1 TITLE 2 Comparative proximity biotinylation produces an inventory of RAB18-interactions and 3 implicates RAB18 in cholesterol mobilization 4 5 **RUNNING TITLE** 6 **GEF-dependent RAB18 interactions** 7 8 **AUTHORS** Robert S. Kiss*1, Jarred Chicoine1, Robert Sladek1, He Chen1, Alessandro Pisaturo1, 9 Cyril Martin¹, Jessica D. Dale², Tegan A. Brudenell², Archith Kamath^{3,4}, Jimi C. 10 Wills³, Alex von Kriegsheim³, Tommy Nilsson¹, Eamonn Sheridan², Mark T. 11 Handley*2 12 13 14 **AFFILIATIONS** ¹Research Institute of the McGill University Health Centre 15 16 1001 boul Decarie 17 Glen Site Block E 18 Montreal, QC 19 H4A 3J1 20 Canada 21 ² Leeds Institute of Medical Research (LIMR) 22 23 St James's University Hospital 24 Leeds LS9 7TF

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45 KEYWORDS
46 RAB18, BioID, Cholesterol, SPG20, SEC22A, TMCO4
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48 SUMMARY STATEMENT
49 We used proximity biotinylation together with guanine nucleotide exchange factor
50 (GEF)-null cell lines to discriminate functional RAB18-interactions. We anticipate that
51 this approach will be broadly applicable in small GTPase research.

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ABSTRACT Loss of functional RAB18 causes the autosomal recessive condition Warburg Micro syndrome. To better understand this disease, we used proximity biotinylation in HEK293 and HeLa cells to generate an inventory of potential RAB18 effectors. In HeLa cells, we expressed BirA*-RAB18 in cells in which RAB18-quanine nucleotide exchange factor (GEF) activity was disrupted with CRISPR. We found that most RAB18-interactions are regulated independently by its different GEFs; the binary RAB3GAP1-RAB3GAP2 complex and the TRAPPC9-containing TRAPPII complex. RAB3GAP-dependent RAB18 interactions included a group of microtubuleinteracting/membrane shaping proteins, a group of proteins involved in membrane tethering and docking, and a group of lipid-modifying/lipid transport proteins. We demonstrate that GEF-dependent Rab-interactions are highly amenable to interrogation by proximity biotinylation. Further, we provide confirmatory evidence for several of the interactors (SPG20/SPART, SEC22A and TMCO4) as well as functional evidence supporting a role for RAB18 in modulating the close apposition of membranes and in cholesterol mobilization.

70 INTRODUCTION 71 Rab Proteins are a large subfamily of small GTPases with discrete roles in 72 coordinating membrane trafficking (Zhen and Stenmark, 2015). They associate with 73 cellular membranes as a result of their C-terminal prenylation and like other small 74 GTPases, adopt different conformations and enter into different protein-protein 75 interactions according to whether they are GDP-, or GTP-bound. For Rab proteins, 76 cycles of GTP binding and hydrolysis are accompanied by cycles of membrane 77 association and dissociation that serve to promote the targeting of particular Rab protein isoforms to particular membrane compartments. Although they possess 78 79 some intrinsic GTP-hydrolysis activity, their *in vivo* nucleotide-bound state is tightly 80 governed in cells by two classes of regulatory proteins. Guanine-nucleotide exchange factors (GEFs) catalyse the exchange of bound GDP for GTP while 81 82 GTPase-activating proteins (GAPs) promote the hydrolysis of bound GTP to GDP 83 (Lamber et al., 2019, Barr and Lambright, 2010). The dissociation of Rab proteins 84 from membranes is mediated by GDP-dissociation inhibitor (GDI) proteins. GDIs sequester GDP-bound Rabs in the cytosol, and are also involved in their 85 86 reassociation with membranes (Zhen and Stenmark, 2015). 87 Rab proteins have a variety of roles in the regulation of processes required to confer 88 compositional identity to membranous organelles and to subdomains within them. 89 These include membrane remodelling and the establishment of membrane contact 90 sites (Bui et al., 2010, Raiborg et al., 2015, Rocha et al., 2009, Sobajima et al., 2018). Where exchange between organelles is mediated by carrier vesicles, these 91 92 processes are important in vesicle budding and transport, tethering at the 93 appropriate target membrane, and fusion (Cai et al., 2007). Under other 94 circumstances, membrane contact sites may be established to mediate direct fusion 95 between organelles or to facilitate the transfer of lipids and ions (Wu et al., 2018, 96 Langemeyer et al., 2018, Wickner, 2010). Rab proteins fulfil their roles by way of 97 protein-protein interactions with interacting partners termed 'effectors'. These 98 comprise an array of phylogenetically unrelated protein classes and can serve a 99 range of molecular functions. As such, they are most usually identified biochemically. 100 Biochemical identification of Rab effectors is challenging; Rab-effector interactions 101 are usually GTP-dependent and are often highly transient. Immunoprecipitation,

affinity purification and yeast-2-hybrid approaches have each been used but may be

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103 more or less effective depending on the Rab isoform studied (Christoforidis et al., 104 1999, Fukuda et al., 2008). One newer approach that has yielded identification of a 105 number of novel interactions is 'BioID' proximity biotinylation utilizing Rab proteins 106 fused to mutant forms of the promiscuous biotin ligase BirA*; the Rab fusion protein 107 biotinylates proximal proteins which are then purified on streptavidin and identified 108 through mass spectrometry (Gillingham et al., 2019, Liu et al., 2018, Roux et al., 109 2012). Biotin labelling occurs in a relatively physiological context, and prospective 110 effectors can be purified under high stringency conditions. However, a drawback of 111 the technique is that it does not discriminate between close associations resulting 112 from functional protein-protein interactions and those resulting from overlapping 113 localizations. 114 RAB18 is a ubiquitously expressed ancestral Rab protein that localizes to the cis-115 Golgi, endoplasmic reticulum (ER) and lipid droplets (LDs)(Martin et al., 2005, Ozeki 116 et al., 2005, Gerondopoulos et al., 2014, Handley et al., 2015). Previous work has 117 suggested that it functions in the regulation of lipolysis, and lipogenesis (Martin et al., 118 2005, Ozeki et al., 2005, Pulido et al., 2011), trafficking between the Golgi and 119 endoplasmic reticulum (ER) (Dejgaard et al., 2008, Handley et al., 2015), ER 120 structure (Gerondopoulos et al., 2014), exocytosis (Vazquez-Martinez et al., 2007) 121 and autophagy (Bekbulat et al., 2018, Feldmann et al., 2017). RAB18 deficiency or 122 its dysregulation cause the autosomal recessive condition Warburg Micro syndrome 123 (Aligianis et al., 2005, Bem et al., 2011, Borck et al., 2011, Handley and Sheridan, 124 2018, Liegel et al., 2013)(MIMs 600118, 614222, 614225, 615663, 212720). 125 In order to address how RAB18 coordinates and performs its roles, we used BioID to 126 generate an inventory of its potential effectors. To discriminate functional 127 interactions, we used complementary comparative analyses in HEK293 and HeLa 128 cells. Known RAB18 effectors were more strongly labelled by a fusion of BirA* and 129 wild-type RAB18 than by one incorporating an inactive form, RAB18(Ser22Asn). 130 which is deficient in nucleotide-binding. Similarly, BirA*-RAB18 labelled known 131 effectors more strongly in wild-type cells than in cells in which RAB18-GEF activity 132 was disrupted with CRISPR. Interestingly, disruption of different GEF complexes 133 largely affected different sets of RAB18 interactions. 134 A restricted set of interactions were dependent on the binary RAB18-GEF complex 135 made up of RAB3GAP1 and RAB3GAP2 (Gerondopoulos et al., 2014). These

included known and novel interactors in discrete functional groups. We present direct validation for several examples including the microtubule-binding protein SPG20/SPART, the SNARE protein homologue SEC22A and an orphan lipase TMCO4. Our data strongly support a previous suggestion (Xu et al., 2018) that RAB18 effectors act collectively in lipid mobilization. The combined data elaborate a model for RAB18 function and provide groundwork for future investigations. In particular, a putative role in cholesterol mobilization and biosynthesis is highlighted.

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RESULTS An inventory of nucleotide-binding-dependent RAB18-associated proteins in HEK293 cells We first generated HEK293 cells stably expressing BirA*-tagged fusion proteins of wild-type RAB18, nucleotide-binding deficient RAB18(Ser22Asn) or GTP-hydrolysis deficient RAB18(Gln67Leu) using the Flp-In system. We then carried out proximitylabelling, affinity purification and mass spectrometry of biotinylated proteins as previously described (Roux et al., 2018, Roux et al., 2012). Following removal from the dataset of previously identified non-specific binders from an in-house database, a total of 98 proteins were identified as associating with RAB18 across all samples (see Table S1). The most comprehensive annotation of candidate RAB18 effectors thus far was made in the 2014 paper by Gillingham et al., which utilized an affinity purificationmass spectrometry (AP-MS) approach and the *Drosophila* RAB18 orthologue (Gillingham et al., 2014). In that study, a total of 456 proteins were identified as interacting with RAB18. However, only 14 of these were well represented in terms of spectral counts, exhibited low non-specific binding to GST/Sepharose and showed low binding to other Rab protein isoforms. We took these 14 proteins as the most plausible physiological RAB18 interactors. Orthologues of 7 of these 14 proteins were also represented in our BioID data. The number of spectral counts recorded for these 7 proteins did not clearly distinguish them from the remainder of this BioID dataset. We therefore proceeded by comparing their differential labelling by the different RAB18 fusion proteins. We began analyses by exploring the possibility that BirA*-RAB18(Gln67Leu) fusion protein would produce enhanced biotinylation of RAB18-effectors (as compared to BirA*-RAB18(WT)). We normalized total spectral counts between the BirA*-RAB18(Gln67Leu) and BirA*-RAB18(WT) datasets and then calculated mutant:wildtype ratios for each RAB18-associated protein (Table S1). Association ratios for known RAB18-interactors ranged from 0.1-1.49 indicating that RAB18 associations were altered by the Gln67Leu variant, but not predictably so.

175 We next compared the BirA*-RAB18(Ser22Asn) and BirA*-RAB18(WT) datasets 176 (Figure 1, Table S1). The RAB18-GEF subunits RAB3GAP1 and RAB3GAP2 177 showed association ratios >1 consistent with the high affinity of Rab-GEFs for 178 cognate Rabs in their nucleotide-free state. Since most effector-interactions are 179 GTP-dependent, we ranked other prospective RAB18-interactors according to those 180 that showed the lowest association ratios. Figure 1B shows the 28 of 98 proteins 181 with BirA*-RAB18:BirA*-RAB18(Ser22Asn) association ratios <0.5. These include 182 the remaining 5 common interactors from the Gillingham et al. study, several of 183 which have association ratios of zero, indicating that labelling by BirA*-184 RAB18(Ser22Asn) was absent. These data suggest that comparison of labelling by 185 wild-type and nucleotide-binding-deficient BirA*-Rab fusion proteins can be an 186 effective means to distinguish putative effectors. 187 188 An inventory of RAB18-GEF-dependent RAB18-associated proteins in HeLa cells 189 We had previously used CRISPR to generate a panel of clonal, otherwise isogenic, 190 HeLa cell lines, null for RAB18 and a number of its regulators (see Figure S1). 191 Having shown that the BirA*-RAB18(WT):BirA*-RAB18(Ser22Asn) comparison in 192 HEK293 cells was informative, we carried out similar comparisons between BioID-193 labelling in wild-type and RAB18-GEF deficient HeLa cells (Figure 2A). Since GEF 194 activity promotes Rab GTP binding, and this is usually necessary for effector 195 interactions, these interactions would be attenuated in GEF-null cells. RAB3GAP1 196 and RAB3GAP2 are each essential subunits of a binary RAB18-GEF complex 197 whereas TRAPPC9 is reported to be essential for the RAB18-GEF activity of a 198 different GEF, the multisubunit TRAPPII complex (Gerondopoulos et al., 2014, Li et 199 al., 2017). We therefore carried out proximity labelling using transient expression of 200 the same exogenous BirA*-RAB18 construct in wild-type cells and in RAB3GAP1-, 201 RAB3GAP2- and TRAPPC9-null cell lines. 202 Prior to mass-spec analysis, samples from each of the streptavidin pull-downs were 203 subjected to Western blotting to ensure comparable BirA*-RAB18 expression (Figure 204 S2A). Label-free quantitative proteomics analyses were used to calculate 'LFQ 205 intensities' (Cox et al., 2014) for each RAB18-associated protein, which were then 206 normalized in each experiment according to the quantity of RAB18 found in each

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sample. Samples from three independent experiments were analysed, and pull-down samples from untransfected biotin-treated cells were used as controls in each case. After filtering the data to remove known mass-spec contaminants, and any protein identified at a high level in control samples, a total of 584, 483 and 506 RAB18associated proteins were identified in each experiment. A total of 457 proteins were present in two or more of the replicate experiments (see Table S2). Orthologues of 10 of the 14 putative RAB18-interacting proteins identified by Gillingham et al. were identified in the HeLa cell BioID dataset including all 7 of those also identified in the HEK293 cell dataset. However, as in that dataset, these were not distinguished by their comparative abundance. Different Rab-GEF complexes may operate in distinct subcellular localizations and coordinate associations with different effectors (Carney et al., 2006). Therefore, we assessed whether non-zero intensities for each RAB18-associated protein correlated between samples (Figure 2B, Figure S2B). Very strong correlations between protein intensities from RAB3GAP1- and RAB3GAP2-null cells indicated that loss of either protein had a functionally equivalent effect (R²=0.99, see Figure 2B). In contrast, intensities from RAB3GAP1- and TRAPPC9-null cells were much more poorly correlated (R²=0.73, see Figure S2B). We therefore considered RAB3GAP- and TRAPPC9-dependent RAB18-associations separately. Of the 457 proteins identified in two or more independent experiments, only 25 showed an association ratio <0.5 in the absence of functional RAB3GAP (Figure 2C-D). These included orthologues of 9 of the 10 proteins identified in the Gillingham et al. study. Thus, our approach was extremely powerful in discriminating putative effector proteins. 133 proteins showed an association ratio <0.5 in the absence of functional TRAPPC9 including the remaining common RAB18-interactor from the Gillingham et al. study (see Table S2). There was only limited overlap between RAB3GAP- and TRAPPC9-dependent associations (Figure 2C). Indeed, among the 28 nucleotide-binding-dependent RAB18 associations identified in HEK293 cells, 6 were also RAB3GAP-dependent, and 7 were TRAPPC9-dependent in HeLa cells respectively, but none were both (Figure 1B). Among the 25 RAB3GAP-dependent associations in HeLa cells, only 5 were also TRAPPC9 dependent (Figure 2C, Table S2).

One of the TRAPPC9-dependent RAB18 associations was that with TBC1D5, a Tre-2/Bub/Cdc16 (TBC) domain-containing RAB-GAP with a well characterised role in regulation of RAB7 (Jia et al., 2016, Jimenez-Orgaz et al., 2018, Seaman et al., 2009). This was a strong candidate as a RAB18 regulator or effector since it had also been identified in the HEK293 dataset as well as in several previous studies (Gillingham et al., 2019, Gillingham et al., 2014). We generated TBC1D5-null HeLa cells, and first tested whether RAB7 and RAB18 dynamics were altered in these cells using fluorescence recovery after photobleaching (FRAP). RAB7 dynamics were substantially different in the TBC1D5-null cells as compared to those in wild-type cells, consistent with reduced RAB7 GTP-hydrolysis resulting in its reduced GDImediated exchange between membrane and cytosolic compartments (Figure S3A). In contrast, RAB18 dynamics were unchanged in TBC1D5-null cells compared to controls (Figure S3B). Further, RAB7 dynamics were unchanged in RAB18-null cells compared to controls, indicating that RAB18 is not required for TBC1D5 activity (Figure S3A). These data do not exclude TBC1D5 as a potential RAB18 effector, but argue against a role for RAB18 in RAB7-regulation under resting conditions.

Validation screening of RAB3GAP-dependent RAB18 associations

Our continued study focused on the 25 RAB3GAP-dependent RAB18 associations identified in HeLa cells on the basis that these included the majority of known RAB18 effectors together with a number of promising candidate effectors not previously identified. Encouragingly, many of them also appear to share interconnected functions and fell into three main groups (Figure 2D). A group of proteins involved in membrane shaping, cytoskeletal remodelling and in membrane-microtubule contacts included SPG20, BICD2, REEP4, CAMSAP1 and FAM134B. Of these, SPG20 was previously found to interact with RAB18 (Gillingham et al., 2014), BICD2 interacts with RAB18 among a number of Rab isoforms (Gillingham et al., 2019, Gillingham et al., 2014) and REEP4 was previously shown to interact with RAB3GAP (Tinti et al., 2012). Although not previously linked to RAB18, CAMSAP1 and FAM134B were each identified in both HEK293 and HeLa datasets. Next, a group of proteins involved in establishing membrane contacts included components of the NRZ/Dsl1 membrane-tethering complex, ZW10, RINT1 and NBAS, the Sec1/Munc18 (SM) protein SCFD2 and SNAP-REceptor (SNARE) proteins STX18 and BNIP1. These

272 proteins have been previously studied in the context of RAB18 (Gillingham et al., 273 2019, Gillingham et al., 2014, Li et al., 2019, Xu et al., 2018, Zhao and Imperiale, 274 2017). Also in this group were the SNARE protein homologue SEC22A and the ER-275 resident multispanning transmembrane protein WFS1, which regulates membrane 276 contacts between the ER and mitochondria (Angebault et al., 2018). These latter 277 proteins had not been linked to RAB18 previously. The third group of proteins 278 associated with RAB18 were a number of lipid transport, exchange and modifying 279 proteins, C2CD2L, C2CD2, ORP2/OSBPL2, INPP5B, EBP and TMCO4. 280 Interestingly, all but TMCO4 have related lipid species as their known substrates. 281 C2CD2 and C2CD2L are thought to mediate phosphatidylinositol (PI) transfer 282 between apposed membranes (Lees et al., 2017), ORP2 has recently been shown to 283 exchange PI(4,5)P₂ and cholesterol and INPP5B hydrolyses PI(4,5)P₂ to PI(4)P 284 (Wang et al., 2019). EBP is a D8-D7 Sterol Isomerase involved in cholesterol 285 biosynthesis (Silve et al., 1996). Various members of the OSBP family interact with 286 Rab proteins (Gillingham et al., 2019, Johansson et al., 2005), and a number of Rab 287 isoforms interact with INPP5B (Fukuda et al., 2008, Williams et al., 2007), though 288 associations with RAB18 have not been previously reported. 289 For initial validation of our HeLa dataset, we first carried out an additional 290 independent BioID experiment with wild-type and RAB3GAP1-null cells and 291 subjected the resulting samples to Western blotting for selected RAB18-associated 292 proteins (Figure 3A). As with the mass spectrometry, these proteins showed either 293 complete (RAB3GAP1, RAB3GAP2, ZW10) or partial (SPG20, STX18) dependence 294 on RAB3GAP for their RAB18 association. 295 Since RAB18 and its regulators are linked to Warburg Micro syndrome, we next 296 asked whether the putative RAB18 effectors were linked to any diseases with 297 overlapping clinical features (Figure 3B). Micro syndrome is a clinically distinctive 298 disorder characterised by intellectual disability (ID), postnatal microcephaly, brain 299 malformations, ascending spastic paraplegia, neuropathy, hypogonadism and eye 300 abnormalities that include congenital bilateral cataracts, microphthalmia, 301 microcornea and optic atrophy (Handley and Sheridan, 2018). Among the RAB3GAP-dependent RAB18 associations identified in the BioID screen, we noted 302 303 that several are encoded by disease-associated genes or their homologues. In 304 common with Micro syndrome, diseases linked to SPG20, BICD2, and the REEP4-

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homologues REEP1 and REEP2 are associated with ascending paraplegia (hereditary spastic paraplegia/HSP). FAM134B is associated with sensory and autonomic neuropathy, while the eye features of Micro syndrome overlap a different set of genes. EBP is linked to microphthalmia, microcornea and cataracts as well as to brain malformations like those in Micro syndrome. Both WFS1 and NBAS have been linked to conditions associated with optic atrophy, and both WFS1 and the INPP5B homologue OCRL1 are linked to conditions associated with congenital cataracts. Given the suggestive convergences in protein function and gene-diseaseassociations, we proceeded to examine the subcellular localizations of 12 putative effectors for which antibodies were available (Figure 3C-D). To determine whether the localization of these proteins was altered in cells lacking RAB18, we analysed wild type and RAB18-null lines in each case. In order to directly compare cells of different genotypes under otherwise identical conditions, we labelled them with CellTrace-Violet and CellTrace-Far Red reagents before seeding, immunostaining and imaging them together. Since RAB18 can localize to LDs, we analysed both untreated cells (Figure 3C) and cells loaded with oleic acid and labelled with BODIPY-558/568-C12 (Figure 3D). We observed a variety of staining patterns for the different putative effector proteins. These ranged from staining that was enriched at the perinuclear region of cells, to staining that appeared reticular, to staining that appeared more diffuse (Figure 3C). Each staining pattern was compatible with the known localization of RAB18, which is distributed between cis-Golgi, ER and cytosolic compartments (Handley et al., 2015). Staining patterns for individual proteins were similar in the HeLa cells and also in wild-type and RAB18-null RPE1 cells generated to provide biological replicates (Figure S4). In lipid-loaded cells, we observed that the localizations of proteins with reticular staining patterns overlapped with LDs, but they did not obviously shift to adopt an LD localization. However, the two proteins that showed the most diffuse staining patterns in untreated cells - ZW10 and SPG20 - appeared enriched in the vicinity of LDs in lipid-loaded cells (Figure 3D). We saw no evidence for dramatic changes in protein localizations in RAB18-null cells as compared to their wild-type counterparts. Fluorescence intensities in RAB18null and wild-type cells were also generally similar, except in the case of staining for

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SPG20, in which it appeared lower in RAB18-null HeLa cells than in wild-type cells (Figure 3C). Levels of SPG20 are significantly reduced in RAB18-null and TBC1D20-null cells In a RAB18-null mouse model of Warburg Micro syndrome, peripheral nerves have a disordered cytoskeleton, and there is a striking accumulation of microtubules at motor nerve terminals (Carpanini et al., 2014). Therefore, interactions between RAB18 and microtubule-binding proteins is of particular interest. To confirm the reduction in levels of SPG20 we observed in RAB18-null HeLa cells, we used quantitative fluorescence microscopy (Figure 4A-C). The SPG20 antibody used in this study has previously been used for this purpose (Nicholson et al., 2015). However, to confirm its specificity and also to determine the levels of non-specific background produced in our experiments, we first analysed SPG20-null cells (Figure 4A-B). Measured fluorescence intensity of SPG20-null cells provided a baseline level, above which fluorescence levels are proportional to levels of SPG20. In RAB18-null cells, SPG20 fluorescence was reduced to 67.16±3.77% (s.e.m., p<0.001) of that in wild-type cells (Figure 4C). To determine whether levels of SPG20 were altered by disrupted RAB18 regulation. we next compared SPG20 fluorescence of wild type, RAB3GAP1-, RAB3GAP2- and TBC1D20-null cells. Loss of the RAB18-GEF subunits RAB3GAP1 or RAB3GAP2 did not significantly affect levels of SPG20, whereas loss of the RAB18-GAP TBC1D20 led to a reduction comparable to that in RAB18-null cells (57.48%±2.57 (s.e.m., p<0.00005). To rule-out the possibility that reduced SPG20 levels in RAB18and TBC1D20-null HeLa cells were the result of clonal variation, we analysed the corresponding panel of RPE1 cell lines. However, because the RPE1 cells were less amenable to comparative immunofluorescence experiments than HeLa cells, we used LFQ analysis of whole cell lysates. As in the HeLa cells, levels of SPG20 were significantly reduced in RAB18- and TBC1D20-null RPE1 cells compared to wildtype controls (p<0.05 following FDR correction), but not in the other genotypes tested (Figure 4D, Table S3). RAB18, TBC1D20 and the RAB3GAP complex have all been linked to roles in proteostasis and autophagy (Bekbulat et al., 2018, Feldmann et al., 2017, Sidjanin et al., 2016, Spang et al., 2014). It was therefore possible that reduced SPG20 levels in RAB18- and TBC1D20-null cells were the result of widespread dysregulation of proteostasis. To assess this possibility, we compared LFQ data from wild-type and TBC1D20-null RPE1 and HeLa cells (Tables S3 and S4). Following FDR correction, only a small number of proteins showed significantly altered levels in each cell type and there was limited overlap between cell types. This is in-line with a recent study showing a compensatory mechanism maintains levels of basal autophagy when RAB18 is absent or dysregulated (Bekbulat et al., 2018). Thus, dysregulation of SPG20 levels most likely arises from a discrete mechanism. Discrete changes in SPG20 levels in RAB18- and TBC1D20-null cells together with the previous report of a RAB18-SPG20 interaction (Gillingham et al., 2014) provided strong evidence for a functional relationship between these proteins. We therefore carried out co-expression experiments to determine whether they colocalize in cells. Coexpression of mCherry-RAB18 and mEmerald-SPG20 in HeLa cells showed that, as previously reported, mCherry-RAB18 adopts a largely reticular localization (Gerondopoulos et al., 2014) whereas mEmerald-SPG20 appears largely diffuse (Eastman et al., 2009)(Figure 4E). We did not observe any clear colocalization between the proteins or any relocalization of mEmerald-SPG20 in cells expressing mCherry-RAB18. Since both RAB18 and SPG20 localize to LDs, we next explored whether the localization of either protein to LDs was dependent on the other. In oleic acid/BODIPY-558/568-C12-loaded HeLa cells, we found that mEmerald-SPG20 became concentrated around LDs, but that its localization was similar in both wildtype and RAB18-null cells (Figure 4F, left panels). Similarly, EGFP-RAB18 was enriched around LDs in both wild-type and SPG20-null cells (Figure 4F, right panels). Collectively, our findings suggest that RAB18 and SPG20 become localized to LDs independently, and that any RAB18-SPG20 interaction is likely to be transient. The reduced levels of SPG20 in RAB18- and TBC1D20-null cells seem likely to result from its reduced stability, and it is intriguing that a transient interaction could influence this. Nevertheless, this is consistent with the involvement of TBC1D20catalysed RAB18 GTP-hydrolysis and accompanying conformational change.

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402 SEC22A associates with RAB18 and its knockdown causes altered LD morphology 403 The most studied group of RAB18 effector proteins to date are the tethering factors ZW10, NBAS and RINT1, which together comprise the NRZ/Dsl1 complex 404 405 (Gillingham et al., 2014, Li et al., 2019, Xu et al., 2018, Zhao and Imperiale, 2017). 406 The NRZ complex regulates assembly of an ER SNARE complex containing STX18. 407 The canonical, fusogenic, form of this complex contains STX18, BNIP1, USE1 and 408 SEC22B (Spang, 2012, Tagaya et al., 2014). However, it has been proposed that 409 SEC22B is dispensable for the functions of RAB18, the NRZ complex, STX18, 410 BNIP1 and USE1 in regulating LDs (Xu et al., 2018). Rather than promoting 411 membrane fusion it is suggested that, in the absence of SEC22B, these proteins can 412 mediate the close apposition of membranes to facilitate lipid transfer. 413 In the HEK293 and HeLa BioID datasets, RAB18-associations with all three NRZ 414 components were nucleotide-binding-dependent and RAB3GAP-dependent 415 respectively (Figures 1B, 2D, Tables S1, S2). In the HeLa dataset, the ER SNARE 416 proteins STX18 and BNIP1 were also identified as associating with RAB18 in a 417 RAB3GAP-dependent manner (Figure 2D, Table S2). Interestingly, also among the 418 RAB3GAP-dependent RAB18 associations was SEC22A, a poorly studied 419 homologue of SEC22B. 420 SEC22A is one of two SEC22B homologues in humans, the other being SEC22C. 421 Like SEC22B, SEC22A and SEC22C possess N-terminal Longin domains and C-422 terminal transmembrane (TM) domains. However, they lack the central coiled-coil 423 SNARE domain through which SEC22B mediates membrane fusion as part of the 424 STX18 complex. SEC22B localizes to the ER-Golgi intermediate compartment. 425 whereas different isoforms of SEC22C localize to the ER or cis-Golgi (Yamamoto et 426 al., 2017, Ge et al., 2013, Zhang et al., 1999). The localization of SEC22A had not 427 been addressed. In the absence of commercially available antibodies for SEC22A, 428 we examined its localization through expression of an mEmerald-SEC22A fusion 429 protein (Figure 5A). mEmerald-SEC22A produced a characteristic reticular staining 430 pattern and colocalized with an exogenous ER marker suggesting that SEC22A 431 localizes to the ER. 432 We next sought to compare the localization of SEC22A and RAB18 and to determine whether they interact. However, coexpression of mEmerald-SEC22A and mCherry-433

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RAB18 appeared to disrupt normal ER morphology and to produce vesicular structures and inclusions positive for both proteins (Figure S5). Although this was not inconsistent with a protein-protein interaction, it precluded the use of coexpressed exogenous proteins to test such an interaction. We therefore used a BirA*-SEC22A fusion protein to verify the RAB18-SEC22A association, to identify other SEC22Aassociations, and to determine whether these associations were influenced by the absence of RAB18 or its regulators. To minimize potential toxicity while increasing biotin-ligase activity, we used BioID2 (Kim et al., 2016) with a p.Gly40Ser active site modification (Branon et al., 2018) and reduced biotin incubation time to 6 hours. LFQ analysis following streptavidin pull-down and mass spectrometry indicated that BioID2(Gly40S)-SEC22A was present at much lower levels than the BirA*-RAB18 in the prior experiments (3.79±0.96%). However, after adjusting for non-specific binding and normalizing the data according to the quantity of BioID2(Gly40S)-SEC22A in each sample, the construct appeared to label RAB18 in a RAB3GAP-dependent manner (Figure 5B). RAB3GAP1 and RAB3GAP2, as well as the putative RAB18associated proteins REEP4 and BICD2, were among 55 SEC22A-associated proteins present in samples from wild-type cells in >2 replicate experiments and represented by >3 spectral counts (Table S5). Furthermore, also among these proteins, a subset of 9 SEC22A-associations were attenuated (association ratios <0.5) in samples from both RAB18-null and RAB3GAP-null cells. Broadly, these data were consistent with a functional SEC22A-RAB18 interaction. Given the involvement of RAB18, the NRZ complex, and a STX18 complex lacking SEC22B in the regulation of LDs, we asked whether SEC22A might also be involved. Multiple studies have shown that LD morphology is altered in lipid-loaded cells in which RAB18 expression - or that of its regulators - is disrupted or silenced, with fewer and/or larger LDs are observed in these cells compared to controls (Bekbulat et al., 2018, Carpanini et al., 2014, Gerondopoulos et al., 2014, Li et al., 2017, Liegel et al., 2013, Xu et al., 2018). Similar observations have been made in ZW10-, NBAS-, STX18-, BNIP1- and USE1-null cells, whereas LD size distribution is unaltered when SEC22B expression is silenced (Xu et al., 2018). We examined the effects of silencing ZW10, NBAS and SEC22A in oleic acid-loaded induced human hepatocyte (IHH) cells (Figure 5C). ZW10 and NBAS silencing provided positive controls. ZW10 silencing led to a significant reduction in LD number (p<0.001) compared to controls,

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whereas NBAS silencing led to both a significant reduction in LD number and a significant increase in LD size (p<0.001 in each case). The effects of SEC22A silencing mirrored those of NBAS silencing, producing a significant reduction in LD number (p<0.001) and a significant increase in LD size (p<0.001). Together, these data implicate SEC22A in RAB18-mediated LD regulation. RAB18 recruits the orphan lipase TMCO4 to the ER membrane in a RAB3GAPdependent manner Among the lipid modifying/mobilizing proteins identified as potential RAB18 effectors in HeLa cells, TMCO4 was identified in all three replicate experiments and its association with RAB18 was highly RAB3GAP-dependent (association ratio 0.06). TMCO4 (transmembrane and coiled-coil domains 4) is annotated as containing transmembrane and coiled-coil domains, but is orthologous to the Yeast protein Mil1/ Yfl034w. Mil1 was found to be peripherally membrane-associated/soluble, and is thought to be an α/β hydrolase and a probable lipase based on structural modelling (Whitfield et al., 2016). Consistent with this, a catalytic triad within its predicted active site was shown to be required to confer tolerance to the membrane-intercalating cationic amphipathic drug sertraline (Whitfield et al., 2016). The best template match produced for TMCO4 by the structural modelling tool Phyre2 (Kelley et al., 2015) is an acylglycerol/diacyclglycerol lipase (Figure S6). To explore the localization of TMCO4, we expressed TMCO4-EGFP in HeLa cells (Figure 6). This construct showed a diffuse appearance consistent with a largely cytosolic localization. In contrast, EGFP-RAB18 partly localizes to the ER, as shown by its colocalization with an ER marker (Figure 6A). To assess the potential interaction between RAB18 and TMCO4, we coexpressed mCherry-RAB18 and TMCO4-EGFP (Figure 6B). As in our previous experiments, we used Celltrace reagents to distinguish cells of wild-type and mutant genotypes and imaged these on the same dishes. In wild-type HeLa cells, we found that the coexpression of mCherry-RAB18 led to a dramatic redistribution of TMCO4-EGFP to the ER membrane suggesting that RAB18 is involved in the recruitment of TMCO4 to this compartment (Figure 6B, upper panels). This redistribution was completely absent in RAB3GAP1- and RAB3GAP2-null cells but unaffected in TRAPPC9-null

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cells consistent with the BioID data. As a means of verifying the interaction, we carried out immunoprecipitation experiments using exogenous HA-RAB18 and TMCO4-EGFP (Figure 6C). As expected, TMCO4-EGFP copurified with HA-RAB18 when expressed in wild-type or TRAPPC9-null cells, but not when expressed in RAB3GAP1-null cells. These data indicate that RAB18 and TMCO4 interact directly or indirectly as part of a protein complex in a RAB3GAP-dependent manner. Further, both the microscopy and the immunoprecipitation data support the suggestion that different GEFs can promote different RAB18-interactions. RAB18 is involved in the mobilization of cholesterol The lipid-related RAB3GAP-dependent RAB18-associated proteins in the HeLa BioID dataset included EBP, which is involved in cholesterol biosynthesis (Silve et al., 1996). ORP2/OSBPL2 and INPP5B are also robustly linked to cholesterol, functioning in cholesterol mobilization. ORP2 is thought to function as a lipid transfer protein that delivers cholesterol to the plasma membrane (PM) in exchange for PI(4,5)P₂, whereas INPP5B is implicated in the hydrolysis of ORP2-bound PI(4,5)P₂, presumably driving the exchange process (Wang et al., 2019). Two other proteins in this group, C2CD2L/TMEM24 and C2CD2, have not been linked to cholesterol, but C2CD2L is characterised as a PI transporter and found to promote PM PI(4,5)P2 synthesis (Lees et al., 2017). Limited information is available about the function of TMCO4 in mammals. However, it was identified in one study as one of a number of genes with upregulated expression in rabbit cerebral arteries under conditions of hypercholesterolemia (Ong et al., 2013). Another study suggests that TMCO4 is present on lipid rafts (Jin et al., 2012). On the basis of these combined findings, we investigated the potential role of RAB18 in cholesterol uptake and efflux. We performed loading and efflux experiments to demonstrate the flux of cholesterol/cholesteryl ester (CE) based on the activity of RAB18. Chinese hamster ovary (CHO) cells were generated to stably express RAB18(WT), RAB18(Gln67Leu), or RAB18(Ser22Asn). These cells were then preincubated with lipoprotein deficient serum (LPDS) before addition of [14C]-oleate for 24 hours. [14C]-oleate was added in the presence of LPDS (Figure 7A, left panel) or 10% FBS (Figure 7A, right panel). [14C]-CE levels were measured at t=0, and efflux was assessed by measuring CE

531 levels 4 and 8 hours following the addition of high density lipoprotein (HDL) to the cells. In each case, [14C]-CE was isolated by thin layer chromatography quantified by 532 533 scintillation counting. Efflux was also assessed directly by loading the cells with [3H]-534 cholesterol, then incubating them with apolipoprotein (apo) A-I. [3H]-cholesterol 535 associated with apoA-I in the medium, as a percentage of total cellular radioactivity. 536 is shown in Figure 7B. 537 In cells loaded with [14C]-oleate/LPDS, levels of CE were comparable in 538 RAB18(Ser22Asn) and RAB18(WT) cells, whereas RAB18(Gln67Leu) cells stored 539 significantly more (Figure 7A, left panel). In cells loaded with [14C]-oleate/FBS, levels 540 of CE in RAB18(Ser22Asn) remained unchanged, whereas storage its storage was 541 elevated in RAB18(WT) cells and RAB18(Gln67Leu) cells (Figure 7A, right panel). 542 Interestingly, in both [14C]-oleate/LPDS-loaded and [14C]-oleate/FBS-loaded cells, 543 the addition of HDL led to rapid depletion of CE in RAB18(Gln67Leu) cells, but not in 544 RAB18(Ser22Asn) or RAB18(WT) cells (Figure 7A). Consistently, [3H]-cholesterol 545 also underwent significantly more rapid efflux from these cells (Figure 7B). 546 Collectively, these data indicate that 'activated' GTP-bound RAB18 strongly 547 promotes the turnover and mobilization of CE. 548

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DISCUSSION In this study, we have complemented previous work showing that proximity biotinylation is a powerful means of identifying candidate Rab effectors (Gillingham et al., 2019). We have shown that comparisons between wild-type and nucleotidebinding deficient BirA*-RAB18 fusions can help to distinguish putative effector proteins, recapitulating findings from conventional affinity purification. Further, we have found that comparing biotin-labelling produced by a BirA*-Rab in wild-type and GEF-deficient cells can be particularly informative. For RAB18, we found that the disruption of different GEF complexes affected largely distinct subsets of its associations. Encouragingly, in the case of RAB3GAP-null cells, we found that marked changes in RAB18-associations were restricted to a relatively small number of proteins and that these comprised known and/or plausible interactors. By identifying these, we were able to exclude ~95% of RAB18-associations from consideration as more likely to represent 'noise' from bystander proteins. The limited overlap between RAB3GAP- and TRAPPC9-dependent RAB18 associations is interesting because it may indicate that different RAB18-GEFs control different aspects of RAB18 function. However, the large number of RAB18associations affected in TRAPPC9-null cells may not all represent functional interactions. Loss of TRAPPC9 does not appear to affect RAB18 localization (Figure S2C). Nevertheless, it could affect RAB18-associations indirectly. TRAPP complexes regulate multiple Rab isoforms, and so broad disruption of membrane trafficking

For validation of our BioID screening, we focused on the subset of RAB3GAP-dependent candidate RAB18 effectors. These 25 proteins largely fell into three apparently interrelated functional groups: microtubule-interacting/membrane-remodelling proteins; proteins involved in bringing membranes into close apposition; and proteins involved in lipid modification and mobilization. It has been proposed that RAB18 functions to coordinate lipid transfer between apposed membranes, and our data are consistent with this model (Xu et al., 2018). Though not definitively

might alter the compliment of RAB18-proximal proteins. Nevertheless, it seems likely

that genuine TRAPPII-dependent RAB18-associations are represented in our

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demonstrated in the present study, it seems likely that membrane remodelling precedes the engagement of tethers and SNAREs and the establishment of membrane-contacts. Lipid transporter/exchange proteins are required for lipid transfer to occur at these contact points, while lipid-modifying proteins may serve to generate concentration gradients necessary to promote directional flow. We have presented preliminary findings supporting functional interactions between RAB18 and one protein from each group; two of these interactors, SEC22A and TMCO4, had not been reported previously. In initial screening of 12 of the candidate effectors by IF, we observed that levels of SPG20 were reduced in the absence of RAB18 or TBC1D20 in two different cell types. SPG20 is a protein of enigmatic function, having been ascribed roles in regulating microtubule stability and bone morphogenic protein (BMP) signalling, chromosome segregation and cytokinesis, LD turnover and ubiquitylation of LD proteins, and in mitochondrial energy metabolism and calcium homeostasis (Eastman et al., 2009, Edwards et al., 2009, Joshi and Bakowska, 2011, Lind et al., 2011, Nahm et al., 2013, Nicholson et al., 2015, Renvoise et al., 2012, Ring et al., 2017, Tsang et al., 2009). Of particular interest in the context of RAB18 and its association with Warburg Micro syndrome, biallelic loss-of-function variants in SPG20 cause Troyer syndrome, which has overlapping features including ID and ascending paraplegia (Baple and Crosby, 1993). One possible avenue for future research could be the potential relationship between SPG20 and the other RAB18associated microtubule (MT)-binding proteins identified in this study. Like SPG20, CAMSAP1 is found to associate with poles of the mitotic spindle (Hueschen et al., 2017, Lind et al., 2011). CAMSAP1 is a MT minus-end binding protein whereas BICD2 is a component of a minus-end-directed dynein-dynactin motor complex (Hendershott and Vale, 2014, Hueschen et al., 2017, Urnavicius et al., 2015). REEP4 is localized to the ER, and like RAB18 is linked to the regulation of ER structure (Kumar et al., 2019, Gerondopoulos et al., 2014). However, it also contributes to the clearance of ER from metaphase chromatin (Schlaitz et al., 2013). This role is dependent on its MT-binding, and therefore spindle-binding during metaphase. Thus, these four MT-binding proteins all function in the same spatiotemporally defined compartment during mitosis.

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Among the membrane tethering and docking proteins associated with RAB18, we investigated SEC22A on the basis of its homology to SEC22B. SEC22B is a component of the canonical syntaxin 18 SNARE complex, and other components of this complex interact with RAB18 to regulate close apposition of membranes, though SEC22B itself is not necessary for this activity (Xu et al., 2018). Reciprocal proximity biotinylation with a BioID2-SEC22A fusion was consistent with a RAB3GAPdependent RAB18-SEC22A interaction, and suggested that other SEC22A interactions are altered when RAB18 is absent or dysregulated. We also found that the silencing of SEC22A led to altered LD morphology, similar to that previously observed when expression of other RAB18-associated tethering/docking proteins was disrupted. It is tempting to speculate that SEC22A contributes to a nonfusogenic SNARE complex, since it lacks the SNARE domain usually required to provide the mechanical force for fusion. SNARE proteins can be relatively promiscuous in their interactions, and there are numerous examples of individual SNAREs contributing to distinct SNARE complexes (Wang et al., 2017, Petkovic et al., 2014). There are also examples in which SNARE complexes mediate the stable association of membranes rather than their fusion (Petkovic et al., 2014). Rab proteins, SM proteins and tethering factors can contribute to compartmental specificity and the assembly of specific SNARE complexes. In Yeast, the RAB1 orthologue Ypt1p and the SM protein Sly1p are implicated in the assembly of the canonical syntaxin 18 (Ufe1p) SNARE complex (Kamena et al., 2008, Yamaquchi et al., 2002, Dilcher et al., 2003, Lewis and Pelham, 1996, Lewis et al., 1997). In mammals, multiple Rab isoform(s) and the Sly1p orthologue SCFD1 may fulfil this role (Nakajima et al., 2004, Hirose et al., 2004, Galea et al., 2015, Tagaya et al., 2014). The suggestion that RAB18 regulates assembly of a distinct SNARE complex (Xu et al., 2018) is consistent both with our identification of SEC22A as a RAB18associated protein, and also with our finding that SCFD2 but not SCFD1 associates with RAB18 in a RAB3GAP-dependent manner. SCFD2 is a poorly characterised Sly1p orthologue, but dramatic effects of SCFD1 silencing on the secretory pathway suggest that SCFD1 and SCFD2 do not function redundantly (Bassaganyas et al., 2019). Although we did not validate the RAB18-SCFD2 association directly, it has recently been reported in another study (Gillingham et al., 2019).

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The lipid modifying/mobilizing proteins that we found associated with RAB18 included the poorly characterised orphan lipase TMCO4. We have presented clear evidence for the RAB3GAP-dependent recruitment of TMCO4 to the ER by RAB18, further validating the BioID approach and dataset. We have not yet conclusively proven the link between RAB18 and the other lipid-related proteins in this group. ORP2, INPP5B, C2CD2, C2CD2L and EBP. Any conjecture based on these prospective interactions therefore remains tentative. Nevertheless, we have shown that RAB18 influences cellular cholesterol mobilization, a process in which TMCO4 may be involved, and in which ORP2 and INPP5B appear to have a central role (Lees et al., 2017). ORP2 is thought to act as cholesterol-PI(4,5)P2 exchanger, whereas INPP5B is thought to drive the exchange process by catalysing PI(4,5)P2 hydrolysis. Therefore, these proteins are strong candidates for future investigation. Interestingly, INPP5B has been shown in several studies to have a relatively broad Rab-binding specificity (Fukuda et al., 2008, Williams et al., 2007). Further, many other phosphoinositide phosphatase enzymes, including members from distinct protein families, bind to Rab proteins (Gillingham et al., 2019). Multiple OSBPhomologues also bind to Rabs (Gillingham et al., 2019, Rocha et al., 2009, Sobajima et al., 2018), raising the possibility that phosphoinositide-unloading from OSBPs constitutes a conserved role for RAB proteins generally: different Rab isoforms may specify intermembrane transfer of different lipid species by association with different combinations of phosphatase and OSBP proteins. One interesting possibility we are currently pursuing is that RAB18 may contribute to cholesterol biosynthesis though the delivery of substrate to the EBP enzyme, or through the relocalization of the lipid species it generates. To summarise, we have used complementary proximity ligation approaches together with CRISPR gene-editing to inventory RAB18-associated proteins. The RAB3GAPdependent RAB18-associations we have highlighted are supported by additional validation of three examples, as well as by correlative information from the literature. Broadly, the latter suggests a role for Rab proteins distinct from that in mediating vesicular membrane traffic. Another role of Rabs may be to assemble a network of diverse membrane-membrane contact sites and then orchestrate highly regulated metabolism and flow of lipid species within a discontinuous lipid phase. We have shown that RAB18 influences cholesterol mobilization, and priorities for future work

will be to categorically define the molecular mechanism involved and to determine whether altered cholesterol flux underlies the pathology of Warburg Micro syndrome. Since several of the putative RAB18 effectors we have uncovered are also encoded by disease-associated genes, a further aim will be to determine the extent to which the molecular pathology of these conditions overlap.

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MATERIALS AND METHODS <u>Plasmids</u> Generation of recombinant pcDNA5 FRT/TO FLAG-BirA(Arg118Gly) vectors for preparation of stable T-Rex-293 cell lines is described below. Generation of recombinant pX461 and pX462 plasmids for CRISPR gene-editing is described below. Generation of recombinant pCMV vectors for preparation of stable CHO cell lines is described below. The EGFP-RAB18 construct has been described previously (Gerondopoulos et al., 2014). The RAB18 sequence was excised from this construct using BamHI and HindIII restriction enzymes (New England Biolabs, Hitchin, UK), and used to generate constructs encoding mEmerald-RAB18 and mCherry-RAB18 by ligation into mEmerald-C1 and mCherry-C1 vectors (Addgene, Watertown, MA) using HC T4 Ligase and rapid ligation buffer (Promega, Southampton, UK). Constructs encoding BirA*-RAB18, BioID2(Gly40Ser)-SEC22A, mEmerald-SEC22A, mEmerald-SPG20 and mEmerald-RAB7A were generated following PCR amplification from template and subcloning into an intermediate pCR-Blunt II-TOPO vector using a Zero Blunt TOPO PCR Cloning Kit (ThermoFisher Scientific, Waltham, MA) according to manufacturer's instructions. Fragments were excised from intermediate vectors and then subcloned into target vectors using restrictionligation, as above. A construct encoding mCherry-ER was obtained from Addgene, and a construct encoding TMCO4-EGFP was synthesised and cloned by GeneWiz (Leipzig, Germany). Details of PCR templates, primers and target vectors are listed in Table S6. Antibodies and reagents A custom polyclonal antibody to RAB18 generated by Eurogentec (Southampton, UK) has been described previously (Handley et al., 2015). An antibody to RAB3GAP1 was obtained from Bethyl Labs (Montgomery, TX), an antibody to GFP was obtained from Takara Bio (Saint-Germain-en-Laye, France), an antibody to β-Tubulin was obtained from Abcam (Cambridge, UK) and an antibody to β-Actin was obtained from ThermoFisher. Antibodies to hemagglutinin (HA), RAB3GAP2 and TBC1D20 were obtained from Merck (Gillingham, UK). Antibodies to ZW10, STX18,

718 SPG20, RINT1, REEP4, BNIP1, C2CD2, TRIM13, WFS1, INPP5B, OSBPL2 and 719 NBAS were obtained from Proteintech (Manchester, UK). Antibody catalogue 720 numbers and the dilutions used in this study are listed in Table S6. 721 722 Cell culture 723 T-REx-293, HeLa and IHH cells were maintained in DMEM media, RPE1 cells in 724 DMEM/F12 media and CHO cells in alpha-MEM media (ThermoFisher). In each 725 case, media was supplemented with 10% foetal calf serum (FCS) and 1% penicillin-726 streptomycin (PS). Cells were maintained at 37°C and 5% CO₂. 727 728 Generation of stable T-Rex-293 and CHO cell lines 729 PCR products encoding mouse RAB18, RAB18(Gln67Leu) and RAB18(Ser22Asn) 730 were subcloned into Notl-linearized pcDNA5 FRT/TO FLAG-BirA(Arg118Gly) vector 731 using the In-Fusion HD EcoDry Cloning Plus kit (Takara Bio) according to 732 manufacturer's instructions. Details of PCR templates, primers and target vectors are 733 listed in Table S6. 1.5ug of each recombinant vector together with 13.5ug of pOG44 734 plasmid (ThermoFisher) were used in cotransfections of T-REx-293 cells. in 10cm 735 dishes, with TransIT-LT1 Transfection Reagent (Mirus Bio, Madison, WI).16 hours 736 following transfection, media was replaced and cells were allowed to recover for 24 737 hours. Each dish was then split to 4x 10cm dishes in selection media containing 10 738 ug/ml Blasticidin and 50 ug/ml Hygromycin B. Resistant clones were pooled and 739 passaged once prior to use. 740 A PCR product encoding mouse RAB18 was subcloned into an intermediate TOPO 741 vector using a TOPO PCR Cloning Kit (ThermoFisher) according to manufacturer's 742 instructions. The RAB18 fragment was then excised and subcloned into the pCMV 743 vector. PCR-based site-directed mutagenesis using a GeneArt kit (ThermoFisher) 744 was then used to generate pCMV-RAB18(Gln67Leu) and pCMV-RAB18(Ser22Asn) 745 constructs. CHO cells were transfected using Lipofectamine 2000 reagent 746 (ThermoFisher) and cells stably-expressing each construct were selected-for with 747 blasticidin. Under continued selection, clonal cell-lines were grown from single cells 748 and then RAB18 protein expression was assessed. Cell lines comparably expressing

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RAB18 constructs at levels 2.5-5x higher than those wild-type cells were used in subsequent experiments. Generation of clonal 'knockout' HeLa and RPE1 cell lines CRISPR/Cas9 gene-editing was carried out essentially as described in Ran et al., 2013 (Ran et al., 2013). Guide RNA (gRNA) sequences are shown in (Table S6). A list of the clonal cell lines generated for this study, together with the loss-of-function variants they carry is shown in (Figure S1A). Western blot validation is shown in (Figure S1B-E). Briefly, for each targeted exon, pairs of gRNA sequences were selected using the online CRISPR design tool (http://crispr.mit.edu/). Oligonucleotide pairs incorporating these sequences (Sigma) were annealed (at 50mM ea.) in 10mM Tris pH8, 50mM NaCl and 1mM EDTA by incubation at 95°C for 10 minutes followed by cooling to room temperature. Annealed oligonucleotides were diluted and ligated into BbsI-digested pX461 and pX462 plasmids (Addgene) using HC T4 Ligase and rapid ligation buffer (Promega). Sequences of all recombinant plasmids were verified by direct sequencing. Pairs of plasmids were contransfected into cells using Lipofectamine 2000 reagent according to manufacturer's instructions. Cells were selected for puromycin resistance (conferred by pX462) using 24 hours puromycintreatment. Following 12 hours recovery, they were selected for GFP fluorescence (conferred by pX461) and cloned using FACSAria2 SORP, Influx or FACSMelody instruments (BD, Wokingham, UK). After sufficient growth, clones were analysed by PCR of the targeted exons (Primers are listed in Table S6). In order to sequence individual gene edited-alleles, PCR products from each clone were first cloned into ZeroBlunt TOPO vector (ThermoFisher) and then subjected to colony PCR. These PCR products were then analysed by direct sequencing. Sequencing data was assessed using BioEdit software (http://www.mbio.ncsu.edu/BioEdit/bioedit.html). BirA/BioID proximity labelling (T-REx-293 cells) The T-REx-293 Cell Lines (described above) were seeded onto 3x 15cm plates each and allowed to adhere. Expression of BirA*-RAB18 fusion proteins was induced by treatment with 20ng/ml Tetracycline for 16 hours. Media was then replaced with media containing 20% FBS, 20 ng/ml Tetracycline and 50 uM Biotin and the cells

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were incubated for a further 8 hours, washed with warmed PBS and pelleted in ice-782 cold PBS. Cell pellets were snap-frozen and stored at -80°C prior to lysis. Lysis was 783 carried out in 3ml of ice-cold RIPA buffer (150 mM NaCl, 1% NP40, 0.5% Sodium Deoxycholate, 0.1% SDS, 1mM EDTA, 50mM Tris, pH 7.4) supplemented with 784 785 complete-mini protease inhibitor cocktail (Roche, Basel, Switzerland), 1mM PMSF, and 62.5 U/ml Benzonase (Merck). Lysates were incubated for 1 hour at 4°C then 786 787 sonicated in an ice bath (four 10 second bursts on low power). They were then 788 clarified by centrifugation, and the supernatants transferred to tubes containing pre-789 washed streptavidin-sepharose (30µl bed-volume)(Merck). The beads were 790 incubated for 3 hours at 4°C, then washed x5 in RIPA buffer and x4 in buffer containing 100mM NaCl, 0.025% SDS and 25 mM Tris, pH7.4. 792 793 BirA*/BioID proximity labelling (HeLa cells) 794 Proximity-labelling in HeLa cells was carried out largely as described by Roux et al. 795 (Roux et al., 2012), but with minor modifications. HeLa cells were grown to 80% 796 confluence in T75 flasks and then each flask was transfected with 1-1.5µg of the 797 BirA*-RAB18 construct or 1µg of the BioID2(Gly40Ser)-SEC22A construct using 798 Lipofectamine 2000 reagent in Optimem serum-free medium (ThermoFisher) for 4 799 hours, according to manufacturer's instructions. 24 hours post-transfection, media 800 was replaced with fresh media containing 50µM Biotin (Merck) and the cells were incubated for a further 24 or 6 hours (for BirA*-RAB18 and BioID2(Gly40Ser)-802 SEC22A experiments respectively). Cells were then trypsinised and washed twice in 803 PBS before pellets were transferred to 2ml microcentrifuge tubes and snap-frozen. 804 For each pellet, lysis was carried out in 420µl of a buffer containing 0.2% SDS, 6% 805 Triton-X-100, 500mM NaCl, 1mM DTT, EDTA-free protease-inhibitor solution 806 (Expedeon, Cambridge, UK), 50mM Tris pH7.4. Lysates were sonicated for 10 807 minutes using a Bioruptor device together with protein extraction beads (Diagenode, 808 Denville, NJ). Each lysate was diluted with 1080µl 50mM Tris pH7.4, and they were 809 then clarified by centrifugation at 20 000xg for 30 minutes at 4°C. Affinity purification 810 of biotinylated proteins was carried out by incubation of clarified lysates with streptavidin-coated magnetic Dynabeads (ThermoFisher) for 24 hours at 4°C. Note that a mixture of Dynabeads - MyOne C1, MyOne T1, M270 and M280 - was used 812

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to overcome a problem with bead-clumping observed when MyOne C1 beads were used alone. Successive washes were carried out at room temperature with 2% SDS, a buffer containing 1% Triton-X-100, 1mM EDTA, 500mM NaCl, 50mM HEPES pH7.5, a buffer containing 0.5% NP40, 1mM EDTA, 250mM LiCl, 10mM Tris pH7.4, 50mM Tris pH7.4 and 50mM ammonium bicarbonate. Preparation of cell lysates for label-free quantitative proteomics RPE1 and HeLa cells were grown to confluence in T75 flasks. They were then trypsinised, and cell pellets were washed with PBS and snap-frozen prior to use. RPE1 pellets were resuspended in 300µl 6M GnHCl, 75mM Tris, pH=8.5. HeLa pellets were resuspended in 300µl 8M urea, 75mM NaCl, 50mM Tris, pH=8.4. In each case, samples were sonicated for 10 minutes using a Bioruptor device together with protein extraction beads (Diagenode). RPE1 samples were heated for 5 minutes at 95°C. Samples were clarified by centrifugation. Mass spectrometry Washed beads from BioID experiments with T-Rex-293 cell lines were resuspended in 50µl 6M urea, 2M thiourea, 10mM Tris, pH=8.5 and DTT was added to 1mM. After 30 minutes incubation at 37°C, samples were alkylated with 5mM iodoacetamide (IAA) in the dark for 20minutes. DTT was increased to 5mM and 1µg lysC was added, then samples were incubated at 37°C for 6 hours. Samples were diluted to 1.4M urea, then digested with trypsin (Promega), overnight at 37°C, according to manufacturer's instructions. Samples were acidified by the addition of 0.9% formic acid and 5% acetonitrile. Washed beads from BioID experiments with HeLa cells were subjected to limited proteolysis by trypsin (0.3 ug) at 27°C for 6.5hours in 2mM urea, 1mM DTT, 75mM Tris, pH=8.5, then supernatants were incubated overnight at 37°C. Samples were alkylated with 50mM IAA in the dark for 20minutes, then acidified by addition of 8ul 10% trifluoroacetic acid (TFA). RPE1 lysates were reduced and alkylated through addition of tris(2-carboxyethyl)phosphine (TCEP) and 2-chloroacetamide (CAA) to 5mM and 10mM respectively and then incubated at 95°C for 5 minutes. After

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cooling, samples were diluted to 3M quanidine and 0.5µg lysC added with incubation overnight at 37°C. A further dilution to 1M guanidine was followed by digest with 0.3µg trypsin at 37°C for 4 hours. Samples were acidified with TFA. HeLa lysates were reduced and alkylated by addition of DTT to 10mM, then by addition of IAA to 25mM, then further addition of DTT to 25mM, with incubation at room temperature for 30-60 minutes following each step. Samples were digested with lvsC, overnight at 37°C. They were then diluted to 2M urea, and further digested, overnight at 37°C. Samples were acidified with TFA. For BioID samples from T-REx-293 cells, LC-MS was carried out as previously described (Brunet et al., 2016). Briefly, peptides in an aqueous solution containing 5% acetonitrile and 0.1% formic acid were loaded onto a 3 µm PepMap100, 2 cm, 75 µm diameter sample column using an Easy nLC 1000 ultrahigh pressure liquid chromatography system (ThermoFisher). They were eluted with acetonitrile/formic acid into an in-line 50 cm separating column (2 µm PepMap C18, 75 µm diameter) at 40°C. Separated peptides were ionized using an Easy Spray nano source and subjected to MS/MS analysis using a Velos Orbitrap instrument (ThermoFisher). Following acquisition, data were analysed using SEAQUEST software. For other samples, peptides were loaded on to activated (methanol), equilibrated (0.1% TFA) C18 stage tips before being washed with 0.1% TFA and eluted with 0.1% TFA/80 acetonitrile. The organic was dried off, 0.1% TFA added to 15 µl and 5 ul injected onto LC-MS. Peptides were separated on an Ultimate nano HPLC instrument (ThermoFisher), and analysed on either an Orbitrap Lumos or a Q Exactive Plus instrument (ThermoFisher). After data-dependent acquisition of HCD fragmentation spectra, data were analysed using MaxQuant and the uniprot human reference proteome. Versions, releases, parameters and gradients used for separation are provided in table S6. Cell labelling In order to distinguish cells of different genotypes within the same well/on the same coverslip, CellTrace Violet and CellTrace Far Red reagents (ThermoFisher) were used to label cells before they were seeded. Cells of different genotypes were first trypsinised and washed with PBS separately. They were then stained in suspension

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by incubation with either 1µM CellTrace Violet or 200nM CellTrace Far Red for 20 minutes at 37°C. Remaining dye was removed by addition of a ten-fold excess of full media, incubation for a further 5 minutes, and then by centrifugation and resuspension of the resulting pellets in fresh media. Differently-labelled cells were combined prior to seeding. Immunofluorescence microscopy HeLa or RPE1 cells were seeded in 96-well glass-bottom plates (PerkinElmer, Waltham, MA) coated with Matrigel (Corning, Amsterdam, Netherlands) according to manufacturer's instructions, and allowed to adhere for 48 hours prior to fixation. In lipid-loading experiments, cells were treated with 200µM oleic acid complexed to albumin (Merck) and 1µg/ml BODIPY-558/568-C12 (ThermoFisher) for 15 hours prior to fixation. Cells were fixed using a solution of 3% deionised Glyoxal, 20% EtOH, 0.75% acetic acid, pH=5 (Richter et al., 2018), for 20 minutes at room temperature. They were then washed with PBS containing 0.9mM CaCl₂ and 0.5mM MgCl₂ and blocked with a sterile-filtered buffer containing 1% Milk, 2% donkey serum (Merck), 0.05% Triton-X-100 (Merck), 0.9mM CaCl₂ and 0.5mM MqCl₂ in PBS pH=7.4 for at least 1 hour prior to incubation with primary antibody. Primary antibodies were added in blocking buffer without Triton-X-100, and plates were incubated overnight at 4°C. Antibody dilutions are listed in Table S6. Following washing in PBS, cells were incubated with 1:2000 Alexa 488-conjugated secondary antibody (ThermoFisher) in blocking buffer at room temperature for 1-2 hours. Following further washing in PBS, cells were imaged using an Operetta High Content Imaging System (PerkinElmer) equipped with Harmony software. In comparative fluorescence quantitation experiments, at least 18 frames – each containing >5 wildtype and >5 mutant cells – were analysed per genotype. ImageJ software was used to produce regions of interest (ROIs) corresponding to each cell using thresholding tools and images from the 405nm and 645nm channels. Median 490nm fluorescence intensity was measured for each cell and mutant fluorescence intensity (as %wild-

Confocal microscopy – Live cell imaging

type) was calculated for each frame and combined for each genotype.

HeLa or RPE1 cells were seeded on glass-bottom dishes (World Precision Instruments, Hitchin, UK) coated with Matrigel (Corning) and allowed to adhere for 24 hours prior to transfection. Transfections and cotransfections were carried out with 0.5µg of each of the indicated constructs using Lipofectamine 2000 reagent in Optimem serum-free medium for 4 hours, according to manufacturer's instructions. Media were replaced and cells were allowed to recover for at least 18 hours prior to imaging. Imaging was carried out on a Nikon A1R confocal microscope equipped with the Nikon Perfect Focus System using a 60x oil immersion objective with a 1.4 numerical aperture. In immunofluorescence experiments, the pinhole was set to airy1. CellTrace Violet was excited using a 403.5nm laser, and emitted light was collected at 425–475nm. EGFP and mEmerald were excited using a 488 nm laser, and emitted light was collected at 500-550 nm. BODIPY-558/568-C12 and mCherry were excited using a 561.3 nm laser, and emitted light was collected at 570-620 nm. CellTrace Far Red was excited using a 638nm laser, and emitted light was collected at 663-738nm. In fluorescence recovery after photobleaching (FRAP) experiments, the pinhole was set to airy2 and digital zoom parameters were kept constant. Bleaching was carried out using 90% laser power.

<u>Immunoprecipitation</u>

HeLa cells were seeded onto 10cm dishes and allowed to adhere for 24 hours prior to transfection. Transfections and cotransfections were carried out with 0.5µg of each of the indicated constructs using Lipofectamine 2000 reagent in Optimem serum-free medium for 4 hours, according to manufacturer's instructions. 24 hours post-transfection cells were trypsinised, washed with PBS, then lysed in a buffer containing 150mM NaCl, 0.5% Triton-X-100 and EDTA-free protease-inhibitor solution (Expedeon), 10mM Tris, pH=7.4. Lysates were clarified by centrifugation, input samples taken, and the remaining supernatants then added to 4µg rabbit anti-HA antibody (Merck). After 30 minutes incubation at 4°C on a rotator, 100µl washed protein G-coupled Dynabeads (ThermoFisher) were added and samples were incubated for a further 1 hour. The Dynabeads were washed x3 with buffer containing 150mM NaCl, 0.1% Triton-X-100, 10mM Tris, pH=7.4, then combined with a reducing loading buffer and subjected to SDS–PAGE.

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<u>Lipid loading experiments</u> For LD number and diameter measurements, IHH cells were seeded onto glass coverslips. siRNA transfections were carried out using FuGene reagent (Promega) according to manufacturer's instructions. siRNAs targeting ZW10 and NBAS were obtained from IDT, Coralville, IA; siRNA targeting SEC22A was obtained from Horizon Discovery, Cambridge, UK. 48 hours following transfection, cells were treated with 200nM BSA conjugated oleate for 24 hours. Coverslips were washed, fixed with 3% paraformaldehyde and stained with 1ug/mL BODIPY and 300nM DAPI. Fluorescence images were captured on a Zeiss LSM 780 confocal microscope equipped with a 100x objective. Images were analysed using ImageJ software. Data are derived from measurements from >100 cells/condition and are representative of three independent experiments. For cholesterol storage and efflux experiments with [14C]-oleate, CHO cell lines (described above) were seeded onto 12-well plates and then grown to 60-75% confluence in Alpha media supplemented with 10% LPDS. Cells were grown in the presence of 10% LPDS for at least 24 hours prior to the addition of oleate. 1 µCi/ml [14C]-oleate (Perkin Elmer) was added in the presence of 10% LPDS or 10% FBS for 24 hours. Cells were then washed and incubated with 50µg/ml HDL for 0, 4 or 8 hours. Cellular lipids were extracted with hexane. Lipids were then dried-down and separated by thin layer chromatography (TLC) in a hexane: diethyl ether: acetic acid (80:20:2) solvent system. TLC plates were obtained from Analtech, Newark, NJ. Bands corresponding to cholesteryl ester (CE) were scraped from the TLC plate, and radioactivity was determined by scintillation counting in a Beckman Coulter LS6500 Scintillation Counter using BetaMax ES Liquid Scintillation Cocktail (ThermoFisher). Three independent experiments were carried out, each with four replicates of each condition. Data from a representative experiment are shown. For cholesterol efflux experiments with [3H]-cholesterol, CHO cells were seeded onto 12-well plates and then grown to 60% confluence in Alpha media supplemented with 10% FBS. 5 µCi/ml [3H]-cholesterol (PerkinElmer) was added in the presence of 10% FBS. After 3x PBS washes, cells were incubated with serum-free media containing 25µg/ml of human apolipoprotein A-I (ApoA-I) for 5 hours. ApoA-I was a kind gift of Dr. Paul Weers (California State University, Long Beach). Radioactivity in aliquots of

972 media were determined by scintillation counting in a Beckman Coulter LS6500 973 Scintillation Counter using LSC Cocktail (PerkinElmer). Cell lysates were produced 974 by addition of 0.1N NaOH for 1 hour, and their radioactivity was determined as 975 above. Cholesterol efflux was calculated as an average (+/- SD) of the % cholesterol 976 efflux (as a ratio of the media cpm/(media + cellular cpm) x 100%). 977 Western blotting 978 Cell lysates were made with a buffer containing 150mM NaCl, 0.5% Triton-X-100 979 and EDTA-free protease-inhibitor solution (Expedeon), 50mM Tris, pH=7.4. Cell 980 lysates and input samples from BioID and immunoprecipitation experiments were 981 combined 1:1 with a 2x reducing loading buffer; a reducing loading buffer containing 982 10mM EDTA was added directly to Dynabead samples. SDS-PAGE and Western 983 blotting were carried out according to standard methods. 984 985 **ACKNOWLEDGEMENTS** 986 We thank the Warburg Micro syndrome children and their families. We thank 987 Professor C. A. Johnson and Dr J. A. Poulter for a critical reading of the manuscript. 988 989 **COMPETING INTERESTS** 990 No competing interests declared. 991 992 FUNDING 993 MH is supported by the University of Leeds and by a Programme Grant from the 994 Newlife Foundation for Disabled Children (Grant Reference Number: 17-18/23). 995 996 DATA AVAILABILITY 997 The mass spectrometry proteomics data have been deposited to the 998 ProteomeXchange Consortium via the PRIDE (Perez-Riverol et al., 2019) partner 999 repository with the dataset identifiers PXD016631, PXD016336, PXD016326, 1000 PXD016233 and PXD016404.

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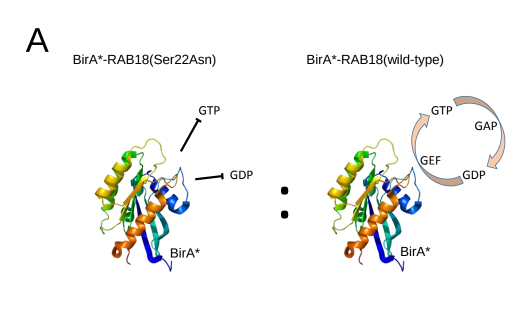
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wild-type HEK293 cells

wild-type HEK293 cells

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Protein	Ratio	Orthologue PPI (Gillingham et al., 2014)	Additional evidence	_
SCFD2 ZW10 NBAS CAMSAP1 FAM134B	0.00 0.00 0.00 0.04 0.36	Slh mit(1)15 rod	Gillingham et al., 2019 Xu et al., 2018; Gillingham et al., 2019 Xu et al., 2018; Gillingham et al., 2019	RAB3GAP- dependent association in HeLa
RINT1 RAB3GAP2 RAB3GAP1	0.36 0.41 1.58 1.59	CG8605 rab3-GAP CG31935	Xu et al., 2018 Gerondopoulos et al., 2014 Gerondopoulos et al., 2014	cells
TBC1D5 c15orf38 SEC23IP ATP6AP2 GORASP2 GIGYF2 TPR	0.00 0.00 0.00 0.29 0.36 0.41 0.41	CG8449	Gillingham et al., 2019	TRAPPII-dependent association in HeLa cells
USP15 TMPO ARFGAP3 ARFGAP2 SLK NUP153 ATG2B SCFD1 USE1 TMX1 CORO1B PREB	0.00 0.00 0.20 0.31 0.34 0.39 0.43 0.45 0.45 0.45	Slh	Gillingham et al., 2019 Xu et al., 2018	Present in HeLa dataset (n≥2)
SLC25A4 LUZP1 DPYSL2	0.00 0.00 0.34		Nakamura et al., 2016	Absent in HeLa dataset (n≥2)

Figure 1. Nucleotide-binding-dependent RAB18-associations in HEK293 cells.

(A) Schematic to show BirA*-RAB18(Ser22Asn):BirA*-RAB18(WT) comparison.

RAB18 crystal structure from RCSB PDB code 1X3S. (B) Table to show putative nucleotide-binding-dependent RAB18-associations with BirA*
RAB18(Ser22Asn):BirA*-RAB18(WT) association ratios <0.5. Proteins orthologous to interactors identified by Gillingham et al. (2014) are indicated. Previous studies providing supporting evidence for interactions are indicated. Proteins are grouped according to their attributes in the HeLa cell dataset (Figure 2 and Table S2).

Association ratios were derived individually following normalization by total spectral counts per condition. The full dataset is provided in table S1.

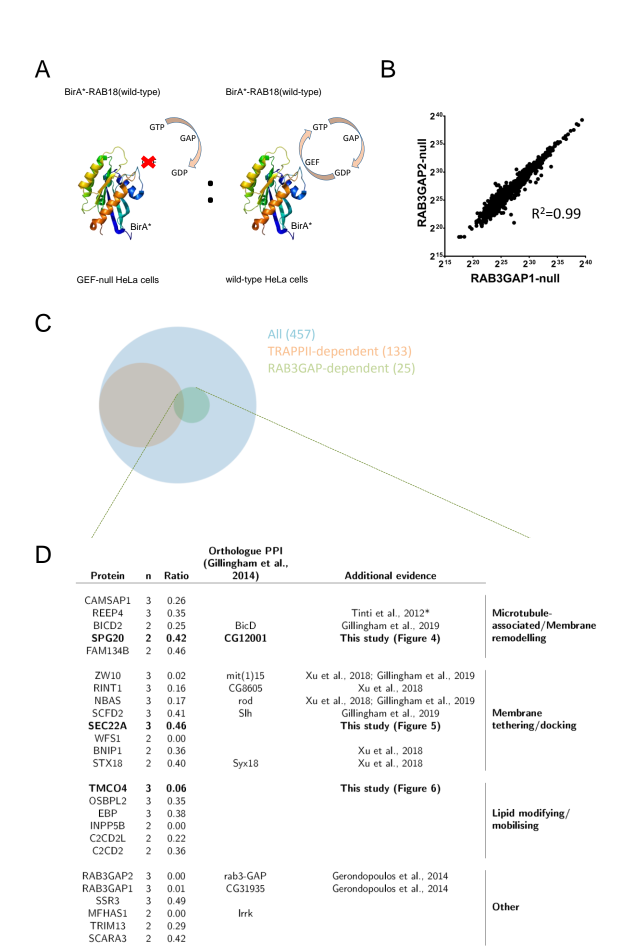


Figure 2. RAB3GAP-dependent RAB18-associations in HeLa cells. (A) Schematic to show comparison of BirA*-RAB18(WT) in wild-type and guanine nucleotide exchange factor (GEF)-null cells. RAB18 crystal structure from RCSB PDB code 1X3S. (B) Plot to show correlation between non-zero LFQ intensities of individual proteins identified in samples purified from RAB3GAP1- and RAB3GAP2-null cells. (C) Venn diagram illustrating all RAB18-associations, TRAPPII-dependent interactions (TRAPPC9-null:wild-type association ratios <0.5) and RAB3GAP-dependent associations (RAB3GAP1/2-null:wild-type association ratios <0.5). (D) Table to show putative RAB18-associations with RAB3GAP1/2-null:wild-type association ratios <0.5. Proteins orthologous to interactors identified by Gillingham et al. (2014) are indicated. Previous studies providing supporting evidence for interactions are indicated. Proteins are grouped according to their reported functions. Association ratios were derived individually following normalization by RAB18 LFQ intensity in each replicate experiment. The full dataset is provided in table S2.

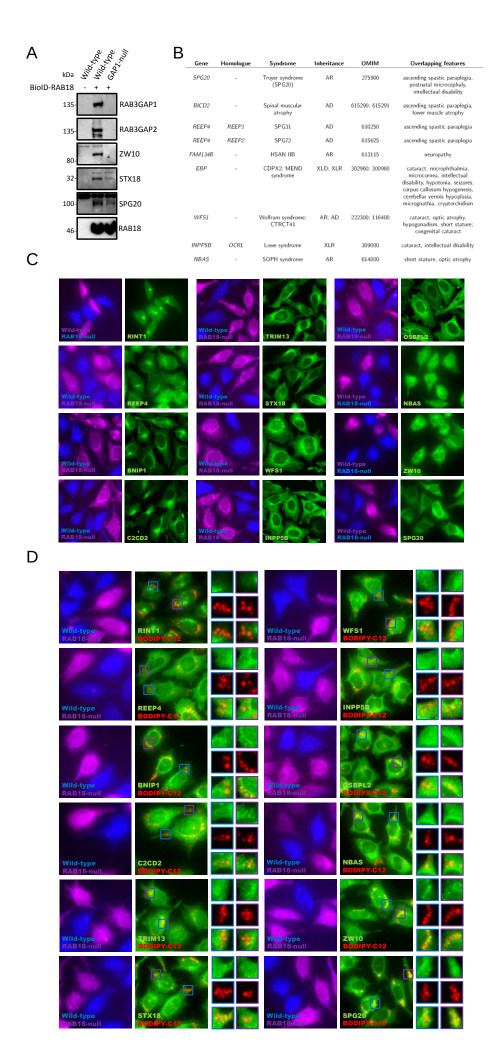
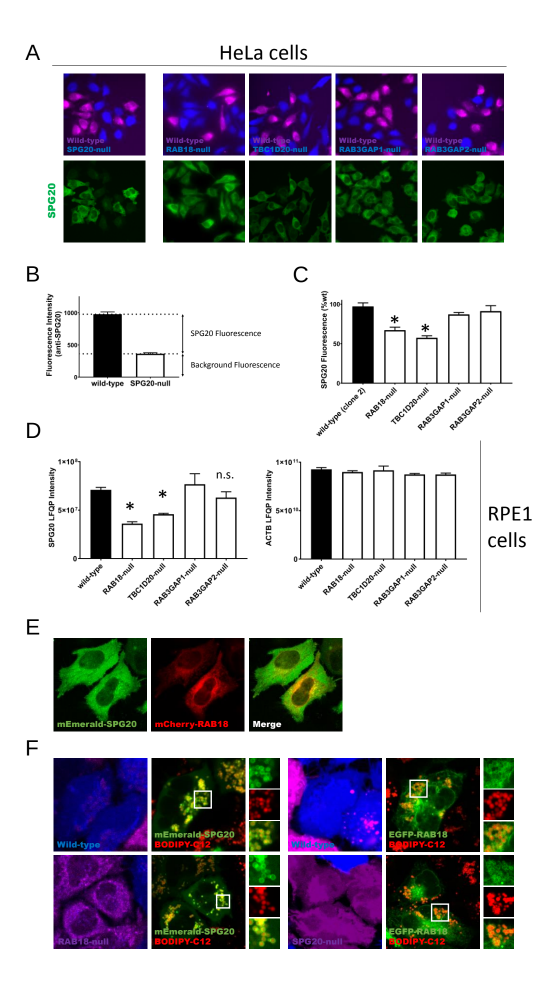


Figure 3. Initial validation of RAB3GAP-dependent RAB18-associations in HeLa cells. (A) Western blotting of samples purified from wild-type and RAB3GAP1-null cells in an independent BioID experiment. Levels of selected proteins are consistent with association ratios in Figure 1C. (B) Table to show diseases with features overlapping those of Warburg Micro syndrome linked to genes encoding putative effector proteins or their homologues. (C) Comparative fluorescence microscopy of selected RAB18-associated proteins in wild-type and RAB18-null HeLa cells. Cells of different genotypes were labelled with CellTrace-Violet and CellTrace-Far Red reagents, corresponding to blue and magenta channels respectively. Cells were stained with antibodies against indicated proteins in green channel panels. (D) Comparative fluorescence microscopy of selected RAB18-associated proteins in lipid-loaded wild-type and RAB18-null HeLa cells. Cells were stained as above but were treated for 15 hours with 200μM oleic acid, 1μg/ml BODIPY-558/568-C12 (Red channel) prior to fixation.



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Figure 4. Levels of SPG20 are significantly reduced in RAB18-null and TBC1D20-null HeLa and RPE1 cells. (A) Comparative fluorescence microscopy of SPG20 in wild-type and mutant HeLa cell lines. Wild-type and mutant cells of the indicated genotypes were labelled with CellTrace-Far Red and CellTrace-Violet and reagents respectively (magenta and blue channels). Cells were stained with an antibody against SPG20 (green channel) (B) Quantification of SPG20 fluorescence in wild-type cells by direct comparison with SPG20-null cells. (C) Quantification of SPG20 fluorescence (%wt) in cells of different genotypes. Data were derived from analysis of at least 18 frames – each containing >5 wild-type and >5 mutant cells – per genotype. *p<0.001. (D) LFQ intensities for SPG20 (Q8N0X7) and β-Actin (P60709) in whole-cell lysates of RPE1 cells of the indicated genotypes. n=3; *p<0.05 following FDR correction. Full dataset provided in table S3. Error bars represent s.e.m. (E) Confocal micrograph to show localization of exogenous mEmerald-SPG20 (Green) and mCherry-RAB18 (Red) in HeLa cells. (F) Confocal micrographs to show localization of exogenous mEmerald-SPG20 (Green; left panels) and EGFP-RAB18 (Green; right panels) in HeLa cells loaded with 200µM oleic acid, 1µg/ml BODIPY-558/568-C12 (Red channel). Wild-type and mutant cells of the indicated genotypes were labelled with CellTrace-Violet and CellTrace-Far Red reagents respectively (magenta and blue channels).

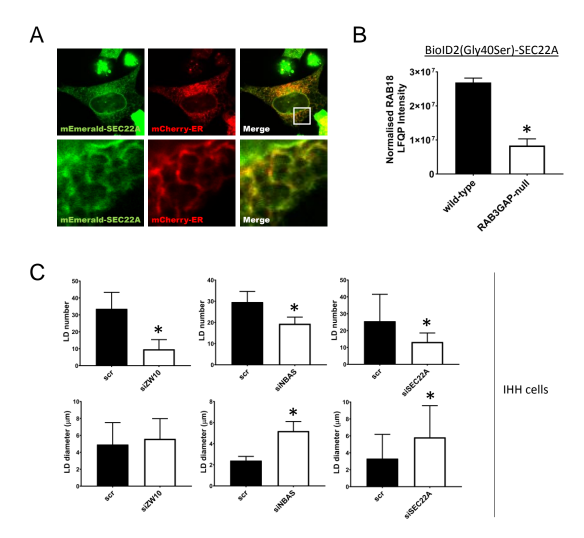
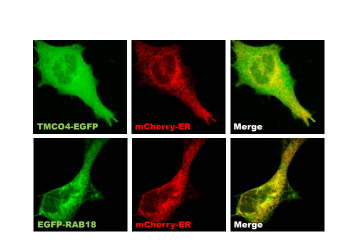
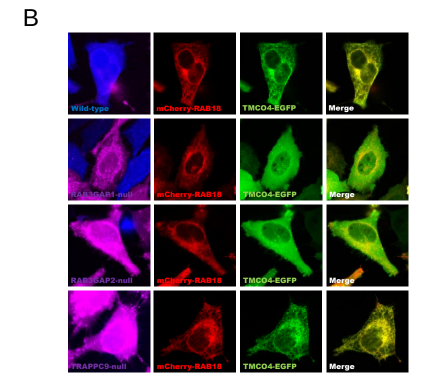


Figure 5. SEC22A associates with RAB18 and influences LD morphology. (A) Confocal micrograph to show overlapping localization of exogenous mEmerald-SEC22A (Green) and mCherry-ER (Red) in HeLa cells. (B) RAB18 LFQ intensities from a reciprocal BioID experiment showing a reduced association between BioID2(Gly40Ser)-SEC22A and endogenous RAB18 in RAB3GAP-null compared to wild-type HeLa cells. Data were adjusted to account for non-specific binding of RAB18 to beads and normalized by SEC22A LFQ intensities in each replicate experiment. Error bars represent s.e.m. Data for other BioID2(Gly40Ser)-SEC22A-associated proteins are provided in table S5. (C) Bar graphs to show effects of ZW10, NBAS and SEC22A knockdowns on lipid droplet number and diameter. siRNA-treated IHH cells were loaded with 200nM BSA-conjugated oleate, fixed and stained with BODIPY and DAPI, and imaged. Images were analysed using ImageJ. Data are derived from measurements from >100 cells/condition and are representative of three independent experiments. Error bars represent SD. *p<0.001



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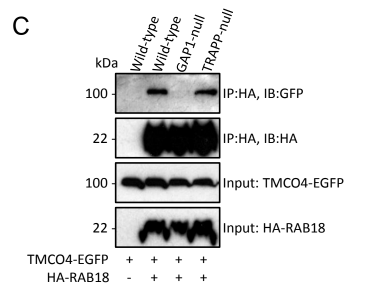


Figure 6. mCherry-RAB18 recruits TMCO4-EGFP to the ER membrane in a RAB3GAP-dependent manner. (A) Confocal micrographs to show diffuse localization of exogenous TMCO4-EGFP (Green) compared to mCherry-ER (Red) and overlapping localization of exogenous EGFP-RAB18 (Green) and mCherry-ER in HeLa cells. (B) Confocal micrographs to show localization of exogenous mCherry-RAB18 and TMCO4-EGFP in wild-type cells and in mutant cells of different genotypes. Wild-type and mutant cells of the indicated genotypes were labelled with CellTrace-Violet and CellTrace-Far Red reagents respectively (magenta and blue channels). (C) Immunoprecipitation of exogenous HA-RAB18 from HeLa cells of different genotypes. Cells were transfected with the indicated constructs and lysed 24 hours post-transfection. Anti-HA immunoprecipitates and input samples were subjected to SDS-PAGE and immunostaining for HA and GFP.

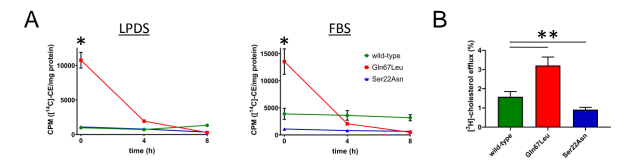


Figure 7. RAB18 is involved in the mobilization of cholesterol. (A) Plots to show cholesteryl ester (CE) loading and efflux. CHO cells, stably expressing RAB18(WT), RAB18(Gln67Leu) and RAB18(Ser22Asn), were incubated with [¹⁴C]-oleate, for 24 hours, in the presence of lipoprotein depleted serum (LPDS)(Left panel) or FBS (Right panel). Following lipid extraction, thin layer chromatography (TLC) was used to separate CE, and radioactivity was measured by scintillation counting. Measurements were made at t=0 and at 4 and 8 hours following the addition of 50μg/ml high density lipoprotein (HDL) to the cells. (B) Bar graph to show cholesterol efflux. The CHO cells were incubated with [³H]-cholesterol, for 24 hours, in the presence of FBS. After washing, they were then incubated with 25μg/ml apolipoprotein A-I for 5 hours. The quantity of [³H]-cholesterol in the media is shown as a percentage of the total cellular radioactivity (mean+/-SD). *p<0.01, **p<0.001.