., 2011, 2012; Martinez-Seidel et al., 2020; Thomas et al., 2006; Xue & Barna, 2012). Recently, many studies suggested that a number of individual animal RPs have wideranging extraribosomal functions in processes such as transcription, translation, mRNA processing, DNA repair, apoptosis, and tumorigenesis (Aseev & Boni, 2011; Lindstrom, 2009; Lu et al., 2015; Naora, 1999; Warner & Mcintosh, 2009; Yang et al., 2019). Although RPs were normally targeted to the nucleolus for cytoplasmic ribosomal subunit assembly with rRNAs (Degenhardt & Bonham-Smith, 2008; Kruger et al., 2007; Lam et al., 2007; Savada & Bonham-Smith, 2014), some RPs could be secreted out of the cell (Dai et al., 2010), all of which may suggest extraribosomal functions of RPs.

The complexity of the plant ribosome biogenesis together with extraribsomal functions of RP genes raise an important question: how the duplicated RP genes are regulated at the transcriptional level to coordinate the needs of cells for specific RPs or specific RP paralogues? By analyzing the expressed sequence tag (EST) data and the complete genomic sequence of Arabidopsis, previous studies have identified 249 genes (including some pseudogenes) corresponding to different cytoplasmic RP types, and found only 52 RP genes lack a matching EST accession and 19 of these contain incomplete open reading frames. These results confirm that most RP genes are expressed (Barakat et al., 2001). A group of researchers identified 996 putative RP genes spanning 79 distinct RPs in Brassica napus using EST data from 16 tissue collections (Whittle & Krochko, 2009). Comparative analysis of the transcript levels according to EST data for Brassica napus RPs revealed that a large fraction of RP genes is differentially expressed and that the number of paralogues expressed for each RP varied extensively with tissue types. Using Genevestigator (Hruz et al., 2008) to analyze Arabidopsis 22k microarray data, another group of researchers studied transcript levels of 192 of the 254 Arabidopsis RP genes and revealed that transcript levels from different RP genes show up to a 300-fold difference across the RP population (Savada & Bonham-Smith, 2014).

Despite these studies on the steady state levels of RP transcripts, so far, it is still not clear about how coordinated expression of RPs from multigene families is accomplished. The genome-wide expression profiles across 79 different tissues and developmental stages using high-throughput transcriptome sequencing (Klepikova et al., 2016) provide the opportunity to comprehensively understand the expression of the duplicated RP genes in *Arabidopsis thaliana*. In this study, we analyzed these data and found that duplicated RP genes are transcribed dynamically in different cell types with some degree of function diversity and co-expression patterns. Furthermore, our analysis results suggested that transcript insufficiency of several RP families brings the possibility of the second layer of heterogeneous ribosomes, which may also have divergent functions.

2 | EXPERIMENTAL PROCEDURES

2.1 | RP gene list

The RP sequences of Arabidopsis thaliana were compiled based on the Arabidopsis Genome Initiative (AGI) identification numbers provided by previous researches (Barakat et al., 2001; Browning & Bailey-Serres, 2015; Whittle & Krochko, 2009). We focused on a total number of 240 RP genes which have definite AGI numbers or have detected transcripts ($N_{RP\,genes} = 256$, $N_{Pseudogenes} = 16$).

2.2 | RNA-seq data analysis

The RNA- seq data were analyzed using the pRNASeqTools. Briefly, raw reads were mapped to the Araport11 genome using STAR v2.6 (Dobin et al., 2013) with the parameters "--alignIn- tronMax 5000 --outSAMmultNmax 1 --outFilterMultimapNmax 50 --outFilterMismatchNoverLmax 0.1". Mapped reads were counted by feature Counts v1.6.4 (Liao et al., 2014). The read counts were normalized by calculating the RPKM (Reads Per Kilobase per Million mapped reads) value using the R package edgeR (Robinson et al., 2010). RPKM= $10^6 \times (\text{reads count})/(\text{total mapped reads}) \times 10^3/(\text{gene length}; Mortazavi et al., 2008). The cDNA length of RP genes were obtained from Araport11.$

2.3 | Heatmap and hierarchical clustering

The original RPKM was \log_2 transformed. A heatmap with the hierarchical cluster tree was made using the R package pheatmap with \log_2 (RPKM) value of RP genes with the parameters "clustering_method =complete", "clustering_distance_rows =euclidean". (pheatmap: Pretty Heatmaps. R package version 1.0.12. https://CRAN.R-project.org/package=pheatmap). The complete linkage method was used to cluster different samples and different RP genes. Euclidean distance was used to calculate the distance between different samples and different RP genes. Pearson correlation coefficient matrix are



presented to show the correlation among 69 different organs and 10 different developmental stages.

2.4 | Boxplot

The boxplot figure showing the overall expression level of RP genes across different organs and developmental stages was generated by the R packages ggpubr. ggpubr: 'ggplot2' Based Publication Ready Plots. R package version 0.2.5. https://CRAN.R-project.org/package=ggpubr).

2.5 | Calculating the transcript levels of different RP gene families

The transcript levels (RPKM value) of different RP gene families from different tissues were combined (sum of individual family members) respectively. The transcripts stoichiometry of 81 RP gene families was calculated with RPKM value of each RP gene family divided by the median RPKM value of total RP gene families of each examined tissue.

2.6 | Expression pattern analysis

Four type of tissues (1-day seedling(hypocotyl, cotyledons, hpical meristem with adjacent tissues), seed germination (SD.g1~SD.g3), meristem (M1~M10), flower (F1~F19.)) which represent samples from important growth and development stages were chosen to investigate the expression pattern of RP genes. To analyse the expression pattern, RPKM of the RP genes were transformed to z-score then clustered in six clusters by fuzzy cmeans (Futschik & Carlisle, 2005; Kumara & Futschik, 2007). R packages TCseq with the parameters algo = cm, k = 6, standardize = TRUE was used to obtain clustering algorithm. TCseq: Time course sequencing data analysis. R package version 1.6.1. https://bioconductor.org/packages/release/bioc/html/TCseq.html).

2.7 | Code availability

RNA-seq data bioinformatic analyses in this study were performed by an integrated pipeline for next-generation sequencing analysis, pRNASeqTools v0.6 [https://github.com/grubbybio/pRNASeqTools/] in its mapping only mode to get the reads count. This pipeline can be used freely under the MIT license.

3 | RESULTS

We downloaded the original RNA-seq data of 79 different Arabidopsis thaliana tissues and developmental stages from NCBI Sequence Read Archive (SRA) (Klepikova et al., 2016). In order to maximize the representation of different organs and stages, and to provide insights into the dynamics of gene expression in the most important processes in the plant life cycle, the sequenced samples were selected from different developmental stages including seed germination, seed development, silique development, transition to flowering, flower development, ovule development. Furthermore, the samples were also chosen with special focus on tissues and stages not sampled in microarray-based transcriptome map (Schmid et al., 2005), for example, detailed shoot apical/inflorescence meristem series and leaf development series (Klepikova et al., 2016). Each sample had two biological replicates. Description of sequenced samples is listed in Data S1. The Arabidopsis RP genes (Table S1) used in this research is based on previous publications (Barakat et al., 2001; Browning & Bailey-Serres, 2015; Whittle & Krochko, 2009). Some of the RP gene AGI numbers have been updated/adjusted based on data from The Arabidopsis Information Resource (TAIR).

3.1 | The expression levels of RP genes across different organs and developmental stages vary substantially in *Arabidopsis thaliana*

In order to understand the overall expression levels of RP genes during temporal and spatial developmental processes, we calculated the median value of RPKM (Reads Per Kilobase per Million mapped reads) of 240 RP genes and total genes across 79 different tissues or developmental stages in Arabidopsis thaliana. As shown in Figure 1, the median value of log₂(RPKM) of 240 RP genes was much higher than that of total genes in all examined tissues, suggesting RP genes are transcribed at relatively high level. Comparison of the median value of log₂(RPKM) of RP genes from different tissues (Figure 1), we found transcripts accumulation level of RP genes is varied substantially across different tissues or developmental stages. The first two lowest transcripts accumulation levels of RPs among the 79 examined tissues were observed in anthers of the mature flower (F.AN.ad) and opened anthers (F.AN), of which the median number of log₂(RPKM) are only around 3.5. Whereas, the accumulation level of RPs from anthers of the young flower (F.AN.y), of which the median number of log₂(RPKM) is 8, was much higher than those from the other two stages of anther development. The first two highest transcripts accumulation level of RPs among the 79 examined tissues were observed in seeds at first day after soaking (SD.g1) and meristem of 1-day seedling (S.M), of which the median number of log₂(RPKM) is around 10. Meristems (M), young seeds (SD.y), flowers (S), axis of the inflorescence (AX), and leaf petiole of the young leaf (L.PET.y) have relatively higher RP transcript accumulation levels than other examined tissues. In the contrast, transcript accumulation level of pod of the siliques (POD), dry seeds (SD,d), petals of the mature flower (F.PT.ad), and sepals of the mature flower (F.SP.ad) are relatively lower than those of other examined tissues. Transcript accumulation levels of RPs were correlated with the development activities of the examined tissues, suggesting that different expression

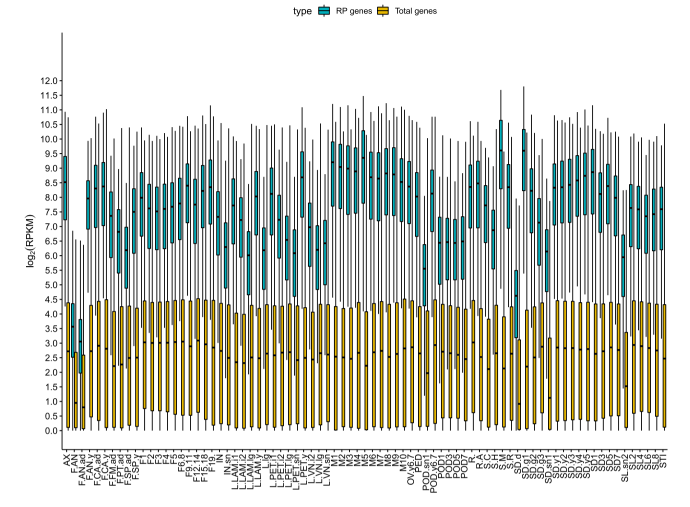


FIGURE 1 A boxplot figure showing the overall transcripts accumulation level of 240 RP genes and total genes across 79 different tissues and developmental stages in *Arabidopsis thaliana*. The average median value of RPKM (Reads Per Kilobase per Million mapped reads) of 240 RP genes and total genes were calculated from two biological replicates for each sample. The median value of log₂(RPKM) together with upper quartile and lower quartile were plotted against different samples. Blue, RP genes; yellow, total genes

level of RPs in different tissues may reflect different demands of ribosomes along with growth and development.

3.2 | Most RP genes are ubiquitously expressed

In order to investigate the transcription spatiotemporal specificity of RP genes, we made a heatmap with hierarchical cluster trees of transcript accumulation intensity of RP genes across different organs and developmental stages (Figure 2). We found that the majority of RP genes are ubiquitously expressed with exceptions that some RP genes are expressed with obviously spatiotemporal specificity. RP genes with extremely low transcripts accumulation levels or with obviously spatiotemporal specificities are shown in Figure S1. Take RPS15 family which contains six members (RPS15A, RPS15B, RPS15C, RPS15D, RPS15E, and RPS15F) for example (Figure 3), RPS15A, and RPS15D are transcribed in all examined tissues, whereas the other four members are specifically transcribed in certain reproductive

organs: *RPS15B* are only transcribed in developing seeds ((SD3), (SD5), (SD7)) and developing siliques ((SL4), (SL6)), *RPS15C* are only transcribed in anthers of the young flower (F.AN.y), several stages of flower development ((F4), (F5), (F6-8), (F9-11), (F12-14)) and two stages of developing seeds ((SD5), (SD7)), *RPS15E* are only transcribed in anthers of the young flower (F.AN.y), only one stage of flower development (F9-11), *RPS15F* are only transcribed in two stages of flower development ((F5), (F6-8)) and one stage of developing seeds (SD1). Divergence in spatiotemporal specificity of paralogous RP genes transcription suggests they could have undergone functional specializations among them. Spatiotemporally specific expression of paralogous RP genes lead to the formation of spatiotemporally specific ribosomes which may be needed for certain mRNA translation, thus could play a role in specific organ establishment and/or tissue-specific physiology.

The hierarchical clustering tree of transcripts accumulation intensity of 240 different RP genes revealed that distinct members from different RP families cluster together, implying that these

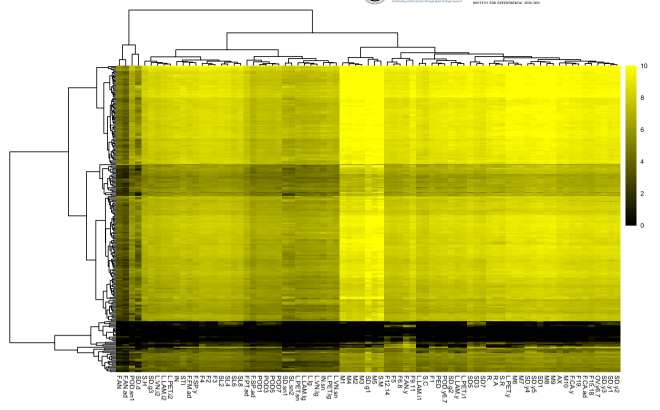


FIGURE 2 A heatmap with hierarchical cluster trees of transcripts accumulation intensity of RP genes across different organs and developmental stages. The value of $\log_2(RPKM)$ of each RP gene was plotted against each examined sample. Yellow, relative high expression; black, relative low expression. The vertical hierarchical cluster tree shows the Euclidean distance of examined samples and the horizontal hierarchical cluster tree shows the Euclidean distance of the log₂(RPKM) value

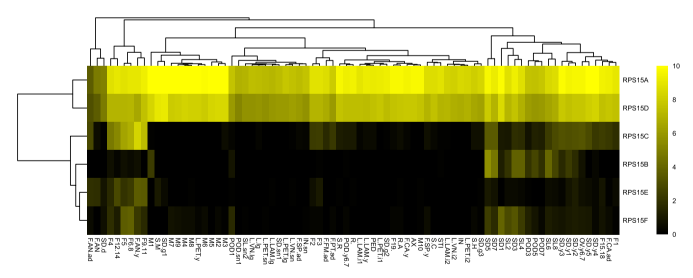


FIGURE 3 Diversified spatiotemporal transcripts accumulation pattern within RPS15 family. A heat map with hierarchical cluster trees of transcripts accumulation intensity of RPS15 family containing six members (RPS15A, RPS15B, RPS15C, RPS15D, RPS15E, and RPS15F) was made. Yellow, relative high expression; black, relative low expression. The vertical hierarchical cluster tree shows the Euclidean distance of examined samples whereas the horizontal hierarchical cluster tree shows the Euclidean distance of different RP genes based on the log₂(RPKM) value

non-paralogous RPs have similar expression intensity crossing different organs and developmental stages. Meanwhile, the hierarchical clustering tree of the examined tissues reflected an organ-specific

and developmental stage-specific structure, as the different tissues series are found to be organized into distinct clades. For example, different parts of root, different parts of leaf, different stages of



meistems are clustered together, respectively. Thus, different tissues and developmental stages have distinct population of RP gene transcripts. Ribosomes are highly heterogenous in *Arabidopsis*, each organ or developmental stage may need different association of non-paralogous RPs.

3.3 | Stoichiometric analysis of steady-state level of mRNAs between different RP families

Any given ribosome contains only a single polypeptide of most RPs except acidic ribosomal proteins P1 and P2 which form a heterodimer and two heterodimers are present per 60S subunit (Armache et al., 2010; Inglis et al., 2019). However, each one of the 81 Arabidopsis RPs is encoded by two or more paralogues. In order to maintain the appropriate stoichiometry of ribosome components. the amount of each RP family must be regulated at comparable level. Therefore, the transcript levels of RP gene families may be coordinately controlled. In this study, we combined transcript levels of each RP gene family (sum of individual family members) of each different tissue. The stoichiometry of 81-RP-family transcripts was calculated with RPKM value of each RP gene divided by the median RPKM value of total RP genes of each examined tissue. As shown in Data S2, the values of most RP gene families are around 1 (>0.5 meanwhile <2) but the value of several RP genes are largely deviated from 1 (<0.5 or >2), indicating that the expression level of most RP gene families are well-matched but some are obviously higher or lower. For instance, the value of the RPL9 gene families is less than 0.5 in almost all examined tissues (Data S2); the value of the gene family RPL10 is greater than 2 in opened anthers (F.AN), pods of the senescent silique (POD.sn1), dry Seeds (SD.d), senescent siliques (SL.sn2), (t-test, p < .05); the value of the gene family RPL14 is greater than 2 in senescent internodes (IN.sn), germinating seeds (SD.g2), veins of the mature leaves (L.VN.lg), carpels of 6th and 7th flowers (POD.y6-7), petioles of the mature leaves (L.PET.lg), petiole of the senescent leaves (L.PET.sn), pedicels (PED), developing flowers ((F3), (F4), (F5), (F6.8), (F12.14), (F19.), (t-test, p < .05)); the value of the gene family RPL19 is greater than 2 in pods of the senescent siliques (POD.sn1), senescent internodes (IN.sn), anthers of the mature flowers (F.AN. ad), opened anthers (F.AN), petiole of the senescent leaves (L.PET. sn), mature leaves (L.Ig), veins of the mature leaves (L.VN.Ig), lamina of the mature leaves (L.LAM.lg), (t-test, p < .05); the value of the gene family RPS17 is greater than 2 in anthers of the mature flowers (F.AN.ad) but is significantly less than 0.5 in seedling meristems (S.M), developing seeds (SD.y1, SD.y2, SD.y5), senescent internodes (IN.sn), (t-test, p < .05), (Data S2, and Figure S2). As for the acidic ribosomal proteins, the value of the gene family RPLP1 is greater than 2 in the mature flowers (F.AN.ad), seedlings meristems (S.M), meristems at 10 days after germination (M5), opened anthers (F.AN) and dry seeds (SD.d), and the value is greater than 1.5 in most examined samples (t-test, p < .05), (Data S2, and Figure S2). The value of the gene family RPLP2 is greater than 1 in all samples and is greater than 1.5 in near half of the examined tissues (t-test, p < .05), (Data S2,

and Figure S2). The value of the plant specific gene family *RPLP3* is >0.5 meanwhile <1 in all of the examined tissues (*t*-test, p<.05), (Data S2).

Interestingly, we found numbers of RP gene families, of which the values are greatly deviated from 1 (<0.5 or >2), are much larger in dry seeds (SD.d), senescent internodes (IN.sn), anthers of the mature flowers (F.AN.ad), and opened anthers (F.AN), where destructive metabolisms are the major biochemical reactions, than other examined tissues (Figure 4), suggesting that the stoichiometries of RP genes transcript accumulation levels are highly unequal in these tissues. The translation efficiencies of RP gene transcripts maybe different from each other (Fernie & Stitt, 2012). On what degree the stoichiometry inequality of RP gene transcripts affect RP stoichiometry at protein level needs to be investigated. RP families with higher transcript accumulation levels may have more RP proteins being translated, thus the amount of them maybe more than enough for the assembly of ribosome (the proposed hypothesis is shown in Figure 4g), suggesting these free RPs may act outside ribosomes. In contrast, RP families with extremely lower transcripts accumulation levels may have less RP proteins being translated, therefore they maybe not sufficient to be incorporated into ribosomes, causing a portion of ribosomes lacking these RPs (the proposed hypothesis is shown in Figure 4g). Apart from the heterogeneity of ribosomes contributed by different combination of non-paralogous RPs, substoichiometry of RP genes causes another layer of ribosome heterogeneity. Ribosomes lacking one or more RP(s) may not recognize some kinds of mRNAs or may have lower translating efficiency toward specific subpools of mRNAs as reported in mammalian cells (Kondrashov et al., 2011; Shi et al., 2017). Together, heterogeneous ribosomes contributed by some substoichiometric RPs and free RPs resulted from several superfluous RPs that are not incorporated into ribosomes, may play roles in regulating specific developmental stages or participating the establishment of specific organs.

3.4 | Synchronous and discrepant expression patterns coexist for RP genes

Former works investigating expression patterns of paralogous RP genes only compared the expression levels among them, RP families in *Arabidopsis* have been classified into two groups based on the transcript levels of their individual members (Type I, RP families with members having similar levels of transcript accumulation; Type II, RP families with members having varied levels of transcript accumulation; Savada & Bonham-Smith, 2014). In this study, in order to further understand the expression patterns of paralogous PR genes, we drew line charts comparing RPKM values of RP genes over different tissues or the same tissue over different developmental stages. Four types of tissues at the same developmental stage (parts of 1-day seedling, hypocotyl, cotyledons, and apical meristem with adjacent tissues); same tissues at different developmental stage (germinating seeds (SD.g1-SD.g3)); shoot meristems (M1-M10); flowers (F1-F19.)), which represent samples from important growth and developmental

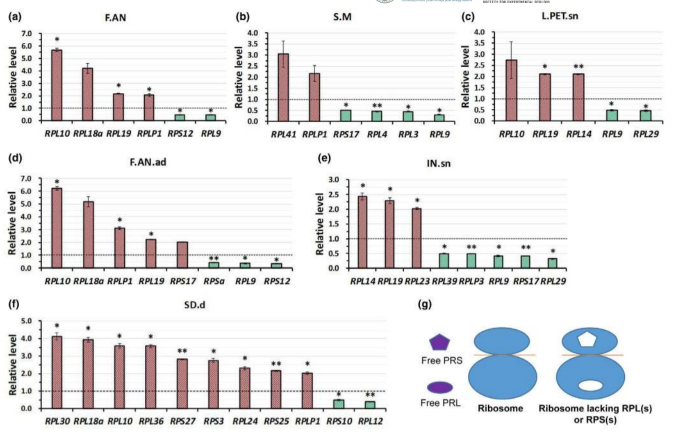


FIGURE 4 Samples of which the number of RP gene families largely deviating from 1 (<-0.5 or >2) is more than 5. Examined tissues such as (a) F.AN; (b) S.M; (c) L.PET.sn; (d) F.AN.ad; (e) IN.sn; (f) SD.d, of which the number of RP gene families largely deviating from 1 (<-0.5 or >2) is more than 5. (g), proposed hypothesis of free RPs which may act outside ribosomes and heterogeneous ribosomes contributed by some substoichiometric RPs. Relative level was calculated with RPKM value of each RP gene divided by the median RPKM value of total RP genes in each examined tissue. Error bars indicate SD from two independent experiments; asterisk indicates significant difference (t-test, p < .05)

stages, were chosen. For the majority of RP families (Table 1), their paralogous PR genes are transcribed in concerted manners. Take RPL34 and RPS29 families for example, although the expression levels of these paralogous PR genes are different, the expression fluctuation trends of them are well matched across different tissues and developmental stages (Figure 5), suggesting they are regulated in concerted manners, thus may demonstrate that these paralogous RP genes are functional redundant to maintain adequate dosages of their families. Meanwhile, for some RP families (Table 1), their members are transcribed in different manners. For example: RPLPO, RPL3, RPL7, RPL15, RPS2, RPS9, RPS15a, and RPS16 are found to have discrepant expression fluctuation trends (Figure 6), suggesting they are transcribed in distinctive manners with the possibility of diversified functions. Compensatory drift model envisions that although dosage-balance selection constrains total expression of paralogous genes to the optimum level, expression of each paralogue can diverge by drifting from its original level (Thompson et al., 2016). Our analysis results suggested that a portion of paralogous RP genes might have undergone functional specialization whereas the majority of paralogous RP genes seemed to remain the same function.

We also compared the expression fluctuation trend between different RP gene families. Interestingly, we found that the expression fluctuation trends of most RP genes from different families are similar (Figure 7, Figures S3–S5), indicating they are regulated in highly coordinated manners. Thus, synchronous expression of RP genes may play an important role in ribosome biogenesis. It was reported that almost half of all *Arabidopsis* RP genes carry several clustered GCCCR motifs in their proximal promoters, which could be recognized by a transcription factor called TCP20 (Li et al., 2005). TCP20 mediated transcriptional regulation of RP gene expression is likely to contribute to the synchronous expression of RP genes.

4 | DISCUSSION

4.1 | The complexity of RP gene expression patterns in *Arabidopsis*

mRNAs are translated into proteins by ribosomes which are composed of rRNAs and RPs in all living creatures. Genes encoding RPs are generally regarded as the housekeeping components of the cell to accommodate translational needs in development stages or growth situations, and are widely used for the standardization of transcript analysis data (Nicot et al., 2005; Sterky et al., 2004). In *E.*





TABLE 1 Expression patterns within RP families

Similar pattern Different pattern RPL4 RPS6 RPL3 RPL5 RPS7 RPL7 RPL6 RPS10 RPL8 RPL7a RPS11 RPL10 RPL9 RPS12 RPL11 RPL10a RPS13 RPL13a RPL12 RPS14 RPL14 RPL13 RPS17 RPL17 RPL13 RPS17 RPL17 RPL13 RPS18 RPL18 RPL13 RPS18 RPL18 RPL14 RPS20 RPL19 RPL22 RPS21 RPL21 RPL23 RPS24 RPL23 RPL23a RPS26 RPL27 RPL28 RPS27 RPL27 RPL28 RPS29 RPL00 RPL29 RPS29 RPL00 RPL36 RPL30 RPL91 RPL37 RPL32 RPL32 RPL37 RPL35 RPL35 RPL37 RPL36 RPL30			
RPL5 RPS7 RPL7 RPL6 RPS10 RPL8 RPL7a RPS11 RPL10 RPL9 RPS12 RPL11 RPL10a RPS13 RPL112 RPL12 RPS14 RPL14 RPL12 RPS14 RPL14 RPL13 RPS17 RPL17 RPL13 RPS17 RPL17 RPL13 RPS18 RPL18 RPL14 RPS18 RPL18 RPL15 RPS18 RPL18 RPL18 RPS20 RPL19 RPL22 RPS21 RPL21 RPL23 RPS24 RPL23 RPL23 RPS26 RPL24 RPL23 RPS27 RPL27 RPL28 RPS28 RPL27a RPL29 RPS29 RPL00 RPL36 RPL30 RPL11 RPL36 RPL31 RPL32 RPL37 RPL32 RPL32 RPL37 RPL32 RPS3 RPL37 RPS3 RPS4 RPL37 RPS8 RPS2	Similar pattern		
RPL6 RPS10 RPL8 RPL7a RPS11 RPL10 RPL9 RPS12 RPL11 RPL10a RPS13 RPL13a RPL12 RPS14 RPL14 RPL12 RPS14 RPL14 RPL13 RPS17 RPL17 RPL13 RPS17 RPL17 RPL13 RPS18 RPL18 RPL13 RPS18 RPL18 RPL18 RPS20 RPL19 RPL22 RPS21 RPL21 RPL23 RPS24 RPL23 RPL23 RPS24 RPL23 RPL24 RPL23 RPL27 RPL28 RPS28 RPL27a RPL29 RPS29 RPL90 RPL34 RPS30 RPL91 RPL35a RPL92 RPL90 RPL36a RPL30 RPL31 RPL37 RPL32 RPL32 RPL37 RPL35 RPL35 RPL39 RPS8 RPS2 RPL39 RPS8 RPS9 RPL41 RPS15a RPS15a	RPL4	RPS6	RPL3
RPL7a RPS11 RPL10 RPL9 RPS12 RPL11 RPL10a RPS13 RPL13a RPL12 RPS14 RPL14 RPL13 RPS17 RPL17 RPL15 RPS18 RPL18 RPL18a RPS20 RPL19 RPL22 RPS21 RPL21 RPL23 RPS24 RPL23 RPL23 RPS24 RPL23 RPL23 RPS24 RPL23 RPL23 RPS26 RPL24 RPL28 RPS27 RPL27 RPL28 RPS28 RPL27a RPL29 RPS29 RPL00 RPL34 RPS30 RPL19 RPL35a RPS30 RPL19 RPL35a RPL36 RPL36 RPL37 RPL37 RPL37 RPL37 RPL37 RPL37 RPL32 RPL37 RPL32 RPL37 RPL32 RPL39 RPS9 RPL39 RPS9 RPL30 RPS9 RPL31 RPS15 RPL37 RPS2 RPL37 RPL35 RPL39 RPS3 RPS4 RPS55 RPS53 RPS54 RPS515 RPS53 RPS53 RPS515 RPS53 RPS53 RPS52 RPS53 RPS53 RPS525	RPL5	RPS7	RPL7
RPL19 RPS12 RPL11 RPL10a RPS13 RPL13a RPL12 RPS14 RPL14 RPL13 RPS17 RPL17 RPL13 RPS18 RPL18 RPL15 RPS18 RPL18 RPL15 RPS18 RPL18 RPL16 RPS20 RPL19 RPL22 RPS21 RPL21 RPL23 RPS24 RPL23 RPL23 RPS24 RPL23 RPL23a RPS26 RPL24 RPL23a RPS26 RPL24 RPL26 RPS27 RPL27 RPL27 RPL27 RPL27 RPL28 RPS28 RPL27a RPL29 RPS29 RPLP0 RPL34 RPS30 RPLP1 RPL35a RPL30 RPL91 RPL36 RPL31 RPL32 RPL37 RPL32 RPL35 RPL37 RPS3 RPS4 RPL41 RPS15 RPS15 RPS3 RPS16 RPS3 RPS19 RPS3a	RPL6	RPS10	RPL8
RPL10a RPS13 RPL13a RPL12 RPS14 RPL14 RPL13 RPS17 RPL17 RPL15 RPS18 RPL18 RPL15 RPS18 RPL18 RPL15 RPS18 RPL18 RPL15 RPS18 RPL18 RPL18a RPS20 RPL19 RPL22 RPS21 RPL21 RPL23 RPS24 RPL23 RPL23a RPS24 RPL23 RPL23a RPS26 RPL24 RPL26 RPS27 RPL27 RPL28 RPS28 RPL27a RPL29 RPS29 RPLP0 RPL34 RPS30 RPLP1 RPL35a RPL30 RPL91 RPL36a RPL30 RPL31 RPL37 RPL32 RPL35 RPL37 RPL32 RPL35 RPL39 RPS8 RPL39 RPL40 RPS9 RPS15a RPS15 RPS15a RPS3 RPS15a RPS3 RPS15a RPS3	RPL7a	RPS11	RPL10
RPL12 RPS14 RPL14 RPL13 RPS17 RPL17 RPL15 RPS18 RPL18 RPL18a RPS20 RPL19 RPL22 RPS21 RPL21 RPL23 RPS24 RPL23 RPL23a RPS26 RPL24 RPL23a RPS26 RPL24 RPL26 RPS27 RPL27 RPL28 RPS28 RPL27a RPL29 RPS29 RPLP0 RPL34 RPS30 RPL91 RPL35a RPL92 RPL90 RPL36a RPL30 RPL31 RPL37 RPL32 RPL32 RPL37a RPL32 RPL35 RPL37a RPS2 RPS2 RPL39 RPS8 RPS40 RPL39 RPS8 RPS40 RPS15a RPS15a RPS3 RPS16 RPS3 RPS19 RPS3a RPS23 RPS4 RPS23 RPS4 RPS23	RPL9	RPS12	RPL11
RPL13 RPS17 RPL17 RPL15 RPS18 RPL18 RPL18 RPL18 RPL18 RPL19 RPL20 RPL19 RPL22 RPS21 RPL21 RPL23 RPS24 RPL23 RPL23 RPS26 RPL24 RPL23 RPS26 RPL27 RPL24 RPS27 RPL27 RPL28 RPS28 RPL27a RPL29 RPS29 RPLP0 RPL34 RPS30 RPLP1 RPL35a RPL92 RPL90 RPL36 RPL30 RPL91 RPL36 RPL30 RPL31 RPL37 RPL32 RPL32 RPL37 RPL32 RPL32 RPL37 RPS3 RPS4 RPL39 RPS4 RPS4 RPL40 RPS9 RPS15 RPS4 RPS16 RPS3 RPS3 RPS19 RPS23 RPS3 RPS19 RPS3a RPS23 RPS4 RPS25	RPL10a	RPS13	RPL13a
RPL15 RPS18 RPL18 RPL18a RPS20 RPL19 RPL22 RPS21 RPL21 RPL23 RPS24 RPL23 RPL23a RPS26 RPL24 RPL26 RPS27 RPL27 RPL28 RPS28 RPL27a RPL29 RPS29 RPLP0 RPL34 RPS30 RPLP1 RPL35a RPL30 RPL91 RPL36a RPL30 RPL30 RPL36a RPL30 RPL31 RPL37 RPL32 RPL32 RPL37a RPL35 RPL35 RPL39 RPS3 RPS3 RPL40 RPS9 RPS15 RPS4 RPS15 RPS15 RPS3 RPS16 RPS19 RPS3 RPS19 RPS23 RPS3 RPS23 RPS25 RPS4 RPS25 RPS25	RPL12	RPS14	RPL14
RPL18a RPS20 RPL19 RPL22 RPS21 RPL21 RPL23 RPS24 RPL23 RPL23a RPS26 RPL24 RPL26a RPS27 RPL27 RPL28 RPS28 RPL27a RPL29 RPS29 RPLP0 RPL34 RPS30 RPLP1 RPL35a RPL92 RPL30 RPL36 RPL30 RPL31 RPL36a RPL31 RPL31 RPL37a RPL35 RPL35 RPL38 RPS2 RPS3 RPL39 RPS8 RPS2 RPL39 RPS8 RPS9 RPL40 RPS9 RPS15 RPS9 RPS15a RPS15a RPS3 RPS16 RPS19 RPS3 RPS19 RPS23 RPS3 RPS23 RPS25	RPL13	RPS17	RPL17
RPL22 RPS21 RPL21 RPL23 RPS24 RPL23 RPL23a RPS26 RPL24 RPL26 RPS27 RPL27 RPL28 RPS28 RPL27a RPL29 RPS29 RPLP0 RPL34 RPS30 RPLP1 RPL35a RPL92 RPL92 RPL36a RPL30 RPL30 RPL37 RPL31 RPL31 RPL37 RPL32 RPL35 RPL38 RPS2 RPL35 RPL39 RPS8 RPS2 RPL40 RPS9 RPS41 RPS15 RPS15 RPS15 RPL93 RPS15a RPS15a RPS3 RPS19 RPS23 RPS3a RPS23 RPS23 RPS4 RPS25 RPS25	RPL15	RPS18	RPL18
RPL23 RPS24 RPL23 RPL23a RPS26 RPL24 RPL26 RPS27 RPL27 RPL28 RPS28 RPL27a RPL29 RPS29 RPLP0 RPL34 RPS30 RPLP1 RPL35a RPLP2 RPL36 RPL36a RPL30 RPL31 RPL37 RPL32 RPL32 RPL37a RPL35 RPS2 RPL39 RPS8 RPS2 RPL39 RPS8 RPS9 RPL40 RPS9 RPS15 RPL93 RPS15a RPS15a RPS3 RPS16 RPS19 RPS3 RPS19 RPS23 RPS3a RPS23 RPS23 RPS4 RPS25 RPS25	RPL18a	RPS20	RPL19
RPL23a RPS26 RPL24 RPL26 RPS27 RPL27 RPL28 RPS28 RPL27a RPL29 RPS29 RPLP0 RPL34 RPS30 RPLP1 RPL35a RPL92 RPL92 RPL36 RPL30 RPL30 RPL36a RPL31 RPL31 RPL37 RPL32 RPL32 RPL37a RPS2 RPS2 RPL39 RPS8 RPS8 RPL40 RPS9 RPL41 RPS15 RPL93 RPS15a RPS15a RPS3 RPS16 RPS19 RPS3a RPS23 RPS23 RPS4 RPS25 RPS25	RPL22	RPS21	RPL21
RPL26 RPS27 RPL27 RPL28 RPS28 RPL27a RPL29 RPS29 RPLP0 RPL34 RPS30 RPLP1 RPL35a RPL92 RPL30 RPL36 RPL30 RPL30 RPL36a RPL31 RPL31 RPL37 RPL32 RPL32 RPL37a RPL35 RPS2 RPL38 RPS2 RPS3 RPL40 RPS9 RPS41 RPS15a RPS15a RPS15a RPS3 RPS16 RPS19 RPS3a RPS23 RPS23 RPS4 RPS25 RPS25	RPL23	RPS24	RPL23
RPL28 RPS28 RPL27a RPL29 RPS29 RPLP0 RPL34 RPS30 RPLP1 RPL35a RPL92 RPL36 RPL36 RPL30 RPL30 RPL37a RPL32 RPL32 RPL37a RPS2 RPS2 RPL39 RPS8 RPS8 RPL40 RPS9 RPS15 RPL93 RPS15a RPS15a RPS3 RPS19 RPS19 RPS3a RPS23 RPS23 RPS4 RPS25 RPS25	RPL23a	RPS26	RPL24
RPL29 RPS29 RPLP0 RPL34 RPS30 RPLP1 RPL35a RPLP2 RPL30 RPL36 RPL30 RPL31 RPL37 RPL32 RPL32 RPL37a RPL35 RPL35 RPL38 RPS2 RPS8 RPL39 RPS8 RPS8 RPL40 RPS9 RPL41 RPS15 RPL93 RPS15a RPS15a RPS3 RPS19 RPS19 RPS3a RPS23 RPS23 RPS4 RPS25 RPS25	RPL26	RPS27	RPL27
RPL34 RPS30 RPLP1 RPL35a RPLP2 RPL36 RPL30 RPL36a RPL31 RPL37 RPL32 RPL37a RPL35 RPL38 RPS2 RPL39 RPS8 RPL40 RPS9 RPL41 RPS15 RPLP3 RPS15a RPS3 RPS16 RPS3 RPS19 RPS3a RPS23 RPS4 RPS25	RPL28	RPS28	RPL27a
RPL35a RPL92 RPL36 RPL30 RPL36a RPL31 RPL37 RPL32 RPL37a RPL35 RPL38 RPS2 RPL39 RPS8 RPL40 RPS9 RPL41 RPS15 RPLP3 RPS15a RPSa RPS16a RPS3 RPS19 RPS3a RPS23 RPS4 RPS25	RPL29	RPS29	RPLP0
RPL36 RPL30 RPL36a RPL31 RPL37 RPL32 RPL37a RPL35 RPL38 RPS2 RPL39 RPS8 RPL40 RPS9 RPL41 RPS15 RPLP3 RPS15a RPS3 RPS16 RPS3 RPS19 RPS3a RPS23 RPS4 RPS25	RPL34	RPS30	RPLP1
RPL36a RPL31 RPL37 RPL32 RPL37a RPL35 RPL38 RPS2 RPL39 RPS8 RPL40 RPS9 RPL41 RPS15 RPLP3 RPS15a RPSa RPS16a RPS3 RPS19 RPS3a RPS23 RPS4 RPS25	RPL35a		RPLP2
RPL37 RPL32 RPL37a RPL35 RPL38 RPS2 RPL39 RPS8 RPL40 RPS9 RPL41 RPS15 RPLP3 RPS15a RPSa RPS16a RPS3 RPS19 RPS3a RPS23 RPS4 RPS25	RPL36		RPL30
RPL37a RPL35 RPL38 RPS2 RPL39 RPS8 RPL40 RPS9 RPL41 RPS15 RPLP3 RPS15a RPSa RPS16 RPS3 RPS19 RPS3a RPS23 RPS4 RPS25	RPL36a		RPL31
RPL38 RPS2 RPL39 RPS8 RPL40 RPS9 RPL41 RPS15 RPLP3 RPS15a RPSa RPS16 RPS3 RPS19 RPS3a RPS23 RPS4 RPS25	RPL37		RPL32
RPL39 RPS8 RPL40 RPS9 RPL41 RPS15 RPLP3 RPS15a RPSa RPS16 RPS3 RPS19 RPS3a RPS23 RPS4 RPS25	RPL37a		RPL35
RPL40 RPS9 RPL41 RPS15 RPLP3 RPS15a RPSa RPS16 RPS3 RPS19 RPS3a RPS23 RPS4 RPS25	RPL38		RPS2
RPL41 RPS15 RPLP3 RPS15a RPSa RPS16 RPS3 RPS19 RPS3a RPS23 RPS4 RPS25	RPL39		RPS8
RPLP3 RPS15a RPSa RPS16 RPS3 RPS19 RPS3a RPS23 RPS4 RPS25	RPL40		RPS9
RPSa RPS16 RPS3 RPS19 RPS3a RPS23 RPS4 RPS25	RPL41		RPS15
RPS3 RPS19 RPS3a RPS23 RPS4 RPS25	RPLP3		RPS15a
RPS3a RPS23 RPS4 RPS25	RPSa		RPS16
RPS4 RPS25	RPS3		RPS19
	RPS3a		RPS23
RPS5 RPS27a	RPS4		RPS25
	RPS5		RPS27a

Note: RP family, with all members are transcribed in concerted manner, is listed in the "Similar pattern" group; whereas RP family, with one or more member(s) has/have different transcription manner(s), is listed in the "Different pattern" group.

coli, RP genes are clustered together and are arranged into 20 operons with approximately half of the genes mapping to a single locus (Mager, 1988). The arrangement of prokaryotic RP genes in operons provides a regulatory strategy which ensures coordinated expression and simultaneous regulation of groups of RP genes. Under normal growth situations, coordinated expression of prokaryotic RPs ensures ribosome biogenesis proportional to the growth rate, without significant accumulation of unincorporated ribosomal constituents. The expression of prokaryotic RP genes could be rapidly changed in response to stimuli such as nutrient availability (Nomura, 1999;

Schmid et al., 2005). In yeast *Saccharomyces cerevisiae*, three-fourths (59/79) of the RPs are encoded by functionally duplicated genes of which are not transcribed at the same level Jimenez et al., 2003; Warner et al., 1985). In mammals, most functional RPs are encoded by a single gene although there are about 2,000 pseudogenes that maybe related to RP genes (Balasubramanian et al., 2009). It should be noted that few mammalian RP genes are encoded by more than one functional paralogues. For example, in human there are three paralogous genes encoding *RPS4*, namely *RPS4X*, *RPS4Y1*, and *RPS4Y2*, which are located on the X chromosome and the Y chromosome respectively (Ellis et al., 2010; Fisher et al., 1990). *RPS4X* and *RPS4Y1* are found to be ubiquitously expressed; in contrast, the expression of *RPS4Y2* is restricted to the testis and prostate, suggesting functional specialization of these paralogues.

The existence of multigenes for each RP family in plants presents a picture of RP expression that may be more complex than those that were previously described for other species. The expression patterns of some RP genes have been investigated, primarily at the level of transcript abundance (Dresselhaus et al., 1999; Hulm et al., 2005; McIntosh & Bonham-Smith, 2005). However, the overall profiles of the coordinate response of RP genes to cell differentiation, growth, development remains to be comprehensively investigated. Achievement of such objective requires the development of new genetic and genomic resources. The genome-wide expression profiles across different tissues and developmental stages obtained from high-throughput transcriptome sequencing allow us to visualize and accurately understand RP gene expression across different organs and developmental stages.

4.2 | Different organs and developmental stages demand different amount of ribosomes

Our analysis results revealed that transcripts accumulation levels of RP genes vary substantially across different organs and developmental stages. Anthers of mature flower (F.AN.ad) and opened anthers (F.AN), which are considered as highly differentiated organs, were found to have the lowest transcripts accumulation level of RP genes among the 79 examined samples with the median number of log₂(RPKM) being only around 3.5. Whereas, anthers of the young flower (F.AN.y), of which the median number of log₂(RPKM) is 8, have much higher transcripts accumulation levels of RP genes than those two development stages. RP transcripts accumulation level from dry seeds (SD.d) at dormant state, was the 3rd lowest among the examined samples. Thus, it is reasonable that the expression level of RPs from established and quiescent tissues are relatively low, a reduced level of ribosome is still sufficient to support the living of them. After soaking, seeds begin to germinate, the expression level of RPs from seeds at first day after soaking (SD.g1) was the first highest among the 79 examined samples cross different organs and different developmental stages. During the early stage of seed germination, initial protein synthesis is dependent on residual ribosomes within the cells of

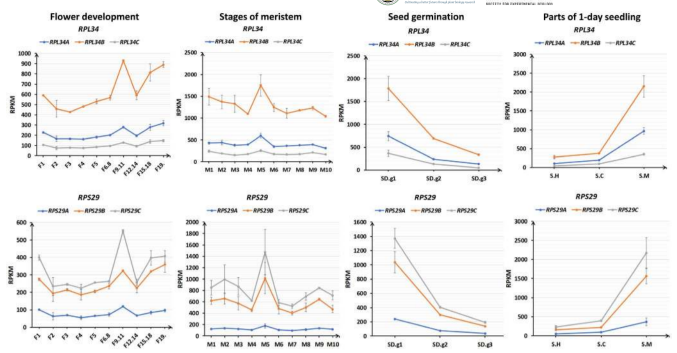


FIGURE 5 Concerted expression patterns of paralogous RP genes. The expression fluctuation trend of RP genes was investigated from four types of tissues representing important growth and development stages (Parts of 1-day seedling (hypocotyl, cotyledons, apical meristem with adjacent tissues); seed germination (SD.g1~SD.g3); meristem (M1~M10), and flower development (F1~19.)). RPL34 and RPS29 families were used as examples of RP gene paralogues with concerted expression pattern

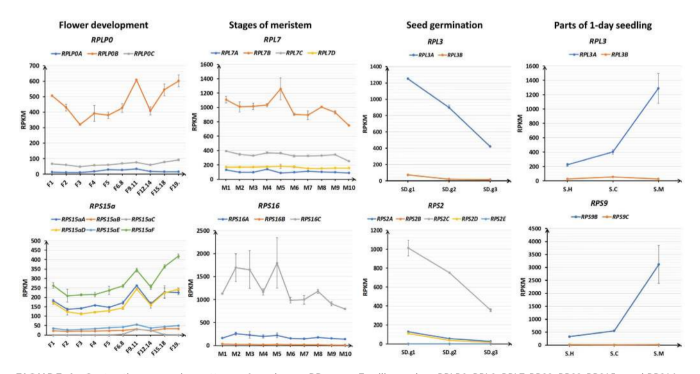


FIGURE 6 Contrasting expression patterns of paralogous RP genes. Families such as RPLP0, RPL3, RPL7, RPS2, RPS9, RPS15a, and RPS16 were used as examples of RP gene paralogues with contrasting expression pattern

mature dry embryos, but newly translated RPs are incorporated into ribosomes within hours of initial protein synthesis (Dommes & Walle, 1990). This developmental stage must require substantial

amount of newly synthesized proteins such as transcription factors, enzymes, et al., thus must demand extensive ribosomes, which well explains the highest expression levels of RP genes.

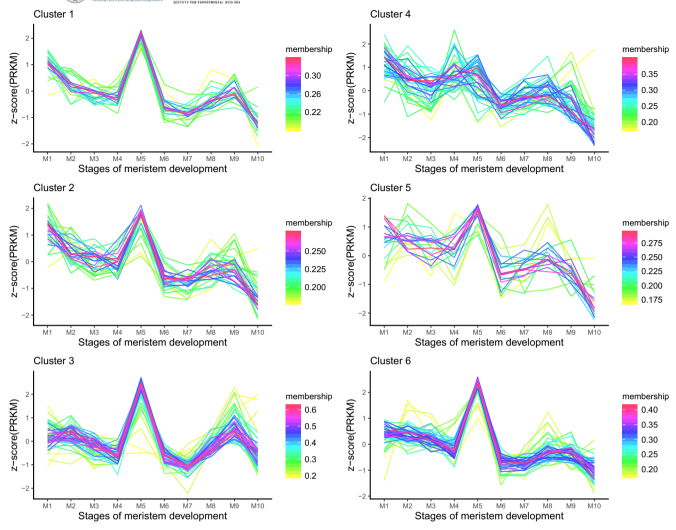


FIGURE 7 Consensus expression patterns of non-paralogous RP genes in different developmental stages of meristem. The RPKM value of paralogous and non-paralogous RP genes from different developmental stages of meristem was treated in accordance with Z-score using R packages TCseq with the parameters "algo = "cm", k = 6"s

4.3 | Varied combination of RP paralogues in different cell types may regulate the selection of mRNA translation subpools

The heat map with hierarchical clustering trees of RP genes across the 79 different tissues and different developmental stages based on the intensity of high-throughput transcriptome sequencing suggests that RP gene transcript population changes dramatically with tissue types. A number of RP genes are ubiquitously transcribed, whereas some RP genes are only transcribed in specific tissues. Tissue specific RPs could be divided into two groups: one group of RPs are only transcribed in limited tissues; another group of RPs are transcribed in most examined tissues but not transcribed in few tissues. We speculate that ubiquitously expressed RPs may assemble into ribosomes which satisfy the basic protein translation demands for cell survival. Tissue specific RPs may play a regulatory role in gene/protein expression through heterogeneous ribosomes lacking or containing different RP paralogues. The participation of heterogeneous ribosomes in gene regulation has been reported recently among

yeast strains (Komili et al., 2007). Furthermore, highly diverse and distinctive RP paralogue combinations observed in many examined tissues indicates that duplicated RP genes may have developed into functionally distinct paralogues associated with distinct tissues, and differentially expressed during plant development. This observation is consistent with the ribosome filter hypothesis, which postulates that rRNAs, and the combination of different RP paralogues, regulate the selection of translating supools of transcripts in different cell types, participating in cell differentiation and organ establishment in an elaborate way (Mauro & Edelman, 2002).

4.4 | Potential ribosome heterogeneity resulted from substoichiometry of RPs

The appropriate stoichiometry of ribosomal proteins in the nucleolus is very important for efficient ribosome biogenesis. It is generally considered that equal molar ratio of each RP family exists in assembled ribsomes. Our analysis results reveal that transcript

accumulation levels of most RP gene families are well-matched but some are obviously higher or lower. What needs to be investigated is whether RPs with lower accumulated transcripts are limiting factors in ribosome biogenesis or whether there is a mechanism to compensate the lower transcript level by higher translational efficiency. However, it is notable that recently a group of researchers applied a quantitative mass spectrometry approach to measure the absolute abundance of subsets of core RPs and identified four RPs (RPL10A, RPL38/eL38, RPS7/eS7, and RPS25/eS25) with an 30%-40% depletion within mouse embryonic stem cells (Shi et al., 2017), which are significantly substoichiometric in polysomes. In addition to the plant ribosome heterogeneity caused by different combination of nonparalogous RPs, substoichiometry of RP genes bring another layer of ribosome heterogeneity. Heterogeneity results from substoichiometry of RPs endows ribosomes with different selectivity for translating subpools of transcripts, including those controlling metabolism, the cell cycle, and development in animals (Shi et al., 2017). We suspect that substoichiometry of some RPs may also exist in Arabidopsis ribosomes according to our findings that transcripts accumulation levels of some RPs are obviously lower. Alternatively, RP families with higher transcripts may have more protein products and may be required for extraribosomal functions in addition to their conventional role in ribosome biogenesis and function. For example, it was reported that RPL24 play an important role in miRNA biogenesis in Arabidopsis (Li et al., 2017). According to our analysis result, the expression level of RPL24 family was a little bit higher than the median level of total RP families in most examined tissues (Data S2), which could be the evidence supporting the existence and action of free RPL24 outside the ribosome. RPL10, which was demonstrated to act in antivirus defense (Zorzatto et al., 2015), was another example of extraribosomal function of RPs. Our analysis results showed that expression level of RPL10 family are obviously higher than expected from a stoichiometric view of ribosome organization (Figure S2 and Data S2).

4.5 | Functional redundant and functional specialization co-exist within paralogous RPs

Duplication of RP genes is originated from gene or genome duplication events (Blanc & Wolfe, 2004). Gene balance pressures together with dosage effects could explain the high rates of retention of duplicated RP genes in the genome (Birchler et al., 2001, 2007; Conant & Wolfe, 2008; Veitia et al., 2008). The non-allelic noncomplementation phenomenon, which may indicate dosage effects between paralogous genes, was found in members within RP families, such as *RPL5* (Fujikura et al., 2009), *RPS6* (Creff et al., 2010), *RPL4* (Rosado et al., 2010), *RPL36a* (Casanova-Saez et al., 2014), and *RPL23a* (Xiong et al., 2020). The transcriptional fluctuation trends of some paralogous RP genes across different tissues and developmental stages was found well-matched although the transcript accumulation levels between them maybe different, suggesting some

paralogous RP genes are regulated in the concerted manner. Studies demonstrated that the loss-of-function rpl36ab and rpl36aa mutations have similar phenotypes and combined effects of haploinsufficiency and purifying selection drive retention of these two paralogs in Arabidopsis (Casanova-Saez et al., 2014). Our analysis results showed that paralogous RPL36a genes are transcribed in a concerted manner. Knockdown of RPL23aA resulted in severely developmental phenotypes whereas knockout of RPL23aB has no obvious phenotype (Degenhardt & Bonhamsmith, 2008a, 2008b). Interestingly, RPL23aA and RPL23aB were found to be transcribed in a concerted manner with much higher expression level of RPL23aA than that of RPL23aB, and over-expression of RPL23aB in rpl23aa could rescue the phenotype of rpl23aa (Xiong et al., 2020). We thought paralogous RP genes with concerted expression patterns may be the evidence supporting the dosage hypothesis which posits that the presence of duplicated RP genes might be necessary to maintain adequate RP dosages with functional redundant. Meanwhile, we also found a lot of paralogous RP genes are transcribed in highly contrasting manners, indicating function divergences exist within these paralogous RP genes. Plant paralogous RPL10 genes were found to have different roles during development and responds differently to ultraviolet-B stress (Ferreyra, Biarc, et al., 2010; Ferreyra, Pezza, et al., 2010). Consistent with genetic and functional studies of paralogous RPL10 genes, our analysis results revealed that paralogous RPL10 genes have contrary expression patterns (Table 1). Divergence in expression patterns is believed to be an important evidence proving functional specialization of duplicated genes (Adams et al., 2003; Blanc & Wolfe, 2004; Casneuf et al., 2006). Together, our analysis results suggested that functional redundant and functional speciali-

ACCESSION NUMBERS

zation may co-exist within paralogous RPs.

The original RNA-seq data of *A. thaliana* different organs and developmental stages were downloaded from NCBI Sequence Read Archive (project ID PRJNA314076 for samples except meristem and project ID PRJNA268115 for the meristem samples).

ACKNOWLEDGMENTS

We thank Dr. Lei Gao, Dr. Chenjiang You, and Dr. Yang Liu from Shenzhen University for their help with data analysis. This work was supported by the Guangdong Innovation Team Project (2014ZT05S078), National Natural Science Foundation of China (31870287) and National Key R&D Program of China Grant (2019YFA0903902), and Shenzhen Science & Technology Program (JCYJ20190808115005598). The funding body played no role in the design of the study, the collection, analysis and interpretation of the data and in writing the manuscript.

CONFLICT OF INTEREST

The authors have declared no conflict of interest.





AUTHOR CONTRIBUTIONS

BM, XC, and WX designed this work; JZ, TL, WK, and XW analyzed the RNA-seq data; WX and LL analyzed the results; WX, JZ, BM, and XC wrote the manuscript. All authors agreed to be accountable for the content of the work.

ORCID

Xuemei Chen https://orcid.org/0000-0002-5209-1157

Beixin Mo https://orcid.org/0000-0001-7279-9788

REFERENCES

- Adams, K. L., Cronn, R., Percifield, R., & Wendel, J. F. (2003). Genes duplicated by polyploidy show unequal contributions to the transcriptome and organ-specific reciprocal silencing. *Proceedings of the National Academy of Sciences of the United States of America*, 100, 4649–4654. https://doi.org/10.1073/pnas.0630618100
- Armache, J.-P., Jarasch, A., Anger, A. M., Villa, E., Becker, T., Bhushan, S., Jossinet, F., Habeck, M., Dindar, G., Franckenberg, S., Marquez, V., Mielke, T., Thomm, M., Berninghausen, O., Beatrix, B., Söding, J., Westhof, E., Wilson, D. N., & Beckmann, R. (2010). Localization of eukaryote-specific ribosomal proteins in a 5.5-Å cryo-EM map of the 80S eukaryotic ribosome. Proceedings of the National Academy of Sciences of the United States of America, 107, 19754–19759.
- Aseev, L. V., & Boni, I. V. (2011). Extraribosomal functions of bacterial ribosomal proteins. *Molecular Biology*, 45, 739. https://doi.org/10.1134/S0026893311050025
- Balasubramanian, S., Zheng, D., Liu, Y. J., Fang, G., Frankish, A., Carriero, N., Robilotto, R., Cayting, P., & Gerstein, M. (2009). Comparative analysis of processed ribosomal protein pseudogenes in four mammalian genomes. *Genome Biology*, 10, R2. https://doi.org/10.1186/gb-2009-10-1-r2
- Barakat, A., Szick-Miranda, K., Chang, I. F., Guyot, R., Blanc, G., Cooke, R., Delseny, M., & Bailey-Serres, J. (2001). The organization of cytoplasmic ribosomal protein genes in the arabidopsis genome. *Plant Physiology*, 127, 398–415. https://doi.org/10.1104/pp.010265
- Birchler, J. A., Bhadra, U., Bhadra, M. P., & Auger, D. L. (2001). Dosage-dependent gene regulation in multicellular eukaryotes: Implications for dosage compensation, aneuploid syndromes, and quantitative traits. *Developmental Biology*, 234, 275–288. https://doi.org/10.1006/dbio.2001.0262
- Birchler, J. A., & Veitia, R. A. (2007). The gene balance hypothesis: From classical genetics to modern genomics. *The Plant Cell*, 19, 395–402. https://doi.org/10.1105/tpc.106.049338
- Blanc, G., & Wolfe, K. H. (2004). Functional divergence of duplicated genes formed by polyploidy during Arabidopsis evolution. *The Plant Cell*, 16, 1679–1691. https://doi.org/10.1105/tpc.021410
- Browning, K. S., & Bailey-Serres, J. (2015). Mechanism of cytoplasmic mRNA translation. *The Arabidopsis Book*, 13, e0176. https://doi.org/10.1199/tab.0176
- Casanova-Saez, R., Candela, H., & Micol, J. L. (2014). Combined haploinsufficiency and purifying selection drive retention of RPL36a paralogs in Arabidopsis. Scientific Reports, 4, 4122. https://doi. org/10.1038/srep04122
- Casneuf, T., De Bodt, S., Raes, J., Maere, S., & Van de Peer, Y. (2006). Nonrandom divergence of gene expression following gene and genome duplications in the flowering plant Arabidopsis thaliana. *Genome Biology*, 7, R13.
- Chang, I. F., Szick-Miranda, K., Pan, S. Q., & Bailey-Serres, J. (2005). Proteomic characterization of evolutionarily conserved and variable proteins of arabidopsis cytosolic ribosomes. *Plant Physiology*, 137, 848–862. https://doi.org/10.1104/pp.104.053637

- Conant, G. C., & Wolfe, K. H. (2008). Turning a hobby into a job: How duplicated genes find new functions. *Nature Reviews Genetics*, *9*, 938–950. https://doi.org/10.1038/nrg2482
- Creff, A., Sormani, R., & Desnos, T. (2010). The two Arabidopsis RPS6 genes, encoding for cytoplasmic ribosomal proteins S6, are functionally equivalent. *Plant Molecular Biology*, 73, 533–546. https://doi.org/10.1007/s11103-010-9639-y
- Dai, Y., Pierson, S. E., Dudney, W. C., & Stack, B. C. Jr (2010). Extraribosomal function of metallopanstimulin-1: Reducing paxillin in head and neck squamous cell carcinoma and inhibiting tumor growth. *International Journal of Cancer*, 126, 611–619. https://doi.org/10.1002/jic.24791
- Degenhardt, R. F., & Bonham-Smith, P. C. (2008). Arabidopsis Ribosomal Proteins RPL23aA and RPL23aB Are Differentially Targeted to the Nucleolus and Are Disparately Required for Normal Development. *Plant Physiology*, 147, 128–142. https://doi.org/10.1104/pp.107.111799
- Degenhardt, R., & Bonhamsmith, P. C. (2008a). Arabidopsis Ribosomal Proteins RPL23aA and RPL23aB Are differentially targeted to the nucleolus and are disparately required for normal development. *Plant Physiology*, 147, 128–142. https://doi.org/10.1104/pp.107.111799
- Degenhardt, R., & Bonhamsmith, P. C. (2008b). Transcript profiling demonstrates absence of dosage compensation in Arabidopsis following loss of a single RPL23a paralog. *Planta*, 228, 627–640. https://doi.org/10.1007/s00425-008-0765-6
- Dobin, A., Davis, C. A., Schlesinger, F., Drenkow, J., Zaleski, C., Jha, S., Batut, P., Chaisson, M., & Gingeras, T. R. (2013). STAR: Ultrafast universal RNA-seq aligner. *Bioinformatics*, 29, 15–21. https://doi. org/10.1093/bioinformatics/bts635
- Dommes, J., & Walle, C. V. D. (1990). Polysome formation and incorporation of new ribosomes into polysomes during germination of the embryonic axis of maize. *Physiologia Plantarum*, 79, 289–296. https://doi.org/10.1111/j.1399-3054.1990.tb06744.x
- Dresselhaus, T., Cordts, S., Heuer, S., Sauter, M., Lörz, H., & Kranz, E. (1999). Novel ribosomal genes from maize are differentially expressed in the zygotic and somatic cell cycles. *Molecular and General Genetics MGG*, *261*, 416–427. https://doi.org/10.1007/s004380050983
- Ellis, P. J., Sargent, C. A., Miguel, R. N., Lopes, A. M., António, A., & Affara, N. A. (2010). The human RPS4 paralogue on Yq11.223 encodes a structurally conserved ribosomal protein and is preferentially expressed during spermatogenesis. BMC Molecular Biology, 11, 1–12.
- Fernie, A. R., & Stitt, M. (2012). On the Discordance of Metabolomics with Proteomics and Transcriptomics: Coping with Increasing Complexity in Logic, Chemistry, and Network Interactions. *Plant Physiology*, 158, 1139–1145.
- Ferreyra, M. L., Biarc, J., Burlingame, A. L., & Casati, P. (2010). Arabidopsis L10 ribosomal proteins in UV-B responses. *Plant Signaling & Behavior*, 5, 1222–1225. https://doi.org/10.4161/psb.5.10.12758
- Ferreyra, M. L. F., Pezza, A., Biarc, J., Burlingame, A. L., & Casati, P. (2010). Plant L10 ribosomal proteins have different roles during development and translation under ultraviolet-B stress. *Plant Physiology*, 153, 1878–1894. https://doi.org/10.1104/pp.110.157057
- Fisher, E. M., Beer-Romero, P., Brown, L. G., Ridley, A., McNeil, J. A., Lawrence, J. B., Willard, H. F., Bieber, F. R., & Page, D. C. (1990). Homologous ribosomal protein genes on the human X and Y chromosomes: Escape from X inactivation and possible implications for Turner syndrome. *Cell*, *63*, 1205–1218. https://doi.org/10.1016/0092-8674(90)90416-C
- Fujikura, U., Horiguchi, G., Ponce, M. R., Micol, J. L., & Tsukaya, H. (2009). Coordination of cell proliferation and cell expansion mediated by ribosome-related processes in the leaves of Arabidopsis thaliana. *Plant Journal*, 59, 499–508.
- Futschik, M. E., & Carlisle, B. (2005). Noise-robust soft clustering of gene expression time-course data. *Journal of Bioinformatics and*

13 of 14

- Computational Biology, 3, 965-988. https://doi.org/10.1142/S0219720005001375
- Giavalisco, P., Wilson, D., Kreitler, T., Lehrach, H., Klose, J., Gobom, J., & Fucini, P. (2005). High heterogeneity within the ribosomal proteins of the Arabidopsis thaliana 80S ribosome. *Plant Molecular Biology*, 57, 577–591. https://doi.org/10.1007/s11103-005-0699-3
- Horiguchi, G., Lijsebettens, M. V., Candela, H., Micol, J. L., & Tsukaya, H. (2012). Ribosomes and translation in plant developmental control. *Plant Science*, 191-192, 24–34. https://doi.org/10.1016/j.plant sci.2012.04.008
- Horiguchi, G., Mollamorales, A., Perezperez, J. M., Kojima, K., Robles, P., Ponce, M. R., Micol, J. L., & Tsukaya, H. (2011). Differential contributions of ribosomal protein genes to Arabidopsis thaliana leaf development. *Plant Journal*, 65, 724–736. https://doi.org/10.1111/j.1365-313X.2010.04457.x
- Hruz, T., Laule, O., Szabo, G., Wessendorp, F., Bleuler, S., Oertle, L., Widmayer, P., Gruissem, W., & Zimmermann, P. (2008). Genevestigator v3: A reference expression database for the meta-analysis of transcriptomes. Adv Bioinformatics, 2008, 420747. https://doi.org/10.1155/2008/420747
- Hulm, J. L., McIntosh, K. B., & Bonham-Smith, P. C. (2005). Variation in transcript abundance among the four members of the Arabidopsis thaliana RIBOSOMAL PROTEIN S15a gene family. *Plant Science*, 169, 267–278. https://doi.org/10.1016/j.plantsci.2005.04.001
- Inglis, A. J., Masson, G. R., Shao, S., Perisic, O., McLaughlin, S. H., Hegde, R. S., & Williams, R. L. (2019). Activation of GCN2 by the ribosomal P-stalk. Proceedings of the National Academy of Sciences of the United States of America, 116, 4946–4954. https://doi.org/10.1073/pnas.1813352116
- Jimenez, J. L., Mitchell, M. P., & Sgouros, J. G. (2003). Microarray analysis of orthologous genes: Conservation of the translational machinery across species at the sequence and expression level. *Genome Biology*, 4. R4.
- Klepikova, A. V., Kasianov, A. S., Gerasimov, E. S., Logacheva, M. D., & Penin, A. A. (2016). A High resolution map of the arabidopsis thaliana developmental transcriptome based on RNA:eq profiling. *The Plant Journal*, 88, 1058.
- Komili, S., Farny, N. G., Roth, F. P., & Silver, P. A. (2007). Functional specificity among ribosomal proteins regulates gene expression. *Cell*, 131, 557–571. https://doi.org/10.1016/j.cell.2007.08.037
- Kondrashov, N., Pusic, A., Stumpf, C. R., Shimizu, K., Hsieh, A. C., Xue, S., Ishijima, J., Shiroishi, T., & Barna, M. (2011). Ribosome-Mediated Specificity in Hox mRNA Translation and Vertebrate Tissue Patterning. *Cell*, 145, 383–397. https://doi.org/10.1016/j. cell.2011.03.028
- Kruger, T., Zentgraf, H., & Scheer, U. (2007). Intranucleolar sites of ribosome biogenesis defined by the localization of early binding ribosomal proteins. *Journal of Cell Biology*, 177, 573–578. https://doi. org/10.1083/jcb.200612048
- Kumar, L., & Futschik, M. E. (2007). Mfuzz: A software package for soft clustering of microarray data. *Bioinformation*, 2, 5–7. https://doi. org/10.6026/97320630002005
- Lam, Y. W., Lamond, A. I., Mann, M., & Andersen, J. S. (2007). Analysis of nucleolar protein dynamics reveals the nuclear degradation of ribosomal proteins. *Current Biology*, 17, 749–760. https://doi. org/10.1016/j.cub.2007.03.064
- Li, C., Potuschak, T., Colón-Carmona, A., Gutiérrez, R. A., & Doerner, P. (2005). Arabidopsis TCP20 links regulation of growth and cell division control pathways. Proceedings of the National Academy of Sciences of the United States of America, 102, 12978–12983.
- Li, S., Liu, K., Zhang, S., Wang, X., & Yu, B. (2017). STV1, a ribosomal protein, binds primary microRNA transcripts to promote their interaction with the processing complex in Arabidopsis. Proceedings of the National Academy of Sciences of the United States of America, 114, 201613069.

- Liao, Y., Smyth, G. K., & Shi, W. (2014). featureCounts: An efficient general purpose program for assigning sequence reads to genomic features. *Bioinformatics*, 30, 923–930. https://doi.org/10.1093/bioinformatics/btt656
- Lindstrom, M. S. (2009). Emerging functions of ribosomal proteins in gene-specific transcription and translation. *Biochemical and Biophysical Research Communications*, 379, 167–170. https://doi. org/10.1016/j.bbrc.2008.12.083
- Lu, H., Zhu, Y. F., Xiong, J., Wang, R., & Jia, Z. (2015). Potential extraribosomal functions of ribosomal proteins in Saccharomyces cerevisiae. *Microbiological Research*, 177, 28–33. https://doi.org/10.1016/j. micres.2015.05.004
- Mager, W. H. (1988). Control of ribosomal-protein geneexpression. *Biochimica Et Biophysica Acta*, 949, 1-15. https://doi. org/10.1016/0167-4781(88)90048-6
- Martinez-Seidel, F., Beine-Golovchuk, O., Hsieh, Y.-C., Kopka, J. (2020). Systematic review of plant ribosome heterogeneity and specialization. Frontiers in Plant Science, 11. https://doi.org/10.3389/fpls.2020.00948
- Mauro, V. P., & Edelman, G. M. (2002). The ribosome filter hypothesis.
 Proceedings of the National Academy of Sciences of the United States
 America, 99, 12031–12036. https://doi.org/10.1073/pnas.19244
 2499
- McIntosh, K. B., & Bonham-Smith, P. C. (2005). The two ribosomal protein L23A genes are differentially transcribed in Arabidopsis thaliana. *Genome*, 48, 443–454.
- Mortazavi, A., Williams, B. A., McCue, K., Schaeffer, L., & Wold, B. (2008). Mapping and quantifying mammalian transcriptomes by RNA-Seq. *Nature Methods*, 5, 621. https://doi.org/10.1038/nmeth.1226
- Naora, H., & Naora, H. (1999). Involvement of ribosomal proteins in regulating cell growth and apoptosis: Translational modulation or recruitment for extraribosomal activity? *Immunology and Cell Biology*, 77, 197–205. https://doi.org/10.1046/j.1440-1711.1999.00816.x
- Nicot, N., Hausman, J. F., Hoffmann, L., & Evers, D. (2005). Housekeeping gene selection for real-time RT-PCR normalization in potato during biotic and abiotic stress. *Journal of Experimental Botany*, 56, 2907– 2914. https://doi.org/10.1093/jxb/eri285
- Nomura, M. (1999). Regulation of ribosome biosynthesis in Escherichia coli and Saccharomyces cerevisiae: Diversity and common principles. *Journal of Bacteriology*, 181, 6857–6864. https://doi.org/10.1128/ JB.181.22.6857-6864.1999
- Robinson, M. D., McCarthy, D. J., & Smyth, G. K. (2010). edgeR: A Bioconductor package for differential expression analysis of digital gene expression data. *Bioinformatics*, 26, 139–140. https://doi. org/10.1093/bioinformatics/btp616
- Rosado, A., Sohn, E. J., Drakakaki, G., Pan, S. Q., Swidergal, A., Xiong, Y. Q., Kang, B. H., Bressan, R. A., & Raikhel, N. V. (2010). Auxinmediated ribosomal biogenesis regulates vacuolar trafficking in arabidopsis. *The Plant Cell*, 22, 143–158.
- Saez-Vasquez, J., & Delseny, M. (2019). Ribosome biogenesis in plants: from functional 45S ribosomal DNA organization to ribosome assembly factors. The Plant Cell, 31, 1945–1967. https://doi.org/10.1105/ tpc.18.00874
- Savada, R. P., & Bonham-Smith, P. C. (2014). Differential transcript accumulation and subcellular localization of Arabidopsis ribosomal proteins. *Plant Science*, 223, 134–145. https://doi.org/10.1016/j.plantsci.2014.03.011
- Schmid, M., Davison, T. S., Henz, S. R., Pape, U. J., Demar, M., Vingron, M., Scholkopf, B., Weigel, D., & Lohmann, J. U. (2005). A gene expression map of Arabidopsis thaliana development. *Nature Genetics*, 37, 501–506. https://doi.org/10.1038/ng1543
- Shi, Z., Fujii, K., Kovary, K. M., Genuth, N. R., Röst, H. L., Teruel, M. N., & Barna, M. (2017). Heterogeneous ribosomes preferentially translate distinct subpools of mRNAs genome-wide. *Molecular Cell*, 67(71–83), e77. https://doi.org/10.1016/j.molcel.2017.05.021





- Sterky, F., Bhalerao, R. R., Unneberg, P., Segerman, B., Nilsson, P., Brunner, A. M., Charbonnel-Campaa, L., Lindvall, J. J., Tandre, K., Strauss, S. H., Sundberg, B., Gustafsson, P., Uhlen, M., Bhalerao, R. P., Nilsson, O., Sandberg, G., Karlsson, J., Lundeberg, J., & Jansson, S. (2004). A Populus EST resource for plant functional genomics. Proceedings of the National Academy of Sciences of the United States, 101, 13951–13956. https://doi.org/10.1073/pnas.0401641101
- Thomas, B. C., Pedersen, B., & Freeling, M. (2006). Following tetraploidy in an Arabidopsis ancestor, genes were removed preferentially from one homeolog leaving clusters enriched in dose-sensitive genes. *Genome Research*, 16, 934–946. https://doi.org/10.1101/gr.4708406
- Thompson, A., Zakon, H. H., & Kirkpatrick, M. (2016). Compensatory drift and the evolutionary dynamics of dosage-sensitive duplicate genes. *Genetics*, 202, 765–774. https://doi.org/10.1534/genet ics.115.178137
- Veitia, R. A., Bottani, S., & Birchler, J. A. (2008). Cellular reactions to gene dosage imbalance: Genomic, transcriptomic and proteomic effects. *Trends in Genetics*, 24, 390–397. https://doi.org/10.1016/j. tig.2008.05.005
- Warner, J. R., & Mcintosh, K. B. (2009). How common are extraribosomal functions of ribosomal proteins? *Molecular Cell*, 34, 3–11. https://doi.org/10.1016/j.molcel.2009.03.006
- Warner, J. R., Mitra, G., Schwindinger, W. F., Studeny, M., & Fried, H. M. (1985). Saccharomyces cerevisiae coordinates accumulation of yeast ribosomal proteins by modulating mRNA splicing, translational initiation, and protein turnover. *Molecular and Cellular Biology*, 5, 1512– 1521. https://doi.org/10.1128/MCB.5.6.1512
- Whittle, C. A., & Krochko, J. E. (2009). Transcript profiling provides evidence of functional divergence and expression networks among ribosomal protein gene paralogs in Brassica napus. The Plant Cell, 21, 2203–2219.
- Xiong, W., Chen, X., Zhu, C., Zhang, J., Lan, T., Liu, L., Mo, B., & Chen, X. (2020). Arabidopsis paralogous genes RPL23aA and RPL23aB

- encode functionally equivalent proteins. *BMC Plant Biology*, 20, 463. https://doi.org/10.1186/s12870-020-02672-1
- Xue, S. F., & Barna, M. (2012). Specialized ribosomes: a new frontier in gene regulation and organismal biology. *Nature Reviews Molecular Cell Biology*, 13(6), 355–369. https://doi.org/10.1038/nrm3359
- Yang, C., Zang, W., Ji, Y., Li, T., Yang, Y., & Zheng, X. (2019). Ribosomal protein L6 (RPL6) is recruited to DNA damage sites in a poly(ADPribose) polymerase-dependent manner and regulates the DNA damage response. *Journal of Biological Chemistry*, 294(8), 2827–5664. https://doi.org/10.1074/jbc.RA118.007009
- Zorzatto, C., Machado, J. P. B., Lopes, K. V. G., Nascimento, K. J. T., Pereira, W. A., Brustolini, O. J. B., Reis, P. A. B., Calil, I. P., Deguchi, M., Sachetto-Martins, G., Gouveia, B. C., Loriato, V. A. P., Silva, M. A. C., Silva, F. F., Santos, A. A., Chory, J., & Fontes, E. P. B. (2015). NIK1-mediated translation suppression functions as a plant antiviral immunity mechanism. *Nature*, 520, 679–682. https://doi.org/10.1038/nature14171

SUPPORTING INFORMATION

Additional Supporting Information may be found online in the Supporting Information section.

How to cite this article: Xiong W, Zhang J, Lan T, et al. High resolution RNA-seq profiling of genes encoding ribosomal proteins across different organs and developmental stages in *Arabidopsis thaliana*. *Plant Direct*. 2021;5:e00320. https://doi.org/10.1002/pld3.320