Immediate Translation of Formin DIAPH1 mRNA after Its Exiting the Nucleus Is Required for Its Perinuclear Localization in Fibroblasts

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Abstract

DIAPH1 is a formin protein which promotes actin polymerization, stabilizes microtubules and consequently is involved in cytoskeleton dynamics, cell migration and differentiation. In contrast to the relatively well-understood signaling cascades that regulate DIAPH1 activity, its spatial regulation of biogenesis is not understood. A recent report showed that synthesis of DIAPH1 is confined in the perinuclear ER compartment through translation-dependent mRNA targeting. However, the underlying mechanism of DIAPH1 local synthesis is yet to be elucidated. Here, we provide evidence to demonstrate that the 5'-cap-mediated immediate translation of DIAPH1 mRNA upon exiting nucleus is required for localizing the mRNA in the perinuclear ER compartment. This is supported by data: 1) Delayed translation of DIAPH1 mRNA resulted in loss of perinuclear localization of the mRNA; 2) Once delocalized, DIAPH1 mRNA could not be retargeted to the perinuclear region; and 3) The translation of DIAPH1 mRNA is 5'-cap dependent. These results provide new insights into the novel mechanism of DIAPH1 local synthesis. In addition, these findings have led to the development of new approaches for manipulating DIAPH1 mRNA localization and local protein synthesis in cells for functional studies. Furthermore, a correlation of DIAPH1 mRNA and DIAPH1 protein localization has been demonstrated using a new method to quantify the intracellular distribution of protein.

Introduction

The importance of localized protein interaction in cellular functional regulation has been well established [1]. In addition to intracellular protein transport [2–5], local protein synthesis through mRNA targeting emerges as an important mechanism to confine a protein at a specific site of function and avoids inappropriate interactions with other proteins in other compartments [6–9]. In contrast to most of the investigated cytoplasmic protein-encoding mRNAs, which are localized through a localization signal sequence (zip-code) within the RNA molecules [6;8;10;11], a new class of cytoplasmic protein-encoding mRNAs employs a zip-code independent strategy for localization to the ER [12–15]. However, the mechanism for the localization of these mRNAs is poorly understood. Interestingly, two recent reports indicate that mRNAs encoding cytoplasmic protein XBP1u and DIAPH1 are targeted to the ER compartment through translation and their nascent peptides [13–15]. These findings add a new dimension to the conventional concept that only mRNAs encoding secreted and membrane proteins are targeted to the ER in a translation and nascent peptide dependent manner [16–18]. DIAPH1 is the one of the most studied formin proteins which stimulate formation of unbranched actin filaments [19–22], bind and stabilize microtubule [23;24] and link actin and microtubule cytoskeleton systems [25;26]. In cultured cells and knockout mice, DIAPH1 has been shown to play an important role in cell adhesion, migration, differentiation, signaling and gene expression [19–23;27–33]. In contrast to these advances, how DIAPH1 is spatially regulated is unclear. Previously, we demonstrated that DIAPH1 mRNA is enriched in the perinuclear compartment in fibroblasts, suggesting a spatial regulation of DIAPH1 protein biogenesis [15]. Our data also show that ongoing translation of DIAPH1 mRNA is required for the mRNA localization to the perinuclear ER compartment [15]. However, how the translation of DIAPH1 mRNA is regulated is not understood.

The vast majority of mRNAs are translated via 5’-cap-mediated initiation [34]. On the contrary, viral mRNA translation is mainly through internal ribosome entry site (IRES) mediated translation initiation [35]. The first IRES was characterized in poliovirus which is used for translation of viral protein, independent of cap-mediated translation [36]. This mechanism was soon found widely used by viruses for translation of their mRNAs while inhibiting the cellular 5’-cap-mediated translation [35;37]. Recently, a portion of cellular proteins has been found to be synthesized through cellular IRES which is in the cellular mRNA. Although both 5’-cap and cellular IRES mediated translational initiations share some common initiation factors, they do require different initiation factors which can be specifically inhibited [34;35]. For example, a small molecule 4E1RCat specifically inhibits 5’-cap mediated mRNA translational initiation whereas has minimal effect on IRES-mediated translational initiation [38]. In this report, we...
have taken advantage of this inhibitor and the differences between 5′-cap and IRES mediated mRNA translational initiation to dissect the mechanism of DIAPH1 mRNA translation and localization.

In this article, we examine the regulatory mechanism of DIAPH1 mRNA translation in the context of perinuclear DIAPH1 mRNA localization. Our data suggest that in order to localize in the perinuclear ER compartment, DIAPH1 mRNA is immediately translated upon being transported out of the nucleus through a 5′-cap mediated initiation. Additionally, unlike the mRNAs encoding membrane and secreted proteins, which are first translated for the signal peptides in the cytoplasm and then translocated to the ER compartment, we find that delocalized DIAPH1 mRNA cannot be translocated to the perinuclear compartment.

**Results**

Delocalized DIAPH1 mRNA cannot be Re-targeted to the Perinuclear Compartement

It was previously demonstrated that DIAPH1 mRNA is localized to the perinuclear ER in fibroblasts [15]. This localization is specific because mRNAs encoding subunit of Arp2/3 complex is localized to the cell protrusions in the same cells [15]. Furthermore, DIAPH1 mRNA is enriched in ER fraction in fractionation assay and co-localized with ER protein marker [15]. Translation is required for DIAPH1 mRNA localization to the perinuclear ER and active translation sites for the DIAPH1 mRNA are located in this perinuclear compartment [15]. However, it is not clear how translation regulates the perinuclear ER localization of the mRNA. We reasoned that there are two possible modes through which translation regulates DIAPH1 mRNA localization: 1) DIAPH1 mRNA is immediately translated after exiting the nucleus and the resulting nascent peptide helps to anchor the ribosome/mRNA complex around the nucleus by the interactions of the GBD-DID domains of the nascent peptide with unknown factor(s) on the ER. 2) Alternatively the mRNA might first enter the cytoplasm and is initially translated there before being translocated to the perinuclear compartment in a DIAPH1 nascent peptide dependent manner for continuous translation. The latter mode is somewhat analogous to the well-known mechanism for ER-translocation of mRNAs encoding membrane and secreted proteins, in which the mRNAs are first translated for the signal peptides in the cytoplasm and then translocated to the ER through signal peptides binding to specific receptors on the ER [16–18]. To distinguish these two modes for DIAPH1 mRNA localization, we tested whether delocalized DIAPH1 mRNA could be translocated to the perinuclear compartment in chicken embryo fibroblasts (CEF). To this end, DIAPH1 mRNA was first delocalized using protein synthesis inhibitor puromycin as previously demonstrated [15]. Puromycin inhibits protein translation by prematurely dissociating the nascent peptide from the ribosome/mRNA complex [39;40], which disrupts the DIAPH1 mRNA perinuclear localization [15]. To ensure that under our experimental conditions protein translation would be resumed after puromycin wash-off, we tested the relative amount and rate of new protein synthesis. This was done by using a Click-IT assay (Invitrogen) to detect newly synthesized proteins in a high signal/noise ratio and synchronized manner. As shown in Figure 1 (A–I), after puromycin wash-off, the relative amount and rate of newly synthesized proteins in the cells are similar to those of the control. We then asked if already delocalized DIAPH1 mRNA could be re-localized to the perinuclear compartment upon translation resumption by puromycin wash-off. As shown in Figure 1 (N & O), treatment with puromycin led to DIAPH1 mRNA delocalization, consistent with previous report [15]. It is unlikely that puromycin-induced DIAPH1 mRNA delocalization was caused by other non-specific effects of puromycin on general mRNA localization as previous studies demonstrated that puromycin treatment of CEF did not have any impact on cell protrusion localization of mRNAs encoding β-actin and the Arp2/3 complex [41;42]. In cells which were first treated with puromycin to delocalize DIAPH1 mRNA and then washed to remove puromycin, the DIAPH1 mRNA was still delocalized (Fig. 1, P & Q). To avoid potential interference for mRNA localization scoring from newly transcribed DIAPH1 mRNA molecules which are expected to localize at the perinuclear compartment, transcription inhibitor actinomycin D was used after puromycin wash-off. Actinomycin D itself had no effect on DIAPH1 mRNA localization (Fig. 1, L & M). These results indicate that delocalized DIAPH1 mRNA cannot be re-localized to the perinuclear compartment, suggesting that DIAPH1 mRNA localization in the perinuclear ER compartment is likely the result of immediate translation of DIAPH1 mRNA after its exiting the nucleus.

**Cap-mediated Translation is Required for DIAPH1 mRNA Localization**

Although the above results suggest immediate translation of DIAPH1 mRNA upon its exit of the nucleus is required for DIAPH1 mRNA perinuclear localization, it is not clear what translational initiation mechanism is involved in and responsible for this localization. Accumulating evidence indicates that although most mRNAs are translated using the well-documented 5′-cap-mediated translation initiation, a subset of cellular mRNAs use internal ribosome entry site (IRES) mediated initiation for their translation in the cell [34;35]. To address the question whether the 5′-cap-mediated or the IRES-mediated initiation is responsible for the translation of DIAPH1 mRNA in the perinuclear compartment, we used a small molecule inhibitor 4E1RCat to block 5′-cap mediated translation and asked if this is sufficient to delocalize DIAPH1 mRNA. 4E1RCat is a specific inhibitor which blocks 5′-cap-mediated translational initiation whereas has minimal effect on IRES-mediated translation initiation [38]. We first confirmed the inhibitory effect of 4E1RCat on protein synthesis in CEF using the Click-IT assay (Fig. 2A–I). To test if 4E1RCat selectively inhibits 5′-cap mediated but not IRES-mediated mRNA translation in these cells, we made a construct which expresses a bi-cistronic mRNA encoding red fluorescence protein mCherry and HA-tagged DIAPH1, respectively (named as M-I-D for mCherry-IRES-DIAPH1. see Fig. 2J). As shown in Figure 2K–O, 4E1RCat significantly inhibited 5′-cap mediated mCherry synthesis while had little effect on the IRES-mediated DIAPH1-HA synthesis. These results confirm the specific inhibitory effect of 4E1RCat on 5′-cap-mediated translation as previously reported [38]. We further asked whether inhibition of 5′-cap-mediated translation is sufficient to delocalize DIAPH1 mRNA. Indeed, treatment of CEF with 4E1RCat resulted in loss of DIAPH1 mRNA localization in the perinuclear compartment (Fig. 2P–T). Thus, 5′-cap mediated translation of DIAPH1 mRNA is required for its perinuclear localization.

**Manipulation of DIAPH1 mRNA Localization by Controlling Cap-mediated Translation using a Riboswitch iron Response Element (IRE)**

The requirement of 5′-cap-mediated translation for DIAPH1 mRNA localization suggests that such localization can be manipulated by controlling 5′-cap-mediated translation initiation. A ribo-switch, iron response element (IRE), has been used to
control 5’-cap mediated translation of mRNA [43–45]. The IRE is an RNA stem-loop which naturally exists in the 5’-UTR of mRNA encoding proteins involved in iron metabolism [46,47]. At low level of iron, an IRE binding protein (FP) binds to the IRE and

Figure 1. Delocalized DIAPH1 mRNA cannot be re-localized. A–I. Resumption of translation after puromycin wash-off. CEF grown on cover slips were treated with DMSO or 10 μg/ml of puromycin in methionine-free DMEM for 90 min and then followed by 2×10 min washes with Hank’s balanced saline. Newly synthesized proteins were detected using the Click-iT kit (Invitrogen) as described in Materials and Methods. A-H. Representative cells showing the fluorescence signal (red) of the newly synthesized proteins. I. Quantitative results of newly synthesized proteins indicate resumption of protein translation after puromycin wash-off (fluorescence per cell, normalized to that of time zero, representing ~80 cells at each time point per condition from two independent experiments). J-Q. Representative cells for DIAPH1 mRNA distribution after the indicated treatments. Images in the left column are gray scale for better display the DIAPH1 mRNA signal. CEF were transfected with HA-tagged DIAPH1 expression plasmid for 24 hr and then treated with DMSO (control, J & K), or 5 μg/ml of transcription inhibitor actinomycin D (Act-D) (L & M), or 10 μg/ml of puromycin (N & O) for 90 min before fixed for FISH detection of DIAPH1 mRNA localization. In P & Q, the cells were first treated with 10 μg/ml of puromycin for 90 min then followed by 2×10 min washes with growth medium plus 5 μg/ml of Act-D then incubated in normal growth medium for 90 min before fixed for FISH and DIAPH1 mRNA localization score. Note that Act-D at this concentration did not affect the normal localization of already transcribed DIAPH1 mRNA. In right column, Red: DIAPH1 mRNA; green: HA-tagged Dia1 protein; Blue: nucleus. Dotted lines show cell border. Arrows indicate localizing DIAPH1 mRNA molecules. Scale bar: 10 μm. R. Quantitative results of DIAPH1 mRNA localization. 300–500 cells were scored for each condition. Error bars: sem. n = 3. **, P<0.01.
prevents ribosome read-through thereby inhibiting translation (Fig. 3A). At high concentration of iron, the FP binds to the iron and dissociates from the IRE thereby allowing the ribosome read-through the 5'-mRNA sequence and resuming normal translation. By inserting the IRE into the 5'-UTR of an mRNA, one can control the translation of this mRNA in the cell by modulating the iron concentration in the culture medium [43–45]. Using the same approach, we generated an IRE-regulated expression construct to control the translation of \textit{DIAPH1} mRNA in transfected cells (Fig. 3A). Transfected CEF incubated in medium containing 100 μM of iron showed normal perinuclear \textit{DIAPH1} mRNA localization whereas those incubated in medium containing 100 μM of iron chelator showed loss of perinuclear \textit{DIAPH1} mRNA localization (Fig. 3 C–I). These results further support the requirement of 5'-cap-mediated translation for \textit{DIAPH1} mRNA localization and demonstrate that \textit{DIAPH1} mRNA localization can be manipulated by controlling its translation.

IRES-mediated Translation Leads to the Loss of \textit{DIAPH1} mRNA Localization

During the above study (Fig. 2), we unexpectedly found that in cells transfected with the construct M-I-D in which the \textit{DIAPH1} mRNA translation was under the control of IRES (see Fig. 2J or Fig. 4A for the structure of the construct), the \textit{DIAPH1} mRNA became diffuse (Fig. 4 E–G). We further compared the intracellular distribution of \textit{DIAPH1} mRNAs whose translation is initiated by the 5'-cap and the IRES, respectively by using the M-I-D and another construct D-I-M (for \textit{DIAPH1}-IRES-mCherry, see Fig. 4A). The results clearly demonstrate that under the same promoter control of mRNA transcription, \textit{DIAPH1} mRNA molecules whose translation was initiated by the 5'-cap localized normally around the perinuclear region whereas those initiated by the IRES were diffuse (delocalized) (Fig. 4). This is intriguing as it suggests that translation initiated by the 5'-cap or by the IRES has different impacts on \textit{DIAPH1} mRNA localization. Again, these results further support the idea that immediate and 5'-cap-mediated translation is required for \textit{DIAPH1} mRNA localization. Although how IRES-mediated translation leads to the loss of \textit{DIAPH1} mRNA localization has yet to be elucidated, this finding has provided a very useful approach for manipulating \textit{DIAPH1} mRNA localization for future functional study. In addition to the CEF, we have also tested D-I-M and M-I-D bicistronic mRNA expression constructs in NIH3T3 fibroblasts and observed similar differential mRNA localizations mediated by the 5'-cap and the IRES, respectively (Fig. 5 A–G). During the analysis of mRNA localization, we noticed that there may be a correlation of corresponding protein distribution with the mRNA. As a test,
instead of detecting the mRNA, we detected the mCherry and HA-tag signal in the cells transfected with the M-I-D and D-I-M constructs, respectively. In general, the protein signal is more diffuse which makes visual scoring difficult. To objectively analyze protein distribution in the cells, instead of analyzing the HA signal directly, we used the ratio of HA versus mCherry to correct the volume effect because the perinuclear region tends to be thicker than the cell periphery. Furthermore, we have developed a computer script to objectively quantify the intracellular distribution of protein (Fig. 5 H–N. see Materials and Methods for detailed description of the method). This script divides the cytoplasmic area into 15 equal area zones according to their relative distance from the edge of the nucleus (Fig. 5 N). The DIAPH1 protein signal was first corrected for cell volume effect and then quantified in a cell as IDI (Intracellular Distribution Index). As shown in Figure 5 O and P, DIAPH1 protein translated from the D-I-M mRNA exhibited perinuclear localization whereas that from the M-I-D mRNA showed more diffuse distribution (Fig. 5 H–M). These quantitative results confirm our observation that there is a correlation of DIAPH1 mRNA and DIAPH1 protein localization in fibroblasts.

Figure 3. Manipulation of DIAPH1 mRNA localization using an Iron ribo-switch. A. Schematic diagram of the IRE riboswitch (See Materials and Methods for details). Red balls represent 5'-cap. FB: iron binding protein which also binds to the IRE stem-loop. Green arrow indicates translation permission. B. Western blotting result of mCherry reporter for the effect of IRE in fibroblasts. A construct consisting of IRE-mCherry was transfected into CEF. 3 hr post transfection, ferric ammonium citrate (final 100 μM) or iron chelator desferrioxamine mesylate (final 100 μM) was added into the growth medium. 16 hr after transfection, the cells were collected for Western blotting. Quantitative results of Western blotting (n = 4), * p<0.05. C–H. IRE-mediated control of DIAPH1 mRNA localization. CEF were transfected with a construct consisting of IRE-DIAPH1 and then treated similarly as in B. 16 hr after transfection, the cells were fixed and processed for FISH detection of mRNA localization. C–H. Representative cells. Red: DIAPH1 mRNA signal; Green: HA-tagged DIAPH1 protein signal; Blue: nucleus. C, E & G are gray scale images for better presentation of DIAPH1 mRNA in the cells. Dotted lines show cell border. Arrows indicate localizing DIAPH1 mRNA. I. Quantitative results of DIAPH1 mRNA localization from analysis of 300–500 cells from three independent experiments for each condition. Error bars: sem. ** p<0.01. doi:10.1371/journal.pone.0068190.g003

Discussion

We previously demonstrated that DIAPH1 mRNA is anchored on the perinuclear ER in a translation dependent manner and the newly translated DIAPH1 protein (indicating the translation site) is located in a narrow zone around the nucleus in comparison to the relatively older DIAPH1 proteins [15]. In this report, we provide evidence to show that delocalized DIAPH1 mRNA cannot be re-localized to the perinuclear compartment. It has been reported that mRNA is transported out of the nuclear pores in a 5'-to-3' direction and translation of an mRNA could be initiated even before it is fully out of the nuclear pore [48;49]. Furthermore, using multiple independent and complementary approaches, we have also demonstrated that DIAPH1 mRNA translational initiation is mediated by the 5'-cap. Taken together, these lines of evidence strongly suggest that DIAPH1 mRNA is immediately translated upon exiting the nuclear pore and the DIAPH1 mRNA in the perinuclear region are the most actively translated, resulting in the perinuclear localization of the DIAPH1 mRNA and localized biogenesis of the DIAPH1 protein.

It is interesting to note that the general distribution of expressed DIAPH1 protein (as detected with HA-tag) in the cell is correlated...
with the location of the DIAPH1 mRNA, even though the protein distribution is more diffuse. This suggests that location of protein biogenesis will affect protein localization. This is consistent with our previous report that mis-targeting Arp2 mRNA, which encodes the Arp2 subunit of the actin polymerization nucleator Arp2/3 complex, to the perinucleus region led to reduced assembly of the Arp2/3 complex as compared to wild type cell with similar total Arp2 protein expression level [50]. This delocalization of Arp2 mRNA resulted in reduction of cell migration speed and the loss of directionality, demonstrating the functional importance of local protein synthesis, perhaps local co-translational assembly of the Arp2/3 complex [50;51]. Since the DIAPH1 protein is involved in cell migration and differentiation [19–22;27;30;32;33], it will be of great interest to investigate whether the manipulation of intracellular localization of DIAPH1 mRNA has functional consequences on these activities.

A question has been raised is why there is only 60% of the cells showing perinuclear DIAPH1 mRNA localization. The underlying mechanism is currently unclear but it may involve several possibilities. It could be the heterogeneous nature of a cell population. With the technical advancement in single cell analysis for proteomics and genomics, it has been known that individual cells in a supposed homogeneous population actually show very different gene expression patterns, morphologies and behaviors [52–55]. The heterogeneity of gene expression alone may play an important role in determining the cell behavior. Another possibility for only a portion of the cells showed intracellular localization of a particular mRNA is the cellular state such as phase of cell locomotion. It is known that mRNAs encoding β-actin and the actin polymerization nucleation complex Arp2/3 (with seven protein subunits) are localized to the protrusion of fibroblasts [41;42;56]. In a population of cells, on average, only about 30% of these cells showed protrusion associated mRNA localization. Using the MS2 system that was originally developed in the Singer laboratory [57], we observed Arp2 mRNA (encoding a subunit of the Arp2/3 complex) in live cells. The Arp2 mRNA was strongly enriched at the leading protrusion of migrating fibroblast with persistent direction (Mingle and Liu, unpublished). The same cells could show very little protrusion Arp2 mRNA localization when they withdrew the leading protrusion, paused or were in the process of turning to the opposition direction. Thus, this cell migratory state may explain why only a fraction of cells show protrusion. Whether cell migratory state and other cellular activities affect DIAPH1 mRNA localization, and vice versa, remains to be studied.

It remains unclear if and how the cap-mediated prompt DIAPH1 mRNA translation is regulated. General inhibition of cap-mediated translation is expected to affect DIAPH1 mRNA translation. It might be possible that DIAPH1 mRNA translational initiation upon exiting the nucleus is autonomous by default without any specific activation required. This is different from many other localizing mRNAs whose translation is suppressed during transport to their intracellular destinations [6–9]. For example, zip-code binding protein 1 (ZBP-1 or IMP-1) binds to the 3′-UTR of β-actin mRNA and suppresses its translation during transport [10;58]. We previously tested whether replacing the 3′-UTR of DIAPH1 with a β-actin zip-code containing sequence would inhibit DIAPH1 mRNA localization to the perinuclear compartment, and our results showed that such swapping of 3′-UTR did not affect DIAPH1 mRNA localization [15]. It remains to be determined whether there is a DIAPH1 mRNA specific inhibition/activation mechanism for its translation. In addition to translation initiation, other processes of translation may also play a role in DIAPH1 mRNA localization. For example, translation pausing which may provide time for the nascent peptide to fold and to maintain the number of ribosome associated with the mRNA as there are several putative translation pausing motifs in the coding region of DIAPH1 mRNA [14].

Figure 4. Internal Ribosome entry site mediated translation leads to delocalization of DIAPH1 mRNA. A. Illustration of bicistronic DIAPH1 expression constructs. D-I-M for DIAPH1-RES-mCherry and M-I-D for mCherry-RES-DIAPH1. CEF were transfected for 24 hr and processed for DIAPH1 mRNA and HA-tag detection. B–G. Representative transfected cells show localizing DIAPH1 mRNA (green in D, indicated by arrows) and delocalizing DIAPH1 mRNA (green in G, indicated by arrowheads), respectively. Red: mCherry. B–C and E–F are gray scale images for better presentation of the distribution of mCherry protein and DIAPH1 mRNA in cells transfected with the localizing and delocalizing constructs, respectively. H. Quantitative results of DIAPH1 mRNA localization from analysis of 300–500 cells from three independent experiments for each expression construct. Error bars: sem. ** p<0.01. doi:10.1371/journal.pone.0068190.g004
It is interesting that IRES-mediated translation results in loss of \(\text{DIAPH1}\) mRNA localization in the perinuclear compartment. It is unlikely that this is caused by the absence of translation of the delocalized mRNA (see representative cells in Figure 2K-N for HA-tagged \(\text{DIAPH1}\) protein expression). There are several possibilities for why IRES-mediated translation leads to delocalization of \(\text{DIAPH1}\) mRNA. First, because IRES- and cap-mediated translation initiation requires different factors, the IRES-specific factors may not be readily available in the perinuclear compartment for immediate translation. Second, because the efficiency of IRES-mediated translation is usually lower than that of 5'-cap-mediated translation, this may compromise the rate of nascent peptide production hence reducing the number of nascent peptide for the anchoring of the ribosome/mRNA/nascent peptide complex on the perinuclear ER. In this regard, drugs reducing cap-mediated translation may affect localized \(\text{DIAPH1}\) protein synthesis and generating adverse effects to the cell and organism.

The unexpected finding that IRES-mediated translation led to loss of \(\text{DIAPH1}\) mRNA localization provides a new means to manipulate \(\text{DIAPH1}\) mRNA localization for functional study. Even though we previously identified the nascent peptide motif that is critical for \(\text{DIAPH1}\) mRNA localization and created single point mutation mutants to delocalize \(\text{DIAPH1}\) mRNA [15], these mutants are not suitable for testing the functional importance of \(\text{DIAPH1}\) local biogenesis in the cell. This is because these mutations not only cause the delocalization of the \(\text{DIAPH1}\) mRNA, but also disrupt the known functions of \(\text{DIAPH1}\) protein, which makes the interpretation of the delocalization difficult. In contrast, the IRES-mediated translation provides a "clean" method to alter the localization of \(\text{DIAPH1}\) mRNA without any mutation in the \(\text{DIAPH1}\) sequence, facilitating the functional study for \(\text{DIAPH1}\) local biogenesis. It could be a useful approach for manipulating other mRNAs for their local translation. In fact, in a previous study using this approach of bicistronic mRNA with \(\text{DIAPH1}\) mRNA, we successfully mis-targeted \(\text{Arp2}\) mRNA to the...
perinuclear compartment without making any mutation in the Arp2 for functional investigation [30].

Materials and Methods

Ethics Statement

Primary chicken embryo fibroblasts (CEF) are a widely used cell type as reported in many publications [59,60]. They were isolated from the breast muscle of 12-day chicken embryos as described in details (52) (also see Cell culture and transfection). The tiny, partially developed, hairless, featherless, motionless embryo was carefully removed from the egg and decapped for euthanasia and convenience of subsequent tissue dissection. The Albany Medical College Institutional Animal Care and Use Committee (IACUC) was consulted and no protocol was required for this work.

Materials

Digoxigenin-11-dUTP (DIG-11-dUTP) and sheep anti-DIG antibody (peroxidase conjugated) were from Roche (Indianapolis, IN). Mouse anti-GAPDH antibody was from Ambion (Austin, TX). Rabbit anti-hemagglutinin (HA) antibody and Click-IT Protein Analysis kit were from Invitrogen (Grand Island, NY). Tyramide signal amplification (TSA) reagents were purchased from Perkin Elmer (Boston, MA). Actinomycin D (Act-D). 4E1RCat, ferric ammonium citrate and its chelator deferoxamine mesylate were from Sigma-Aldrich (Milwaukee, WI). Other general chemicals were from Sigma-Aldrich and Fisher (Pittsburgh, PA).

Cell Culture and Transfection

Standard quality fertilized chicken eggs were purchased from Charles River SPAFAS (North Franklin, CT). These eggs were incubated at 37°C for 12 days. They were then transferred to biosafety cabinet and sterilized by wiping with 70% alcohol. The tiny, partially developed, motionless, hairless and featherless embryos were then removed from the eggs which were still largely filled up with egg white and egg yolk at this stage. The embryos were decapitated for humane reason and convenience of tissue dissection. Breast muscle was dissected from the embryo and digested with trypsin at 37°C for 3 min then centrifuged at 1,000 × g for 5 min to remove the trypsin. The cell pellet was suspended in MEM with 10% fetal bovine serum and then either incubated with methionine-free DMEM (with DMSO or 10 μM puromycin) for 50 min and then followed by 2×10 min washes with Hank's balanced saline. Click-IT AHA (final 50 μM) was then added. At preset time points after Click-IT AHA addition, samples were fixed and processed for IF and/or FISH. In cell samples for DIAPH1 mRNA localization, to ensure only the “old” DIAPH1 mRNA molecules were detected and scored, we used Act-D to inhibit new DIAPH1 transcripts after puromycin wash-off.

Plasmid Construction

Standard molecular biology techniques were used in cloning and plasmid construction. Accession numbers for the cDNA sequences used in this study are: AB025226 (chicken DIAPH1), NM_205086.1 (the IRE element of chicken ferritin heavy chain) and NC_001479.1 (the IRES of encephalomyocarditis virus). For construction of iron/IRE mediated translation control of mCherry (red fluorescence protein) or DIAPH1, a pRL expression plasmid was used, which is under the control of a viral SV40 promoter (courtesy of Dr. Andrew Aplin). IRES was first inserted to a proper site within the SV40 promoter and then mCherry or DIAPH1 fused with HA tag (for protein detection) was cloned to the vector followed by a fragment of LacZ in the 3′-UTR for mRNA detection. For construction of IRES mediated translation of DIAPH1 or mCherry, the pNEW expression plasmid was used, which is under the control of a chicken β-actin promoter (courtesy of Dr. Stefan Kindler, Hamburg). To compare the localization of DIAPH1 mRNA whose translation is initiated by 5′-cap or the IRES, we first replaced the GFP in the pNEW plasmid with a cassette that contains an HA-tag at the end of the coding region for protein detection and a fragment of LacZ in the 3′-UTR for mRNA detection, a fragment of IRES for its translation initiation, and then sequentially inserted DIAPH1 or mCherry to either the upstream or downstream of the IRES to make two types of DIAPH1 expression constructs. All the resulted expression plasmids were verified by DNA sequencing.

Inhibition of 5′-cap Mediated Translated using 4E1RCat

4E1RCat is an inhibitor for 5′-cap mediated translation initiation but has little effect on IRES-mediated translation [38]. Because the vast majority of the mRNAs are translated through 5′-cap-mediated initiation, we first tested if 4E1RCat could inhibit new protein synthesis in the CEF. Cells grown on cover slips were incubated with methionine-free DMEM (with DMSO or 10 μM 4E1RCat) for 30 min, and then Click-IT AHA (final 50 μM) was added. At preset time points after Click-IT AHA addition, samples were fixed and processed for immunofluorescence staining. To test the specificity of this inhibitor, CEF were first transfected with the bicistronic plasmid for 2 hr and then incubated with DMSO or 10 μM 4E1RCat for 11 hr. This long time incubation is for better presentation of the differential effects of 4E1RCat on cap- and IRES-mediated translation, respectively.

The cells were then fixed and processed for immunofluorescence staining.

Using Iron Response Element (IRE) to Control DIAPH1 mRNA Translation and Localization

IRE is a structured RNA motif found in the 5′-UTR of mRNA encoded for proteins involved in iron metabolism [46,47]. At low iron concentration, an IRE binding protein binds to the IRE,
which blocks translation. At higher iron concentration the IRE binding protein dissociates from the IRE and translation starts. By inserting the IRE into other mRNAs and manipulating iron concentration, translation has been controlled successfully in a variety of cell types [43–45].

Probe Preparation and FISH

Nucleotides 62-1470 of chicken DIAPH1 (accession AB025226), and 388642-388413 of LacZ (accession CP002291.1) were cloned into pGEM-T Easy plasmids (Promega). These plasmids were linearized and transcribed in vitro in the presence of DIG-labelled dUTP for RNA probes using a Maxiscript transcription kit (Ambion, Austin, TX). Corresponding sense probes were also prepared similarly and used for specificity control tests. FISH with TSA was used to detect mRNA in the cells as previously described [15]. Briefly, RNA probes were hybridized to the fixed and permeabilized cells overnight at 60°C and then washed extensive-ly. Sheep anti-DIG antibody (peroxidase-conjugated) was used and the fluorescence signal was amplified with TSA (using tetramethylrhodamine-tyramide or fluorescein-tyramide).

Microscopic Image Acquisition, Data Analysis and Statistics

Fluorescence images were acquired using an Olympus microscope BX61 with an UPlanApo 40x oil objective (NA 1.0), a cooled CCD camera (SenCam from Cooke) and IPLab software (version 3.6.5, Scanalytics Inc. Fairfax, VA). Images were acquired using identical parameters and quantified for fluorescence per cell. Additional image processing was performed using Adobe Photoshop (version 7.0, Adobe Systems, Mountain View, CA) and ImageJ (version 1.43u, NIH). Statistical analysis was performed using the Student’s t-test for two samples with normal data distribution. For data with unequal distribution, Kruskal-Wallis test was used. Two samples with a P value <0.05 are regarded as significantly different.

Quantification of mRNA Localization

DIAPH1 mRNA localization was scored as described previously [15]. Briefly, cells were scored visually with sample identity concealed (single blind). A cell with ≥80% of the total mRNA signal in the perinuclear region was scored as perinuclearly localized otherwise will be scored as not perinuclearly localized. About 300-500 cells for each condition from three independent experiments were scored.

Quantification of Intracellular Protein Distribution

A new method for quantifying protein distribution between the perinuclear region and the cell periphery in the cytoplasm has been developed in our laboratory and used for the analysis of DIAPH1 protein distribution in this study. Cells transfected with the D-I-M or M-I-D construct were fixed and processed for IF to detect mCherry and HA-tagged DIAPH1 protein (note that under these conditions, mCherry keeps its fluorescence). Fluorescence images were acquired as described above. Because the protein signal is more diffuse than the punctate mRNA signal therefore it is difficult to score localization visually, we developed a computer script (within the IPlab software package) to objectively quantify protein distribution in the cell. Since the protein examined produce a relatively diffuse signal in the cell, if directly measured this can lead to a quantification artifact due to the volume-effect, as the perinuclear region of a cell is generally thicker than the cell periphery. To correct such cell volume effect, we first calculated the fluorescence signal ratio of DIAPH1-HA versus mCherry pixel by pixel from the original images and generated ratio images as corrected DIAPH1 protein signal. These images were then analyzed with a custom written computer script (Zone quant) for the relative distribution of DIAPH1 protein in each cell. The working principle and major steps of the script are as follows: 1) Dividing the cytoplasmic area of each cell into 15 equal area zones according to their relative distance to the nucleus border (see Fig. 5 for a simplified illustration which is with 5 zones). This was achieved by first obtaining the total cell area and the nuclear area using separate segments. The cytoplasmic area was derived from subtracting the nuclear area from the total cell area and was divided by 15 into fifteen equal area zones. At this point, the area of each zone was known but the location of each zone was not determined. 2) Creating and defining zones. This started from the border of the nucleus. A “dilate” function was used to add one pixel layer around the nuclear border which was defined by a segment. These dilation steps were repeated until the area of this segment was equal to the area of pre-defined for one zone in the cell. To prevent dilation from occurring beyond the cell border, during the dilation, if a dilating pixel met a ROI pixel which was used to define the cell border, the dilation of this pixel would be abolished while the dilation of other pixels continued. This created the first zone and the DIAPH1 protein signal in this zone was quantified and saved to a database table. The dilation would be then resumed by adding pixels to the outer edge of the previous zone until the area of this new zone was equal to the pre-determined zone area and DIAPH1 protein signal in this new zone was quantified. By reiterating the above processes, the DIAPH1 protein signals in the 15 zones were quantified. 3) Scatter plot graph for curve-fitting and generation of Intracellular Distribution Index (IDI). To minimize the impacts of differences in protein expression level, cell size and shape among the cells in a population on the results, the value of DIAPH1 protein signal in each zone was divided by the mean of the total 15 zones to generate a series of ratio values for each cell. The resulting ratio data from each cell were plotted as scatter plot graph using Sigma Plot (version 10.0, Systat Software Inc. San Jose, California) and curve-fitted with linear regression. The value of slope was used as IDI. If an IDI = 0, it suggests that the protein is uniformly distributed in the cells. If an IDI >0, it means that there is an ascending gradient from the nucleus to the cell periphery while IDI <0 indicates a descending gradient from the nucleus to the cell periphery.

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Author Contributions

Conceived and designed the experiments: GL GNL. Performed the experiments: GL GNL. Analyzed the data: GL GNL. Wrote the paper: GL GNL.

References

1. Hung MC, Link W (2011) Protein localization in disease and therapy. J Cell Sci 124: 3381–3392.

2. Boumen E, Simons K (1998) Protein and lipid sorting from the trans-Golgi network to the plasma membrane in polarized cells. Semin Cell Dev Biol 9: 503–509.
3. Mostov KE, Veves M, Alschuler Y (2000) Membrane traffic in polarized epithelial cells. Curr Opin Cell Biol 12: 483–490.
4. Paknikar KM (2007) Landmark discoveries in intracellular transport and secretion. J Cell Mol Med 11: 393–397.
5. Benham AM (2012) Protein secretion and the endoplasmic reticulum. Cold Spring Harb Perspect Biol 4: a012872.
6. St Johnstone D (2005) Moving messages: the intracellular localization of mRNAs. Nat Rev Mol Cell Biol 6: 363–375.
7. Lecuyer E, Yoshida H, Parthasarathy N, Alm C, Babak T, et al. (2007) Global analysis of mRNA localization reveals a prominent role in organizing cellular architecture and function. Cell 131: 174–187.
8. Holt CE, Bullock SL (2009) Subcellular mRNA localization in animal cells and why it matters. Science 326: 1212–1216.
9. Meignin C, Davis I (2010) Transmitting the message: intracellular mRNA localization. Curr Opin Cell Biol 22: 112–119.
10. Orenyakov Y, Singer RH (1998) RNA localization: different zipcodes, same postmark? Trends Cell Biol 8: 381–383.
11. Du TG, Schmid M, Jansen RP (2007) Why cells move messages: the biological functions of mRNA localization. Semin Cell Dev Biol 18: 171–177.
12. Stephens SB, Niewchich GV (2008) Divergent Regulation of Protein Synthesis in the Cytosol and Endoplasmic Reticulum Compartments of Mammalian Cells. Mol Cell Biol 19: 623–632.
13. Yanagita K, Kimata Y, Kodakura H, Kono K (2011) Translational pausing ensures membrane targeting and cytoplasmic splicing of XBPI mRNA. Science 331: 586–589.
14. Liao G, Liu G (2011) Why and how does DIAPH mRNA localize? Commun Integr Biol 4: 360–362.
15. Gao L, Ma X, Liu G (2011) An RNA-zipcode-independent mechanism that localizes DIAPH mRNA to the perinuclear ER through interactions between DIAPH nascent peptide and Rho-GTP. J Cell Sci 124: 589–599.
16. Walter P, Blobel G (1981) Translocation of proteins across the endoplasmic reticulum. II. Signal recognition particle (SRP) mediates the selective binding to microsomal membranes of in-vitro-assembled poly-somes synthesizing secretory protein. J Cell Biol 91: 551–556.
17. Blobel G, Dobberstein B (1975) Transfer of proteins across membranes. II. Reconstitution of functional rough microsomes from heterologous components. J Cell Biol 67: 852–862.
18. Blobel G, Dobberstein B (1975) Transfer of proteins across membranes. I. Presence of proteolytically processed and unprocessed nascent immunoglobulin light chains on membrane-bound ribosomes of murine myeloma. J Cell Biol 67: 835–851.
19. Pruyne D, Evangelista M, Yang C, Bi E, Zigmond S, et al. (2002) Role of formins in actin assembly: nucleation and barbed-end association. Science 297: 835–851.
20. Mingle LA, Yu EH, Tullio AN, Adelstein RS, Singer RH (2001) A Rho-Rock signaling, ROCK and mDia1, in the Cytosol and Endoplasmic Reticulum Compartments of Mammalian Cells. Mol Cell Biol 21: 623–632.
21. Goulimari P, Kitzing TM, Knieling H, Brandt DT, Offermanns S, et al. (2005) mDia2 stabilizes microtubules independently of its actin nucleation activity. J Biol Chem 280: 42242–42251.
22. Schonichen A, Geyer M (2010) Fifteen formins for an actin filament: a molecular function of mRNA directed by a sequence derived from poliovirus RNA. Nature 334: 320–325.
23. Bertrand E, Chartrand P, Schaefer M, Shenoy SM, Singer RH, et al. (1998) Identification of newly synthesized proteins in mammalian cells using bioorthogonal noncanonical amino acid tagging (BONCAT). Proc Natl Acad Sci U S A 95: 5118–5122.
24. Lin AH, Van Deursen JM, Feig M, Holmgren J, White J, et al. (2007) Impaired T lymphocyte trafficking in mice deficient in an actin-nucleating protein, mDia1. J Exp Med 204: 2011–2024.
25. Fusi P, Tullio AN, Chen Z, Singer RH, Condeelis J, et al. (2005) Formin mDia1 mediates vascular remodeling via integration of oxidative and signal transduction pathways. Curr Biol 15: 1279–1293.
26. Sonnenberg N, Hinnebusch AG (2009) Translation regulation of eukaryotic mechanisms and biological targets. Cell 136: 731–745.
27. Kornar AA, Hataouoglou M (2013) Cellular IRES-mediated translation: the war of chemoresistance by small molecule inhibition of the translation initiation complex eIF4F. Proc Natl Acad Sci U S A 106: 1046–1051.
28. Pelleiter J, Sonnenberg N (1989) Internal initiation of translation of eukaryotic mRNA directed by a sequence derived from poliovirus RNA. Nature 334: 320–325.
29. Balyov L, Soto R, Ribi EC, Decimo D, Ohlmann T (2009) Structural and functional diversity of viral IRESes. Biochem Biophys Acta 1789: 542–557.
30. Cencic R, Hall DR, Robert F, Du Y, Min J, et al. (2011) Reversing chemoresistance by small molecule inhibition of the translation initiation complex eIF4F. Proc Natl Acad Sci U S A 108: 2551–2555.
31. Bartolini F, Gundersen GG (2010) Formins and microtubules. Biochim Biophys Acta 1798: 966–973.
32. Tanizaki H, Egawa G, Inaba K, Honda T, Nakajima S, et al. (2010) Rho-mDia1 pathway is required for adhesion, migration, and T-cell stimulation in dendritic cells. Blood 116: 5837–5848.
33. Toure F, Fritz G, Li Q, Xavi R, Diwu G, et al. (2012) Formin mDia1 mediates vascular remodeling via integration of oxidative and signal transduction pathways. Circ Res 110: 1279–1293.
34. Schonichen A, Geyer M (2010) Fifteen formins for an actin filament: a molecular function of mRNA directed by a sequence derived from poliovirus RNA. Nature 334: 320–325.