



Friedrich Miescher Institute
for Biomedical Research

Master Thesis

Investigating the stress granule protein G3BP1 with single-molecule imaging of mRNA

Author:

Bastian Th. Eichenberger

Supervisor:

Jeffrey A. Chao and Susan E. Mango

UNIVERSITY OF BASEL
Basel, Switzerland

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Abstract

Stress granules are cytoplasmic, membrane-less organelles containing a plethora of proteins involved in translation initiation and repression and are exclusively formed in stress conditions. This study aimed to better understand the role of stress granules by focussing on a core protein component: Ras GTPase-Activating Protein-Binding Protein 1 (G3BP1). By tethering G3BP1 to specific mRNAs, various single-molecule imaging tools were used to investigate G3BP1's effect on translation (SunTag) and mRNA stability (TREAT). G3BP1 tethering to mRNAs encoding *Renilla* luciferase and SunTag yielded an increase in bulk *Renilla* Luciferase activity but SunTag imaging of translation showed that translation sites had similar intensities, which measures ribosome occupancy, compared to control cells. However, as shown by a TREAT assay to measure 5'-3' mRNA degradation, mRNA stability was significantly increased upon G3BP1 tethering. Lastly, an optimized version of the SunTag translation site imaging tool using a spaghetti-monster-based structural support is presented that can be used in future studies to address many unanswered questions.

Keywords: Stress granules, G3BP1, SunTag Imaging, TREAT, Luciferase assay, mRNA Tethering.

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1

Introduction

1.1 mRNA life cycle

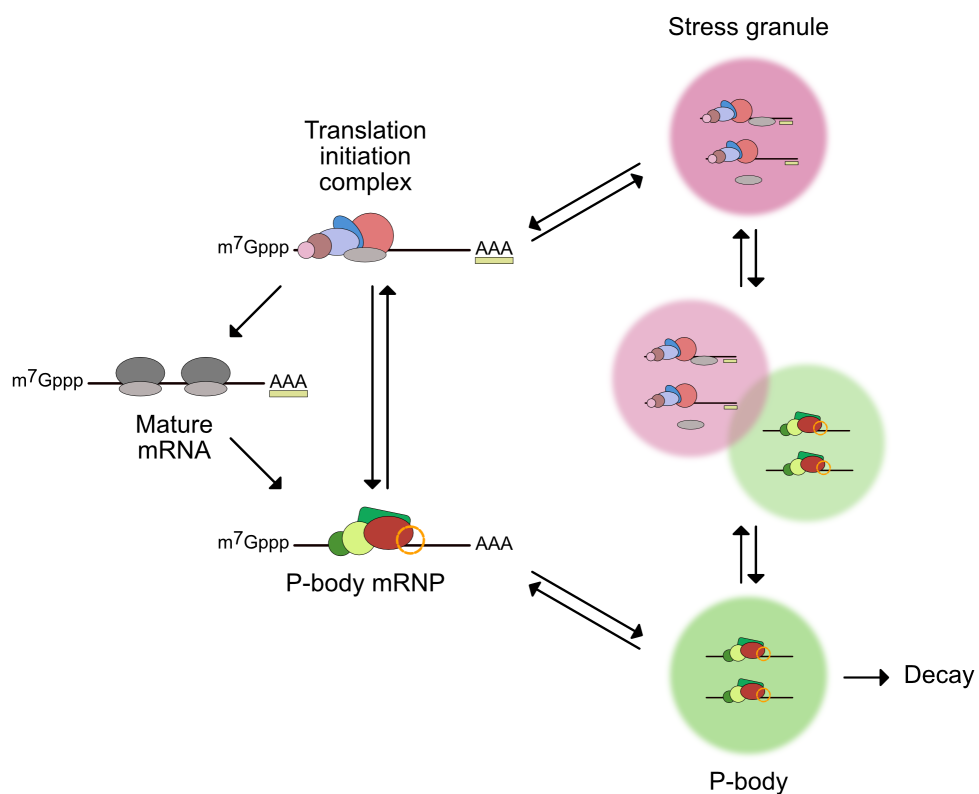
Transcription is the first step in gene expression in which a premature RNA molecule is made from a gene's DNA template. In eukaryotes, splicing, adding a 5' cap and poly-A-tail, as well as binding of cap- and RNA-binding proteins yields mature messenger RNAs (mRNAs). As transcription and translation are compartmentally separated, these mRNAs are then typically exported through nuclear pore complexes, which perforate the nuclear envelope. Once in the cytoplasm, the acquired protein coat determines the fate of the mRNA [1]. One of these fates is translation to produce proteins. This process involves polysome (multiple ribosomes translating an mRNA) assembly and messenger ribonucleoprotein complex (mRNP) remodeling and can occur immediately or delayed, such as in the case of various developmental transcripts (reviewed in [2]). If a protein no longer needs to be produced polysomes might get disassembled. Based on the interactions of mRNA-binding proteins, mRNAs can now be targeted for degradation (reviewed in [3]). Alternatively, subsets of these mRNAs can be packaged into RNA granules (Figure 1.1A). RNA granules are cytoplasmic, membrane-less, and evolutionarily conserved aggregates containing various proteins involved in translation initiation and repression (reviewed in [4]).

Stress granules (SGs) are a prominent type of RNA granules that exclusively

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form under stress. Cellular stress arises from various external or internal factors and can trigger the integrated stress response pathway. Typical extrinsic factors range from starvation, infection, hypoxia, the presence of oxidants, and many more. The largest intrinsic factor is endoplasmic reticulum (ER) stress through the accumulation of unfolded or misfolded proteins. All of these acti-

A



B

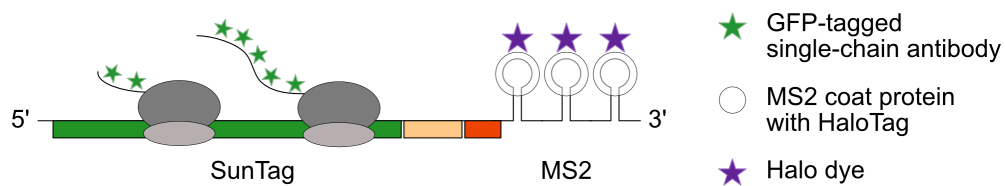


Figure 1.1: (Continued on the following page.)

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Figure 1.1: mRNA life cycle and translation site imaging. (A) Overview of a mature, translating mRNAs life cycle. Protein components of the translation initiation complex together with their bound mRNAs can aggregate into membrane-less organelles termed stress granules. A similar type of RNA granule, the P-bodies, have been associated with mRNA decay and a different set of proteins. Both granules can interact and possibly exchange components. (B) Schematic description of SunTag imaging to visualize and quantify translational activity on a single-molecule level. mRNA building blocks (from left to right): SunTag cassette, *Renilla* luciferase, proteotuner tag, and MS2 stem-loops.

vations converge to the phosphorylation of S52 on the α subunit on eukaryotic translation initiation factor 2 (eIF2 α). This phosphorylation deactivates eIF2 α and inhibits global protein synthesis which can lead to an up-regulation of stress-response genes (reviewed in [5]). Multiple immunofluorescence-based studies have shown that SGs showed an enrichment in initiation factors including eIF2 but not for other translation-related machinery components such as the 60S ribosomal subunit [6]. Some proteins found in SGs such as the T-cell intracellular antigen-1 (TIA-1) [7] and Ras GTPase-Activating Protein-Binding Protein 1 (G3BP1) [8] have been found to induce SG formation upon overexpression. The exact nature of SG formation is unknown but various mechanisms from prion-like assembly [9, 10], maturation of mRNA Processing Bodies (P-bodies) (described below) [11], and microtubule involvement have been proposed [12]. While SG formation seems to be a conserved phenomenon throughout eukaryotes, their function also remains unclear. Previous studies have suggested an involvement of SGs in mRNA stability and translational repression (reviewed in [13]). Furthermore, SG assembly coincides with an accumulation of stalled translation preinitiation complexes in conjunction with polysome disassembly and an increase in uncoated cytoplasmic mRNAs [14]. This, along with the observed enrichment of SGs in initiation factors leads to the assumption that SGs exclusively contain translationally repressed mRNAs (reviewed in [15]). However, recent findings from our laboratory have reported

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that some SG-localized mRNAs can still be translated [16].

A different type of RNA granules, the P-bodies, were originally described as mouse XRN1p (a highly conserved 5'-3' exonuclease) foci in the cytoplasm [17]. P-bodies, unlike SGs, are present in unstressed physiology. However, stress can further increase their formation (reviewed in [18]). Contrasting SGs, they contain non-overlapping protein components including multiple nucleases that have made them a proposed site of mRNA decay [19].

Many questions regarding RNA granules, and especially SGs, remain largely unanswered. For example, their dynamic formation and disintegration as well as what determines their composition remains unclear. Several large scale analyses of SGs using spatial proteomics [20, 21, 22] and RNA sequencing [23] approaches have given important insights into the content of SGs but not their function. In a recently published trio of papers [24, 25, 26] a combination of experiments and computational modeling provided more insights into the formation of SGs. They hinted at G3BP1 being a key hub in the SG protein interaction network playing a central role during SG formation by RNA-induced conformational changes. None of these studies, however, gave insights into the biological function that G3BP1 or other SG components exert on single mRNAs. Recently established single-molecule techniques allow for real-time analyses in live cells. Compared with extensively used bulk assays, these give a higher spatio-temporal resolution of cellular processes. In this study, single-molecule imaging techniques will be used to address the effects of the SG-protein G3BP1 on an mRNA's life cycle.

1.2 Translation site imaging

The stellar explosion SUpernova (SunTag) imaging approach allows for live cell imaging-based analysis to visualize and quantify translation sites [27]. The SunTag system relies on three core components: a reporter RNA containing a SunTag cassette and MS2 stem-loops, a green fluorescent protein (GFP)-

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tagged single-chain antibody (scFv) against GCN4, and an MS2 coat protein fused to a HaloTag (Figure 1.1B).

The MS2 system has been widely adopted to tag and visualize specific mRNAs. MS2 stem-loops serve as structural motifs and are recognized by the MS2 bacteriophage capsid RNA-binding protein MS2 (MCP) (reviewed in [28]). Typical MS2 tagging involves the visualization of single mRNAs encoding an array MS2 stem-loops by an MCP-GFP fusion. Multiple MS2 stem-loops are used to increase the local concentration of MCP bound fluorescent molecules allowing for significant signal amplification. Previous studies have used this method to investigate mRNA metabolism by tethering mRNAs to membranes [29], to proteins [30], or to themselves as shown in a recent optogenetic mRNA clustering approach [31].

Currently available standard SunTag cassettes consist of 24 GCN4 epitope repeats that, upon translation and exposure of the GCN4 epitopes, can be recognized and bound by the co-expressed GFP-tagged scFv. As translation continues, more epitopes get synthesized allowing more scFvs to bind. This increases the local concentration and thereby brightens the fluorescent signal. Once translation is terminated, SunTag peptides together with attached scFvs are released from the mRNA causing a decrease in signal intensity. Degradation of the mature protein is increased due to a proteotuner tag that is located C-terminal of the SunTag cassette and that targets the newly formed protein for proteasomal degradation [32]. The MS2 coat protein-HaloTag fusion in conjunction with a soluble Halo dye is used to visualize the MS2 stem loops on the reporter RNA described above. Taken together, the system allows for accurate tracking of single RNAs and simultaneous quantification of translation.

This study focuses on the understanding of how SGs form and how they function. Using modified components of the SunTag imaging system, G3BP1 is used as the main antagonist to provide interesting insights into the unknown world of SGs. G3BP1 will be tethered to a reporter mRNA by fusion to MCP-Halo. Single-molecule imaging tools including SunTag will be used to investi-

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gate the effects of an increased local concentration of SG proteins on mRNA dynamics. Lastly, an improvement to the existing SunTag system is proposed.

2

Methods

2.1 Key resources

Table 2.1: Antibodies

| Reagent or Resource | Source | Identifier |
|---------------------|--------|------------|
| Alexa Fluor 647 | Abcam | ab150075 |
| TIA-1 | Abcam | ab40693 |

Table 2.2: Experimental Models: Cell lines

| Reagent or Resource | Source | Identifier |
|---|------------------------------|------------|
| Hela 11ht | Weidenfeld et al., 2009 [33] | N/A |
| Hela 11ht + scAB-GFP + Renilla-MS2 | This study | N/A |
| Hela 11ht + scAB-GFP + smGCN4-Renilla-MS2 | This study | N/A |
| Hela 11ht + scAB-GFP + SunTag-Renilla-MS2 | This study | N/A |
| Hela 11ht + TREAT | This study | N/A |

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Table 2.3: Chemicals and Peptides

| Reagent or Resource | Source | Identifier |
|--------------------------------|-------------------------------------|-------------|
| Amino-11-ddUTP | Lumiprobe | 15040 |
| Atto 647N NHS ester | Sigma-Aldrich | AD 647N-31 |
| Bovine serum albumin | Sigma-Aldrich | A2153-50G |
| Dextran sulfate | Sigma | D6001-50G |
| Doxycycline | Sigma | D3891-1G |
| FluoroBrite DMEM | Life Technologies | A1896703 |
| Formamide (deionized) | Chemicon | S4117 |
| Ganciclovir | Sigma-Aldrich | G2536-100MG |
| JF585 HaloTag ligand | Grimm et al., 2015, 2017 [34, 35] | N/A |
| Lipofectamine 2000 | Invitrogen | 11668019 |
| Opti-MEM™ | Gibco | 31985070 |
| Paraformaldehyde (aqueous) | 20% Electron Microscopy Sciences | 15713 |
| ProLong Gold Antifade Mountant | Molecular Probes, Life Technologies | P36935 |
| Puromycin | Invivogen | ant-pr-1 |
| Sodium Arsenite solution | Sigma | 35000-1L-R |

Table 2.4: Critical Commercial Assays

| Reagent or Resource | Source | Identifier |
|-------------------------------|--------|------------|
| <i>continues on next page</i> | | |

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| Reagent or Resource | Source | Identifier |
|---------------------------------|---------|------------|
| Bradford Protein Assay | Biorad | 5000006 |
| Renilla Luciferase Assay System | Promega | E2810 |

Table 2.5: Oligonucleotides

| Reagent or Resource | Source | Identifier |
|---|--------|------------|
| Oligodeoxyribonucleotides and primers are listed in Appendix A. | N/A | N/A |

Table 2.6: Recombinant DNA

| Reagent or Resource | Source | Identifier |
|--------------------------|----------------------------|-----------------|
| G3BP1-GFP-GFP | Wilbertz et al., 2019 [36] | Addgene #119950 |
| NLS-stdMCP-stdHalo | Voigt et al., 2017 [37] | Addgene #104999 |
| NLS-stdMCP-stdHalo-G3BP1 | This Study | N/A |
| NLS-stdMCP-stdHalo-Rh1 | This Study | N/A |
| pCAGGS-FLPe-IRESpuro | Beard et al., 2006 [38] | Addgene #20733 |
| Renilla-MS2v5 | This Study | N/A |
| scAB-GFP | Voigt et al., 2017 [37] | Addgene #104998 |

continues on next page

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| Reagent or Resource | Source | Identifier |
|--------------------------|------------------------------|--------------------|
| smGCN4-Renilla-MS2v5 | This study | N/A |
| SunTag-Renilla-MS2v5 | Wilbertz et al., 2019 [36] | Addgene #119945 |
| SunTag-Renilla-PP7-MS2v4 | Horvathova et al., 2017 [39] | N/A |

Table 2.7: Critical Equipment

| Reagent or Resource | Source | Identifier |
|--|----------------------|------------|
| 405 iBeam Smart | Toptica Photonics | N/A |
| 488 iBeam Smart | Toptica Photonics | N/A |
| 561 Cobolt Jive | Cobolt | N/A |
| 639 iBeam Smart | Toptica Photonics | N/A |
| 96-Well White Polystyrene Microplates | Costar | 07-200-589 |
| CFI Plan Apochromat Lambda 100x Oil/1.45 Objective | Nikon | N/A |
| CSU-W1 Confocal Scanner Unit | Yokogawa | N/A |
| Glass Coverslips | Paul Marienfeld GmbH | 117580 |
| iXon-Ultra-888 EMCCD Cameras | Andor | N/A |
| Mithras Multimode Microplate Reader LB 940 | Berthold | 38099 |

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| Reagent or Resource | Source | Identifier |
|---|------------------------------------|------------|
| MS-2000 Motorized Stage | Applied Scientific Instrumentation | N/A |
| TetraSpeck™ Fluorescent Microspheres Size Kit | Thermo Fisher Scientific | T14792 |
| Ti2-E Eclipse Inverted Microscope | Nikon | N/A |
| VS-Homogenizer | Visitron Systems GmbH | N/A |

Table 2.8: Software and Algorithms

| Reagent or Resource | Source | Identifier |
|-------------------------|------------------------------|--|
| Affinity Designer 1.8.2 | Serif (Europe) Ltd | affinity.serif.com/designer/ |
| Benchling 2020 | Benchling | benchling.com |
| Fiji 2.0.0-rc-69/1.52p | Schindelin et al., 2012 [40] | fiji.sc |
| Fluffy 0.2.2 | Eichenberger, 2020 [41] | github.com/BBQuercus/fluffy |
| KNIME 3.7.2 | Berthold et al., 2009 [42] | knime.com/knime-analytics-platform |
| PyMOL 2.3.3 | Schrodinger LLC. | pymol.org |
| Python 3.7.4 | Python Software Foundation | python.org |
| TrackMate v5.2.0 | Tinevez et al., 2017 [43] | imagej.net/TrackMate |

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| Reagent or Resource | Source | Identifier |
|---------------------|-----------------------|---|
| VisiView 4.4.0 | Visitron Systems GmbH | visitron.de/ products/ visiviewr-software |

2.2 Method details

Cell lines and culture details

The previously described HeLa-11ht cell line [33] was used for this study. The integrated Flp-RMCE (recombinase-mediated cassette exchange) site allows for controlled, single-copy, genomic integration of a target gene. In addition, to induce inserted target genes reversibly with doxycycline, cells are expressing a reverse tetracycline controlled transactivator (rtTA2S-M2). Dulbecco's Modified Eagle Medium (DMEM) containing 4.5 g/l glucose, 100 U/ml Penicillin, 100 µg/ml Streptomycin, 4 mM L-Glutamine, and 10% v/v Fetal Bovine Serum (FBS) was used to culture HeLa cells. Cells were maintained at 37°C with 5% CO₂. Transient transfections were performed with Lipofectamine 2000 transfection reagent (Invitrogen) with Opti-MEM reduced serum medium (Gibco) according to manufacturer's instructions but scaled down to 0.5 µg plasmid DNA and 2 µl Lipofectamine 2000 per 1 ml growth medium.

Plasmid construction

To generate the G3BP1 coat protein fusion, the genomic sequence encoding the full-length G3BP1 Isoform 1 (Q13283-1) was inserted in-frame, downstream of the HaloTag sequence. A 33 nt linker sequence was kept between HaloTag and G3BP1 to promote flexibility between the domains. Assembly was performed by PCR amplification of G3BP1 and the MCP-Halo-linker backbone, followed by Gibson cloning [44]. The resulting construct contains a constitutive UbiC promoter, SV40 NLS (nuclear localization signal), stdMCP (MS2 coat protein), HaloTag, 33 nt linker sequence, G3BP1 with a stop codon, and

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a Woodchuck Hepatitis Virus Posttranscriptional Regulatory Element (WPRE) improving expression in lentiviral transfection (Figure B.3).

The smGCN4 reporter sequence containing GCN4 epitopes was synthesized into a pUC57-Kan expression vector. Assembly was performed by PCR amplification of smGCN4 and the *Renilla*-MS2v5 backbone followed by Gibson cloning. The resulting plasmid contains a Tet-CMV (cytomegalovirus) promoter followed by smGCN4, *Renilla* luciferase, FKBP domain with a stop codon, MS2v5 stem-loop cassette (24x), CTE (constitutive transport element, and SV40 polyA tail (Figure B.4).

Reporter cell line generation

A day before selection, HeLa cells were seeded into a 6-well plate. The targeting plasmid containing the reporter (2 µg) and pCAGGS-FLPe-IRESpuo plasmid (2 µg) were transfected using Lipofectamine 2000 (Invitrogen) according to manufacturers' protocol [38]. On the next day, cells were preselected with 5 µl/ml puromycin (Invivogen). Two days later, the puromycin-containing medium was removed and replaced with fresh growth medium containing 50 µM ganciclovir (Sigma-Aldrich). The selection was performed for 10-14 days to get resistant colonies that had undergone RCME. Single-cell sorting into a 96-well plate was performed. Clones were tested for reporter expression using luciferase assays.

***Renilla* luciferase assays**

Reporter production was measured with Promega's *Renilla* Luciferase Assay System. The described cell lines were seeded on 12-well plates. The next day, plasmid DNA was transfected once cells reached approximately 70% confluence as described above. On the subsequent day, the expression of reporters was induced with 1 µg/ml doxycycline (Sigma) for 3 hours unless otherwise specified. Stress was induced by adding 1 mM/ml sodium arsenite (Sigma) for the last 1 hour of induction. To measure recovery, cells were washed twice with

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PBS and replaced with fresh culturing medium. At the specified time points, cells were washed once with PBS and lysed with 250 µl Passive Lysis Buffer per well. To ensure full cellular lysis, plates were gently shaken for 15 minutes at room temperature. 30 µl lysate was transferred to 96-well EIA/RIA Plate (Costar) in triplicates. Bioluminescence was measured 2 seconds after injecting 12 µl *Renilla* Luciferase Assay Reagent per well. Measurements were performed on the Mithras Multimode Microplate Reader LB 940 (Berthold). Data were normalized by protein concentration measured by standard Bradford Protein Assay (Biorad). Means of at least 3 biological replicates (each with 3 described technical replicates) were calculated.

Single-molecule fluorescence in situ hybridization (FISH)

Single-molecule RNA detection against the *Renilla* luciferase coding sequence and MS2 (v5) was performed using Stellaris FISH probes (Biosearch Technologies). Probes targeting MS2 (v4) were made by enzymatic oligonucleotide labeling [45] with Amino-11-ddUTP (Lumiprobe) and Atto647-NHS (ATTO-TEC).

Cells were seeded on glass coverslips (Paul Marienfeld GmbH) placed in 12-well plates. The next day, cells were transfected as indicated. On the subsequent day, cells were treated as indicated, followed by two washes with PBS and fixation in 4% paraformaldehyde (Electron Microscopy Sciences) diluted in PBS for 5 minutes. Cells were washed thrice with PBS and permeabilized with 0.5% Triton X-100 diluted in PBS for 5 minutes (if immunofluorescence was performed) or overnight at 4°C in 70% Ethanol. Cells were washed two more times with PBS and prehybridized with wash buffer (2x SSC (Invitrogen), 10% v/v formamide (Abcam)) twice for 5 minutes. Coverslips were then incubated between 4 and 16 hours with hybridization solution (2x SSC, 10% v/v formamide, 10% v/v dextran sulfate, 0.5% v/v BSA, 200 nM FISH probes) at 37°C in humidified chambers. Cells were washed once more with wash buffer for 30 minutes and twice with PBS. Immunofluorescence staining was performed according to the "Immunofluorescence" section from BSA blocking

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to mounting. Samples were then mounted on ProLong Gold Antifade Mountant with DAPI. Imaging was performed as described in the “Live-cell imaging” section using sequential, single-camera acquisition.

Immunofluorescence

Cells were seeded on glass coverslips (Paul Marienfeld GmbH) placed in 12-well plates. The next day, cells were transfected as indicated. On the subsequent day, cells were treated as indicated, followed by two washes with PBS and fixation in 4% v/v paraformaldehyde (Electron Microscopy Sciences) diluted in PBS for 5 minutes. Cells were washed thrice with PBS and permeabilized with 0.5% v/v Triton X-100 diluted in PBS for 5 minutes. Cells were washed two more times in PBS and incubated in 3% w/v bovine serum albumin (BSA) (Sigma-Aldrich) diluted in PBS for 1 hour. Primary antibodies (diluted in 1% w/v BSA in PBS) were incubated for 1 hour at room temperature or overnight at 4°C. After washing three times with PBS, the cells were incubated with secondary antibodies (diluted in 1% w/v BSA in PBS) for 1 hour at room temperature. Cells were washed twice with PBS and mounted on glass slides in ProLong Gold Antifade Mountant with DAPI. Imaging was performed as described in the “Live-cell imaging” section using sequential, single-camera acquisition.

Live-cell imaging

Cells were seeded on 35 mm glass-bottom μ -Dish (ibidi GmbH). The next day, cells were transfected as indicated. On the subsequent day, cells were treated as indicated. During the last 15 minutes of treatment, the medium was supplemented with JF585 HaloTag ligand, obtained from L. Lavis (Janelia Research Campus) [34, 35], at 100 nM final concentration. After incubation, cells were washed once with PBS and kept in FluoroBrite DMEM (Life Technologies) containing 10% v/v FBS and 4 mM L-glutamine. Cells were imaged within 15 minutes of medium exchange on an inverted Ti2-E Eclipse microscope (Nikon)

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with a CSU-W1 Confocal Scanner Unit (Yokogawa), two back-illuminated EM-CCD cameras iXon-Ultra-888 (Andor), an MS-2000 motorized stage (Applied Scientific Instrumentation), and VisiView imaging software (Visitron Systems GmbH). Specimens were illuminated with 561 Cobolt Jive (Cobolt), 405 iBeam Smart, 488 iBeam Smart, and 639 iBeam Smart lasers (Toptica Photonics) and a VS-Homogenizer (Visitron Systems GmbH). All images were acquired with a CFI Plan Apochromat Lambda 100X Oil/1.45 objective (Nikon). This setup results in a pixel size of 130 nm. Unless otherwise indicated, excitation was performed with a 50 ms exposure time in a single plane. The second camera was used to detect the 488 nm channel. To ensure proper camera alignment, TetraSpeck™ Fluorescent Microspheres Size Kit (Thermo Fisher Scientific) was used to image 0.5 μ m fluorescent beads after each imaging session. Cells were maintained at 37°C and 5% CO₂ within an incubation box.

2.3 Quantification and statistical analysis

Detection of mRNA spots from smFISH and colocalization

Using custom-built python scripts, images were registered and maximum intensity projected in all channels. Cells were segmented in two steps. First, a sequential workflow comprising Gaussian filtering steps, Otsu thresholding [46], and distance transform calculations of the DAPI channel (405 nm) was used to segment individual nuclei. Subsequently, the nuclei were used as seeds to perform a watershed segmentation into a previously thresholded cytoplasmic map. Cytoplasmic channels were chosen based on their signal uniformity (561 nm *Renilla* luciferase or MS2v5 probe channel). SGs were segmented in Fluffy [41]. mRNA spots were detected with a Laplacian of Gaussian filter. Thresholds for cellular segmentation and spot detection were kept constant throughout a dataset. To colocalize mRNA spots, Euclidean distances were calculated between spots across both channels. To ensure

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proper channel-to-channel association of mRNA spots, cells with a high spot density were filtered out. Two mRNAs were considered as colocalized when their coordinates are less than 2 px (240 nm) apart.

Processing of live-cell imaging data

Images of the fluorescent beads were used to perform channel alignment. Using the `pystackreg` python package [47] an affine transformation was registered. This model was subsequently re-applied to all images acquired with the secondary camera. Fiji [40] was used to create representative movies and images via cropping, brightness adjustments, channel merging, and scale bar annotation. TrackMate [43] was used to display tracked particles.

Quantification of translational status

Tracking-based quantification of translation was performed using a KNIME [42] workflow described in Mateju et al. [16] in section “Track-based analysis of translational status and colocalization” without including the described SG association component.

Statistics

Results are presented as the mean \pm confidence interval (95%) of independent experiments. Significant differences between variables are based on independent sample t-tests. P-values are indicated using stars in each figure. Each star corresponds to the following p-values:

- ns: $5e^{-2} < p \leq 1$
- *: $e^{-2} < p \leq 5e^{-2}$
- **: $e^{-3} < p \leq e^{-2}$
- ***: $e^{-4} < p \leq e^{-3}$
- ****: $p \leq e^{-4}$

2.4 Data and code availability

The code used to analyze and visualize the data together with plasmid maps is available on GitHub¹. All data supporting the findings of this study are available on request.

¹ <https://github.com/BBQuercus/master-thesis>

3

Results

3.1 G3BP1 tethering affects mRNA localization

Proteins of the G3BP RNA-binding family are key components of SGs. In mammals, G3BP1 (and its paralog G3BP2) were reported to be essential in nucleating the formation of SGs [48]. SGs can not only be induced by stress (e.g. sodium arsenite) but also by the overexpression of G3BPs [8]. In addition, a recent proteomics-based study by Markmiller et al. [20] showed that "many well-characterized SG proteins (e.g., G3BP1, TIA1, CAPRIN1, PABPC1, FMR1, and ATXN2) were identified as highly significant interactors" suggesting an interplay of SG proteins even in unstressed conditions. This finding might suggest the formation of an mRNP complex containing SG proteins allowing for a rapid assembly of SGs.

The exact role of G3BPs in SG assembly as well as in unstressed cells is still largely unknown. As reviewed in Alam and Kennedy [49] various functions have been attributed to this protein family. These range from transcript destabilization and repression to the polar opposite in transcript stabilization. Furthermore, G3BPs also showed effects on transcript localization and sequestration to virus-induced foci (reviewed in [50]). The only consensus is G3BP's involvement in mRNA translational control. The subsequent experiments attempt to clarify this disunity by analyzing G3BP1's effect on mRNA transcripts at a single-molecule level. For this reason, I designed an assay to directly tether G3BP1 to reporter mRNAs and thereby allow its effect on trans-

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lation and mRNA degradation to be measured as well as to promote mRNA accumulation into SGs.

This study focussed on a reporter RNA containing a SunTag cassette, the gene encoding for *Renilla* luciferase, a proteotuner tag, and MS2 stem-loops (Figure 3.1A). In order to promote mRNA recruitment to SGs, I fused G3BP1 MCP-Halo (abbreviated as MCPG3). The construct was validated by FISH against the reporter RNA followed by immunofluorescence for the endogenous

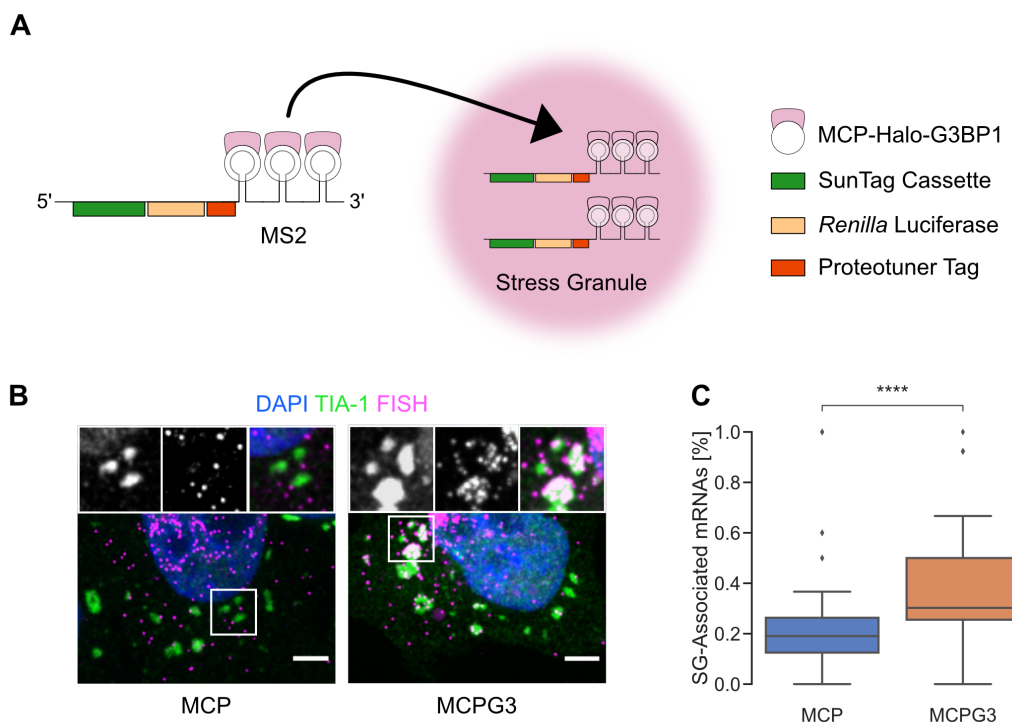


Figure 3.1: Design of MCPG3. (A) Schematic depiction of MCPG3 function. (B) Representative fluorescence images of the SG marker TIA-1 and the tethered RNAs. To induce SGs, cells were treated with sodium arsenite (1 mM) for 1 hour. Panels show TIA-1, FISH, Merge respectively. Scale bars, 10 μ m. (C) Quantification of the mRNA recruitment to SGs. Number of cells quantified (left to right): 27, 21. Two replicates.

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SG marker TIA-1 [7] (Figure 3.1B). SGs were induced by sodium arsenite treatment (1 mM for 1 hour). As a control, an identical construct only lacking G3BP1 fusion was used. The G3BP1 fusion construct resulted in a significant difference in SG association (Figure 3.1C). MCPG3 shows an approximately two-fold increase in SG associated mRNAs compared to the control while not causing mRNA aggregation in unstressed conditions (Supplementary Figure B.1). In addition, both the area and intensity of SGs seem to not be affected by the G3BP1 fusion (Supplementary Figures B.2A and B.2B respectively). Taken together, fusing G3BP1 to MCP seems to be a reliable way to recruit functional G3BP1 to the local vicinity of mRNAs and thereby promote localization to SGs.

3.1.1 Reporter expression is impacted by G3BP1

Having shown that G3BP1 can be recruited to mRNAs, I wanted to investigate the effect on translational activity. *Renilla* luciferase assays are used to measure reporter expression on a global level. As G3BP1 was shown to be involved in stress-induced events as well as in unstressed conditions, the luciferase activity at multiple time points in unstressed cells as well as in cells recovering from oxidative stress was measured.

In unstressed cells, G3BP1 recruitment resulted in increased luciferase readings over time (Figure 3.2A). This suggests that the presence of G3BP1 on a transcript might promote protein synthesis. Comparing both constructs during stress recovery, one can observe a slightly less significant but still noticeable increase in luciferase activity for MCPG3 expressing cells (Figure 3.2B). During stress conditions, most mRNAs get translationally silenced [36] which explains the comparative dropoff in luciferase activity directly after stress. Interestingly, the largest effects are at later time points and during non-stress conditions while not being immediately after stress recovery. Preliminary experiments using different durations of stress have yielded similar results. SGs have been observed to persist between a few minutes to hours after removal

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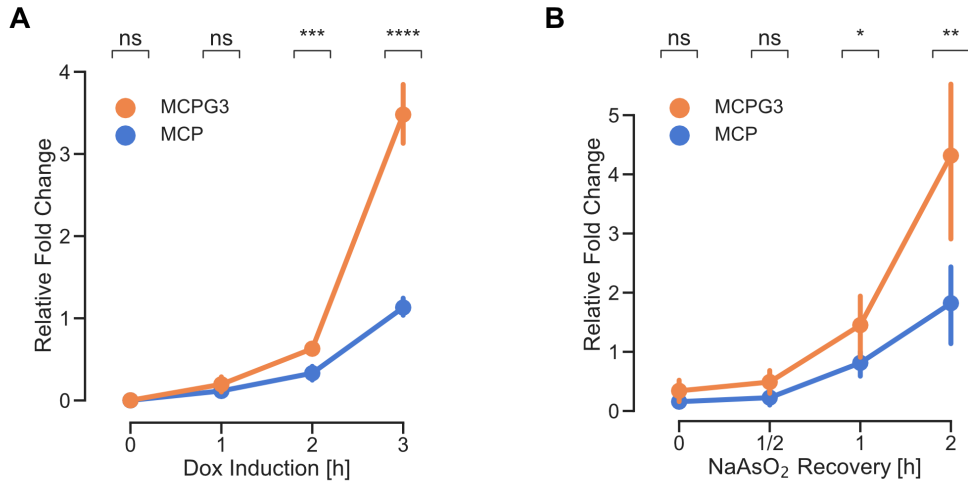


Figure 3.2: G3BP1 tethering increases reporter translational activity. (A) Luciferase assays at different time points during induction. (B) Sodium arsenite stress recovery time points after induction. Cells were induced for 3 hours and stressed with sodium arsenite (1 mM) in the last hour. Timepoint capture started immediately after washing out the stressor. (A and B) Data normalized to the 3-hour induction condition of MCP. Five replicates.

of the stressor [51, 36]. This correlates with the time when an increase in luciferase activity can be observed again.

3.1.2 G3BP1 does not alter mRNA translation

The observed differences in luciferase activity described in Section 3.1.1 can result from various causes. These differences in reporter protein synthesis levels could arise from an increase in translational activity of mRNAs or an increase in mRNA stability. To investigate the former possibility, I decided to use SunTag imaging, to directly quantify translational activity (see Section 1.2).

In SunTag imaging, the fluorescence intensity is a direct measure of translation rate. Therefore, to analyze the translation activity of MCPG3-bound mRNAs, the intensity of translation site spots was quantified over at least four

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consecutive timeframes with a maximum of one gap frame.

When comparing SunTag tracks of single mRNAs in unstressed cells expressing MCPG3 or control MCP (Figure 3.3A), both conditions show similar fluorescence intensities suggesting G3BP1 does not have a major effect on mRNA translation rates. Similarly, the grouping of all SunTag tracks measured within a single cell shows comparable mean intensities in cells expressing MCPG3 or control MCP. Equivalently to the image-level analysis, the mean cellular intensities (Figure 3.3B) do not yield different results. The average intensity of all tracks per cell does not appear to be affected by G3BP1 fusion.

As previously mentioned, most mRNAs get translationally silenced during stress conditions [36]. To image translation sites despite this silencing and to investigate if increased recruitment of mRNA to SGs by MCPG3 has an effect on translation rates, images were acquired 15 to 30 minutes after removing the stressor. Nonetheless, cells did not fully recover the translational silencing leading to lower readings of fluorescent intensity in the stress condition compared to unstressed cells. However, in this case, as well, G3BP1 tethering does not lead to significant differences in translational activity.

Lastly, translation can also increase due to a higher number of translating transcripts per cell. However, as is evident from Figure 3.3C the number of translating tracks stays consistent in all experiments. This suggests that G3BP1 does not have a profound impact on translation activity.

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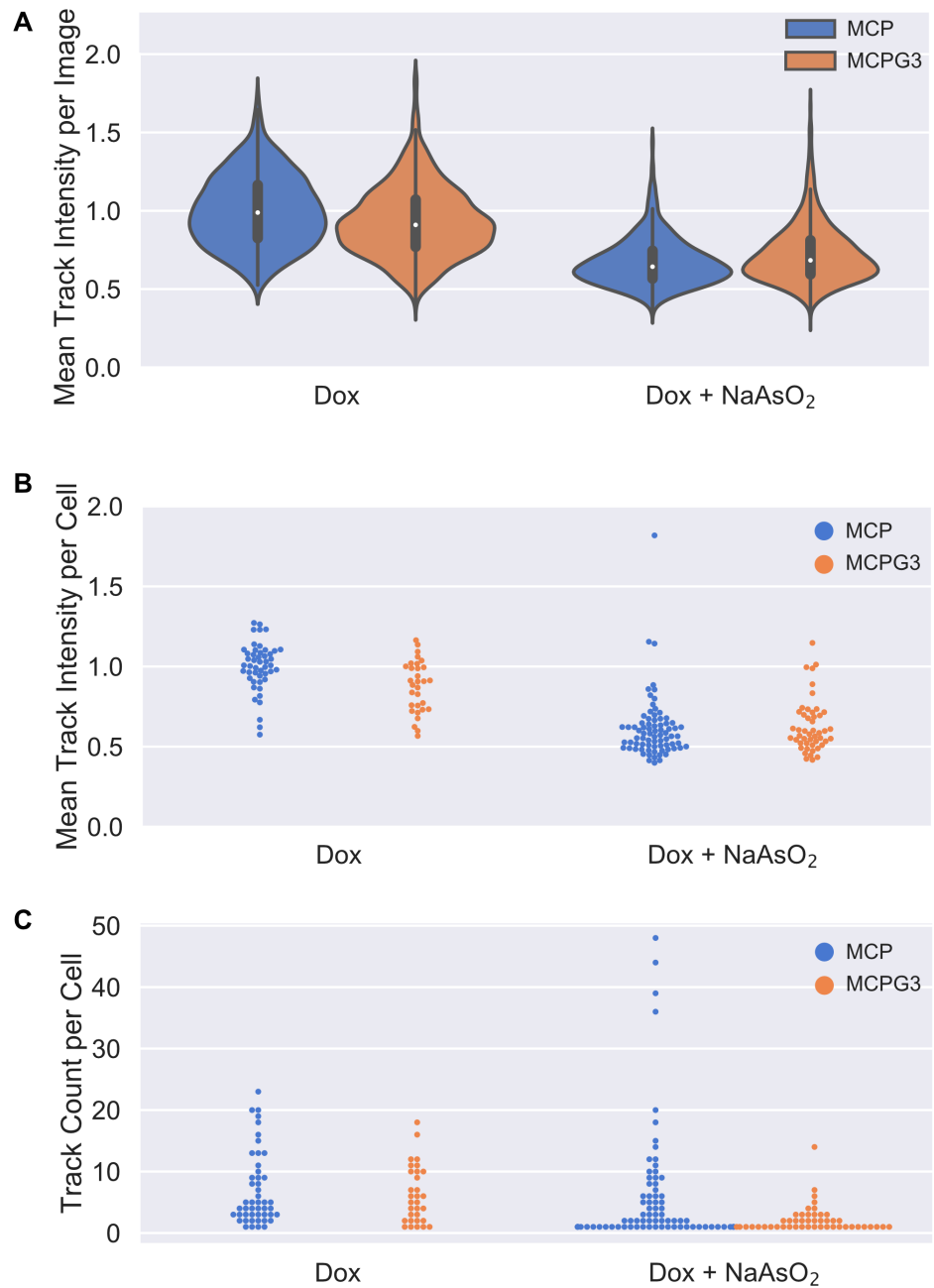


Figure 3.3: (Continued on the following page.)

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Figure 3.3: G3BP1 does not affect translational activity. (A) Distribution of the mean fluorescence intensity of tracked mRNAs in the full-sized image. The total number of tracks (left to right): 715, 1064, 997, 3442. (B) Track intensity averaged per cell. (A and B) Data normalized to the Dox induction condition for MCP. (C) The number of tracks registered per cell. (B and C) The total number of cells (left to right): 48, 77, 31, 48. Single replicate.

3.1.3 Transcript stability is affected by G3BP1 tethering

SunTag measurements (see Section 3.1.2) did not suggest any effect of G3BP1 on translational activity. Alternatively, a decrease in mRNA degradation could also explain the increased *Renilla* luciferase readings. To ask whether G3BP1 has an mRNA stabilizing effect, I compared the half-lives of transcripts between MCPG3 and the control using 3(three)'-RNA end accumulation during turnover (TREAT) [39].

TREAT uses a slightly modified reporter mRNA containing viral RNA pseudo-knot (PK) structures upstream of the MS2 stem-loops. These PKs can block Xrn1 mediated 5'-3' degradation of a transcript by sequestering the 5' phosphate group [52]. Subsequently, by using FISH probes against *Renilla* luciferase upstream of PKs (showing intact mRNAs) and FISH probes against MS2 stem-loops downstream of PKs (showing both intact and degraded mRNAs) one can quantify the number of degradation intermediates.

To look at transcript stabilization, a cell line expressing the aforementioned TREAT mRNA reporter was used. Five time points were chosen to match TREAT half-life after a 1 hour induction followed by thorough washing to remove doxycycline to halt any further transcription. As can be seen in Figure 3.4A, transcripts are exported from the nucleus into the cytosol over the course of the experiment. This suggests that no new transcripts are produced in the nucleus and that MCPG3 does not negatively affect nuclear export.

When comparing the percentage of intact mRNAs, i.e. transcripts still containing FISH spots against both *Renilla* luciferase and MS2, between MCPG3

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and the control, I observed a significant difference (Figure 3.4B). The G3BP1 fusion shows a significant reduction in degraded transcripts compared to the control suggesting an increase in mRNA stability. This effect is not reflected in the translation site-based mRNA count from the SunTag measurements (see Section 3.1.2) which might arise from the nature of the SunTag cassette that will be discussed in the following Section 3.2.

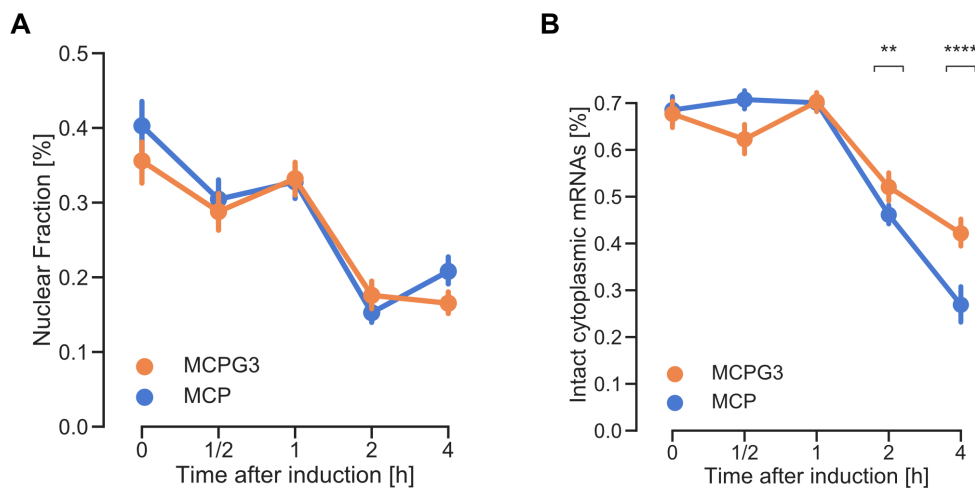


Figure 3.4: mRNA stability increases by G3BP1 tethering. (A) The fraction of nuclear mRNAs (of total cellular mRNAs) at different time points after induction. (B) Percentage of intact cytoplasmic mRNAs at different time points. (A and B) Number of cells quantified (ascending, MCP then MCPG3): 117, 154, 200, 175, 55, 114, 148, 167, 111, 132. Single replicate.

3.2 How can the SunTag reporter be improved?

The original design of the SunTag cassette consists of 24 GCN4 repeats. These completely disordered repeats have previously led to the observations of decreased *Renilla* luciferase activity and lower induction rates than *Renilla* luciferase without SunTag. In an attempt to improve the SunTag based translation imaging system, a spaghetti monster reporter was made. Spaghetti monster green fluorescent proteins (smGFP) are essentially a non-fluorescent green fluorescent protein's (GFP) beta-barrel used to attach small epitope tags [53]. Here, three GCN4 repeats each were placed at the N-terminus, C-terminus, and 10/11-loop of the GFP beta-barrel as shown in Figure 3.5A. Previously, other tags, such as influenza hemagglutinin (HA), were used and visualized with frankenbodies (single-chain antibodies targeting various tags) [54]. In this study, I combined SunTag imaging with smGFP constructs.

Initially, while testing the reporter functionality the new reporter showed an approximative 4-fold increase in luciferase activity compared to the standard SunTag reporter (Figure 3.5B). A *Renilla* reporter was used as control which does not contain any GCN4 repeats. While the smGFP reporter drastically increases activity, it does not yet reach the levels of the control. It must be noted that the control also construct does not contain a proteotuner tag which has lowered the luciferase activity in previous experiments by enhancing the degradation of the tagged protein [55].

To test whether this new reporter allows the visualization of translation sites in live cells, stable cell lines were created and observed after 1 hour of induction. From a representative image in Figure 3.5C, one can see that cells typically have more, slightly dimmer translation sites. Quantification of images showed a significant increase in the number of translation sites per cell compared to the cells expressing the standard SunTag reporter (Figure 3.5D). The brightness of individual spots decreased around 2.7 fold (Figure 3.5E). This decrease can be explained by a lower number of GCN4 repeats available for the scFv to bind. Whereas the standard SunTag has 24 GCN4 repeats,

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smGFP only has 9 leading to a binding site difference of $2.6\bar{6}$ to 1. This suggests that the scFv binds to both tags with full occupancy.

Taken together, this study proposes a new SunTag reporter which improves some of the drawbacks of fusing a largely disordered polypeptide to a protein of interest. This new reporter allows for shorter induction times with higher mRNA numbers.

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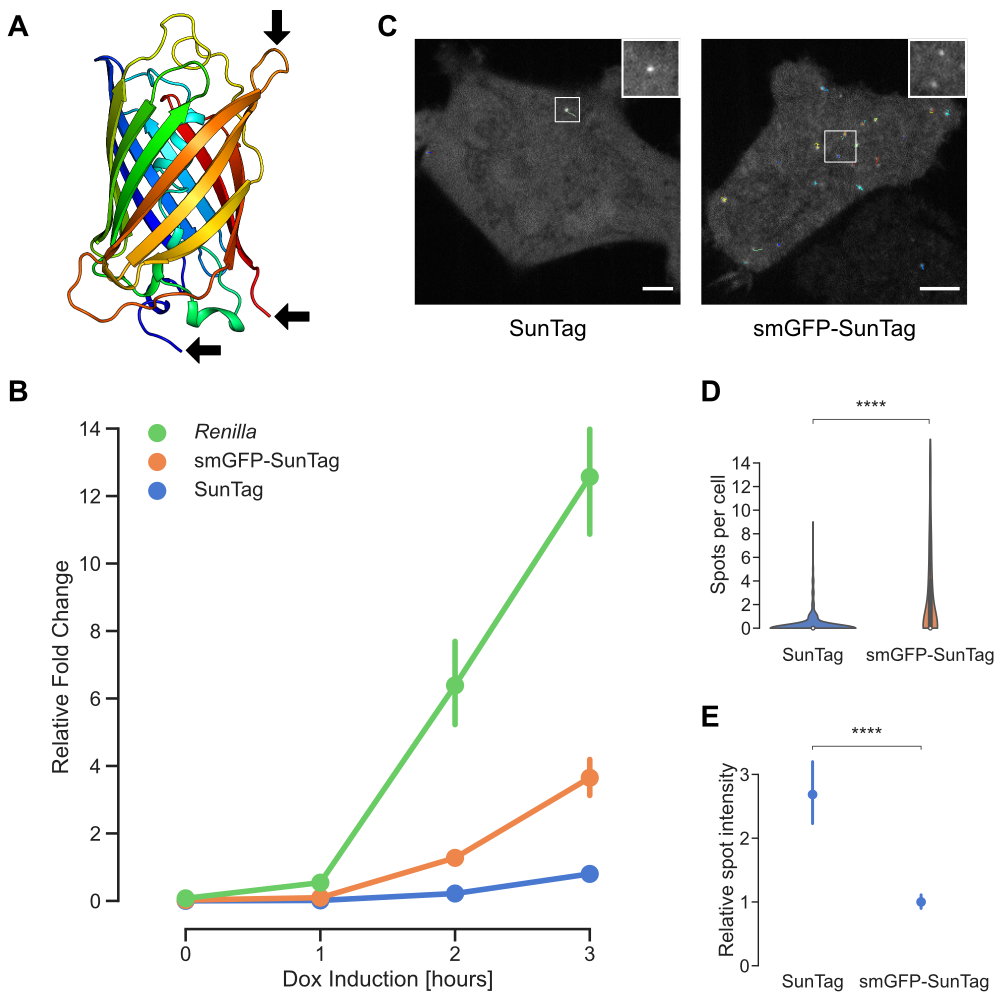


Figure 3.5: smGFP SunTag reporter increases transcript numbers. (A) Structural overview of GCN4 repeat placement on the smGFP beta-barrel (PDB ID: 1GFL [56]). (B) Relative luciferase activity comparing the standard SunTag reporter, the smGFP SunTag reporter (smGFP-SunTag), and a *Renilla* reporter without SunTag cassette. (C) Representative fluorescence images. Each recorded mRNA track is visualized with a unique color. Scale bar, 10 μ m. The inset shows spots from raw images without brightness corrections. (D) Quantification of translation site count per cell. The number of cells quantified (left to right): 286, 233. (E) Average spot intensity normalized to the smGFP reporter. The number of spots quantified (left to right): 36, 39. Two replicates. (C, D, and E) Live imaging experiment with stable cell lines imaged after 1 hour of induction.

4

Discussion

The role of many SG-proteins in unstressed cells is still unclear. An understanding of these proteins, such as G3BP1 could provide important insights into how SGs form and what functional purpose they serve. This study approaches this mystery experimentally.

My results show that fusing G3BP1 to MCP has profound effects on their associated reporter transcripts. First, mRNAs are around two-fold more likely to get recruited to SGs. The nature of transient transfections used for these sets of experiments typically does not achieve complete cellular coverage. Therefore, this likely lowers the measured numbers compared to if all cells were expressing the constructs. Furthermore, due to highly clustered mRNAs in SGs, accurate quantification was proven to be rather difficult, also decreasing the number. Second, G3BP1 has a significant effect on protein expression irrespective of cellular stress. This interesting finding showed that the role of G3BP1 in SGs might just be secondary while its more important function occurs in normal cell homeostasis. As previously proposed, SGs might arise from the necessity to sequester proteins like G3BP1 from mRNA targets in the cytosol [57]. SunTag imaging experiments were performed to observe differences in translational activity. These, however, did not show major differences suggesting that G3BP1 might only have minor roles in translational activity. Interestingly, TREAT assays looking at mRNA stability showed significant differences. This suggests that G3BP1 tethering reduces 5'-3' mRNA degradation. While some recent publications have provided evidence for a decay promoting role of G3BP1 [57, 58], others including this study have sug-

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gested the opposite [59, 60, 61]. It must be noted that this mRNA stabilizing effect, while significant, still does not entirely prevent mRNAs from degradation. The molecular mechanisms behind this finding remain unknown but could arise from a microscopic SG-like environment forming around single mRNAs. Even though G3BP1 tethering does not form visible SGs in unstressed cells, the recruitment of SG proteins by G3BP1 could still be sufficient to reduce mRNA degradation. Similarly, G3BP1 could also actively be preventing the recruitment of decay enzymes to their tethered target mRNAs. To follow up on these proposals, it would be necessary to investigate whether the local protein composition changes proximally to tethered mRNAs. A similar experimental setup could also be used to test the mRNA stabilizing effect of other SG-proteins like TIA-1, single G3BP1 domains, or domains enriched in the SG transcriptome such as DEAD-box helicases. Lastly, the direct involvement of G3BP1 on mRNA stabilization during stress conditions has not been investigated and must be addressed to understand the overall role of G3BP1 in mRNA metabolism.

In an effort to increase the expression levels of the currently used SunTag reporter cassette, a novel cassette relying on structural support by smGFP was created. The preliminary results shown here suggest that single-chain antibody occupancies are comparable with the previous version while greatly increasing transcript numbers. This seems to mainly be achieved through more translating transcripts per cell. However, it is still unclear what the underlying cause of the poor induction of the SunTag reporter is. The lower signal intensity has shown to make a track based analysis slightly harder. To counteract this, one could, however, use two or more cassettes, or add more GCN4 repeats at other locations within the structural scaffold. Looking forward, it would be interesting to see if mRNA stability is also affected and how the system can be evolved. Recent progress in single-chain antibodies leads the way to a highly interchangeable system with different tags and fluorescent colors.

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A

Supplementary Methods

NLS-stdMCP-stdHalo-G3BP1 cloning primers:

- gattaggatctatcgattactgccgtggcgcaagcccc
- aagtctgtttcagggcccgatggatggagaagcctagtccc
- ctaggcttcacatcaccatcggccctgaaacagaactt
- ggggcttgcgccacggcagtaatcgatagatctaatacacc

smGCN4-Renilla-MS2v5 cloning primers:

- tacacgcggcccaatcatcggtccgggtggatctggagggtggaggttctggaggagaaga
- cctcttccgagccagttccctggcccttattccggttt

Renilla luciferase FISH probes:

- tcgtacacccgttggaaagccat
- agtgatcatgcgttgtcggt
- gttcattgttgcagcgag
- agttgatgaaggaggccagc
- gcgtgttccgttccgttccgtt
- cagcgttaccatgcagaaaa
- tcgatgtgaggcacgacgt
- gatgtatgttccgttccgtt
- ttaccattccgttccgttccgtt
- gatccaggaggcgatatgag
- caagcggtgagggtacttgc
- ttggaaagggttccgttccgtt

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- gtggcccacaaagatgattt
- gagtagtgaaaggccagaca
- ttgatcttgtctggtgctc
- cactctcagcatggacgatg
- caggactcgatcacgtccac
- atatccctcgatgtcagg
- ctcttcgctctgtatcaggg
- attctcaaggcaccatttct
- gcatggtctcgacgaagaag
- tttccgcatgatctgcttg
- cttaaatggctccaggtagg
- gagagggtaggccgtctaacc
- ttaacgagagggatctcgcg
- gacaatctggacgacgtcgg
- gaaggtaggcgtttagttg
- gaacatcttaggcagatcgt
- tggaaaagaaccagggtcg
- cttagctccctcgacaatag
- ttcacgaactcggtttagg
- cttaaccatttcatctggag
- gctccacgaagctttagt
- tactgctcgttcttcagcac

MS2v4 FISH probes:

- tgccgttttaggttaggatc
- cgcttgaagattggacagtgc
- gactgtaatgacagtggagc
- ctgatgctgctggagttga
- atgtcctgatgttagtcggag
- atcgtcgagcgttgaatgat
- atagtgtctgaggcatgctg
- gtgatgtcgagccgttga

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- cgataacgtagaggcagtag
- ctgatgctcgtcgcagaaga
- atctggtatgtccgatgttq

MS2v5 FISH probes:

- tgattgtgaagtgtcgggtg
- tccacccttgttatgtac
- ctgtgtaatgtgtctggagg
- gtgcttctgtttgattggat
- aaatttggtagcagaagccc
- cttgtttagtttttgag
- caggaaatccctgatggta
- ccctcgttgacgtatattg
- gatattcgggaggcgtgatc
- acgcactgaattcgaaagcc
- attcgactctgattggctgc
- ctcttcgcaaagtgcactt
- taagaatggcgcaaggctg
- gtagggagagtgtggttt
- caggaacgctgatgctgttc
- ttcttgagttgggtactg
- tcatgctgcatggggacata
- ttggggatgtattctgggg
- ttggtgctcgatgtgattt
- aagaaaacaacaactccgagcc
- atggagggttgtccagttg
- ttgtcttgtggtagagat
- ctgatgctgcttcgagaaga
- gtatgctcgagtgtttcgaa
- gatcgccacccaagaata
- aattcgtagagatgggtg
- tcgtattggacgtggaacga

Appendix A. Supplementary Methods

B

Supplementary Figures

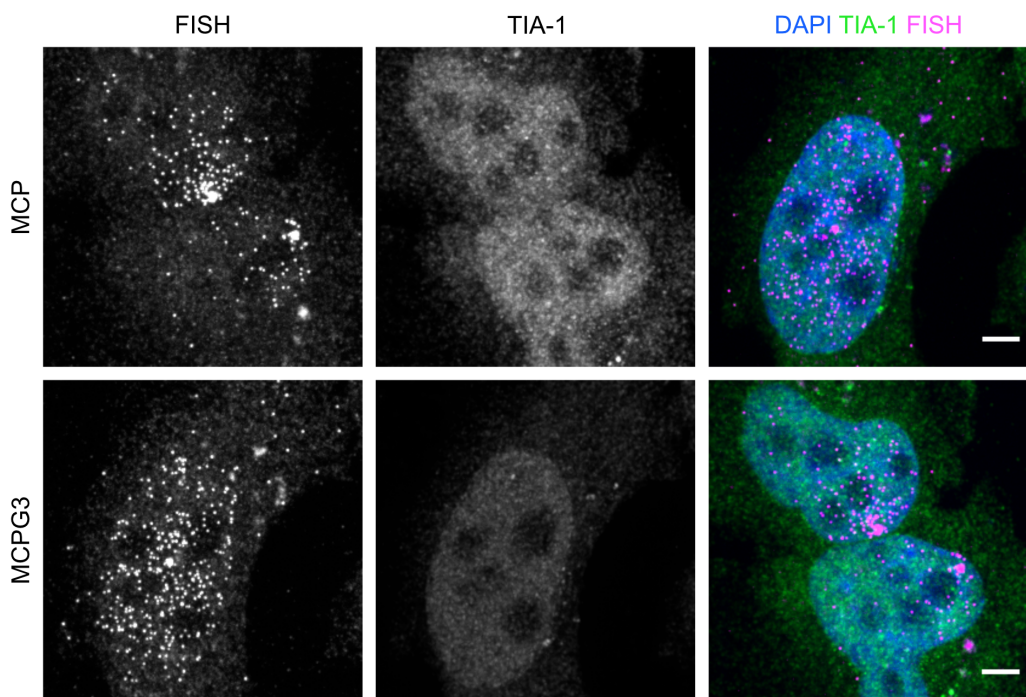


Figure B.1: mRNA localization in non-stressed cells. Representative fluorescence images of the SG marker TIA-1 and the tethered RNAs in unstressed cells. Scale bars, 10 μ m.

Appendix B. Supplementary Figures

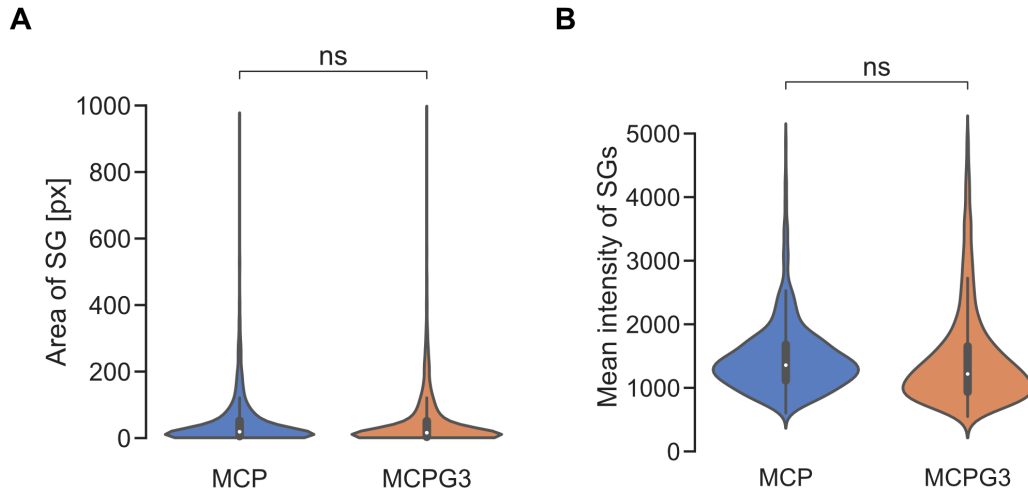


Figure B.2: Comparison of SG properties. (A) SG size comparison between both constructs using a total of 66 cells each. (B) Mean intensity values of single SGs in the TIA-1 SG marker channel.

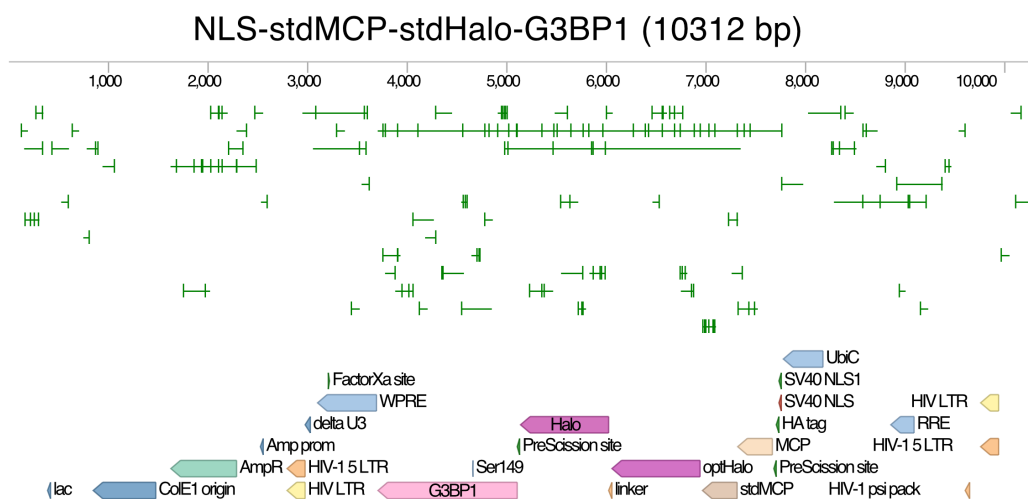


Figure B.3: Design of MCPG3.

Appendix B. Supplementary Figures

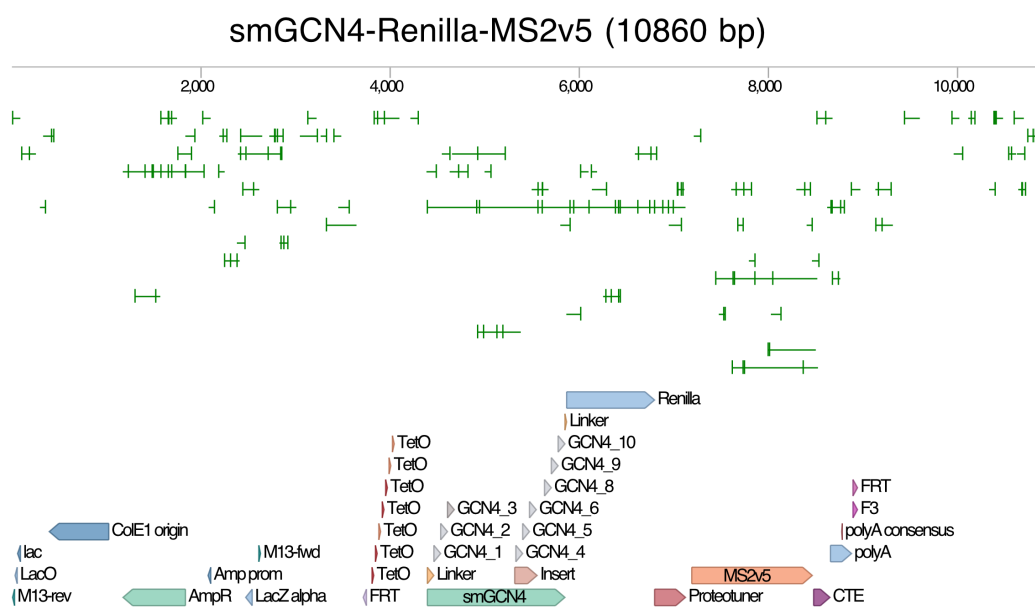


Figure B.4: Design of smGFP SunTag.

Colophon

This document was created using \LaTeX and Bib \TeX typesetting originally developed by Leslie Lamport, based on \TeX created by Donald Knuth.

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The text is set in Helvetica.