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Sungkaworn, Titiwat; Jobin, Marie-Lise; Burnecki, Krzysztof; Weron, Aleksander; Lohse, Martin J.; Calebiro, Davide

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1	Single-molecule imaging reveals receptor-G protein interactions at cell surface hot spots
2	Titiwat Sungkaworn ^{1,2} , Marie-Lise Jobin ^{1,2} , Krzysztof Burnecki ³ , Aleksander Weron ³ ,
3	Martin J. Lohse ^{1,2,†} , Davide Calebiro ^{1,2,4,5*}
4	
5	¹ Institute of Pharmacology and Toxicology and ² Bio-Imaging Center/Rudolf Virchow Center,
6	University of Würzburg, Germany
7	³ Faculty of Pure and Applied Mathematics, Hugo Steinhaus Center, Wroclaw University of
8	Science and Technology, Poland
9	⁴ Institute of Metabolism and Systems Research, University of Birmingham, Birmingham, UK
10	⁵ Centre of Membrane Proteins and Receptors (COMPARE), Universities of Birmingham and
11	Nottingham, UK
12	
13	[†] Present address: Max Delbrück Center for Molecular Medicine, Berlin, Germany
14	
15	* address correspondence to:
16	Davide Calebiro, Institute of Pharmacology and Toxicology, Versbacher Str. 9,
17	97078 Würzburg, Germany – E-mail: <u>davide.calebiro@toxi.uni-wuerzburg.de</u>
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19 G protein-coupled receptors (GPCRs) mediate the biological effects of many hormones and neurotransmitters and are major pharmacological targets¹. They 20 21 transmit their signals to the cell interior via interaction with G proteins. However, how receptors and G proteins meet, interact and couple is still ill understood. 22 23 Here, we analyse the concerted motion of GPCRs and G proteins on the plasma 24 membrane and provide a quantitative model that reveals the key factors at the 25 basis of the high spatiotemporal complexity of their interactions. Using two-26 colour, single-molecule imaging we visualize interactions between individual receptors and G proteins at the surface of living cells. Under basal conditions, 27 receptors and G proteins form activity-dependent complexes that last for ~1 28 29 second. Different agonists specifically regulate the kinetics of receptor-G protein interactions, mostly increasing their association rate. We find hot spots on the 30 31 plasma membrane, at least partially defined by the cytoskeleton and clathrin-32 coated pits, where receptors and G proteins are confined and preferentially couple. Imaging with the nanobody Nb37 suggests that signalling preferentially 33 occurs at these hot spots. These findings shed new light on the dynamic 34 interactions governing GPCR signalling. 35

Different scenarios have been developed to explain how receptors and G proteins interact^{2,3}. 36 37 However, key questions concerning the stability of these interactions and the occurrence of 38 GPCR signalling subdomains at the plasma membrane are still open (see also Supplementary 39 Discussion). To address these questions, we visualized individual receptors and G proteins at 40 the surface of living cells with high spatial (≈ 20 nm) and temporal (≈ 30 ms) resolution⁴. As a model, we chose the α_{2A} -adrenergic receptor (α_{2A} -AR), a prototypical family-A GPCR with 41 42 strong G_i coupling². The α_{24} -AR and a pertussis toxin (PTX)-insensitive G α_{i1} construct were specifically labelled with two different organic fluorophores via a SNAP⁵ or CLIP⁶ tag, 43 44 respectively (Fig. 1a); both constructs were fully functional (Extended Data Fig. 1a, b). These 45 constructs were transiently expressed at low physiological densities (0.55±0.10 and 0.51±0.09 molecule µm⁻², respectively) in CHO cells - cultured with PTX to inactivate endogenous G_{i/o} 46 47 proteins - and simultaneously imaged by fast two-colour single-molecule microscopy combined 48 with single-particle tracking⁴ (Fig. 1b and Supplementary Videos 1 and 2). Labelling efficiencies 49 were ~90% (extracellular) and ~80% (intracellular); non-specific labelling was <1% (Extended 50 Data Fig. 1c-e).

51 Individual α_{2A} -AR trajectories were evaluated by mean square displacement (MSD) analysis 52 (corrected for localization error, see Supplementary Methods), which revealed a high 53 heterogeneity and features of anomalous diffusion⁷. Under basal conditions, 11% of the 54 receptors were virtually immobile, while 38% had sub-, 45% normal and 6% super-diffusion (i.e. 55 directional motion) (Extended Data Fig. 2). $G\alpha_i$ had a significantly different diffusion pattern, with 56 a larger immobile fraction (37%) (Extended Data Fig. 2c, d). Stimulation with the full agonist 57 norepinephrine (NE) or brimonidine (UK-14,304) caused a small, significant change in the 58 overall diffusion pattern of $G\alpha_i$ but not of α_{2A} -AR (Extended Data Fig. 2d). Similar results were 59 obtained for a second receptor/G protein pair, i.e. β_2 -adrenergic receptor (β_2 -AR) and G α_s , but 60 no significant differences were observed upon stimulation with the full agonist isoproterenol (Iso; 61 Extended Data Fig. 2d).

We then analysed the trajectories with an algorithm based on hidden Markov models (HMMs)⁸, 62 63 which assumes that particles switch among discrete diffusive states following a stochastic 64 process. We found that both receptors and G proteins frequently switched among at least four 65 distinguishable states (S1 to S4), characterized by distinct diffusion coefficients (D) and ranging from a virtually immobile (S1) to a fast diffusive (S4) state (Fig. 1c-f and Extended Data Fig. 3). 66 67 The results were overall consistent with those of the MSD analysis. We hypothesized that the 68 two slowest states (S1 and S2) were due to trapping in small membrane compartments and, 69 based on the corresponding D and average dwell times, we estimated compartment radiuses of 70 <50 nm and ~270 nm, respectively (see Supplementary Methods). Although with some 71 differences, a similar picture was observed for the integral membrane protein CD86 – used as control⁴ (Fig. 1f, Extended Data Figs. 2d and 3), indicating that such diffusion behaviour is not 72 73 unique to GPCRs/G proteins.

74 Density maps of single-molecule localizations revealed areas that were either preferentially 75 explored or avoided by α_{2A} -ARs and $G\alpha_i$ subunits (Extended Data Fig. 4a and Supplementary 76 Video 3). To better characterize these areas, we generate dynamic maps from the trajectories, 77 reporting local D and potential energy (V) values⁹. This analysis revealed a complex dynamic 78 landscape at the plasma membrane, with high-potential areas, which were rapidly left by α_{2A} -79 ARs/G α_i subunits, and low-potential areas, where they tended to be trapped (Extended Data 80 Fig. 4b, dark areas; see Extended Data Fig. 1f for control). There was a partial but consistent 81 overlap between the potential energy maps of α_{2A} -ARs and $G\alpha_i$ subunits (Extended Data Fig. 82 4c). To quantify this, we measured the relative potential energy values ($V_{i rel}$) of $G\alpha_i$ at the sites 83 of α_{2A} -AR localization and vice versa, which were both significantly lower than for random

localizations or compared to CD86 (Fig. 2a). Importantly, receptor:G protein interactions preferentially occurred at the shared low potential energy areas ("hot spots"), as indicated by negative $V_{L,rel}$ values (Fig. 2b). Similar results were obtained for β_2 -AR and G α_s (Extended Data Fig. 5a, b).

88 To investigate possible factors responsible for this complex diffusion dynamics, we imaged both 89 the cytoskeleton and clathrin-coated pits (CCPs) underneath the plasma membrane. α_{2A} -ARs 90 trajectories tended to avoid microtubules and actin fibres, as suggested by negative 91 colocalization index values (Fig 2c, d, Supplementary Videos 4 and 5), in agreement with the fence-and-picket model¹⁰. Moreover, the majority of trajectories tended to avoid CCPs (Fig. 2e, 92 93 Supplementary Video 6). This coexisted with a minor fraction of α_{2A} -ARs that either transiently 94 stopped at CCPs (Fig. 2f, arrowheads) - consistent with receptor recruitment to pre-existing $CCPs^{11}$ – or were immobile and localized at CCPs. The fraction of CCPs occupied by α_{2A} -ARs 95 96 increased upon agonist stimulation (Fig. 2f, right), Overlays of single particle trajectories with 97 superresolved actin images, obtained by photoactivated localization microscopy (PALM)¹², 98 suggested that the actin mesh underneath the plasma membrane created sub-micrometre 99 compartments in which α_{2A} -ARs were apparently loosely trapped (Fig. 2g and Supplementary 100 Video 7; radius ~100-300 nm, in agreement with estimation based on HMM analysis). 101 Consistently, superimposition of PALM images with potential energy maps showed that the low 102 potential areas were often at least partially delimited by actin fibres (Fig. 2h and Extended Data Fig. 4d). Similar results were obtained for β_2 -AR (Extended Data Fig. 5c-f) and G α_i (Extended 103 104 Data Fig. 6 and Supplementary Video 8).

105 Next, we developed a mathematical analysis to estimate the duration of receptor: G protein 106 interactions based on their trajectories. We reasoned that, on average, for two particles undergoing a true interaction, their observed colocalization time (Δt_{obs}) should correspond to the 107 108 average duration of true interactions (Δt_{true}) plus the average duration of random colocalizations 109 (Δt_{random}) (Fig. 3a and Extended Data Fig. 7). Thus, we deconvolved the observed colocalization 110 times with those of random colocalizations (obtained with CD86 and $G\alpha_i$) to estimate the 111 distribution of the underlying true receptor: G protein interactions (Fig. 3b and Supplementary 112 Methods). The results were subsequently expressed as normalized relaxation curves, showing 113 the fraction of the interactions that are still ongoing at time t from the beginning of each 114 interaction (Fig. 3c). The very fast component in Fig. 3b, seen also with the control CD86, 115 corresponds to non-productive interactions plus random colocalizations, the rate of which (k_{np+rc}) 116 did not differ among the conditions tested (Fig. 3d, left), while we considered the remainder to

117 be productive interactions, i.e. interactions that result in the formation of a "true" complex (see 118 Supplementary Discussion). α_{2A} -ARs and $G\alpha_i$ underwent some productive interactions already 119 under basal conditions (Fig. 3b, c). A major fraction of these interactions terminated following an 120 exponential decay, while a very small fraction (approximately 3.10^{-4}) was stable over the 121 observation time (Fig. 3c). From a fitting of the major component in Fig. 3c and the particle densities we estimated an association rate constant (k_{on}) of ~0.015 μ m² molecule⁻¹ s⁻¹ and a 122 dissociation rate constant (k_{off}) of ~0.8 s⁻¹ for the productive interactions under basal conditions 123 124 (Fig. 3d, middle/right). Treatment with an inverse agonist (yohimbine) or $G\alpha_i$ inactivation (using 125 a PTX-sensitive construct) suppressed the major fraction of transient productive interactions, 126 suggesting that they resulted from constitutive α_{2A} -AR activity and required a functional G α_i 127 subunit; in contrast, the small fraction of stable productive interactions was not affected (Fig. 128 3e). Stimulation with norepinephrine caused a concentration-dependent increase of k_{on} up to ~0.2 μ m² molecule⁻¹ s⁻¹, while k_{off} was only marginally affected (Fig. 3b-d). This translates into 2-129 dimensional equilibrium dissociation constants (K_{d}) of ~50 and 6 molecule μm^{-2} for basal and 130 131 stimulated conditions, respectively. Based on these results, we estimated that, at the tested 132 densities, ~0.5% (basal) or 5% (stimulated) of all α_{2A} -ARs were in complex with $G\alpha_i$ at any given time. Similar results were obtained for β_2 -AR:G α_s interactions, although with 10-fold lower k_{on} 133 values and no long-lived interactions (Fig. 3f, g). A panel of α_{2A} -AR agonists with varying 134 efficacy and affinity revealed statistically significant differences in the estimated k_{on} and, to a 135 136 lesser extent, k_{off} values (Fig. 3d). Overall, there was a positive correlation between k_{on} and 137 efficacy (Fig. 3h). However, there was also a trend towards smaller k_{on} values for higher affinity 138 agonists, both considering full (UK-14,304 vs. norepinephrine) or partial (clonidine vs. 139 oxymetazoline) agonists with comparable efficacies and dissimilar affinities (Fig. 3d, h).

140 By visually inspecting the trajectories, we observed that several α_{2A} -ARs and $G\alpha_i$ subunits 141 slowed down or stopped during apparent interactions to then either remain confined or resume 142 their motion (Fig. 3i, Fig 3j, left and Supplementary Video 9), while the remainder retained their 143 mobility (Fig. 3j, right). A quantitative analysis of the HMM states of α_{2A} -AR and $G\alpha_i$ trajectories 144 showed that, during the time of interaction, higher fractions of receptors/G proteins were in 145 states S1 (virtually immobile) and S2 (slowly diffusing) (Fig. 3k). These mobility changes 146 occurring during the short interaction times and the global changes in $G\alpha_i$ diffusion described in 147 Extended Data Fig. 2d and 3 likely represent distinct phenomena.

148 To further validate our results, we performed deterministic simulations of GPCR signalling using 149 the estimated microscopic k_{on} and k_{off} values for receptor:G protein interactions. The results

150 were in very good agreement with ensemble (FRET) measurements of α_{2A} -AR/G_i 151 association/dissociation (Extended Data Fig. 8). These simulations also suggested that G 152 protein signalling can be fast only if it occurs while the G protein is still bound to the receptor 153 (Supplementary Data). Moreover, we performed particle-based stochastic simulations of 154 receptors and G proteins diffusing and interacting on a 2D surface (Fig 4a). Introducing the 155 experimentally measured potential energy (V) landscapes (as in Fig. 2b) in these simulations 156 doubled the probability of receptor: G protein interactions compared to conditions of simple 157 Brownian motion (Fig. 4a).

To investigate whether hot spots for receptor:G protein interactions also occur in a more physiological context, we studied primary human endothelial cells (HUVEC), where both α_{2A} and β_2 -ARs are endogenously expressed and regulate vascular tone¹³. We found that in these cells both α_{2A} -AR:G α_i and β_2 -AR:G α_s interactions were preferentially occurring at low potential energy areas (Extended Data Fig. 9a, b). We also found that receptors and G proteins slowed down or stopped moving during their interactions (Extended Data Fig. 9c), further strengthening our observations in CHO cells.

Finally, we assessed G_s activation using the conformation-sensitive nanobody Nb37^{14,15}, which recognizes the active (nucleotide-free) state of $G\alpha_s$. In HUVEC transfected with Nb37 fused to a fluorescent protein (EYFP), Nb37 preferentially localized at the sites where β_2 -ARs were concentrated (Fig. 4b).

169 The main findings of our study are summarized in Extended Data Figure 10. First, our results 170 reveal a complex picture, whereby barriers, at least partially constituted by actin fibres. 171 microtubules and CCPs, contribute to the formation of hot spots where receptors and G proteins 172 are both concentrated, and where G protein coupling as well as signalling preferentially occur. This provides a direct visualization of previously postulated GPCR signalling nanodomains^{16,17}. 173 174 Based on our results and simulations, we hypothesize that this complex organization increases 175 both the speed and efficiency of receptor: G protein coupling, while allowing G protein signalling 176 to occur locally.

Second, our data provide direct estimates of the frequency and duration of receptor:G protein interactions in living cells. We find that most receptor:G protein interactions are short-lived (lifetime ~1-2 s). The dependency of these complexes on receptor activation suggests that they are linked to signalling, which is further supported by the observation that G protein activation occurs preferentially at the sites of interaction. In addition, we observe a very small fraction of

182 long-lived complexes (lifetime >> 4s), possibly corresponding to those reported in previous 183 studies^{3,18}. The coexistence of short- and long-lived complexes might reconcile earlier 184 contrasting data. Intriguingly, the estimated duration of the short-lived interactions is much 185 longer than the time required for effector activation, which can happen in ~40 ms². Thus, as 186 suggested by our deterministic simulations, it is conceivable that fast effector activation might 187 occur while the G protein is still bound to the receptor¹⁹.

188 Third, our results reveal that receptor: G protein interactions are regulated by agonists largely at 189 the level of k_{on} . The low k_{on} values measured here also indicate that random collisions only 190 seldom lead to the formation of productive receptor: G protein complexes. The fact that k_{on} is 191 regulated by agonists and the low k_{on} values suggest that receptor: G protein interactions are not 192 limited by diffusion, but rather by the major conformational changes occurring during the formation of receptor: G protein complexes^{14,20-22} (see also Supplementary Discussion). 193 194 Interestingly, different agonists induce substantially different k_{on} values, which correlate at least 195 partially with their efficacies. Together with small differences in the k_{off} values, these findings 196 suggest the possibility of fine-tuning receptor signalling using drugs with tailored effects on the 197 kinetics of receptor: G protein interactions. Finally, our finding of lower k_{on} values for β_2 -AR: G α_s than for α_{2A} -AR:G α_i interactions is consistent with the view that coupling to G_s might require a 198 larger conformational change than coupling to G_i^{23} . 199

200 In summary, our single-molecule results reveal new key factors involved in the regulation of 201 receptor: G protein interactions, which may allow modifying receptor signalling in ways that far 202 exceed simple receptor blockade or activation achieved with currently available drugs, for 203 example by modulating the on/off rates of receptor: G protein interactions or manipulating 204 receptor/G protein mobility and coupling at the hot spots. They further illustrate how GPCR 205 signalling results from dynamic interactions among receptors, G proteins and the complex 206 surrounding membrane environment, which confers flexibility and versatility to this fundamental 207 biological process.

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- 264 **Supplementary Information** is linked to the online version of the paper at 265 <u>www.nature.com/nature</u>.
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275

Author contributions T.S. and M-L.J. performed the experiments. T.S., M-L.J., K.B. and D.C. analysed the data. K.B., A.W. and D.C. developed the mathematical analyses. D.C. wrote the software. D.C. and T.S. wrote the manuscript with contribution of M.J.L. All authors discussed the results. D.C. conceived and supervised the study.

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285

286 Figure Legends

287 Figure 1 | Single-molecule imaging of receptors/G proteins. a, Overall strategy. The α_{2A} -AR 288 was labelled with S549-BG via a SNAP tag fused to its N-terminus. The G_i protein was labelled 289 with SiR-BC via a CLIP tag inserted in an internal loop of the $G\alpha_i$ subunit. **b**, Selected frame 290 from a fast single-molecule image sequence (left) and corresponding trajectories (right). c-e, 291 HMM analysis of diffusive states. Shown is a representative α_{2A} -AR trajectory (c), with its 292 displacement (r) over time (d) and the result of the global HMM analysis (e) revealing 4 states (S1-S4, labelled with different colours). **f**, Model and diffusion coefficients (D: $\mu m^2 s^{-1}$) derived 293 294 from the HMM analysis. Each state is represented by a solid circle; circle area and arrow 295 thickness proportional to occupancy and transition probability, respectively. Differences were 296 statistically significant by two-way ANOVA. *, P<0.05, **, P<0.01, ***, P<0.001, ****, P<0.001, ****, P<0.0001 297 vs. corresponding state of α_{2A} -AR by Tukey's multiple comparison test. Data are mean±s.d. 298 n=22 (85,475), 13 (47,062) and 28 (110,907) cells (trajectories) for α_{2A} -AR, $G\alpha_i$ and CD86, 299 respectively. Images (**b**, **c**) represent 5 independent experiments.

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Figure 2 | Complex diffusion dynamics generates hot spots for receptor:G protein interactions. **a**, α_{2A} -AR localizations over $G\alpha_i$ potential energy map (left) and quantifications of relative potential energy at the localizations ($V_{L,rel}$; right). **b**, α_{2A} -AR:G α_i interactions over merged α_{2A} -AR and $G\alpha_i$ potential energy (V) maps (left) and corresponding quantifications (right).

305 Arrowheads, localizations concentrated at hot spots. **c-e**, α_{2A} -AR trajectories over images of 306 tubulin (c), actin (d) or CCPs (e) (left) and corresponding colocalization analyses (right). f, 307 Trajectories of α_{2A} -ARs (colour-coded according to HMM states) stopping at CCPs (arrowheads; 308 left) and corresponding quantitative analysis (right). **g**, α_{2A} -AR trajectories over actin PALM 309 image (left) and corresponding colocalization analysis (right). Arrowheads, α_{2A} -ARs crossing 310 over actin fibres. **h**, α_{2A} -AR potential energy map over actin PALM image (left) and zoom-in view 311 showing hot spot surrounded by actin fibres (right). α_{2A} -ARs in **g** and **h** were labelled with S647-312 BG. Results (a-e, g) were compared to random localizations. See Supplementary Methods for 313 details. Data are mean±s.d. n, number of cells. #, P<0.05, ##, P<0.01, ###, P<0.001, ####, P<0.0001 vs. random localizations by two-sided paired t-test. *, P<0.05, ****, P<0.0001 vs. α_{2A} -314 315 AR (basal) by two-sided unpaired t-test. Images (h) represent 2 independent experiments.

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317 Figure 3 | Analysis of receptor: G protein interactions. a, Schematic comparison between 318 random colocalizations and true interactions. On average, the observed duration of true 319 interactions (Δt_{obs}) corresponds to their true duration (Δt_{true}) plus that of random colocalizations 320 (Δt_{random}) . The distribution of true durations can then be estimated via deconvolution. **b**, 321 Distributions of the estimated durations of α_{2A} -AR:G α_i interactions under basal and stimulated 322 (NE, 100 μ M) conditions, based on deconvolution. CD86 was used as non-interacting control. c, 323 Relaxation curves calculated from the data in **b**, showing the dissociation kinetics of α_{2A} -AR:G α_i 324 complexes (left, linear; right, semilogarithmic plot). d, Estimated rate constants of non-325 productive interactions plus random colocalizations (k_{nn+rc} ; left), α_{2A} -AR:G α_i association (k_{nn+rc} ; 326 middle) and dissociation (k_{off} ; mean, 95%CI; right) for the indicated ligands. Differences in k_{on} 327 values are statistically significant by one-way ANOVA. ****, P<0.0001 vs. NE 100 µM by Tukey's 328 multiple comparison test. Differences in k_{off} values vs. NE 100 μ M were assessed by two-sided 329 unpaired t-test with Bonferroni correction (****, P<0.0001). **e**, Relaxation curves of α_{2A} -AR:G α_i 330 interactions obtained with an inverse agonist (yohimbine) or using a PTX-sensitive $G\alpha_i$ 331 construct. **f**, Relaxation curves of β_2 -AR:G α_s interactions. **g**, Estimated k_{on} and k_{off} (mean, 332 95%CI) for β_2 -AR:G α_s interactions. ****, P<0.0001 vs. Iso by two-sided unpaired t-test. **h**, 333 Relationship between measured k_{on} and efficacy on α_{2A} -AR activation (mean±s.e.m.). Brackets, 334 affinity values (pK). Efficacy and affinity values are from ref. 24. i, Apparent interaction between 335 α_{2A} -AR and $G\alpha_i$ lasting for 1.2 s. After the interaction, the receptor resumes moving, whereas 336 the G protein remains immobile. **j**, α_{2A} -AR and $G\alpha_i$ trajectories stopping (left) or continuing

337 moving (right) during apparent interactions. k, Distribution of diffusive states (based on HMM 338 analysis) of α_{2A} -AR and $G\alpha_i$ (NE, 100 μ M) during apparent interactions (colocalization duration \geq 339 1.1 s) compared to time outside interactions. Differences are statistically significant by chi-340 square test (****, P<0.0001; n=1,265,634 and 527,058 data points for α_{2A} -AR and $G\alpha_i$, 341 respectively). All ligands were used at saturating concentrations, unless otherwise indicated. 342 See Supplementary Methods for details. Data are mean±s.d., unless otherwise indicated. n, 343 number of cells (d, g). Images (i, j) represent 5 independent experiments. N.D., not 344 determinable.

345

346 Figure 4 | Hot spots for receptor-G protein signalling. a, Stochastic simulations of 347 receptor: G protein interactions. Left, simulated trajectories. Right, fraction of interacting 348 molecules over time. Compared are results with experimentally measured potential energy (V) 349 landscapes vs. simple Brownian motion. **b**, Visualization of local G_s protein activation at the 350 plasma membrane of primary human endothelial cells. Cells were transfected with a fluorescent 351 sensor (Nb37-EYFP) recognizing active, nucleotide-free $G\alpha_s$. Left, β_2 -AR localizations over the 352 obtained spatial map of G_s protein activity. Right, quantification. Data are mean±s.d. n, number 353 of cells. ####, P<0.0001 vs. random localizations by two-sided paired t-test.

354

355 Methods

356 Materials

357 Cell culture reagents, Lipofectamine 2000, Lipofectamine 3000, TetraSpeck fluorescent beads, 358 fluorescein arsenical hairpin binder (FIAsH) and CellMask Green Plasma Membrane Stain were 359 from Thermo Fisher Scientific. The Effectene transfection reagent was from Qiagen. UK-14,304 360 and clonidine were from Tocris Bioscience. All other GPCR ligands, pertussis toxin (PTX), 1,2-361 ethanedithiol (EDT) and guanosine 5'-triphosphate (GTP) were from Sigma-Aldrich. [35 S]GTP γ S 362 was from PerkinElmer. The fluorescent benzyl guanine derivatives SNAP-Surface 549 (S549-363 BG) and SNAP-Surface Alexa Fluor 647 (S647-BG) were from New England Biolabs. Live-cell fluorogenic probes for actin (SiR-Actin) and tubulin (SiR-Tubulin)²⁵ were from Spirochrome. The 364 silicon-rhodamine benzyl cytosine derivative (SiR-BC)²⁶ was kindly provided by Kai Johnsson 365 (Max Planck Institute for Medical Research, Heidelberg, Germany). Ultraclean glass coverslips 366 367 were obtained as previously described⁴.

369 Molecular biology

370 A plasmid coding for the N-terminally SNAP-tagged α_{2A} -adrenergic receptor (SNAP- α_{2A} -AR) 371 was generated by inserting the SNAP tag⁵ before the coding sequence of the murine α_{2A} -372 adrenergic receptor. The generation and functional characterization of the N-terminally SNAP-373 tagged β_2 -adrenergic receptor construct (SNAP- β_2 -AR) have been described in a previous 374 study⁴. A plasmid ($G\alpha_i$ -CLIP) coding for the rat $G\alpha_{i1}$ subunit with the CLIP tag⁶ inserted in the 375 $\alpha A - \alpha B$ loop within the α -helical domain (between positions 91 and 92) was generated by 376 replacing YFP with the CLIP tag in a previously described YFP-tagged $G\alpha_{i1}$ construct²⁷. The 377 construct additionally harboured the C3511 mutation to render it PTX-insensitive²⁸. A plasmid 378 coding for the rat $G\alpha_s$ subunit with the CLIP tag inserted between positions 72 and 85 ($G\alpha_s$ -379 CLIP) was generated by replacing YFP with the CLIP tag in a previously described YFP-tagged $G\alpha_s$ construct²⁹. All tagged receptor and $G\alpha$ subunit constructs behaved like the corresponding 380 381 wild type in functional assays (Extended Data Fig. 1a, b). A construct coding for His-tagged 382 Nb37¹⁴ was kindly provided by Jan Steyaert (VIB, Brussels, Belgium). A plasmid coding for the 383 C-terminally EYFP-tagged Nb37 (Nb37-EYFP) was generated by fusing EYFP to the C-terminus 384 of Nb37. Plasmids coding for CD86 with either one or two SNAP tags fused to its N-terminus have been previously described⁴. Plasmids coding for CD86 with either one or two CLIP tags 385 386 fused to its C-terminus were generated by inserting either one or two copies of the CLIP tag 387 before the stop codon of CD86.

388

389 Cell culture and transfection

390 Chinese hamster ovary K1 (CHO-K1) cells (ATCC) were cultured in phenol red-free Dulbecco's 391 modified Eagle's medium (DMEM)/F-12 supplemented with 5% FCS, 100 U/ml penicillin and 0.1 mg/ml streptomycin at 37 °C, 5% CO₂. For single-molecule experiments, CHO-K1 cells were 392 393 seeded on ultraclean 24-mm glass coverslips in 6-well culture plates at a density of 3.10⁵ 394 cells/well. Cells were treated with 50 ng/ml pertussis toxin (PTX) to inactivate endogenous $G\alpha_{i/o}$ 395 proteins. Transfection was performed 24 h after seeding using Lipofectamine 2000. For each 396 well, 0.8 µg SNAP- α_{2A} -AR or SNAP- β_2 -AR, 0.6 µg G α_i -CLIP or G α_s -CLIP, 0.4 µg G β_1 , 0.2 µg 397 G_{γ_2} , and 6 μ L Lipofectamine 2000 were used. Cells were labelled and imaged by single-398 molecule microscopy 4-6 h after transfection to obtain low physiological expression levels⁴. To 399 label CCPs, cells were transfected 24 h prior to the experiment with GFP-tagged adaptor protein 400 2 (AP2-GFP), kindly provided by Tom Kirchhausen (Harvard Medical School, USA). Human

401 embryonic kidney 293 (HEK293) cells (ATCC) were cultured in DMEM supplemented with 5%
402 FCS, 100 U/ml penicillin and 0.1 mg/ml streptomycin at 37 °C, 5% CO₂. HEK293 cells were
403 transfected with Effectene, following the manufacturer's instructions. Cell lines have not been
404 authenticated. Cells were routinely tested for mycoplasma contamination by PCR using specific
405 primers.

406 For the [³⁵S]GTP γ S binding experiments, HEK293 cells were plated in 10-cm culture dishes and 407 transfected with 3.3 µg α_{2A} -AR, 3.3 µg wild-type or CLIP tagged G α_i , 2.0 µg G β_1 and 1.5 µg G γ_2 408 plasmids.

For FRET experiments, HEK293 cells were seeded on poly-L-lysine–coated 24-mm coverslips and transfected with the indicated constructs. The α_{2A} -AR-Flash/CFP sensor was used to monitor receptor activation³⁰. Co-transfection of α_{2A} -AR-YFP, $G\alpha_i$ -CFP, $G\beta_1$ and $G\gamma_2$ was used to monitor G protein recruitment to the receptor². The $G\beta_1$ -2A-cpV- $G\gamma_2$ -IRES- $G\alpha_{i2}$ -mTq2 sensor³¹ was used to monitor G_i protein activation.

Human umbilical vein endothelial cells (HUVEC) were purchased from Lonza and cultured in complete EGM-2 BulletKit medium (Lonza). HUVEC were plated on ultraclean 24-mm glass coverslips at a density of $3.5 \cdot 10^5$ cells/well and transfected with Lipofectamine 3000 using the same DNA amounts indicated for CHO cells. HUVEC were cultured for a maximum of 10 passages. To visualize local G_s protein activation at the plasma membrane, HUVEC were transfected 24 h prior to the experiment with the Nb37-EYFP construct.

420

421 [35 S]GTP γ S binding assay

422 Membrane preparation and $[^{35}S]GTP\gamma S$ binding assay were performed following a previously 423 described protocol³². Cells were homogenized in lysis buffer (5 mM Tris, 2 mM EDTA, pH 7.4) 424 and then centrifuged at 1,000xg for 10 min. The supernatant was collected and centrifuged at 425 50,000xg for 30 min. The remaining pellet was resuspended in binding buffer (100 mM NaCl, 10 426 mM MgCl₂, 20 mM HEPES, pH 7.4). All procedures were performed at 4 °C. Protein 427 concentrations were determined using the Bradford assay. 10 µg membrane proteins were then 428 incubated with the indicated agonist concentrations and 100 pM [³⁵S]GTP_YS for 15-300 s. Non-429 specific binding was evaluated by adding 10 µM GTP. The samples were then passed through 430 glass fibre filters and radioactivity was determined using a liquid scintillation counter (Beckman 431 LS-1801).

432

433 Live-cell protein labelling

Cells were labelled with a combination of a cell-impermeable SNAP substrate (S549-BG), to label cell-surface receptors, and a highly cell-permeable CLIP substrate (SiR-BC)²⁶, to label intracellular G proteins. Cells were incubated with 4 μ M S549-BG and 8 μ M SiR-BC in complete culture medium for 20 min at 37 °C. Cells were then washed three times using complete culture medium, with 5 min incubation after each wash. This protocol gives labelling efficacy of ~90% and ~80% for extracellular SNAP and intracellular CLIP labelling, respectively (Extended Data Fig. 1c, d).

441 Actin and tubulin labelling were performed using SiR-actin and SiR-tubulin, respectively, 442 following the manufacturer's protocol. Briefly, cells were labelled with 3 μ M SiR-actin or SiR-443 tubulin in the presence of 10 μ M verapamil for 20 min at 37 °C, followed by three washes with 444 complete culture medium.

FIAsH labelling was performed as previously described³⁰. Briefly, cells were incubated with 1 μ M FIAsH and 12.5 μ M EDT in Hank's balanced salt solution (HBSS) for 1 h. The cells were then washed twice with HBSS and incubated with 250 μ M EDT in HBSS for 10 min. The cells were washed a third time with HBSS immediately before the FRET measurement.

449

450 **FRET measurements**

451 Fluorescence resonance energy transfer (FRET) experiments to examine the ensemble kinetics 452 of receptor/G protein signalling in intact cells were done as previously described^{2,33,34}. 453 Measurements were performed on an Axiovert 200 inverted microscope (Zeiss) equipped with 454 an oil immersion 100X objective (Plan-Neofluar 100x, N.A. 1.30), a beamsplitter (DCLP505) and 455 a Polychrome IV monochromator and dual-emission photometric system (Till Photonics). 456 Transfected HEK293 cells were placed in a microscopy chamber filled with imaging buffer (137 457 mM NaCl, 5.4 mM KCl, 2 mM CaCl₂, 1 mM MgCl₂, 10 mM HEPES, pH 7.3). Agonist stimulation 458 was applied using a pressurized rapid superfusion system (ALA-VM8, ALA Scientific 459 Instruments). FRET was monitored as the ratio between YFP (535±15 nm) and CFP (480±20) 460 emission upon CFP excitation at 436±10 nm. The YFP signal was corrected for direct excitation 461 and bleed-through of CFP emission into the YFP channel as previously described².

463 Single-molecule microscopy and PALM

Single-molecule microscopy experiments were performed using total internal reflection 464 465 fluorescence (TIRF) illumination on a custom Nikon Eclipse Ti TIRF microscope equipped with 466 405 nm, 488 nm, 561 nm and 640 nm diode lasers (Coherent), a quadruple band excitation 467 filter, a 100x oil-immersion objective (CFI Apo TIRF 100x, N.A. 1.49), two beam splitters, four 468 separate EMCCD cameras (iXon DU897, Andor), hardware focus stabilization and a 469 temperature control system. Coverslips were mounted in a microscopy chamber filled with 470 imaging buffer. The objective and the sample were maintained at 20 °C by means of a water-471 cooled inset and objective ring connected to a thermostated water bath. Images in the four 472 channels were acquired simultaneously on the four separate EMCCD cameras. Image 473 sequences (400 frames) were taken in crop and frame-transfer mode, resulting in an acquisition 474 speed of 35 frames/s (i.e. one image every 28 ms).

PALM imaging was performed by TIRF microscopy immediately after the acquisition for singleparticle tracking. In this case, cells were additionally transfected 24 h before the experiment with the photoconvertible probe mEOS-LifeAct (a kind gift of Markus Sauer, University of Würzburg, Germany). mEOS was excited at 561 nm, while applying low-intensity 405 nm laser light to induce photoconversion. 10,000 frames were acquired at a speed of 35 frames/s. Superresolved images were then obtained using the rapidSTORM software³⁵.

Images from different channels were registered against each other using a linear piecewise
transformation in *Matlab* based on reference points obtained with multicolour fluorescent beads
(TetraSpeck; 100 nm size).

484

485 Single particle tracking and subsequent analyses

Single particle detection and tracking were performed using the u-track software³⁶ in Matlab 486 487 environment as previously described⁴. The interchannel localization precision after coordinate 488 registration by linear piecewise transformation was ~20 nm. For the analysis of receptor:G 489 protein interactions, a non-related membrane receptor (CD86) with diffusion characteristics 490 comparable to those of the α_{2A} -AR was used as negative control and as reference for random 491 colocalizations⁴. A method based on deconvolution of the observed interaction times with the 492 Lucy-Richardson algorithm^{37,38} was then applied to estimate the underlying duration of 493 receptor: G protein interactions (see Supplementary Methods and Extended Data Fig. 7).

494 To investigate the motion of receptors and G proteins during or immediately before/after an

interaction (Fig. 3i-k and Extended Data Fig. 9c), we considered only apparent interactions with duration \geq 1.1 s, so that random colocalization represented only a small fraction (approximately 15%), based on a comparison between α_{2A} -AR (NE 100 μ M) and CD86 (used as negative control).

499 Detailed information about the computational analyses can be found in Supplementary500 Methods.

501

502 Hidden Markov model (HMM) analysis

A software based on a variational Bayesian treatment of HMMs (vbSPT)⁸ was used to identify discrete diffusive states in the single molecule trajectories and analyse their characteristics. The number of iterations and bootstrapping were set to 25 and 100, respectively. Diffusion coefficients and dwell times derived from the analysis were used to estimate the size of the corresponding nanocompartments on the plasma membrane (see Supplementary Methods).

508

509 Spatial mapping of receptor/G protein dynamics

510 Spatial maps of diffusivity (D) and potential energy (V) were obtained using the Inference MAP 511 software⁹, based on Bayesian inference, considering a physical model of diffusion in a potential 512 field. Only well-adhering cells with a flat plasma membrane were chosen to avoid artefacts due 513 to uneven distance from the coverslip. The flatness of the plasma membrane was verified by 514 staining with a fluorescent phospholipid (CellMask Green). The analysed areas were partitioned 515 in small regions of variable size by Voronoi tessellation⁹. The number of regions was optimized 516 to avoid areas with low number of localizations. The obtained potential energy maps were 517 subsequently used to perform particle-based stochastic simulations of receptor:G protein 518 interactions (see Supplementary Methods).

519

520 Statistics and reproducibility

521 Statistical analyses were performed using the Prism 6 software (GraphPad Software). 522 Differences between two groups were assessed by two-sided Student's t-test. Differences 523 among three or more groups were assessed by one-way or two-way analysis of variance 524 (ANOVA), as appropriate, followed by Tukey's multiple comparison test (with the exception of 525 the data in Figure 3d, right, which were compared by two-sided unpaired t-test with Bonferroni

Data availability The data that support the findings of this study are available from the corresponding author
530 The data that support the findings of this study are available from the corresponding author
 upon reasonable request.
533 Code availability
534 Matlab scripts are available from the corresponding author upon reasonable request.
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568 **Extended Data Figure legends**

569 Extended Data Figure 1 | Control experiments. a-c, Functional characterization of the SNAP/CLIP tagged receptor/G protein constructs. a, FRET measurements of G_i protein 570 571 activation to test the SNAP-tagged α_{2A} -AR construct. HEK-293 cells were co-transfected with a 572 FRET sensor for G_i protein activation (G β_1 -2A-cpV-G γ_2 -IRES-G α_{i2} -mTq2) and either wild-type or SNAP-tagged α_{2A} -AR (n=9, 11 cells). Concentration response relationships were obtained from 573 574 FRET measurements in which the cells were stimulated with increasing concentrations of NE. 575 The SNAP-tagged β_2 -AR construct was tested in a previous study⁴. **b**, GTP_yS binding assay to 576 test the CLIP-tagged $G\alpha_i$ and $G\alpha_s$ constructs. The corresponding wild-type constructs were used as control. Shown are time courses of GTP_γS binding in the presence or absence of 577 578 agonist (left, clonidine, 10 μ M; right, isoproterenol, 10 μ M) (n=3 biological replicates per 579 condition). c, Efficiency of extracellular labelling with the cell impermeable SNAP substrate 580 S549-BG in cells transfected with CD86 carrying two SNAP tags at its N-terminus (n=28, 13, 17, 581 14 and 11 cells). d, Efficiency of intracellular labelling with the cell permeable CLIP substrate 582 SiR-BC in cells transfected with CD86 carrying two CLIP tags at its C-terminus (n=18, 26, 31, 27 583 cells). Labelling efficiencies in **c** and **d** were determined by fitting single-particle intensity data 584 with a mixed Gaussian model⁴. The following concentrations were chosen for subsequent 585 experiments: 4 μ M S549-BG (labelling efficiency 91.1±2.9%) and 8 μ M SiR-BC (labelling

586 efficiency 82.6±2.1%). e, Specificity of labelling. Shown are TIRF images of CHO cells 587 transfected with different combinations of SNAP- α_{2A} -AR and G α_i -CLIP, followed by labelling with 588 S549-BG or SiR-BC (except for the positive control cotransfected with SNAP- α_{2A} -AR and $G\alpha_{i}$ -589 CLIP, which was labelled with both fluorescent substrates). Unspecific labelling in either mock or 590 cross-transfected cells was responsible for 2-3 spots per cell on average against ~300-400 in 591 the positive controls. This very low number of unspecific immobile localizations (<1%) does not 592 significantly interfere with the analyses of this study. f, Representative potential energy (V) maps 593 calculated in mock-transfected cells. Shown are results obtained after adding simulated 594 trajectories with Brownian motion to reproduce a condition of diffusing particles over immobile 595 unspecific localizations. The presence of immobile unspecific localizations was not sufficient to 596 generate local low potential energy areas (see **Extended Data Fig. 4b** for comparison). Data 597 are mean±s.e.m. Images (e, f) represent 3 independent experiments.

598

599 Extended Data Figure 2 | MSD analysis of receptor and G protein trajectories. a, Scatter 600 plot of diffusion coefficient (D) and anomalous diffusion exponent (α) values estimated for 601 simulated trajectories with Brownian motion and characteristics similar to those of α_{2A} -ARs. The 602 results were used to set the cut-offs for classifying the trajectories into four groups according to their motion: immobile (D<0.01 μ m² s⁻¹), sub-diffusion (α <0.75), normal diffusion (0.75≤ α ≤1.25) 603 604 and super-diffusion (α >1.25). **b**, Scatter plot as in **a** for α_{2A} -AR trajectories. **c**, Representative 605 α_{2A} -AR and $G\alpha_i$ trajectories classified in the four groups. **d**. Relative frequency distributions of 606 the trajectories in the four groups (left) and corresponding D values (right) for receptors and G 607 proteins under basal and stimulated conditions. The control CD86 was expressed together with 608 wild-type α_{2A} -AR and stimulated with NE to verify if the effects observed upon agonist 609 stimulation were specific for $G\alpha_i$. Differences in **d** are statistically significant by two-way ANOVA. 610 #, P<0.05, ##, P<0.01 and ####, P<0.0001 vs. the corresponding basal condition and **. P<0.01, ***, P<0.001, ****, P<0.0001, vs. α_{2A} -AR basal (top) or β_2 -AR basal (bottom) by Tukey's 611 612 multiple comparison test. Data are mean±s.e.m. n=30 (9,273), 17 (6,623), 37 (8,309), 30 613 (4,699), 18 (2,182), 36 (5,240), 28 (11,267), 27 (12,697), 29 (10,760), 47 (16,461), 29 (41,079) 614 and 47 (7,585) cells (trajectories) for α_{2A} -AR basal, α_{2A} -AR NE, α_{2A} -AR UK-14,304, G α_i basal, 615 $G\alpha_i$ NE, $G\alpha_i$ UK-14,304, CD86 basal, CD86 NE, β_2 -AR basal, β_2 -AR lso, $G\alpha_s$ basal, $G\alpha_s$ lso, 616 respectively. Images (c) represent 5 independent experiments.

618 Extended Data Figure 3 | Complete results of the hidden Markov model (HMM) analysis. 619 Differences are statistically significant by two-way ANOVA. #, P<0.05, ##, P<0.01, ###, P<0.001 and ####, P<0.0001 vs. the corresponding basal condition, and *, P<0.05, **,P<0.01, ***, 620 P<0.001 and ****: P<0.0001 vs. α_{2A} -AR basal (for $G\alpha_i$ and CD86) or β_2 -AR basal (for $G\alpha_s$) by 621 622 Tukey's multiple comparison test. Data are mean±s.d. n=22 (85.475), 11 (44.797), 31 (153.072), 623 13 (47,062), 9 (15,161), 30 (88,397), 28 (110,907), 30 (147,222), 27 (142,243), 44 (229,815), 28 624 (84,668) and 44 (171,623) cells (trajectories) for α_{2A} -AR basal, α_{2A} -AR NE, α_{2A} -AR UK-14,304, 625 $G\alpha_i$ basal, $G\alpha_i$ NE, $G\alpha_i$ UK-14,304, CD86 basal, CD86 NE, β_2 -AR basal, β_2 -AR lso, $G\alpha_s$ basal, 626 $G\alpha_s$ Iso, respectively. N.D., not determinable.

627

Extended Data Figure 4 | Complex diffusion dynamics of α_{2A} -AR and $G\alpha_i$. **a**, density maps of α_{2A} -AR and $G\alpha_i$ localizations (selected trajectories overlaid in different colours; arrowheads, areas of high density). **b**, Potential energy (*V*) maps for α_{2A} -AR and $G\alpha_i$, calculated for the same membrane region. **c**, Merge of potential energy maps in **b** (top) and line-profile plot along the dashed line (bottom). Arrowheads, hot spots where *V* is low for both α_{2A} -AR and $G\alpha_i$. **d**, Additional examples of α_{2A} -AR potential energy (*V*) maps over actin PALM images and corresponding zoom-in views. Images represent 3 (**a-c**) and 2 (**d**) independent experiments.

635

636 Extended Data Figure 5 | Complex diffusion dynamics of β_2 -AR and $G\alpha_s$. a, β_{2A} -AR 637 localizations over $G\alpha_s$ potential energy (V) map and vice versa (top) and quantifications of 638 relative potential energy at the localizations ($V_{\rm l}$ rel; bottom). A negative value indicates relatively 639 lower potential energy at the localizations. **b**, β_2 -AR:G α_s interactions over merged β_2 -AR and 640 $G\alpha_s$ potential energy map (top) and corresponding quantifications (bottom). Arrowheads, 641 localizations concentrated at hot spots. **c-e**, β_2 -AR trajectories over images of tubulin (**c**), actin 642 (d) or CCPs (e) (top) and corresponding colocalization analyses (bottom). Negative 643 colocalization index (1) values indicate preferential avoidance of the imaged structures by the 644 receptors. f, Trajectory of β_2 -AR stopping at CCP (arrowhead; top) and corresponding 645 quantitative analysis (bottom). The trajectory is colour-coded according to the HMM states. 646 Results in **a-e** were compared to those obtained with random localizations. See Supplementary 647 Methods for details. Data are mean±s.d. n, number of cells. #, P<0.05, ##, P<0.01, ###, 648 P<0.001, ####, P<0.0001 vs. random localizations by two-sided paired t-test. *, P<0.05, **, 649 P<0.01, ***, P<0.001, ****, P<0.0001 vs. β_2 -AR (basal) by two-sided unpaired t-test.

650

Extended Data Figure 6 | Complex diffusion dynamics of $G\alpha_i$. **a**, $G\alpha_i$ trajectories over actin PALM image. Arrowheads, $G\alpha_i$ subunits crossing over actin fibres. **b**, $G\alpha_i$ trajectories over image of CCPs. **c**, $G\alpha_i$ potential energy (*V*) map over PALM image of actin fibres (left) and corresponding zoom-in view (right). Images represent 2 (**a**, **c**) and 3 (**b**) independent experiments.

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657 Extended Data Figure 7 | Validation of the analysis of receptor: G protein interactions 658 using simulated data. a-d, Test of the Lucy-Richardson (LR) deconvolution algorithm on data 659 obtained with numerical simulations. A numerical simulation of particles interacting and dissociating following an exponential law (n=20,000 particles; $k_{off}=0.7 \text{ s}^{-1}$) was performed. In 660 addition, we considered particles undergoing random colocalizations (also terminating following 661 662 an exponential law; n=20,000 particles; $k=1.75 \text{ s}^{-1}$). a, Underlying distribution of the interaction times for the true interactions. b, Distribution of the colocalization times for the random 663 664 colocalizations. c, Convolution of the distribution in a with that in b, corresponding to the 665 observed colocalization times. d, Deconvolution of the distribution in c with that in b. Note that 666 the algorithm was capable of correctly retrieving the distribution of the true interaction times. e. 667 f, Simulated two-channel image sequences of particles undergoing transient interactions. A 668 defined fraction of particles in the first channel was simulated to undergo interactions with 669 particles in the second channel. The synthetic image sequences were then analysed using 670 automated particle detection and tracking as for the experimental ones. e. Representative frame 671 of a simulated two-colour image sequence. f, Trajectories obtained by automated single-particle 672 detection and tracking. g, Relaxation curve obtained from simulations of non-interacting 673 particles. h, Relaxation curve obtained from simulations of interacting and non-dissociating 674 particles. i, Result of the LR deconvolution analysis of the data in h with those in g. Data were 675 fitted with an exponential decay, used to estimate the rate of premature termination (see 676 Supplementary Methods). j, Results of the LR deconvolution analysis on simulated image 677 sequences comparing the input dissociation rate constants (k_{off}) and the ones estimated by the 678 analysis. The results were corrected for premature termination of the interactions as described 679 in Supplementary Methods. All simulations were repeated 3 times with similar results.

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681 Extended Data Figure 8 | Simulations with deterministic model of GPCR signalling. a,

682 Schematic representation of the model. r, inactive receptor. R, active receptor. L, ligand 683 (agonist). b, Model reactions and kinetics parameters used in the simulations. c, d, FRET 684 measurements of the ensemble kinetics of α_{2A} -AR activation/deactivation in response to 685 transient agonist stimulation. A sensor consisting in the α_{2A} -AR carrying a FIAsH tag in the third 686 intracellular loop and CFP at the C-terminus was used. Shown are a representative FRET 687 measurement (c) and the normalized average of the FRET data (d; n=4 cells). e, Fitting of 688 model parameters using the measured FRET data for receptor activation/deactivation. f, 689 Concentration-response relationships for ligand binding, receptor activation and G protein 690 activation generated with the model. g, h, Simulations of GPCR signalling in response to 691 transient agonist stimulation, applying the estimated k_{on} and k_{off} for receptor:G protein 692 interactions to the model. Simulations were performed both considering low (g) and high (h) 693 receptor/G protein expression levels. i, j, FRET measurements of the ensemble kinetics of α_{2A^-} 694 AR:G α_i association/dissociation in response to transient agonist stimulation. A sensor consisting 695 in the α_{2A} -AR carrying YFP at the C-terminus and the $G\alpha_{i1}$ subunit carrying CFP in the αA - αB 696 loop within the α -helical domain was used. Shown are a representative FRET measurement (i) 697 and the normalized average (n=16 cells) of the FRET data (i). Association and dissociation time 698 constants (mean, 95%CI) were 44.4 (38.3-52.9) ms and 1.22 (1.16-1.29) s. k, Comparison 699 between the FRET data in j and the result of simulation with the mathematical model. Data are 700 mean±s.e.m.

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702 Extended Data Figure 9 | Hot spots for receptor: G protein interaction in primary human 703 endothelial cells. a, α_{2A} -AR:G α_i interactions over merged α_{2A} -AR and G α_i potential energy (V) 704 map (left) and corresponding quantifications (right). Arrowheads, α_{2A} -AR:G α_i interactions 705 concentrated at hot spots. **b**, Same analysis as in **a** for β_2 -AR and G α_s . ####, P<0.0001 vs. 706 random localizations by two-sided paired t-test. c. Distribution of diffusion states (based on 707 HMM analysis) of receptor/G protein trajectories during apparent interactions. Differences are 708 statistically significant by chi-square test (****, P<0.0001; n=2,488,438 and 1,382,193 data 709 points for α_{2A} -AR and G α_i with NE stimulation; n=1,992,190 and 874,317 data points for α_{2A} -AR 710 and $G\alpha_i$ with UK-14,304 stimulation; n=5,073,163 and 3,959,938 data points for β_2 -AR and $G\alpha_s$ 711 with Iso stimulation, respectively). Data (**a**, **b**) are mean±s.d. n, number of cells (**a**, **b**).

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713 **Extended Data Figure 10 | Schematic summary. a,** The complex organization of the plasma

714 membrane, including barriers provided by actin fibres, microtubules and CCPs, generates hot 715 spots for receptor: G protein interaction and signalling. b, Receptors and G proteins undergo 716 random collisions (preferentially within these hot spots), which, via very short-lived encounter 717 complexes, only seldom lead to the formation of productive receptor: G protein (R:G) complexes 718 (low k_{on}). Most of these complexes dissociate with a lifetime of ~1-2 s, while very few are long-719 lived. Agonists mainly act by increasing the k_{on} for receptor: G protein interactions in a ligand-720 specific manner. These data suggest that most receptor:G protein complexes are transient and 721 that receptor: G protein interactions are not diffusion limited but rather controlled by the large 722 conformational rearrangements occurring during the formation of productive receptor:G protein 723 complexes. See also Supplementary Discussion.







