A Conserved Di-Basic Motif of Drosophila Crumbs Contributes to Efficient ER Export

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Abstract

The Drosophila type I transmembrane protein Crumbs is an apical determinant required for the maintenance of apico-basal epithelial cell polarity. The level of Crumbs at the plasma membrane is crucial, but how it is regulated is poorly understood. In a genetic screen for regulators of Crumbs protein trafficking we identified Sar1, the core component of the coat protein complex II transport vesicles. sar1 mutant embryos show a reduced plasma membrane localization of Crumbs, a defect similar to that observed in haunted and ghost mutant embryos, which lack Sec23 and Sec24CD, respectively. By pulse-chase assays in Drosophila Schneider cells and analysis of protein transport kinetics based on Endoglycosidase H resistance we identified an RNKR motif in Crumbs, which contributes to efficient ER export. The motif identified fits the highly conserved di-basic RxKR motif and mediates interaction with Sar1. The RNKR motif is also required for plasma membrane delivery of transgene-encoded Crumbs in epithelial cells of Drosophila embryos. Our data are the first to show that a di-basic motif acts as a signal for ER exit of a type I plasma membrane protein in a metazoan organism.

Keywords COPII, Crumbs, di-basic motif, Drosophila, ER export, Sar1

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Crumbs (Crb) is an evolutionarily conserved type I transmembrane (TM) protein, required for apico-basal polarity, integrity of epithelial tissues, and prevention of retinal degeneration from flies to mammals (1,2). Crb proteins localize apically in epithelia and photoreceptor cells, where they organize a membrane-associated protein scaffold (1,3). The large extracellular region of most Crb proteins contains an array of epidermal growth factor (EGF)-like repeats and several laminin A G-like domains. The short, highly conserved cytoplasmic tail is characterized by a 4.1/Ezrin/Radixin/Moesin (FERM) domain-binding and a carboxy-terminal (C-terminal) PSD-95/Dlg/ZO-1 (PDZ) domain-binding site (4), mediating various protein interactions. In Drosophila, the amount of Crb at the apical surface is crucial for cell polarity and growth regulation via the Hippo pathway. Loss of Crb can result in loss/reduction of the apical surface or enhanced tissue growth, while overexpression can lead to an increase of the apical surface (5), the induction of a second apical pole (6) or deregulated growth (7). Several mechanisms regulate Crb trafficking and stability, and thus ensure correct Crb levels at the plasma membrane. These include trafficking by exocyst (Exo84)-dependent exocytosis as well as Cdc42-, Rab11- and retromer-dependent recycling (8–13).

Trafficking of TM proteins to the surface involves the packaging into ER-derived coat protein complex II (COPII)-coated vesicles. This is mediated by specific sorting signals in the cytoplasmic regions of cargo proteins and their interaction with highly conserved COPII components (14,15). The COPII complex consists of the small GTPase secretion-associated RAS-related 1 (Sar1) as well as the heterodimeric Sec23/24 and Sec13/31 complexes. COPII coat assembly at the ER membrane and subsequent vesicle budding is initiated by the GDP/GTP exchange on Sar1 through its guanine nucleotide exchange factor Sec12. Thereby Sar1 is activated and recruits the Sec23/24 and Sec13/31 complexes, which form the inner
and outer vesicle coat, respectively (16–18). Various ER export signals in the cytoplasmic regions of cargo proteins have been identified, including di-acidic, hydrophobic and di-basic motifs (14). In most cases, cargo molecules interact with the COPII subunit Sec24 (19,20), while few cargo molecules, such as Golgi resident glycosyltransferases, directly interact with Sar1 (21).

Here, we show by in vitro and in vivo assays that Drosophila Crb uses a di-basic RNKR motif for efficient COPII-mediated export. This motif is sufficient to improve ER export of a reporter protein, while a di-acidic motif and the PDZ domain-binding motif are dispensable. Our results further suggest that the RNKR motif, which is highly conserved in most other Crb proteins, binds Sar1 to mediate efficient COPII-dependent ER export.

**Results and Discussion**

**Mutations in components of the COPII complex lead to defects in Crb trafficking in embryonic epithelia**

To identify regulators of Crb protein trafficking to the apical plasma membrane in Drosophila embryonic epithelial cells, we performed a genetic screen using a collection of deficiency lines covering the third chromosome (unpublished data). One candidate identified as a crucial component for Crb transport in many embryonic epithelial cells turned out to be the small GTPase Sar1. In wild-type epithelial cells, such as the trachea, Crb localizes to the apical plasma membrane (22) (Figure 1A-A′′ arrowheads). In contrast, the majority of Crb is diffusely distributed in the cytoplasm of sar1 mutant embryos from late stage 15 onward (Figure 1B-B′′, orange arrowheads). Due to the strong maternal contribution of Sar1, zygotic sar1 mutant embryos show a reduction, but not a complete removal of Sar1 from the trachea (23,24). This explains the residual Crb protein detected at the apical surface (Figure 1B-B′′, white arrowheads) and the lack of defects in tracheal tissue morphogenesis, which is usually observed in embryos completely lacking crb (22). The maternal contribution of Sar1 is also sufficient for normal development of other embryonic epithelia. As a COPII component, Sar1 is involved in protein secretion and tracheal tube expansion, and its loss causes defects in cuticle formation and airway maturation as published by Tsarouhas et al. (23). Indeed, we also observe a reduction in tracheal tube diameter (Figure 1B). Similar but less severe defects are also seen in embryos mutant for haunted (hau) and ghost (gho) (also called stenosis [sten]), the Drosophila genes encoding the COPII components Sec23 and Sec24CD, respectively (25,26). As in sar1 mutants, Crb protein accumulates in the cytoplasm of tracheal cells of stage 17 hau and gho mutant embryos (Figure 1C-C′′, orange arrowheads). The normal localization of Crb in stage 15 mutant hau embryos described previously (25) can be explained by the strong maternal contribution of Sec23, which allows for normal trafficking early in development, but no longer sustains Crb trafficking at later stages as shown here. For gho mutant embryos, our results are in agreement with published data showing a cytoplasmic accumulation of Crb in the epidermis of stage 16 gho mutant embryos (25). The weaker phenotype observed in gho mutants compared to sar1 mutants can also be explained by the maternal contribution of Sec24CD. In addition, a second sec24 gene, i.e. sec24AB, has been described in Drosophila (25), whose expression could also contribute to the weaker phenotype of gho mutants. However, to our knowledge, Sec24AB is not yet described at the protein level, especially since a well characterized mutant allele is missing. In conclusion, our results imply that Crb is trafficked from the ER in a COPII-dependent manner.

**Mutation of the RNKR motif leads to trafficking defects of Crb**

Classical COPII-dependent ER export motifs within the cytoplasmic tails of TM proteins include di-acidic and hydrophobic motifs as well as a di-basic motif (14,27). The inspection of the 37-amino acid long cytoplasmic tail of Crb revealed two putative classical ER export signals, a di-basic motif (RNKR) in the juxtamembrane region and di-acidic motif (EMD) located more distally. Furthermore, the short cytoplasmic tail contains a FERM domain and a PDZ domain-binding site (Figure 2A, marked blue and orange, respectively), which have known interaction partners, e.g. Stardust for the PDZ domain-binding site. PDZ domain-containing proteins have been discussed to be involved in ER exit of their binding partners, e.g. the TM protein NR1, a subunit of the N-methyl-D-aspartate receptor (28), or Na+/H+ exchanger regulatory factor 1 and −2, which bind to the glutamate transporter GLAST at different steps during its transport to the plasma membrane (29). Thus, beside the two classical ER export motifs, the PDZ
domain-binding site might also be involved in ER export. Interestingly, all three motifs are highly conserved in most Crb proteins from *Drosophila* to vertebrates (Figure 2B).

To determine which of the motifs is required for ER export of Crb, we analyzed protein transport kinetics. In mammalian cell culture systems this approach is often based on the Endoglycosidase H (Endo H) resistance of overexpressed wild-type and mutant variants. Endo H cleaves high mannose N-glycans (containing at least three mannose moieties) and some hybrid type but not complex N-glycans as found in the late Golgi. Thus, Endo H resistance is characteristic for proteins that have already passed the early Golgi (30,31). *Drosophila* is known to form predominantly high mannose and paucimannose glycans, which might be core-fucosylated (32). Endo H resistance of fly proteins has been shown to occur, e.g. for VIP36 expressed in *Drosophila*-derived SL-3 cells (33). Here, we adopted a pulse-chase experiment using *Drosophila* Schneider (S2R+) cells overexpressing wild-type Crb as well as variants carrying mutations in the putative ER exit motifs, and analyzed Endo H resistance.

The constructs used encode a minimized version of Crb, named Crb-short, in which the extracellular region is reduced to the signal sequence and the two C-terminal EGF-like repeats, which carry four potential N-glycosylation sites, followed by the TM region and the cytoplasmic tail. To detect these proteins, an HA-tag was introduced N-terminal to the EGF-like repeats. Based on a wild-type version, called HA-Crb-short\textsuperscript{RNKR}, single and double mutants in the putative export motifs were generated, called HA-Crb-short\textsuperscript{RNKR>MNEP}, HA-Crb-short\textsuperscript{EMD>AMA}, HA-Crb-short\textsuperscript{double} and HA-Crb-short\textsuperscript{ΔERLI} (Figure 2C). To assess transport kinetics in the early secretory pathway of wild-type and mutant HA-Crb-short proteins, S2R+ cells expressing the respective proteins were metabolically labeled, chased for different time intervals and lysed. Proteins were treated with Endo H and analyzed by gel electrophoresis and autoradiography. At time point 0, a portion of pulse labeled HA-Crb-short\textsuperscript{RNKR} (wild-type) exhibits Endo H resistance (Figure 2D, the two protein fractions are termed resistant [r] and sensitive [s], respectively). After 90 and 180 min chase intervals, metabolically labeled HA-Crb-short\textsuperscript{RNKR>MNEP} achieves almost full Endo H resistance, demonstrating that most of the labeled protein has left the ER and reached the Golgi. In contrast, mutant HA-Crb-short\textsuperscript{RNKR>MNEP} shows reduced Endo H resistance after 90 or 180 min chase intervals (Figure 2D). Quantification of Endo H resistance in six independent experiments revealed that only about 67% of mutant HA-Crb-short\textsuperscript{RNKR>MNEP} protein is Endo H resistant after 180 min chase, in comparison to about...
Figure 2: The RNKR motif is necessary for efficient ER export of HA-Crb-shortRNKR. (A) Schematic representation of the Crb cytoplasmic tail with the FERM domain-binding motif in blue and the PDZ domain-binding motif in orange. (A–C) The underlined letters correspond to the RNKR, the EMD and the ERLI motif. (B) A ClustalW multiple sequence alignment of Crb cytoplasmic tails of Drosophila melanogaster (DmCrb, P10040) together with different Crb proteins from Homo sapiens (HsCRB1, P82279; HsCRB2, Q5IJ48; HsCRB3, Q9BUF7) and Danio rerio (DrCRB1, Q1A5L3; DrCRB2b, Q1A5L1; DrCRB3a, Q1A5L0). (C) The Crb variants used in this study. Brown rectangle: HA-tag, blue rectangles: EGF-like repeats, gray bar: TM region. Red letters in the amino acid sequences indicate exchanges. (D) S2R+ cells expressing HA-Crb-shortRNKR or HA-Crb-shortRNKR>MNEP were pulse labeled for 60 min and chased for the times indicated. Transport to the Golgi was measured by appearance of Endo H resistant protein (r) in relation to Endo H sensitive protein (s). (E) Quantification of Endo H resistance of HA-Crb-shortRNKR and HA-Crb-shortRNKR>MNEP. The values are means ± standard deviation of six independent experiments. ***p ≤ 0.001 versus wild-type t-test. *p ≤ 0.01 versus wild-type t-test. (F) Quantification of Endo H resistance of HA-Crb-shortRNKR, HA-Crb-shortEMD>AMA and HA-Crb-shortERLI as well as HA-Crb-short-double and HA-Crb-shortRNKR>MNEP. The values are means ± standard deviation of three independent experiments. The t-test comparing wild-type versus HA-Crb-shortRNKR>MNEP revealed p ≤ 0.01 at each time point. The t-test comparing wild-type versus HA-Crb-short-double revealed p ≤ 0.05 at 0 and 90 min and p ≤ 0.01 at 180 min.
80% of wild-type HA-Crb-short\textsuperscript{RNKR} protein (Figure 2E). To analyze the role of the di-acidic motif (EMD), the di-acidic motif in combination with the di-basic motif (double mutant), and the PDZ domain-binding site ERLI, we expressed HA-Crb-short\textsuperscript{EMD\textgreater\textgreater AMA}, HA-Crb-short\textsuperscript{double} and HA-Crb-short\textsuperscript{\textgreater\textgreater AERLI} (Figure 2C) and measured their Endo H resistance in pulse-chase experiments. Strikingly, no decreased Endo H resistance is observed for HA-Crb-short\textsuperscript{EMD\textgreater\textgreater AMA} or HA-Crb-short\textsuperscript{\textgreater\textgreater AERLI} compared to wild-type HA-Crb-short\textsuperscript{RNKR}. At 180 min chase, only the mature, Endo H-resistant proteins are detected after Endo H treatment, similar as described for the wild-type protein (Figure 2F). In contrast, HA-Crb-short\textsuperscript{double} shows the same reduced Endo H resistance as the singly mutated protein HA-Crb-short\textsuperscript{\textgreater\textgreater RNKR\textgreater\textgreater MNEP} (Figure 2F).

To our knowledge, these results are the first obtained using Endo H resistance of a \textit{Drosophila} protein as a tool to measure transport kinetics. By this, we could demonstrate that the conserved di-basic RNKR motif plays a vital role for the efficient exit of the type I plasma membrane protein Crb from the ER, while the di-acidic motif and the PDZ domain-binding site are not required under these experimental conditions. The high conservation of this di-basic RxKR motif in most vertebrate Crb proteins (Figure 2B) suggests an evolutionary conserved function. So far, a di-basic motif was shown to act as an ER export signal for Golgi-resident glycosyltransferases, which are type II membrane proteins (21). A di-basic sorting motif together with a di-leucine motif is required for the COPII-dependent ER export of a phosphotransferase subunit, which is a type III membrane protein (34). Finally, an RRR motif is required for the ER exit of the \textalpha\textsubscript{2B} adrenergic receptor, a seven TM protein (35). Thus, our finding of the influence of a di-basic motif for the exit of a type I plasma membrane protein is a further example for a more general role of this motif in ER export.

The observation that a large fraction of the mutant HA-Crb-short\textsuperscript{\textgreater\textgreater RNKR\textgreater\textgreater MNEP} protein was still transported out of the ER suggests that the RNKR motif is not the only export signal required under these experimental conditions. However, our results excluded the di-acidic motif and the PDZ-domain binding region. Additional, conserved or novel, motifs in the TM region or the cytoplasmic tail may be required for efficient Crb export.

A possible motif is an RL-dipeptide, a sequence, which was recently identified as an ER export signal in the rat \textgamma \textalpha\textdelta\textbetaaminobutyricacid transporter-1 (36). Two RL-dipeptides are also present in the Crb cytoplasmic tail. Other proteins use unique signals, such as the Kit ligand, whose efficient ER export relies on a valine residue in the C-terminus of its cytoplasmic domain (37). Therefore, it is not unlikely that more than one ER exit signal is involved in the efficient ER export of Crb.

\textbf{A mutated RNKR motif affects efficient Crb trafficking in embryonic epithelial cells}

In order to test whether the RNKR motif in the HA-Crb-short cytoplasmic tail is also essential for ER export in \textit{Drosophila}, we generated transgenic flies carrying UAS-constructs encoding HA-Crb-short\textsuperscript{RNKR} or HA-Crb-short\textsuperscript{RNKR\textgreater\textgreater MNEP}. First, we expressed both proteins in embryonic tracheal cells using the \textit{btl}-\textit{GAL4} driver line in otherwise wild-type embryos. Immunohistochemistry of stage 16 embryos revealed that the majority of HA-Crb-short\textsuperscript{RNKR} localizes at the plasma membrane together with the membrane marker \textalpha\text-Spectrin (Figure 3A-A′′ white arrowheads). In contrast, HA-Crb-short\textsuperscript{RNKR\textgreater\textgreater MNEP} is predominantly visible in the cytoplasm of the tracheal cells (Figure 3B-B′′ orange arrowheads), but hardly detectable at the plasma membrane (Figure 3B and B′′ white arrowheads). To confirm our data in a different cell type, we also analyzed both transgene-encoded proteins in the amnioserosa cells of stage 13–14 embryos, using the ubiquitously expressing \textit{daG32-GAL4} line (Figure 3C-F). In these relatively big cells, the majority of HA-Crb-short\textsuperscript{RNKR} localizes to the plasma membrane, where it colocalizes with the membrane marker \textalpha\text-Spectrin (Figure 3C-C′′ white arrowheads). Only some protein is detected intracellularly (Figure 3C and C′′ orange arrowheads). In contrast, HA-Crb-short\textsuperscript{RNKR\textgreater\textgreater MNEP} is predominantly visible in the cytoplasm of the amnioserosa cells (Figure 3D-D′′ orange arrowheads), but hardly visible at the plasma membrane (Figure 3D and D′′ white arrowheads). To identify the intracellular compartment, to which both proteins localize, we used an antibody that recognizes the KDEL signal, necessary for retrograde protein transport from the Golgi and therefore considered as an ER marker. Most of the intracellular pool of both HA-Crb-short\textsuperscript{RNKR} and HA-Crb-short\textsuperscript{RNKR\textgreater\textgreater MNEP} colocalize with the KDEL signal.
ER Export of Crb

(Figure 3E-F′′ orange arrowheads), confirming their localization in the ER. The presence of some HA-Crb-short\textsuperscript{RNKR} protein in the ER could be explained by GAL4-mediated overexpression of the protein.

To summarize, the RNKR motif of the Crb cytoplasmic tail is necessary for efficient ER export not only in S2\textsuperscript{R+} cells, but also in embryonic epithelia, such as tracheal and amnioserosa cells. Thus, the function of a di-basic ER export motif in a Drosophila type I TM protein can be shown in the physiological context of the organism. The results are supported by the observation that the truncated Crb protein encoded by the allele \textit{crb}\textsuperscript{BF105} localizes at the apical plasma membrane in some epithelial cells that maintain their polarity, e.g. the boundary cells in the embryonic hindgut (38). The protein encoded by this allele retains the N-terminal 14 amino acids of the cytoplasmic domain, including the RNKR motif, but lacks the EMD and the ERLI motif (including one RL di-peptide) as well as an additional RL-dipeptide harbored between the FERM- and PDZ-domain-binding sites (4). Therefore, the di-acidic EMD-, the RL- and the ERLI motif can be excluded as crucial ER export signals, not only in S2\textsuperscript{R+} cells but also in embryonic epithelial cells. Furthermore, a Crb-short version with the entire cytoplasmic region replaced by GFP, thus missing the RNKR motif, predominantly localizes to the ER upon overexpression in S2\textsuperscript{R+} cells and amnioserosa cells (data not shown), which strengthens the role of the RNKR motif for efficient ER export.

The RNKR motif is also functional in a reporter protein context

In order to test whether the identified ER export motif is also functional in a different protein context, we designed two reporter constructs. They encode the extracellular and TM-region of CD2, followed by the wild-type or the mutated RNKR motif fused to GFP, and are named CD2\textsuperscript{RNKR-GFP} and CD2\textsuperscript{MNEP-GFP}, respectively (Figure 4A). CD2, as a heterologous membrane marker (39), contains three potential N-glycosylation sites. The proteins were expressed in S2\textsuperscript{R+} cells, and Endo H resistance of metabolically labeled CD2 proteins was monitored at two different chase time points (30 and 180 min; Figure 4B and C). Without Endo H treatment, four bands are detected in cells expressing CD2\textsuperscript{RNKR-GFP} and CD2\textsuperscript{MNEP-GFP} after 0, 30 and 180 min chase. The fastest migrating band probably corresponds to the non-glycosylated protein, as it co-migrates with fully deglycosylated protein after Endo H treatment (Figure 4B; [s]). The slower migrating bands (Figure 4B; 1–3) probably correspond to mono-, di- and tri-glycosylated proteins, respectively. Upon Endo H treatment of either protein, predominantly fully deglycosylated proteins were detected, indicating that the majority of CD2\textsuperscript{RNKR-GFP} and CD2\textsuperscript{MNEP-GFP} has not yet obtained a Golgi-dependent modification of their glycans (Figure 4B, time point 0 and 30 min). After 180 min chase less CD2\textsuperscript{MNEP-GFP} is Endo H resistant compared to CD2\textsuperscript{RNKR-GFP} (Figure 4B). The difference observed between the Endo H resistance of CD2\textsuperscript{RNKR-GFP} and CD2\textsuperscript{MNEP-GFP} after a 180-min chase is very small, but significant (Figure 4C). A similar difference in Endo H resistance was observed after a 90- and 360-min chase period (data not shown). Interestingly, the Endo H-resistant protein (r) migrates faster than the tri-glycosylated one (Figure 4B, 180 min chase, band 3). This may be due to strong trimming of the N-glycans in the Golgi. This is supported by the observation that after 360 min chase, the majority of non-treated proteins co-migrate with Endo H resistant proteins (data not shown). The results obtained suggest that the RNKR motif introduced into a reporter protein also improves ER exit. The rather small effect of the RNKR motif on the transport of the reporter is either due to the fact that this motif is not fully recognized in this particular context, or suggests that the RNKR motif alone is not a very strong signal, or a combination of both.

The cytoplasmic tail of Crb interacts with Sec24CD

In most cases studied, cargo selection by the COPII machinery is mediated by the Sec24 subunit (19,20). In order to find out whether the RNKR motif of Crb is required for the interaction with the COPII complex, we co-expressed an EGFP-tagged version of Sec24CD together with either HA-Crb-short\textsuperscript{RNKR} or any of the HA-Crb-short mutant proteins described above (Figure 2C). Following a successful co-overexpression in S2\textsuperscript{R+} cells, complexes were immunoprecipitated from cell lysates with an anti-HA antibody and analyzed by Western Blot using an anti-GFP-antibody (Figure 5A, top panel). The wild-type HA-Crb-short\textsuperscript{RNKR} as well as all mutant versions of the protein could precipitate EGFP-Sec24CD, showing that Sec24CD can interact with...
Figure 3: HA-Crb-short^RNKR shows impaired traffic to the plasma membrane in embryonic epithelial tissues. (A-B′′) Confocal microscopy images of tracheal epithelial cells of stage 16 embryos expressing HA-crb-short^RNKR (A-A′′) or HA-crb-short^RNKR>MNEP (B-B′′) under the control of blt-GAL4. Embryos were stained with anti Crb-intra (green) and anti α-Spectrin (magenta), marking the cell outlines. (C-F′′) Confocal microscopy images of amnioserosa epithelial cells of stage 13–14 embryos expressing HA-crb-short^RNKR (C-C′′ and E-E′′) or HA-crb-short^RNKR>MNEP (D-D′′ and F-F′′) under the control of daG32-GAL4. (C-D′′) Embryos were stained with anti Crb-intra (green) and anti α-Spectrin (magenta) (C′, C′′, D′, D′′) and the ER marker KDEL (magenta) (E′, E′′, F′, F′′). Scale: 10 μm (A-B′′); 5 μm (C-F′′). White arrowheads mark the cell outlines, orange arrowheads the cytoplasm.
Figure 4: The RNKR motif is functional in the CD2-reporter construct. (A) Schematic representation of the reporter constructs consisting of the extracellular (brown rectangle) and TM (gray bar) region of CD2, followed by the wild-type or mutated motif and GFP (green rectangle). (B) S2R+ cells expressing CD2RNKR-GFP and CD2MNEP-GFP were pulse labeled for 30 min and chased for the times indicated. Transport to the Golgi was measured by the presence of Endo H resistant protein (r) in relation to Endo H sensitive protein (s). (D) Quantification of Endo H resistance of CD2RNKR-GFP and CD2MNEP-GFP in three independent experiments. *p ≤ 0.01 versus wild-type t-test.

the cytoplasmic tail of Crb regardless of the mutations in the putative ER export motifs. Interestingly and comparable to our results, the above mentioned RRR motif of the mammalian α2B-adrenergic receptor was shown to preferentially interact with Sec24C and Sec24D rather than Sec24A or Sec24B (35).

The RNKR motif mediates the interaction with GST-Sar1

As binding of Sec24CD was not largely affected by mutations in the RNKR motif, we further analyzed the role of the di-basic motif for a possible COPII interaction. In vertebrate glycosyltransferases a di-basic ER export motif has been reported to directly bind to the small GTPase Sar1 (21). To test whether the di-basic ER export motif of Crb interacts with Drosophila Sar1, S2R+ cell lysates containing either HA-Crb-shortRNKR or HA-Crb-shortRNKR>MNEP were incubated with recombinant GST-tagged Sar1H74G, which is the active GTP-locked mutant (40). Protein complexes were analyzed by glutathione-sepharose pull-down and Western blotting. GST-Sar1H74G pulled down some HA-Crb-shortRNKR (Figure 5B, lane 5), but not HA-Crb-shortRNKR>MNEP (Figure 5B, lane 6). Thus, the RNKR motif in the cytoplasmic tail of Crb could potentially mediate the interaction between Crb and the COPII complex via Sar1 in order to increase transport efficiency.

Beside the vertebrate glycosyltransferases mentioned above, Sar1 is known to directly interact with cargo proteins such as yeast Bet1 (21,41), suggesting that direct binding is a more common phenomenon. The rather weak interaction between GST-Sar1H74G and HA-Crb-shortRNKR could be due to the fact that the interaction requires co-factors whose amounts and/or activity are limited in this experimental system. Alternatively, GST-Sar1H74G or HA-Crb-shortRNKR may bind strongly to other proteins present in the lysate. Beyond that, the weak interaction might result from the lack of membranes, which usually provide the physiological environment and promote interactions.

In summary, we have shown that the RNKR motif in Crb contributes to the ER export by interactions with Sar1. This is the first example of the role of a di-basic motif as ER export signal in a Drosophila type I membrane protein. We further show that this sequence is functional in a
The cytoplasmic tail of Crb interacts with the COPII machinery. (A) EGFP-Sec24CD interacts with HA-Crb-short RNKR, HA-Crb-shortRNKR>MNEP, HA-Crb-shortEMD>AMA, HA-Crb-shortDouble and HA-Crb-shortΔERLI, shown by co-immunoprecipitation and Western Blot analyses. S2R+ cells expressing EGFP-Sec24CD together with GFP (control), HA-tagged Crb-shortRNKR, Crb-shortRNKR>MNEP, Crb-shortEMD>AMA, HA-Crb-shortDouble or Crb-shortΔERLI were lysed and the Crb variants were immunoprecipitated with anti HA-antibody. Immunoprecipitates were separated by SDS-PAGE and Sec24CD was determined by immunoblotting using anti GFP-antibody (upper panel). Lower panels show the input controls of EGFP-Sec24CD and the HA-tagged Crb variants, probed with anti-EGFP and anti-HA antibody, respectively. (B) Lysates from S2R+ cells expressing HA-Crb-short RNKR, HA-Crb-shortRNKR>MNEP or mock-transfected (control) were incubated with glutathione-sepharose beads with GST (lane 1–3) or in the presence of recombinant GST-Sar1H74G (lane 4–6). After incubation, beads were washed and bound complexes analyzed by SDS-PAGE and Western blotting with anti-HA antibody. Lanes 7–9 serve as input control. The asterisk in lane 5 marks the band.

Materials and Methods

Fly stocks

Flies were kept at 18 or 25°C and staged according to the time of development described in Campos-Ortega and Hartenstein (45). The following stocks were obtained from the Bloomington stock center: OregonR as wild type control, sar105712, sec239G, gho1, daG32-GAL4, and btlGAL4. The mutant stocks were balanced over TM3, twist-GAL4, UAS-EGFP or Cyo, twist-GAL4, UAS-EGFP. The following stocks were generated in this study: UAS-HA-crb-shortRNKR, UAS-HA-crb-shortRNKR>MNEP. Transgenic flies were generated using the phiC31 integrase mediated site-specific integration into attP landing-sites (46). For the injection and establishment of transgenic lines, standard protocols were followed (47). UAS-HA-crb-shortRNKR and UAS-HA-crb-shortRNKR>MNEP were integrated into the landing site attP3A of the stock y M(vas-int.DM)ZH-2A w; PBac (y+ attP-3B) VK00001 (Bloomington No. 24861).

Antibodies

Primary antibodies used: rabbit anti-Crb-intra2662, (1:400, raised against the peptide NKRATRGTYSPSAQE), rat anti-Crb2.8 (1:1000) (48), rabbit anti-GFP (Life Technologies), mouse anti-HA antibodies (HA.11 Clone 16B12, Covance), mouse anti-KDEL (1:300) (Stressgen), mouse anti-αSpectrin 3A9 (1:400) (DSHB).

Plasmids

To clone the Crb-short construct, the cDNA sequence representing the 28th and 29th EGF-like repeat, the TM region, the cytoplasmic tail and the 3’UTR of Crbwt (5) was amplified by PCR using the primers (5’TATAGTCTGACATCGAGTGCAAC-3’ and 5’GGTCTAGAGCAAAATATGTTTTTTATTTG-3’), and (5’TTGCTAGATCGGACGAGTGCAAC-3’ and 5’GGTCTAGAGCAAAATATGTTTTTTATTTG-3’), including the restriction sites BglII and XbaI. The PCR product was cloned into the Crbwt construct (5), replacing the TM region, cytoplasmic tail and 3’UTR of this construct. The HA-tag was inserted by overlap-extension PCR using the primers (5’TATAGTCTGACATCGAGTGCAAC-3’ and 5’AGCTGTAATCGGAACATCGGACGACGAGTGCAAC-3’).
5'-TACCCCATAGTTCTGCCATTACGGTAGGCAGGGTACATT
5'-TTTTGTTCCAAATATTGCT-3', respectively.

The CD22RNKR-GFP and CD22MNEP-GFP fusion constructs, the
CD2-CrbintA (4) construct was used as matrix and was amplified with
the primers 5'-GGACATTGCGATGAGCAACCT-3' and either 5'-CCTGAAT
CCATGGACCTAGGG-3' or 5'-CTCAATGCGATGAGCAACCT-3'. The
second mutation was introduced by overlap-extension PCR with
the primer used for cDNA amplification. Each construct was
PCR using the primer 5'-ACTTGGGTTGGCGCCTACTCAGGCACG-3'
and 5'-CGTGCGTGGAGTGCGGCCCGCCAACTCAGG-3', each in combina-
tion with the primer used for cDNA amplification. Each construct
was sequenced for confirmation of its identity.

Embryo collection and immunohistochemistry
Embryo collection, fixation and antibody staining were conducted as
described in Ref. 51. The primary antibodies used are stated above.
The secondary antibodies used in this study were conjugated to Alexa
Fluor-488, −568 and −647 (Life Technologies). Stained embryos were
mounted in glycerol-propyl gallate (75% glycerol, 50mg/mL propyl
gallate) and imaged at room temperature on a confocal microscope with a
Zeiss Plan-Neofluar 100×/1.3 NA oil objective (LSM 510 controlled by
LENS software, Carl Zeiss). Images were further processed with Fiji and
Adobe Illustrator CS5.

Cell culture and transfection
S2R+ cells (Drosophila Genomics Resource Center, stock no. 150) were
grown in Schneider's insect medium (Sigma-Aldrich) supplemented with
heat-inactivated fetal calf serum (Sigma-Aldrich) at 24°C. Cells were
controlled for the absence of mycoplasma using standard PCR. Trans-
fection of S2R+ was done with FuGENE® HD Transfection Reagent
(Promega) as suggested by the manufacturer applying the plasmids
pAct5C-GAL4 together with pUAST-based, Crb-encoding plasmids. As
control, pAct5C-GAL4 was used alone. Cells were harvested 72 h after
transfection.

Metabolic labeling and enzymatic removal of glycans
Cells were depleted of methionine and cysteine for 2 h (Schneider's
Drosophila medium without methionine and glutamine (PAN Biotech)
supplemented with 2mM-glutamine and 10% fetal calf serum dialyzed
against PBS to remove small molecules up to 12 kDa). Labeling
was done with 25μCi/mL Met-[35S]-label (Hartmann) for 1 h, followed
by various periods of time as detailed in the figure legends. Cells were
lysed in 1% Triton X-100 containing lysis buffer [100mM Pipes, pH6.8;
120mM NaCl; 1.5mM MgCl2; 3mM CaCl2; 1% (v/v) Triton X-100;
oncanelutionbuffer,GEHealthcareLifeSciences).Proteinswereboiled
withdithiothreitol(50mM,5°C)andseparatedbySDS–PAGE. Gels were
fixed (30min × 10min,4°C). For immunoprecipitation, cell
lysates were diluted 1:1 with PNTG buffer [50mM Pipes, pH6.8; 120mM
NaCl; 1.5mM MgCl2; 3mM CaCl2; 1% (v/v) Triton X-100; 10% (v/v)
glycerol; 1mM phenylmethylsulfonyl fluoride; 1×CompleteTM
(Roche)] for 10 min at 4°C. Non-solubilized material was separated by
centrifugation (10,000 × g, 10 min, 4°C). For immunoprecipitation, cell
lysates were diluted 1:1 with PNTG buffer [50mM Pipes, pH6.8; 120mM
NaCl; 1.5mM MgCl2; 3mM CaCl2; 1% (v/v) Triton X-100, 3mM CaCl2],
and protein A/G-agarose beads (Roche) as well as anti-HA/GFAP antibodies
were added. Samples were rotated for 16 h, and the beads were washed
twice with PNTG buffer and once with PN buffer (50mM Pipes, pH6.8;
120mM NaCl). Endoglobosidase H (Endo H) treatment was done as sug-
gested by the manufacturer (New England Biolabs). Proteins were boiled
and separated by SDS–PAGE. Gels were fixed (30 min in 40% methanol,
10% acetic acid), dried and analyzed by phosphorimaging (Typhoon FLA
9500; GE Healthcare Life Sciences).

Co-immunoprecipitation
To measure the interaction between Sec24CD and the different
HA-Crb-short versions, S2R+ cells expressing the respective proteins

Traffic 2015; 16: 604–616

613

ER Export of Crb

PCRs were also performed with
the primers 5'-TACCCCATAGTTCTGCCATTACGGTAGGCAGGGTACATT
5'-TTTTGTTCCAAATATTGCT-3', respectively. The overlap-extension product was
amplified with the primers 5'-GGACATTGCGATGAGCAACCT-3' and either 5'-CTCAATGCGATGAGCAACCT-3' or
5'-CCTGAATCCATGGACCTAGGG-3' to insert an isoleucine residue followed by a premature
stop codon. Thus, the construct resembles the construct Myc-IntraΔERLI used in Klebes & Knust (4).

For the double mutant, the single mutant was used to insert the
second mutation by a second round of overlap-extension PCR. The con-
struct 5'-TACCCCATAGTTCTGCCATTACGGTAGGCAGGGTACATT
5'-TTTTGTTCCAAATATTGCT-3' was subsequently ligated into pUAST
vector. For expression in S2R+ cells (Drosophila Genomics Resource Center, stock no. 150) were
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heat-inactivated fetal calf serum (Sigma-Aldrich) at 24°C. Cells were
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lysates were diluted 1:1 with PNTG buffer [50mM Pipes, pH6.8; 120mM
NaCl; 1.5mM MgCl2; 3mM CaCl2; 1% (v/v) Triton X-100; 10% (v/v)
glycerol; 1mM phenylmethylsulfonyl fluoride; 1×CompleteTM
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NaCl; 1.5mM MgCl2; 3mM CaCl2, and protein A/G-agarose beads (Roche) as well as anti-HA/GFAP antibodies
were added. Samples were rotated for 16 h, and the beads were washed
twice with PNTG buffer and once with PN buffer (50mM Pipes, pH6.8;
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gested by the manufacturer (New England Biolabs). Proteins were boiled
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Co-immunoprecipitation
To measure the interaction between Sec24CD and the different
HA-Crb-short versions, S2R+ cells expressing the respective proteins

Traffic 2015; 16: 604–616

613

ER Export of Crb

PCRs were also performed with
were lysed as described above. The lysates were used for co-immunoprecipitation with 2 μg anti-HA antibodies and protein G-agarose beads (Roche) for 16 h at 4°C. The beads were washed three times with PNTG buffer (see above) and once with HN buffer (50 mM Hepes, pH 7.5; 150 mM NaCl). Immunoprecipitated proteins were boiled with SDS gel loading buffer for 10 min at 65°C, separated by SDS–PAGE, and blotted on nitrocellulose. Blots were blocked and incubated with antibody with 5% BSA (Serva) in 1× TBS–T (50 mM Tris–HCl, pH 7.5; 150 mM NaCl; 5 mM EDTA, pH 8.0; 0.2% Tween). For detection, an anti-GFP antibody (1:5000) and an anti-HA antibody (1:2000) were used as primary and a goat anti-rabbit and anti-mouse antibody conjugated with peroxidase (1:10 000; Sigma-Aldrich) were used as secondary antibodies. Immunodetection was done with ECL chemiluminescence blotting substrate (GE Healthcare Life Sciences).

**Sar1 expression and purification**

Expression constructs were transformed into *E. coli* strain BL21 DE3. Cells were grown to OD$_{600}$ 0.6–0.8 at 37°C, induced using 0.1 mM IPTG for 4 h at 30°C and lysed with an EmulsiFlex Homogenizer. Lysates were centrifuged for 30 min at 4°C and 12 000 × g, and the supernatant was loaded on a glutathione sepharose column for gravity flow. GST-Sar1$^{H74G}$ was eluted with 10 mM reduced glutathione (GE Healthcare Life Sciences) in 50 mM Tris, pH 8.0. Elution fractions were analyzed by SDS–PAGE and pooled.

**Interaction between Sar1$^{H74G}$ and HA-Crb-short**

S2R$^+$ cells expressing HA-Crb-short or HA-Crb-short$^{ENKRS-MNPEP}$ were lysed as described before, and incubated with GST-Sar1 (2 μg) and glutathione-sepharose 4B beads (GE Healthcare Life Science). Bound proteins were analyzed by SDS–PAGE and immunoblotting as described above. For detection an anti-HA antibody (1:5000) was used as primary and a goat anti-mouse antibody conjugated with peroxidase (1:10 000) (Sigma-Aldrich) was used as secondary antibody.

**In silico analyses**

The annotated Crb sequences were retrieved from FlyBase (http://flybase.org) and the UniProt accession numbers are given in the figure legend. The alignment was obtained using ClustalW. Den-sitometric analyses were performed with the ImageQuant TL software for 1D gel analysis v8.1 (GE Healthcare Life Sciences). For statistical analyses, two-tailed t-tests for unpaired samples were applied using Excel (Microsoft Office 2011, version 14.01.3).

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**GLOSSARY**

- **COPII**: coat protein complex II
- **Crb**: Crumbs
- **C-terminal**: carboxy-terminal
- **EGF**: epidermal growth factor
- **Endo H**: Endoglycosidase H
- **FERM**: 4.1/Ezrin/Radixin/Moesin
- **N-terminal**: amino-terminal
- **PDZ**: (PSD-95/Dlg/ZO-1)
- **S2R$$^+$$**: Schneider cells
- **TM**: transmembrane
- **UAS**: upstream activating sequence

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