One-way membrane trafficking of SOS in receptor-triggered Ras activation

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SOS is a key activator of the small GTPase Ras. In cells, SOS-Ras signaling is thought to be initiated predominantly by membrane recruitment of SOS via the adaptor Grb2 and balanced by rapidly reversible Grb2-SOS binding kinetics. However, SOS has multiple protein and lipid interactions that provide linkage to the membrane. In reconstituted-membrane experiments, these Grb2-independent interactions were sufficient to retain human SOS on the membrane for many minutes, during which a single SOS molecule could processively activate thousands of Ras molecules. These observations raised questions concerning how receptors maintain control of SOS in cells and how membrane-recruited SOS is ultimately released. We addressed these questions in quantitative assays of reconstituted SOS-deficient chicken B-cell signaling systems combined with single-molecule measurements in supported membranes. These studies revealed an essentially one-way trafficking process in which membrane-recruited SOS remains trapped on the membrane and continuously activates Ras until being actively removed via endocytosis.

Ras is a membrane-anchored small GTPase that plays a central role in many signaling pathways. Ras can exist in an inactive (GDP-bound) or active (GTP-bound) state. Ras activation is mediated by a variety of Ras guanine-nucleotide-exchange factors (RasGEFs) that catalyze the exchange of Ras-bound nucleotide with cytoplasmic GTP. This process is opposed by Ras-GTPase-activating proteins (RasGAPs) that enhance the intrinsic GTPase activity of Ras and thus promote Ras deactivation. Ras activation must be tightly regulated; aberrant activation of Ras is responsible for many human cancers.

Son of Sevenless (SOS) is a widely distributed RasGEF whose full activation through an allosteric mechanism results in large (i.e., bimodal) patterns of receptor-induced Ras kinase signaling. The activation of Ras by SOS is critical for diverse processes such as cell growth, T-cell activation and development, early B-cell development, embryogenesis, and differentiation of embryonic stem cells.

Receptor-triggered activation of SOS is a multilayered process involving membrane recruitment, release of autoinhibition, and allosteric modulation by Ras. The initial membrane recruitment of SOS is thought to occur via association of PxxP motifs in the C-terminal proline-rich (PR) domain with Grb2, which in turn binds phosphotyrosine motifs on activated receptors or transmembrane adaptor proteins. SOS additionally contains a series of N-terminal domains with Dbl homology (DH) and pleckstrin homology (PH) as well as a histone-fold (HF) domain, which autoinhibits SOS activity when assayed in solution. On membranes, this autoinhibition is released through interactions with various membrane lipids (reviewed in ref. 9). Full activation of SOS is contingent on binding of Ras to an allosteric pocket situated at the rim of the REM and CDC25 domains. The REM and CDC25 domains in SOS1 together form the catalytic core, which we denote SOScat herein (Fig. 1a). Mutations in SOS1 that perturb these regulatory functions result in altered signaling behavior and have been implicated in developmental disorders such as Noonan, Costello, and CFC syndromes. SOS2 has a very similar domain makeup but appears to be somewhat redundant with SOS1 in cells; in this study, we focused solely on SOS1.

Historically, SOS activation has been rationalized in terms of a simple membrane-recruitment model based on substrate accessibility (Fig. 1b). Grb2 binding to activated receptors recruits the SOS–Grb2 complex from the cytosol, thereby positioning SOS in proximity to membrane-anchored Ras and consequently promoting nucleotide exchange. However, the importance of Grb2-mediated membrane recruitment is challenged by observations that truncated SOS constructs lacking the PR domain still localize to the membrane after receptor stimulation and are fully signaling competent or even exhibit increased responsiveness compared with that of the full-length enzyme. Recent work with mouse embryonic stem cells has demonstrated that, beyond Grb2-facilitated membrane recruitment, SOS activity is governed by a combination of weak-to-moderate interactions involving multiple protein and lipid interactions that provide linkage to the membrane.
protein–protein and protein–lipid interactions mediated by the multiple domains of SOS. These studies suggest that the recruitment to membrane integral receptors via Grb2 is an oversimplified model for SOS function (Supplementary Note 1).

We observed that SOS constructs lacking the Grb2-binding PR domain are successfully recruited to reconstituted Ras-functionalized membranes through Ras- and lipid-binding interactions. Additionally, using a micropatterned fluid supported-lipid-bilayer platform in which the catalytic activity of individual SOS molecules can be directly resolved, we found that a single SOS molecule has the capacity to processively activate thousands of Ras proteins during a single membrane residency period (Fig. 1c,d). Such high degrees of processivity and essentially irreversible membrane recruitment in the activation of Ras by SOS have not been captured in earlier mechanistic and computational models of SOS activity, or in synthetic-biology approaches using Grb2-SOS1 fusion proteins.

Does such extreme processivity of SOS occur in cells, and, if so, how is it regulated? To address this question, we mapped the individual contributions of the different domains of SOS1 to membrane association, through a series of single-molecule dwell-time measurements and bulk kinetic observations. These studies used a reconstituted-membrane system in combination with quantitative cell-based signaling assays (details in Supplementary Note 1). Altogether, our results reveal an essentially one-way trafficking process in which membrane-recruited SOS1 remains trapped on the membrane and continuously activates Ras until being actively removed, such as by endocytosis. This mechanism differs substantially from the reversible Grb2-dependent process that has been generally assumed.

Nucleotide-dependent recruitment was preserved regardless of the triggering state of the receptor that initiated the signal; this phenomenon substantially affects the quantitative input–response function for Ras activation by receptor triggering and underscores the importance of strong inhibition of spontaneous SOS activation.

RESULTS

Supported-lipid-bilayer SOS-activation assay

We developed an imaging assay to study the interaction of SOS with Ras on supported lipid bilayers (SLBs; Fig. 2a). In this experimental configuration, we coupled H-Ras (residues 1–181, C118S mutant, henceforth referred to as Ras) to the bilayer at C181 via a maleimide-functionalized lipid (Online Methods), thus yielding permanently bound and laterally mobile Ras that was fully functional with respect to SOS activity (Fig. 2a) (Supplementary Fig. 1a). A calibration curve obtained with fluorescence correlation spectroscopy provided access to the local surface density of Ras via epifluorescence imaging of Ras-bound fluorescent nucleotide labels (GDP- and GTP-BODIPY; Supplementary Fig. 1b and ref. 38). Labeling of SOS with a photostable and bright fluorophore (ATTO 647N) facilitated reliable counting and tracking of individual SOS molecules at the membrane surface by total internal reflection fluorescence microscopy (TIRFM). Control experiments showed that labeling did not perturb the observed activity of SOS (Supplementary Fig. 1c).

In this system, we initiated measurements by flowing purified SOS1 over the Ras-functionalized SLBs in a transient pulse with a defined concentration profile (Fig. 2a). During such a pulse, SOS1 interacts with membrane-bound Ras and, in the absence of free nucleotide in solution, becomes trapped after binding Ras at the catalytic site. This method provided a convenient means of quantifying the probability of SOS1 engaging Ras by directly counting the number of SOS1 molecules remaining at the bilayer after a pulse (Fig. 2b and Online Methods). Chasing with unlabeled nucleotide initiated the exchange reaction and resulted in processive (i.e., sustained) turnover of Ras by the recruited and successfully activated SOS1 molecules (Fig. 2a and Supplementary Fig. 1d). A constant flow during the experiment ensured that dissociated SOS1 was removed from the reaction chamber, thus permitting measurement of the desorption kinetics.

Allosteric activation of SOS via altered membrane recruitment

An important functional aspect of SOS1 in the cellular context is its activation by RasGTP binding to an allosteric site, located between the CDC25 and Ras-exchanger motif (REM) domains in the catalytic core, termed SOSCat (ref. 25). This allosteric activation depends on the nucleotide state of Ras and is thought to enable a RasGTP positive feedback loop operating at the membrane.

Allosteric binding of Ras by SOS also provides an alternate mechanism to recruit SOS to the membrane. Here, we first quantitatively analyzed recruitment by examining the SOSCat module, which contains both the active site and the allosteric Ras-binding pocket but lacks any lipid-binding domains. Labeling of SOS with a photostable and bright fluorophore (ATTO 647N) facilitated reliable counting and tracking of individual SOS molecules at the membrane surface by total internal reflection fluorescence microscopy (TIRFM). Control experiments showed that labeling did not perturb the observed activity of SOS (Supplementary Fig. 1c).

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allosteric Ras-binding pocket, as observed in crystal structures\textsuperscript{23,43}. The PH domain interacts with phosphatidylinositol 4,5-bisphosphate (PIP\textsubscript{2}) lipids\textsuperscript{35,44} and phosphatidic acid\textsuperscript{33}, and the HF domain contains several additional interaction sites for negatively charged lipids\textsuperscript{23,24}. These lipid interactions are generally believed to play a role in the release of autoinhibition, but the underlying mechanisms are unclear.

We observed a pronounced damping effect on initial membrane recruitment of SOS1 after adding the N-terminal domains to SOS\textsuperscript{Cat}. Appendixing the DH-PH unit to the catalytic core (SOS\textsuperscript{DPC}) reduced recruitment to the membrane by approximately threefold. Inclusion of the full N terminus (construct comprising HF-DH-PH-PR domains (SOS\textsuperscript{HDPC})) damped recruitment by \sim 66-fold relative to that of SOS\textsuperscript{Cat} (Fig. 3b). Even in the case of the highly autoinhibited HDPC construct, Ras-specific binding was evident (Supplementary Fig. 2e). These observations clearly demonstrated that a major property of the N terminus is the down-modulation of spontaneous SOS1 activation by hindering its initial recruitment to the membrane, as evidenced by the steric hindrance of the allosteric Ras-binding site observed in structures\textsuperscript{23,43}. Interestingly, a gain-of-function R552G point mutation associated with Noonan syndrome (SOS\textsuperscript{HDPC R552G})\textsuperscript{6}, compared with SOS\textsuperscript{HDPC}, caused a slight relief of such inhibition (Fig. 3b and Supplementary Fig. 3a), thus emphasizing the importance of a tightly regulated membrane recruitment step. As observed for SOS\textsuperscript{Cat} (Fig. 2c,d), the longer constructs also exhibited increased recruitment on bilayers displaying RasGTP (Supplementary Fig. 3a).

Although the N-terminal domains inhibited initial recruitment, SOS\textsuperscript{DPC} and SOS\textsuperscript{HDPC} exhibited extremely long dwell times on Ras-functionalized bilayers (with a mean residency period on the hour scale; Fig. 3c, Supplementary Fig. 3b,c and Online Methods). The N-terminal domains thus mediate two major functions: inhibition of the initial recruitment probability and enhancement of the dwell time in the active membrane-bound state. This anticorrelation between membrane recruitment probability and dwell time gives rise to an interesting dual functionality in which rare activation events are coupled to a potent response (Supplementary Fig. 3d,e).

**Multicomponent analysis of SOS-Ras-ERK signaling**

To establish the effects of intrinsic chemical SOS1 properties—as determined from reconstituted-SLB assays—on cellular SOS1-Ras signaling, we optimized a SOS1- and SOS2-double-deficient (SOS1-2−) DT40 chicken B-cell system that we have previously used to characterize digital SOS1-Ras-MAPK ERK signal transduction after B-cell receptor (BCR) ligation\textsuperscript{8,45,46}. Here, we introduced...
EGFP-tagged variants of human SOS1 (hSOS1) into these cells entirely devoid of endogenous SOS1 and SOS2, left the cells unstimulated or induced BCR ligation, and monitored EGFP-SOS localization by fluorescence microscopy or activation of the ERK kinase by using an antibody to phospho-ERK (pERK) and flow cytometry$^9,46$ (Fig. 4a,b). The latter experimental platform, henceforth denoted the p-FLOW assay (Online Methods), revealed the quantitative magnitudes of Ras-ERK responses at the individual-cell level along with SOS1 expression levels. We depicted 3D representations of the data by mapping the time evolution of pERK after BCR stimulation as a function of SOS1 expression level (Fig. 4c,d). pERK traces corresponding to specific SOS1 levels represent 2D slices through the data (Fig. 4f,g,i,j).

Timely signaling requires SOS$^{Cat}$ flanking domains

Transient transfection of EGFP-tagged full-length human SOS1 (SOSFL) rescued the characteristic BCR-induced pERK patterns in SOS-deficient DT40 cells (Fig. 4b and Supplementary Fig. 4a,b).

Figure 3  The N terminus of SOS suppresses bilayer recruitment while prolonging dwell time in the active membrane-bound state. (a) SOS constructs tested in the stopped-flow SLB assay. All experiments shown were conducted with RasGDP on the bilayer. (b) Recruitment probability of SOS constructs obtained from the stopped-flow assay. Each bar represents the average of data collected for the following number of SLB samples, except for HDPC, for which each bar reflects data from one SLB: SOS$^{Cat}$, n = 4; DPC, n = 4; HDPC, n = 2; HDPC R552G, n = 3. Each sample was imaged in at least 15 different positions. Error bars, s.e.m. (data for SOS$^{Cat}$ are replotted from Supplementary Fig. 3b for comparison). (c) Membrane residence time of SOS constructs obtained from the stopped-flow assay. The mean residency period for each construct was obtained by fitting desorption traces (Supplementary Fig. 3b) from the following number of SLB samples: SOS$^{Cat}$, n = 5; DPC, n = 4; HDPC, n = 2; HDPC R552G, n = 3. Error bars, s.e.m. (data for plots and graphs are available online.

Figure 4  Multiparameter assay of SOS-RAS-ERK pathway activity reveals the functional importance of SOS flanking domains in a cell-signaling context. (a) p-FLOW assay of pERK in transiently transfected SOS1-- DT40 B cells. [SOS], SOS concentration. (b) Multiparameter analysis of the SOS-RAS-ERK pathway in model B cells expressing human SOS1-- CT-terminally fused to an EGFP label. A.U., arbitrary units; int., intensity. (c,d) BCR-induced SOS-RAS-ERK pathway activation as a function of increasing SOS expression level and time after stimulation of BCR for SOS$^{Cat}$-expressing (c) and SOSFL-expressing (d) cells. Arrowheads indicate the time of BCR activation. The pERK level is reported as mean fluorescence intensity (MFI). (e) Comparison of basal pERK level across increasing protein concentrations of SOS$^{Cat}$ and SOSFL. The yellow plane on the cube indicates the subspace of the 3D parameter space of the assay corresponding to the shown traces. (f,g) Comparative plots representing the dynamic change in BCR-induced pERK as a function of stimulation time in cells expressing superphysiological levels (f) and intermediate levels (g) of SOS$^{Cat}$ and SOSFL. (h-j) Ratios of pERK observed in SOS$^{Cat}$- and SOSFL-transfected cells, corresponding to traces in e–g. Red fill indicates increased activity of SOS$^{Cat}$ as compared to SOSFL whereas blue fill highlights decreased relative activity. Data are based on seven independent cell cultures and p-FLOW experiments. Error bars, s.e.m. Source data for plots and graphs are available online.
SOS\textsuperscript{Cat}, lacking the Grb2-binding domain as well as the N-terminal lipid-interacting domains, triggered Ras-ERK signaling patterns that differed substantially from those triggered by SOS\textsuperscript{FL} (Fig. 4c,d and Supplementary Fig. 4c,d). Cells expressing high levels of SOS\textsuperscript{Cat}, compared with cells expressing SOS\textsuperscript{FL}, exhibited more spontaneous activation of ERK in the absence of receptor stimulation (Fig. 4e,h). Even under these conditions, BCR stimulation further increased ERK activation in SOS\textsuperscript{Cat}-containing cells (Fig. 4f,i). Another notable difference was the signal attenuation. Whereas SOS\textsuperscript{FL}-induced pERK signals decreased at later time points after BCR stimulation (10–20 min.), SOS\textsuperscript{Cat} continued to signal in a sustained manner, and SOS\textsuperscript{Cat} outperformed SOS\textsuperscript{FL} (Fig. 4f,i). The sustained signaling from SOS\textsuperscript{Cat} cells suggests that the essentially irreversible membrane anchoring of SOS\textsuperscript{Cat} observed in reconstituted assays may exist in cells as well, but not for SOS\textsuperscript{FL}.

Domains flanking SOS\textsuperscript{Cat} might initially appear to merely dampen signal output. However, selective examination of cells expressing intermediate SOS levels revealed that SOS\textsuperscript{FL} signaled more efficiently than SOS\textsuperscript{Cat} in response to BCR stimulation (Fig. 4g,i). Moreover, this intermediate SOS\textsuperscript{FL} level resulted in rescued pERK responses that were nearly identical to those observed for wild-type DT40 cells, thus suggesting that reconstitution with intermediate hSOS1 levels matches the physiological level expressed in wild-type DT40 cells (Supplementary Fig. 4b). The data revealed that domains flanking SOS\textsuperscript{Cat} have both positive and negative regulatory roles.

**SOS autoinhibition prevents spontaneous activation**

A number of structural and cellular studies have established regulatory mechanisms that affect SOS1 activity, but several proposed mechanisms appear to be contradictory\textsuperscript{22–24,43}. To understand how

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**Figure 5** SOS\textsuperscript{Cat} flanking domains block spontaneous activation in the basal state but promote RAS-ERK signal transduction after receptor stimulation. (a–c) p-FLOW assays showing inhibition of spontaneous ERK activation after addition of SOS\textsuperscript{Cat} flanking domains: DH-PH domain (a), HF domain (b), and PR domain (c) (depicted schematically in domain diagrams at the top). The ratio of pERK MFI for longer to shorter SOS variants is plotted against increasing SOS concentration for unstimulated cells (basal state). (d–f) Time dependence of the pERK MFI ratio after BCR stimulation, plotted for the DH-PH domain (d), HF domain (e), and PR domain (f) (depicted schematically in domain diagrams at the top). (g–i) Comparison of BCR-induced ERK activation in the KR-EE PH-domain mutant (**) and wild-type SOS1 variants, showing disruption of membrane lipid interaction in the KR-EE mutant. Yellow planes on the cubes to the left indicate the corresponding subspace of the 3D parameter space in the p-FLOW assay (c.f. Fig. 4c,d). Data are based on three independent experiments. Error bars, s.e.m. Source data for plots and graphs are available online.
SOS1 restricts spontaneous signaling in cells yet allows for controlled allosteric activation near the membrane interface, we first focused on SOS\textsuperscript{Cat} flanking domains in the basal state (Fig. 5a–c), i.e., in resting cells\textsuperscript{6,7}.

Addition of N-terminal domains to SOS\textsuperscript{Cat} blocked the spontaneous activation of Ras-ERK in cells expressing high levels of SOS (Fig. 5a,b and Supplementary Fig. 5a–d). The inhibitory potential scaled in an incremental manner with the number of domains flanking the catalytic core; i.e., SOS\textsuperscript{DPC} signaling was more restrained than SOS\textsuperscript{Cat} signaling (Fig. 5a), and SOS\textsuperscript{HDPC} was more inhibited than SOS\textsuperscript{DPC} (Fig. 5b). These results corroborate the SLB results in Figure 3b. Structural and biochemical studies on SOS1 demonstrated that the DH domain limits Ras binding at the allosteric pocket, and without removal of DH-mediated autoinhibition and allosteric activation, the catalytic pocket cannot fully accommodate RasGDP or dislodge GDP from Ras\textsuperscript{23,47}. The HF strengthens SOS autoinhibition by blocking allosteric activation and by stabilizing a closed conformation of SOS\textsuperscript{23,47}. These structural findings are consistent with our p-FLOW results for the resting cell state (Fig. 5a–c). Notably, despite considerable effort, it has not been feasible to purify functional full-length SOS1 including the PR domain, thus preventing its examination in our earlier SLB assays\textsuperscript{48}.

The C-terminal PR domain is most noted for its positive regulatory role in connecting SOS to activated receptors via Grb2. Grafting the only the PR domain onto SOS\textsuperscript{Cat} revealed an inhibitory effect of this domain in restricting ligand-independent activation of SOS1 (Fig. 5c); this effect was independent of the autoinhibitory effect of the HF and DH-PH domains. The magnitude of inhibition conferred by the PR domain was comparable to that of the DH-PH domain relative to SOS\textsuperscript{Cat} (Fig. 5a,c), thus demonstrating that the N- and C-terminal domains have similar potency in curbing the activity of the catalytic SOS\textsuperscript{Cat} core in resting cells.

Positive regulation of SOS activity in stimulated cells

Next we investigated SOS1 regulation in BCR-stimulated cells expressing intermediate SOS1-EGFP levels (Fig. 5d–f). It has been reported that autoinhibition by the DH domain can be released by electrostatic interaction of the PH domain with membrane lipids, thus allowing allosteric Ras binding\textsuperscript{15,22,33}. In our p-FLOW assay, we found that the DH-PH domain alone had a purely inhibitory effect relative to that of SOS\textsuperscript{Cat} under conditions of BCR stimulation (Fig. 5d). In contrast with inclusion of the DH-PH, inclusion of the HF domain in SOS\textsuperscript{DPC} resulted in increased signaling output (Fig. 5e). We also observed a positive regulatory role of HF after BCR stimulation for SOS containing the PR domain (Supplementary Fig. 5e–g). These findings are in agreement with the in vitro observation that HF enhances the residence time of membrane-recruited SOS (Fig. 3c).

For the DH-PH, our results from stimulated cells conflicted with the increased dwell time observed in the SLB assays (Fig. 3c). The inhibitory effect of DH-PH was unexpected because PH-lipid interaction has been reported to positively regulate GTP loading of Ras in COS-1 cells and in mouse embryonic-stem-cell differentiation\textsuperscript{15,22,33}. This disparity may arise from the HF truncation counteracting the phospholipid binding of PH in the cell system. To test this possibility, we introduced combined mutation of K456E and R459E (KR-EE mutation) within the PH domain, thereby disrupting the PIP\textsubscript{2}-PH interaction\textsuperscript{15,35}, and compared the BCR-stimulated ERK activation associated with the mutant and wild-type SOS1 variants (Fig. 5g–i). The KR-EE mutation in DPC format had a relatively small effect, resulting in a small decrease in pERK (Fig. 5g). However, the KR-EE mutation in HDPC markedly antagonized SOS1 activation throughout the entire assay duration, thus supporting the requirement of HF in stabilizing membrane-targeted SOS1 through phospholipid-PH interaction\textsuperscript{23} (Fig. 5f). The KR-EE HDPC signals were comparable to those of the shorter wild-type DPC, thereby negating the positive regulatory effect of HF domain (Fig. 5i). These observations collectively indicate that the HF and PH domains, through lipid interactions, cooperatively stabilize active SOS1 at the membrane.

In sum, p-FLOW results (Fig. 5) combined with single-molecule measurements in our SLB assays (Figs. 2 and 3) indicated that the flanking domains on both sides of SOS\textsuperscript{Cat} have evolved the ability to simultaneously dampen SOS activity in the basal
state but enhance SOS activity after receptor stimulation (further discussed in Supplementary Note 3).

Regulation of superprocessive SOS by endocytosis
SOS\textsuperscript{Cat}, SOS\textsuperscript{DPC}, and SOS\textsuperscript{HDPC} are all highly processive in SLB assays and in cellular p-FLOW assays are less sensitive than SOS\textsuperscript{FL} to attenuation at late time points of induced signaling. Interestingly, SOS\textsuperscript{FL} mimics these characteristics of SOS truncation when functionalized with a C-terminally grafted farnesylation signal sequence from H-Ras, which artificially targets SOS1 to the membrane\textsuperscript{48} (Supplementary Fig. 6). Deletion of the Grb2-binding domain of SOS1, its putative primary mode of membrane recruitment, thus produces a molecular and cellular phenotype resembling artificial membrane targeting.

To further investigate membrane recruitment and subsequent trafficking of SOS1, we imaged SOS1-EGFP in living cells by TIRFM and spinning-disc confocal microscopy. For this experiment, we used the hybrid live-cell SLB platform\textsuperscript{49–52} to simulate the native signaling geometry of B cells interacting with antigen-presenting cells (Supplementary Fig. 7a). SOS-deficient DT40 B cells expressing human SOS1-EGFP were spread on SLBs functionalized with antibody that recognizes and activates the BCR\textsuperscript{53}, thereby triggering activation of SOS\textsuperscript{FL} (Online Methods).

B-cell activation from the supported membrane led to formation of BCR microclusters, as observed through TIRFM imaging of a Cy5 label on the antibody (Fig. 6a). SOS\textsuperscript{FL} was efficiently recruited to sites of BCR clusters, whereas SOS\textsuperscript{Cat} did not colocalize with BCR clusters, although it did localize to the membrane, presumably through binding allosteric Ras (Fig. 6a,b). SOS\textsuperscript{HDPC} also did not colocalize with the BCR clusters (Fig. 6b and Supplementary Fig. 7b). Contrasting reports have addressed the roles of signaling complexes and SOS1 function. In our B-cell system devoid of any endogenous SOS expression, chimeric SOS\textsuperscript{HDPC}-SH2, with a single SH2 domain of Grb2 grafted onto SOS\textsuperscript{HDPC}, did not colocalize with sites of BCR microclusters (Fig. 6b and Supplementary Fig. 7b). In contrast, addition of the PR domain to SOS\textsuperscript{Cat} or to SOS\textsuperscript{DPC} enabled SOS1-BCR colocalization (Fig. 6b, Supplementary Fig. 7b and Supplementary Note 4).

Over time, the initially scattered BCR clusters concatenated and moved toward the center of the synapses formed between the B cells and the SLB. Approximately 15–20 min after cell landing, a large central cluster appeared, a phenomenon commonly referred to as ‘BCR cAPPING’ (Fig. 6c and Supplementary Movie 1). SOS\textsuperscript{FL} initially moved with the activated BCR, but at later time points we found that it was depleted from the central BCR cluster (Fig. 6d,e). Thus, SOS\textsuperscript{FL} leaves the plasma membrane at the site of the central BCR cluster, and this occurrence also correlates with attenuation of SOS\textsuperscript{FL}-driven ERK signaling at later time points (Fig. 4g).

Confocal fluorescence microscopy revealed the appearance of punctate SOS structures, which were located inside the cells and were reminiscent of endocytic vesicles (Fig. 7a). Moreover, these vesicle-like structures appeared away from the cell-bilayer contact zone at late time points (~10–30 min after cell landing). Scale bar, 10 μm. Source data for plots and graphs are available online.
We found that the kinetics of SOS1 endocytosis was influenced by the allosteric Ras-binding pocket. A SOS1 mutant impaired in allosteric Ras binding (SOS\(^{EL L887E R688A}\)) exhibited accelerated endocytosis (Fig. 7e). Binding of SOS1 to Ras via its allosteric pocket thus appears to counteract the endocytosis of SOS1.

**DISCUSSION**

Signal propagation from receptors to the Ras pathway is commonly accepted to involve recruitment of SOS from the cytosol to the plasma membrane via the adaptor protein Grb2. In its classical interpretation, the increased membrane localization of SOS is presumed to tip the RasGEF-RasGAP balance at the membrane in favor of Ras activation, thus explaining how signals are relayed downstream. However, several results have challenged this classical model, particularly the recurring observation that SOS-truncation mutants lacking the Grb2-binding PR domain remain signaling competent in cells\(^{29–34}\). More recently, we have shown that SOS stably associates with a lipid-membrane surface by engaging Ras at the allosteric binding pocket. In reconstituted-membrane systems, this mechanism alone (i.e., independently of other mechanisms of SOS membrane anchoring) is sufficient for sustained association of SOS with the membrane, where it can processively activate thousands of Ras molecules\(^{35}\). Strikingly, essentially no dynamic equilibrium is present; membrane recruitment of SOS is quasi-irreversible at signaling-relevant timescales.

Here we demonstrated that the membrane recruitment probability of SOS by allosteric Ras is strongly accelerated by RasGTP relative to RasGDP, thereby explaining how SOS constructs lacking the Grb2-binding PR domain are capable of sensing receptor triggering. In a cellular context, RasGTP levels are primed after receptor activation, for example, because of the activity of RasGRP or other exchange factors that produce RasGTP and facilitate SOS recruitment; this process is fueled by strong positive feedback as the recruited SOS produces increasingly more RasGTP. This ability to respond to receptor stimuli independently of Grb2 is further augmented by the lipid-interacting PH and HF domains, which bind lipidic second messengers such as PIP\(_3\) and phosphatidic acid.

In light of the spontaneous and nearly irreversible activating characteristics of SOS, the question shifts to how receptor-mediated signals maintain control of SOS via Grb2 binding. The literature abounds with apparently conflicting results on this matter. In particular, it has been unclear whether the C-terminal PR domain plays a positive, redundant, or even negative regulatory role in SOS signaling. Our p-FLOW assay, which considers the multifactorial aspects of signal transduction (i.e., expression level, pathway activity, and time after receptor stimulation), revealed that the PR domain performs dual functions in receptor-stimulated cells, acting as either a signal facilitator or a signal terminator, depending on the phase of the signaling process. In addition, the PR domain contributes to inhibition of SOS in the basal state.

From the perspective of receptor-mediated activation of SOS, Grb2 binding by the PR domain clearly increases the rate of activation. Our multiparameter mapping of the activity of the SOS-Ras-ERK cascade, enabled by reconstitution of SOS1 in SOS-deficient B cells, revealed that spontaneous activation of SOS scales with SOS expression level. Essentially, the spontaneous activation of SOS is driven by Le Chatelier’s principle and is simply a probabilistic event that scales with concentration. Under endogenous expression levels, this spontaneous activation must be sufficiently slow as to be inconsequential in the context of background GAP activity, thus requiring the additional boost from receptor-mediated Grb2 recruitment to trigger a productive Ras signal (extended discussion in **Supplementary Note 5**; **Supplementary Fig. 8a–d**). We propose endocytosis as a method of signal attenuation that provides an actively regulated mechanism to remove SOS from the plasma membrane, effectively cutting off access to new Ras molecules. SOS constructs lacking the PR domain are not endocytosed, and they exhibit sustained ERK activation levels (further discussed in **Supplementary Note 6**). Thus, in its natural state, SOS activation follows a one-way trafficking circuit with active removal from the membrane via the PR domain as the shutdown mechanism.

Recently, it has become clear that single–amino acid variants in RasGEFs have a profound biological effect. We established that the EF hands in RasGRP1 play a dual role in keeping this RasGEF in the autoinhibited state while simultaneously allowing for calcium-induced activation\(^{38}\). A single–amino acid variant allele, Rasgrp1\(^{Amaef}\), with a point-mutated EF hand, perturbs both regulatory roles of this domain and leads to autoimmune features in Rasgrp1\(^{Amaef}\) mice\(^{59}\). The structural basis for PR-domain-facilitated autoinhibition and the transition to the activated state of SOS is unknown, because efforts to produce functional full-length SOS1 protein including the PR domain have been unsuccessful to date. Mining public databases, we found several SOS1 variants with point mutations or stop codons in the PR domain, which are linked to Noonan developmental syndrome, hyperplastic syndromes such as hereditary gingival fibromatoses\(^{60}\), and various cancers (**Supplementary Fig. 8e**). It is plausible that subtle point mutations in the PR domain may have substantial biological effects and contribute to human disease.

**METHODS**

Methods and any associated references are available in the online version of the paper.

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**AUTHOR CONTRIBUTIONS**

S.M.C., H.-L.T., and J.E.J. performed experiments and analyzed data. S.A. and M.G.T. assisted with live-cell experiments. J.S.I. purified proteins. K.K.Y. performed COS1 cell experiments under supervision of D.B.-S, J.T.G., I.P.R., S.M.C., H.-L.T., and J.E.J. conceptualized and designed experiments. S.M.C., H.-L.T., J.E.J., J.T.G., and J.E.J. wrote the paper. J.T.G. and J.P.R. supervised the project. All authors discussed and commented on the results.

**COMPETING FINANCIAL INTERESTS**

The authors declare no competing financial interests.

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ONLINE METHODS

Proteins and reagents. H-RasC118S C181 (H-Ras construct containing residues 1–181 with a single cysteine at position C181 used for coupling to the bilayer, termed Ras herein), SOS2Cys–Lys (residues 566–1049 with the following mutations: C386A, C655A, C980S, and E718C), SOS2PM (residues 198–1049), SOS2HP (residues 1–1049), and SOS2HP–EV (residues 1–1049 with RS52G) of human SOS1 were expressed in Escherichia coli and purified as previously described22,38. Lipids were purchased from Avanti. TR-DHPE, BODIPY-GDP, and BODIPY-GTP were purchased from Invitrogen. ATTO 488–labeled guanosine diphosphate (EDDA–GDP–ATTO 488) and EDA–GppNp–ATTO 488 (nonhydrolyzable analog of GTP were purchased from Jena Bioscience. GTP was purchased from Sigma-Aldrich, and GDP was purchased from MP Biomedicals. Biotinylated anti-chicken IgM was purchased from Sigma (SAB3700240), and Cy5-labeled streptavidin was from Life Technologies (43–4316). Validation information for commercial antibodies is available on the manufacturers’ websites.

Protein labeling and benchmarking. SOS constructs were fluorescently labeled by reaction of 1:10 molar ratio of unlabeled protein with ATTO 487/N–maleimide for 2 h at 23 °C. Unreacted fluorophores were removed with PD–10 columns (GE Healthcare). The degree of labeling was determined by UV–vis spectroscopy (NanoDrop 2000, Thermo Scientific), yielding 98% for SOS2Cys–Lys, 119% for SOS2PM, 106% for SOS2HP, and 118% for SOS2HP–EV. SOS2PM, SOS2HP, and SOS2HP–EV contained multiple cysteines, thus explaining why labeling efficiencies exceeded 100%.

Dye labeling can potentially alter protein behavior, and caution is always needed in the interpretation of related results. Here, nucleotide-exchange experiments were conducted to ascertain that labeling did not alter enzyme behavior; comparison of unlabeled and labeled constructs in the stopped-flow assay indicated that labeling had a negligible effect on the in vitro activity of SOS in our system (Supplementary Fig. 1c).

Ras-decorated supported lipid bilayers for in vitro assays. Ras-decorated bilayers were prepared as previously described22,38. Lipids dissolved in chloroform were mixed in a round-bottomed flask. Solvent was evaporated by rotary evaporation (40 °C, 10 min) followed by N2 flow (20 min). Small unilamellar vesicles (SUVs) were formed by rehydration of the dried lipid film in PBS (pH 7.45). The vesicle suspension was extruded 11 times (Avestin Miniextruder, 30-nm-pore-diameter polycarbonate membranes; Millipore). The lipid composition was 3% DOPS, 3% MCT–DOPE, 0.01% TxRed–DHPE, and the remainder Egg–PC.

SLBs were formed by incubation of the SUV suspension for 30 min on cleaned piranha-etched glass slides mounted in flow chambers (FCS2 flow chambers, Biotech). The sample was incubated with casein in PBS (2.5 mg/mL) for 10 min and was then incubated for 2.5 h with Ras in PBS (1 mg/mL). After Ras incubation, unreacted MCC was quenched by treatment with 2–β-mercaptoethanol in PBS (5 mM) for 10 min. A motorized syringe pump (PHD 2000, Harvard Apparatus) was used throughout the sample preparation for liquid injections and washing steps.

For loading of fluorescent nucleotide onto Ras, samples were equilibrated at 4 °C and washed with 3 mL loading buffer (40 mM HEPES and 150 mM NaCl, pH 7.4); the native nucleotide bound to Ras was stripped in a 20-min incubation with EDTA in loading buffer (50 mM EDTA, 40 mM HEPES, and 150 mM NaCl, pH 7.4). This step was immediately followed by overnight incubation of samples with 10 μM fluorescent nucleotide analog in reaction buffer (40 mM HEPES, 100 mM NaCl, and 5 mM MgCl2, pH 7.4). Fluorescent nucleotides used in this study included BODIPY-GDP, BODIPY-GTP, ATTO 488–GDP, and ATTO 488–GppNp. A control experiment in which samples underwent all steps except Ras incubation showed no detectable nonspecific binding of the applied fluorescent nucleotides to the SLB.

Immediately before microscopy, samples were brought to room temperature, and any unbound fluorescent nucleotide was removed by washing with 3 mL reaction buffer (40 mM HEPES, 100 mM NaCl, 5 mM MgCl2, and 1 mM TCEP, pH 7.4) under constant flow. The two-dimensional fluidity of lipids and Ras was confirmed for each sample with fluorescence recovery after photobleaching (FRAP).

Antibody-functionalized supported lipid bilayers for live-cell imaging. Bilayers for live-cell experiments were prepared as described above, with a lipid composition of 5% DOPS, 0.1% biotinyl cap PE, 0.005% TxRed–DHPE, and the remainder Egg–PC. A piranha-etched glass slide (1, Fisher Scientific) mounted in a microscopy chamber (A–7816, Life Technologies) was incubated with SUV suspension (1 mg/mL) for 30 min. The sample was then treated with Cy5-labeled streptavidin (18.8 nM) for 30 min and then was incubated with biotinylated anti-chicken IgM (62 nM; SAB700240, Sigma) for 30 min. Each incubation step was followed by copious washing with PBS.

Stopped-flow supported lipid bilayer assay. Labeled and unlabeled SOS constructs were mixed at the desired ratio (typically 1:20) at a total concentration of 100 nM and flowed over the bilayer as a transient pulse. The number of labeled SOS molecules remaining on the bilayer after the pulse (due to capture by catalytic Ras in the absence of free nucleotide)30,40 was counted at the single-molecule level and used to infer the recruitment probability (Supplementary Note 7). We experimentally confirmed that SOS in our system was indeed stably tethered to the bilayer via Ras in the absence of free nucleotide. For Y64A experiments, SOS engaged the membrane in a transient manner (Fig. 2c), and the extent of binding was inferred from the observed peak binding during the SOS pulse instead of from the plateau (data in Fig. 2d).

The nucleotide-exchange reaction was initiated by providing a continuous flow of nucleotide (120 μM GDP or GTP). SOS desorption and nucleotide-exchange kinetics were quantified at different time points by acquisition of an image of the fluorescent nucleotide on Ras and ten images of the labeled SOS. For each time point, we imaged a different position in the flow chamber to avoid bleaching. The ten images of SOS at each position allowed us to discard immobile SOS in the analysis (i.e., SOS bound to defects in the bilayer). This is a crucial aspect of the experimental design because it avoids bias from sample-to-sample variation in the number of defects as well as possible differential tendencies of various protein constructs to adhere to bilayer defects. By counting membrane-bound SOS through single-molecule tracking, we were able to focus entirely on species that are laterally mobile.

A clear demonstration that the assay probed specific interactions between Ras and SOS came from the observation that all SOS constructs tested exhibited sensitivity to the nucleotide state of Ras with consistently increased recruitment probability as well as a prolonged residency period on membranes displaying RasGTP (Fig. 2d and Supplementary Fig. 3a–c).

For specific comparison of desorption for successfully activated SOS constructs (Fig. 3c), traces were normalized to the SOS count at the membrane observed at the initiation of the nucleotide chase. For SOS2Cys–Lys, we observed a fraction of rapidly desorbing species during the first few seconds of the chase (Fig. 2c). This fast-desorbing fraction did not contribute to processive Ras turnover (Supplementary Fig. 1d), and for the comparisons with other constructs (Fig. 3c and Supplementary Fig. 3b,c), we cropped the first 10 s of the trace.

Maintenance and transfection of DT40 and Jurkat cell lines. Culture maintenance, plasmid transfection, and BCR stimulation of chicken DT40 B cell lines were carried out as previously described46. Jurkat cell culture and transfection techniques were also performed as previously described6. The SOS1−2− DT40 B cells were generated in T. Kurosaki’s laboratory (RIKEN). Both wild-type and SOS1−2− DT40 B cells were gifts from T. Kurosaki. The obtained cell lines were confirmed to be free of mycoplasma contamination. For routine cell functional authentication, surface expression of B-cell receptor (BCR) was confirmed by flow cytometry and by BCR-induced eERK2 measurement similar to the experiment shown in Supplementary Figure 4. Jurkat T cells were obtained from the ACCC and were maintained according to the provided guidelines.

To generate EGFP-tagged hSOS1 variants, EGFP coding sequence (CDS) was PCR-amplified with Xia f- and Not I-flanked primers from pEFPN–N1 plasmid (Clontech). The resulting SOS1–EGFP construct bears a five–amino acid linker (SRGGR) between SOS1 and EGFP CDS. Expression was confirmed by western blotting with anti-EGFP antibody (Cell Signaling, 2956) (Supplementary Fig. 4a).

Live-cell imaging. For live-cell microscopy, 2.5 million cells were exchanged from cell culture medium to 1 mL of serum-free RPMI by pelleting cells through 5-min centrifugation at 500g; this was followed by 30-min incubation in serum-free RPMI at 37 °C. Cells were imaged in pH 7.40, 10 mM HEPES, 68 mM NaCl, 2.5 mM KCl, 0.35 mM Na2HPO4, 3 mM n-glucose, 1 mM CaCl2, 2 mM MgCl2, and 0.1% BSA.
Live-cell imaging was performed with a stage-top incubator and an objective heater (Chamlide TC-A, Quorum Technology). Experiments were initiated by addition of cells to SLBs functionalized with anti-BCR (Sigma, SAB3700240). The bilayer was heated to 37°C before addition of cells. The 488-nm channel was used for SOS-EGFP, and the 640-nm channel was used for BCR-engaged antibody on SLBs. For a few selected cells, TIRF images were acquired every ~1–5 min to follow the kinetics of the signaling reaction. Approximately 30 min after cells were added to the chamber, 488-nm and 640-nm TIRF together with bright field and RICM micrographs were acquired at a number of positions in the microscope chamber.

**Flow cytometry and data analysis.** Jurkat T cells were transiently transfected for 20 h with 10 μg of wild-type or allosteric mutant (W729E) SOSCat-encoding plasmid together with 10 μg of GFP plasmid. The activity of the Ras-ERK pathway was measured by FACS staining of surface CD69 (sCD69, BD Pharmingen, 555531) together with GFP intensity measurements. GFP-positive cells were subgated into nine fractions. The geometric mean fluorescence of CD69 was determined for each fraction.

For quantitative and qualitative assays of the RAS-ERK signal module, intracellular staining of BCR-induced ERK phosphorylation was performed according to established procedures. In brief, cells were stimulated with BCR cross-linking mouse IgM (clone M4, Southern Biotech, 8300-01) for the desired time period. Stimulation was then stopped by addition of 4% paraformaldehyde-PBS, and cells were fixed for 20 min at room temperature. Fixed cells were washed three times with FACS wash buffer (PBS, 1% BSA, and 10 mM EDTA) and subsequently permeabilized with prechilled 90% methanol overnight. Cells were then washed three times with FACS wash buffer and stained for pERK with rabbit antiserum (Cell Signaling, 9101). pERK was visualized by secondary staining with goat anti-rabbit IgG conjugated with APC (Jackson Immunochemicals, 711-136-152).

For FACS acquisition, the microscope body (Nikon Eclipse Ti (HUBC/A), Technical Instruments) equipped with a Nikon Apo TIRF 100× oil immersion objective (1.49 NA). The microscope had a custom-built laser launch with 488-nm, 561-nm, and 633-nm lasers (all from the OBIS product line, Coherent) controlled via a laser control module (OBIS scientific remote). The TIRF setup was operated through the objective mode, and images were collected on an EMCCD (iXon ultra 897, Andor). The microscope was controlled with μManager.

Confocal microscopy was performed on a custom-built spinning-disk confocal system. Briefly, images were captured with a Nikon Apo TIRF 100× oil-immersion objective (1.49 NA) and an EMCCD (Andor iXon3 888), and the microscope was controlled with μManager. The axial slice step size was 0.5 μm.

**Data analysis.** A detailed description of the data analysis procedures relating to imaging experiments can be found in Supplementary Note 7.

**Code availability.** Supplementary Note 7 provides a detailed description of the data analysis procedures that can be implemented in a given coding language.

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