The Ras family G-proteins RalA and RalB make critical non-overlapping contributions to the generation of a tumorigenic regulatory network, supporting bypass of the normal restraints on both cell proliferation and survival. The Sec6/8 complex, or exocyst, has emerged as a principal direct effector complex for Ral GTPases. Here, we show that RalA and RalB support mitotic progression through mobilization of the exocyst for two spatially and kinetically distinct steps of cytokinesis. RalA is required to tether the exocyst to the cytokinetic furrow in early cytokinesis. RalB is then required for recruitment of the exocyst to the midbody of this bridge to drive abscission and completion of cytokinesis. The collaborative action of RalA and RalB is specified by discrete subcellular compartmentalization and unique pairs of RalGEF proteins that provide inputs from both Ras-family protein-dependent and protein-independent regulatory cues. This suggests that Ral GTPases integrate diverse upstream signals to choreograph multiple roles for the exocyst in mitotic progression.

Introduction

The Ral GTPases belong to the Ras branch of the superfamily of monomeric GTPases (Valencia et al., 1991). Interest in Ral function increased following the discovery that Ral proteins acted as proximal effectors of oncogenic Ras, that their activity is essential to support tumorigenesis of human epithelial cells of diverse origin and can be even sufficient in some genetic backgrounds to drive oncogenic transformation (Hamad et al., 2002; Chien and White, 2003; Rangarajan et al., 2004; Sablina et al., 2007).

The two Ral genes in mammals, RalA and RalB, encode highly homologous proteins (81% identity, 89% similarity), which make distinct and sometimes collaborative contributions to diverse cellular functions. RalA drives exocyst assembly and polarized exocytosis in epithelial cells (Moskalenko et al., 2002; Shipitsin and Feig, 2004), whereas RalB drives exocyst assembly during polarized cell migration (Rosse et al., 2006). RalA and B cooperate to support oncogenesis, as RalA is required for anchorage-independent proliferation and tumorigenesis, and RalB for metastasis formation and cell survival (Chien et al., 2003; Chien et al., 2006; Lim et al., 2006).

In contrast to the diversity of their biological functions (Feig, 2003; Camonis and White, 2005), Ral GTPases exert an effect through a limited number of well-documented Ral-GTP functional effectors, RalBP1/RLIP76 and the exocyst complex, being the best documented. RalBP1/RLIP76 participates in the regulation of a subset of endocytic pathways (Nakashima et al., 1999; Jullien-Flores et al., 2000; Rosse et al., 2003). The exocyst complex, in addition to its structural role in vesicle–plasma membrane tethering, functions as a signalling complex through direct interactions with effector kinases (Moskalenko et al., 2002; Feig, 2003; Camonis and White, 2005; Balakireva et al., 2006; Chien et al., 2006).

The exocyst complex is required for appropriate completion of cytokinesis (Wang et al., 2002; Dobbelaere and Barral, 2004; Gromley et al., 2005; VerPlank and Li, 2005). The correct course of cytokinesis, the ultimate step of cell division, is essential for high-fidelity chromosomal segregation. In metazoans, cytokinesis can be divided into two steps. Actino-myosin ring formation and cleavage furrow ingestion generate a cytoplasmic bridge, whose abscission requires mobilization of secretory vesicles and exocytosis (Echard et al., 2004; Eggert et al., 2004; Schweitzer and D’Souza-Schorey, 2004; Skop et al., 2004; Albertson et al., 2005; Glotzer, 2005; Gromley et al., 2005; Schweitzer et al., 2005). In metazoans, the exocyst complex is required for late cytokinesis and is recruited to the midbody for the resolution of the intracellular bridge (Gromley et al., 2005).

Here, we have examined the role of Ral GTPases, their effectors and their activators, the RalGEFs, during cytokinesis. Although RalA has been shown to be required for the completion of cytokinesis (Chen et al., 2006), we found that RalA and RalB make collaborative exocyst-dependent contributions to cytokinesis that are distinct in space and time. The non-overlapping roles of RalA and RalB are specified by unique pairs of RalGEF proteins that provide inputs from both Ras-family protein-dependent and protein-independent regulatory cues, suggesting that each Ral GTPase functions as a signalling integration node from several transduction pathways. In this model, the concomitant correct functioning of these pathways would be required to permit progression of cytokinesis beyond RalA- and RalB-controlled specific steps.
Results

**RaIA and RaIB make distinct contributions to cytokinesis**

Mitotic failure as a consequence of cytokinesis defects can manifest as the fusion of nascent ‘daughter cells’ leading to binucleate cells or in prolonged daughter cell attachments as a consequence of delayed or failed abscission (Echard et al., 2004; Fielding et al., 2005; Gromley et al., 2005). To examine the explicit contributions of RaIA and RaIB to cytokinesis, we monitored mitotic progression upon siRNA-mediated depletion. Real-time imaging of control cells demonstrated complete cleavage furrow ingression 5 min post-telophase initiation and the appearance of intracellular bridges around 10–20 min post-telophase initiation. Cytokinesis appeared complete by approximately 5 h after telophase start (Figure 1A, upper panel, and Supplementary Movie S1). By contrast, depletion of RaIA resulted in cytokinesis regression, whereas depletion of RaIB resulted in abscission failure. RaIA-depleted cells (Figure 1A, middle panel, and Supplementary Movie S2) often did not spread properly, and in 16% of cells entering mitosis, furrows regressed, resulting in binucleate cells (Figure 1A). In siRaIA-treated cells (Figure 1A, lower panel, and Supplementary Movie S3), the intracellular bridge appeared correctly (00:20, arrow), but approximately 20% of the cells failed to complete abscission.

The RaIA- and RaIB-dependent cytokinesis phenotypes were quantified in fixed samples (Figure 1B). Depleting RaIA significantly increased the percentage of binucleate cells and not intracellular bridges, whereas depleting RaIB had no effect on the percentage of binucleate cells but strongly induced accumulation of bridged cells. When RaIA and RaIB were depleted together, the RaIA phenotype dominated (Figure 1B). These results were reproduced in a different cell line, the telomerase-immortalized human bronchial epithelial cell line (Chien et al., 2006) (Supplementary Figure S4A).

We confirmed the specificity of the RaIA and RaIB phenotypes with a second independent set of specific siRNAs and with rescue experiments (Figure 1C). siRaIA-treated cells were transfected with plasmids expressing Myc-RaIA made resistant to the siRaIA by three conservative point mutations, Myc-RaIB or control plasmid. siRaIA-treated cells transfected with empty plasmid displayed the expected RaIA-dependent phenotype. Expressing Myc–RaIA suppressed the phenotype, whereas expressing Myc–RaIB did not (Figure 1C). The same approach was applied to siRaIB-treated cells using an siRaIB targeted to the 3′-UTR. siRaIB-transfected cells displayed the ‘bridge’ phenotype (Figure 1C), which was suppressed by expression of Myc–RaIB and not of Myc–RaIA. These data demonstrate that the binucleate phenotype is due to the specific depletion of RaIA and that the bridge phenotype is due to the specific depletion of RaIB. Thus, RaIA participates in the progress of cytokinesis beyond a point where fusion is no longer possible, whereas RaIB contributes to the termination of cytokinesis.

To examine the consequences of RaIB gain of function, HeLa cells overexpressing dominant active mutants of RaIA (RaIA(D)) or RaIB (RaIB(D)) was followed by time-lapse videomicroscopy and cells entering mitosis were tracked. Expressions of RaIA(D) and RaIB(D) caused cytokinesis failure: 15% of transfected cells that entered mitosis resulted in binucleate cells (data not shown). Statistical analysis of the accumulation of binucleate cells after 48 h of transfection but in fixed samples confirmed that RaIA(D)- or RaIB(D)-expressing cells significantly accumulated binucleate cells (Supplementary Figure S4B and C). The RaIA binding domain (RaIBD) of RaIB interacts with the effector loop of both activated RaIA and RaIB and blocks Ra downstream signaling by sequestering active Ra proteins away from their endogenous effectors (Wolthuis et al., 1998; Moskalenko et al., 2002). RaIBD expression induced accumulation of binucleate cells (Supplementary Figure S4B), suggesting that Ra GTPases participate in cytokinesis through the activation of one or several effectors.

RaIB variants selectively uncoupled from downstream effectors were employed to examine the G-protein/effector relationships required for Ra-GTP to engage cytokinesis machinery. D49N uncouples Ra from RLI76/RaIBP1 and D49E from both Exo84 and Sec5 as described previously (Jullien-Flores et al., 2000; Moskalenko et al., 2002, 2003). To generate variants selectively uncoupled from Exo84 versus Sec5, we examined the consequences of point mutations defined previously as making selective contributions to Ra-GTP/Exo84 and Ra-GTP/Sec5 complex formation in vitro (Jin et al., 2005). As shown in Supplementary Figure S4D, A48W and E38R dramatically and specifically decrease affinity of RaIB G23V for Exo84 and Sec5, respectively. Similar results were obtained for RaA G23V (not shown). We also employed a deletion of the NH2-terminal 11 amino acids (A11) of Ra, which are required for association with active Phospholipase D (PLD) (Jiang et al., 1995). RaA(D) and RaB(D) uncoupled from RLI76/RaIBP1 or PLD still induced a binucleate phenotype. In contrast, impairing interaction with any exocyst component abolished the capacity of RaIA(D) and RaIB(D) to impair cytokinesis (Supplementary Figure S4B and D).

**RaIA and RaIB use common and distinct effectors to progress cytokinesis**

The above-mentioned data show that constitutively activated Ra impacts cytokinesis through its effector, the exocyst complex. Previous observations from our group and others strongly suggest that overexpression of constitutively active Ra GTPases obscures selectivity of biological control observed with the endogenous proteins. To circumvent this problem in the context of complementation experiments, we employed effector-selective Ra variants that are otherwise wild type rather than dominant active. Figure 2 shows that RaA uncoupled from RLI76 (D49N) can rescue cytokinesis upon depletion of native RaA, whereas variants uncoupled from PLD, Exo84 or Sec5 (A48W and E38R, respectively) cannot. In contrast, only RaIB variants uncoupled from Exo84 and PLD fail to support abscission. All alleles were expressed at similar levels (Supplementary Figure S4E). The RaIB variant uncoupled from both Sec5 and Exo84 (D49E) failed to support abscission (data not shown). These results show that RaA and RaIB share Exo84 and PLD1 as effectors for cytokinesis, and Sec5 is a specific effector of RaA. They do not imply that no other partner of Ra has an important function in cytokinesis.
Figure 1 Depletion of RalA induces accumulation of binucleate cells, whereas depletion of RalB induces accumulation of cells with an intracellular bridge. (A) Selected phase-contrast time-lapse videomicroscopy frames of representative cytokinesis steps from HeLa cells transfected with control siRNA (upper panel) or siRNA targeting RalA (central panel) or RalB (lower panel). In control cells, the intracellular bridge appeared 20 min after the beginning of the process (arrows). In RalA-depleted cells, the intracellular bridge did not appear and cells fused back giving binucleate cells. In RalB-depleted cells, the intracellular bridge formed timely and stayed for an extended period without abscission. In some cases, one of the daughter cells re-entered mitosis still bound to its sister cell (Figure 1A, arrow in last image of third panel). For every video, 50 cells entering mitosis were tracked. In RalA-depleted populations, we counted the frequency of binucleate cells (lower left graph), and in RalB-depleted population, the frequency of cells failing to resolve their intracellular bridge at the end of the video (lower right graph). (B) HeLa cells were depleted of RalA or RalB or both by siRNA (10 nM). Cells were fixed after 72 h and stained (lower panels) with a nuclear marker (DAPI) (blue) and an anti-tubulin FITC mAb to mark the intracellular bridge (green). Binucleate cells (arrows in the left picture) or cells with an intracellular bridge (arrows in the right picture) were counted. In the graph, the percentages of binucleate or bridged cells are shown, with an indicated statistical significance (**P<0.01). (C) HeLa cells were depleted of RalA or RalB by a second independent siRNA, and transfected with plasmids expressing Myc–RalA or Myc–RalB, or an empty vector. Myc–RalA was made insensitive to siRalA by making silent mutations in three nucleotides within the siRNA target sequence. Myc–RalB was not mutagenized, as the siRalB target sequence is within the 3'-UTR and as pRK5–Myc–RalB carries only RalB coding region. Binucleate cells or cells with an intracellular bridge were counted as in (B). In the graphs, the percentage of binucleate or bridged cells are shown, and statistical significance of the difference with control siRNA (**P<0.01) is indicated, as is the case for rescue compared with siRal (P<0.01). In all experiments, at least 1000 cells were counted, and results are from at least three independent experiments. Efficiency and specificity of depletion of RalA and RalB by the siRNAs were tested by western blots (lower panel).
Figure 2. RalA and RalB use common and different effectors. HeLa cells were transfected with siRNA for RalA or RalB, and plasmids expressing Ral proteins resistant to these siRNA (see legend of Figure 1C). These variants carry a mutation in the effector loop as indicated in the figure. A11 refers to a deletion of the first 11 amino acids of Ral. Binucleate (A) and bridged (B) cells were counted as described in Figure 1C. Percentage of binucleate and bridged cells are indicated on top of each bar. In ordinate are indicated the percentages of rescue by each allele calculated versus the siRalA or siRalB phenotypes. Statistical significance of differences is indicated.

**RalA is localized to the cleavage furrow, whereas RalB is at the midbody**

Overexpressed RalA has been reported to localize to the cleavage furrow region (Chen et al., 2006). We have examined the localization of Ral proteins in early and late cytokinesis in synchronized cells by immunofluorescence. Mitotic-kinesin like protein-1 (MKLP1) was used as a marker of the midbody ring in early and late stages of cytokinesis (Gromley et al., 2005). The early midbody ring appears during acto-myosin ring constriction, which occurs 8 h post-release from G1-M cell-cycle blockade. The late midbody ring encircles the midbody per se, which appears as a dense structure in phase-contrast microscopy in elongated intracellular bridges 10 h post-release.

We found that, in early cytokinesis, RalA was concentrated at the cleavage furrow region surrounding MKLP1 (Figure 3). Globally, RalA localized mostly to the plasma membrane delimited by the cortical cytoskeleton revealed by β-spectrin (Supplementary Figure S5A). This accumulation was not a ‘passive’ effect from membrane apposition, as RalA was not enriched at the double membrane of two just mitotic adjacent cells (arrowheads in Supplementary Figure S5A). The staining of RalA was specific as verified by loss of signal in RalA-depleted cells (Supplementary Figure S5B). This localization is in contrast to that reported previously employing overexpression of green fluorescent protein (GFP)–RalA fusions (Chen et al., 2006). We did not find any discrete localization of endogenous RalA during the late stage of cytokinesis (Figure 3).

Given the absence of appropriate antibodies to detect endogenous RalB, we had to turn to Myc- and Flag-fusions to localize RalB. Results are shown for Flag–RalB. During early cytokinesis, we could not detect any discrete localization of RalB. Later in cytokinesis, Flag–RalB was enriched at the midbody (Figure 3). This is not a Flag-dependent effect, as Myc–RalB displayed the same localization as Flag–RalB, nor a tag-fusion effect, as both Myc–RalA and Flag–RalA were enriched at the cleavage furrow (data not shown), displaying the same localization as endogenous RalA at an early stage, and were never detected at the midbody. We could not analyse the localization of RalA and tagged-RalB in the same cells due to different fixation protocols.

**Exocyst components are already mobilized at early cytokinesis**

Sec5 together with other components of the exocyst complex is localized at the midbody ring in late cytokinesis (Gromley et al., 2005), which is what we observed here (Figure 4A and B). As Sec5 is an effector of RalA in cytokinesis, we asked whether Sec5 was specifically localized in early cytokinesis. We detected a discrete localization of Sec5 encircling the microtubule bundles as soon as furrow ingestion terminated, suggesting that the exocyst complex is already engaged at this early step (Figure 4A). Three-dimensional reconstruction of deconvolution microscope images revealed that Sec5 was present in a ring-like structure, which is presumably the early midbody ring, which starts to organize at this step (Figure 4A, lower panel). This signal was lost in Sec5-depleted cells (Supplementary Figure S5B).

In PC12 cells, Sec5 and Sec6 are present in the same heavy plasma membrane fraction (Moskalenko et al., 2003). We examined whether they localize similarly at the early midbody ring. Whereas Sec5 was localized at the early midbody ring, Sec6 could only be detected at the Golgi apparatus during its post-mitotic reassembly (data not shown). However, following furrow ingestion termination, Sec6 displayed a discrete pattern of accumulation (arrowheads in Figure 4C, left panel), which appeared to be on both sides of the early midbody ring and the midbody as indicated by MKLP1 staining (Figure 4D, upper panel) and phase-contrast images, respectively (arrowheads in Figure 4C, right panel; arrow to the midbody). Specificity of Sec6 mAb labelling was validated by Sec6 depletion (Supplementary Figure S5B).

**Ral GTPases control Sec6 but not Sec5 localization during cytokinesis**

As RalA participates in early cytokinesis events and is localized to the cleavage furrow (Figure 3), we examined if the early Sec5 and Sec6 recruitment were dependent on RalA. As shown in Figure 4B, recruitment of Sec5 to the early midbody ring was not perturbed by siRNA-mediated depletion of RalA. In contrast, the early recruitment of Sec6 was blocked by RalA depletion; perturbing RalB had no effect (Figure 4D). RalB appears to make an obligate contribution to abscission. RNAi-mediated depletion of RalB induced an accumulation of elongated bridges (Figure 2) with normal Sec5 localization (Figure 4B); however, Sec6 was absent (Figure 4D). We observed an increase of endobrevin/
VAMP8 staining (Figure 4D, lower panels). Elongated bridges of the RalB-depleted but not control cells overexpressing GFP-endobrevin accumulated GFP flanking the midbody, but we were unable to discriminate single vesicles (Figure 4D, lower panels).

The loss of Sec6 recruitment to cytokinesis machinery in RalA- and RalB-depleted cells was not due to a general defect of central spindle organization, as bridge architecture was globally unaffected (Figure 4E). However, when the intracellular diameter channels were measured in the various populations, we found that the average sizes of the diameter were different according to the presence of Ral. When RalA was depleted, the intracellular diameters were on average wider, and when RalB was depleted, they were thinner, as compared with cells with full complements of RalA and RalB (Figure 4F). We did not find any defect in microtubule morphology (data not shown).

**RalA and Sec5 are both required for recruitment of exocyst components at early stages of cytokinesis**

RalB drives late cytokinesis events as has been described for the exocyst (Gromley et al., 2005), but our observations indicate that Sec5 and RalA are already mobilized at early steps, and they colocalize at the early midbody ring during furrow ingression (Figure 5B and Supplementary Movie S6). Depletion of Sec5 induced cytokinesis defects (Gromley et al., 2005) and accumulation of binucleate cells similar to the depletion of RalA (Supplementary Figure S7). To further investigate the role of RalA in early cytokinesis, we depleted cells of RalA, Sec5 or Sec6 and tracked the consequences on the localization of exocyst components in early cytokinesis. Sec6 depletion led to the same cytokinesis defects as those caused by Sec5 or RalA depletion (Supplementary Figure S7) but did not perturb Sec5 or RalA localization (Figure 5A and data not shown, respectively). In contrast, Sec5 depletion blocked recruitment of Sec6, whereas RalA was unaffected (Figure 5A). Sec10 is a component of the exocyst present on vesicles and assembles with the core Sec6/8 exocyst complex in a Ral-dependent manner (Moskalenko et al., 2002; Rosse et al., 2006). During cytokinesis, Sec10 localized similarly to Sec5 at the midbody ring, and this localization was lost when RalA, Sec5 or Sec6 were depleted (Figure 5A, lower row, and graph on the right). Thus, both Sec5 and RalA constitute a spatial landmark for further early localization or stabilization of exocyst holocomplexes. These results also show that in addition to its role in abscission, Sec5 has a previous role in early cytokinesis.

**Sec5 and RalA interact functionally and colocalize at the midbody ring**

A RalA mutant (E38R) unable to interact with Sec5 is unable to support cytokinesis (Figure 2), and RalA and Sec5 colocalize at the early midbody ring (Figure 5B). To confirm that a physical interaction between RalA and Sec5 is required to mediate RalA-dependent progression of cytokinesis, we employed a Sec5 variant (T11A) defective for RalA binding (Fukai et al., 2003). As shown in Figure 5C, both wild-type and mutant Sec5 correctly localized to the midbody ring...
during cytokinesis, confirming that the localization of Sec5 is independent of RalA. RNAi-mediated depletion of endogenous Sec5 resulted in cytokinesis failures similar to those observed upon depletion of RalA (Supplementary Figure S7). These phenotypes were rescued upon expression of siRNA-resistant wild-type Sec5 but not Sec5T11A (Figure 5C, right panel). These observations indicate that a physical interaction between RalA and Sec5 is required for progression of cytokinesis.

Specific Ras-dependent and Ras-independent RalGEFs are involved at different steps of cytokinesis

There are six RalGEFs encoded by the human genome, which can mediate Ral-GTP loading. Four of them (RalGDS, RGL, RGL2 and RGL3) are themselves activated by the Ras GTPases and two of them (RalGPS1 and 2) activate Ral in a Ras-independent manner (Rebhun et al., 2000; Ceriani et al., 2007).

We individually depleted all RalGEFs by siRNAs, as validated by qRT-PCR (Supplementary Figure S8). Remarkably,
we found that depletion of RalGDS or RalGPS2 caused the accumulation of binucleate cells similarly to the depletion of RalA (Figure 6A). In contrast, the depletion of RGL or RalGPS2 caused the accumulation of cells connected by an intracellular bridge, similarly to the depletion of RalB (Figure 6A). These phenotypes were confirmed with an independent siRNA for each of these genes (Supplementary Figure S8). We never observed both phenotypes upon depletion of any RalGEF. The depletion of RGL2 or RGL3 had no significant effect despite effective depletion (Figure 6A and Supplementary Figure S8). These results suggest that some RalGEFs, during cytokinesis, display specificity towards RalA or RalB.

We used time-lapse videomicroscopy to evaluate the stage of cytokinesis impacted by these RalGEFs. In cells treated with siRNA-targeting RalGDS or RalGPS2, furrow ingression took place but cells failed to spread and form intracellular bridges, and daughter cells ultimately fused and formed binucleate cells. Cells treated with siRNA-targeting RGL or RalGPS1 were defective in abscission and maintained intracellular bridges for extended periods (Figure 6B). The depletion of RalGDS or RalGPS2 impacted cytokinesis dynamics very similarly to what was observed upon depletion of RalA, and the depletion of RGL or RalGPS1 impacted cytokinesis dynamics very similarly to the depletion of RalB: these data suggest that RalGDS and RalGPS2 might have RalA as substrate during cytokinesis, and RGL and RalGPS1 would have RalB. Notably, one member of each pair of RalGEF is a Ras-responsive RalGEF, whereas the other member is not and contains a PH domain that may associate with discrete membrane microdomains and transduce signals whose nature remains elusive.

**RalGEFs localize differently**

RalGEFs are able to induce the activation of RalA and RalB (Albright et al., 1993), but their in vivo specificity for RalA or RalB under physiological conditions is unknown. In interphase cells, RalA and RalB have been reported to have overlapping but not identical localizations: RalA was found more at the plasma membrane and RalB was found more on endomembranes (Shipitsin and Feig, 2004). We also found distinct and exclusive localizations of RalA and B during cytokinesis.

We questioned whether RalGEFs might also display specific intracellular localizations, which might be the basis of a preferential activation of RalA or B. Owing to the lack of antibodies to localize endogenous proteins, we used tagged versions of the RalGEFs. In HeLa cells, Flag–RalGDS and Myc–RalGPS2 localized to the plasma membrane (Supplementary Figure S8), which is enriched in RalA, and Flag–RGL and Flag–RalGPS1 displayed a punctuate localization through the cytoplasm (Supplementary Figure S8), similar to RalB (Shipitsin and Feig, 2004). We examined the localization of these RalGEFs during cytokinesis in synchronized cells. We found that Flag–RalGDS localized to the early midbody ring (Figure 7), where RalA and Sec5 colocalized (Figure 5), and persisted there till maturation of the midbody (Figure 7). Myc–RalGPS2 was found, also like RalA, on the ingressed cleavage furrow membrane, and was lost later in cytokinesis (Figure 7). Upon depletion of RalGDS or RalGPS2, Sec6 failed to accumulate at the midbody ring (Figure 8, upper panels) as was observed upon RalA depletion. Similar to RalB, Flag–RGL and Flag–RalGPS1 did not display any discrete localization at the early stages of cytokinesis but were enriched on the midbody during late-stage cytokinesis (Figure 7). When RGL or RalGPS1 were depleted, Sec6 was still localized normally (Figure 8).

**Discussion**

Ral GTPases RalA and RalB are effectors of the Ras GT Pases, and Ral in turn signals through a small set of effectors, the best documented being the exocyst complex (Feig, 2003; Camonis and White, 2005). The latter is involved in exocytosis and is required for the abscission of the bridge joining the sister cells emerging from mitosis (Echard et al., 2004; Gromley et al., 2005). Here, we show that the multistep process of cytokinesis integrates the contribution of both

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**Figure 4** RalA and RalB regulate Sec6 localizations. (A) Synchronized HeLa cells were fixed at early (upper left panel) and late (upper right panel) cytokinesis as shown in Figure 3 and stained with anti-Sec5 Ab (green) and co-stained with anti-tubulin mAb (upper row). Sec5 (arrows) can be observed in late as well as early cytokinesis on the midbody ring. Cells stained with anti-Sec5 Ab (green) were imaged with a deconvolution microscope. Images show a 180° 3D reconstruction: the staining of Sec5 is in a ring-like structure (arrow). (B) Synchronized HeLa cells transfected with siRalA, siRalB or control siRNA were stained with anti-Sec5 mAb (green). Early (8 h after release of the thymidine block) and late (10 h after release of the thymidine block) localisations of Sec5 did not appear sensitive to a depletion of RalA or B.

**Figure 5** RalA and RalB have distinct and exclusive localizations during cytokinesis. (A) Synchronized HeLa cells were stained with anti-Sec6 Ab at early (left panel) and late (right panel) cytokinesis as shown in Figure 3. Sec6 (arrow heads) accumulated on two discrete regions in early cytokinesis and on both sides of the midbody in late cytokinesis on elongated bridges. On these phase-contrast microscopic pictures, a dense midbody can be seen (arrow) at late times (right panel) and not at early times (left panel). (B) Synchronized HeLa cells were transfected with an siRalB or control siRNA, and stained for Sec6 (green) and MKLP1 (red). The early localization of Sec6 on both sides of the early midbody ring marked by the anti-MKLP1 disappeared in RalA- but not in RalB-depleted cells. The late localization of Sec6 was lost in RalB-depleted cells. Graphs on the right give a quantification of the rate of Sec6-positive cells in the various situations. Data are from at least three independent experiments where at least 40 cells have been counted. Elongated bridges of RalB-depleted cells stained for endoA3, which marks exocytotic vesicle, displayed an accumulation compared with the control (low row) as was the case in cells where the exocyst was disrupted (Gromley et al., 2005). Elongated bridges of the RalB-depleted cells overexpressing GFP–endoA3 (Martineau et al., 2008) accumulated GFP (n = 40). The physiological late intracellular bridges of the siA3-treated cells overexpressing GFP–endoA3 did not show any accumulation of the GFP (n = 10).

(E, F) The absence of RalA and RalB does not impact formation of the spindle per se but modifies bridges. The organization of the central spindle and of the intracellular bridge was controlled by co-staining synchronized cells with an anti-AuroraB mAb (green) and an anti-MKLP1 (red). (F) In controls, RalA-depleted cells and RalB-depleted cells, AuroraB concentrated in the region of the bundled microtubules on both sides of the midbody ring stained by MKLP1, showing that cytokinetic spindle was correctly formed (E). Cells were stained with FITC-coupled phospholipids, which stains plasma membranes. This allowed to measure the diameter of the intracellular channel (as in the given example) in cells transfected with siA3, siB3 and a control siRNA. We defined five categories (0–1, 1–2, 2–3, 3–4 and 4–5 μm) and distributed the number of diameters per category. The distribution is shown on the graph, which reveals that the intracellular diameters are rather wider in RalA-depleted cells and are rather thinner in RalB-depleted cells, as opposed to the distribution in control cells.
Ral GTPase family members to distinct spatial and temporal mobilization of exocyst components and that this integration is specified by distinct RalGEFs.

We find that the exocyst is not only required for abscission, but it is also already engaged upon the formation of the cleavage furrow. Second, we find that RalA and RalB drive distinct tasks: RalA supports the stabilization and elongation of the intracellular bridge, whereas RalB supports abscission. RalA cooperates with Sec5 to allow tethering of other components of the exocyst to the early midbody ring; upon completion of bridge formation, RalB takes over to mediate exocyst tethering and exocytosis for abscission. The specificity of engagement of the RalA-exocyst and RalB-exocyst modules appears to be driven by different Ral activators; RalGDS and RalGPS2 for RalA, RGL and RalGPS1 for RalB. The diversity of regulatory inputs made available through these GEF family members suggests that RalA and RalB may integrate multiple signals to allow transition through distinct phases of cytokinesis and function as signal integration nodes.

The exocyst complex is required for abscission, which involves endobrevin/VAMP8 secretory vesicle tethering to the midbody (Gromley et al., 2005). We found that Sec5 is already recruited at the early midbody ring during actomyosin-ring constriction where it colocalizes with RalA. At this step, Sec6 is at the Golgi apparatus during its post-mitotic reassembly (data not shown) but soon displays a discrete pattern on both sides of the early midbody ring, a region of active exocytic and endocytic processes (Albertson et al., 2005). This exocyst localization might correspond to the symmetric flow of exocytosis that has been documented recently as taking place during cytokinesis (Goss and Toomre, 2008), which might be under the control of RalA. Although Sec5 or Sec6 have a late role in cytokinesis (Gromley et al., 2005), we have also observed earlier failures upon Sec5 and Sec6 depletions, which led to binucleate cells. This suggests that at least some exocyst components have timely distinct functions during cytokinesis. During cytokinesis, Sec5 localization appears to orchestrate recruitment of the other subcomplexes. As opposed to its ‘late’ function, this...
early function of Sec5 is not dependent on centriolin. Upon centriolin depletion by siRNA, we observed a ‘bridge phenotype’ reflecting the defect in abscission but not in furrow formation (Supplementary Figure S5D), and Sec5 was properly recruited to the early midbody ring, although it was lost in the midbody ring of matured intracellular bridges as published (Supplementary Figure S5D; Gromley et al., 2005). Sec5 localization at the midbody ring was Ral independent. Similarly, centriolin localization was not modified by depletion of RalA or B (Supplementary Figure S5C). However, the Sec5–RalA interaction was required for cytokinesis to proceed, demonstrating that RalA does not control Sec5 localization but does control its function. Hence, RalA and Sec5 cooperate to mediate and control exocyst assembly as soon as the early midbody ring is formed.

In RalA-perturbed cells, the cleavage furrow and the central spindle organization progresses normally, but the intracellular diameter of daughter cells after cleavage furrow ingression is wider, suggesting that stability and elongation, rather than formation of the intracellular bridge, was affected. Control cells show that the maturation of the intracellular bridge and cell spreading are concomitant, and both events were impaired in RalA-perturbed cells. A role of RalA could be to tether vesicles carrying a Rab11/FIP3-4/exocyst subcomplex to Sec5-carrying receptor membranes. RalA would assemble the exocyst complex by interacting with...
Exo84 and Sec5 (Moskalenko et al., 2003), allowing tethering of these vesicles before fusion. Our data with RalA are consistent with the binucleate phenotype observed upon depletion of Rab11, FIP3 and ARF6 together with Rab11 (Wilson et al., 2005; Yu et al., 2007). We also found that both RalA and RalB may need to interact with PLD1 to support cytokinesis. This interaction might be an integrating node for signals coming from other GTPases, like ARF6. ARF6 and Ral are able to coactivate PLD1 (Xu et al., 2003), and on the other hand, ARF6 is involved in cytokinesis, where it supports correct localization of exocyst subunits (Fielding et al., 2005). Bridge stability requires signals carried by the trafficking machinery (Echard et al., 2004): complementary or alternatively to its function as a ‘membrane supplier’, the exocyst complex could also be in charge of targeting signalling molecules during cytokinesis, as observed for cell survival and innate immunity (Balakireva et al., 2006; Chien et al., 2006).

Similarly, the participation of RalB in abscission further highlights functional coupling of Ral GTPases and the exocyst, and RalB appears as a required conductor of the exocyst-dependent membrane dynamics that drive abscission. However, the process is complex, with RalB potentially engaging an Exo84-dependent but Sec5-independent activity. The fact that, in RalB-depleted cells, endobrevin-carrying vesicles display a pattern similar to the one observed when exocyst function was impaired (Gromley et al., 2005) suggests that RalB and the exocyst complex are involved in exocytosis of the same pool of vesicles. However, this requires more investigation, as depleting RGL1 and RalB, although leading to the same failure in abscission and accumulation of endo-

Figure 7 Different RalGEFs localize in different cellular compartments. Synchronized HeLa cells were transfected with plasmids expressing Flag-fusions of RalGDS, RGL, RalGPS1 or Myc-fusion of RalGPS2. Cells were fixed at different times as shown in Figure 3. RalGEFs were detected by indirect immunofluorescence using anti-Flag mAb (green), co-stained with anti-MKLP1 Ab (red) and DAPI (blue).
brevin-positive vesicles near the midbody, did not have the same impact on Sec6: in the former depletion, Sec6 was correctly localized, in the latter it was not. This suggests that RalB even under its GDP bound form might have a function which may be related to PLD activation (Xu et al., 2003).

A role of RalA in abscission and its localization at the midbody has been reported (Chen et al., 2006), which does not fit with our observations. We do not know the source of this discrepancy. The localization of RalA at the midbody might be due to the use of a GFP-fused RalA, and GFP fusions in fixed cells can get mislocalized (Schmitz and Bereiter-Hahn, 2001). Our data are supported by tracking endogenous RalA as well as Myc-tagged and Flag-tagged RalA. As for the role in abscission, our data are supported by the use of several different siRNA as well as by siRNA-complementation experiments.

Ral activation of effector complexes requires GTP loading through the action of RalGEFS. We found that depletion of RalGDS or RalGPS2 phenocopied depletion of RalA, whereas depletion of RGL and RalGPS1 phenocopied depletion of RalB. This suggests pathways where RalA activation requires both Ras-dependent (RalGDS) and Ras-independent (RalGPS2) GEFs, and RalB activation requires a distinct Ras-dependent (RGL) and Ras-independent (RalGPS1) pair. RalGTPases could therefore integrate information from two pathways, which must be active to mobilize the exocyst for both progression and termination of cytokinesis. The required concomitant actions of two GEFs on one GTPase for functional activation is reminiscent of the participation of RhoA in cytokinesis, which also requires two GEFs to function properly: one, Ect2, localizes RhoA in the vicinity of the other, GEF-H1, in charge of the activating GDP/GTP exchange per se (Birkenfeld et al., 2007). The fact that the RalGEFs are responsible for timely engagement of RalA or RalB during cytokinesis might explain the observation that activated mutants of RalA and RalB display nearly the same phenotype: in these mutants, RalGEFs are no longer required and thus activated RalA and RalB would be similarly activated, without the time separation provided by their specific GEFs.

How the specificity for RalA and RalB is achieved by RalGEFs is unknown. Although it might be an intrinsic in vivo biochemical property of each RalGEF, it is possibly (and more likely) the consequence of differential localizations, and indeed we show that RalGEFs can have specific localizations. Taking into account the recently described capacity of Ras to impact cytokinesis during tumorigenesis (Thullberg et al., 2007), an unified theory would be that a RalGPS, perhaps through its interaction with specific membrane domains, localizes its partnering Ras in the proximity of its activating Ras-dependent RalGEF.

Materials and methods

Antibodies, plasmids and siRNA oligonucleotides
RalA, RalB, Myc–RalB, RalBDA (RalBG23V), RalBDA/D49N, RalBDA/D49E, RalBDA/A48E and RalBAN11 (deleted of their 11 first codons) (Jullien-Flores et al., 1995, 2000; Bauer et al., 1999; Moskalenko et al., 2002), and RalADA (RalAQ72L), RalADA/D49N, RalADA/D49E and AN1RalA (Lalli and Hall, 2005) were expressed using pRK5. RalA G23V/E38R, RalA G23V/A48W, RalB G23V/E38R and RalB G23V/A48W were expressed using pCMV10–3FLAG as their controls RalA G23V and RalB G23V. Wild-type RalA and RalB, and RalA and RalB carrying D49N, D49E, A48W and E38R in an otherwise wild-type background were expressed from the pRK5–Myc or 3XFlag–CMV10 plasmids. Myc–Sec5wt and Myc–Sec5T11A were expressed in pCMV.

siRai_I resistant RalA was obtained by PCR mutagenesis using oligonucleotides GAAGATAAAAGGcAtAGTGTAGAAGAGGCA with the shorter primer and GAAAGATAAAAGGCaaGtaAGTGTAGAAGAGGCA with the longer primer.

Figure 8 Impact of RalGEFs on Sec6 localization. Synchronized HeLa cells were transfected with siRNA targeting the indicated genes, fixed and stained for MKLP1 (red) and Sec6 (green). Nuclei were visualized by DAPI staining (blue). Graphs on the right represent the percentage of cells in cytokinesis where Sec6 was correctly localized. Depleting RalGDS and RalGPS2 impacted Sec6 localization (upper graph), whereas depleting RGL and RalGPS1 did not.

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AAAACAGACC and GCCCTTCTCACATGCGCCTTTTATCCTC TAAACTCT. PCR product was cloned in pRK5-Myc and sequenced. Synthetic siRNAs targeting Ral GTPases and RalGEFs were designed by standard methods (sequences in Supplementary Methods S9). Exocyst subunits were depleted by published siRNA (Chien and White, 2003; Chien et al., 2006; Lim et al., 2006).

Antibodies: anti-RalA and anti-Aurora B kinase (AM1) were obtained from Beckton Dickinson; anti-tubulin FITC-conjugated mAb from Sigma; anti-Myc mAb from Roche; anti-Sec5 and anti- Sec10 Abs as previously described (Gromley et al., 2005). Anti-Sec6 was obtained from Stressgen Bioreagents; anti-MKL1p from Santa-Cruz; anti-PRC1 was a kind gift of W Jiang (Burnham Institute, La Jolla, CA, USA). Anti-GSD-1A and -1B spectrin was a gift of N Nicolas and MC Lecomte (Inserm, U665, Paris); anti-endobrevin was a gift of T Galli (UJM, Paris, France).

**Cell culture**

HeLa cells were cultivated in Dulbecco’s modified Eagle medium (DMEM) and 10% fetal calf serum at 37°C in humidified chamber on Falcon plastic dishes. Cells (60–120 × 10^5/ml) were plated on treated slides in 24-wells (Falcon) synchronized by double thymidine block (Bootsma et al., 1964), maintained in 2.5 mM thymidine (Sigma) for 18 h, washed three times in DMEM and incubated in DMEM containing 10% fetal calf serum and 24 μM 2′-deoxyxytidine hydrochloride (Sigma) for 8 h. Cells were then blocked again in 2.5 mM thymidine for 16 h. The second block was released as described above. Cells entered G2/M ~ 6.5 h later.

**Transfection and biochemistry**

HeLa cells (12 × 10^5/ml) were plated in 24-wells (Falcon) and synchronized as described above. They were transfected with plasmids (0.4 μg) between the first and the second thymidine block with Lipofectamine PLUS reagents (Invitrogen) in OptiMEM for 3 h according to the manufacturer’s instructions. Where indicated, cells were co-transfected with a plasmid encoding for the GFP, used as a tracer.

siRNA transfection for immunolocalization studies was performed using the Hiperfect forward-protocol (Qiagen) on the first day of synchronization. A total of 60 × 10^5 cells per ml were plated on treated slides in 24-well microplate (Falcon) in growth medium containing the indicated concentrations of siRNA and 3 μl of Hiperfect. For phenotyping, 20 × 10^5 cells per ml were plated on black 96-well microplate with clear bottom (Perkin Elmer) in growth medium containing 5 nM of pooled siRNAs and 1 μl of Hiperfect for 72 h. In rescue experiments, cells were transfected with siRNA in 96-well plates as described above and the next day they were transfected with plasmids using JETPEI reagents (Polyplus Transfection). mRNA levels were measured by quantitative RT-PCR using TaqMan Universal PCR or SYBR Green PCR Master Mix (Applied Biosystems) and 7500 Real-Time PCR System (see Tagman probes and primers used with SYBR Green in Supplementary Methods S9).

**Immunofluorescence and time-lapse videomicroscopy**

HeLa cells were fixed and stained, and images were acquired and analysed using classical immunofluorescence methods. Specific treatments according to antibodies and aims are described in Supplementary Methods S9.

For time-lapse videomicroscopy, HeLa cells were synchronized and transfected on 35-mm glass dishes (Iwaki) and maintained at 37°C in an open chamber (Life imaging) equilibrated in 5% CO_2_. When GFP was used as a tracer, cells were transfected with a ratio of 5:1 between the plasmid expressing the studied gene and the plasmid expressing GFP. Time-lapse sequences were recorded at 5-min intervals on a Leica DMIRBE microscope using a ×20 objective controlled by the Metamorph software. This microscope was equipped with a cooled CCD camera (Micro Max 5MHz; Ropper Scientific). Video analysis was performed by the Meta- morph software.

**Supplementary data**

Supplementary data are available at The EMBO Journal Online (http://www.embojournal.org).

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