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**Ras-Raf signalosome structures** 

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# A structural model of a Ras-Raf signalosome

Venkatesh P. Mysore<sup>1,10</sup>, Zhi-Wei Zhou<sup>2,10</sup>, Chiara Ambrogio<sup>3,4,10</sup>, Lianbo Li<sup>2</sup>, Jonas N. Kapp<sup>5</sup>, Chunya Lu<sup>6,7</sup>, Qi Wang<sup>1</sup>, Maxwell R. Tucker<sup>1</sup>, Jeffrey J. Okoro<sup>3</sup>, Gabriela Nagy-Davidescu<sup>5</sup>, Xiaochen Bai<sup>6</sup>, Andreas Plückthun<sup>5</sup>, Pasi A. Jänne<sup>3,8</sup>, Kenneth D. Westover<sup>2</sup>, Yibing Shan<sup>1</sup> and David E. Shaw<sup>1,9</sup>

The protein K-Ras functions as a molecular switch in signaling pathways regulating cell growth. In the human mitogen-activated protein kinase (MAPK) pathway, which is implicated in many cancers, multiple K-Ras proteins are thought to assemble at the cell membrane with Ras effector proteins from the Raf family. Here we propose an atomistic structural model for such an assembly. Our starting point was an asymmetric guanosine triphosphate-mediated K-Ras dimer model, which we generated using unbiased molecular dynamics simulations and verified with mutagenesis experiments. Adding further K-Ras monomers in a head-to-tail fashion led to a compact helical assembly, a model we validated using electron microscopy and cell-based experiments. This assembly stabilizes K-Ras in its active state and presents composite interfaces to facilitate Raf binding. Guided by existing experimental data, we then positioned C-Raf, the downstream kinase MEK1 and accessory proteins (Galectin-3 and 14-3-3 $\sigma$ ) on and around the helical assembly. The resulting Ras-Raf signalosome model offers an explanation for a large body of data on MAPK signaling.

he protein K-Ras is a crucial molecular switch in signaling pathways that regulate cell proliferation, differentiation and survival<sup>1</sup>. Gain-of-function mutations in three closely related members of the Ras family, K-Ras, H-Ras and N-Ras, are found in approximately 25% of all human cancers, with mutations in K-Ras being particularly common<sup>2</sup>. In many cases, oncogenic mutations in Ras are associated with hyperactivation of the MAPK signaling pathway<sup>3</sup>. Dysregulation of the MAPK pathway has also been implicated in a host of hyperproliferative developmental disorders termed RASopathies<sup>4</sup>. Inhibiting oncogenic or aberrant Ras is thus of great clinical interest<sup>5,6</sup>. Our structural understanding of how K-Ras activates downstream MAPK signaling, however, has been incomplete.

Ras is active when guanosine triphosphate (GTP) bound and inactive when guanosine diphosphate (GDP) bound; conversions between these states are catalyzed by guanine nucleotide-exchange factors (GEFs) and GTPase-activating proteins (GAPs), respectively<sup>7</sup>. Active and inactive Ras conformations differ primarily at the 'switch' regions involved in effector binding<sup>8,9</sup>. Despite some early evidence that Ras can form oligomers<sup>10</sup>, activated Ras has long been thought to act in MAPK signaling as a monomeric membrane anchor for C-Raf or other effectors from the Raf kinase family (with Ras itself being anchored to the plasma membrane by its lipidated C-terminal tail). A growing body of data now indicate that Ras proteins dimerize<sup>11–15</sup>, that their dimerization may depend on GTP binding<sup>16</sup> and that Ras dimerization may be critical for Raf activation<sup>17–20</sup>.

A notable fraction of membrane-bound, GTP-bound Ras proteins have been found to organize into relatively immobile 'nanoclusters', each with about eight members and a radius of roughly 100 Å (refs. <sup>21,22</sup>). Ras dimerization appears to be crucial to nanocluster formation<sup>19</sup>, and it is possible that nanoclusters comprise networks of loosely interacting, lower-order Ras structures (such as monomers and dimers); structurally well-defined, higher-order oligomers; or a mixture of both. It is known that Galectin-3 (Gal-3) is crucial for K-Ras nanoclustering and signaling<sup>23</sup>, as is Galectin-1 (Gal-1) for H-Ras<sup>24</sup>. Raf kinases colocalize with nanoclustered Ras proteins; it has been shown that such colocalized effectors generate most of the downstream MAPK signal<sup>22,25</sup>, and that Ras activity can be suppressed by proteins that disrupt this clustering<sup>19,26</sup>. Structural information about nanoclusters is sparse, however, because they are difficult to reconstitute in vitro and the resolution of cellular imaging is limited.

In the work reported here, we used unbiased molecular dynamics simulations to inform the construction of an atomistic structural model of a K-Ras nanocluster. We term our proposed structure a Ras-Raf signalosome because it is a higher-order hetero-oligomer (containing multiple monomers of K-Ras, C-Raf, Gal-3 and other proteins) with well-defined structural features and is reminiscent of 'signalosomes' that underlie the activities of other signaling systems<sup>27</sup>. We first generated a GTP-mediated asymmetric (GMA) dimer model by directly simulating the association of two K-Ras monomers; subsequent mutagenesis experiments strongly supported this model. The asymmetric dimer model was then extended to a higher-order oligomer model by adding monomers in a head-to-tail fashion, resulting in a compact helical K-Ras assembly. This assembly promotes the stability and accessibility of active K-Ras and creates composite interfaces that facilitate Raf binding. Guided by existing experimental data and further simulations, we then positioned C-Raf dimers, the downstream kinase MEK1, and

<sup>&</sup>lt;sup>1</sup>D. E. Shaw Research, New York, NY, USA. <sup>2</sup>Departments of Biochemistry and Radiation Oncology, University of Texas Southwestern Medical Center, Dallas, TX, USA. <sup>3</sup>Department of Medical Oncology, Dana-Farber Cancer Institute, Boston, MA, USA. <sup>4</sup>Department of Molecular Biotechnology and Health Sciences, Molecular Biotechnology Center, University of Turin, Turin, Italy. <sup>5</sup>Department of Biochemistry, University of Zürich, Zürich, Switzerland. <sup>6</sup>Department of Biophysics, University of Texas Southwestern Medical Center, Dallas, TX, USA. <sup>7</sup>Department of Respiratory Medicine, The First Affiliated Hospital of Zhengzhou University, Zhengzhou, China. <sup>8</sup>Belfer Center for Applied Cancer Science, Dana-Farber Cancer Institute, Boston, MA, USA. <sup>9</sup>Department of Biochemistry and Molecular Biophysics, Columbia University, New York, NY, USA. <sup>10</sup>These authors contributed equally: Venkatesh P. Mysore, Zhi-Wei Zhou, Chiara Ambrogio. <sup>58</sup>e-mail: Yibing.Shan@DEShawResearch.com; David.Shaw@DEShawResearch.com



**Fig. 1 Outline of the structural model of a Ras-Raf signalosome. a**, The GMA K-Ras dimer model. Residues contacting the membrane (mesh) are shown in purple; the GTP at the interface is in red. **b**, The (eight-protomer) K-Ras helical assembly with illustrative cartoons. **c**, The K-Ras helical assembly (pink), decorated with C-Raf RBD and CRD domains (green) and Gal-3 proteins (yellow). Two scenarios—one with Gal-3 at the base tier (left) and one without (right)—are shown, with the same orientation. **d**, The signalosome model on the membrane. The eight-protomer model contains four catalytic units at the periphery. **e**, Diagram of the signalosome model with a helical wheel representing the Ras assembly at the center. The Gal-3 molecules associated with the base-tier K-Ras (1-4) are outlined by dashed lines to indicate that Gal-3 is not structurally required at the base tier. To illustrate how the signalosome could grow beyond eight protomers, 16 protomers are shown. **f**, One catalytic unit centered on a Raf kinase dimer. ATP and phosphorylated Raf residues (pSer338, pTyr341 and pSer621) are in red. **g**, Previously resolved structures used as building blocks in the modeling: the K-Ras monomer<sup>85</sup> (PDB 4DSN), H-Ras-Raf RBD complex<sup>47</sup> (PDB 4GON), C-Raf CRD<sup>86</sup> (PDB 1FAR), Gal-3 (ref. <sup>87</sup>) (PDB 3ZSM), C-Raf kinase domain dimer<sup>64</sup> (PDB 3OMV), B-Raf-MEK1 complex<sup>65</sup> (PDB 4MNE) and 14-3-3σ dimer bound to C-Raf phosphopeptide<sup>63</sup> (PDB 4IEA).

the accessory proteins Gal-3 and 14-3-3 $\sigma$  around the K-Ras core. Our Ras–Raf signalosome model, which synthesizes a large body of data on the MAPK pathway, may extend to other Ras and Raf isoforms and provides a structural framework to inform further investigations of MAPK signaling.

#### Results

We first summarize key structural features of our Ras–Raf signalosome model for convenience before detailing its construction and resulting findings.

The signalosome model centers on a K-Ras helical assembly. In brief, we obtained our Ras–Raf signalosome model by proposing a GMA K-Ras dimer model (Fig. 1a), extending it to a K-Ras helical assembly (Fig. 1b) and adding further components. The final model is anchored to the plasma membrane, and the Ras-binding domains (RBDs) and cysteine-rich domains (CRDs) of C-Raf interact directly with the K-Ras helical assembly (Fig. 1c), essentially covering the outside of the assembly. The four K-Ras molecules at the base of the helical assembly contact the membrane. We considered two possible scenarios with respect to the farnesylated Cys185 (fCys185) residues<sup>28</sup> of the K-Ras proteins at the base tier. In one, the farnesyl groups at the base tier are capped by membrane-bound Gal-3 molecules (we will discuss in detail below the motivation for this aspect of the model) and in another, which is more consistent with the current understanding of the role fCys185 in membrane-anchoring K-Ras, these fCys185 residues are inserted into the membrane. Our model was stable in both scenarios in our simulations. In both scenarios, Gal-3 molecules cap the farnesyl groups of all the K-Ras molecules beyond the base tier, in line with experimental data<sup>23</sup>.

Unstructured linkers connect the CRDs of C-Raf proteins to their kinase domains, which dimerize at the periphery of the structure (Fig. 1d,e), in complex with MEK1 kinases and 14-3-3 dimers (Fig. 1f). The signalosome model, which is structurally open-ended, can accommodate a variable number of K-Ras and Raf molecules. We anticipate that the signalosome is regulated by GAP and GEF at the cell membrane and thus cannot grow beyond the reach of membrane-anchored GEF.

In this report, we mostly focus on an eight-protomer signalosome as an example system. The signalosome (Supplementary Video 1, Fig. 1d and Extended Data Fig. 1a; atomic coordinates in Supplementary Dataset 1) produces a millimolar local concentration of C-Raf (Supplementary Note and Analyses), which would be sufficient to ensure the dimerization-dependent activation of C-Raf<sup>29</sup>. The model incorporates seven previously resolved single- or double-domain structures of the constituent proteins (Fig. 1g). The following sections describe in greater detail our construction of this signalosome model, which was informed by existing experimental data, new experiments conducted as part of the current study, and previous experience with molecular dynamics simulations of protein–protein and protein–small molecule association<sup>30–32</sup>.

A GMA dimer of active K-Ras. We first performed 20 simulations (680 µs total simulation time) of two GTP-bound K-Ras proteins (Protein Data Bank (PDB) ID 4DSN) in aqueous solvent (Extended Data Fig. 2a, left). In one of the simulations, two K-Ras proteins formed a symmetric dimer that is almost identical (Extended Data Fig. 2b) to the crystal structure of a widely discussed symmetric Ras dimer that uses the  $\alpha 4-\alpha 5$  helices as the dimer interface<sup>13,20</sup>, illustrating the ability of the simulations to recapitulate experimentally observed interactions for this system. (We will discuss this and other existing Ras dimer models in relation to our signalosome model later in this article.) In another simulation, two K-Ras proteins formed stable interactions mediated in part by a bound GTP (Supplementary Video 2). This model is new and compelling because it provides a direct explanation for the GTP dependence of K-Ras dimerization<sup>16</sup>. Hereafter we will refer to this model as the GMA dimer model.

Because K-Ras dimerization occurs at the membrane, we then performed 23 simulations ( $363 \mu s$  total simulation time) of two GTP-bound K-Ras proteins anchored to the membrane by their fCys185 residues (Extended Data Fig. 2a, right). In one of these membrane simulations (Fig. 2a and Supplementary Video 3), the K-Ras proteins also formed the GMA dimer; the structure is virtually identical to that obtained from the solvent simulations (Fig. 2b, upper panel). The GMA dimer (atomic coordinates in Supplementary Dataset 2) remained stable (Fig. 2b).

At the GMA dimer interface, a key interaction occurs between the GTP  $\gamma$ -phosphate of one K-Ras protein and Arg135 or Lys128 of the other (Fig. 2b,c and Extended Data Fig. 2c). Hereafter, we will refer to the former K-Ras protein as the GTP donor (or, more briefly, the donor) and the latter as the GTP acceptor (or simply the acceptor). Unlike the 'arginine finger' interaction between Ras and GAP, whereby an arginine mediates electron transfer from the  $\gamma$ -phosphate to the  $\beta$ -phosphate<sup>33</sup> and catalyzes GTP hydrolysis<sup>34</sup>, the GMA dimer model has Arg135 or Lys128 interacting only with the  $\gamma$ -phosphate (Fig. 2c), hindering electron transfer and thus GTP hydrolysis.

The GMA dimer interface has a buried surface area of roughly 1,750 Å<sup>2</sup> and pronounced electrostatic complementarity (Extended Data Fig. 2j). The donor interface involves parts of the switch I region and the  $\beta$ 4– $\alpha$ 3,  $\beta$ 5– $\alpha$ 4 and  $\beta$ 6– $\alpha$ 5 loops. The acceptor interface primarily involves the  $\alpha$ 4 and  $\alpha$ 5 helices and the  $\beta$ 6 strand (Fig. 2d) and largely overlaps with the interface for NS1 (Extended Data Fig. 2i), a synthetic protein that disrupts Ras dimerization<sup>19</sup>. The GMA dimer is consistent with existing nuclear magnetic resonance (NMR) spectroscopy data: residues that exhibited broadening of <sup>15</sup>N-HSQC spectra on sample dilution<sup>16</sup> are predominantly located at the dimer interface (Fig. 2e and Supplementary Analyses). The interface is mostly polar (Extended Data Fig. 2e), suggesting modest dimerization affinity. Several salt bridges (for example, Asp154–Lys147, Arg161–Asp30/Glu31 and Glu62–Lys128/Arg135) are present at the interface (Fig. 3a,b).

**The GMA dimer stabilizes the active state of K-Ras.** The GMA dimer may promote RBD binding by stabilizing the active Ras conformation and favoring membrane orientations that accommodate the RBD. It has been shown that GTP binding favors the active conformation<sup>35</sup> but does not fully stabilize it<sup>9,36-39</sup>. In our simulations, although the switch I region visited the active conformation with greater frequency in a GTP-bound K-Ras monomer than in a GDP-bound one (Extended Data Fig. 3a), the inactive conformation

was prevalent in both (Extended Data Fig. 3b,c). In contrast, in simulations of the GMA dimer, the switch I region of the donor (but not that of the acceptor) was largely stabilized in active-like conformations (Fig. 2f and Extended Data Fig. 3a).

Our GMA dimer model also constrains the membrane orientation of K-Ras in a way that is consistent with previous NMR and simulation data<sup>40–42</sup>. In the GMA dimer formed on the membrane (Fig. 1a), the donor is in the  $\alpha$  orientation<sup>40</sup> and the acceptor is in a similar orientation (Fig. 2g). Both the donor and acceptor orientations are compatible with RBD binding (Extended Data Fig. 2g). This contrasts with monomeric K-Ras, which mostly adopted the  $\beta$  membrane orientation that occludes RBD binding (Fig. 2g and Extended Data Fig. 2f).

The common oncogenic K-Ras mutations are at or adjacent to the GMA dimer interface (Fig. 3c). We simulated GMA dimers with G12V, G12D, G12C, G13C, G13S, Q61L and Q61R and found that the dimer was not disrupted by these mutations (Fig. 3d and Extended Data Fig. 2j), likely because the GMA dimer interface can sterically accommodate the mutations. Although not quantified by the simulations, these mutations likely exert subtle differential effects on GMA dimerization. By contrast, the G13R and G13D mutations to bulkier residues did disrupt the GMA dimer in simulations (Extended Data Fig. 2k); this suggests that these two mutants may differ from the G12 and Q61 mutants in their signaling mechanism. Our bioluminescence resonance energy transfer (BRET) experiment showed that indeed the wild type and the G12C, G12V and G12D mutants form assemblies in cells (Fig. 3e), but the G13D mutant does not (Extended Data Fig. 5f).

Experimental validation of the GMA dimer model. A K-Raslox/ K-RAS<sup>MUT</sup> inducible system—wherein cell lines generated from Ras-less mouse embryonic fibroblasts (MEFs)43 are dependent on exogenous K-Ras expression for proliferation-has been reported in a previous study<sup>20</sup>, along with a cell-based fluorescence resonance energy transfer (FRET) platform, to probe K-Ras-K-Ras interactions. In both experiments, K-Ras signaling is constitutive with introduction of oncogenic K-Ras mutants such as G12D or G12C. Because the G12D and G12C mutations are compatible with the dimer (Fig. 3d,e), we used these two systems to test the GMA dimer. The MEF and FRET experiments showed that D154Q and R161E mutations in the background of either G12C or G12D impair cell fitness by disrupting K-Ras-K-Ras interactions. The GMA dimer model is also consistent with those findings, because Asp154 and Arg161 are involved in key salt bridges at the GMA dimer interface (Fig. 3a).

We used the MEF and FRET systems to test residues (Asp30, Glu31, Glu62 and Lys147) involved in the GMA dimer interface (Fig. 3a,b). We found that D30R, E31R, E62R and K147D mutations each impaired cell growth (Fig. 3f and Extended Data Fig. 4b) and reduced ERK phosphorylation (Fig. 3g and Extended Data Fig. 4c). The mutations also disrupted the FRET signal of K-Ras-K-Ras interactions (Fig. 3h and Extended Data Fig. 5a). We further found that a D154Q/K147D double mutation, which was predicted to restore the interaction, indeed recovered the FRET signal (Fig. 3h and Extended Data Fig. 5a). FRET analysis also showed that mutations disrupting the active conformation of Ras, such as T35A, T35S<sup>38</sup> and G60A<sup>44</sup>, also disrupted K-Ras-K-Ras interactions (Extended Data Fig. 5b,c), supporting the connection between GMA dimerization and the active conformation. The substantial set of FRET and MEF results collectively are highly consistent with and provide strong support for the GMA dimer model. Of these results, we find the observed compensatory effect of D154Q and K147D mutations especially compelling.

**Extrapolation of the GMA dimer into a K-Ras helical assembly.** Adding K-Ras proteins to the GMA dimer in a head-to-tail fashion

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**Fig. 2 | The GMA dimer model of K-Ras. a**, Snapshots, taken at the simulation times shown, of the GTP-mediated dimerization of membrane-anchored K-Ras proteins (top view, membrane shown as a mesh). **b**, Upper panel, the root mean squared deviation (r.m.s.d.) of the two K-Ras proteins in the solvent and in the membranous simulation of K-Ras dimerization with respect to the final GMA K-Ras dimer model (the snapshot at 100 µs from the simulation shown in **a**). Lower panel, the Arg135 distance to the GTP  $\gamma$ -phosphate ( $\gamma$ P) at the dimer interface in the simulations. **c**, Comparison of the GTP-Arg135 (or Lys128) interaction with the GAP 'arginine finger'. **d**, The dimer interface; only part of the acceptor is shown. **e**, All 12 NMR-identified residues<sup>16</sup> involved in K-Ras dimerization, shown with respect to the GMA dimer interface. **f**, The conformations of the switch I and II regions in simulations of the GMA dimer. **g**, Distributions of K-Ras membrane orientations in simulations of a K-Ras monomer, an RBD-bound K-Ras monomer and the GMA dimer in the membrane environment; the orientations are mapped to a 2D space defined by the distances of the GTP  $\beta$ -phosphate ( $\beta$ P) and the GTP amine nitrogen of the guanine ring (N2) to the membrane plane (Extended Data Fig. 2d,e). The  $\alpha$  and  $\beta$  regions correspond to the  $\alpha$  and  $\beta$  orientations<sup>40</sup>, respectively; in the former, the  $\alpha$ 4 and  $\alpha$ 5 helices contact the membrane and, in the latter, the  $\beta$  strands do.

extends it into a left-handed helical assembly, which is highly compact and yet does not exhibit any steric clashes (Fig. 1b). In this assembly, each K-Ras protein serves as both a donor and an acceptor (with the exception of the 'head' serving only as a donor and the 'tail' serving only as an acceptor). For convenience, we number the K-Ras proteins in the assembly from 1 to n starting from the head (and incrementing by one from a donor to its acceptor). The helical assembly consists of four K-Ras proteins per turn and has a radius of roughly 50 Å and a pitch of roughly 40 Å, with virtually no unoccupied space between turns or in the center. We placed the K-Ras assembly on the membrane, with the axis of the helical assembly at an approximately 60° angle to the membrane surface (Fig. 1b, left), which maximizes membrane contact with the base tier. We refer to the tier comprising K-Ras 1-4 as the base tier and the four highest-numbered K-Ras proteins as the top tier. The helical assembly can grow in a straight tower shape, but the base tier alone, without further assembly, is stable on the membrane in our simulations (Extended Data Fig. 3i). In the helical assembly, the RBD interface of each K-Ras protein is exposed and available for Raf recruitment. Further, the C and N termini of the K-Ras proteins are positioned outward at the surface of the assembly, allowing the C-terminal tail to interact with other proteins and for fluorescent proteins to fuse with the N termini<sup>45</sup> without disrupting the assembly.

A K-Ras protein at position *n* in our helical assembly model engages in multivalent interactions (Fig. 4a), making contact with up to six other K-Ras proteins: GTP-mediated primary interactions with K-Ras n - 1 and n + 1, secondary (stacking) interactions with K-Ras n - 4 and n + 4 along the axis of the helical assembly, and tertiary interactions with K-Ras n - 3 and n + 3 (Fig. 4c and Extended Data Fig. 6a,b). The secondary and tertiary interactions are predominantly polar, with a buried surface area of roughly 1,450 Å<sup>2</sup> and 660 Å<sup>2</sup>, respectively (Extended Data Fig. 6a,b). Crystallographic

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**Fig. 3 | Mutagenesis validation of the GMA dimer interface. a**, The residues involved in the salt bridges at the dimer interface. **b**, Residue contacts at the dimer interface (black and red lines). Red lines highlight E62-K128 and GTP- $\gamma$ P-R135 contacts, which alternate with E62-R135 and  $\gamma$ P-K128 contacts. **c**, G12, G13 and Q61 oncogenic mutation sites at the GMA dimer interface. **d**, The stability of the G12C, G12D and G12V GMA dimers in simulations in terms of C $\alpha$  r.m.s.d. **e**, BRET signals, which provide an indication of K-Ras assembly, for wild-type (WT) K-Ras and three oncogenic mutants at three different FBS concentrations. **f**, Growth rates of *K*-Ras<sup>for</sup>/*K*-RAS<sup>MUT</sup> cells shown as confluence values measured by IncuCyte. G12C or G12D was in the background of the tested mutations. Representative images at the endpoint are shown in Extended Data Fig. 4a. Cells were kept in 0.1% FBS (see Extended Data Fig. 4b for curves obtained in 0.5%, 1% and 10% FBS). Data are shown as mean ± s.d. (*n* = 3 biologically independent experiments). **g**, Phosphorylation of ERK and AKT in *K*-Ras<sup>for</sup>/*K*-RAS<sup>MUT</sup> cells. Cells were lysed after incubation for 48 h. See Extended Data Fig. 4c for the phosphorylation at other FBS levels. These results are representative of three independent experiments with similar results. **h**, CFP emission in HEK293T cells cotransfected with CFP- and YFP-fused K-Ras (wild type and various mutants), where binding between the CFP- and YFP-fused proteins would bring the fluorescent tags into close proximity, leading to an increased CFP emission after YFP bleaching. Assays were repeated three times. Data are shown as mean ± s.e.m. WT, *n*=35; D154Q, *n*=22; G12C/D154Q, *n*=18; G12D/D154Q, *n*=20; D30R, *n*=30; G12C/D30R, *n*=26; G12D/D30R, *n*=23; E31R, *n*=32; G12C/E31R, *n*=27; G12C/E31R, *n*=23; E62R, *n*=34; G12C/E62R, *n*=33; G12D/E62R, *n*=36; K147D, *n*=25; G12C/K147D, *n*=26; G12D/K147D, *n*=22 and K147D/D154Q, *n*=31 (\*\*\*\*P<0.0001 by two-way ANOVA; NS, not significant).

and simulation data provide independent support for the secondary interaction. In one of the 20 simulations of the association of two K-Ras molecules (Extended Data Fig. 2a), the secondary (stacking) dimer formed and remained stable (Fig. 4d). Further, the secondary dimer structure closely resembles a K-Ras dimer seen in crystal packing (for example, in PDB 5UQW; Extended Data Fig. 6f). The secondary interaction places the  $\beta$ 2– $\beta$ 3 hairpin of K-Ras *n*, which is exploited by covalent inhibitors of K-Ras G12C<sup>46</sup>.

Experimental validation of the helical assembly. To experimentally validate the K-Ras helical assembly, we first used negative-stain electron microscopy to image K-Ras particles in a reconstituted system. In these experiments, full-length K-Ras proteins were tethered to a lipid monolayer by Cys185, forming chemical bonds with maleimide lipids. The K-Ras proteins were then applied to a grid, stained and imaged (Fig. 5h). The K-Ras particles form in a maleimide lipid-dependent manner (Fig. 5a-d), consistent with the notion that K-Ras assembly requires K-Ras membrane localization. Classification of 73,282 visualized particles into 100 classes revealed a range of sizes (Fig. 5e). The variability of particle size is consistent with the notion that the helical assembly can accommodate different numbers of K-Ras molecules, but particle heterogeneity prevented reconstruction of the 3D structures. Consistent with the MEF FRET (Fig. 3f-h) and BRET (Fig. 6d) analyses, the D154Q mutation disrupted particle formation (Fig. 5f). Particle formation was also disrupted by K88D (Fig. 5g), a mutation that disrupted the FRET and BRET signals of K-Ras assembly in cells (Fig. 6d and Extended Data Fig. 5g). These mutagenesis data indicate that, at least partially, the helical assemblies underlie the K-Ras particles we visualized. Based on our model, a subtle yet important difference between the effects of D154Q and K88D is expected: D154Q is located at the primary interface, and it should thus disrupt GMA dimers, whereas K88D is at the secondary interface, and it should thus not interfere with GMA dimerization. This prediction is consistent with the negative-stain images, in which the D154Q particles generally appear to be smaller than the K88D ones (Fig. 5f,g) under identical conditions.

We also evaluated mutations predicted to disrupt the K-Ras helical assembly using the MEF BRET and FRET systems. K-Ras-dependent MEFs with K88D, which was predicted to disrupt the secondary interface, and Q129L and R149D, which were predicted to disrupt the tertiary K-Ras-K-Ras interface (Fig. 6a), slowed proliferation (Fig. 6b and Extended Data Fig. 4f) and impaired ERK phosphorylation (Fig. 6c and Extended Data Fig. 4e). To supplement the FRET experiment, we confirmed using BRET, which allows better quantification, that K88D, Q129L and R149D mutations in the background of G12D disrupt K-Ras assembly (Fig. 6d). Although they are distal to the Raf interface of K-Ras, K88D, Q129L and R149D mutations also hindered K-Ras recruitment of C-Raf to the cell membrane (Fig. 6e), suggesting that the K-Ras assemblies play an important role in K-Ras-Raf interaction.

**Multivalent interactions between C-Raf and the K-Ras assembly.** Using a cocrystal structure of the C-Raf RBD and H-Ras (PDB 4G0N)<sup>47</sup> as a template, we docked a C-Raf RBD to each K-Ras of the



**Fig. 4 | K-Ras-K-Ras and K-Ras-RBD interactions in the signalosome model. a**, Cartoon representation of the relative positions of K-Ras proteins and RBD domains in the model; the membrane is shown as a mesh and GTP is shown in red. **b**, Left, the primary (with K-Ras 5), secondary (with K-Ras 6) and tertiary (with K-Ras 2) RBD-Ras interactions (of RBD 5); a fourth K-Ras (K-Ras 1) is shown for geometric context. The secondary and tertiary interface residues (and GTP) are colored in red and orange. Right, close-up of the Ras-RBD interactions. **c**, Center, three K-Ras proteins engaged in primary, secondary and tertiary interactions with one another; the interface residues are shown in yellow and GTP is shown in red. Left and right, close-ups of the secondary (stacking) and tertiary Ras-Ras interactions, respectively. **d**, Superposition of a K-Ras dimer generated by simulation of K-Ras dimerization in aqueous solution and the secondary K-Ras dimer interactions in the helical assembly model.



**Fig. 5 | Electron microscopy images with negative stain of K-Ras assembly. a-d**, K-Ras wild type at a 10  $\mu$ M concentration; maleimide lipids were used at the following percentages: 0% (**a**), 5% (**b**), 10% (**c**) and 20% (**d**). **e**, 2D class average of the K-Ras wild-type particles. **f**, D154Q K-Ras at a 10  $\mu$ M concentration with 20% maleimide lipids. **g**, K88D K-Ras at a 10  $\mu$ M concentration with 20% maleimide lipids. **g**, K88D K-Ras at a 10  $\mu$ M concentration with 20% maleimide lipids. **h**, Workflow of the experiment. After incubation with the monolayer lipid overnight, the K-Ras particles were moved to a carbon-coated copper grid, washed intensively, negatively stained and examined by electron microscopy. Scale bars, 200 nm.

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**Fig. 6 | Mutagenesis validation of the secondary and tertiary K-Ras-K-Ras interactions. a**, Illustration of the locations of the tested mutations, including those at the primary interface shown in Fig. 3f-h (green) and those at the secondary and tertiary interfaces (yellow). **b**, Growth rates of *K-Ras<sup>lov</sup>/K-RAS<sup>MUT</sup>* cells with K88D, Q129L and R149D mutations in *cis* with either G12C or G12D. Representative images at the endpoint are shown in Extended Data Fig. 4d. See Extended Data Fig. 4f for results obtained in 0.5% and 10% FBS. Data are shown as mean  $\pm$  s.d. (n = 3 biologically independent experiments). **c**, Phosphorylation of ERK and AKT in *K-Ras<sup>lov</sup>/K-RAS<sup>MUT</sup>* cells. See Extended Data Fig. 4e for the phosphorylation in 0.5% and 10% FBS. These results are representative of three independent experiments with similar results. **d**, Left, endpoint BRET measurements for cells with K88D, Q129L and R149D in the background of G12D, compared with the G12D mutation only; D154Q is included as a positive control. K-Ras construct (residues 1–166) lacking the membrane-anchoring hypervariable region was also used as a control (Extended Data Fig. 5f). The BRET signals were normalized to the maximum signal of cells with only G12D. An acceptor/donor expression plasmid ratio of 200:1 was chosen for comparing the K-Ras mutants. The average of three technical replicates was calculated for each of the biological triplicates and subjected to one-way ANOVA followed by Dunnett's post tests (\*\*P < 0.01, \*\*\*P < 0.001). Right, cotransfection of increasing ratios of donor and acceptor plasmids, which allows discrimination between specific and nonspecific (random collision) protein-protein interactions. **e**, Fluorescence imaging of Raf localization of K-Ras mutants K88D, Q129L and R149D; K-Ras wild type and Q61H are used as positive controls, and E37G, C185S and K-Ras-free cells are used as negative controls. Assays were repeated three times and 8-10 regions of interest were imaged each time.

helical assembly. The docked RBDs did not clash with any K-Ras proteins, nor with one another. The RBD that primarily interacts with K-Ras n, which we refer to as RBD n, forms a secondary contact with K-Ras n + 1 and a tertiary contact with K-Ras n - 3 (Fig. 4b and Extended Data Fig. 6c,d). These secondary and tertiary Ras-RBD interactions may confer an advantage on the helical assembly over a K-Ras monomer in terms of RBD binding, and, conversely, RBD binding may help stabilize the K-Ras assembly.

The 52-residue zinc-coordinated Raf CRD also interacts with Ras<sup>48</sup> and imparts specificity to Ras–Raf binding<sup>49</sup>. Informed by experimental data and extensive simulations of a CRD tethered to a structure of a K-Ras–RBD complex (Supplementary Analyses), we positioned the

CRD at the C terminus of the  $\alpha$ 5 helix and  $\beta$  strands of K-Ras (Fig. 7a), with an interface of roughly 1,400 Å<sup>2</sup>. The CRD also contacts the switch II region of K-Ras (Extended Data Fig. 7f); this is consistent with switch II mutations (G60A and Y64W) hindering the CRD binding of H-Ras<sup>50</sup>. NMR and other analyses have identified a set of CRD and Ras residues involved in the Ras–CRD interaction<sup>48,51–53</sup>; these residues are either part of the modeled Ras–CRD interface or near it (Extended Data Fig. 7f,g). By our model, a CRD interacts with a K-Ras and an adjacent RBD with extensive electrostatic complementarity (Extended Data Fig. 7h). We added a CRD to each K-Ras protein in the helical assembly. In the base tier, both the RBDs and CRDs interact extensively with the membrane (Fig. 7a).



**Fig. 7 | Ras interactions with Gal-3 and Raf RBD and CRD domains. a**, Side views of the interactions of two stacking K-Ras proteins with Gal-3 and C-Raf RBD and CRD domains. Center, relative positions of K-Ras, Gal-3, and Raf RBD and CRD domains. K-Ras and RBDs are shown as spheres, CRDs are shown as ribbons and Gal-3 proteins are shown as ribbons surrounded by mesh. Left, the K-Ras proteins with RBDs and Gal-3 proteins in cartoon representation; the galectin-bound fCys185 residues of K-Ras are shown in blue and purple, and Gal-3-RBD interfaces are marked in red. Right, positions of CRD domains and Gal-3 proteins with respect to RBD-bound K-Ras. Ras-CRD interfaces are marked in purple. Also labeled is the  $\alpha$ 2 helix in the switch II region of K-Ras involved in the interaction with the CRD. **b**, Top view of the RBD interaction with Gal-3. Two K-Ras proteins, which form a GTP-mediated dimer, are shown as faded images. The fCys185 of K-Ras 2 is shown. **c**, K-Ras membrane-contacting residues (yellow) in the base tier. **d**, Membrane imprint of the K-Ras and Gal-3 proteins of the base tier, which is enriched in POPS lipids. **e**, GAP interacting with the tail K-Ras (K-Ras 8) of the helical assembly. **f**, SOS1 interacting with the tail K-Ras. In the secondary SOS1-K-Ras interaction, the SOS1 helical hairpin (yellow) is inserted into the switch II pocket of K-Ras 5. **g**, Three interactions using the switch II pocket: the secondary SOS1-K-Ras interaction, the K-Ras-K-Ras stacking involving the  $\beta$ 2- $\beta$ 3 turn (green) and covalent G12C inhibitor binding (cyan).

**Gal-3 plays important roles in the signalosome.** The mechanism of K-Ras membrane localization is commonly thought to be insertion of fCys185. Consistent with this notion, the C-terminal tails of K-Ras molecules at the base tier of the helical assembly contact the membrane, and thus their fCys185 residues can be readily buried in the membrane. For the higher tiers, the C-terminal tails are lifted from the membrane, and the fCys185 residues in our model are capped by Gal-3 molecules. This is motivated by the notion that Gal-3, which can cap the farnesyl group of fCys185, mediates membrane localization of K-Ras<sup>54</sup> and is essential for K-Ras nanoclustering<sup>23,55,56</sup>, as is Gal-1 for H-Ras nanoclustering<sup>24,57,58</sup>.

Since galectins associate with the lipid bilayer<sup>59</sup> and have been shown to anchor Ras to the membrane<sup>57</sup>, we also considered another

scenario in which all fCys185 residues, including those in the base tier, interact with Gal-3 proteins and Gal-3 proteins interact with and localize the K-Ras assembly to the membrane. Although this scenario departs from the current understanding of how fCys185 interacts with the membrane, we find it compelling in that it provides an explanation for the importance of Gal-3 dimerization in facilitating K-Ras nanoclustering<sup>60</sup>. The Ras–Raf signalosome model was stable in simulations with or without base-tier Gal-3 molecules (Extended Data Fig. 1c). Further experimental elucidation will be needed to fully establish the role of Gal-3 in the signalosome.

In the presence of C-Raf RBDs and CRDs surrounding the K-Ras helical assembly, the unoccupied space is limited but can accommodate the Gal-3 molecules. Existing mutagenesis and dimerization data on Gal-3 and Gal-1 with respect to Ras nanoclustering, and extensive simulations (Supplementary Note and Analyses), led us to a model in which Gal-3 n (the Gal-3 protein bound to the fCys185 residue of K-Ras *n*) interacts with RBD n-1 (Fig. 7a,b). Any pair of Gal-3 n and n+4 proteins in the model are stacked parallel to the K-Ras stacking, with the C-terminal tail of K-Ras n+4 positioned near the interface, contacting both Gal-3 proteins (Fig. 7a,d and Extended Data Fig. 7b). As in the GMA dimer, K-Ras proteins in the base tier of the helical assembly adopt the  $\alpha$ -like orientation, contacting the membrane with their  $\alpha$ 5 helices and  $\beta$ 2- $\beta$ 3 hairpins (Fig. 7c), and Gal-3 proteins interact with the membrane extensively (Fig. 7a and Extended Data Fig. 8a,b). This model of K-Ras-Gal-3 interaction resembles resolved K-Ras-PDES structures<sup>61</sup> (see the Supplementary Analyses for a functional comparison of Gal-3 and PDE\delta). From this model, by simply removing the four Gal-3 molecules at the base tier and embedding the fCys185 residues, we obtained a model that is consistent with the generally accepted view that fCvs185 embedding in membrane is crucial to the membrane anchoring of K-Ras. With or without Gal-3 at the base tier, the model was stable overall in our simulation (Extended Data Fig. 1b).

The C-Raf linker and a fuzzy signalosome structure. In C-Raf activation in the MAPK pathway, a 14-3-3 dimer acts as a scaffold that binds to and stabilizes a C-Raf kinase domain dimer<sup>62</sup>. C-Raf contains a loop C-terminal to the kinase domain that is phosphorylated at Ser621 and binds to 14-3-3 in C-Raf activation. There are multiple proteins in the 14-3-3 family; in the modeling, we incorporated 14-3-30 into the Ras-Raf signalosome. Each C-Raf kinase domain also binds to and phosphorylates MEK1 kinase. We separately modeled the C-Raf kinase domain dimer complexed with a 14-3-3 $\sigma$  dimer and two MEK1 kinase domains (Fig. 1f) using crystal structures<sup>63-65</sup> that were available at the time we were constructing the model (Fig. 1g) (Supplementary Note and Analyses). The resulting model is overall highly consistent with subsequently reported crvo-EM structures of the Raf-14-3-3 heterotetramer (Extended Data Fig. 9a)<sup>66,67</sup>. C-Raf protein contains a roughly 200-residue-long, unstructured linker between its CRD and kinase domains. We modeled such linkers to connect the RBD and CRD domains to the kinase domains (Supplementary Note and Analyses). We obtained a diverse set of conformations with transient secondary structures (Extended Data Fig. 9e). We then grafted a linker in a simulation-generated conformation onto each CRD in the model and then grafted onto each linker pair a C-Raf kinase domain dimer bound to 14-3-3 $\sigma$  and MEK1 (Fig. 1f). Incorporating this model into the larger model then produced our full signalosome model. The unstructured linkers introduced 'fuzziness' to the signalosome, which is often seen in signalosome complexes<sup>68</sup>.

GEF and GAP regulate the size of the Ras-Raf signalosome. GAPs, such as RasGAP, catalyze the hydrolysis of GTP in Ras to revert it to its GDP-bound state<sup>7</sup>. Ras nanoclustering has been suggested to attenuate RasGAP-catalyzed GTP hydrolysis<sup>55,56</sup> and to sequester Ras from GAP binding<sup>23,57</sup>. Consistent with these results, RasGAP is sterically prevented from interacting with all but the tail K-Ras protein in the helical assembly. Based on an H-Ras-p120GAP structure<sup>33</sup>, GAP can dock to a tail K-Ras protein without steric clashes, provided that the K-Ras protein is not bound to Raf (Fig. 7e). GAP can then convert the tail K-Ras protein to the GDP-bound state, preventing any additional K-Raf protein from joining the helical assembly and promoting dissolution of the signalosome. Compared to the other K-Ras molecules in the signalosome, the tail K-Ras is most exposed and likely the least stable. We thus anticipate that, under GAP regulation, a signalosome will eventually disassemble by one tail-position K-Ras falling off at a time.

Based on a structure of H-Ras in complex with the GEF protein SOS1 (ref.  $^{69}$ ), an SOS1 protein in its active conformation can also

dock to the tail K-Ras protein (Fig. 7f). By ensuring that the tail K-Ras protein is in the GTP-bound state, ready to receive another K-Ras protein into the signalosome, SOS1 may help maintain and grow the signalosome. SOS1 is activated in part by an allosteric GTP-bound Ras protein<sup>70</sup>, which our model accommodates. The model predicts, moreover, that if an active SOS1 protein docks to the tail K-Ras protein *n*, it additionally engages K-Ras n-3 in a secondary interaction in which a helical hairpin of SOS1 is inserted into the switch II pocket of K-Ras n-3 (Fig. 7f,g). This secondary Ras–SOS1 interaction may contribute to the specificity of SOS1 in regulating MAPK signaling. One consequence of this SOS1 interaction is that it may place an upper limit on the height (and thus overall size) of the signalosome, since SOS1 is membrane-anchored by its pleckstrin homology domain and thus has only limited reach from the membrane.

#### Discussion

The Ras–Raf signalosome model we propose here entails a host of multivalent interactions. Although each interaction is weak on its own, together they give rise to a stable signalosome that provides a favorable composite interface for Raf recruitment. Signalosome formation is likely a nucleation-like process, which should yield a switch-like signal. The low affinity of the GMA dimer means that the signalosome cannot form without the membrane, which promotes GMA dimerization by raising the local concentration of K-Ras and restricting its orientation in favor of dimerization and signalosome formation (Fig. 2g).

Two (the  $\alpha$  and  $\beta$ ) membrane orientations of monomeric K-Ras have been observed (Fig. 2g)<sup>40</sup>, and they appear to be associated with different K-Ras assemblies. In addition to the widely discussed symmetric dimer<sup>13,20</sup> that has been observed using crystallography and was also generated by our simulation (Extended Data Fig. 2b), a number of Ras dimer models<sup>71–73</sup> and a pentamer model have been previously proposed<sup>15</sup>. Several crystal and NMR structures of a K-Ras–CRD complex, which differ from our model, have also recently been reported<sup>74–76</sup>. Notably, whereas the GMA dimer model imposes the  $\alpha$  orientation of K-Ras on the membrane, these alternative dimer and pentamer models all impose the  $\beta$  orientation. We speculate that these differences may result from differences in the local environment—such as membrane composition and Gal-3 concentration—and may lead to distinct downstream signals.

Proteins downstream of Ras may also affect signalosome formation. Certain small-molecule inhibitors of B-Raf give rise to 'paradoxical activation' of Raf by inducing C-Raf–B-Raf heterodimerization and even C-Raf homodimerization<sup>64</sup> and also promote Ras nanoclustering by inducing more, but not larger, Ras nanoclusters in cells<sup>77</sup>. Consistent with this surprising B-Raf inhibitor effect, our model suggests a scenario in which enhanced Raf dimerization promotes nucleus formation and, in turn, signalosome formation. Similarly, 14-3-3 proteins may promote Raf dimerization and in turn Ras assembly, enabling a feedback loop in MAPK signaling.

The K-Ras helical assembly provides a possible explanation for why wild-type K-Ras acts as a tumor suppressor in cells driven by oncogenic K-Ras mutants<sup>78,79</sup>. Wild-type K-Ras, which is predominantly GDP bound and cannot serve as a GTP donor, may inhibit the growth of a helical assembly by capping it at the tail position. This scenario is consistent with the D154Q K-Ras mutant being inactive as a tumor suppressor<sup>20</sup>, since the D154Q mutation at the acceptor interface may prevent the mutant from assuming the tail position. This rationale leads us to predict that mutations at the donor interface (for example, K147D and R161E) should not disrupt the tumor suppressor activity, as these mutants should still be able to assume the tail position and cap helical assemblies of oncogenic K-Ras mutants.

It is worth noting that the K-Ras helical assembly is almost certainly not applicable to PI3K activation (Supplementary Analyses) or to Raf

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signaling involving KSR pseudokinase as a scaffold, as in this case K-Ras dimerization may not be required<sup>80</sup>. Moreover, many proteins not incorporated in our signalosome model (for example, SHOC2, ref. <sup>81</sup>; SUR-8, ref. <sup>82</sup>; Aurora kinase A<sup>83</sup>; and SHP2 phosphatase<sup>84</sup>) are involved in Ras signaling and may, for example, prime K-Ras for the signalosome without being part of it. In many respects, however, findings on H-Ras nanoclustering echo those on K-Ras<sup>45</sup>, and our signalosome model is consistent with the structures of other Ras and Raf proteins (such as N-Ras, H-Ras, A-Raf and B-Raf) involved in MAPK signaling. With local alterations, our model may be extendable to those Ras and Raf proteins and provide a framework for understanding their overlapping, yet distinct roles in MAPK signaling.

#### **Online content**

Any methods, additional references, Nature Research reporting summaries, source data, extended data, supplementary information, acknowledgements, peer review information; details of author contributions and competing interests; and statements of data and code availability are available at https://doi.org/10.1038/ s41594-021-00667-6.

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#### Methods

Molecular dynamics simulations. Simulation protocol and force field parameters. Molecular dynamics simulations were performed on the special-purpose supercomputer Anton 2 (ref. 88). The simulated systems ranged in size from around 37,000 to 1,740,000 atoms, and were parameterized using the TIP3P model<sup>8</sup> for water molecules, the Amber99SB\*-ILDN force field<sup>90-95</sup> for proteins and the CHARMM36 force field<sup>96</sup> for lipids. Specialized force field parameters were used for farnesylated cysteine97, phosphorylated serine and tyrosine98, and GDP and GTP<sup>99</sup>. Simulations of the disordered C-Raf linker by itself were exceptional in that the simulation systems were parameterized using a99SB-disp<sup>100</sup> and the TIP4P water model<sup>101</sup>. a99SB-disp is a variant of the Amber force field developed to work together with TIP4P for accurate simulations of disordered (as well as ordered) protein states. Due to concerns about force field inaccuracy for Zn2+ ions, distance restraints were applied between CRD zinc finger residues and Zn2+ ions in simulations that included the zinc-coordinated Raf CRD domain. Molecular dynamics simulations were performed in the NPT ensemble with constant pressure (1 bar) and constant temperature (310 K) imposed by a Martyna-Tuckerman-Klein Nosé-Hoover chain coupling scheme<sup>102</sup>, which was implemented using a multigrator scheme<sup>103</sup> with a relaxation time of 10 ps. Initial velocities were sampled from the Boltzmann distribution. Water molecules and all bond lengths to hydrogen atoms were constrained using an in-house implementation<sup>104</sup> of M-SHAKE<sup>105</sup>. Production simulations (for data collection) were launched after energy minimization and nanosecond-timescale molecular dynamics simulation with harmonic position restraints on backbone atoms.

The protein backbone atoms were restrained to their initial positions using an initial harmonic potential with a force constant of  $10 \text{ kcal mol}^{-1} \text{Å}^{-2}$  for 10-500 ns as an equilibration step; the force constant was linearly reduced to zero over the course of the equilibration step. The van der Waals and short-range electrostatic interactions were cut off at 10 Å for simulations without membrane and 12 Å for simulations with membrane. Long-range electrostatic forces were calculated in k-space using a grid-based method with Gaussian spreading<sup>106</sup> to the grid every 7.5 fs. The simulation time step was 1 fs for the equilibration stage and 2.5 fs for production simulations; the r-RESPA integration method<sup>107</sup> was used, with long-range electrostatics evaluated every 7.5 fs.

*System preparation.* Structures derived from the PDB were back-mutated into the wild type unless stated otherwise, and missing atoms and residues were built in. The system of interest was placed at the center of a simulation box that was cubic for solvent simulations and orthorhombic for membranous simulations, with a separation greater than 20 Å from any periodic image. Explicitly represented water molecules were added to fill the system, and Na<sup>+</sup> and Cl<sup>-</sup> ions were included to maintain physiological salinity (150 mM) and to obtain a neutral total charge for the system. Residue protonation states corresponded to pH7.

In membranous simulations, a phosphotidylcholine (POPC) lipid bilayer with roughly 30% phosphotidylserine (POPS) in the inner layer proximal to the proteins was used. The model membrane was built from a neutral POPC lipid membrane by replacing 15% (molar) POPC with negatively charged POPS lipids. (The POPC lipids to replace were all taken from the inner bilayer but otherwise chosen at random.) This POPS fraction was chosen to mimic the abundance of anionic lipids in the mammalian plasma membrane<sup>20,108,109</sup>, in which approximately 10% of all lipids are POPS species, with other anionic species, such as phosphoinositides, bringing the total content of anionic lipids up to 15%. The POPS lipids were introduced only in the intracellular leaflet, which is where anionic lipids are almost exclusively found in cell membranes. The lipid content was thus around 30% POPS and 70% POPC in the intracellular leaflet and 100% POPC in the extracellular leaflet.

Simulations of K-Ras dimerization. We performed 20 simulations, each 10  $\mu$ s long, of two GTP-bound K-Ras molecules (PDB 4DSN) in aqueous solvent. In addition, we performed 23 unbiased simulations, each at least 2  $\mu$ s long, of two GTP-bound K-Ras proteins anchored to the membrane by their fCys185 residues.

*FRET assays.* Full details of our FRET assay are available elsewhere<sup>20</sup>. Briefly, human embryonic kidney 293T (HEK293T) cells were cotransfected with paired cyan fluorescent protein (CFP)- and yellow fluorescent protein (YFP)-fused *KRAS* constructs under the conditions of 0.5 and 10% FBS in four-well chambered cover glass (Lab-Tek). After 36–48 h, live cell imaging was performed using a Confocal/ Multiphoton Zeiss LSM880 microscope. To examine the GTP-dependent Ras-Raf interaction, cotransfection of CFP-K-RasWT and YFP-C-Raf, CFP-K-RasT35A and YFP-C-Raf, CFP-K-RasG60A and YFP-C-Raf, or CFP-K-RasG13D and YFP-C-Raf plasmids was conducted. After 24 h of transfection, cells underwent 22 h of serum starvation, followed by epidermal growth factor (EGF) (10 ng ml<sup>-1</sup>) exposure for 30 min, and then were subjected to microscopy. Data were collected from three biological repeats, and 10–12 different cells in different fields from the same coverslip were selected for microscopy. Quantitation was done using ZEN software (Zeiss).

Preparation of GMP-PNP-loaded K-Ras protein for electron microscopy studies. K-Ras (1–188) wild-type protein was expressed and purified as described

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previously<sup>110</sup>. Point mutations were generated using the GeneArt Site-Directed Mutagenesis System (Life Technologies). K-Ras proteins (100  $\mu$ M) in buffer (20 mM HEPES pH 8.0, 150 mM NaCl, 15 mM EDTA and 200 mM (NH<sub>4</sub>)<sub>2</sub>SO<sub>4</sub>) were loaded with 200  $\mu$ M GMP-PNP at 4°C overnight. The reaction was terminated by buffer exchange to 20 mM HEPES pH 8.0, 150 mM NaCl, 200  $\mu$ M GMP-PNP, 10 mM MgCl<sub>2</sub> with Zeba Spin Desalting Columns (Thermo Scientific). GMP-PNP loading was verified by back extraction of nucleotide using 6 M urea and evaluation of nucleotide peaks by high-performance liquid chromatography using an ion-exchange column as described previously<sup>110</sup>.

Preparation of lipid monolayers. Phospholipids obtained from Avanti Polar Lipids were mixed according to the following proportions in chloroform:methanol (3:1, vol/vol): 10% DOPS, 10% DOPE, 60% Egg-PE and 20% PE-MCC<sup>111</sup>. Assembly of protein on lipid monolayers was patterned following previous methods<sup>112</sup>. Briefly, droplets (15 µl) of K-Ras (10 µM) in buffer were added to Teflon depression wells (3 mm in diameter and 0.5 mm in depth). 0.5 µl of 1 mM lipid solution was then delivered to the surface of the droplet. The wells were incubated in an airtight humidified chamber at 4°C overnight. The lipid monolayers at the air–liquid interface were collected by hydrophobic carbon-coated grids (catalog no. CF400-Cu, Electron Microscopy Sciences). After washing with buffer three times, the grids were blotted and stained with uranyl formate solution (1% wt/vol) three times.

Transmission electron microscopy data collection and processing. Stained grids were examined in an FEI Tecnai G2 Spirit Biotwin Transmission Electron Microscope operated at 120 kV with a magnification of 13,000 times. Images were collected with a pixel size of 0.78 nm. RELION 3.0 was used for all image processing<sup>113</sup>. For wild-type K-Ras reconstructions, 73,282 particles were computationally selected from 185 micrographs and subjected to 2D classification into 100 classes.

*Raf localization imaging.* Raf localization was evaluated by cotransfection of plasmids expressing CFP–K-Ras<sup>R489D</sup> and YFP–C-Raf, CFP–K-Ras<sup>R129L</sup> and YFP–C-Raf, and CFP–K-Ras<sup>R149D</sup> and YFP–C-Raf. Constructs were designed as previously reported<sup>23</sup>. Cells were cultured in 10% FBS. Then, 36h after transfection, cells were subjected to microscopy. Images were selected from three biological replicates, and 10–20 different cells in different fields from the same coverslip were used for analysis. Imaging was done using ZEN software (Zeiss).

BRET assays. Construct design. N-terminal fusions of *KRAS* to either mNeonGreen or NanoLuc were constructed in a pcDNA3.1(+) vector (Invitrogen) as reporters for BRET assays. *KRAS* point mutations were introduced either by site-directed mutagenesis (QuikChange, Agilent) or by replacing the *KRAS* gene with a synthetic DNA string (Geneart) carrying the desired mutation. All sequences were confirmed by sequencing.

Transfection. HEK293T cells were cultured in DMEM complete medium (10% FBS) and seeded in white 96-well clear-bottom plates (Corning) 24 h before transfection. Cotransfections of the reporter plasmids were carried out with TransIT-293 (Mirus) using the manufacturer's suggested protocol, using ratios as indicated in the figures. Then, 24 h after transfection, medium was exchanged to DMEM with no phenol red (Life Technologies), containing varying amounts of FCS (0.1%, 4.0% and 10.0%).

<u>BRET measurements and data analysis</u>. Here, 48 h after transfection, BRET measurements were taken on a Victor 3 Multilabel Plate Reader after the addition of 10µl of 32µM coelenterazine 400a (Cayman Chemical) resulting in a 2.9µM final concentration. Emission of mNeonGreen and NanoLuc was observed for 2 s at 535 $\pm$ 25 and 460 $\pm$ 25 nm, respectively. Expression of both reporters was monitored by measuring mNeonGreen (excitation 485 nm, emission 535 nm) before the addition of luciferase substrate and measuring total luminescence directly after the BRET measurements for 0.3 s.

BRET ratios were calculated based on the following equation for each transfected well:

BRET ratio = Em<sub>535 nm</sub>/Em<sub>420 nm</sub> - Cf

 $Cf = Em_{535 nm}/Em_{420 nm}$  (donor only samples)

The average of three technical replicates was calculated for each of the biological duplicates or triplicates after removal of obvious outliers and subjected to statistical analysis. Statistical analyses were performed using one-way ANOVA followed by Dunnett's post tests (\*P < 0.05, \*\*P < 0.01, \*\*\*P < 0.001, \*\*\*P < 0.0001).

Plasmids used in the FRET experiments. pcDNA3-CFP (catalog no. 13030), pcDNA3-YFP (catalog no. 13033) and pBABEpuro-CRAF (catalog no. 51124) plasmids were purchased from Addgene. Full-length fragments of KRASWT and CRAF were inserted into vectors containing CFP or YFP to obtain CFP-KRASWT, YFP-KRASWT and YFP-CRAF constructs. KRASG12C, KRASG12D, KRASG13D, KRASD30R, KRASE31R, KRAST35A, KRAST35S, KRASG60A, KRASE62R,

KRASR135A, KRASK147D, KRASD154Q, KRASG12C/D30R, KRASG12D/D30R, KRASG12C/E31R, KRASG12D/E31R, KRASG12C/E62R, KRASG12D/E62R, KRASG12C/R135A, KRASG12D/R135A, KRASR135A/D154Q, KRASG12C/ K147D, KRASG12D/K147D and KRASK147D/D154Q mutants were generated by site-directed mutagenesis using PfuUltra II Hotstart PCR Master Mix (catalog no. 600850-51). The sequences were confirmed by sequencing.

*Generation of K-Ras<sup>kax</sup>/K-RAS<sup>MUT</sup> cells.* Full details of our generation of *K-Ras<sup>lax</sup>/K-RAS<sup>MUT</sup>* cells are available elsewhere<sup>20</sup>. Briefly, K-RAS<sup>D154Q</sup>, K-RAS<sup>D30R</sup>, K-RAS<sup>E31R</sup>, K-RAS<sup>E62R</sup>, K-RAS<sup>K147D</sup> and K-RAS<sup>A135R</sup> mutations, in *cis* with either a G12C or G12D mutation, were created by point mutagenesis from pBABE HA-tagged K-RAS<sup>WT</sup> retroviral plasmid (provided by C. Der, Addgene plasmid no. 75282). Retroviruses were generated by cotransfection of pBABE plasmids together with pAmpho plasmid into HEK293T cells using FuGENE HD Transfection Reagent (Promega). The retroviruses were transduced into *H-Ras<sup>-/-</sup>; K-Ras<sup>lax/kx</sup>* MEFs<sup>43</sup> followed by 2 weeks of puromycin selection (1 µg ml<sup>-1</sup>) in DMEM supplemented with 10% FBS, 100 µg ml<sup>-1</sup> penicillin and 100 units ml<sup>-1</sup> streptomycin. To obtain *K-Ras<sup>lax</sup>/K-RAS<sup>MUT</sup>* clones, we then cultured cells in the presence of 4-hydroxytamoxifen (4OHT) (Sigma, 600 nM) for another 2 weeks to achieve complete deletion of endogenous *K-Ras* alleles.

Growth assessment by IncuCyte. Cells  $(1 \times 10^3)$  were seeded in 96-well plates in 150 µl of DMEM complete medium. The following day, 10% FBS medium was replaced by cell starvation medium (1%, 0.5% or 0.1% FBS) after two washes with PBS. Plates were incubated in the IncuCyte Zoom for real-time imaging, with three fields imaged per well under ×10 magnification every 2h. Data were analyzed using the IncuCyte Confluence v.1.5 software, which quantified cell surface area coverage as confluence values. IncuCyte experiments were performed in triplicate. A single representative growth curve is shown for each condition.

Western blot analysis. Cells were lysed in RIPA lysis buffer (catalog no. 89900, Thermo Fisher) supplemented with protease and phosphatase inhibitor cocktail tablets (Roche). The antibodies used for western blotting included those against HA-Tag (6E2) (Cell Signaling, catalog no. 2367, 1:1,000), HSP90 (H114) (Santa Cruz Biotech, catalog no. sc-7947, 1:1,000), phosphorylated Akt (Ser473) (Cell Signaling, catalog no. 4060, 1:1,000), Akt (Cell Signaling, catalog no. 9272, 1:1,000), phosphorylated ERK1/2 (Cell Signaling, catalog no. 4370, 1:1,000), ERK1/2 (Cell Signaling, catalog no. 4665, 1:1,000), phosphorylated S6 (Ser235/236) (Cell Signaling, catalog no. 4858, 1:1,000), S6 ribosomal protein (Cell Signaling, catalog no. 2217, 1:1,000), antirabbit IgG, HRP-linked secondary antibody (Cell Signaling, catalog no. 7074P2, 1:2,000), ECL Sheep anti-Mouse IgG, HRP-linked secondary antibody (GE Healthcare, catalog no. NA931V, 1:4,000) and ECL donkey antirabbit IgG and HRP-linked secondary antibody (GE Healthcare, catalog no. NA934V, 1:2,000).

*Nucleotide-exchange assay.* Our SOS1-mediated Ras nucleotide-exchange assay was performed at Icagen. The purified K-Ras mutant R135A was diluted in assay buffer (40 mM HEPES pH7.5, 1.5µM Mant-GDP, 10 mM MgCl<sub>2</sub>, 0.05% CHAPS and 0.01% NP-40) to a final concentration of 1µM. The exchange reaction was triggered by adding purified human SOS1 (residues 564–1049, 0.05µM) to the reaction mix. Kinetic readings were taken to measure fluorescence (excitation at 360 nm, emission at 450 nm) for 30 min at 30-s intervals using a 384-well plate reader.

Bioinformatics analysis. The K-Ras sequences used for evolution analysis were compiled using protein–protein BLAST searches in the NCBI nonredundant protein database, with the human K-Ras sequence (residues 1–166) as reference. The initial search generated 20,000 sequences with a minimal sequence identity of 29.84%. K-Ras sequences of mammals, birds and fish were then extracted from the initial search results based on taxonomy IDs. The sequence pool was then filtered based on the following rules: (1) duplicate entries or near-identical (sequence identity greater than 95%) entries were removed; (2) sequences with large (greater than 25 amino acids) insertions or deletions compared to human K-Ras were removed; and (3) sequences labeled as 'partial, 'synthetic' or 'predicted' were removed. After the filtering, 334 sequences remained and were used for multiple-sequence alignment. Sequence alignment of these sequences was obtained from the NCBI server using default settings, and sequence logos were generated using the online tool WebLogo3. The final figures were manually adjusted from the WebLogo3 result to show only the residues corresponding to human K-Ras (residues 1–166).

**Reporting Summary.** Further information on research design is available in the Nature Research Reporting Summary linked to this article.

#### Data availability

Atomic coordinates of the structural model of the eight-protomer Ras–Raf signalosome are available as Supplementary Dataset 1, and atomic coordinates of the GMA K-Ras dimer on a membrane are available as Supplementary Dataset 2. Due to the large size of the molecular dynamics trajectories (listed in Supplementary Table 1), they are available upon request (for noncommercial use) by contacting trajectories@deshawresearch.com. Source data are provided with this paper.

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#### Code availability

The molecular dynamics simulations were performed using the Anton 2 supercomputer (the simulation code we used is specialized to Anton 2, but codes for performing molecular dynamics simulations are widely available).

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#### Author contributions

Y.S. conceived this study and, with D.E.S., oversaw molecular dynamics simulations and analysis. V.P.M., Q.W. and M.R.T. performed and analyzed molecular dynamics simulations. Y.S. and V.P.M. designed molecular dynamics simulations, built the structural model of the signalosome and, with K.D.W., C.A., P.A.J. and J.N.K., selected mutations for experimental testing. K.D.W. and Z.W.Z. designed and analyzed FRET assays. Z.W.Z. performed the FRET assays. C.A. designed and analyzed cell proliferation and western blot assays. C.A. and J.J.O. performed these assays. J.N.K. and A.P. designed and analyzed BRET assays. J.N.K and G.N.-D. performed these assays. K.D.W., X.B., L.L. and C.L. designed and analyzed negative-stain assays. L.L. and C.L. performed these assays. Y.S., V.P.M., K.D.W., C.A., P.A.J. and D.E.S. wrote the paper. Y.S., C.A., K.D.W., P.A.J., A.P. and D.E.S. supervised the overall research.

#### **Competing interests**

P.A.J. reports grants from AstraZeneca, Boehringer Ingelheim, Daiichi Sankyo, Eli-Lilly, Takeda Oncology, Astellas, PUMA and Revolution Medicines and personal fees from AstraZeneca, Boehringer Ingelheim, Pfizer, Roche/Genentech, Eli-Lilly, Chugai, Ignyta, Loxo Oncology, SFJ Pharmaceuticals, Voronoi, Daiichi Sankyo, Biocartis, Novartis, Sanofi Oncology, Takeda Oncology, Mirati Therapeutics, Trasncenta, Silicon Therapeutics, Syndax, Bayer, Esai, Allorion Therapeutics, Accutar Biotech and AbbVie outside the submitted work; P.A.J. also receives postmarketing royalties from a DFCI-owned patent on EGFR mutations issued and licensed to Lab Corp. K.D.W. receives or has received grant funding from Revolution Medicines and Astellas outside the submitted work. He also is on the scientific advisory board for Vibliome Therapeutics. The remaining authors declare no competing interests.

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Correspondence and requests for materials should be addressed to Yibing Shan or David E. Shaw.

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#### ARTICLES **NATURE STRUCTURAL & MOLECULAR BIOLOGY** А В С 16-protomer signalosome model (Ras, Gal, Raf, 14-3-30 and MEK) with Gal-3 at the base tier without Gal-3 at the base tier RMSD (Å) RMSD (Å) CRD 0 80 20 40 60 100 25 7 9 10 Time (µs) Time (µs)

**Extended Data Fig. 1 Construction of the Ras-Raf signalosome model in 16-protomer form. a**. Top and side views of the stepwise addition (left to right) of Gal-3 (yellow), C-Raf (green), 14-3-3 $\sigma$  (blue), and MEK1 (violet) to the K-Ras helical assembly (red), producing the 16-protomer signalosome model. The membrane is shown as a mesh. Each C-Raf KD dimer binds to a 14-3-3 $\sigma$  dimer and two MEK1 kinases. **b**. The C $\alpha$  atom RMSD (w.r.t. the starting structure) of a membrane-anchored 8-protomer Ras-Raf signalosome and its K-Ras octamer core in a 100- $\mu$ s simulation. As shown, the signalosome—especially the K-Ras core—was stable in the course of the simulation. **c**. Similar to B, the C $\alpha$  atom RMSDs of a membrane-anchored 8-protomer Ras-Raf signalosome without Gal-3 at the base tier.

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Extended Data Fig. 2 | See next page for caption.

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Extended Data Fig. 2 | The GTP-mediated K-Ras dimer model and its orientation on the membrane. a. Left panel: A collection of snapshots of the 20 simulations starting from two separated K-Ras proteins in solvent. The snapshots are aligned to one K-Ras protein therein (cyan). The GMA dimer model is highlighted. The  $\alpha$ 4 helix is shown in cartoon representation. Snapshots 10  $\mu$ s apart were taken from the 20 simulations (which have an aggregate simulation time of 680 µs). Right panel: A collection of snapshots of the 23 simulations of two GTP-bound K-Ras proteins on the membrane; snapshots 10 µs apart were taken from the 23 simulations (which have an aggregate simulation time of 210 µs). Only in one solvent and one membrane simulation was the GMA dimer reached, and in both cases the dimer remained stable. b. A stable dimer (orange) generated from one of the 20 simulations of K-Ras/K-Ras association in solvent, which is consistent with the crystal structures of Ras symmetric dimer using the  $\alpha$ 4- $\alpha$ 5 interface (for example PDB 1QRA, cyan and green). c. Upper panel: Acceptor (yellow) interaction with the γ-phosphate of the GTP donor (cyan) in a simulation of a GMA K-Ras dimer. The Arg135 of the acceptor interacts with the  $\gamma$ -phosphate, while the Lys128 interacts with the Glu62 of the donor, and vice versa. Snapshots at 40 µs and 60 µs from the simulation of K-Ras dimerization on the membrane are shown; the motions of Arg135 and Lys128 are shown by dashed red arrows. Lower panel: Distances from Arg135 (black) and Lys128 (red) to the y-phosphate. d. Comparison of GTP interactions and GMPPNP interactions with the acceptor residues at the GMA dimer interface. e. Interface residues of the GMA dimer (left) and water occupancy map (right, blue) at the interface. f. Snapshots at 10 µs from four independent runs of free GTP-bound K-Ras monomer (cyan) on the membrane. To illustrate the relationship between the membrane orientation and RBD binding of K-Ras, the C-Raf RBD (green) is positioned on each snapshot based on the Ras-RBD structure (PDB 4G0N). Shown are the distance of the  $\beta$  phosphorus ( $\beta$ P) and the amine nitrogen of the guanine ring (N2) from the plane of the membrane surface; we use these distances to describe the membrane orientation of a Ras protein (Fig. 2g). The Run 1 snapshot is compatible with RBD binding, which corresponds to the Ras membrane orientation of the lower red contours in Fig. 2g in the main text. The snapshot from Run 2 corresponds to the right center of the top red contours; this orientation of K-Ras on the membrane positions the RBD away from the membrane. In the Run 3 and Run 4 snapshots, RBD binding leads to a severe steric clash with the membrane; this corresponds to the left center of the top red contour. g. Snapshot at 100 µs from the simulation of the GMA dimer formation on the membrane, with two C-Raf RBDs (green) added based on the Ras-RBD pose; the RBDs do not clash with the membrane. The membrane orientation of the donor and acceptor Ras proteins correspond to the lower and upper black contours, respectively, in Fig. 2g in the main text. h. Examples of G nucleotide-mediated dimerization, in which an arginine interacts with a nucleotide phosphate at the dimerization interface. The Toc34 homodimer<sup>114</sup> (PDB 3BB1), the Ffh-FtsY heterodimer<sup>115</sup> (PDB 1RJ9), and the adenylosuccinate synthetase homodimer (PDB 4M9D) are compared to the GMA K-Ras dimer model (cyan); the host G proteins (not shown) were aligned in this comparison. i. The binding site of the synthetic monobody NS1<sup>19</sup> (red) and the donor site (cyan) of an acceptor in the GMA dimer are shown on a Ras protein (yellow). The two binding sites largely overlap. j. The electrostatic complementarity at the GMA dimer interface, with contacting areas at the interface connected by dashed lines. k. Cα RMSD of the GMA dimer of the wild type and various oncogenic mutants in simulations. I. Sequence conservation of K-Ras proteins in various species (see Materials and Methods). A graphical representation of sequence alignment of representative K-Ras proteins in evolution. The residues involved in the K-Ras/K-Ras interactions in the K-Ras helical assembly are labeled. The 32 K-Ras sequences by GenBank IDs are: OLS17184.1, OLS30914.1, OLS23071.1, XP\_020603162.1, XP\_023347765.1, XP\_023347766.1, XP\_003378992.1, XP\_003377451.1, GBC14959.1, XP\_027484897.1, QBM87817.1, XP\_013758736.1, EHB13737.1, OQV11758.1, XP\_027289840.1, NP\_001356715.1, XP\_027290137.1, XP\_027290057.1, XP\_027289839.1, XP\_027289566.1, XP\_029436565.1, ELK24704.1, XP\_027290375.1, XP\_023664743.1, XP\_021777811.1, XP\_021247616.1, XP\_025064324.1, XP\_006127724.1, NP\_001243091.1, RDD38924.1, KPM02442.1, OTF69949.1.

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Extended Data Fig. 3 | Stability of the active switch I and switch II conformations. a. Distributions of the switch I conformation of K-Ras in various simulations. Normalized histograms of the switch I backbone root-mean-square deviation (RMSD) with respect to the active switch I conformation in PDB 4DSN are shown. b. Conformations of the switch I (residues 30-38, cyan) and switch II (residue 60-76, yellow) regions in simulations of the GTP-bound K-Ras monomer. c. Conformations of the switch I and II regions in simulations of GDP-bound K-Ras monomer. d. Conformations of the switch I and II regions in simulations of GTP-bound K-Ras bound with C-Raf RBD. e. Crystal structures of Ras bound with GTP or GTP analogs, with the switch I and switch II regions highlighted. The PDB entries included are 1AGP, 1CLU, 1CTQ, 1GNP, 1GNR, 1HE8, 1JAH, 1JAH, 1K8R, 1LFO, 1LFD, 1NVU, 1NVV, 1NVV, 1NVX, 1P2S, 1P2T, 1P2U, 1P2V, 1PLJ, 1PLK, 1QRA, 1RVD, 1ZW6, 2C5L, 2RGA, 2RGB, 2RGC, 2RGD, 2RGE, 2RGG, 2UZI, 2VH5, 3DDC, 3GFT, 3I3S, 3K8Y, 3L8Y, 3L8Z, 3LBH, 3LBI, 3LBN, 3OIU, 3OIV, 3OIW, 3RRY, 3RRZ, 3RS0, 3RS2, 3RS3, 3RS4, 3RS5, 3RS7, 3RS0, 3TGP, 3V4F, 4DLR, 4DLS, 4DLT, 4DLU, 4DLV, 4DLW, 4DLX, 4DLY, 4DLZ, 4DSN, 4DSO, 4DST, 4EFL, 4EFM, 4EFN, 4GON, 4G3X, 4K81, 4L9W, 4NMM, 4NYI, 4NYJ, 4NYM, 4RSG, 4XVQ, 4XVR, 5B2Z, 5B30, 5P21, 6Q21, 121P, 421P, 521P, 621P, 721P, 821P, and 221P. f. Crystal structures of Ras loaded with GDP. The PDB entries included are 1AA9, 1CRP, 1CRQ, 1CRR, 3L05, 11OZ, 1LF5, 1PLL, 1Q21, 1WQ1, 1XD2, 1XJ0, 1ZVQ, 2CE2, 2CLD, 2Q21, 2QUZ, 2X1V, 3CON, 3KUD, 4DSU, 4EPR, 4EPT, 4EPV, 4EPW, 4EPX, 3EPY, 4L8G, 4L9S, 4LDJ, 4LPK, 4LRW, 4LUC, 4LV6, 4LYF, 4LYH, 4LYJ, 4M10, 4M15, 4M1T, 4M1W, 4M1Y, 4M21, 4M22, 4OBE, 4PZY, 4PZZ, 4Q01, 4Q02, 4Q03, 4Q21, 4QL3, 4TQ9, 4TQA, 4WA7, and 5F2E. g. Crystal structures of the T35S Ras mutant loaded with GTP or GTP analog, with the switch I and switch II regions highlighted. The PDB entries included are 1IAQ, 2LCF, 2LWI, 3KKM, and 3KKN. The hydroxyl of Thr35 in the switch I region in wild-type Ras coordinates the Mg<sup>2+</sup>ion bound to the GTP, and the T35S mutation is known to disrupt the switch I active conformation. The switch I inactive conformations sampled by the simulations (Supplementary Fig. 3B and C) are broadly consistent with the inactive conformations in T35S structures<sup>116</sup>. h. Conformations of the switch I and II regions in simulations of 24 copies of GTP-bound K-Ras in a crystal lattice (of PDB 3GFT). i. The Ca atom RMSDs of a GMA K-Ras tetramer (w.r.t. the starting structure) in a 10-µs simulation with membrane and in a 10-µs simulation in water solvent. As shown, the membrane helps stabilize the tetramer structure.

#### С А 0.1% EBS 10% FBS 0.5% FBS G12D G120 G12C G12F 015430231262214273 016430E31E82E14EP351 0154398348281478354 0154308318628142935 X-G12C X/D1540 X/D30R · G12D X/D1540 X/D30E DERK ..... 42 44 42 40 60 60 25 90 ======= FRK ----pAK1 AK1 HA ( ) X/K147D X/R135A MW (KDa X/D31R X/E62 X/K147D X/D31R X/E62 В 10% FBS 1% FBS 0.5% FBS X: G12C . X: G12D X. G12D X: G12C X- G12D X: G12C Growth • X • X/D154Q • X/D30R • X/E31R X/E62R Relative in in it. X/K1470 8 24 30 36 42 Hours 18, 24, 30, 36, 42, 48 Hours 18 24 30 36 42 48 Hours 8 24 30 36 42 48 Hours 18 24 30 36 42 Hours <sup>8 24 30</sup> Hours X X/D154Q X/D30R X X/D154Q X/D30R X/D154Q X/D30E X X/D154Q X/D30R X X/D154Q X/D30E X X/D154Q X/D30R X/E31R X/E62R X/K147D X/R135A X/R135A X/R135A X/R135A X/R135A X/R135A 0.1% FBS D Е 10% FB 0.5% EB9 X: G120 X: G12E pERK FRK 🚍 = 42 nAKT -60 AKT 60 HA -- 25 K-Ras 20 HSP90 - - - - -- 90 X/Q12 MW (KDa) 0.5% FBS 10% FBS F X X/K88D X/Q129L X/R149D X: G12C X: G12D X: G12C X: G12D 4 3 growth S Relative 5 \*\*\*\*\*\*\*\*\* 0+ 0 0<del>1</del> 0 0**+** 0 0**+** 0 36 48 48 12 36 48 12 24 Hours 36 48 12 24 Hours 12 24 36 24 X/K88E X/R149D X/01291

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**Extended Data Fig. 4 | Cellular validation of the K-Ras signalosome interfaces.** Panels A-C here refer to the experiments testing mutations at the GMA dimer interface, and panels D-F refer to the experiments testing secondary/tertiary K-Ras interfaces. **a**. Representative cell images at the endpoint of the experiment testing mutations at the GMA dimer interface (Fig. 3f); scale bar: 50 µm. **b**. Growth rates of *K-Ras<sup>lox</sup>/K-RAS<sup>MUT</sup>* cells expressing the indicated mutations in cis with either G12C or G12D mutations in 10%, 1%, and 0.5% FBS medium, represented by the confluence value assessed by IncuCyte. Representative pictures at the endpoint are shown in the bottom panels (scale bar: 50 µm). Data are shown as mean +/- standard deviation (n = 3 biologically independent experiments). **c**. Phosphorylation of ERK and AKT in *K-Ras<sup>lox</sup>/K-RAS<sup>MUT</sup>* cells expressing the indicated mutations in cis with either G12C or G12D mutations. Cells were lysed after 48 hours incubation in 0.5% or 10% FBS, as indicated, and analyzed by Western blot. These results are representative of three independent experiments with similar results. **d**. Representative cell images at the endpoint of the expressing the indicated and analyzed by Western blot. These results are representative of three independent experiments with similar results. **d**. Representative cell images at the endpoint of FBS as indicated and analyzed by Western blot. These results are representative of three independent experiments with similar results. **d**. Representative cell images at the endpoint of the experising mutations at the secondary and tertiary K-Ras/K-Ras interfaces (Fig. 6b); scale bar: 50 µm. **e**. Phosphorylation of ERK and AKT in *K-Ras<sup>lox</sup>/K-RAS<sup>MUT</sup>* cells expressing the indicated mutations in cis with either G12C or G12D mutations. Cells were lysed after 48 hours incubation in 0.5% or 10% FBS as indicated and analyzed by Western blot. These results are representative of three independent experiments with similar results. **f**. Growth rates of *K-Ra* 

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Extended Data Fig. 5 | See next page for caption.

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Extended Data Fig. 5 | FRET and BRET validation of K-Ras signalosome interfaces. a. CFP emission of wild-type K-Ras and various K-Ras mutants to test the K-Ras/K-Ras GMA dimer. HEK293T cells were co-transfected with CFP-K-Ras and YFP-K-Ras, serum starved, and stimulated with 10% FBS. Each K-Ras construct is labeled under its respective plot. Assays were repeated three times. Error bars represent mean ± S.E.M. WT: n = 42; D154Q: n = 33; G12C/D154Q: n = 35; G12D/D154Q: n = 20; D30R: n = 32; G12C/D30R: n = 30; G12D/D30R: n = 28; E31R: n = 42; G12C/E31R: n = 26; G12D/E31R: n = 28; E31R: n = 42; G12C/E31R: n = 42; G12D/E31R: n = 42; G12D/E3 E62R: n = 33; G12C/E62R: n = 32; G12D/E62R: n = 22; K147D: n = 36; G12C/K147D: n = 34; G12D/K147D: n = 38; K147D/D154Q: n = 29. (\*\*\*\* denotes P < 0.0001 by two-way ANOVA; ns stands for not significant). b. CFP emission of wild-type K-Ras and T35 and G60 mutants to test K-Ras association. Assays were repeated three times. Error bars represent mean ± S.E.M. WT: n=14; T35A: n=18; T35S: n=11; G60A: n=17. (P value was calculated by two-way ANOVA; ns stands for not significant). c. CFP emission of wild-type K-Ras and T35 and G60 mutants with C-Raf to monitor Ras-Raf binding at 10% FBS or 10 ng mL<sup>-1</sup> EGF. Assays were repeated three times. Error bars represent mean ± S.E.M. T35A (FBS): n = 36; T35A (No FBS): n = 27; T35A (No FBS+EGF): n=21; G60A (FBS): n=55; G60A (No FBS): n=38; G60A (No FBS+EGF): n=37; WT (FBS): n=12; WT (No FBS): n=37; WT (No FBS+EGF): n=11. (\*\*\*\* denotes P < 0.0001 by two-way ANOVA; ns stands for not significant). d. Nucleotide exchange assay of wild-type K-Ras with (orange) and without (gray) SOS1, and R135A without SOS1 (blue). e. CFP emission of wild-type K-Ras and R135A mutants. Assays were repeated three times. Error bars represent mean ± S.E.M. In 10% FBS condition, WT: n = 44; R135A: n = 82; G12C/R135A: n = 50; G12D/R135A: n = 34; D154Q/R135A: n=51; D154Q: n=33. In 0.5% FBS condition, WT: n=25; R135A: n=47; G12C/R135A: n=44; G12D/R135A: n=34; D154Q/R135A: n=42; D154Q: n=26. (\* denotes P < 0.05, \*\* denotes P < 0.01, \*\*\* denotes P < 0.001, \*\*\*\* denotes P < 0.0001 by two-way ANOVA; ns stands for not significant). **f.** BRET signal as an indicator of K-Ras assembly for G12C and G13D mutants, and for the K-Ras construct lacking the membrane-anchoring HVR tail (residues 1-166). Co-transfection of increasing ratios of donor and acceptor plasmids enables discrimination between specific and non-specific (random collision) proteinprotein interactions. G. CFP emission of wild-type K-Ras and K88D, Q129L, and R149D mutants. Assays were repeated three times. Error bars represent mean ± S.E.M. In 0.5% FBS condition, WT: n = 43; K88D: n = 38; Q129L: n = 34; R149D: n = 37; G12C: n = 22; G12C/K88D: n = 29; G12C/Q129L: n = 34; G12C/R149D: n = 23; G12D: n = 37; G12D/K88D: n = 34; G12D/Q129L: n = 44; G12D/R149D: n = 38. In 10% FBS condition, WT: n = 40; K88D: n = 35; Q129L: n = 40; R149D: n = 42; G12C: n = 27; G12C/K88D: n = 41; G12C/Q129L: n = 46; G12C/R149D: n = 57; G12D: n = 39; G12D/K88D: n = 67; G12D/ Q129L: n = 49; G12D/R149D: n = 42. (\* denotes P < 0.05, \*\* denotes P < 0.01, \*\*\* denotes P < 0.001, \*\*\*\* denotes P < 0.001 by two-way ANOVA; ns stands for not significant).

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**Extended Data Fig. 6 | Structural details of secondary and tertiary Ras-Ras and Ras-RBD interactions. a.** Key contacts of the secondary (stacking) Ras-Ras interaction of K-Ras *n* (blue) and K-Ras n + 4 (green). A series of salt bridges form intermittently in MD simulations between the Lys88, Glu91, Glu98, Arg102, and Asp105 residues on the  $\alpha$ 3 helix of K-Ras *n* and the Asp154, Arg161, Lys165, Glu168, and Lys172 residues, respectively, on the  $\alpha$ 5 helix of K-Ras n + 4. **b**. Key contacts at the interface of K-Ras n + 3 (purple) and K-Ras *n* (blue). **c**. Key contacts of the secondary Ras-RBD interaction of C-Raf RBD *n* (yellow) with K-Ras n + 1 (green), which is the GTP acceptor of K-Ras *n* (purple), the primary binding partner of the RBD. The RBD also interacts with the HVR of K-Ras n + 1. Residues 1-53 of C-Raf are structurally unresolved but are likely to make additional interactions at this secondary Ras-RBD interface. **d**. Key contacts of the tertiary Ras-RBD interaction of Raf RBD n + 3 (yellow) and K-Ras *n* (blue). **e**. Unoccupied Switch-II pocket in a crystal structure of K-Ras (PDB 4LDJ) compared with the Switch-II pocket of K-Ras *n* in the stacking interaction. **f**. Superposition of stacking (n and n + 4) K-Ras proteins from the signalosome model with a crystal dimer of K-Ras (PDB 5UQW); in both cases the dimer interaction is mediated by a  $\beta^2$ - $\beta^3$  turn inserting into a Switch-II pocket. G. Comparison of D154Q/R161E with the WT in K-Ras oligomer simulations. As shown, the double mutation disrupts the D154-K147 salt bridge at the GMA dimer (n/n-1), and as compensation, allows an E161-R102 salt bridge in stacking (n/n + 4) by shifting the relative position of the  $\alpha^3$  and  $\alpha^5$  helices.

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Extended Data Fig. 7 | Structural details of interactions of K-Ras, Gal-3 and the C-Raf CRD. a. Gal-3 n (olive) binds to the HVR fCys185 of Ras n (blue) and to C-Raf RBD n – 1 (orange). The thiodigalactoside inhibitor binding site of Gal-3<sup>117</sup> (PDB 4JC1) is shown in red. **b**. Gal-3 stacking; Gal-3 n (green) and Gal-3 n + 4 (yellow) are stacked in parallel with K-Ras stacking. c. Position of Gal-3 (olive) with respect to K-Ras (blue), in comparison with that of PDE66 (orange) with respect to farnesylated K-Ras<sup>61</sup> (PDB 5TAR), PDE66 (yellow) with respect to farnesylated Rheb<sup>118</sup> (PDB 3T5G), and RhoGDI (cyan) with respect to geranylgeranylated CDC42<sup>19</sup> (PDB 1DOA). The Gal-3 position somewhat resembles the PDE66 complex in the overall pose. The insert shows the simulation-generated Gal-3 binding pose of fCys with the Gal-3 crystal structure (PDB 3ZSM) overlaid for comparison. d. Left: snapshots of the simulation in which fCys enters the Gal-3 (yellow) pocket, with color coding on fCys indicating simulation time; right: the final Gal-3/fCys model superimposed on the Gal-3 structure used to initiate the simulation (PDB 3ZSM), e. Final snapshots from 55 simulations of the three-domain system of the CRD (light orange) tethered to a K-Ras-bound RBD (orange). The snapshots were aligned with respect to the K-Ras protein. Of the 55 simulations, 24 were in solvent (80 µs in aggregate), and the other 31 were with the RBD and the membrane (117 µs in aggregate). The simulation-generated CRD poses are grouped into three clusters: 1) clashing with the GMA K-Ras dimer; 2) clashing with the membrane if applied to the GMA K-Ras dimer on the membrane (Fig. 1a); and 3) near the C-terminal of the  $\alpha$ 5 helix and the  $\beta$ 1,  $\beta$ 2, and  $\beta$ 3 strands of K-Ras. Only cluster 3 is compatible with the helical assembly of K-Ras; cluster 2 is similar to recently reported structures<sup>74-76</sup>. f. Details of the CRD-Ras interface of a CRD pose selected from the third cluster. NMR-identified Ras residues involved in Ras-CRD interaction are shown. The Glu3 residue on the β1 strand of the K-Ras protein interacts stably with a CRD-bound Zn<sup>2+</sup>ion (lower right panel). The Lys157(CRD)/Glu76(K-Ras) distance in simulations is shown to indicate the stability of the Ras-CRD pose (lower left panel). The upper right panel shows the relative positions of K-Ras (blue), CRD (orange), Gal-3 (olive), and GTP (red) in a top view. g. CRD residues identified by an NMR study<sup>51</sup> as part of the K-Ras interface. K-Ras M170, fCys185, and CRD Zn<sup>2+</sup>ions are also shown. h. The surfaces of RBD-bound K-Ras and CRD colored by their electrostatic potential (calculated using PyMOL). A cartoon image is used to show the orientation of the Ras-CRD complex. The CRD-bound Zn<sup>2+</sup>ions are shown in purple. I. A C-Raf CRD at the base tier with RBD and K-Ras. Arg143, Lys144, Lys148, and the two coordinating Zn<sup>2+</sup>ions of the CRD are shown, together with the occupancy map of POPS lipids in contact with the three amino acids shown in red mesh.

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**Extended Data Fig. 8 | Membrane interface of K-Ras, Gal-3, and C-Raf. a**. Positively charged residues of a base-tier K-Ras protein proximal to the membrane in the signalosome model. The inset shows the direct interaction between the HVR lysines of a K-Ras protein at the base tier and the phosphotidylserine lipids. **b**. Positively charged residues and prolines of Gal-3 proximal to the membrane at the base tier. A Val126 residue that has been suggested to be involved in Gal-3 membrane interaction<sup>23</sup> is also shown. **c**. Positively charged residues of the C-Raf RBD and CRD and the two CRD-bound Zn<sup>2+</sup> ions proximal to the membrane.

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**Extended Data Fig. 9 | Additional interactions between Raf, 14-3-3** $\sigma$ , and MEK1. **a**. Superposition of the structural model (gray) of the Raf/MEK/14-3-3 complex with the separately resolved cryo-EM structure (PDB 6Q0J). **b**. Interaction between the Raf kinase domain dimer (green) and 14-3-3 $\sigma$  dimer (cyan). The linker residues pSer338 and pTyr341 are located between the two  $\alpha$ C helices of the kinase dimer, and pSer621 is bound to a 14-3-3 $\sigma$  protein. **c**. Electrostatic potential surface of the C-Raf kinase domain dimer; the region of the two  $\alpha$ C helices, which is where pSer339 and pTyr341 (shown) are located, is highly positively charged. **d**. Interaction of phosphorylated residues pSer338 and pTyr341 with the positively charged residues of the two  $\alpha$ C helices of the C-Raf KD dimer. **e**. Samples of conformations of the C-Raf linker (rainbow colors) in the Ras-Raf signalosome model. The C-Raf KD (gray) is attached to the linker for reference. **f**. Top and side views of an alternative eight-protomer signalosome model, wherein C-Raf *n* and C-Raf *n* + 1 form a dimer, as opposed to the model shown in Fig. 1d and e, wherein C-Raf *n* and C-Raf *n* + 4 form a dimer. The color scheme and protein representation are similar to those in Fig. 1d. The inset shows that in this alternative model a C-Raf linker (linker 2) has to circumvent a Gal-3 protein (Gal-3 2) to engage the other C-Raf linker in a C-Raf dimer. G. An illustration of how the GEF domain of SOS1 can reach the K-Ras 8. The modeling was based on PDB entry 1NVU for the K-Ras/SOS1 pose, and PDB entry 1XD4 for the structures of the PH domain and the PH-GEF linker; the linker orientation with respect to the PH and the GEF domains was manually adjusted and energetically minimized in the modeling. The helical hairpin of SOS1 is shown in yellow.

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Corresponding author(s): Yibing Shan and David E. Shaw

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# **Reporting Summary**

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## Statistics

For	all st	atistical analyses, confirm that the following items are present in the figure legend, table legend, main text, or Methods section.
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	$\boxtimes$	The exact sample size ( $n$ ) for each experimental group/condition, given as a discrete number and unit of measurement
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$\boxtimes$		For Bayesian analysis, information on the choice of priors and Markov chain Monte Carlo settings
$\boxtimes$		For hierarchical and complex designs, identification of the appropriate level for tests and full reporting of outcomes
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		Our web collection on <u>statistics for biologists</u> contains articles on many of the points above.

## Software and code

Policy information	about <u>availability of computer code</u>
Data collection	The molecular dynamics (MD) simulations were performed using the Anton 2 supercomputer. (The simulation code we used is specialized to Anton 2, but codes for performing MD simulation are widely available.) The software used to capture images and fluorescent signal was ZEN software (Version 3.4) from ZEISS. The negative staining TEM was performed using FEI Tecnai G2 Spirit Biotwin transmission electron microscope. Cell growth data were acquired using the IncuCyte Confluence version 1.5 software
Data analysis	We used Visual Molecular Dynamics (Version 1.9.2) developed by the University of Illinois at Urbana-Champaign) and Pymol (Version 2.3, developed by Schrodinger, Inc.) for analyses of the simulation data. FRET data was analyzed using ImageJ (Version 1.52r) and ZEN (version 3.4) software. The negative staining images were processed using RELION 3.0. Cell growth data were analyzed using the IncuCyte Confluence version 1.5 software

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Data

Policy information about availability of data

All manuscripts must include a <u>data availability statement</u>. This statement should provide the following information, where applicable:

- Accession codes, unique identifiers, or web links for publicly available datasets
- A description of any restrictions on data availability
- For clinical datasets or third party data, please ensure that the statement adheres to our policy

Due to the large size of the molecular dynamics trajectories described in this work, they are available (for non-commercial use) by contacting trajectories@deshawresearch.com.

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# Life sciences study design

 All studies must disclose on these points even when the disclosure is negative.

 Sample size
 We used the sample sizes conventional to the in vitro and cell assays we performed.

 Data exclusions
 No data were excluded from the analysis.

 Replication
 All the in vitro and cell assays were replicated at least three times in order to ensure statistically significant power. All attempts at replication were successful.

 Randomization
 Randomization is not applicable to this research, which does not involve patients or animals.

 Blinding
 Blinding is not applicable to this research, which does not entail randomization and does not involve patients or animals.

# Reporting for specific materials, systems and methods

We require information from authors about some types of materials, experimental systems and methods used in many studies. Here, indicate whether each material, system or method listed is relevant to your study. If you are not sure if a list item applies to your research, read the appropriate section before selecting a response.

#### Materials & experimental systems

N/	Δt	ho	de
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n/a	Involved in the study	n/a	Involved in the study
	X Antibodies	$\boxtimes$	ChIP-seq
	Eukaryotic cell lines	$\boxtimes$	Flow cytometry
$\boxtimes$	Palaeontology and archaeology	$\boxtimes$	MRI-based neuroimaging
$\boxtimes$	Animals and other organisms		
$\boxtimes$	Human research participants		
$\boxtimes$	Clinical data		
$\boxtimes$	Dual use research of concern		

## Antibodies

Antibodies used	The antibodies used for western blotting included those against: HA-Tag (6E2) (Cell Signaling Cat#2367, 1:1000), HSP90 (H114) (Santa Cruz Biotech Cat#sc-7947, 1:1000), phosphorylated Akt (Ser473) (Cell Signaling Cat#4060, 1:1000), Akt (Cell Signaling Cat#9272, 1:1000), phosphorylated ERK1/2 (Cell Signaling Cat#4370, 1:1000), ERK1/2 (Cell Signaling Cat#4695, 1:1000), phosphorylated S6 (Ser235/236) (Cell Signaling Cat#4858, 1:1000), S6 ribosomal protein (Cell Signaling Cat#2217, 1:1000), anti-rabbit IgG, HRP-linked secondary antibody (Cell Signaling Cat#7074P2, 1:2000), ECL Sheep anti- Mouse IgG, HRP-linked secondary antibody (GE Healthcare Cat#NA931V, 1:4000), ECL Donkey anti-Rabbit IgG,
	and HRP-linked secondary antibody (GE Healthcare Cat#NA934V, 1:2000).
Validation	Phosphorylated Akt (Ser473) #4060 Application Key:
Validation	Phosphorylated Akt (Ser473) #4060 Application Key:

WB-Western Blot IP-Immunoprecipitation IHC-Immunohistochemistry ChIP-Chromatin Immunoprecipitation IF-Immunofluorescence F-Flow Cytometry E-P-ELISA-Peptide Species Cross-Reactivity Key:

H-Human M-Mouse R-Rat Hm-Hamster Mk-Monkey Vir-Virus Mi-Mink C-Chicken Dm-D. melanogaster X-Xenopus Z-Zebrafish B-Bovine Dg-Dog Pg-Pig Sc-S. cerevisiae Ce-C. elegans Hr-Horse All-All Species Expected

AKT (Cat #9272) Application Key: WB-Western Blot IP-Immunoprecipitation IHC-Immunohistochemistry ChIP-Chromatin Immunoprecipitation IF-Immunofluorescence F-Flow Cytometry E-P-ELISA-Peptide Species Cross-Reactivity Key: H-Human M-Mouse R-Rat Hm-Hamster Mk-Monkey Vir-Virus Mi-Mink C-Chicken Dm-D. melanogaster X-Xenopus Z-Zebrafish B-Bovine Dg-Dog Pg-Pig Sc-S. cerevisiae Ce-C. elegans Hr-Horse All-All Species Expected ERK1/2 (Cell Signaling Cat#4695 Application Key: WB-Western Blot IP-Immunoprecipitation IHC-Immunohistochemistry ChIP-Chromatin Immunoprecipitation IF-Immunofluorescence F-Flow Cytometry E-P-ELISA-Peptide Species Cross-Reactivity Key: H-Human M-Mouse R-Rat Hm-Hamster Mk-Monkey Vir-Virus Mi-Mink C-Chicken Dm-D. melanogaster X-Xenopus Z-Zebrafish B-Bovine Dg-Dog Pg-Pig Sc-S. cerevisiae Ce-C. elegans Hr-Horse All-All Species Expected phosphorylated ERK1/2 Application Key: WB-Western Blot IP-Immunoprecipitation IHC-Immunohistochemistry ChIP-Chromatin Immunoprecipitation IF-Immunofluorescence F-Flow Cytometry E-P-ELISA-Peptide Species Cross-Reactivity Key: H-Human M-Mouse R-Rat Hm-Hamster Mk-Monkey Vir-Virus Mi-Mink C-Chicken Dm-D. melanogaster X-Xenopus Z-Zebrafish B-Bovine Dg-Dog Pg-Pig Sc-S. cerevisiae Ce-C. elegans Hr-Horse All-All Species Expected

## Eukaryotic cell lines

Policy information about <u>cell lines</u>	
Cell line source(s)	K-Raslox K-RASMUT cells generation was described in Ambrogio et al, Cell, 2018. HEK293T cell line was purchased from the ATCC.
Authentication	K-Raslox K-RASMUT cells were generated from primary MEFs (Murine Embryonic Fibroblasts), no authentication is required. HEK293T cells were generated and authenticated in ATCC.
Mycoplasma contamination	All cell lines tested negative for Mycoplasma.
Commonly misidentified lines (See <u>ICLAC</u> register)	No commonly misidentified cell lines were used in the study.