Bao, Leyuan, Fearnley, Gareth W., Lin, Chi-Chuan, Odell, Adam ORCID logoORCID: https://orcid.org/0000-0002-6855-7214, Redondo, Ana C., Kinsella, Gemma K., Findlay, John B. C., Ladbury, John E. ORCID logoORCID: https://orcid.org/0000-0002-6328-7200, Harrison, Michael A. and Ponnambalam, Sreenivasan ORCID logoORCID: https://orcid.org/0000-0002-4452-7619 (2021) Calcium-binding protein S100A6 interaction with VEGF receptor integrates signaling and trafficking pathways. BioRxiv. (Unpublished)

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2	Calcium-binding protein S100A6 interaction with VEGF receptors	
3	integrates signaling and trafficking pathways	
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;	Leyuan Bao ^{1#} , Gareth W. Fearnley ^{1#} , Chi-Chuan Lin ¹ , Adam F. Odell ¹ , Ana C. Redondo ² ,	
)	Gemma K. Kinsella ³ , John B. C. Findlay ² , John E. Ladbury ¹ , Michael A. Harrison ² ,	
,	Sreenivasan Ponnambalam ^{1*}	
	School of Molecular & Cellular Biology, ² School of Biomedical Sciences, University of Leeds, Leeds LS2 9JT, U	K;
	School of Food Science and Environmental Health, College of Sciences and Health, Technological University Dubli	in,
	Dublin D07 ADY7, Eire.	
	Co-first authors; L.B., G.W.F.	
	Corresponding author: Sreenivasan Ponnambalam, School of Molecular and Cellular Biology, University of Leed	ls,
	.eeds LS2 9JT, U.K. E-mail: <u>s.ponnambalam@leeds.ac.uk</u>	
,	Abstract	

The mammalian endothelium which lines all blood vessels responds to soluble factors which 18 19 control vascular development and sprouting. Endothelial cells bind to vascular endothelial 20 growth factor A via two different receptor tyrosine kinases (VEGFR1, VEGFR2) which regulate 21 such cellular responses. The integration of VEGFR signal transduction and membrane 22 trafficking is not well understood. Here, we used a yeast-based membrane protein screen to 23 identify VEGFR-interacting factor(s) which modulate endothelial cell function. By screening 24 a human endothelial cDNA library, we identified a calcium-binding protein, S100A6, which 25 can interact with either VEGFR. We found that S100A6 binds in a calcium-dependent manner 26 to either VEGFR1 or VEGFR2. S100A6 binding was mapped to the VEGFR2 tyrosine kinase 27 domain. Depletion of S100A6 impacts on VEGF-A-regulated signaling through the canonical 28 mitogen-activated protein kinase (MAPK) pathway. Furthermore, S100A6 depletion caused 29 contrasting effects on biosynthetic VEGFR delivery to the plasma membrane. Co-distribution 30 of S100A6 and VEGFRs on tubular profiles suggest the presence of transport carriers that 31 facilitate VEGFR trafficking. We propose a mechanism whereby S100A6 acts as a calcium-32 regulated switch which facilitates biosynthetic VEGFR trafficking from the TGN-to-plasma 33 membrane. VEGFR-S100A6 interactions thus enable integration of signaling and trafficking 34 pathways in controlling the endothelial response to VEGF-A. (197 words) 35 36 Keywords: Endothelial, VEGFR, S100A6, calcium, signaling, trafficking

37

38 Abbreviations: Vascular endothelial growth factor A, VEGF-A; Vascular endothelial growth

39 factor receptor, VEGFR; Calcium-binding S100 protein A6, S100A6;

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41 Introduction

42 Receptor tyrosine kinases (RTKs) are integral membrane proteins and enzymes which regulate 43 essential features of cell, organ, tissue and animal function (Lemmon et al., 2016). RTK binding to 44 exogenous ligands enables the transmission of signals into the cell interior through activation of 45 multiple signal transduction pathways. In spite of numerous studies on different RTKs over the past 50 yrs, we still lack an understanding of how different cells integrate RTK activation, 46 47 signaling and cellular responses. This is further complicated by discovery that post-translational 48 modifications such as phosphorylation and ubiquitination can also modulate trafficking and 49 proteolysis. This is important as the presence of activated RTK complexes at different intracellular 50 locations could activate different signal transduction pathways.

51 An archetypal RTK is a Type I membrane protein with a glycosylated, extracellular N-52 terminus which is used to 'sense' exogenous soluble and membrane-bound ligands (Lemmon and 53 Schlessinger, 2010). Ligand binding transmits conformational changes through the single 54 transmembrane region to the cytoplasmic domain, which activates an ~300 residue tyrosine 55 kinase module comprised of a N- and C-lobes around a central cleft which binds ATP and protein 56 substrates (Endres et al., 2014; Maruyama, 2015; Tatulian, 2015). RTK-mediated tyrosine 57 phosphorylation of multiple substrates activates multiple signal transduction pathways which 58 control different cellular responses such as migration, survival, proliferation and differentiation. 59 Although many RTK phospho-substrates have been identified with specific roles in different 60 aspects of cellular physiology, there is no mechanism to adequately explain how RTK trafficking 61 is regulated in resting and ligand-stimulated conditions to meter RTK bioavailability.

62 One RTK model is the vascular endothelial growth receptor (VEGFRs) comprising VEGFR1, 63 VEGFR2 and VEGFR3 (Bates et al., 2018; Simons et al., 2016). VEGFR2 is a major pro-angiogenic 64 switch which also contributes to tumor angiogenesis (Apte et al., 2019). VEGF-A binds to both 65 VEGFR1 and VEGFR2 with different outcomes and physiological responses (Koch and Claesson-66 Welsh, 2012; Simons et al., 2016; Smith et al., 2015). One of the best characterized factors which 67 interact with VEGFR2 is phospholipase Cy1 (PLy1), whose plasma membrane recruitment 68 promotes PIP₂ hydrolysis leading to cytosolic calcium ion flux and protein kinase C activation 69 (Takahashi et al., 2001). VEGFR2 binds a number of adaptors such as TsAd (Matsumoto et al., 70 2005; Sun et al., 2012), Shc, Grb2, Nck (Guo et al., 1995; Kroll and Waltenberger, 1997), Crk 71 (Stoletov et al., 2001), Shb, Sck, SHP-1, and p66Shc (Simons et al., 2016). VEGFR2 binding to epsin 72 (Rahman et al., 2016) and synectin (Lanahan et al., 2013; Salikhova et al., 2008) suggests that 73 interactions with endocytic regulators facilitates VEGFR2 internalization and delivery to 74 endosomes. However, we were still lacking a mechanism to explain how VEGFR trafficking 75 controls receptor bioavailability for exogenous VEGF-A ligand.

76 In this study, we explored the idea that VEGFR interaction with novel cytosolic factor(s) 77 facilitates integration of signaling and trafficking pathways. We employed a membrane protein-78 based genetic screen to identify VEGFR-interacting cytosolic factors. One such protein was 79 S100A6, a calcium-binding protein which binds both VEGFR1 and VEGFR2. Calcium-dependent 80 S100A6 binding to VEGFR1 and VEGFR2 regulates membrane trafficking and VEGF-A-regulated 81 signal transduction. Our model postulates a feedback circuit involving calcium-dependent 82 protein-protein interactions which modulate VEGFR trafficking and bioavailability at the plasma 83 membrane.

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84 **Results**

85 S100A6 identified as a VEGFR2-binding protein using a membrane Y2H screen

86 There has been a lack of genetic screens using a native VEGFR membrane protein to identify new 87 binding partners and potential regulators. To address this, we used the split ubiquitin membrane 88 yeast two-hybrid (Y2H) system (Johnsson and Varshavsky, 1994; Stagljar et al., 1998) which 89 enables the use of native membrane proteins as 'baits' to screen genome-wide libraries. In this 90 membrane Y2H system, a split ubiquitin polypeptide and the LexA-VP16 transactivator were 91 used to control nuclear yeast gene expression (*Figure 1A*). The 'bait' and 'prey' are tagged with 92 different halves of the ubiquitin molecule, which when brought together in the cytosol, allows 93 activation of a cytosolic ubiquitin-specific protease which cleaves the LexA-VP16 transactivator 94 from the hybrid prey (Figure 1A). Cleaved LexA-VP16 can now translocate into the yeast nucleus, 95 and stimulates auxotrophic gene expression to enable cell growth on defined media (*Figure 1A*). In this study, we fused the complete human VEGFR2 coding sequence to Cub and LexA-VP16 to 96 97 form a 'bait' hybrid protein to screen for binding to interacting factors (protein X) fused to the

98 NubG 'prey' hybrid protein (*Figure 1B*).

99 We then assessed the expression of VEGFR2 hybrid proteins using either the 'prey' or 'bait' 100 plasmid vectors in transformed yeast cells (Figure 1C). Yeast expression of either VEGFR2-Cub-101 LexA-VP16 or VEGFR2-NubG revealed high molecular weight bands ~200-250 kDa 102 corresponding to the predicted size of hybrid proteins (Figure 1C). Probing yeast cells expressing 103 Alg5-LexA-VP16 or VEGFR2-LexA-VP16 hybrid proteins using anti-VP16 antibodies again 104 detected bands of expected sizes (Figure 1D). A positive control PLCy1-SH2 domain (known to 105 interact with VEGFR2), with an engineered HA tag was fused to NubG, expressed in yeast cells, 106 revealing a hybrid protein of expected size (Figure 1E). This PLCy1-SH2-NubG prey construct 107 could now be used as a positive control in subsequent yeast genetic screens.

108 We checked for yeast reporter gene expression comparing bait VEGFR2 with positive and 109 negative controls (Supplement Figure S1). We used the PLCy1-SH2 domain that binds the 110 phosphotyrosine epitope in the VEGFR2 cytoplasmic tail (Guo et al., 1995; Takahashi et al., 2001); 111 (Larose et al., 1995). When fused to the Nub-G prey construct, PLCy1-SH2-Nub promoted a 4-5-112 fold increase (vs. controls) in LacZ (β-galactosidase) activity (Supplement Figure S1), suggesting 113 interaction between VEGFR2 'bait' and PLCy1-SH2 'prey'. We also tested whether VEGFR2 could 114 form homodimers by co-expressing a VEGFR2-NubG 'prey' construct. Again, there was a 5-fold 115 increase in LacZ activity (Supplement Figure S1), indicating that VEGFR2-VEGFR2 homodimers 116 were formed. When control yeast proteins such as Fur4, Ost1 and Alg5 were fused to NubG, LacZ 117 expression was relatively low but higher than 'empty' prey vector (Nub-G) alone. However, when 118 the same proteins are fused to NubI, which causes self-association with the Cub moiety on the 119 bait protein, LacZ expression was increased 5-fold (Supplement Figure S1). 120





123 Figure 1. VEGFR2 expression and interaction analysis in a membrane yeast two-hybrid system. (A) In the 124 split-ubiquitin system, C-terminal of ubiquitin (Cub) and N-terminal of ubiquitin (Nub) spontaneously 125 reconstitute into a native ubiquitin fold that can be recognized by ubiquitin-specific proteases (UBPs). This 126 UBP cleaves the artificial fusion protein to release LexA-VP16 which in turn translocates to the yeast 127 nucleus to activate reporter gene expression. Ile>Gly substitution in Nub (Nub-G) reduces affinity for Cub, 128 blocking spontaneous Nub-Cub re-assembly. UBP-mediated proteolysis and release of VP16 promotes 129 transcription of auxotrophic marker genes which enable survival on histidine or adenine-deficient media. 130 Bacterial β -galactosidase (LacZ) is an additional reporter controlled by LexA-VP16 nuclear translocation. 131 (B) Schematic of the VEGFR2 bait with the full-length human VEGFR2 fused to Cub and LexA-VP16. (C) 132 Expression and detection of VEGFR2-Cub-LexA-VP16 and VEGFR2-NubG (compared to control Alg5-133 LexA-VP16) in yeast by immunoblotting using anti-VEGFR2 antibodies. (D) Expression of Alg5-Cub-LexA-134 VP16 (control) and VEGFR2-Cub-LexA-VP16 hybrid proteins in yeast detected by immunoblotting using 135 anti-VP16 antibodies. Arrowhead indicates full-length VEGFR2 fusion protein. (E) Expression of PLCy1-136 SH2-NubG prey protein in yeast detected by immunoblotting using anti-HA tag antibody. 137

138 We constructed and screened a 'prey' cDNA library of human endothelial proteins fused 139 to NubG-LexA-VP16 (see Materials and Methods). From this screen, we identified a calcium-140 binding protein S100A6 as a potential binding partner for VEGFR2. The S100A6 prey plasmid 141 construct was isolated, re-transformed into yeast cells and compared to a range of controls under 142 defined growth conditions (Supplement Table 1). The S100A6 prey showed yeast growth when co-143 expressed with VEGFR2 bait (Supplement Table 1), indicating protein-protein interactions between 144 the two molecules. LacZ activity assay showed that co-expression of VEGFR2 bait and S100A6 145 prey caused a 5-fold rise in LacZ activity (Supplement Table 1), indicating protein-protein 146 interactions between VEGFR2 and S100A6. Biochemical analysis of S100A6-NubG in yeast cells

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147 revealed a fusion protein of the expected size that contained an engineered HA tag (*Figure S2A*)

and cross-reactive with anti-human S100A6 antibodies (Figure S2B). Probing human endothelial

cells with anti-S100A6 antibodies revealed a low molecular weight band of ~10 kDa (Figure S2C).

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151 S100A6 binds to the VEGFR2 cytoplasmic domain

152 Endothelial cells express both VEGFR1 and VEGFR2, two closely related but distinct gene 153 products with distinct functional roles (Shibuya, 2015; Simons et al., 2016; Smith et al., 2015). To 154 investigate the association of endogenously expressed VEGFRs and S100A6, we immunoisolated 155 detergent-solubilized complexes from endothelial cells and probed for different proteins 156 including using the transferrin receptor as a control (Figure 2A). As expected, VEGFR2 complexes 157 contained S100A6; surprisingly, VEGFR1 complexes also contained S100A6 (Figure 2A). 158 Immunoisolation of S100A6 complexes from endothelial cells revealed the presence of both 159 VEGFR1 and VEGFR2 (Figure 2A). Another membrane protein, the transferrin receptor, was 160 absent from immunoisolated complexes of VEGFR or S100A6 (Figure 2A), indicating specificity 161 in the interaction between S100A6 and VEGFRs.

162 S100A6 belongs to a family of relatively small (~10 kDa) proteins which undergo calcium-163 dependent conformational changes which modulate protein-protein interactions (Rezvanpour 164 and Shaw, 2009; Santamaria-Kisiel et al., 2006). One likelihood is that S100A6 interacts with the 165 VEGFR2 cytoplasmic domain (Figure 1B). To test this possibility, we expressed and purified a 166 recombinant soluble VEGFR2 cytoplasmic domain fragment to assess binding to purified 167 recombinant S100A6 (Figure 2B). Two potential regulatory aspects of VEGFR2-S100A6 168 interactions are calcium ion binding to S100A6 (Donato et al., 2017; Santamaria-Kisiel et al., 2006) 169 and VEGFR2 tyrosine autophosphorylation (Simons et al., 2016; Smith et al., 2015). Interestingly, 170 recombinant VEGFR2 exhibits phosphorylation on residue Y1175, indicating functional tyrosine 171 kinase activity (Figure 2B). We investigated the biochemistry of VEGFR2-S100A6 interactions: 172 VEGFR2 cytoplasmic domain bound to S100A6 in the presence of calcium ions (Figure 2A). 173 VEGFR2-S100A6 complex formation was blocked in the presence of EGTA with no evidence for 174 VEGFR2 association with immobilized S100A6 (Figure 2B). We then evaluated requirement for 175 VEGFR2 tyrosine phosphorylation in binding to S100A6: de-phosphorylated VEGFR2 still bound 176 immobilized S100A6 in the presence of calcium ions similar to phosphorylated VEGFR2 (Figure 177 2B). Such VEGFR2-S100A6 interactions are thus calcium-dependent but do not require VEGFR2 178 phosphorylation.

179 As our data suggested that the VEGFR2 cytoplasmic domain binds to S100A6 (Figure 1), we 180 asked whether the VEGFR1 cytoplasmic domain protein could also bind to S100A6 (Figure 2B). 181 This VEGFR1 cytoplasmic domain protein bound to immobilized S100A6 in the presence of 182 calcium ions (Figure 2C). Again, VEGFR1-S100A6 complex formation was inhibited by the 183 addition of EGTA (Figure 2C). To investigate this further, we used the membrane Y2H system to 184 assess whether VEGFR1 displayed interaction with S100A6 (Supplement Table 1). Yeast cells co-185 expressing the VEGFR1-Cub-LexA-VP16 'bait' and the S100A6-Nub-G 'prey' showed auxotrophic 186 growth and LacZ expression (Supplement Table 1). These findings are consistent with calcium-187 dependent VEGFR1-S100A6 complex formation.



189

190 Figure 2. VEGFR and S100A6 complexes exhibit calcium-dependence. (A) Immunoisolation of VEGFR or 191 S100A6 complexes followed by immunoblot analysis. Whole cell lysates (WCL) were subjected to detergent 192 lysis (see Materials and Methods) and VEGFR2, VEGFR1 and S100A6 complexes isolated before SDS-PAGE 193 and immunoblotting (IB). Goat anti-VEGFR2, goat anti-VEGFR1 or rabbit anti-S100A6 antibodies were 194 used to isolate VEGFR or S100A6 complexes respectively. Molecular weight markers (kDa) and respective 195 proteins are indicated in the panel. Transferrin receptor (TfR) was used as a negative control. (B) 196 Recombinant proteins comprising soluble VEGFR2 cytoplasmic domain or de-phosphorylated VEGFR2 197 cytoplasmic domain was incubated with soluble GST-S100A6 or GST in the presence of 1 mM calcium ions 198 or 1 mM EGTA. This was followed by incubation with glutathione-agarose beads, centrifugation and brief 199 washes with buffer. Bound proteins were analyzed on 12% SDS-PAGE together with purified 200 phosphorylated VEGFR2 (lane 9) or de-phosphorylated VEGFR2 (lane 10). Immunoblotting was carried 201 out using mouse anti-phosphotyrosine (pY20), rabbit anti-VEGFR2-pY1175 or sheep anti-VEGFR2 202 cytoplasmic domain antibodies. (C) Similar experiments carried out using the soluble VEGFR1 cytoplasmic 203 domain incubated with soluble GST-S100A6 or GST in the presence of 1 mM calcium ions or 1 mM EGTA. 204 This was followed by incubation with glutathione-agarose beads, centrifugation and brief washes with 205 buffer. Bound proteins were analyzed on 12% SDS-PAGE and immunoblotting using sheep anti-VEGFR1 206 cytoplasmic domain antibodies.

207

208 Biochemistry of VEGFR2-S100A6 interactions

209 The interaction between two molecules can be described by biochemical parameters. We explored 210 the interactions between S100A6 and VEGFR2 using different assays (Figure 3). First, surface 211 plasmon resonance (SPR) was used to measure S100A6 binding to the VEGFR2 cytoplasmic 212 domain in the presence of calcium ions (Figure 3A). SPR data showed that titration of S100A6 213 displayed dose-dependent kinetics of binding to immobilized VEGFR2 in the presence of calcium 214 ions (Figure 3A). Based on these SPR data, the VEGFR2-S100A6 dissociation constant (K_d) was 215 calculated to be ~0.2 µM. S100A6 binding to immobilized VEGFR2 was abolished in the presence 216 of EGTA (Figure 3A).

The S100A6 protein (90 residues) binding to the larger VEGFR2 cytoplasmic domain (568 residues) was mapped using deletion analysis (*Figure 3B*). We separated the 568 VEGFR2 cytoplasmic domain into the juxtamembrane (JM) region, the tyrosine kinase domain comprising the N- and C- lobes (TK-N, TK-C) including a kinase insert region, and a flexible C-terminal tail (*Figure 3B*). These different portions of the VEGFR2 cytoplasmic domain were fused to the maltose-binding protein (MBP), and MBP-VEGFR2 hybrid proteins were tested for binding to

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223	GST-S100A6 (Figure 3C). S100A6 protein strongly bound	to the 568 residue VEGFR2 cytoplasmic
224	domain (Figure 3C). Deletion analysis showed that the	VEGFR2 kinase domain (329 residues)
225	bound S100A6 (<i>Figure 3C</i>). Neither the juxtamembrane re	egion nor the C-terminal tail showed any

significant binding to S100A6 (*Figure 3C*).



227

228 Figure 3. Interaction of the VEGFR2 cytoplasmic domain with S100A6. (A) SPR analysis of recombinant 229 immobilized soluble VEGFR2 cytoplasmic domain binding to S100A6 the presence of 1 mM divalent 230 calcium ions, or 1 mM EGTA (no Ca2+). Soluble VEGFR2 cytoplasmic domain was immobilized on the chip 231 as described in Materials and Methods. GST-S100A6 solutions of 10 µM, 5 µM, 2 µM or 1 µM in buffer 232 supplied with 1 mM calcium ions or 1 mM EGTA was flowed over immobilized VEGFR2. The response 233 unit (RU) was recorded using evaluation software. (B) Schematic view of the VEGFR2 protein showing the 234 various regions. The juxtamembrane (JM), tyrosine kinase domain (KD) and C-terminal tail (CT) are 235 indicated on the line diagram. Sequences corresponding to residues from the VEGFR2 juxtamembrane 236 (JM), kinase domain (KD) and C-terminal tail (CT) were fused to MBP and used in binding studies. (C) 237 Interaction of MBP-VEGFR2 proteins with S100A6. The VEGFR2 cytoplasmic domain fused to MBP (Cyto), 238 N-proximal juxtamembrane region (JM), kinase domain alone (KD) and cytoplasmic tail (CT) were tested 239 for their ability to bind either GST or GST-S100A6 in a pull-down assay. The upper panel shows an 240 immunoblot for GST to detect GST-S100A6 fusion; lower panel shows the Ponceau S stain showing the 241 MBP-VEGFR2 proteins used in the assay. Molecular weight markers are indicated.

242

243 S100A6 modulates VEGFR1 and VEGFR2 trafficking, modification and turnover

VEGFR1 and VEGFR2 display complex patterns of steady-state and ligand-stimulated distribution in endothelial cells (Ewan et al., 2006; Gampel et al., 2006; Lampugnani et al., 2006; Mittar et al., 2009). One possibility is that calcium-dependent S100A6 interactions with VEGFR2 and/or VEGFR1 modulates trafficking and turnover. To assess this, we compared S100A6 with another S100 family member (S100A10) also expressed in endothelial cells (Bao et al., 2012), using protein knockdown using RNAi followed by analyses of VEGFR trafficking and cellular distribution (*Figure 4*).

251 Depletion of S100A6 caused a significant rise in overall VEGFR2 levels but this did not affect 252 other membrane proteins including VEGFR1 or VEGF co-receptors, the neuropilins (NRP1, 253 NRP2) (Figure 4A). In contrast, depletion of S100A10 did not significantly affect VEGFR or control 254 membrane protein levels (Figure 4A). Quantification showed that S100A6 depletion caused ~80% 255 rise in total VEGFR2 levels, with little or no significant effects on membrane-bound or soluble 256 VEGFR1 levels (*Figure 4B*). We used cell surface biotinylation to assess plasma membrane VEGFR 257 pools under these conditions (Figure 4C). Increased mature plasma membrane VEGFR2 levels 258 were detected upon S100A6 knockdown (*Figure 4C*). Quantification showed ~2.5-fold increase in 259 mature plasma membrane VEGFR2 levels compared to controls (Figure 4D). Under basal or 260 resting conditions, S100A6-depleted cells showed no significant change in cell surface membrane-

bound or soluble VEGFR1 compared to controls (*Figure 4D*).



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263 Figure 4. S100A6 requirement for VEGFR1 and VEGFR2 trafficking. Endothelial cells were subjected to 264 RNAi on S100 proteins and analyzed for VEGFR2 trafficking. (A) Endothelial cells subjected to control, scrambled, S100A6 or S100A10 siRNA treatments were lysed and immunoblotted for various proteins 265 266 indicated. Molecular weights of markers are indicated. (B) Quantification of relative protein levels under 267 different conditions of RNAi. Color coding indicates non-transfected (red), scrambled siRNA (blue), 268 S100A6 knockdown (purple) and S100A10 knockdown (grey). Error bars indicate ±SEM (n>3).*, p<0.05. (C) 269 Endothelial cells subjected to control, scrambled, S100A6 or S100A10 siRNA treatments followed by cell 270 surface biotinylation, cell lysis and purification of biotinylated proteins. Whole cell lysates and purified 271 proteins were immunoblotted for the various proteins indicated. Molecular weights of markers are also 272 indicated. (D) Quantification of relative protein levels under different conditions of RNAi. Color coding 273 indicates non-transfected (red), scrambled siRNA (blue), S100A6 knockdown (purple) and S100A10 274 knockdown (grey). Error bars indicate +SEM (n>3). ***, p<0.001. (E) Quantification of VEGF-A-regulated 275 mature VEGFR1 trafficking to the plasma membrane using cell surface biotinylation. Endothelial cells 276 subjected to control, scrambled or S100A6 siRNA treatments followed by VEGF-A (10 ng/ml) stimulation, 277 followed by cell surface biotinylation, cell lysis and purification of biotinylated proteins before 278 immunoblotting. Color coding indicates non-transfected (red), scrambled siRNA (blue) and S100A6 279 knockdown (purple).

280

281 A previous study showed that biosynthetic VEGFR1 undergoes VEGF-A-stimulated and 282 calcium-dependent trafficking from the distal Golgi to the plasma membrane (Mittar et al., 2009). 283 One possibility was that S100A6 is involved in this calcium-dependent Golgi-to-plasma 284 membrane trafficking step. To test this idea, we used cell surface biotinylation of S100A6-285 depleted cells to assess VEGFR1 plasma membrane levels in resting or VEGF-A-stimulated cells 286 (Figure 4E). Upon VEGF-A stimulation, we detected a time-dependent, ~2.5-fold increase in 287 plasma membrane VEGFR1 levels over a 60 min time period (Figure 4E). However, S100A6 288 knockdown caused a complete block in VEGF-A-stimulated VEGFR trafficking to the plasma 289 membrane (Figure 4E). Depletion of S100A6, but not S100A10, thus modulates both VEGFR1 and 290 VEGFR2 trafficking and plasma membrane levels.

S100A6 and VEGFR Trafficking in Endothelial Cells Bao et al. S100A6 regulates VEGFR2 bioavailability and VEGF-A-stimulated signal transduction 292 293 VEGFR2 trafficking influences VEGF-A-stimulated signaling from the cell surface (Bruns et al., 294 2010; Ewan et al., 2006; Lampugnani et al., 2006; Manickam et al., 2011; Yamada et al., 2014). Based 295 on our findings in this study, we then asked whether S100A6 regulation of plasma membrane 296 VEGFR2 levels regulates VEGF-A-stimulated signal transduction events (Figure 5). We monitored 297 2 different VEGFR2 phosphotyrosine epitopes (pY1175, pY1214) which exhibit different kinetics 298 upon VEGF-A stimulation (Fearnley et al., 2016). As expected, VEGFR2-pY1175 is rapidly 299 generated in response to VEGF-A stimulation (Figure 5A). Upon S100A6 knockdown, VEGFR2-300 pY1175 levels are elevated (Figure 5A), corresponding to ~5-fold magnitude increase compared to 301 controls (Figure 5B). However, the kinetics of VEGFR2-pY1175 appearance, time to peak and 302 decline were similar in both control and S100A6-depleted endothelial cells (Figure 5B). 303 Interestingly, VEGFR2-pY1214 is detected under basal conditions in line with previous studies 304 (Fearnley et al., 2016); such signaling is also substantially elevated upon S100A6 knockdown in 305 VEGF-A-stimulated cells (Figure 5A).



306

Figure 5. Modulation of VEGFR2 signaling by S100A6. Endothelial cells were subjected to RNA interference
 and analyzed by (A) immunoblotting, (B) quantification of immunoblot data. Endothelial cells subjected to
 control, scrambled or S100A6 siRNA treatments were stimulated with VEGF-A (10 ng/ml), lysed and
 immunoblotted for the VEGFR2-pY1175 epitope and relative levels quantified. The blots were also probed
 with anti-ERK1/2 and anti-phospho-ERK1/2 antibodies to check for canonical MAPK signaling. Blotting for
 tubulin was used as an additional loading control in these experiments. Error bars indicate ±SEM (n>3). *,
 p<0.05.

314

315 We also carried out confocal microscopy to ascertain subcellular VEGFR and S100A6 316 localization (Figure 6A, 6B). Steady-state VEGFR2 distribution shows localization to the plasma 317 membrane, endosomes and juxtanuclear Golgi region (Figure 6A). In contrast, S100A6 is widely 318 distributed in the cytosol and nucleus, with occasional staining of tubular profiles emanating 319 from the Golgi region (Figure 6A, short arrows). Overlay images suggests co-distribution and 320 close proximity of VEGFR2 and S100A6 (Figure 6A, boxed). Analysis of the VEGFR1 and S100A6 321 (Figure 6B) showed also showed co-distribution in the Golgi region (Figure 6B, long arrows). Co-322 labeling of VEGFR1 and S100A6 elongated tubular profiles emanating from the Golgi region was

also detected (*Figure 6B*, boxed).





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Figure 6. Co-distribution of VEGFRs and S100A6. Endothelial cells were fixed and processed for wide-field
 deconvolution microscopy analysis (see Materials and Methods) of (A) VEGFR2, or (B) VEGFR1 vs. S100A6
 on fixed endothelial cells. Triple color overlay shown in extreme right-hand panel. Bar, 10 µm. Arrows
 and arrowheads denote co-distribution of VEGFR and S100A6 along tubular profiles. Each image shown
 comprises 20-35 optical sections to better visualize tubular profiles.

We then evaluated the 3-D structures of the S100A6 and VEGFR2 cytoplasmic domain using *in silico* modelling (*Figure 7*). Comparison of the structures of free and calcium-bound S100A6 shows relatively large movements of helix H3 (*Figure 7A*). *In silico* docking studies using the VEGFR2 tyrosine kinase domain suggests that calcium-bound S100A6 binds close to the cleft of the tyrosine kinase module (*Figure 7B*). It is unclear whether VEGFR tyrosine kinase activity and calcium-S100A6 binding are functionally coupled.

337

338 Discussion

339 How does a cell integrate membrane receptor bioavailability for a specific ligand? In the case of 340 VEGF-A binding to two different membrane receptors, VEGFR1 and VEGFR2, different 341 pathways of signaling, trafficking and turnover need to be integrated to control cellular responses 342 such as cell migration, proliferation and tubulogenesis. Up to now, we lacked molecules that 343 could bridge VEGFR signaling and trafficking. Herein, we now present evidence that a calcium-344 dependent cytosolic protein, S100A6, binds both VEGFR1 and VEGFR2 to integrate signaling and 345 trafficking pathways. Five lines of evidence support this conclusion. Firstly, a genetic screen of 346 human endothelial proteins identified S100A6 as a binding partner for VEGFR2. Second, 347 membrane Y2H assay shows that either VEGFR2 or VEGFR1 can interact with S100A6. This was 348 confirmed by the detection of stable complexes of S100A6 with either VEGFR1 or VEGFR2 in 349 endothelial cells. Third, S100A6 binds the VEGFR2 cytoplasmic domain in vitro, with sub-350 micromolar binding affinity (Kd) and displays calcium-dependence. Furthermore, the VEGFR1 351 cytoplasmic domain also binds to S100A6 in a calcium-dependent manner. S100A6 binding to 352 VEGFR2 maps to the tyrosine kinase module. Fourth, S100A6 regulates VEGFR1 and VEGFR2 353 trafficking with different functional outcomes. Whereas S100A6 depletion causes dysregulated

Bao et al.S100A6 and VEGFR Trafficking in Endothelial CellsVEGFR2 trafficking and increased plasma membrane levels, loss of S100A6 completely blocks

- 355 VEGFR1 Golgi-to-plasma membrane trafficking. Finally, S100A6 influences VEGFR2 plasma
- 356 membrane bioavailability by modulating VEGF-A-regulated VEGFR2 tyrosine phosphorylation
- 357 and downstream canonical MAPK signaling.



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359 Figure 7. Models of VEGFR2/S100A6 interaction and membrane trafficking. (A) Crystal structures of 360 calcium-free (grey: PDB ID 1K9P) and calcium-bound (blue: PDB ID 1K96) forms of S100A6. There is a large 361 movement of Helix III (brown arrow) upon binding Ca²⁺ (red spheres). (B) Energy minimized model of 362 VEGFR2 kinase domain (PDB ID: 4AGD) bound to calcium-bound S100A6 (PDB ID: 1K96) using in silico 363 docking with HADDOCK 2.2 (see Materials and Methods). The kinase domain insert between the N-364 terminal (NTD: light blue) and C-terminal (CTD: grey) segments of the kinase domain is not resolved in 365 VEGFR2 tyrosine kinase domain structure (PDB ID: 4AGD). The regulatory loop that contains the 366 autophosphorylation sites Y1054 and Y1059 is shown green, with the conserved DFG motif highlighted 367 (purple). Sunitinib (Sutent: orange) is bound to the kinase domain. The major contacts contributed by 368 S100A6 are via the interhelical Ca²⁺-binding loops. (C) Regulation of TGN-to-plasma membrane trafficking 369 of VEGFR1 and VEGFR2 requiring S100A6. Depicted are 3 parallel transport routes: a constitutive transport 370 step accessed by VEGFR2, a calcium-dependent trafficking route from the TGN which is dependent on 371 S100A6 binding to the cytoplasmic domains of VEGFR1 and VEGFR2 cargo for inclusion into a new class 372 of transport carriers, and the dense-core secretory granule route.

373

374 Our study supports a mechanism where at least two trafficking routes regulate VEGFR 375 delivery from the trans-Golgi network (TGN) to the plasma membrane (Figure 7C). In higher 376 eukaryotes, a constitutive TGN-to-plasma membrane anterograde trafficking step is utilized by 377 a majority of soluble and membrane-bound secretory proteins (Guo et al., 2014; Pakdel and von 378 Blume, 2018). Our findings that S100A6 depletion leading to elevated VEGFR2 levels at the 379 plasma membrane suggests dysregulation in TGN-to-plasma membrane trafficking. One 380 explanation is that VEGFR2 utilizes both constitutive and calcium-regulated trafficking routes to 381 exit the TGN (Figure 7C). Newly synthesized VEGFR2 also accumulates within the Golgi 382 (Manickam et al., 2011); it was postulated constitutive TGN-to-plasma membrane trafficking 383 enables replenishment of the cell surface VEGFR2 pool undergoing endocytosis, recycling or 384 degradation (Ewan et al., 2006; Jopling et al., 2011). Our studies now suggest that plasma 385 membrane VEGFR2 bioavailability is also dependent on calcium-regulated TGN-to-plasma 386 membrane trafficking event (Figure 7C). VEGFR2 levels are thus 'metered' by 2 parallel 387 anterograde trafficking steps to ensure that biosynthetic VEGFR2 delivery is synchronized with 388 endocytosis of plasma membrane VEGFR2 for delivery to endosomes.

Endothelial cells display VEGFR2 localization to Golgi, plasma membrane and endosomes
(Ewan et al., 2006; Gampel et al., 2006; Manickam et al., 2011). Co-distribution of VEGFR1,
VEGFR2 and S100A6 on tubular profiles emanating from a juxtanuclear Golgi region in

Bao et al.S100A6 and VEGFR Trafficking in Endothelial Cells392endothelial cells indicate transport carriers which mediate VEGFR cargo delivery to the plasma393membrane. VEGFR2 Golgi trafficking shows dependence on cytosolic factors such as t-SNARE394syntaxin 6 (Manickam et al., 2011), the KIF13B microtubule motor (Yamada et al., 2014) and the395Myo1c actomyosin motor (Tiwari et al., 2013). However, the cytoskeletal machinery involved in396this calcium-regulated TGN-to-plasma membrane trafficking step (Figure 7C) is at present ill-397defined.

398 VEGFR1 intracellular localization is complicated by a lack of clear functional roles for this 399 RTK. Generally, VEGFR1 is thought to act as a 'VEGF sink' which sequesters ligand and acts as a 400 negative regulator of angiogenesis, but also modulates some aspects of cancer cell proliferation 401 (Autiero et al., 2003; Jones et al., 2009; Lichtenberger et al., 2010; Yang et al., 2006). Different 402 studies suggest VEGFR1 localization to Golgi (Mittar et al., 2009) and nuclear (Lee et al., 2007; 403 Zhang et al., 2010) compartments. However, VEGFR1 plasma membrane levels are elevated by 404 cytosolic calcium ion flux in endothelial cells (Mittar et al., 2009) and cardiomyocytes (Yang et al., 405 2015), suggesting that VEGFR1 can undergo calcium-dependent TGN-to-plasma membrane 406 trafficking in different cell types. Our study now provides a mechanism to explain this 407 phenomenon with the finding that calcium-dependent binding of S100A6 to the VEGFR1 408 cytoplasmic domain. Depletion of S100A6 levels blocked VEGF-A-stimulated VEGFR1 delivery 409 to the plasma membrane, consistent with VEGFR1 cargo sequestration into calcium-regulated 410 TGN-to-plasma membrane transport carriers (Figure 7C). Interestingly, the VEGFR1 Golgi pool 411 shows partial co-distribution with TGN46, a standard marker for the human TGN (Mittar et al., 412 2009). One explanation is that the mammalian TGN is more extensive than currently postulated, 413 with steady-state VEGFR1 residence within a TGN-like subcompartment. From this location 414 VEGFR1 membrane cargo is delivered to the plasma membrane via this S100A6-regulated 415 trafficking step.

416 How can our proposed model (Figure 7C) be reconciled with other TGN trafficking events? 417 The TGN is a site of multiple sorting, packaging and transport events, including constitutive and 418 regulated secretion (Guo et al., 2014). In endothelial cells, the biogenesis of electron dense 419 cylindrical Weibel-Palade bodies (WBPs) from the TGN precedes requirement for a calcium-420 regulated stimulus to undergo docking and fusion with the plasma membrane (McCormack et 421 al., 2017). However, there is little or no evidence of VEGFR1 or VEGFR2 association with WBP 422 trafficking. This then raises the question whether such a calcium-regulated trafficking event 423 (*Figure 7C*) is endothelial-specific or exists in other cell types. Interestingly, there are similarities 424 to calcium-regulated TGN dynamics in immortalized cell lines (Pakdel and von Blume, 2018; von 425 Blume et al., 2012) and neurons (Mikhaylova et al., 2010; Mundhenk et al., 2019). The TGN 426 resident and calcium-binding protein Cab45 is in close proximity to a calcium pump, SPCA1 427 (Deng et al., 2018; von Blume et al., 2012), but how this is linked to constitutive or regulated 428 protein secretion was unknown. Recent studies suggest another member of the S100 family, 429 S100A10, is involved in Weibel-Palade body exocytosis (Chehab et al., 2017). Our studies now 430 suggest existence of a specialized calcium-regulated trafficking route from the TGN-to-plasma 431 membrane in higher eukaryotes.

432 S100A6-VEGFR interactions involves binding to the tyrosine kinase region (*Figure 7B*).
433 Deletion analysis of the VEGFR2 cytoplasmic domain maps calcium-S100A6 binding to the
434 tyrosine kinase module. In this context, de-phosphorylation of the VEGFR2 cytoplasmic domain

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435 at Y1175 (within the carboxy-proximal tail region) does not significantly affect VEGFR2-S100A6 436 complex formation. Interestingly, VEGFR2 undergoes tyrosine phosphorylation at 6-8 distinct 437 epitopes, some of which are present within the tyrosine kinase domain. It remains to be 438 determined whether other phosphotyrosines hinder or promote S100A6 recruitment, and thus 439 influence VEGFR TGN-to-plasma membrane trafficking. Recently, it was reported that 440 SUMOylation of the VEGFR2 cytoplasmic domain mediates Golgi targeting (Zhou et al., 2018). 441 SUMOvlation of VEGFR2-K1270 within the flexible carboxy-terminal tail (Zhou et al., 2018) is 442 unlikely to modulate S100A6 binding to the tyrosine kinase region (residues 833-1162; Figure 7B) 443 in this context.

444 Our study provides a new mechanism where newly synthesized membrane cargo trafficking 445 to the plasma membrane is dependent on integration with signal transduction pathways. Here, 446 activation of plasma membrane receptors which trigger rise in second messenger levels such as 447 calcium ions causes conformational changes in S100A6 to enable binding to VEGFRs, and cargo 448 selection for TGN-to-plasma membrane trafficking. Importantly, our findings also provide a 449 mechanistic explanation for how trafficking and secretion of newly synthesized membrane cargo 450 is synergized with plasma membrane signaling for replenishment of membrane receptors. In this 451 context, there is increasing evidence that S100 protein family members e.g. S100A10, regulates 452 biosynthetic trafficking of sodium (Okuse et al., 2002) and potassium (Girard et al., 2002) 453 channels, and regulates Weibel-Palade body exocytosis (Chehab et al., 2017).

454 Our proposed mechanism enables the endothelial cell to integrate plasma membrane 455 VEGFR2 activation, downstream signal transduction with secretion of newly synthesized VEGFRs to regulate plasma membrane VEGFR bioavailability for VEGF-A. This calcium-456 457 regulated mechanism acts as a feedback loop that synchronizes VEGFR2 activation with 458 controlling arrival of both VEGFRs at the cell surface, thus enabling tight control of cellular 459 responses to exogenous VEGF-A. One of the puzzles in understanding VEGF biology is the 460 existence of 2 receptor tyrosine kinases (VEGFR1, VEGFR2) that bind with differing affinity to 461 the same ligand, VEGF-A (Ewan et al., 2006; Vaisman et al., 1990). VEGFR2 plays a major role in 462 angiogenesis but the role of VEGFR1 is less clear (Shibuya and Claesson-Welsh, 2006). VEGFR1 463 is postulated to negatively regulate angiogenesis by acting as a 'VEGF trap'; however, VEGFR1-464 specific ligands such as PIGF are functionally implicated in some cancers (Fischer et al., 2008). 465 One likelihood is that calcium-stimulated VEGFR1 arrival at the cell surface (and higher affinity 466 for VEGF-A) dictates preferential sequestration of VEGF-A, thus reducing VEGF-A 467 bioavailability to VEGFR2. VEGFR1 could thus not only act as a ligand trap under these 468 situations, but have a different signaling role distinct from the pro-angiogenic VEGFR2. In tumor 469 angiogenesis, pathological levels of exogenous VEGF-A could not only promote calcium-470 stimulated translocation of both VEGFR1 and VEGFR2 to the plasma membrane, but continue to 471 promote pro-angiogenic signaling through VEGFR2. Future work is needed to explore this 472 mechanism in the context of health and disease states.

473

474 Materials and Methods

475 Gene Manipulation

476 The full-length human VEGFR2 open reading frame was amplified using PCR using pVEGFR2-

477 EGFP (Jopling et al., 2011). The PLCγ1-N-SH2 domain (Larose et al., 1995) was amplified by using

S100A6 and VEGFR Trafficking in Endothelial Cells Bao et al. 478 pGEX-2T-PLCy1-N-SH2 plasmid. Full-length VEGFR2 was cloned into both the pBT3-SUC 'bait' 479 and pPR3-N 'prey' plasmids. The PLCy1-SH2 domain was cloned into pPR3-N 'prey' plasmid. 480 Control Fur4-NubG, Ost1-NubG, Fur4-NubI, Ost1-NubI, pBT3-SUC2, pPR3-N, Alg5-NubG, Alg-481 NubI plasmids were already provided as controls. All recombinant constructs were checked by 482 DNA sequence analysis. Total endothelial RNA was extracted from confluent early passage 483 endothelial cells using Trizol (Sigma-Aldrich). 1st strand cDNA was synthesized using SMART 484 (Switching Mechanism At 5' end of the RNA Transcript) technique using a kit (Easyclone cDNA 485 Synthesis). The ds cDNA fragments with SfiI restriction enzyme sites was directionally cloned 486 into the pPR3-N 'prey' plasmid which will express NubG-cDNA 'prey' fusion proteins. This 487 cDNA library was transformed into yeast strain NMY51 containing the VEGFR2 bait by the 488 lithium acetate transformation method. The colonies grown on media lacking histidine and 489 adenine (SD-LWHA) plate were tested for β-galactosidase activity using an X-gal filter test and 490 further β -galactosidase activity assay. Plasmids isolated from yeast colonies expressing β -491 galactosidase were transformed into E. coli for plasmid purification and DNA sequencing. DNA 492 sequence analysis was carried out using software packages at the European Bioinformatics 493 Institute (Hinxton, UK).

494

495 Construction of Endothelial cDNA Library and Membrane Y2H analysis

496 Total RNA was extracted from confluent early passage (P0-P2) endothelial cells using Trizol 497 (Sigma-Aldrich). 1st strand cDNA was synthesized using the Easyclone cDNA library kit with 498 SMART (Switching Mechanism At 5' end of the RNA Transcript) technique. The cDNA fragments 499 with SfiI restriction enzyme sites were directionally cloned into the pPR3-N 'prey' plasmid which 500 expressed these NubG-cDNA constructs as 'prey' hybrid proteins. The cDNA library were 501 transformed into NMY51 S.cerevisiae strain containing VEGFR2 bait construct by the lithium 502 acetate transformation. Yeast colonies which grew on media lacking histidine and adenine (SD-503 LWHA) were picked and re-checked for β -galactosidase (LacZ) expression using the X-gal filter 504 test. Plasmids were isolated from blue X-gal positive colonies and retransformed into the XL-1 505 Blue E. coli strain. Positive plasmids were sequenced and re-checked by rounds of transformation, 506 expression and verification in the yeast assay. Quantification of β-galactosidase (LacZ) expression 507 was done by picking several yeast colonies for each interaction pair. A high throughput β -508 galactosidase activity kit was used to evaluate expression of this marker.

509

510 Pull-down assays for protein-protein interactions

511 Bacterial cultures of 50-500 ml expressing GST, MBP or hexahistidine-tagged constructs were 512 lysed in buffer (1% (w/v) Triton X-100, protease cocktail inhibitors,1 mM PMSF in PBS), then 513 subjected to rapid purification by sonication before incubation with glutathione-agarose, amylose 514 resin or nickel-agarose beads at 4°C for 2 h. Beads were washed 3 times, and purified proteins 515 eluted using glutathione, maltose or imidazole respectively. In binding studies, immobilized 516 proteins were not eluted but incubated with purified test protein in non-ionic detergent buffer 517 (150 mM NaCl, 10 mM Tris, 1% (w/v) NP-40, 1 mM Na₃VO₄, 10 mM NaF, 1 mM CaCl₂ or 1 mM 518 EGTA, protease cocktail inhibitor mix, 1 mM PMSF) for 2 h at 4°C. These were then briefly 519 centrifuged and washed with lysis buffer 3 times. 2X Sample buffer containing 5% β-

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523 Surface plasmon resonance (SPR)

524 Surface plasmon resonance was used to analyze protein-protein affinity. EDC (N-ethyl-N'-525 (dimethylaminopropyl)-carbodiimide) / NHS (N-hydroxy- succinimide) was used to activate a 526 CM5 sensor chip flow cells #1-3. Purified soluble VEGFR2 (0.2 mg/ml in 200 mM sodium acetate, 527 pH 5.5) was injected onto flow cell 2 to generate surface densities of 4000 RU. Rabbit anti-GST 528 antibody in 0.2 M sodium acetate buffer was injected onto flow cell #3 to generate surface 529 densities of 19 000 RU. Flow cell 1 was used as an activated blank flow cell. Subsequently 1 M 530 ethanolamine pH 8.5 was injected onto each flow cell to quench unreacted esters. Different 531 concentrations of recombinant GST-S100A6 or GST in running buffer (10 mM Tris-HCl pH 7.3, 532 150 mM NaCl, 0.005% (v/v) surfactant P20) containing 1 mM EDTA or 1 mM CaCl₂ was injected 533 for 5 min at a flow rate of 10 µl/min sequentially over the three flow cells. Association and 534 dissociation phases were recorded for 5 min for each reaction. Every cycle was finished with two 535 injections of 10 mM glycine pH 1.9 (20 sec) to remove non-covalently bound proteins from the 536 chip surface. Non-specific binding was subtracted from the activated blank flow cell for every 537 cycle. The analyses were performed on a Biacore 3000 system and the data were evaluated with 538 BIAevaluation 3.2 software.

539

540 Cell culture, protein knockdown and immunoblotting

541 Human umbilical vein endothelial cells (HUVECs) were cultured and analyzed as previously 542 described (Fearnley et al., 2014). For immunoprecipitation of protein complexes from HUVECs 543 were lysed in buffer containing 1% (w/v) n-dodecyl-β-D-maltoside, 150 mM NaCl, 25 mM Tris 544 pH 7.5, 1 mM Na₃VO₄, 10 mM NaF, 1 mM CaCl₂, protease cocktail inhibitors and 1 mM PMSF. 545 0.5-1 µg of purified goat anti-VEGFR2, goat anti-VEGFR1 or mouse anti-S100A6 antibodies were 546 added to the soluble supernatant. After 16-24 h incubation, protein G-agarose beads were added 547 for another 2 h, briefly centrifuged and washed 3 times with lysis buffer. Samples were 548 resuspended in 2X reducing sample buffer, boiled and subjected to SDS-PAGE and 549 immunobotting. For signaling experiments, HUVECs were starved for 2 h in media lacking 550 growth factors but containing 0.2% (w/v) BSA. VEGF-A (10 ng/ml) was added for different times 551 and cells lysed in 2% SDS, PBS, protease cocktail inhibitors, 1 mM PMSF. 20 µg of total cell lysate 552 was subjected to SDS-PAGE and immunoblotting. Band intensities were quantified using a 553 digital system as previously described(Fearnley et al., 2014). For cells subjected to RNAi, 554 endothelial cells were incubated under mock-transfection conditions or incubated with 10 nM 555 siRNA duplex as previously described (Fearnley et al., 2016; Smith et al., 2017). Cell surface 556 biotinylation and analysis was carried out as previously described (Fearnley et al., 2016).

557

558 **Confocal microscopy**

559 HUVECs were grown on glass coverslips and fixed with 3% (w/v) paraformaldehyde, quenched

560 with 50 mM ammonium chloride, washed with PBS and subjected to a 5 min permeabilization

- 561 with 0.2% (w/v) TX-100. Non-specific binding sites were blocked by incubation with 0.2%
- 562 BSA/PBS blocking buffer before incubation in primary antibodies such as goat anti-VEGFR1 or

S100A6 and VEGFR Trafficking in Endothelial Cells Bao et al. 563 goat anti-VEGFR2 in combination with mouse anti-S100A6 (1 @g/ml) for 16-20 h. After extensive 564 washes, cells were incubated with AlexaFluor 488 donkey anti-goat and AlexaFluor 594 donkey 565 anti-mouse conjugates (1 @g/ml) for 1-2 h. After washes, coverslips were inverted on a drop of 566 mounting medium and sealed on a glass slide. Samples were viewed using a DeltaVision wide-567 field deconvolution microscope and 0.365 µm optical sections collected as previously described 568 (Mittar et al., 2009). Each wide-field image shown is comprises a stack of 20-35 optical sections to 569 better visualize 3-D structures such as tubules. 570

571 S100A6 and VEGFR2 modeling

572 A structural model of the VEGFR2 tyrosine kinase domain (PDB ID:1VR2) (McTigue et al., 1999) 573 or bound to the tyrosine kinase inhibitor Sunitinib (PDB ID: 4AGD) was used to dock the human 574 calcium-bound S100A6 (PDB ID: 1K96) and calcium-free (PDB ID: 1K9P) (Otterbein et al., 2002). 575 Protein structures were prepared for docking using Biovia Discovery Studio Modelling 576 Environment, Release 3.5 (Biovia Software Inc., San Diego). A model for the complex was 577 developed using the high ambiguity-driven protein-protein docking approach, HADDOCK 578 (Dominguez et al., 2003), for in silico docking of the individual protein structures (using the 579 experimentally determined structures). The Naccess program (Hubbard and Thornton, 1993) was 580 utilized to analyze the solvent-accessible residues which were defined as the active residues for 581 the docking protocol. Default Haddock parameters were used. The resultant docked poses were 582 analyzed using Biovia Discovery Studio 3.5 for inter-molecular interactions. Further modelling 583 studies were carried out using HADDOCK 2.2 (van Zundert et al., 2016) and CPORT (de Vries 584 and Bonvin, 2011).

585

586 Quantification and statistical analysis

587 We used one-way analysis of variance (ANOVA) followed by Tukey's post-hoc test or two-way
588 ANOVA followed by Bonferroni multiple comparison test using GraphPad Prism software (La
590 Like MCAA Given if it is a biffer on the test of test of

Jolla, USA). Significant differences between control and test groups were evaluated with *P* values less than 0.05 (*), 0.01 (**), 0.001 (***) and 0.0001 (****) indicated on the graphs. Error bars in

590 less mail 0.05 (), 0.01 (), 0.001 () and 0.0001 () 591 histograms denote mean ±SEM.

592 Data availability

- 593 The study did not generate large datasets. All necessary data is included in the figures in this
- 594 study.
- 595

596 Materials

Reagent type	Designation	Source/reference	Identifiers	Additional
			Cat. No.	information
Goat anti-VEGFR2	Extracellular	R&D Systems	AF357	
Goat anti-VEGFR1	Extracellular	R&D Systems	AF321	
Mouse anti-		Sigma		
S100A6				

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Rabbit anti-			
S100A10			
Rabbit anti-	Cytoplasmic	Cell Signaling	2478
VEGFR2-pY1175		Technology	
Rabbit anti-	Cytoplasmic	Abcam	
VEGFR2-pY1214			
Rabbit anti-VP16		Sigma-Aldrich	V4388
Mouse anti-	Clone H68.4	ThermoFisher	13-6800
transferrin			
receptor			
Rabbit anti-		Cell Signaling	4695
ERK1/2		Technology	
Rabbit anti-		Cell Signaling	4376
ERK1/2-		Technology	
pT02/pY204			
Rabbit anti-GST		Sigma-Aldrich	
Mouse anti-		Sigma-Aldrich	
tubulin			
AlexaFluor	Species-specific	ThermoFisher	
secondary	antibody		
antibodies	conjugates		
Endothelial cells	HUVECs	Promocell	C-12203
Endothelial cell	ECGM	Promocell	C-22110
growth medium			
S. cerevisiae yeast	Yeast	DualSystems AG	
strains			
Laboratory		Sigma-Aldrich	
chemicals			
Yeast nitrogen		BD Biosciences	
base			
Yeast dropout		Europa	
media		Bioproducts	
Easyclone cDNA		DualSystems	
synthesis kit		Biotech AG	
β-galactosidase		DualSystems	
assay		Biotech AG	
TRI Reagent		Sigma-Aldrich	T9424
Complete		Sigma-Aldrich	11873580001
protease cocktail			
inhibitor			

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Glutathione-		Sigma-Aldrich	G4510	
agarose				
Amylose resin		New England	E8021	
		Biolabs		
Nickel-agarose		Qiagen	30210	
resin				
SPR sensor chip	CM5	Biacore	CM5	
S100A6 siRNA		GE Healthcare	L-013463	
S100A6 siRNA		GE Healthcare	L-0011766	
High Fidelity	Q5	New England	M0491	
DNA Polymerase		Biolabs		
VEGF-A		Promocell	C-64423	
pGEX-2T-PLC⊚1-		(Larose et al., 1995)		
N-SH2				
Split-ubiquitin		DualSystems		
plasmids and		Biotech AG		
controls				
HexaHis-VEGFR2		This study		
cytoplasmic				
domain				
HexaHis-VEGFR1		This study		
cytoplasmic				
domain				
MBP-VEGFR2		This study		
cytoplasmic		-		
domain constructs				
DNA sequencing		University of		
		Dundee		
		Sequencing		
		Facility		
Biacore 3000	SPR	Biacore AB		Biacore evaluation
				software version
				3.2
DeltaVision	Wide-field	Applied Precision		
	microscope	Inc.		
Image analysis	1	NIH Image J		Freely available
software		0,2		software package
Statistical analysis		GraphPad	GraphPad Prism	Software package

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-	Supplementary Materials: The following supplementary data and tables are available online a
	XXX.
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:	structural biology and modeling; ACR, GKK, MAH, JEL, SP, drafting and revising the article.
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	connets of merest. The autions declare no connet of merest.
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803	Table 1. Membrane Y2H analysis of VEGFR2 and VEGFR1 interaction with S100A6.					
804						
805	Bait	Prey	-LW	-LWH	-LWHA	

- Sector sector for			and the second	A set of the set of the set of the set	ł
VEGFR2-Cub	pPR3-N (vector only)	+++	-	-	
VEGFR2-Cub	Alg5-Nubl	+++	++	+	
VEGFR2-Cub	Fur4-NubG	+++	-	-	
VEGFR2-Cub	PLCg1-(SH2)-NubG	+++	+	-	
VEGFR2-Cub	S100A6-NubG	+++	+	-	
VEGFR1-Cub	pPR3-N (vector only)	+++	-	-	
VEGFR1-Cub	Alg5-Nubl	+++	++	+	
VEGFR1-Cub	Fur4-NubG	+++	-	-	
VEGFR1-Cub	S100A6-NubG	+++	+	-	
pMBV-Fur4	S100A6-NubG	+++	-	-	

Abbreviations for yeast minimal (SD) culture medium additives: L, Leucine; H, Histidine; A,
Adenine; W, Tryptophan. All bait plasmids were co-transformed with test prey plasmids, either
as empty vector carrying only NubG (pR3-N) or carrying ORFs for denoted NubG or NubI hybrid
proteins. Fur4 yeast plasma membrane protein expression was used as another negative control.

825 +++/++/+ denotes scoring of colony growth 5 days post-transformation.



852 Ost1 or Fur4 (negative controls) or NubI fused to Alg5, Ost1 or Fur4 (non-specific control). Co-

- 853 expression of VEGFR2-Cub bait with VEGFR2, PLCγ1-SH2 or S100A6 test proteins fused to
- 854 NubG. LacZ activity assay representative of 3 or more independent experiments.
- 855







Figure S2. Human S100A6 expression in yeast and primary human endothelial cells. (A) Expression
and detection of S100A6-NubG in yeast by immunoblotting using anti-HA antibodies. (D) Expression and
detection of S100A6-NubG in yeast by immunoblotting using anti-S100A6 antibodies. (E) Expression and
detection of human endothelial S100A6 and S100A6-NubG in yeast by immunoblotting using anti-S100A6
antibodies. Arrowheads denote human endothelial S100A6 and S100A6-NubG fusion proteins
respectively.