

1 **Bifunctional probes reveal the rules of intracellular ether lipid**
2 **transport**

3

4 Kristin Böhlig^[a], Juan M. Iglesias-Artola^[a], H. Mathilda Lennartz^[a], Anna C. Link^[a], Björn
5 Drobot^[b], and André Nadler^{[a]*}

6 ^[a]Max Planck Institute of Molecular Cell Biology and Genetics, Pfotenauerstraße 108,
7 01307 Dresden, Germany

8 ^[b]Helmholtz-Zentrum Dresden-Rossendorf, Institute of Resource Ecology, Bautzner
9 Landstraße 400, 01328 Dresden, Germany

10

11 *Correspondence: nadler@mpi-cbg.de

12 **Abstract**

13 Ether glycerophospholipids bear a long chain alcohol attached via an alkyl or
14 vinyl ether bond at the *sn1* position of the glycerol backbone. Emerging evidence
15 suggests that ether lipids play a significant role in physiology and human health but
16 their precise cellular functions remain largely unknown. Here, we introduce bifunctional
17 ether lipid probes bearing diazirine and alkyne groups to study ether lipid biology. To
18 interrogate the kinetics of intracellular ether lipid transport in mammalian cells we used
19 a combination of fluorescence imaging, machine learning-assisted image analysis and
20 mathematical modelling. We find that alkyl-linked ether lipids are transported up to
21 twofold faster than vinyl-linked plasmalogens, suggesting that the lipid transport
22 machinery can distinguish between linkage types differing by as little as two hydrogen
23 atoms. We find that ether lipid transport predominantly occurs via non-vesicular
24 pathways, with varying contributions from vesicular mechanisms between cell
25 types. Altogether, our results suggest that differential recognition of alkyl- and vinyl
26 ether lipids by lipid transfer proteins contributes to their distinct biological functions.
27 In the future, the probes reported here will enable studying ether lipid biology in much
28 greater detail through identification of interacting proteins and in-depth
29 characterization of intracellular ether lipid dynamics.

30

31 **Main**

32 Eukaryotic ether lipids are primarily phosphatidylcholines (PCs) and
33 phosphatidylethanolamines (PEs) featuring a long-chain alcohol attached via alkyl
34 (plamanyl) or vinyl (plasmenyl) ether bonds at the *sn1* position of the glycerol
35 scaffold.^[1] Between 10-20% of all glycerophospholipids in humans are ether lipids,
36 with increased levels in brain, heart, kidney and skeletal muscle tissues.^[1] Ether lipids
37 are thought to have specific physiological roles despite the rather small structural
38 deviation from the corresponding diacyl ester lipids. For example, disruption of
39 peroxisomal ether lipid biosynthesis causes severe hereditary diseases, which are
40 characterized by skeletal, renal and cerebral abnormalities.^[2] Corresponding mice

41 models show abnormal eye development, male infertility and irregular behavior.^[3–6]
42 Many studies also indicate a correlation between disturbed ether lipid metabolism and
43 neurodegenerative diseases, psychiatric disorders and cancer but the underlying
44 molecular mechanisms are unclear.^[1,7–13]
45 Our knowledge of the cell biology of ether lipids is derived primarily from indirect
46 evidence. Ether lipids have been proposed to play a role in membrane trafficking^[14],
47 the general architecture of the endomembrane system, lipid sorting in the cell,
48 neurotransmitter release^[15] and ferroptosis.^[16–19] Intriguingly, elevated levels of ether
49 lipids were recently shown to be an adaptation of deep sea organisms to the high
50 pressure environment.^[20] Perhaps the clearest evidence points to a role of ether lipids
51 in modulating the export of GPI-anchored proteins via a direct interaction with the
52 transmembrane domain of mammalian p24 proteins, being metabolically co-regulated
53 with sphingolipids.^[21] Recent progress in ether lipid biology was mostly made by
54 combining genetic or other perturbations such as oxidative stress with lipidomic
55 approaches that now allow to distinguish between plasmenyl and plasmanyl lipids.^[22]
56 So far, very few ether lipid functions have been studied in mechanistic detail in
57 biological assays. Thus, the possibility remains that the wide range of observed
58 phenotypes are partially second-order effects of genetic perturbations. The difficulties
59 associated with mechanistic investigations into cellular functions of ether lipids can be
60 explained by an almost complete lack of suitable chemical tools for studying ether
61 lipids *in situ*. Apart from ether lipid analogues labelled with bulky fluorescent
62 moieties^[23,24] which are of limited use in cellular assays, the most promising approach
63 has so far been the use of polyene lyso-ether lipids, which are intrinsically fluorescent
64 due to conjugated trans double bonds.^[25,26] Despite promising early results, their use
65 has been limited, likely due to the relative complexity of the synthetic routes and issues
66 with rapid photobleaching during microscopy. Further development of strategies for
67 directly monitoring intracellular ether lipid localization, trafficking and interaction
68 partners are needed.

69 Among the most versatile tools for studying lipid biology are bifunctional probes
70 equipped with photoactivatable (diazirine) and clickable (alkyne) moieties. After UV-
71 induced crosslinking to proximal proteins, reporter groups can be attached to the
72 alkyne group by copper-mediated click chemistry. Bifunctional lipids have been used
73 to study lipid-protein interactions of fatty acyls, regular (diacyl) glycerophospholipids,
74 sphingolipids and sterols by utilizing affinity tags (e.g. biotin) and subsequent pulldown
75 of lipid-protein conjugates generated by crosslinking.^[27-34] In a limited number of
76 examples, these probes have also been used to report on lipid localization and
77 transport by attaching fluorophores after crosslinking and cell fixation and subsequent
78 analysis by fluorescence microscopy.^[29,34,35] Recently, we expanded upon this
79 methodology by employing bifunctional phospholipids to quantify both lipid transport
80 and metabolism using a combination of ultra-high resolution mass spectrometry and
81 fluorescence imaging.^[36] We reasoned that bifunctional ether lipid probes would be
82 ideal tools to study ether lipid biology.

83 As the primary probe of this study, we synthesized a PC ether lipid (1) featuring a
84 bifunctional, alkyl-ether-linked *sn1* chain in a 15-step synthesis featuring a
85 stereoselective $\text{Sc}(\text{OTf})_3$ -mediated epoxide opening. A complementary set of
86 bifunctional phosphatidylcholines (2-5) that only differ in the *sn1* linkage type via ester,
87 alkyl ether or a vinyl ether bond was generated to enable comparative studies of lipid
88 transport in cells. We performed pulse-chase experiments in two different cell lines
89 (HCT-116 and U2OS) and found that the plasmalogen probe was consistently
90 transported slower compared to both alkyl-ether and ester lipid probes. Ether lipid
91 transport was found to be primarily non-vesicular in both investigated cell lines, but the
92 fraction of material transported through the vesicular route was higher in U2OS vs
93 HCT-116 cells. Our data suggest that lipid transfer proteins distinguish plasmanyl and
94 plasmenyl ether lipids, suggesting a plausible mechanism for the enrichment of distinct
95 ether lipids in cellular membranes.

96

97

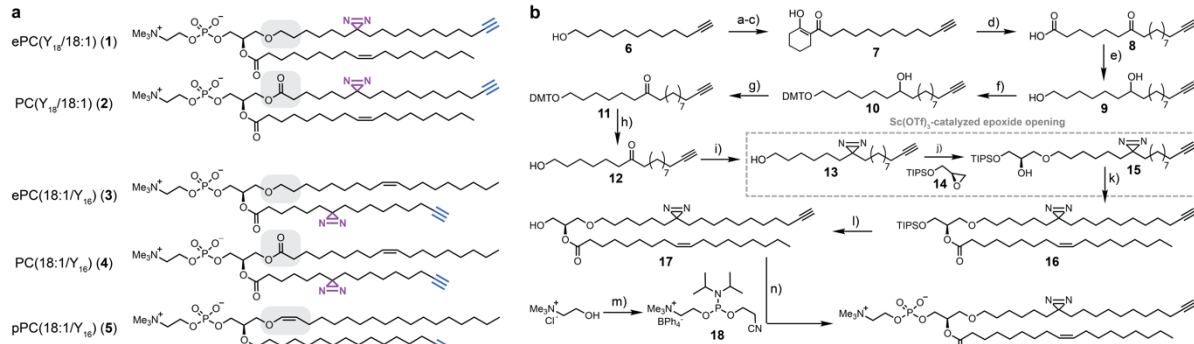
98 **Results**

99 **Synthesis of bifunctional ether lipid probes**

100 To study the influence of the linkage type on lipid transport, we synthesized
101 bifunctional lipids with identical head groups and closely related side chain
102 compositions. We focused on PC species as they are among the most abundant ether
103 lipids in mammalian cells.^[37] As plasmalogens feature one unsaturation in the *sn*1
104 chain we decided to use the monounsaturated oleic acid for the corresponding ester
105 and alkyl ether version. Probes **1** and **2** were designed to assess the influence of alkyl
106 ether vs ester linkage at the *sn*1 position (Fig. 1a), as they are otherwise identical
107 molecules. A complementary set of probes (**3-5**) was designed to perform a three-way
108 comparison of alkyl-ether, vinyl-ether and ester linkages (Fig. 1a). Together, the
109 complete set of probes was designed to study the effects of the following structural
110 elements that can plausibly affect transport mechanisms of ether lipids in cells: (i) side
111 chain linkage types (alkyl ether, vinyl ether, ester) and (ii) positioning of a single double
112 bond (*sn*1/*sn*2 chain, Δ9/Δ1).

113 The synthesis of the bifunctional lipids required two different strategies. For molecules
114 2-5, we first synthesized bifunctional fatty acids according to our recently published
115 protocol (see details in SI)^[36]. We then coupled the bifunctional fatty acids to the
116 respective lyso-lipids via EDC/DMAP-mediated esterification (see SI for details).
117 Probe 1, which bears the diazirine and alkyne functionalities in the ether chain at the
118 *sn*1 position, required a 15-step synthesis (Fig. 1b, see SI for synthetic details and
119 analytical data). We chose a synthetic route via the 1-Alkyl-2-acylglycerol ether 17, as
120 this intermediate offers straightforward access to a range of other ether lipid classes
121 via phosphoramidite chemistry.

122



123
124 **Figure 1 Overview of bifunctional ether lipid probes and synthesis of ePC(Y_{18:18:1}) bearing a bifunctional**
125 **ether linked side chain. a,** Structures of synthesized bifunctional ether and ester PCs 1-5 differing in the linkage
126 type at the *sn*1 position. **b,** Synthesis of the bifunctional ether lipid 1. The bifunctional alcohol 13 was synthesized
127 in 8 steps starting from 11-Dodecynol (6) and coupled to the TIPS-protected (S)-glycidol 14. After oleic acid
128 coupling and deprotection, phosphorylation according to Xu et. al.^[38] gave the final molecule 1. Reaction
129 conditions: a) CrO_3 , H_2SO_4 , H_2O , 0 °C-RT, 1h, 85%; b) $(\text{COCl})_2$, DMF, DCM, 1 h, 0 °C-RT; c) 1-
130 morpholinocyclohexene, NEt_3 , DCM, O.N., RT, 76% over 2 steps; d) aq. KOH, 15 min, 100 °C, 94%; e) LiAlH_4 ,
131 THF, O.N., 0 °C -RT, 83%; f) DMTCI, pyridine, 3.5 h, RT, 95%; g) DMP, pyridine, DCM, O.N., RT, 82%; h) FeCl_3 ,
132 MeOH, CHCl_3 , O.N., RT, 90%; i) 1. ammonia (solution in MeOH), MeOH, 0 °C-RT, 5.5 h. 2. $\text{H}_2\text{NOSO}_3\text{H}$, MeOH,
133 O.N., 0 °C-RT. 3. NEt_3 , iodine, MeOH, 8 h, 16%; j) 14, 5 mol% $\text{Sc}(\text{OTf})_3$, DCM, 2 d, RT, 59%; k) oleic acid, DMAP,
134 EDC-HCl, DCM, O.N., 0 °C-RT, 57%; l) TBAF, acetic acid, THF, O.N., 0 °C-RT, 32%; m) 1. NaBPh_4 , water. 2. 2-
135 cyanoethyl-N,N',N'-tetraisopropylphosphoro-amidite, 1H-tetrazole, MeCN, O.N., RT; n) 1. 18, 1H-tetrazole,
136 MeCN, 3.5 h, RT. 2. tBuOOH , MeCN, 1.5 h, 0 °C-RT. 3. NEt_3 , DCM, 20 h, RT, 47% over 3 steps.

137
138 The key step of the synthesis was the regioselective epoxide opening reaction used
139 to attach the bifunctional long-chain alcohol at the *sn*1 position of the glycerol scaffold.
140 Similar transformations have been reported previously^[39-41], but the utilized reagents
141 gave unsatisfactory yields in our hands, presumably due to the longer chain length
142 and the presence of the diazirine and alkyne groups.
143 The installation of the two functional groups at the ether-linked *sn*1 chain required the
144 synthesis of a long-chain alcohol 13 with an internal ketone and a terminal alkyne.
145 Test reactions indicated that the yields of a Grignard reaction with ϵ -caprolactone^[42]
146 as well as a Rh-catalyzed C-C-coupling reaction^[43] were not sufficient (see SI for
147 details). Thus, we first synthesized a long-chain fatty acid 8 bearing the internal ketone
148 and a terminal alkyne. To generate the alcohol 13, we initially attempted to selectively
149 reduce the carboxylic acid in the presence of the ketone. The use of mild reducing
150 reagents, such as NBu_4BH_4 , after activation of the carboxylic group via an acyl chloride
151 only gave moderate yields (see SI for details). The best overall yield was achieved
152 with a complete reduction of carboxylate and ketone to the respective hydroxy groups,
153 followed by a selective protection of the primary hydroxy group with DMTCI. After

154 oxidation of the secondary hydroxy group with Dess-Martin-Periodinane and
155 deprotection via FeCl_3 , the long-chain alcohol 12 was isolated in a yield of 58% over
156 four steps. We note, that the alternative route via ketone acetalization, reduction of the
157 carboxylate and deprotection was also feasible but gave a lower overall yield. After
158 introducing the diazirine group, the resulting bifunctional alcohol 13 was coupled to a
159 glycerol backbone via an epoxide opening reaction to afford intermediate 15. Many
160 different reagents, both bases and Lewis-acid catalysts, have been described in the
161 literature to afford high yields in similar transformations.^[40,41,44–52] In our hands, the
162 epoxide opening worked best with catalytic amounts of $\text{Sc}(\text{OTf})_3$ (5 mol%) in dry
163 DCM,^[52] whereas most published protocols gave either very low yields or no product
164 at all (see SI for details). The bifunctional 1-alkyl-glycerol ether 15 was isolated in 59%
165 yield under optimized reaction conditions (3.5 equivalents of epoxide 14 added, longer
166 reaction time and addition of a second batch of catalyst after 24 h). After subsequent
167 esterification with oleic acid using EDC/DMAP, the silyl protection group at the *sn*3
168 hydroxy group was removed using TBAF. We note that the addition of acetic acid was
169 necessary to minimize acyl chain migration from the *sn*2 to the *sn*3-position. The
170 synthesis was concluded by attaching the phosphocholine headgroup to intermediate
171 17 using phosphoramidite 18.^[34] After tetrazole-mediated coupling, the phosphor atom
172 was oxidized with $^t\text{BuOOH}$ and the cyanoethyl protecting group was removed with
173 NEt_3 to obtain the bifunctional alkyl ether PC 1 in an overall yield of 2.9%.

174

175 **Ether lipid transport depends on the linkage type and position of the** 176 **bifunctional chain**

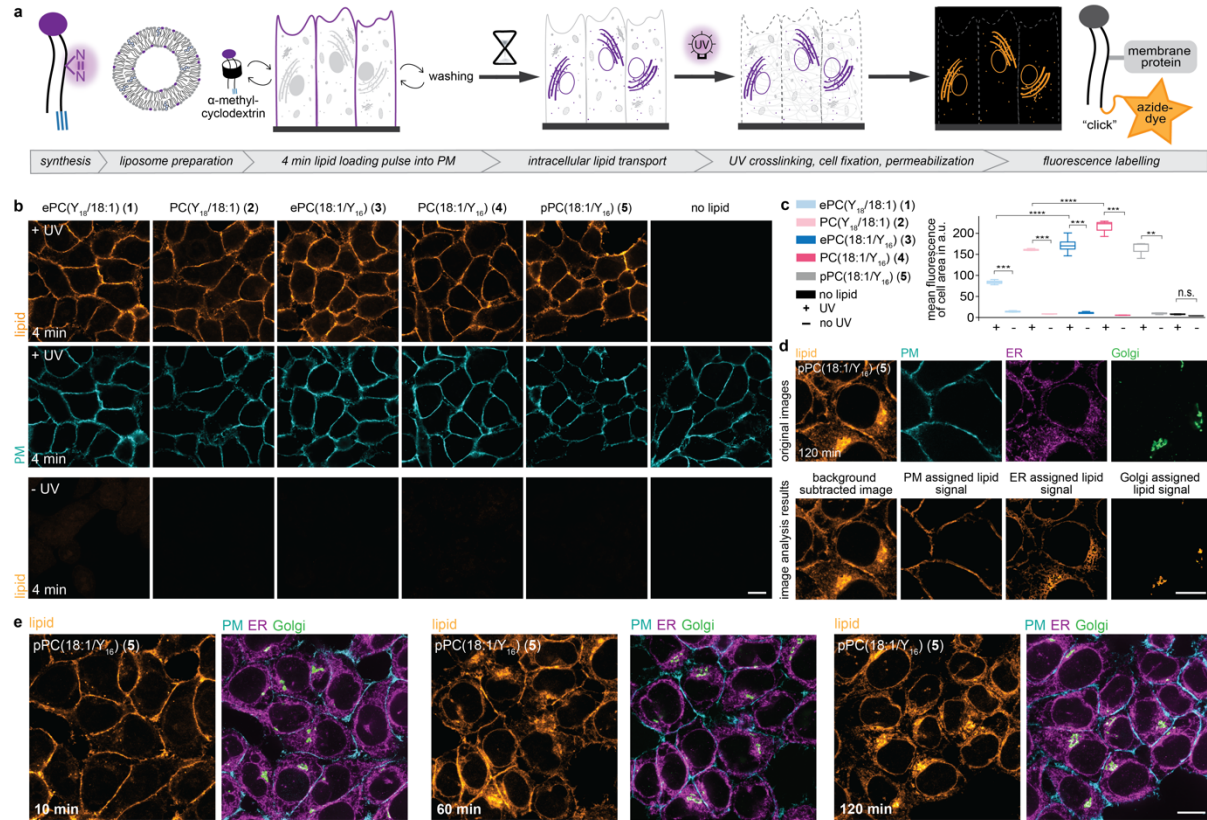
177 With the set of bifunctional ether lipids in-hand, we first asked how the linkage type at
178 the *sn*1 position influences the intracellular transport behaviour of PCs. Bifunctional
179 ether lipid probes were incorporated into the outer leaflet of the plasma membrane of
180 living cells by incubation with probe-containing liposomes and α -methylcyclodextrin.^[36]
181 After the 4 min loading pulse, the loading mix was replaced with cell growth medium
182 and the cells were placed at 37 °C to enable retrograde lipid transport. UV-crosslinking

183 (10 s at 310 nm) after the respective chase times was followed by cell fixation and
184 detergent treatment, achieving both cell permeabilization and the removal of non-
185 crosslinked lipid probes. The generated covalent lipid-protein conjugates were
186 subsequently fluorescently labelled via Cu-mediated click chemistry using an AF594
187 picolyl-azide dye and lipid localization was assessed by fluorescence microscopy (Fig.
188 2a).

189 We first confirmed lipid incorporation into the plasma membrane by analysing the
190 colocalization of lipid signal with a plasma membrane marker in HCT-116 cells after
191 the 4 min loading pulse (Fig. 2b). Control experiments without lipid loading and/or
192 without UV irradiation demonstrate that the observed fluorescence signal is specific to
193 the covalent lipid-protein conjugates, as very low background signal was observed.
194 (Fig. 2b lower panels, Fig. 2c, see supporting information for other timepoints). We
195 observed that both bifunctional lipids modified in the *sn1* position (probes **1+2**)
196 exhibited a significant lower overall fluorescence intensity in contrast to their *sn2*-
197 functionalized counterparts (probes **3-5**) (Fig. 2c). This could be due to different
198 crosslinking probabilities caused by the chain length difference or a higher likelihood
199 of proteins to interact with the *sn2* sidechain.

200 In order to analyze lipid transport processes, we performed four-color fluorescence
201 microscopy experiments with iterative immunofluorescent co-staining for the plasma
202 membrane, the endoplasmic reticulum, the Golgi and endosomes. We quantified the
203 lipid signal in each organelle using our recently reported image analysis pipeline based
204 on machine learning-assisted image segmentation.^[36] Briefly, self-trained Ilastik
205 Models were used to obtain probability masks to assign the fluorescence signal in the
206 lipid channel to individual organelles and quantify lipid distribution in the cell (Fig. 2d,
207 see SI for details of image background correction and image analysis).

208



209
210 **Figure 2 Fluorescence signal derived from bifunctional probes is specific and allows to analyze ether lipid**
211 **localization after intracellular transport. a**, General workflow for pulse-chase experiments to monitor ether lipid
212 transport with fluorescence microscopy. **b**, Cellular localization and signal intensity of bifunctional lipids 1-5 in HCT-
213 116 cells after a 4 min pulse with and without UV irradiation. Lipid-protein conjugates were labeled with an Azide-
214 AF594 dye via copper-catalyzed click chemistry. The plasma membrane was labeled with NHS-PEG4-Biotin. The
215 +UV and -UV images for the same probe are shown using identical settings, +UV image pairs are brightness-
216 contrast adjusted for better comparability as the signal intensity between lipid probes varied. Scale bar is 10 μ m.
217 **c**, Quantification of lipid signal intensity at 4 min for all probes and control conditions. The mean fluorescence
218 intensity of the cell-covered area is shown; images were processed using a Gaussian blur (sigma=4) filter and
219 background corrected. The error bars indicate the three-quartile values of the distribution. A permutation test from
220 the Python library Mlxtend (method=approximate, seed=0, number of rounds=1000) was used to test the
221 significance. **d**, Colocalization of lipid signal with plasma membrane, ER and Golgi markers allows to assign the
222 lipid signal to the respective organelles. The plasma membrane was labelled with MemBrite®Fix 405/430. The ER
223 and Golgi were labeled using immunofluorescence. Upper panels: Raw data of all 4 channels. Lower panels:
224 Background corrected lipid signal and lipid signal assigned to the plasma membrane, the ER and the Golgi,
225 respectively. Scale bar is 10 μ m. **e**, Cellular localization of pPC(18:1/C₁₆) at 10 min, 60 min and 120 min after lipid
226 loading with co-staining of the plasma membrane, ER and Golgi. The plasma membrane was labelled with
227 MemBrite®Fix 405/430. The ER and Golgi were labeled using immunofluorescence. Scale bar is 10 μ m.

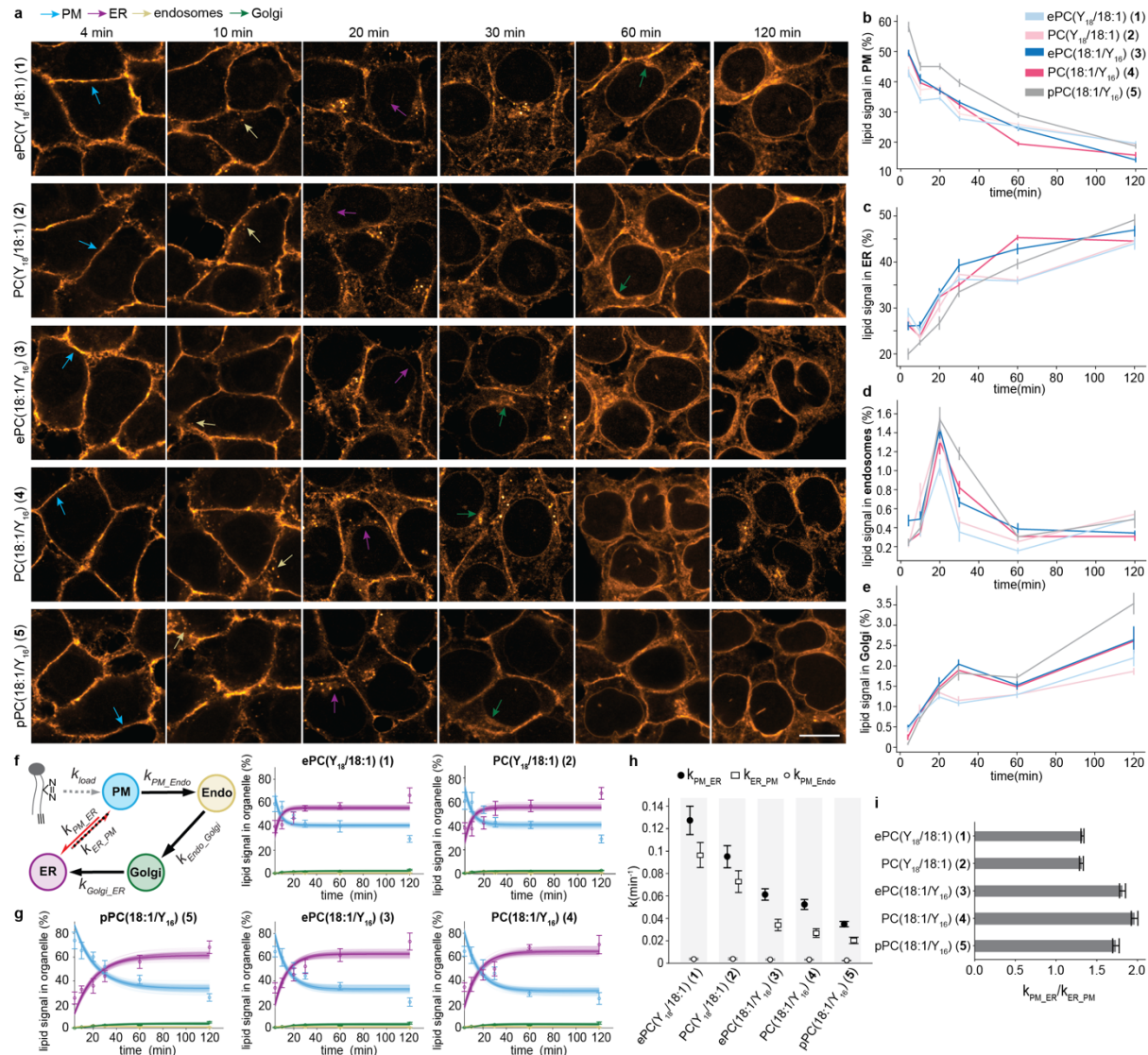
228
229 To quantify changes of lipid distribution over time, we performed pulse-chase
230 experiments with all lipid probes (1-5) covering a time period of 120 min in HCT-116
231 cells (Fig. 2e). Lipid signal intensity in the plasma membrane decreased over time for
232 all lipids, indicating probe internalization via retrograde trafficking (Fig. 3a,b, SI Fig.S2-
233 S8 for -UV controls). A comparison between lipid probes revealed that the

234 plasmalogen probe pPC(18:1/Y₁₆) (5) was retained at the plasma membrane to a
235 larger extent compared to both alkyl ether and ester lipid probes (Fig. 3a,b). Upon
236 internalization most of the signal for all probes was detected within the ER (Fig. 3c).
237 Apart from a brief spike at 20 min, very little fluorescence signal was assigned to
238 endosomal structures (Fig. 3d). Interestingly, all probes bearing the bifunctional fatty
239 acid at the *sn*2 position (probes 3-5) exhibited more Golgi localization after 30 min
240 (Fig. 3a,e, SI Fig.S9). This suggested that intracellular lipid transport routes are
241 specific to the presence of the double bond within the *sn*1 chain. At 120 min the
242 plasmalogen was more enriched in the Golgi (Fig. 3e) implying a differential steady
243 state distribution between the probes.

244 To obtain a quantitative description of the lipid transport kinetics and to test the
245 contribution of vesicular and non-vesicular transport routes, we fitted a kinetic model
246 to the obtained time-course data (Fig. 3f,g see SI for model details). The model
247 distinguishes non-vesicular transport from the plasma membrane to the ER from
248 retrograde vesicular transport (plasma membrane via endosomes and the Golgi to the
249 ER). Anterograde lipid transport is described by a summary rate constant from the ER
250 to the plasma membrane capturing both vesicular and non-vesicular transport modes.
251 We found that transport via the non-vesicular route in the retrograde direction
252 (captured by the rate constant k_{PM_ER}) was much faster than vesicular membrane
253 trafficking (k_{PM_Endo}) for all lipids (Fig. 3h). The alkyl ether and ester lipid probes bearing
254 the bifunctional moieties at the *sn*1 chain (probes 1+2) were transported faster than
255 the corresponding *sn*2-modified probes (probes 3-5), which is likely due to both chain
256 length and positioning. However, the linkage type via alkyl ether or ester had only
257 negligible effects on transport kinetics. In contrast, the vinyl ether linkage of the
258 plasmalogen 5 led to overall much slower transport. The steady-state distribution of
259 the plasmalogen probe pPC(18:1/Y₁₆) (5) was shifted towards the plasma membrane
260 compared to the corresponding alkyl ether (ePC(18:1/Y₁₆) (3) and ester lipid
261 (PC(18:1/Y₁₆) (4) probes (quasi-equilibrium constants k_{PM_ER}/k_{ER_PM} , Fig. 3i). In
262 general, we observed that the effect of plasmalogen-vinyl ether linkage supersedes

263 the effect caused by the placement of the double bond in the *sn1* or *sn2* acyl chain
 264 whereas the latter appears to be stronger than the difference between alkyl ether and
 265 ester linkage types.

266



267
 268 **Figure 3 Plasmalogens exhibit slower retrograde transport kinetics than the corresponding alkyl-ether and**
 269 **ester lipid probes in HCT-116 cells. a**, Cellular localization of bifunctional lipids 1-5 in HCT-116 cells at 4, 10, 20,
 270 30, 60 and 120 min after lipid loading. The lipid was labeled with an Azide-AF594 dye via copper-catalyzed click
 271 chemistry. Scale bar is 10 μ m. **b-e**, Quantification of the relative lipid signal in the plasma membrane, the ER,
 272 endosomes and the Golgi apparatus. Error bars: SE. **f**, Kinetic model for quantifying lipid transport from
 273 fluorescence microscopy data. Black arrows indicate vesicular transport, red arrows non-vesicular transport. **g**,
 274 The corresponding model fits of the lipid transport kinetics for all lipids derived from 100 model runs. **h**, Comparison
 275 of rate constants describing retrograde vesicular transport from the plasma membrane to endosomes, retrograde
 276 non-vesicular transport from the PM to the ER and total transport in the anterograde direction from the ER to
 277 the PM. Error bars: SD. **i**, Ratio of the rate constants for non-vesicular lipid transport between the plasma membrane
 278 and the ER in the retrograde and anterograde direction. Error bars: SD.

279

280

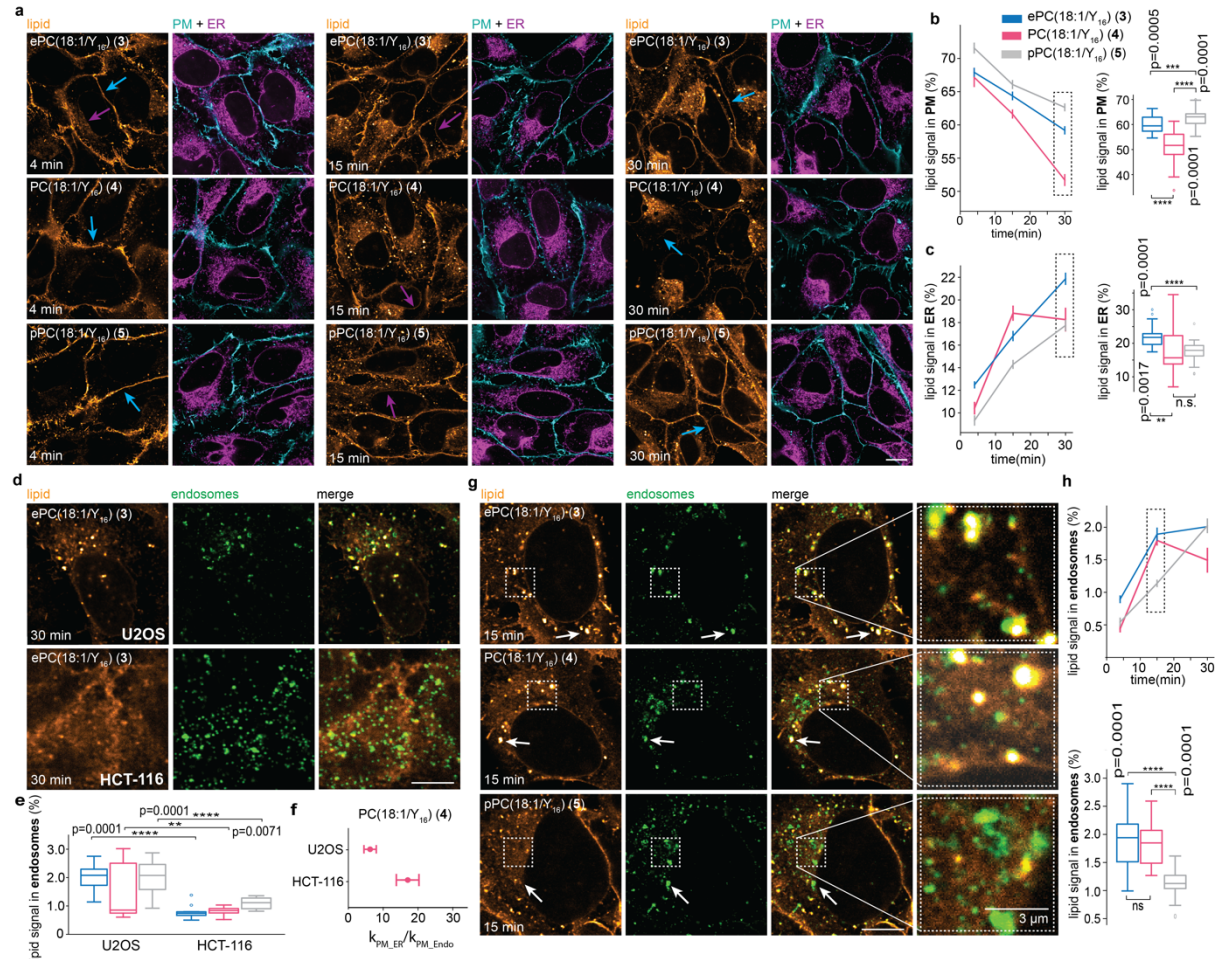
281 **Ether lipid transport is similar in HCT-116 and U2OS cells**

282 To assess whether the trends observed for ether lipid transport in HCT-116 cells are
283 more generally conserved, we conducted lipid transport experiments in U2OS cells.
284 The two cell lines differ significantly in origin (bone marrow and colon) and
285 morphology. U2OS cells are large, flat cells resembling mesenchymal cells, whereas
286 HCT-116 cells are smaller and retain most structural features of their epithelial origin.
287 We used the *sn*2-bifunctional lipid probes in pulse-chase experiments to assess
288 potential differences in lipid transport. Endpoint quantification at 30 min showed that
289 the plasmalogen was retained at the plasma membrane to a larger extent than both
290 the corresponding ester and alkyl ether probes (Fig. 4 a,b). This effect was also
291 reflected by lower plasmalogen lipid signal in the ER at all timepoints (Fig. 4 a,c).
292 Overall, the relative differences between individual lipids were more pronounced for
293 U2OS cells compared to HCT-116 cells.

294 One notable difference between the cell lines were bright vesicular structures in the
295 lipid image observed in U2OS but not in HCT-116 cells. These structures were
296 identified as endosomes with immunofluorescence (Fig. 4 d). The quantification of the
297 lipid signal fraction in endosomes confirmed that a larger amount of lipid material was
298 routed through the endocytic route in U2OS cells (Fig. 4 e). This observation suggests
299 that the non-vesicular transport route is even more dominant in HCT-116 cells
300 compared to U2OS cells which is also reflected in the difference between rate
301 constants of PC(18:1/Y₁₆) (4) describing endocytosis and non-vesicular transport from
302 the plasma membrane to the ER in the retrograde direction (Fig. 4 f).^[36] A closer
303 inspection of lipid incorporation in endosomes in U2OS cells revealed a slower uptake
304 of the plasmalogen probe, which is also reflected in a slower appearance of lipid signal
305 in the Golgi (Fig. 4 g,h and SI Fig. S10). This effect is not caused by a change in the
306 bulk endocytic rate induced by probe loading, as the number and size of endosomes
307 were found to be constant for all probes (SI Fig. S11). Taken together, while some cell
308 line-specific differences were apparent, the major trends for ether lipid transport in

309 both cell lines were similar, in particular with regard to slower internalization kinetics
 310 of the plasmalogen probe and predominant non-vesicular transport.

311



312
 313
 314
 315
 316
 317
 318
 319
 320
 321
 322
 323
 324
 325
 326

327
 328

Figure 4 Retrograde vesicular transport of ether lipids is cell- and linkage type specific. **a**, Cellular localization of *sn*2-bifunctional lipid probes at 4 min, 15 min and 30 min in U2OS cells with co-staining of the plasma membrane and the ER. Scale bar is 10 μ m. **b-c**, Quantification of the relative lipid intensity in the plasma membrane and the ER. A permutation test from the Python library *MLxtend* (method=approximate, seed=0, number of rounds=1000) was used to test the significance at the 30 min timepoint. **d**, Cellular localization of ePC(18:1/Y₁₆) in U2OS and HCT-116 cells at the bottom plane after 30 min with co-staining of the Rab5 and Rab7 proteins. Scale bar is 10 μ m. **e**, Quantification of the relative lipid signal in endosomes of the *sn*2-bifunctional lipid probes in U2OS and HCT-116 cells after 30 min. **f**, Comparison of ratios between the kinetic rates for non-vesicular (k_{PM_ER}) and vesicular (k_{PM_Endo}) transport of PC(18:1/Y₁₆) in HCT-116 (this work) and U2OS cells^[36]. Error bars: SD. **g**, Cellular localization of the *sn*2-bifunctional lipid probes in U2OS cells at 15 min with co-staining of the Rab5 and Rab7 proteins. Scale bar is 10 μ m. **h**, Quantification of the relative lipid signal in endosomes. A permutation test from the Python library *MLxtend* (method=approximate, seed=0, number of rounds=1000) was used to test the significance at the 15 min timepoint.

329 **Discussion**

330 We report a new set of bifunctional photoaffinity probes for studying ether lipid biology.
331 We find notable differences in transport speed for plasmalogen and alkyl ether lipids,
332 suggesting that the lipid transport machinery indeed exhibits specificity for distinct
333 ether lipid types. While very little is known about the cellular roles of individual ether
334 lipid species, our findings imply distinct functions of plasmalogen vs alkyl ether and
335 ester lipids. The effect sizes found here are smaller compared to more structurally
336 diverse lipids with different headgroups and saturation degrees, but nonetheless
337 significant.^[36] These data imply that small structural differences are recognized by the
338 cellular lipid handling machinery, providing further evidence for the biological
339 importance of lipid diversity.^[53]

340 The utilized modular synthesis employing the long chain bifunctional alcohol **13**,
341 stereoselective $\text{Sc}(\text{OTf})_3$ catalyzed epoxide opening, and phosphor amidite
342 headgroup coupling enables straightforward synthetic access to further bifunctional
343 ether lipid classes. This opens the way to study the biology of ether lipids both with
344 regard to detailed analyses of individual lipid species as well as to expand the scope
345 of investigations to the broader ether lipidome. Furthermore, while this study is focused
346 on analyzing ether lipid transport by fluorescence imaging, bifunctional lipids can also
347 be employed for tracing metabolism and identifying lipid-protein interactions.^[29,31,35,36]
348 Taken together, the bifunctional ether lipid probes reported here represent a versatile
349 toolkit for studying ether lipid biology in mechanistic detail. This capability will have
350 significant benefits for understanding the functions of ether lipids in fundamental cell
351 biology and their role in human diseases.

352

353 **Acknowledgements**

354 AN gratefully acknowledges financial support by the European Research Council
355 (ERC) under the European Union's Horizon 2020 research and innovation program
356 (grant agreements no GA 758334 ASYMMEM and AURORA). This research was
357 supported by an Allen Distinguished Investigator Award, a Paul G. Allen Frontiers

358 Group advised grant of the Paul G. Allen Family Foundation to AN. AN and KB
359 acknowledge financial support by the VolkswagenStiftung within the Life? initiative.
360 We thank the following services and facilities at MPI-CBG Dresden for their support:
361 Light Microscopy Facility, Computer Department, in particular Oscar Gonzalez for
362 expert advice with regard to high performance computing, and Mass Spectrometry
363 Facility.

364

365 **Conflict of Interest**

366 AN and JMIA have received a Proof-of-Concept grant from the European Research
367 Council to explore the commercial potential of the lipid imaging methodology.

368

369 **Keywords:** Ether lipids, plasmalogens, lipid transport, bifunctional lipids, lipid
370 imaging

371

372 **References**

373 [1] N. E. Braverman, A. B. Moser, *Biochimica et Biophysica Acta (BBA) - Molecular*
374 *Basis of Disease* **2012**, 1822, 1442–1452.

375 [2] S. Goldfischer, C. L. Moore, A. B. Johnson, A. J. Spiro, M. P. Valsamis, H. K.
376 Wisniewski, R. H. Ritch, W. T. Norton, I. Rapin, L. M. Gartner, *Science* **1973**, 182,
377 62–64.

378 [3] F. Dorninger, A. Gundacker, G. Zeitler, D. D. Pollak, J. Berger, *Int J Mol Sci* **2019**,
379 20, E3929.

380 [4] C. Rodemer, T.-P. Thai, B. Brugger, T. Kaercher, H. Werner, K.-A. Nave, F.
381 Wieland, K. Gorgas, W. W. Just, *Human Molecular Genetics* **2003**, 12, 1881–
382 1895.

383 [5] K. Gorgas, A. Teigler, D. Komljenovic, W. W. Just, *Biochimica et Biophysica Acta*
384 *(BBA) - Molecular Cell Research* **2006**, 1763, 1511–1526.

385 [6] F. Dorninger, R. Herbst, B. Kravic, B. Z. Camurdanoglu, I. Macinkovic, G. Zeitler,
386 S. Forss-Petter, S. Strack, M. M. Khan, H. R. Waterham, R. Rudolf, S.
387 Hashemolhosseini, J. Berger, *Journal of neurochemistry* **2017**, 143, 569.

388 [7] R. Kaddurah-Daouk, J. McEvoy, R. Baillie, H. Zhu, J. K. Yao, V. L. Nimgaonkar,
389 P. F. Buckley, M. S. Keshavan, A. Georgiades, H. A. Nasrallah, *Psychiatry*
390 *Research* **2012**, 198, 347–352.

391 [8] L. D. Spears, S. Adak, G. Dong, X. Wei, G. Spyropoulos, Q. Zhang, L. Yin, C.
392 Feng, D. Hu, I. J. Lodhi, F.-F. Hsu, R. Rajagopal, K. K. Noguchi, C. M. Halabi, L.
393 Brier, A. R. Bice, B. V. Lananna, E. S. Musiek, O. Avraham, V. Cavalli, J. K. Holth,
394 D. M. Holtzman, D. F. Wozniak, J. P. Culver, C. F. Semenkovich, *J Lipid Res*
395 **2021**, 62, 100079.

396 [9] J. Kou, G. G. Kovacs, R. Höftberger, W. Kulik, A. Brodde, S. Forss-Petter, S.
397 Höningschnabl, A. Gleiss, B. Brügger, R. Wanders, W. Just, H. Budka, S.
398 Jungwirth, P. Fischer, J. Berger, *Acta Neuropathol* **2011**, 122, 271–283.

399 [10] A. M. Dickens, P. Sen, M. J. Kempton, N. Barrantes-Vidal, C. Iyegbe, M.
400 Nordentoft, T. Pollak, A. Riecher-Rössler, S. Ruhrmann, G. Sachs, R. Bressan,
401 M.-O. Krebs, G. P. Amminger, L. De Haan, M. Van Der Gaag, L. Valmaggia, T.
402 Hyötyläinen, M. Orešić, P. McGuire, P. McGuire, L. R. Valmaggia, M. J. Kempton,
403 M. Calem, S. Tognin, G. Modinos, L. De Haan, M. Van Der Gaag, E. Velthorst, T.
404 C. Kraan, D. S. Van Dam, N. Burger, B. Nelson, P. McGorry, G. P. Amminger, C.
405 Pantelis, A. Politis, J. Goodall, A. Riecher-Rössler, S. Borgwardt, C. Rapp, S. Ittig,
406 E. Studerus, R. Smieskova, R. Bressan, A. Gadelha, E. Brietzke, G. Asevedo, E.
407 Asevedo, A. Zugman, N. Barrantes-Vidal, T. Domínguez-Martínez, A. Raciopi,
408 T. R. Kwapil, M. Monsonet, A. Rosa, A. Frajerman, B. Chaumette, J. Bourgin, O.
409 Kebir, C. Jantac, M.-O. Krebs, D. Nordholm, L. Randers, K. Krakauer, L. Glenthøj,
410 B. Glenthøj, M. Nordentoft, S. Ruhrmann, D. Gebhard, J. Arnhold, J.
411 Klosterkötter, G. Sachs, I. Lasser, B. Winklbaur, P. A. Delespaul, B. P. Rutten, J.
412 Van Os, *Biological Psychiatry* **2021**, 89, 288–297.

413 [11] D. I. Benjamin, A. Cozzo, X. Ji, L. S. Roberts, S. M. Louie, M. M. Mulvihill, K. Luo,
414 D. K. Nomura, *Proc. Natl. Acad. Sci. U.S.A.* **2013**, *110*, 14912–14917.

415 [12] F. Dorninger, S. Forss-Petter, J. Berger, *FEBS Lett* **2017**, *591*, 2761–2788.

416 [13] F. Dorninger, S. Forss-Petter, I. Wimmer, J. Berger, *Neurobiology of Disease*
417 **2020**, *145*, 105061.

418 [14] T.-P. Thai, *Human Molecular Genetics* **2001**, *10*, 127–136.

419 [15] F. Dorninger, T. König, P. Scholze, M. L. Berger, G. Zeitler, C. Wiesinger, A.
420 Gundacker, D. D. Pollak, S. Huck, W. W. Just, S. Forss-Petter, C. Pifl, J. Berger,
421 *Human Molecular Genetics* **2019**, *28*, 2046–2061.

422 [16] Y. Zou, W. S. Henry, E. L. Ricq, E. T. Graham, V. V. Phadnis, P. Maretich, S.
423 Paradkar, N. Bohnke, A. A. Deik, F. Reinhardt, J. K. Eaton, B. Ferguson, W.
424 Wang, J. Fairman, H. R. Keys, V. Dančík, C. B. Clish, P. A. Clemons, P. T.
425 Hammond, L. A. Boyer, R. A. Weinberg, S. L. Schreiber, *Nature* **2020**, *585*, 603–
426 608.

427 [17] M. A. Perez, A. J. Clostio, I. R. Houston, J. Ruiz, L. Magtanong, S. J. Dixon, J. L.
428 Watts, *PLOS Genetics* **2022**, *18*, e1010436.

429 [18] N. Jiménez-Rojo, H. Riezman, *FEBS Lett* **2019**, *593*, 2378–2389.

430 [19] J. M. Dean, I. J. Lodhi, *Protein Cell* **2018**, *9*, 196–206.

431 [20] J. R. Winnikoff, D. Milshteyn, S. J. Vargas-Urbano, M. A. Pedraza-Joya, A. M.
432 Armando, O. Quehenberger, A. Sodt, R. E. Gillilan, E. A. Dennis, E. Lyman, S.
433 H. D. Haddock, I. Budin, *Science* **2024**, *384*, 1482–1488.

434 [21] N. Jiménez-Rojo, M. D. Leonetti, V. Zoni, A. Colom, S. Feng, N. R. Iyengar, S.
435 Matile, A. Roux, S. Vanni, J. S. Weissman, H. Riezman, *Current Biology* **2020**,
436 30, 3775–3787.

437 [22] J. Koch, K. Lackner, Y. Wohlfarter, S. Sailer, J. Zschocke, E. R. Werner, K.
438 Watschinger, M. A. Keller, *Anal. Chem.* **2020**, *92*, 11268–11276.

439 [23] H. Zheng, R. I. Duclos, C. C. Smith, H. W. Farber, R. A. Zoeller, *Journal of Lipid*
440 *Research* **2006**, *47*, 633–642.

441 [24] E. R. Werner, M. A. Keller, S. Sailer, D. Seppi, G. Golderer, G. Werner-Felmayer,
442 R. A. Zoeller, K. Watschinger, *Journal of Lipid Research* **2018**, 59, 901–909.

443 [25] L. Kuerschner, D. Richter, H. K. Hannibal-Bach, A. Gaebler, A. Shevchenko, C.
444 S. Ejsing, C. Thiele, *PLoS ONE* **2012**, 7, e31342.

445 [26] L. Kuerschner, C. S. Ejsing, K. Ekoos, A. Shevchenko, K. I. Anderson, C. Thiele,
446 *Nat Methods* **2005**, 2, 39–45.

447 [27] T. Peng, X. Yuan, H. C. Hang, *Current Opinion in Chemical Biology* **2014**, 21,
448 144–153.

449 [28] P. Haberkant, J. C. M. Holthuis, *Biochimica et Biophysica Acta (BBA) - Molecular*
450 *and Cell Biology of Lipids* **2014**, 1841, 1022–1030.

451 [29] R. Müller, A. Kojic, M. Citir, C. Schultz, *Angew. Chem. Int. Ed.* **2021**, 60, 19759–
452 19765.

453 [30] P. Hartwig, D. Höglinder, *Int. J. Mol. Sci.* **2021**, 22, 7065.

454 [31] S. Bockelmann, J. G. M. Mina, S. Korneev, D. G. Hassan, D. Müller, A. Hilderink,
455 H. C. Vlieg, R. Raijmakers, A. J. R. Heck, P. Haberkant, J. C. M. Holthuis, *Journal*
456 *of Lipid Research* **2018**, 59, 515–530.

457 [32] M. J. Niphakis, K. M. Lum, A. B. Cognetta, B. E. Correia, T.-A. Ichu, J. Olucha, S.
458 J. Brown, S. Kundu, F. Piscitelli, H. Rosen, B. F. Cravatt, *Cell* **2015**, 161, 1668–
459 1680.

460 [33] P. Haberkant, F. Stein, D. Höglinder, M. J. Gerl, B. Brügger, P. P. Van Veldhoven,
461 J. Krijgsveld, A.-C. Gavin, C. Schultz, *ACS Chem. Biol.* **2016**, 11, 222–230.

462 [34] D. Höglinder, A. Nadler, P. Haberkant, J. Kirkpatrick, M. Schifferer, F. Stein, S.
463 Hauke, F. D. Porter, C. Schultz, *Proceedings of the National Academy of*
464 *Sciences* **2017**, 114, 1566–1571.

465 [35] P. Haberkant, R. Raijmakers, M. Wildwater, T. Sachsenheimer, B. Brügger, K.
466 Maeda, M. Houweling, A.-C. Gavin, C. Schultz, G. van Meer, A. J. R. Heck, J. C.
467 M. Holthuis, *Angew. Chem. Int. Ed.* **2013**, 52, 4033–4038.

468 [36] J. M. Iglesias-Artola, K. Schuhmann, K. Böhlig, H. M. Lennartz, M. Schuhmacher,
469 P. Barahtjan, C. Jiménez López, R. Šachl, K. Pombo-Garcia, A. Lohmann, P.

470 Riegerová, M. Hof, B. Drobot, A. Shevchenko, A. Honigmann, A. Nadler, **2024**,
471 bioRxiv preprint DOI: 10.1101/2024.05.14.594078.

472 [37] N. Nagan, R. A. Zoeller, *Progress in Lipid Research* **2001**, *40*, 199–229.

473 [38] L. L. Xu, L. J. Berg, D. Jamin Keith, S. D. Townsend, *Org. Biomol. Chem.* **2020**,
474 *18*, 767–770.

475 [39] T. L. Andresen, J. Davidsen, M. Begtrup, O. G. Mouritsen, K. Jørgensen, *J. Med.*
476 *Chem.* **2004**, *47*, 1694–1703.

477 [40] P. N. Guivisalsky, R. Bittman, *J. Org. Chem.* **1989**, *54*, 4637–4642.

478 [41] S. D. Stamatov, J. Stawinski, *Org. Biomol. Chem.* **2007**, *5*, 3787.

479 [42] J. Davies, S. G. Booth, S. Essafi, R. A. W. Dryfe, D. Leonori, *Angew. Chem. Int.*
480 *Ed.* **2015**, *54*, 14017–14021.

481 [43] C.-H. Jun, H. Lee, J.-B. Hong, *J. Org. Chem.* **1997**, *62*, 1200–1201.

482 [44] S. Hoppen, S. Bärle, U. Koert, *Chemistry – A European Journal* **2000**, *6*, 2382–
483 2396.

484 [45] J. Barluenga, H. Vázquez-Villa, A. Ballesteros, J. M. González, *Org. Lett.* **2002**,
485 *4*, 2817–2819.

486 [46] T. L. Andresen, J. Davidsen, M. Begtrup, O. G. Mouritsen, K. Jørgensen, *J. Med.*
487 *Chem.* **2004**, *47*, 1694–1703.

488 [47] P. N. Guivisalsky, R. Bittman, *J. Am. Chem. Soc.* **1989**, *111*, 3077–3079.

489 [48] M. Moghadam, S. Tangestaninejad, V. Mirkhani, I. Mohammadpoor-Baltork, S. A.
490 Taghavi, *Catalysis Communications* **2007**, *8*, 2087–2095.

491 [49] N. Iranpoor, M. Shekarriz, F. Shiriny, *Synthetic Communications* **1998**, *28*, 347–
492 366.

493 [50] M. Fallah-Mehrjardi, A. R. Kiasat, K. Niknam, *J. Iran. Chem. Soc.* **2018**, *15*, 2033–
494 2081.

495 [51] T. Hansen, P. Vermeeren, A. Haim, M. J. H. van Dorp, J. D. C. Codée, F. M.
496 Bickelhaupt, T. A. Hamlin, *Eur. J. Org. Chem.* **2020**, *25*, 3822–3828.

497 [52] C. Schneider, A. R. Sreekanth, E. Mai, *Angew. Chem. Int. Ed.* **2004**, *43*, 5691–
498 5694.

499 [53] A. Shevchenko, K. Simons, *Nat Rev Mol Cell Biol* **2010**, *11*, 593–598.

500