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To cite this version:

Xavier Pichon, Konstadinos Moissoglu, Emeline Coleno, Tianhong Wang, Arthur Imbert, et al.. The kinesin KIF1C transports APC-dependent mRNAs to cell protrusions. RNA, Cold Spring Harbor Laboratory Press, 2021, pp.rna.078576.120. 10.1261/rna.078576.120. hal-03371977

HAL Id: hal-03371977
https://hal.archives-ouvertes.fr/hal-03371977
Submitted on 9 Oct 2021

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The kinesin KIF1C transports APC-dependent mRNAs to cell protrusions

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Running title: mRNA localization and transport

Keywords: RNA localization, local translation, RNA transport, cytoplasmic protrusions

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Abstract

RNA localization and local translation are important for numerous cellular functions. In mammals, a class of mRNAs localize to cytoplasmic protrusions in an APC-dependent manner, with roles during cell migration. Here, we investigated this localization mechanism. We found that the KIF1C motor interacts with APC-dependent mRNAs and is required for their localization. Live cell imaging revealed rapid, active transport of single mRNAs over long distances that requires both microtubules and KIF1C. Two color imaging directly revealed single mRNAs transported by single KIF1C motors, with the 3’UTR being sufficient to trigger KIF1C-dependent RNA transport and localization. Moreover, KIF1C remained associated with peripheral, multimeric RNA clusters and was required for their formation. These results reveal a widespread RNA transport pathway in mammalian cells, in which the KIF1C motor has a dual role in transporting RNAs and clustering them within cytoplasmic protrusions. Interestingly, KIF1C also transports its own mRNA suggesting a possible feedback loop acting at the level of mRNA transport.
Introduction

Localization of mRNA to specific subcellular compartments is an important mechanism for the spatio-temporal regulation of gene expression in diverse cell types and organisms (Chin and Lécuyer 2017; Eliscovich and Singer 2017). Subcellular mRNA localization allows localized protein synthesis and this is important for many biological functions such as cell fate determination (Berleth et al. 1988), cell polarization (Condeelis and Singer 2005), cell division (Chouaib et al. 2020; Ryder et al. 2020), cell migration (Katz et al. 2012; Wang et al. 2017; Moissoglu et al. 2020), embryonic patterning (Forrest and Gavis 2003) and synaptic plasticity (Martin and Zukin 2006; Lin and Holt 2007). One of the best characterized examples is the yeast *Ash1* mRNA that localizes specifically in the bud of the daughter cells and encodes a transcriptional repressor protein involved in suppressing mating-type switching (Paquin and Chartrand 2008). Studies of this and other models revealed that the subcellular localization of mRNA relies on three main mechanisms, acting separately or in combination: random diffusion combined with local entrapment, general transcript degradation coupled to localized protection and directed transport along the cytoskeleton (Cody et al. 2013; Medioni et al. 2012; Bovaird et al. 2018).

Active, motor driven transport of mRNAs along the cytoskeleton is thus far the most common localization mechanism. It generally involves cis-acting elements, also called zipcodes, contained in the 3’UTR sequence of the transcript. This is exemplified by the case of the β-actin mRNA in vertebrates, which accumulates at the leading edge of migrating cells and was among the first localized mRNAs discovered (Singer, 1993). This mRNA contains a zipcode sequence recognized by the RNA Binding Protein ZBP1, allowing the transport of β-actin mRNAs in a motor-driven manner along the cytoskeleton (Kislauskis et al., 1993; Oleynikov and Singer, 2003; Liao et al., 2015; Condeelis and Singer, 2005). Interestingly, transport of β-actin mRNA by ZBP1 involves both microtubules (MTs) and actin filaments.
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(Fusco et al. 2003; Oleynikov and Singer 2003), as well as several motors that display some cell type and compartment specificity. Indeed, MYO5A and KIF5A interact with ZBP1 to transport β-actin mRNAs in dendrites and axons (Ma et al. 2011; Nalavadi et al. 2012), while Myosin IIB (MYH10) and KIF11, which directly binds ZBP1, regulate the transport of β-actin mRNAs in fibroblasts and during cell migration (Song et al. 2015; Latham et al. 2001).

In vertebrate systems, the motors involved in RNA transport have been investigated mostly in neuronal cells. Kinesin-1 (KIF5) was shown to associate with neuronal RNP granules and to be involved in their trafficking (Kanai et al. 2004). Kinesin-1 was also implicated in transport of myelin basic protein (MBP) mRNA in oligodendrocytes (Carson et al. 1997) as well as in shank1 mRNA transport in rat neurons (Falley et al. 2009). Kinesin-2 (KIF3A/B/KAP3) can transport RNAs in vitro (Baumann et al. 2020), but its in vivo relevance is still unclear. The involvement of additional motors, and the means through which they connect to potential RNA cargoes are still largely unexplored, especially in the case of non-neuronal cell types (Gagnon and Mowry 2011; Xing and Bassell 2013).

Localized RNAs are prevalent in non-neuronal, mesenchymal cells. Apart from β-actin, numerous other RNAs are localized at protrusions of mesenchymal cells and their local translation is important for cell migration (Mili et al. 2008; Mardakheh et al. 2015; Moissoglu et al. 2020; Costa et al. 2020). Localization of these RNAs is carried out through at least two distinct pathways. Specifically, a subset of about a hundred RNAs, which include transcripts encoding signaling and cytoskeleton regulators (such as the Rab GTPase RAB13, the RhoA exchange factor NET1, the collagen receptor DDR2, the motor related proteins TRAK2, DYNLL2, and others), require the APC tumor suppressor protein for localization and have been referred to as APC-dependent (Wang et al. 2017). Other protrusion-enriched RNAs, exemplified by RNAs encoding ribosomal proteins, do not require APC and exhibit distinct regulation (Wang et al. 2017).
Similar to what has been described for other localized RNAs, sequences within the 3’UTR of APC-dependent RNAs are necessary and sufficient for targeting to the cell periphery (Mili et al. 2008). Specifically, interfering with or deleting particular GA-rich regions is sufficient to disrupt peripheral localization and perturb cell movement in various systems (Moissoglu et al. 2020; Costa et al. 2020; Chrisafis et al. 2020). Furthermore, localization to the periphery requires the microtubule cytoskeleton and in particular a subset of stable, detyrosinated microtubules (Wang et al. 2017; Moissoglu et al. 2019). Indeed, at least some APC-dependent RNAs exhibit a co-localization with the plus ends of detyrosinated microtubules (Mili et al. 2008). The peripheral complexes also contain APC, a protein that has the ability to directly bind microtubules via its C-terminus (Barth et al. 2008; Bahmanyar et al. 2009; Jimbo et al. 2002; Munemitsu et al. 1994; Zumbrunn et al. 2001), hence suggesting that APC might mediate the interaction of localized mRNAs with microtubules (Mili et al. 2008; Preitner et al. 2014).

An additional feature integrated with the localization of APC-dependent RNAs is their existence in distinct physical states. In particular, RNAs in internal or peripheral, actively extending cytoplasmic regions exist as single molecules that are undergoing translation. However, at some peripheral areas, single RNAs coalesce in multimeric heterogeneous clusters that are composed of multiple distinct RNA species. Interestingly, these clusters preferentially form at retracting protrusions and contain translationally silent mRNAs (Moissoglu et al. 2019). These data indicate the existence of a dynamic regulatory mechanism during cell migration, which coordinates local mRNA translation with protrusion formation and retraction. However, the exact mechanisms and molecular players involved in transport to the periphery and cluster formation for this group of RNAs are still unclear.

In this study, we focused on the kinesin KIF1C, which we recently showed to accumulate and colocalize with its own mRNA in cytoplasmic protrusions (Chouaib et al. 2018).
We show here that KIF1C associates with additional protrusion-localized RNAs belonging to the APC-dependent group. We describe a specific mRNA transport mechanism by which the KIF1C kinesin motor binds APC-dependent mRNAs, including its own, actively transports them to cell protrusions in a 3’UTR dependent manner and additionally participates in promoting and/or maintaining their peripheral clusters.
Results

Identification of a specific mRNA subset associating with KIF1C motor in human cells

High-throughput mRNA-protein cross-linking approaches previously showed that KIF1C directly binds mRNAs (Baltz et al. 2012; Castello et al. 2012), and we recently showed that KIF1C mRNAs and proteins colocalize together in protrusions of HeLa cells (Chouaib et al. 2020; Fig. 1A), suggesting that the KIF1C kinesin might be somehow involved in the metabolism of protrusion mRNAs. To determine the identity of the mRNAs bound by the KIF1C motor, we used a HeLa cell line stably expressing a KIF1C-GFP fusion from a bacterial artificial chromosome containing all the regulatory sequence of the human KIF1C gene, including its 5’ and 3’UTRs (Poser et al. 2008; Chouaib et al. 2020). We immunoprecipitated (IP) KIF1C-GFP with anti-GFP antibodies or uncoated beads as controls, and identified the co-precipitated RNAs using microarrays (Fig. 1B and Table S1). We found that many mRNAs were enriched in the KIF1C-GFP IP as compared to the control IP. To explore in more detail the localization of the mRNAs associated with KIF1C, we performed a small smFISH localization screen in HeLa cells. We tested the 26 most enriched mRNAs in the KIF1C IP (Table S1 and S2) and found four that were peripherally enriched. These included the RAB13 mRNA (5.7 fold enrichment, Table S1), along with the KIF1C mRNA itself (2.6 fold enrichment) and the NET1 and TRAK2 mRNAs (5.7 and 4.9 fold enrichment respectively). We had previously reported that these transcripts localize to protrusions of mouse cells in an APC-dependent manner (Wang et al. 2017), and we thus focused on them. Visual examination of the images revealed that KIF1C, NET1, TRAK2 and RAB13 mRNAs clearly localized also to protrusions of HeLa cells (Fig. S1A). To confirm the link between APC-dependent mRNA localization and binding to KIF1C protein, we performed a correlation analysis of the two metrics (Fig. S1B and Table S3). This indicated that APC-dependent mRNAs indeed preferentially associate with KIF1C protein, while mRNAs coding for ribosomal proteins,
which often localize to protrusion independently of APC (Wang et al. 2017), do not. The IP/microarray data thus show a physical link between KIF1C and mRNAs that localize to protrusions in an APC-dependent manner.

Next, we tested whether these mRNAs colocalize with the KIF1C protein in vivo. To this end, we performed smFISH experiments in a HeLa cell line stably expressing a KIF1C-GFP mRNA from a cDNA and found that indeed, KIF1C, NET1, TRAK2 and RAB13 mRNAs co-localized with the KIF1C-GFP protein in cytoplasmic protrusions (Fig. 1C). In order to show that this colocalization reflected a molecular interaction, we performed single molecule imaging using the SunTag system (Tanenbaum et al. 2014). To this end, we generated a stable HeLa cell line expressing KIF1C-fused to 24 repeats of the GCN4 epitope (KIF1C-SunTag\textsubscript{x24}), together with the single-chain variable fragment fused to sfGFP (scFv-sfGFP). This system enables the detection of single molecules of the KIF1C protein (Tanenbaum et al., 2014; Fig. 1D). We thus combined detection of single KIF1C-SunTag\textsubscript{x24} proteins with single mRNA detection by smFISH, using probes against RAB13 and NET1 mRNA, or CRM1 and RBP1 mRNAs as controls (Fig. 1D and 1E; Fig. S1C-E). KIF1C-SunTag\textsubscript{x24} proteins were found to colocalize with RAB13 and NET1 mRNAs at protrusions as expected, while the control mRNAs did not. In addition, we also observed colocalization of KIF1C-SunTag\textsubscript{x24} with both RAB13 and NET1 mRNAs at the single molecule level at more internal locations in the cytoplasm (Fig. 1E, panels 1, 3-4; Fig. S1D). This confirmed the interaction of single molecules of KIF1C protein with single molecules of RAB13 mRNAs. Taken together, these data raise the possibility that the kinesin KIF1C might be part of a mechanism that localizes APC-dependent mRNAs to cytoplasmic protrusions.

**KIF1C associates with APC and is required for the localization of APC-dependent mRNAs to cytoplasmic protrusions in human and mouse cells**
To further support the connection between KIF1C and APC-dependent RNAs we tested whether KIF1C and APC interact in cells. Indeed, immunoprecipitation of GFP-APC revealed a specific association with KIF1C-mCherry (Fig. 2A). We additionally imaged the two fluorescent proteins in live cells and detected a colocalization between GFP-APC and KIF1C-mCherry in peripheral clusters (Fig. 2B). Therefore, KIF1C exhibits a specific association with both APC protein and peripherally localized APC-dependent RNAs.

To test whether the localization of APC-dependent mRNAs in protrusions depended on the KIF1C protein, we depleted KIF1C by multiple siRNAs in HeLa or MDA-MB-231 cells, and performed smFISH experiments using probes against RAB13 or NET1 mRNAs. Intracellular distributions of mRNAs were quantitatively assessed by calculating a Peripheral Distribution Index (PDI), a metric that distinguishes diffusely distributed from peripherally localized RNAs, by measuring the distance of the RNA signal relative to the centroid of the nucleus (Wang et al. 2017; Stueland et al. 2019). For each cell, the RNA distribution is normalized to a hypothetical uniform distribution such that a PDI value of 1 indicates a uniform, diffuse signal, while values smaller or greater than 1 indicate a perinuclear or peripheral localization, respectively. Remarkably, both RAB13 and NET1 mRNAs became less localized when KIF1C expression was reduced with siRNAs (Fig. S2A-D), demonstrating that the KIF1C kinesin was required for mRNA localization in human cells.

Next, we moved to a mouse system, NIH/3T3 cells, where the localization of mRNAs in protrusions has been extensively studied (Chicurel et al., 1998; Mili et al., 2008; Wang et al., 2017; Moissoglu et al., 2019). To test whether the KIF1C protein has a general role in localizing mRNAs at cell protrusions, we assessed the localization of a series of APC-dependent and APC-independent mRNAs by smFISH, following depletion of KIF1C expression with two different siRNAs. As shown in Fig. 2C-E and S2E-F, several APC-dependent RNAs, including Net1, Rab13, Ddr2, Dynll2 and Cyb5r3, exhibited a protrusion localization pattern that was lost
following KIF1C depletion. Indeed, KIF1C loss led to RNA distributions that were mostly diffuse (PDI values centering around 1; Fig. 2B and Fig. S2F), indicating that the KIF1C motor has an important contribution towards directing peripheral mRNA localization. Interestingly, the localization of two APC-independent mRNAs, \textit{Rps20} and \textit{Rpl27a}, was not affected (Fig. S2G). To ascertain that this effect was not due to altered mRNA expression, we measured their levels following KIF1C depletion (Fig. S2H). This analysis showed no changes in the overall abundance of APC-dependent mRNAs, except for the depleted KIF1C mRNA (Fig. S2H). Therefore, we conclude that KIF1C is required for the localization of APC-dependent mRNAs to cell protrusions, in various human and mouse cells.

**KIF1C is required for active transport of APC-dependent mRNAs on microtubules**

To monitor trafficking of APC-dependent mRNAs, we expressed in NIH/3T3 fibroblasts a reporter carrying the β-globin coding sequence followed by 24 binding sites for the bacteriophage MS2 coat protein (MCP; Fig. 3A schematic). Binding of co-expressed MCP-GFP to these sites allows visualization and tracking of single molecules of the reporter mRNA in living cells (Fusco et al. 2003). To recapitulate the localization of APC-dependent RNAs, the reporter additionally included a control 3’UTR or the 3’UTR of \textit{Net1} or \textit{Rab13} (hereafter referred to as β24bs/Ctrl, β24bs/Net1 and β24bs/Rab13, respectively). As shown previously, these 3’UTR sequences are sufficient to direct peripheral distribution of this reporter transcript in NIH/3T3 cells (Moissoglu et al. 2019). We initially examined trafficking of the reporter during early stages of cell spreading, which mimic conditions in actively protruding cell regions. Indeed, live fluorescence imaging of the reporter containing the Net1 3’UTR revealed a distinct peripheral pattern after plating cells on fibronectin for 30 minutes (Fig. 3A and Movie S1). Because kinesin-dependent mRNA trafficking is expected to occur on the microtubule cytoskeleton, it was important to identify microtubule-dependent events and discriminate them.
from other modes of motion. For this, reporter particles were tracked in cells before and after
15 min of nocodazole treatment. To identify long and linear movements, as those expected to
occur on microtubules, we used two different metrics to quantitatively describe individual
tracks: ‘Linearity of forward progression’ and ‘Track displacement’. To determine the range of
these parameters that define directed microtubule-based tracks, we compared movements
before and after microtubule depolymerization. From this, we separated ‘long/directed’ tracks
(Fig. 3B). These tracks exhibit higher displacement and linearity (net displacement > 4 microns,
linearity > 0.7) and they are absent in cells treated with nocodazole. They correspond to ~3-6%
of the total tracks (Fig. 3B and Movies S1, S2). Consistent with their representing persistent
directed motions, this subset of tracks exhibit high Mean Square Displacements (MSD) over
their lifetime (generally more than 15 μm²) and display positive velocity autocorrelation (Fig.
3C, left panels). By contrast, the remaining tracks, which we classify as ‘short/diffuse’, exhibit
characteristics similar to tracks of nocodazole-treated cells. Specifically, they exhibit low
MSDs and zero velocity autocorrelation, as is characteristic of diffusive Brownian motions
(Fig. 3C, middle and right panels). Both ‘long/directed’ and ‘short/diffuse’ tracks have similar
range of lifetimes, while ‘long/directed’ tracks have higher mean speed (Fig. S3).

Tracking of a reporter carrying the Rab13 3’UTR also exhibited long/directed tracks
with dependence on microtubules (Fig. 3D, E). In contrast, a control reporter lacking a
localizing 3’UTR did not produce tracks with these characteristics (Fig. 3D, E). Importantly,
this subset of tracks was not affected by disruption of the actin cytoskeleton with cytochalasin
D or following treatment with a control vehicle, DMSO (Fig. 3F, Fig. S4 and Movies S3, S4).
Thus, the reporters carrying the 3’UTR of Net1 or Rab13 mimic the localization pattern of APC-
dependent mRNAs and allow the identification of long and linear microtubule- and 3’UTR-
dependent transport events.
To directly test the role of KIF1C in these trafficking events, we visualized fluorescent particles of the reporter carrying the *Net1* 3’UTR and measured the frequency of long, directed microtubule-dependent displacements in actively spreading cells following KIF1C depletion. As previously observed with endogenous transcripts (Fig. 2), reporter mRNAs became less localized when KIF1C expression was reduced with siRNAs (Fig. 4A and Movies S5-S8). Importantly, track analysis showed that KIF1C loss significantly reduced the number of the microtubule-dependent displacements (Fig. 4B and 4C). To assess specificity, we depleted two additional kinesins that have been linked to APC and RNA transport, KIF5B and KIF3A (Dunn et al. 2008; Kanai et al. 2004; Cai et al. 2009; Yasuda et al. 2017; Baumann et al. 2020). Fig. 4A-C shows that depleting these kinesins did not change the overall peripheral accumulation of the reporter and did not result in a reduction of the long, directed transport events. Thus, KIF1C exhibits a specific function in transporting APC-dependent mRNAs via microtubules in actively protruding cell regions.

**Peripheral clustering of APC-dependent mRNAs depends on KIF1C**

Peripheral APC-dependent mRNAs can form large heterogeneous clusters that are translationally silent (Moissoglu et al. 2019). These clusters often associate with retracting protrusions in migrating cells, suggesting that they are part of a spatio-temporal control of protein synthesis (Moissoglu et al. 2019). Formation of these clusters is recapitulated by the reporter constructs carrying the *Net1* or *Rab13* 3’UTR, but not by a control reporter (Fig. S5). These clusters are visible at later time points after plating (ca. 3 hours), when most protrusions are not actively extending, consistent with the appearance of endogenous RNA clusters in non-extending or retracting protrusions (Fig. 5A; Moissoglu et al., 2019). These clusters can be identified as bright particles with intensities higher than those characteristic of single molecules (Fig. 5A). To test whether KIF1C is implicated in the formation of these clusters, we scored the
frequency of bright particles in KIF1C-depleted cells during late stages of spreading (3 hours; Fig. 5B, C; particle brightness > 4950). As shown in Fig. 5, while clusters formed by the Net1 3’UTR-reporter were readily observed in protrusions of control siRNA-treated cells, their frequency was substantially reduced, and mostly single molecules were present, when KIF1C was depleted (Fig. 5A-C). Moreover, cluster formation was only marginally affected by the depletion of KIF5B or KIF3A. Thus, KIF1C specifically controls the clustering of APC-dependent mRNAs. We note that clusters are not detected even in protrusions containing a substantial amount of single RNA molecules (see enlarged KIF1C insets in Fig. 5A), suggesting that cluster loss is not a secondary consequence of reduced number of mRNA molecules arriving at protrusions upon KIF1C depletion. We rather think that these results indicate an additional role of KIF1C in forming higher order RNP complexes at protrusions.

**Single molecule two color imaging provides direct evidence that the KIF1C motor transports protrusion mRNAs**

To provide direct evidence that the protrusion mRNAs are transported by the KIF1C motor, we performed two color single molecule imaging of mRNAs and motors, in order to visualize cotransport of the two types of molecules. To this end, we used the NIH/3T3 cells expressing the Net1 3’UTR-containing reporter and modified them to also stably express a KIF1C protein fused to the SunTag (KIF1C-ST\textsubscript{24}), together with a single-chain variable fragment antibody fused to mScarletI (scFv-mScarletI).

Imaging of fixed cells showed that the KIF1C-SunTag\textsubscript{24} motor and reporter mRNAs accumulated in protrusions as expected (Fig. 6A-B, Fig. S4A-B). Moreover, we could also occasionally detect colocalization events where a single molecule of KIF1C-SunTag\textsubscript{24} would colocalize with a single reporter mRNA at internal cellular areas. To confirm that this colocalization was relevant to mRNA transport, we performed two-color live-cell imaging
using movies recorded at a high frame rate (7.36 fps for 52 seconds). This allowed the detection of co-transport events, in which a single molecule of KIF1C-STx24 moved with a reporter mRNA molecule in a rectilinear manner at high speed (Fig. 6C, D; Movie S9; Fig. S4C, D; Movie S10). Kymographs confirmed that both molecules moved together in an anterograde direction toward protrusion (Fig. 6E and Fig. S4E), traveling an average distance of 22 microns at speeds of 2.6 μm/second (Fig. 6F-G). Taken together, these data demonstrate that the KIF1C kinesin actively transports this Net1 reporter mRNA to cell protrusions along microtubule cables.
Discussion

RNA transport along the cytoskeleton is a well-established mechanism allowing subcellular mRNA localization and local translation. In mammals, a large class of mRNAs localize to cytoplasmic protrusions of many cell types, where they are anchored at the plus-end of detyrosinated microtubules by APC. Here, we show that these mRNAs associate with the microtubule motor KIF1C and interestingly, they include the KIF1C mRNA itself. We show that APC-dependent mRNAs and KIF1C protein co-localize in protrusions and can also be seen co-transported together along directed tracks. Moreover, the peripheral localization of these mRNAs as well as their microtubule-dependent motion depend on KIF1C, demonstrating that it is an essential motor that transports APC-dependent mRNAs to protrusions. Our data provide a striking in vivo visualization of the co-transport of individual RNA molecules with a specific molecular motor, involved in a widespread RNA transport pathway.

The kinesin KIF1C controls a widespread mammalian mRNA transport system

RNA localization controls spatial and temporal aspects of gene expression in a variety of species and cell types. Although its significance is better understood in specialized cells such as neurons, recent reports highlight its widespread prevalence, including in cells with a mesenchymal phenotype (Mardakheh et al. 2015; Wang et al. 2017; Fazal et al. 2019; Costa et al. 2020; Chouaib et al. 2020). Our current understanding of the transport mechanisms includes the requirement of cis-acting sequence elements and trans-acting factors, which work with the actin or microtubule networks and motor proteins to bring mRNAs to their destination (Cody et al. 2013; Medioni et al. 2012; Bovaird et al. 2018). Nevertheless, our knowledge of common RNA transport mechanisms that operate in most, if not all cells, is limited. In the case of protrusion mRNAs, which localize in all cell types examined so far, their localization was shown to require APC and detyrosinated microtubules (Mili et al. 2008; Wang et al. 2017).
is not yet known whether APC is transported together with the KIF1C-RNA complexes or whether it is independently transported and subsequently associated with RNAs at the periphery. *In vitro* studies suggest that motile complexes can be formed between mRNA, APC and KIF3A (Baumann et al. 2020). Our data, however, clearly show the involvement of KIF1C in transporting protrusion mRNAs *in vivo*. Moreover, APC was shown to accumulate at the leading edge of migrating cells using kinesin-1 and kinesin-2 (Mimori-Kiyosue et al. 2000; Nakamura et al. 2001; Ruane et al. 2016). The use of distinct motors suggests an independent transport for APC and protrusion mRNAs. It is also possible that this diversity of motors reflects differences between cell types, as has been described for the β-actin RNA that uses different motors in neurons and fibroblasts (see Introduction; Ma et al., 2011; Nalavadi et al., 2012; Song et al., 2015; Latham et al., 2001). One potential reason for using different motors could pertain to the dynamics and transport speeds required in each case. In this context, the fact that KIF1C appears to be the fastest human cargo transporter (Lipka et al. 2016) might provide an advantage that could underlie its preferential use in the highly dynamic mesenchymal cells. Furthermore, kinesins other than KIF1C, such as KIF5B, contribute to localization of APC-dependent RNAs (Yasuda et al. 2017). We speculate that they may function in specialized cells or affect different aspects of localization that are distinct from transport *per se*. In line with this idea, protrusion mRNAs display a complex translational regulation concomitant to the protrusion dynamics and a local reorganization of RNA clusters (Moissoglu et al. 2019), likely requiring a number of yet uncharacterized actors.

Another important question deals with the adaptors that link KIF1C to mRNAs. On one hand, proteomic data showed that KIF1C protein interacts with the exon junction complex (EJC; Hein et al., 2015). The EJC is assembled on spliced RNAs and serves as an interaction platform for proteins that direct mRNA export, localization, translation and nonsense-mediated mRNA decay (NMD), suggesting a role for the EJC in the transport of protrusion mRNAs. On
the other hand, regions rich in G and A nucleotides are present in the 3’UTR of APC-dependent RNAs and are part of the localization element (Moissoglu et al. 2020; Costa et al. 2019). Furthermore, the 3’UTR is sufficient to direct protrusion localization of intronless exogenous constructs (Moissoglu et al. 2019, 2020), indicating that a potential KIF1C recruitment through exon-exon junctions might not be necessary for protrusion localization. Specific RNA-binding proteins could serve as adaptors that mediate KIF1C recruitment similar to the model suggested for other RNA transport complexes. However, given that high-throughput mRNA-protein cross-linking approaches previously showed that KIF1C directly binds mRNAs (Baltz et al. 2012; Castello et al. 2012), an alternative interesting possibility would be that in this case the motor directly selects and binds to its RNA cargo.

**KIF1C triggers mRNA clustering in cytoplasmic protrusions**

Our live imaging experiments show that reporter mRNAs are transported predominantly as single molecules during the early stages of cell spreading, reminiscing the single molecule appearance of RAB13 mRNA in actively protruding regions in migrating cells (Moissoglu et al. 2019). Remarkably, APC-dependent mRNAs coalesce into higher order clusters at peripheral regions during later time points of cell spreading and this phenotype depends specifically on KIF1C. We think that clustering is not merely a consequence of peripheral mRNA accumulation, because we have not observed it in the absence of KIF1C even at protrusions still containing substantial numbers of single mRNA molecules. We rather favor the explanation that clustering is a separate function of KIF1C that is temporally and spatially regulated. Given that these clusters contain stably anchored mRNAs (Mili et al. 2008), we envision that KIF1C switches from a microtubule motor to an mRNA anchoring module promoting clustering. A similar switch has been observed in Drosophila oocytes, whereby Dynein converts from a motor of *gurken* mRNA to a static anchor at its final destination.
Such a switch on KIF1C may take place on pre-existing motor molecules as they reach the periphery or may be a function of newly-synthesized KIF1C translated from its peripherally localized mRNA. It is still unclear how such a switch would occur and/or whether it might additionally involve a change in the RNA-binding mode of KIF1C (direct or indirect through other RNA-binding proteins). Clusters of APC-dependent mRNAs have been previously reported to be heterogeneous and to contain translationally silent mRNAs (Moissoglu et al. 2019). Thus, overall our results point to a spatially and temporally controlled mRNA clustering role of KIF1C that is separate from its motor function and that might be coordinated with translational regulation.

**RNA transported by KIF1C mediates diverse functions at cell protrusions**

Peripheral localization of APC-dependent RNAs promotes cell migration (Wang et al. 2017). Specifically, approaches targeting the localization elements of these mRNAs, as a group (Wang et al. 2017) or individually (Moissoglu et al. 2020), resulted in inhibition of cell migration. These effects are likely due to a requirement for locally translating these mRNAs for full activation of the encoded proteins (Moissoglu et al. 2020). Interestingly, KIF1C has been shown to control adhesion dynamics and cell migration (Theisen et al. 2012). It was proposed to act via the trafficking of α5β1 integrins. While we cannot completely rule out that KIF1C indirectly affects peripheral RNA localization through altering adhesion dynamics, we consider this highly unlikely given the physical interaction and co-transport of KIF1C and RNAs that we report here. Instead, our results indicate that the transport of APC-dependent mRNAs to the periphery by KIF1C likely itself contributes to the mechanism by which this kinesin controls cell migration. Along this line, the GO terms associated with the top 200 KIF1C-associated mRNAs presented in this study (i.e. organelle organization; plasma membrane bounded cell
projection organization; microtubule-based transport; cilium organization; Table S5) indicate how KIF1C-mediated mRNA transport could impact processes related to cell motility.

**KIF1C protein localizes its own mRNAs to cell protrusions: a transport feedback loop?**

The KIF1C transcript localizes to cytoplasmic protrusions in mammalian cells. Moreover, it co-localizes with KIF1C protein in protrusions (Chouaib et al. 2020), and we show here that the KIF1C protein physically associates with its own mRNA. This local accumulation of KIF1C could be involved in an RNA clustering and anchoring mechanism as discussed above, but it could also serve to transport additional mRNAs by alternating back-and-forth movements on the cytoskeleton. Indeed, locally translated KIF1C protein would allow the motor to explore the local cytoplasm and transport back additional mRNAs to protrusions using the same MT tracks. Such a bidirectional motility has been reported for KIF1C and it is mediated by the scaffold protein Hook3. This protein forms a complex between dynein and KIF1C (Kendrick et al. 2019), and regulates their activities to allow the motor to perform multiple transport cycles while avoiding a tug-of-war between opposite motors (Siddiqui et al. 2019). The fact that KIF1C also brings its own mRNA to protrusions suggests the possible existence of a positive feedback loop in which locally translated KIF1C provides additional motor molecules to sustain the persistent and directional transport of its RNA cargoes, to locally maintain protrusive extensions during cell movement.
Acknowledgements

We thank the staff of MRI imaging facility for their technical support. This project was supported by France BioImaging (ANR-10-INBS-04), the Agence Nationale de la Recherche (ANR-11-BSV8-018-02, ANR-14-CE10-0018-01 and ANR-19-CE12-0007-03), and the Ligue Nationale Contre le Cancer and the Fondation pour la Recherche Médicale. This work was supported by the Labex EpiGenMed, from the framework "Investissements d’avenir". This work was supported in part by the Intramural Research Program of the National Cancer Institute, NIH.

Author contributions statement

EB conceived the study with SM and KZ. Experiments were performed by EB, XP, KM, EC, TiW, MP, MCR and RC. AI, KM, SM, ThW and FM analyzed images. EB, XP, KM, SM, EC, ThW, MP, AI, FM, MCR and RC analyzed the data. EB, XP, KM prepared the Figures. EB, XP, KM and SM wrote the manuscript.
References


Methods

Generation and maintenance of cell lines

The HeLa-Kyoto cells stably transfected with the KIF1C-GFP BAC were previously described (Maliga et al. 2013; Poser et al. 2008; Chouaib et al. 2020). HeLa Flp-in H9 (a kind gift of S. Emiliani) and the BAC-GFP cells were maintained in Dulbecco’s modified Eagle’s Medium (DMEM, Gibco) supplemented with 10% fetal bovine serum (FBS, Sigma), 100 U/mL penicillin/streptomycin (Sigma) and with 400 µg/ml G418 (Gibco) for the HeLa-Kyoto KIF1C-GFP tagged BAC cells. NIH/3T3 cells were maintained in DMEM supplemented with 10% calf serum, sodium pyruvate and penicillin/streptomycin at 37°C, 5% CO2. MDA-MB-231 human breast cancer cells (ATCC) were grown in Leibovitz’s L15 media supplemented with 10% fetal bovine serum and penicillin/streptomycin at 37°C in atmospheric air. Stable HeLa cell lines expressing a KIF1C-GFP cDNA were created using the Flp-in system in HeLa H9 cells. Flp-in integrants were selected on hygromycin (150 µg ml⁻¹). To generate cell lines expressing RNA reporters, NIH/3T3 cells were infected with lentivirus expressing tdMCP-GFP (Addgene plasmid #40649) and GFP-expressing cells with low level of GFP expression were sorted by FACS. This stable population was infected with pInducer20-based reporter constructs expressing β-globin followed by 24xMS2 binding sites and the mouse Net1, Rab13 or control 3’UTRs (pIND20-β24bs/Net1 3’UTR; pIND20-β24bs/Rab13 3’UTR; pIND20-β24bs/Ctrl 3’UTR; Moissoglu et al, 2019). Stable lines were selected with geneticin (Thermo Fisher Scientific) and expression of the reporter was induced by addition of 1 µg/ml doxycycline (Fisher Scientific) 2-3 hours before imaging.

For the two-color tracking experiment, NIH/3T3 cells expressing the pIND20-β24bs/Net1 reporter and tdMCP-GFP described below were modified as follows. Stable expression of scFv-mScarletI-GB1 in NIH/3T3 cells was achieved by lentiviral-mediated integration of a plasmid derived from Addgene (#60906) and sorted by FACS with low level
of mScarletI expression. Then a lentivirus expressing a KIF1C fusion with 24 repeats of the GCN4 peptide array was used to infect these NIH/3T3 cells allowing the detection of KIF1C-STx24 protein with scFv-mScarletI-GB1.

Treatments with siRNAs and drugs

HeLa cells were seeded on 0.17 mm glass coverslips deposited in 6-well plates. Cells were transfected at 70% confluency using JetPrime (Polyplus). Double-stranded siRNAs (30 pmoles) were diluted into 200 μl of JetPrime buffer. JetPrime reagent was added (4 μl) and the mixture was vortexed. After 10 minutes at room temperature (RT), it was added to the cells grown in 2 ml of serum-containing medium. After 24 hours, the transfection medium was replaced with fresh growth medium and cells were fixed 24h later. The sequences of the siRNA were: KIF1C: 5’-CCCAGCCGUCUUUACCAUdCdG; control: 5’-CAACAGAAGGAGAGCGAAAdTdT. For knockdown of human or mouse Kif1c the following additional siRNAs were used: Hs_KIF1C_5 (Qiagen cat# SI02655401), Hs_KIF1C_6 (Qiagen cat# SI02781331), Hs_KIF1C_7 (Qiagen cat# SI02781338), Hs_KIF1C_8 (Qiagen cat# SI03019744), Mm_Kif1c_2 (Qiagen cat# SI00239687), Mm_Kif1c_3 siRNA (Qiagen cat# SI00239694) and AllStars negative control siRNA (Qiagen cat# 1027281). siRNAs were delivered into cells using Lipofectamine RNAiMAX (Thermo Fisher Scientific, cat# 13778-150) according to the manufacturer’s instructions. Cells were assayed 72 hours after siRNA transfection.

For drug treatments, 10 μM nocodazole or Cytochalasin D, or an equal volume of DMSO, were added to the growth media for 15 min.

Plasmid construction
Plasmids were generated with standard molecular biology techniques. The GFP-APC plasmid was a gift from Inke Nathke. The KIF1C-mCherry plasmid was a gift from Anne Straube (Addgene plasmid #130978). The inducible constructs expressing MS2-containing RNA reporters (pIND20-β24bs/Net1 3’UTR; pIND20- β24bs/Rab13 3’UTR) are described in Moissoglu et al, 2019. They contain the human β-globin gene followed by 24xMS2 binding sites and the mouse Net1, Rab13 or control 3’UTRs. The control reporter (pIND20- β24bs/Ctrl 3’UTR) carries a short, random, vector-derived UTR sequence. To generate the KIF1C-SunTagx24 cell lines, the KIF1C cDNA was amplified by PCR and cloned in pHrdSV40-K560-GCN4x24 (Addgene #72229), in place of the K560 cDNA and upstream of the SunTag. Then the KIF1C-STx24 sequence was PCR amplified and cloned into pHAGE-Ubc-MCP-YFP (Addgene #31230) in place of the MCP-GFP sequence. Maps and sequences are available upon request.

**Immunoprecipitation and microarrays**

HeLa cells containing the KIF1C-GFP BAC were grown to near confluence in 10 cm plates, and two plates were used per IP. Cells were rinsed in ice-cold PBS, and all subsequent manipulations were performed at 4°C. Cells were scraped in HTNG buffer (20 mM HEPES-KOH pH 7.9, 150 mM NaCl, 1% Triton X-100, 10% glycerol, 1 mM MgCl₂, 1 mM EGTA), containing an antiprotease cocktail (Roche Diagnostic). Cells were incubated for 20 minutes on a rotating wheel, and cellular debris were removed by centrifugating the extracts 10 minutes at 20,000g. Beads coated with GFP-trap antibody (ChromoTek), or uncoated as control, were washed in HNTG (25 μl of beads per IP). Beads were incubated 1h with a control extract to saturate non-specific binding and then incubated with the proper extract. After 4h of incubation on a rotating wheel, beads were washed four times in HNGT with anti-protease, and twice with PBS. Beads were then incubated with Trizol to extract RNAs, and RNA purification was done.
as recommended by the manufacturer. The resulting RNAs were amplified and converted into cDNAs by the WT PICO kit (Thermo Fisher), and hybridized on HTA 2.0 chip on an Affymetrix platform (Thermo Fisher). Experiments were performed in duplicates, data were normalized and averaged. Data are deposited on GEO with the following accession number (GSE161316).

For GFP-APC immunoprecipitation, cells were lysed with a buffer containing 50mM Tris pH 7.4, 1% Triton X-100, 75mM NaCl, 10mM MgCl₂, 10% glycerol and a cocktail of protease and phosphatase inhibitors (Thermo Fischer Scientific, cat# 1861281). Lysates were cleared by centrifugation and mixed with GFP-Trap Magnetic Agarose beads (Chromotek, cat# gtma-10) for 1.5 h, at 4°C. Immobilized complexes were eluted with Laemmli’s buffer and analyzed by SDS-PAGE and immunoblotting.

**Western blot**

For Western blot detection the following antibodies were used: anti-GFP rabbit polyclonal (Invitrogen, cat# A11122, 1/2,000 dilution), anti-KIF1C rabbit polyclonal (Proteintech, cat# 12760-1-AP), anti-KIF1C rabbit polyclonal (Bethyl Laboratories, cat# A301-070A, 1/2,000 dilution), anti-α-tubulin mouse monoclonal (Sigma-Aldrich, cat# T6199, 1/10,000 dilution), anti-mCherry mouse monoclonal [1C51] (Abcam, cat# ab125096, 1/2,000 dilution).

**RNA analyses**

For total RNA analysis, cells were lysed with Trizol LS reagent (Thermo Fisher Scientific, cat# 10296010) and RNA was extracted according to the manufacturer’s instructions. Isolated RNA was treated with RQ1 RNase-free DNase (M6101, Promega) and analyzed with the nCounter system (NanoString Technologies) using a custom-made codeset. Data were processed using nSolver analysis software (NanoString technologies).
Single molecule FISH

Cells grown on glass coverslips were fixed for 20 min at RT with 4% paraformaldehyde diluted in PBS, and permeabilized with 70% ethanol overnight at 4°C. For smFISH, we used a set of 44 amino-modified oligonucleotide probes against the GFP-IRES-Neo sequence (sequences given in Table S2). Each oligonucleotide probe contained 4 primary amines that were conjugated to Cy3 using the Mono-Reactive Dye Pack (PA23001, GE Healthcare Life Sciences). To this end, the oligos were precipitated with ethanol and resuspended in water. For labelling, 4 μg of each probe was incubated with 6 μl of Cy3 (1/5 of a vial resuspended in 30 μl of DMSO), and 14 μl of carbonate buffer 0.1 M pH 8.8, overnight at RT and in the dark, after extensive vortexing. The next day, 10 μg of yeast tRNAs were added and the probes were precipitated several times with ethanol until the supernatant lost its pink color. For hybridization, fixed cells were washed with PBS and hybridization buffer (15% formamide in 1xSSC), and then incubated overnight at 37°C in the hybridization buffer also containing 130 ng of the probe set for 100 μl of final volume, 0.34 mg/ml tRNA (Sigma), 2 mM VRC (Sigma), 0.2 mg/ml RNAse-free BSA (Roche Diagnostic), and 10% Dextran sulfate. The next day, the samples were washed twice for 30 minutes in the hybridization buffer at 37°C, and rinsed in PBS. Coverslips were then mounted using Vectashield containing DAPI (Vector laboratories, Inc.).

For smiFISH (Tsanov et al. 2016), 24 to 48 unlabeled primary probes were used (sequences given in Table S2). In addition to hybridizing to their targets, these probes contained a FLAP sequence that was hybridized to a secondary fluorescent oligonucleotide. To this end, 40 pmoles of primary probes were pre-hybridized to 50 pmoles of secondary probe in 10 μl of 100 mM NaCl, 50 mM Tris-HCl, 10 mM MgCl2, pH 7.9. Hybridization was performed at 85°C for 3 min, 65°C for 3 min, and 25°C for 5 min. The final hybridization mixture contained the
probe duplexes (2 µl per 100 µl of final volume), with 1X SSC, 0.34 mg/ml tRNA (Sigma), 15% Formamide, 2 mM VRC (Sigma), 0.2 mg/ml RNAse-free BSA, 10% Dextran sulfate. Slides were then processed as above. For Figure S1A, the probes used were RNA and not DNA (sequence in Table S2). The protocol was similar except that hybridization was performed at 48°C and that 50 ng of primary probe (total amount of the pool of probes) and 30 ng of the secondary probes were used per 100 µl of hybridization mix.

For FISH of mouse cells, cells plated on fibronectin-coated coverslips were fixed for 20 min at RT with 4% paraformaldehyde in PBS. FISH was performed with the ViewRNA ISH Cell Assay kit (Thermo Fisher Scientific) according to the manufacturer’s instructions. The flowing probe sets were used: Kif1c (cat# VB6-3200442), Net1 (cat# VB1-3034209), Rab13 (cat# VB1-14374), Ddr2 (cat# VB1-14375), Dynll2 (cat# VB1-18646), Cyb5r3 (cat# VB1-18647). To detect polyA+ RNAs, LNA modified oligodT probes (30 nucleotides) labelled with ATTO 655 were added during hybridization, pre-amplification, amplification and last hybridization steps of the ViewRNA ISH Cell Assay. Cell mask stain (Thermo Fisher Scientific) was used to identify the cell outlines. Samples were mounted with ProLong Gold antifade reagent (Thermo Scientific)

**Imaging of fixed cells**

Microscopy slides were imaged on a Zeiss Axioimager Z1 wide-field microscope equipped with a motorized stage, a camera scMOS ZYLA 4.2 MP, using a 63x or 100x objective (Plan Apochromat; 1.4 NA; oil). Images were taken as z-stacks with one plane every 0.3 µm. The microscope was controlled by MetaMorph and figures were constructed using ImageJ, Adobe Photoshop and Illustrator. For the small smiFISH screen, 96-well plates were imaged on an Opera Phenix High-Content Screening System (PerkinElmer), with a 63x water-immersion objective (NA 1.15). Three-dimensional images were acquired, consisting of 35 slices with a
spacing of 0.3 µm. FISH images of mouse cells were obtained using a Leica SP8 confocal microscope, equipped with a HC PL APO 63x CS2 objective. Z-stacks through the cell volume were obtained and maximum intensity projections were used for subsequent analysis.

**Image analysis and quantifications**

Automated nuclear and cell segmentation was performed with a custom algorithm based on the U-net deep convolutional network (Ronneberger et al. 2015). Nuclear segmentation was performed with the DAPI channel, cell segmentation was performed with the autofluorescence of the actual smFISH image. For segmentation, 3D images were projected into 2D images as described previously (Tsarev et al. 2016). Messenger RNAs were detected with FISH-quant (Mueller et al. 2013) by applying a local maximum detection on LoG filtered images. For calculation of Peripheral Distribution Index (PDI) a custom Matlab script was used. The code is described and is available in (Stueland et al. 2019).

**Imaging of live cells**

Live imaging (for dual visualization of β24bs/Net1 reporter RNA and KIF1C-STx24 protein) was done using a spinning disk confocal microscope (Nikon Ti with a Yokogawa CSU-X1 head) operated by the Andor iQ3 software. Acquisitions were performed using a 100X objective (CF1 PlanApo λ 1.45 NA oil), and an EMCCD iXon897 camera (Andor). For two-color imaging, samples were sequentially excited at 488 and 540nm. We imaged at a rate of 7.36 fps for 52 sec. The power of illuminating light and the exposure time were set to the lowest values that still allowed visualization of the signal. This minimized bleaching, toxicity and maximized the number of frames that were collected. Cells were maintained in anti-bleaching live cell visualization medium (DMEM^gfp; Evrogen), supplemented with 10% fetal bovine serum at 37°C in 5% CO₂ and rutin at a final concentration of 20 mg/l.
Live imaging (for β24bs/Net1 reporter RNA tracking) was done using a Nikon Eclipse Ti2-E inverted microscope, equipped with a motorized stage, a Yokogawa CSU-X1 spinning disk confocal scanner unit, and operated using NIS-elements software. Acquisitions were performed using an Apochromat TIRF 100x oil immersion objective (N.A. 1.49, W.D. 0.12mm, F.O.V. 22mm) and a Photometrics Prime 95B Back-illuminated sCMOS camera with W-view Gemini Image splitter. Constant 37°C temperature and 5% CO2 were maintained using a Tokai Hit incubation system. Cells were plated on fibronectin (2mg/ml)-coated 35mm glass bottom dishes and samples were excited using a 488nm (20mw) laser line and imaged at a rate of 6.66 fps for 60 sec.

**Live cell imaging quantification**

Images were processed for brightness/contrast, cropped and annotated using ImageJ/FIJI. Kymographs were generated using standard ImageJ/Fiji plugins. Film presentation in figures and videos were edited in ImageJ/Fiji. Bicolor tracking of KIF1C-STx24 proteins and β24bs/Net1 mRNAs was performed using the Manual Tracking plugin in ImageJ/Fiji.

Single color tracking of β24bs/Net1 mRNAs was performed using TrackMate plugin in ImageJ/Fiji. For every cell, all tracks lasting for >2.5 secs (ca. 17 consecutive frames) were used for analysis. Values of ‘Track displacement’, ‘Linearity of forward progression’, ‘track duration’ and ‘mean speed’ were extracted and plotted. ‘Track displacement’ is defined as the distance from the first to the last spot of the track. ‘Linearity of forward progression’ is the mean straight line speed divided by the mean speed; where mean straight line speed is defined as the net displacement divided by the total track time. MSD and velocity autocorrelation of tracks was determined using MSDanalyzer (https://github.com/tinevez/msdanalyzer). For RNA cluster analysis, TrackMate was used to identify spots and extract intensity values. Frequency histograms of spot intensities were plotted using GraphPad Prism software.
FIGURES AND FIGURE LEGENDS

Figure 1: Identification of mRNAs associated with the KIF1C motor

A- The KIF1C kinesin colocalizes with its mRNA in protrusions of HeLa cells. Images are micrographs of a H9 FlipIn HeLa cell line stably expressing a KIF1C-GFP cDNA. Top left: KIF1C mRNA detected by smFISH with probes against the endogenous mRNA; top right: KIF1C-GFP protein; bottom left: DNA stained with DAPI, bottom right: merge of the two signals with the KIF1C-GFP protein in green and KIF1C mRNA in red. Orange arrow: a cell protrusion. Blue: DNA stained with DAPI. Scale bar: 10 microns. Insets represent zooms of the boxed areas in the merge panel. White and black arrowheads indicate the colocalization of KIF1C mRNA with KIF1C-GFP protein.

B- Transcripts associating with the KIF1C-GFP protein. The graph depicts the microarray signal intensity of RNAs detected in a KIF1C-GFP pull-down (x-axis), versus the control IP (y-axis). Each dot represents an mRNA. Red dot: KIF1C mRNA; blue dot: mRNAs enriched in the KIF1C-GFP IP. N=2.

C- Colocalization of KIF1C-GFP with KIF1C, NET1, TRAK2 and RAB13 mRNAs. Images are micrographs of HeLa H9 Flip-In cells stably expressing a KIF1C-GFP cDNA, labelled by smiFISH with probes against the indicated mRNAs. Top: Cy3 fluorescent signals corresponding to endogenous KIF1C, NET1, TRAK2 and RAB13 mRNAs. Middle: KIF1C-GFP signal. Lower: merge with the Cy3 signal in red and GFP signal in green. Blue: DNA stained with DAPI. Scale bar: 10 microns. Arrowheads indicate accumulation of single mRNA molecules at cell protrusions.

D- Single molecule colocalization of KIF1C-ST-x24 with RAB13 mRNAs. Images are micrographs of Hela cells stably expressing KIF1C-STx24 and scFv-GFP. Red: Cy3 fluorescent signals corresponding to RAB13 mRNAs labeled by smiFISH with probes against
endogenous RAB13 mRNA. Green: GFP signal corresponding to single molecules of KIF1C protein. Blue: DNA stained with DAPI. Scale bar: 10 microns.

E- Insets represent zooms of the numbered areas from panel D. Legend as in D. Arrowheads indicate molecules of RAB13 mRNA and KIF1C-STx24 protein. Micrographs show cells representative of the population.

**Figure 2: KIF1C associates with APC and is required for the localization of APC-dependent mRNAs to cytoplasmic protrusions**

A- Co-immunoprecipitation of KIF1C with APC. GFP or GFP-APC were immunoprecipitated from cells also expressing KIF1C-mCherry and analyzed by Western blot to detect the indicated proteins. Results are representative of 3 independent experiments.

B- Colocalization of GFP-APC and KIF1C-mCherry at cytoplasmic protrusions (arrows). Images are representative of multiple cells observed in two independent experiments. Scale bar: 10 microns.

C- Depletion of Kif1c prevents mRNA accumulation in protrusions. Images are micrographs of NIH/3T3 cells labelled by smFISH with probes against *Net1, Rab13, Ddr2* and *Dynll2* mRNAs, following treatment with siRNAs against Kif1c (panels si-Kif1c) or a control sequence (panel si-Control). Scale bars: 10 microns. Green: outline of the cells; blue: outline of the nuclei; black: smFISH signals. Insets represent magnifications of the boxed areas.

D- Quantification of mRNA localization of cells described in C. Graphs represent the intracellular distribution of the indicated mRNAs as measured by PDI index, with and without treatment of cells with the indicated siRNAs. Red bars represent the mean and 95% confidence interval. Points indicate individual cells observed in two independent experiments.

E- Detection of Kif1c protein (upper panels) or Kif1c RNA levels (lower graph) from cells treated with the indicated siRNAs.
Stars in D and E are p-values: ****<0.0001, ***<0.001, estimated by analysis of variance with Bonferroni’s multiple comparisons test.

Figure 3: Reporter mRNAs containing Net1 and Rab13 3’UTRs display long, directed microtubule-dependent displacements.

A- Schematic of the mRNA reporter construct containing the β-globin coding sequence followed by 24xMS2 binding sites and the mouse Net1 3’UTR (β24bs/Net1). Images are snapshots of live NIH/3T3 before or after nocodazole treatment, following the experimental scheme detailed on the left. High speed imaging was performed over 1min to track individual RNA movements. See movies S1 and S2 for time lapse imaging. The cells stably expressed the β24bs/Net1 reporter mRNA and MCP-GFP (Green). Scale bar: 5 microns.

B- The graphs plot the displacement over the linearity of forward progression (defined as the mean straight line speed divided by the mean speed) of single RNA tracks from cells treated as described in A. Red lines indicate the thresholds used to separate ‘long/directed’ from ‘short/diffuse’ tracks. N=6-7 cells.

C- Analysis of ‘long/directed’ or ‘short/diffuse’ tracks, from untreated cells, or of all tracks from nocodazole-treated cells, from B. Individual raw tracks from a representative cell are displayed in the upper panels. Mean Square Displacement (MSD) (middle panels) and velocity autocorrelation (bottom panels) of tracks are shown. N=180 (long/directed); 4427 (short/diffuse); 3795 (after nocodazole) tracks from 7 cells. Note that ‘long/directed’ tracks exhibit higher MSD (>15 μm², dashed black line), than ‘short/diffuse’ or nocodazole-treated tracks, as well as positive velocity autocorrelation. Insets of MSD plots present zooms of y-axis scale.

D- Single RNA molecules of the β24bs/Control 3’UTR (Ctrl) or the β24bs/Rab13 3’UTR reporters were tracked over 1 minute period in cells treated or not with nocodazole. Graphs plot
the displacements of individual tracks (x-axis) over the linearity of their forward progression (y-axis). Red lines indicate the thresholds used to filter tracks of molecules undergoing directed movement. N=8-12 cells.

**E-** The graph depicts the percentage of long/directed tracks of the indicated reporters per cell following treatment with nocodazole. Stars represent p-values: ****<0.0001, estimated using one-way analysis of variance with Sidak’s multiple comparisons test. Error bars: standard error of the mean.

**F-** The bar plot depicts the percentage of long/directed tracks per cell before and after treatment with the indicated compounds. Average values of respective ‘Before’ values were set to 1. N=6-7. Stars represent p-values: ****<0.0001, ns: non-significant, estimated using one-way analysis of variance with Tukey’s multiple comparisons test. Error bars: standard error of the mean.

**Figure 4: Reporter mRNAs containing the Net1 3’UTR require the Kif1c motor for long, linear microtubule-dependent displacements.**

**A-** Images are snapshots of live NIH/3T3 cells taken 30 minutes after plating. The cells stably expressed the β24bs/Net1 mRNA reporter and MCP-GFP (Green) and were treated with the indicated siRNAs. The green spots correspond to single mRNAs detected with the MCP-GFP. High speed imaging was performed over 1 min to track individual RNA movements. See movies S5-S8 for time lapse imaging. Scale bars are 5 microns.

**B-** Graphs plot the displacements of individual RNA tracks (x-axis) over the linearity of their forward progression (y-axis), (defined as the mean straight line speed divided by the mean speed), using the movies of cells as shown in A. Red lines indicate the thresholds used to filter tracks of molecules undergoing directed movement (based on Figure 3).

**C-** Graph depicts the percentage of directed tracks per cell following treatment with the indicated siRNAs (see panel B). N=11-14 cells. Stars represent p-values: ***<0.001, ns: non-
significant, estimated using one-way analysis of variance with Dunnett’s multiple comparisons test. Error bars: standard error of the mean.

**Figure 5: Kif1c is required for the peripheral clustering of reporter mRNAs containing the Net1 3’UTR in live mouse fibroblasts.**

A- Images are micrographs of live NIH/3T3 cells taken 3h after plating and expressing MCP-GFP and the β24bs/Net1 reporter mRNA. Cells were treated with the siRNAs indicated on the left. Scale bar is 10 microns. Boxed insets are magnifications of the areas indicated by green arrows.

B- Frequency histograms of the intensities of the β24bs/Net1 reporter mRNA spots following treatment with the indicated siRNAs, measured from images as shown in A, from N=14-20 cells. The majority of molecules fall under a single lower intensity peak, likely indicative of single molecules, while a smaller fraction exhibits higher intensities indicative of higher order clusters.

C- Graph depicts the mRNA cluster frequency per cell following treatment of cells with the indicated siRNAs. Clusters correspond to β24bs/Net1 mRNA spots of intensities higher than 4,950, measured from the graphs shown in B. N=14-20 cells. Stars represent p-values: **<0.01, estimated using one-way analysis of variance with Dunnett’s multiple comparisons test. Error bars: standard error of the mean.

**Figure 6: The KIF1C motor transports mRNAs containing the Net1 3’UTR to cell protrusions.**

A- Images are micrographs of fixed NIH/3T3 cells expressing the β24bs/Net1 reporter mRNA, MCP-GFP (green), KIF1C-STx24 protein and scFv-mScarletI (red). Single molecules of the β24bs/Net1 reporter mRNA are visible in green, while single molecules of KIF1C-STx24 protein
are red. The numbered white boxes are magnified in B. Blue: DNA stained with DAPI. Scale bar is 5 microns.

B- Insets represent magnifications of the boxed areas from the image shown in A. Left: MCP-GFP signals labelling β24bs/Net1 mRNAs; middle: scFv-mScarletI labelling KIF1C-ST<sub>x24</sub> protein; right: merge with mRNAs in green and KIF1C-ST<sub>x24</sub> in red. Black and white arrowheads indicate colocalization of single molecules of β24bs/Net1 mRNAs and KIF1C-ST<sub>x24</sub>. Scale bar is 5 microns.

C- Snapshot of a live NIH/3T3 cell expressing β24bs/Net1 mRNA, MCP-GFP (green), KIF1C-ST<sub>x24</sub> protein and scFv-mScarletI (red). Snapshot is extracted from Movie S9. The white arrowhead indicates a co-transport event of a single molecule of β24bs/Net1 mRNA (green) with a KIF1C-ST<sub>x24</sub> protein (red). The boxed area is magnified in panels D and E. Scale bar is 5 microns.

D- Magnification of the boxed area in panel C, highlighting a co-transport event. Top: KIF1C-ST<sub>x24</sub>; Middle: β24bs/Net1 mRNA; bottom: merged panel with the β24bs/Net1 mRNA in green and KIF1C-ST<sub>x24</sub> in red. Scale bar is 1 micron.

E- Kymograph from Movie S9, showing the trajectory of a single molecule of KIF1C-ST<sub>x24</sub> (top panel), a single molecule of β24bs/Net1 mRNA (middle panel) and the merge (bottom panel). The cellular area shown corresponds to panel D.

F- The graph depicts the distance travelled by co-transported molecules of KIF1C-ST<sub>x24</sub> and β24bs/Net1 mRNA. Each data point is a track (40 tracks in total), and the mean and 95% confidence intervals are shown by horizontal lines. Source data are provided in Table S4.

G- Boxplot depicting the mean speed of co-transported molecules of KIF1C-ST<sub>x24</sub> and β24bs/Net1 mRNA in NIH/3T3 cells. Speed is microns/second. The vertical bars display the first and last quartile, the box corresponds to the second and third quartiles, and the horizontal line to the mean (40 tracks in total). Source data are provided in Table S4.
**Figure S1: Identification of mRNAs associated with the KIF1C motor.**

**A**- Localization of mRNAs in protrusion of human cells. Images are micrographs of HeLa cells labelled by smiFISH with probes against endogenous genes. Top: Cy3 fluorescent signals corresponding to endogenous KIF1C, NET1, TRAK2 and RAB13 mRNAs. Bottom: merge panel with red being the Cy3 fluorescent smiFISH signals and blue corresponding to DNA stained with DAPI. Scale bar: 10 microns. Arrowheads indicate accumulation of single molecule mRNA at cell protrusions.

**B** - The KIF1C motorpreferentially associates with APC-dependent mRNAs. Correlation analysis of mRNA enrichment in protrusions (x axis; Ps/CB is the Protrusion versus Cell Body ratio; data from Wang et al., 2017) versus the enrichment fold of the same mRNAs in KIF1C-GFP IP (y-axis; KIF1C-GFP versus control IP). Red dots depict APC-dependent mRNAs, green dots depict ribosomal protein mRNAs (APC-independent mRNAs), based on data from Wang et al., 2017. See Table S3 for source data.

**C-E** Single molecule colocalization of KIF1C-ST-x24 with NET1 mRNAs. Images are micrographs of Hela cells stably expressing KIF1C-STx24 and scFv-GFP. Red: Cy3 fluorescent signals corresponding to mRNAs labeled by smiFISH with probes against endogenous NET1 mRNA (D), or CRM1 (C) and RPB1 (E) as controls. Green: GFP signal corresponding to KIF1C motor protein. Blue: DNA stained with DAPI. Scale bar: 10 microns in the main panels, 2.5 microns in the zoom. Arrowheads: single mRNA molecules.

**Figure S2:** **KIF1C is required for the localization of APC-dependent mRNAs to cytoplasmic protrusions in mouse and human cells.**

**A**- Efficiency of the KIF1C knock-down measured by smiFISH. The graph depicts the number of single molecules of KIF1C mRNA per HeLa cell, following treatment with control siRNA
(orange bar) or KIF1C siRNA (blue bar). Error bars: standard deviation (n=2); number of cells: 50 per condition.

**B-** The KIF1C motor is required for accumulation of RAB13 mRNA in protrusions of HeLa cells. Images are micrographs of HeLa cells labelled by smiFISH with probes against RAB13 mRNAs. Left: Hela cells treated with a control siRNA; right: HeLa cells treated with siRNA against KIF1C. Top: Cy3 fluorescent signal corresponding to endogenous RAB13 mRNA. Lower panels are the merge with Cy3 fluorescent signals in red and DNA in blue (DAPI staining). Scale bar: 10 microns. The orange arrow indicates a protrusion accumulating RAB13 mRNA.

**C-** Quantification of NET1 or RAB13 mRNA localization in HeLa cells treated with the indicated siRNAs against KIF1C. Bars represent the mean and error bars the 95% confidence interval. Points indicate individual cells (N=30) observed in two independent experiments.

**D-** Treatment of MDA-MB-231 cells with the indicated siRNAs against KIF1C. Left panel: Western blot indicating the efficiency of KIF1C knockdown. Right graphs: Quantification of NET1 or RAB13 mRNA localization. Bars represent the mean and error bars the 95% confidence interval. Points indicate individual cells (N=30) observed in two independent experiments.

**E-** Micrographs of NIH/3T3 cells labelled by smFISH with probes against the indicated mRNAs. Cells were treated with Kif1C or Control siRNAs. Green outline: cell contour; blue outline: DAPI staining; black spots: mRNAs detected by smFISH. Scale bars: 10 microns.

**F-** Intracellular distribution of Cyb5r3 mRNA calculated by their PDI values, after treatment with Control or Kif1C siRNAs. Horizontal lines represent the mean with 95% confidence interval. Points indicate individual cells. $p<0.0001$, estimated by Student’s t-test.

**G-** Localization of APC-independent RNAs is not affected by Kif1c depletion in NIH/3T3 fibroblasts. The graph plots PDI values of Ddr2, Rps20, Rpl27a and polyA+ RNAs, measured
from smFISH images of cells treated with Control or Kif1c siRNAs. Points indicate individual cells. Stars are p-values: ****<0.0001, estimated by analysis of variance with Tukey’s multiple comparisons test.

**H**- Expression of localized (APC-dependent and -independent) and non-localized mRNAs is not affected by Kif1c depletion in NIH/3T3 fibroblasts. The graph depicts Log2 RNA expression ratios (Kif1c depleted versus control cells), measured by Nanostring analysis. n=3. Bars indicate mean +/- standard error.

**Figure S3: Analysis of single molecule RNA tracks**

**A**- Frequency histogram of track duration of long/directed or short/diffuse tracks. Analyzed tracks correspond to tracks of the β24bs/Net1 reporter RNA from untreated cells presented in Figure 3A-C.

**B**- Frequency histogram of mean track speed of the same tracks analyzed in A.

**Figure S4: Long/directed tracks of the β24bs/Net1 reporter RNA do not require the actin cytoskeleton**

Images are snapshots of live NIH/3T3 cells taken 30 minutes after plating. High speed imaging was performed over 1min to track individual RNA movements before and after 15 minutes of Cytochalasin D or vehicle control (DMSO) as indicated. See movies S3 and S4 for time lapse imaging. The cells stably expressed the β24bs/Net1 reporter mRNA and MCP-GFP (Green). Scale bar: 5 microns. The associated graphs plot the displacements of individual tracks (x-axis) over the linearity of their forward progression (y-axis), (defined as the mean straight line speed divided by the mean speed). Red lines indicate the thresholds used to filter tracks of molecules undergoing directed movement. N=6-7 cells.
Figure S5: Reporter mRNAs containing Net1 and Rab13 3’UTRs form clusters at the tips of protrusions in mouse fibroblasts

Micrographs of live NIH/3T3 cells taken 3h after plating and expressing MCP-GFP and the β24bs reporter mRNA carrying the indicated 3’UTRs. Red line: cell contour. Scale bars are 10 microns. Note that reporter mRNAs containing the Net1 or Rab13 3’UTR accumulate in clusters at the tips of protrusions, while the control reporter does not. Boxed insets (right) are magnifications of the areas indicated by the green arrows.

Figure S6: Reporter mRNAs containing the Net1 3’UTR are transported to protrusions by the Kif1c motor.

A- Micrograph of a fixed NIH/3T3 cell expressing β24bs/Net1 reporter mRNA, MCP-GFP (green), KIF1C-STx24 protein and scFv-mScarletI (red). Single molecules of β24bs/Net1 mRNAs are visible in green, while single molecules of KIF1C-STx24 protein are red. The numbered white boxes are magnified in B. Blue: DNA stained with DAPI. Scale bar is 5 microns.

B- Insets represent magnifications of the boxed areas from the image shown in A. Left: MCP-GFP signals labelling β24bs/Net1 mRNAs; middle: scFv-mScarletI labelling KIF1C-STx24 protein; right: merge with mRNAs in green and KIF1C-STx24 in red. Black and white arrowheads indicate colocalization of single molecules of β24bs/Net1 mRNA and KIF1C-STx24. Scale bars are 5 microns.

C- Snapshot of a live NIH/3T3 cell expressing β24bs/Net1 mRNA, MCP-GFP (green), KIF1C-STx24 protein and scFv-mScarletI (red). Snapshot is extracted from Movie S10. The white arrowhead indicates a co-transport event of a single molecule of β24bs/Net1 mRNA (green) with a KIF1C-STx24 protein (Red). The boxed area is zoomed in panels D and E. Scale bar is 5 microns.
D- Magnification of the boxed area in panel C, highlighting a co-transport event. Top: KIF1C-STx24; Middle: β24bs/Net1 mRNA; bottom: merged panel with the β24bs/Net1 mRNA in green and KIF1C-STx24 in red. Scale bar is 1 micron.

E- Kymograph extracted from Movie S10, showing the trajectory of a single molecule of β24bs/Net1 mRNA (left panel), KIF1C-STx24 (middle panel) and the merge (right panel) in NIH/3T3 cells. Total time of the kymograph is 12.9 seconds and corresponds to the event shown in panels C and D.
Supplementary spreadsheets:

**Table S1:** Excel file showing the KIF1C-GFP IP microarray data.

**Table S2:** Human smFISH probe sequences and list of probes used for smFISH localization screen in HeLa cells using 18 of the mRNAs most enriched in the KIF1C IP.

**Table S3:** File presenting the source data for the graph presented in Figure S1E. APC-dependence was specified based on reduced protrusion enrichment upon APC knockdown (described in Wang et al, 2017). For each RNA, the fold-change of protrusion enrichment and significance values upon APC knockdown from the Wang et al study are also indicated.

**Table S4:** File presenting the source data for the graphs presented in Figure 6G-H.

**Table S5:** GO term enrichment for the 200 mRNAs most enriched in the KIF1C IP. Analysis was done with http://pantherdb.org/tools/compareToList.jsp. The terms highlighted in yellow could relate to the function of KIF1C in cell migration.

Movie legends:

**Movie S1:** A live NIH/3T3 cell expressing β24bs/Net1 reporter mRNA and MCP-GFP (green) imaged at 6.67 frames per second for 1 minute, prior to addition of nocodazole. A snapshot of this movie is shown in Figure 3B. Left panel presents the raw signal. Right panel presents the raw signal with overlaid accumulated tracks of individual RNA spots. Only tracks lasting longer than 2.5 seconds are shown. Warmer colors indicate tracks of higher linearity. Scale bar is 5 microns.

**Movie S2:** A live NIH/3T3 cell (same as shown in Movie S1) expressing β24bs/Net1 reporter mRNA and MCP-GFP (green) imaged at 6.67 frames per second for 1 minute, 15 minutes after addition of nocodazole. A snapshot of this movie is shown in Figure 3B. Left panel presents the raw signal. Right panel presents the raw signal with overlaid accumulated tracks of
individual RNA spots. Only tracks lasting longer than 2.5 seconds are shown. Warmer colors indicate tracks of higher linearity. Scale bar is 5 microns.

**Movie S3:** A live NIH/3T3 cell expressing β24bs/Net1 reporter mRNA and MCP-GFP (green) imaged at 6.67 frames per second for 1 minute, prior to addition of cytochalasin D. A snapshot of this movie is shown in Figure 3B. Left panel presents the raw signal. Right panel presents the raw signal with overlaid accumulated tracks of individual RNA spots. Only tracks lasting longer than 2.5 seconds are shown. Warmer colors indicate tracks of higher linearity. Scale bar is 5 microns.

**Movie S4:** A live NIH/3T3 cell (same as shown in Movie S3) expressing β24bs/Net1 reporter mRNA and MCP-GFP (green) imaged at 6.67 frames per second for 1 minute, 15 minutes after addition of cytochalasin D. A snapshot of this movie is shown in Figure 3B. Left panel presents the raw signal. Right panel presents the raw signal with overlaid accumulated tracks of individual RNA spots. Only tracks lasting longer than 2.5 seconds are shown. Warmer colors indicate tracks of higher linearity. Scale bar is 5 microns.

**Movie S5:** A live NIH/3T3 cell expressing β24bs/Net1 reporter mRNA and MCP-GFP (green), transfected with control siRNAs, imaged at 6.67 frames per second for 1 minute. A snapshot of this movie is shown in Figure 4A. Left panel presents the raw signal. Right panel presents the raw signal with overlaid accumulated tracks of individual RNA spots. Only tracks lasting longer than 2.5 seconds are shown. Warmer colors indicate tracks of higher linearity. Scale bar is 5 microns.

**Movie S6:** A live NIH/3T3 cell expressing β24bs/Net1 reporter mRNA and MCP-GFP (green), transfected with KIF1C siRNAs, imaged at 6.67 frames per second for 1 minute. A snapshot of this movie is shown in Figure 4A. Left panel presents the raw signal. Right panel presents the raw signal with overlaid accumulated tracks of individual RNA spots. Only tracks lasting longer
than 2.5 seconds are shown. Warmer colors indicate tracks of higher linearity. Scale bar is 5 microns.

**Movie S7:** A live NIH/3T3 cell expressing β24bs/Net1 reporter mRNA and MCP-GFP (green), transfected with KIF5B siRNAs, imaged at 6.67 frames per second for 1 minute. A snapshot of this movie is shown in Figure 4A. Left panel presents the raw signal. Right panel presents the raw signal with overlaid accumulated tracks of individual RNA spots. Only tracks lasting longer than 2.5 seconds are shown. Warmer colors indicate tracks of higher linearity. Scale bar is 5 microns.

**Movie S8:** A live NIH/3T3 cell expressing β24bs/Net1 reporter mRNA and MCP-GFP (green), transfected with KIF3A siRNAs, imaged at 6.67 frames per second for 1 minute. A snapshot of this movie is shown in Figure 4A. Left panel presents the raw signal. Right panel presents the raw signal with overlaid accumulated tracks of individual RNA spots. Only tracks lasting longer than 2.5 seconds are shown. Warmer colors indicate tracks of higher linearity. Scale bar is 5 microns.

**Movie S9:** A live NIH/3T3 cell expressing β24bs/Net1 reporter mRNA, MCP-GFP (green), KIF1C-ST_{x24} protein and scFv-mScarletI (red) was imaged at 7.36 frames per second during 6.45 seconds. The movie follows a co-directed transport event highlighted by the white arrowhead in Figure 6C-E. Scale bar is 5 microns. Top left insets are magnifications of the yellow boxed area showing separately each fluorescent channel. Scale bar is 1 micron.

**Movie S10:** A live NIH/3T3 cell expressing β24bs/Net1 reporter mRNA, MCP-GFP (green), KIF1C-ST_{x24} protein and scFv-mScarletI (red) was imaged at 7.36 frames per second during 12.9 seconds. The movie follows a co-directed transport event highlighted by the white arrowhead in Figure S4C-E. Scale bar is 5 microns. Top left insets are magnifications of the yellow boxed area showing separately each fluorescent channel. Scale bar is 1 micron.
A

- β-globin gene
- 24s UTR

Plating for 30min
1min imaging
Inhibitor for 15min
1min imaging

15min Nocodazole

B

**Linearity of forward progression**

*(Short/Diffuse)*

Track displacement

15min Nocodazole

C

**Before Nocodazole**

- ‘Long/Directed’ tracks (above cutoffs)
- ‘Short/Diffuse’ tracks (below cutoffs)
- All tracks

Y (µm)

X (µm)

MSD (µm²)

Normalized velocity autocorrelation

D

- Ctrl UTR
- Rab13 UTR

Linearity of forward progression

Track displacement

E

% Long/Directed tracks/cell

UTR:
- Ctrl
- Rab13
- Net1

Nocodazole:
- - + +

**ns**

****
NIH/3T3 cells
plating for 30min
1min imaging

A

si-Control
si-KIF1C
si-KIF5B
si-KIF3A

Reporter:
β-globin/24bs-Net1 UTR

B

Linearity of forward progression

Track displacement

14 cells
11 cells
12 cells
11 cells

C

% Long/Directed tracks/cell

si-Ctrl
si-KIF1C
si-KIF5B
si-KIF3A

ns
ns

***
A β-globin/24xMS2bs-Net1 UTR

B

si-Control

RNA clusters

si-KIF1C

si-KIF5B

si-KIF3A

NIH/3T3 cells
3hrs plating

C

Spot intensity

Cluster frequency

si-KIF1C

si-KIF5B

si-KIF3A

si-KIF5B

3194 spots from 14 cells

4466 spots from 20 cells

3733 spots from 17 cells

3942 spots from 15 cells

**
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RNA published online September 7, 2021

Supplemental Material
http://rnajournal.cshlp.org/content/suppl/2021/09/07/rna.078576.120.DC1

Peer-reviewed and accepted for publication but not copyedited or typeset; accepted manuscript is likely to differ from the final, published version.

Published online September 7, 2021 in advance of the print journal.

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