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Rabs on the fly: Functions of Rab GTPases during development

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ABSTRACT
The organization of intracellular transport processes is adapted specifically to different cell types, developmental stages, and physiologic requirements. Some protein traffic routes are universal to all cells and constitutively active, while other routes are cell-type specific, transient, and induced under particular conditions only. Small GTPases of the Rab (Ras related in brain) subfamily are conserved across eukaryotes and regulate most intracellular transit pathways. The complete sets of Rab proteins have been identified in model organisms, and molecular principles underlying Rab functions have been uncovered. Rabs provide intracellular landmarks that define intracellular transport sequences. Nevertheless, it remains a challenge to systematically map the subcellular distribution of all Rab and their functional interrelations. This task requires novel tools to precisely describe and manipulate the Rab machinery in vivo. Here we discuss recent findings about Rab roles during development and we consider novel approaches to investigate Rab functions in vivo.

Introduction
Intracellular transport is adjusted to the demands and functions of diverse cell types. Differentiating and mature cells within tissues have distinct requirements for membrane and protein transport. However, independent of cell type and organismal developmental stage, cells must maintain Essential Permanent protein/membrane Traffic (EPT) routes. Well-known EPTs are the endocytic pathway or the transport route between the Endoplasmic reticulum (ER) and the Golgi apparatus. Rab proteins that control EPTs are among the first transport regulators to be identified. However, since these Rab proteins regulate early trafficking steps at the basis of the cellular transport machineries, the loss-of-function phenotypes of these EPT regulators are often the sum of perturbations of multiple intracellular pathways, making it difficult to pinpoint the mechanistic role of individual Rabs or transport pathways. Therefore, novel approaches are required to study individual pathways and to perturb traffic regulators in a temporally and spatially controlled manner.

While EPTs are commonly required in most cell types, specialized cellular functions and developmental processes require Dedicated Transient Traffic (DTT) routes. DTTs usually do not control essential cellular functions, but are induced on demand in response to external signals, such as mechanical or immunological stimuli. These signals often trigger intracellular events, such as the release of second messengers or transient increases in intracellular Ca²⁺ levels. Well-studied examples are neurotransmitter release in neurons and pulsed secretion by salivary gland cells. Genetic disruption of DTT transport often results in morphological or functional defects, although cell viability is usually not affected upon loss of Rab controlling DTTs.

Here we discuss recent work on Rab-regulated transport routes in Drosophila melanogaster. Small GTPases are master regulators of intracellular membrane and protein transport and control the spatio-temporally targeted delivery of membrane and protein payloads. In the past decade all Rab family members in Drosophila were identified and their regulative elements characterized. In addition, comprehensive collections of inducible Rab transgenes were generated that allow targeted misexpression of tagged Rab proteins. While these genetic
resources have provided many important insights into the organization of the Rab machinery, they are based on overexpression of Rab proteins. However, excess of certain Rab proteins has been shown to change transport and to generate mutant phenotypes.\textsuperscript{11-13} Therefore, maintaining endogenous expression levels of Rab proteins is a key prerequisite for faithfully charting intracellular traffic. Dunst et al. reported the generation of endogenous YFP-tagged \textit{rab} alleles (\textit{YRabs}) and characterized them in 6 different organs consisting of 23 different cell types.\textsuperscript{14} This resource is suitable to predict and compare traffic in different cell types, distinct cell differentiation stages or allows screening Rabs involved in cargo-specific transport. Caviglia et al. used the \textit{YRab} collection to investigate the identity of the membrane material that mediates the formation of tubular connections during tracheal morphogenesis in \textit{Drosophila}.\textsuperscript{15} They identified Rab7 and Rab39, which, together with Munc13–4/Staccato, act as regulators of lysosome-related organelles (LROs) in tracheal tip cells.

**The Rab activity cycle: hard facts and open questions**

Rabs are switchable protein-binding platforms. Active Rabs are GTP-bound and anchored to membranes, where they recruit effector proteins involved in specific traffic steps\textsuperscript{16,17} or signaling cascades.\textsuperscript{18,19} Rab proteins are directly controlled by GDP/GTP exchange factors (GEFs) and GTPase activating proteins (GAPs). Once Rab-containing vesicles reach their target (delivery) sites, local GAPs trigger hydrolysis of Rab-bound GTP, thus rendering Rabs inactive. Deactivated GDP-bound Rabs are then extracted from membranes through mechanisms that are not yet fully understood. It has been suggested that lipid translocation proteins are involved. Recently, one study using attenuated total reflect-Fourier transform infrared (ATR-FTIR) spectroscopy proposed that GDP-dissociation factors (GDIs)\textsuperscript{20-22} directly mediate membrane extraction.\textsuperscript{23} ATR-FTIR involves the collection of infrared radiation reflected from the interphase between an aqueous phase and a prism, in which the evanescent wave penetrated from the prism into the aqueous solution is absorbed by substances solved in the aqueous phase.\textsuperscript{24}

The subsequent formation of cytosolic Rab-GDP/GDI complexes enables inactive Rab-GDPs to recycle and reintegrate into their original membranes.\textsuperscript{25} How are Rab-GDP/GDI assemblies trafficked? One possibility is that cytosolic Rab-GDP/GDI complexes are present in excess over local membrane-associated Rab-GTP pools. In this scenario no directed transport is required between different Rab pools. Alternatively, Rab-GDP/GDIs bind Rab-GDP effectors that move the complex along retrograde transport routes. Although neither speculation is easy to validate, recent technological advances will aid in resolving Rab-GDP/GDI traffic. Especially the availability of endogenously tagged Rabs provides an opportunity to study and manipulate transport with unprecedented resolution (Fig. 4). Once Rab-GDPs reach the membrane, GDI-displacement factors\textsuperscript{26} or GEFs\textsuperscript{27} promote their membrane integration and conversion into GTP-bound Rabs, and the traffic cycle starts over.\textsuperscript{28}

How is Rab traffic spatially and temporally organized? A sequence of Rab proteins regulates the directed transport of specific cargo species and the Rab target-membrane specificity is controlled by GAPs and GEFs.\textsuperscript{29,30} However, how GEF/GAP specificity and regulatory activity is orchestrated in time and space remains unclear.\textsuperscript{28} Yeast Rabs can recruit GEFs that activate the next Rab in the traffic sequence, as well as GAPs that deactivate the predecessor.\textsuperscript{31,32} Similar regulatory mechanisms have also been reported for Rabs in higher eukaryotes.\textsuperscript{33,34} Other studies show that the hypervariable C-terminal domain of each Rab is instructive for insertion into specific membranes,\textsuperscript{35} and that active Rabs are able to induce self-amplifying loops by generating functional lipid micro-domains.\textsuperscript{16,36} Interestingly, on the same membrane entity multiple different Rab micro-domains can coexist.\textsuperscript{37} Therefore, specific combinations of Rabs may constitute a “Rab code,” defining the identities of intracellular transport vehicles.\textsuperscript{38}

**Rab proteins in drosophila melanogaster**

Genetic resources that allow systematic analyses of protein localization in vivo are established for baker’s yeast,\textsuperscript{39} \textit{C. elegans}\textsuperscript{40} and fruit flies.\textsuperscript{41} However, comprehensive collections of tagged \textit{rab} alleles suitable to study endogenous Rab dependent transport in a complex model organism emerged only recently for \textit{Drosophila}. In \textit{Drosophila melanogaster}, 31 members of the Rab-GTPase family were identified based on sequence comparisons.\textsuperscript{9} Many Rabs are expressed in specific patterns during embryogenesis,\textsuperscript{9} and transcription profiles indicate dynamic expression levels of many Rab transcripts throughout development and FlyBase.org. However, it is important to explore how mRNA expression patterns correlate with the localization, levels and activities of the corresponding proteins. To this aim, Dunst et al. generated a comprehensive set of endogenously modified \textit{rab} alleles (\textit{Yrabs}). The \textit{YRab}-collection comprises 27 strains; each different Rab is N-terminally tagged with YFP\textsuperscript{14} (rablibrary.mpi-cbg.de). The YFP-tag allows comparing the distributions of different Rab proteins directly and provides a common target for RNAi or nanobody-
induced protein depletion approaches. Dunst and colleagues systematically analyzed the expression, localization and abundance of Yrabs in 6 different organs, which harbor a total of 23 different cell types. Each cell type is characterized by a unique composition of the Rab machinery. For example, all cell types sampled have high levels of Rab1 mRNA and protein. In contrast, Rab10 mRNA levels are generally high but only imaginal disk cells show correspondingly high protein levels, whereas all other cell types tested have low Rab10 protein levels. Therefore, some Rabs may have cell type specific mRNA translation or protein turnover rates. In addition, the organization of Rab compartments is dependent on the state of cellular differentiation. For example, Rab3 and Rab27 regulate synaptic activity in a redundant fashion, and their expression is limited to very few cell types. However, their expression and localization differ strikingly. Rab3 is expressed during early axonogenesis and the protein is distributed throughout the cytosol (Fig. 1). Only upon initiation of synaptogenesis does Rab3 become restricted to presynaptic contact sites. In contrast, Rab27 protein is not detectable in embryonic neurons and only appears in interconnected neurons at mature synaptic contacts (rablibrary.mpi-cbg.de).

What is the biologic relevance of the intracellular compartmentalization of Rab proteins? Total Rab protein levels do not reflect the proportion of active GTP-bound Rabs, and visualizing the localization of YRabs does not allow discriminating between different Rab activity states. However, active GTP-Rabs are generally integrated into membranes, whereas inactive GDP-Rabs are GDI-bound and distributed in the cytosol. Biochemical approaches can be used to separate these Rab pools. Isopycnic density gradients can be used to sediment cytosolic Rabs. Consequently, the non-sedimenting Rab protein fraction represents GTP-Rabs engaged in membrane traffic. However, an in-depth understanding of traffic routes requires systematic mapping of the subcellular localization of all Rabs and characterizing their functional interconnections. The most common strategy is to use immunohistochemical approaches. The detection of Rab proteins with fluorescent probes often reveals prominent puncta (rablibrary.mpi-cbg.de). Notably, however, the subcellular distribution of membrane proteins in fixed samples is prone to artifacts, and only live imaging can reveal the dynamic morphologies of intracellular membrane compartments. For instance, compartments that move from the trans-Golgi network toward the plasma membrane often resemble tubular shapes, distinct from spherical vesicles. Thus, fixation and staining methods need to be carefully validated to minimize experimental artifacts. Dunst and colleagues used fixed samples and annotated systematically the localization of each Rab in many different cell types using defined and unbiased terminology for grouping all Rabs into similarity-clusters. For instance, Rab dependent ER traffic is well studied, and Rab1, 2, 10 and 18 are known to regulate traffic between ER and other organelles or maintain ER structure. In Drosophila salivary gland cells these ER-Rabs show similar distribution.

Figure 1. Rab3 expression in embryonic neurons. Rab3 and Rab27 are pan-neuronal presynaptic vesicle identifiers. In contrast to Rab27 (no detectable YRab27 in embryonic neurons, data not shown), Rab3 is expressed in a subset of wiring neurons (before synapse formation). Interestingly, after axogenesis all interconnected neurons show Rab3 and Rab27 restricted to presynaptic membranes. A-C Cartoons (adapted from Fly atlas (Hartenstein, V. (1993) Atlas of Drosophila development and flybase.org) depicting different developmental stages of Drosophila embryos. (A-C) Images from corresponding Yrab3 embryonic ventral nerve cords. Of note, YRab3 (green) is detectable in neural cell bodies (A-C, white arrowheads) and projections (A-C, white arrows). HRP (magenta) labels all neuronal membranes, developmental stages are indicated (A-C, upper left corner), scale bars indicate 20 µm.
and the authors grouped these Rabs into one annotation cluster (Fig. 2, D-D$'''$). Rab6 is linked to Golgi and post-Golgi transport.\textsuperscript{49} Although Rab6 is localized in puncta, the Rab6 annotation tree is different from the ER-Rab cluster (Fig. 2, B-B$'''$). Surprisingly, also Rab7 and 8 clustered together with ER-Rabs. Only Rab7 was found to localize at ER contact sites.\textsuperscript{50} So far, Rab8 is not reported to organize ER transport. Are Rab7 and 8 associated with ER membranes in Drosophila cells? In salivary gland cells Rab7 and Rab8 partially co-localize with Rab1, thus confirming that the annotation correctly grouped Rab7 and Rab8 with Rab1 (Fig. 2, C-C$'''$ and E-E$'''$).

The collection of tagged rab alleles in Drosophila represents a unique resource to systematically chart intracellular transport in vivo. The findings indicate a very complex interplay between protein expression, localization and activity. Thus, membrane and protein traffic pathways are likely to depend to a large extent on the particular cell types, developmental stages and physiologic conditions analyzed in a given study, suggesting that existing models of cellular traffic may in some cases be too superficial.

**Essential permanent transport and dedicated transient traffic crossroads**

Differentiation and tissue morphogenesis are interconnected processes that follow a hard-wired sequence of events. How do cells organize their protein and membrane traffic according to morphogenetic programs, and what kinds of signals control the spatial and temporal constitution of the intracellular transport machinery?

DTT pathways are induced locally, and ensure spatially and temporally targeted delivery of membrane material and cargo. However, the initiation, regulation and composition of DTTs are not well understood. Recently, Caviglia and colleagues reported new insights into one particular DTT route. They studied the
formation of tubular connections in the tracheal system in *Drosophila*, which consists of a network of epithelial tubes. The tracheal network transports oxygen to tissues. It originates from segmentally repeated invaginations of the embryonic ectoderm that extend and undergo a series of branching events. Tracheal tubes consist of an epithelial monolayer that surrounds a lumen lined by apical extracellular matrix (cuticle) that prevents tracheal tubes from collapsing.

In the tracheal system, tubular connections (anastomoses) are built between pre-existing lumenized branches. These connections are made by specialized tip-cells called fusion cells (FCs; Fig. 3). FCs migrate toward each other, and, upon contact, form a new adherens junction at the contact site. The fusion cells then repolarize and form a secondary apical domain at the cell-cell contact site. The original primary apical side of each tip cell faces the invading stalk cell lumen, where the 2 cell types are connected via adherens junctions. The stalk cell 'invades' the tip cell (resembling a finger poking into a balloon), bringing the primary FC apical membrane very close to the secondary apical membrane domain at the contact between the 2 tip cells. The subsequent intracellular fusion of the 2 apical membrane domains leads to the formation of a luminal connection between the adjacent branches (Fig. 3).

Caviglia et al. investigated the mechanism of intracellular membrane fusion in the tip cells and identified lysosome-related secretory organelles (LROs) essential for lumen fusion. LROs are pleomorphic organelles in several specialized cell types (including, among others cytotoxic T-cells, endothelial cells and pigment cells) and have cell-specific origin membranes. These compartments store specific cargo and acquire specific structural proteins/luminal environments. In response to external stimuli LROs release their cargo by fusing with the plasma membrane in a spatiotemporally, tightly regulated manner. Interestingly, early findings identified Rab geranylgeranyl transferases important for successful LRO maturation and secretion. Later, individual Rab proteins were associated with specific LRO maturation and traffic steps. However, only recently Azous et al. published a comprehensive screen aimed to identify all Rabs that regulate granule secretion in mast cells. The authors identified 30 Rabs involved in exocytosis, including Rabs that affect selectively secretion triggered by a Ca²⁺ ionophore. Among these Rabs responsible to control Ca²⁺-induced release is Rab39A. Interestingly, in tracheal fusion cells Caviglia and colleagues found Arl3-dependent recruitment of Rab39 and Munc13–4/Staccato, a Ca²⁺-binding protein, onto Rab7- and LAMP1-positive LROs. Active Rab7 and Rab39 are known to

Figure 3. Overview of membrane trafficking steps during tracheal tube fusion. (A) Scheme depicting a fusion tip cell (blue) invaded by a stalk cell (gray). The primary (1') and secondary (2') apical membranes are in magenta, adherens junctions in bright green, actin/microtubule track in dark green, ER in blue, Golgi apparatus in light green, LROs in orange, late endosomes in dark red, early endosomes in light red and recycling endosomes in yellow. Broken lines represent hypothetical trafficking steps; continuous lines represent experimentally validated steps. Endocytic compartments (a) donate membrane material to apical membranes (b) or mature to give rise to late endosomes (c). Arl3 is recruited to Rab7- and Lamp1-positive compartments, and Arl3 effectors recruit Munc13–4/Staccato and Rab39. Dynemin and kinesin (KIF) motor proteins are effectors of Rab7 and/or Rab39, and link LROs to the actin/microtubule track (d). Once the apical plasma membrane domains are reached, Ca²⁺-bound Munc13–4/Staccato promotes SNARE-dependent membrane fusion (e). Ca²⁺-enriched microdomains are established by localized release from Sec31-positive ER portions close to the fusing membranes. (B) Stages of lumen fusion in the wild type (I-III). (C) In the absence of Stac or Rab39, LROs fail to form. As a consequence the 2 apposed apical membranes fail to fuse, although fusion cells are still able to expand their apical domains. (Modified after Caviglia et al. NCB, 2016).
bind effectors, including Dynein and kinesin-3 KIF1A motor proteins, which link LROs to microtubules and can initiate transport along microtubules toward the apical plasma-membrane. In addition, the authors describe the appearance of local transient intracellular Ca\textsuperscript{2+} sparks specific to the tip cell type that mediates lumen fusion, by triggering the fusion of LROs to each apical plasma membrane domain, thus creating a connection between pre-existing lumina. Thus, in tip cells the Rab machinery may act to position the membrane fusion machinery in the right place to mediate the connection of growing lumina.

**Experimental challenges**

Intracellular transport is tightly regulated. However, the plasticity of such regulative networks is hardly assessed and comprehensive expression studies of endogenous Rabs show large spatial, temporal and quantitative differences. Therefore experimental setups need to be evaluated and standardised. Available genetic tools yielded many important findings. However, misexpression and perturbation of multiple pathways may blur results and make interpretations difficult. Therefore, here we propose novel strategies to manipulate Rab
dependent transport with a high spatio-temporal resolution.

**Closing remarks**

What are the next aims? Although some principles apply to all Rabs, such as GDP/GTP-switch regulation by GEFs and GAPs, other paradigms may not apply generally (e.g., mechanisms of Rab membrane insertion and extraction). A challenging task is to validate existing molecular and mechanistic models for all members of the Rab family. A second major challenge is to systematically chart the Rab machinery in each cell type. Many Rabs are allocated, based on published findings in different cell types, to specific organelles or traffic routes. However, recent findings show cell-type specific complements of Rab proteins, consistent with the idea of a “Rab code” that is characteristic of each cell type. Comprehensive libraries of endogenously tagged rab alleles provide a powerful tool to faithfully characterize cell type specific transport routes and Rab codes.

**Rab protein localization and loss of function rescue strategies**

Many Rabs are small proteins and their surface shows only limited immunogenic activity. Therefore, antibodies that specifically detect the intracellular localization of endogenous Rab proteins are rare. As an alternative to antibodies, expression of tagged transgenes (for instance YFP-tagged wild-type rab alleles) can be used to study Rab protein localization. However, experiments using over- or misexpression of Rab transgenes need to be interpreted with great care. First, increased Rab levels or ectopic Rab activity can lead to mutant phenotypes. Second, large amounts of inactive cytosolic transgenic protein can mask the localization of the limited pool of active Rab proteins. Genomic rescue constructs in mutant backgrounds or manipulation of endogenous rab loci by homologous recombination are powerful approaches that allow targeting the introduced tag for controlled Rab RNA and/or protein knock-down experiments.

**Rab protein depletion**

Mutagenesis, RNAi knock-down or nanobody approaches reduce the entire cellular Rab protein pool. The expression of dominant-negative alleles is another way to impede cellular Rab function. Dominant negative Rabs preferentially bind GDP and outcompete the endogenous Rab pool for the binding of regulatory elements. Therefore, the GTP-bound Rab pool is depleted and all traffic controlled by the affected Rab is jammed. However, overexpressed dominant-negative Rabs may interfere not only with the targeted Rab protein, but also with other Rabs, and therefore the effects may be to some extent ambiguous. Moreover, a given Rab can be involved in multiple traffic steps and the complete Rab depletion can result in phenotypes that may reflect the perturbation of multiple traffic routes. In addition, the plasticity of the transport machinery may be able to compensate for the affected traffic step.

**Genetic tweezers to dissect Rab traffic steps**

To manipulate individual traffic steps we propose a new approach for local deactivation of Rab proteins. Particular cellular membrane compartments and organelles are characterized by distinct Rabs, and membrane maturation/differentiation processes are characterized by a change in the set of Rabs that decorate the corresponding membranes (“Rab identity”). In consequence, for a period of time different Rabs share the very same membrane compartment and therefore come in close spatial proximity. The idea is to use that time frame and local proteolytic activity to perturb the membrane maturation sequence. Our experimental approach uses a 2-component system with one Rab protein tagged with the Tobacco-etch virus Protease (TevP, Fig. 4A) and a second Rab tagged with a TevP cleavage site (TEV, Fig. 4B) inserted between the GTPase domain and the membrane anchor. The correct genetic combination of TevP-Rab and TEV-Rab allows blocking one specific membrane traffic step without impeding Rab functionality in other trafficking steps.

For instance, flies that express TevP-Rab5 and TEV-Rab7 may show hampered endosomal maturation while Lysosome-ER transport, which depends on Rab9 and Rab7, should remain unperturbed (Fig. 4C-E). Systematically probing the functional relationships of Rabs using TevP-Rab/TEV-Rab combinations may prove useful to chart the sequence of Rabs required to shuttle particular protein cargo. In an analogous approach, a protein of interest is tagged with TevP (TevP-cargo) and its intracellular transport can be mapped using TEV-Rabs.

**Disclosure of potential conflicts of interest**

No potential conflicts of interest were disclosed.

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