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Systems-level interactions between insulin–EGF networks amplify mitogenic signaling

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Crosstalk mechanisms have not been studied as thoroughly as individual signaling pathways. We exploit experimental and computational approaches to reveal how a concordant interplay between the insulin and epidermal growth factor (EGF) signaling networks can potenti ate mitogenic signaling. In HEK293 cells, insulin is a poor activator of the Ras/ERK (extracellular signal-regulated kinase) cascade, yet it enhances ERK activation by low EGF doses. We find that major crosstalk mechanisms that amplify ERK signaling are localized upstream of Ras and at the Ras/Raf level. Computational modeling unveils how critical network nodes, the adaptor proteins GAB1 and insulin receptor substrate (IRS), Src kinase, and phosphatase SHP2, convert insulin-induced increase in the phosphatidylinositol-3,4,5-triphosphate (PIP3) concentration into enhanced Ras/ERK activity. The model predicts and experiments confirm that insulin-induced amplification of mitogenic signaling is abolished by disrupting PIP3-mediated positive feedback via GAB1 and IRS. We demonstrate that GAB1 behaves as a non-linear amplifier of mitogenic responses and insulin endows EGF signaling with robustness to GAB1 suppression. Our results show the feasibility of using computational models to identify key target combinations and predict complex cellular responses to a mixture of external cues.

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Introduction

Cells respond to a myriad of external cues using a limited number of signaling pathways that convert multiple inputs into diverse cellular decisions. Although individual receptors and pathways have been extensively studied, it is not understood how signaling networks integrate multiple cues. The epidermal growth factor (EGF) receptor (EGFR) and the insulin receptor (IR) belong to the family of receptors with intrinsic tyrosine kinase activity (referred to as receptor tyrosine kinases, RTKs), which regulate pivotal cellular processes, including proliferation, differentiation, cell metabolism, survival, and apoptosis (Schlessinger, 2000; Taniguchi et al., 2006). The main physiological function of insulin signaling is metabolic, involving the control of glucose metabolism and stimulation of protein and lipid syntheses, whereas EGF induces proliferative responses.

The EGFR and IR networks share many downstream components. Under some conditions, EGF can evoke metabolic responses, e.g., GLUT4 translocation (Ishii et al., 1994; Gogg and Smith, 2002), whereas insulin can be mitogenic, especially in cancer cells (Ish-Shalom et al., 1997; Papa et al., 1997). There is evidence that insulin can enhance EGF-stimulated extracellular signal-regulated kinase (ERK) activation, DNA synthesis, and cell proliferation responses (Crouch et al., 2000; Ediger and Toews, 2000; Chong et al., 2004), whereas growth factors, cytokines, and other hormones can negatively regulate insulin signaling (Gual et al., 2003). Yet, despite experimental evidence of crosstalk between insulin and growth factor pathways, it is unknown how combined
EGF and insulin inputs are processed into integrative cellular response. This is at least in part due to the combinatorial complexity of molecular interactions and a variety of feedback and feed-forward loops, whose concerted operation is difficult to comprehend intuitively.

Ligand binding and subsequent autophosphorylation of tyrosine residues on the EGFR and IR triggers mobilization of multiple adaptors, such as Src (src avian sarcoma viral oncogene homolog) homology and collagen domain protein (Shc), growth factor receptor binding protein 2 (Grb2), insulin receptor substrate family members (IRS1–6, GAB1–3), and enzymes that contain characteristic domains recognizing receptor phosphoryrosines, such as phosphatidylinositol 3-kinase (PI3K), soluble tyrosine kinase c-Src, protein tyrosine phosphatases (e.g., SHP2 and PTP1B) and others (White, 1998; Sebastian et al., 2006). Subsequently, EGF and insulin-induced signals propagate through multiple interacting branches, including the mitogen-activated protein kinase (MAPK) cascade downstream of the small membrane-anchored GTPase Ras and the PI3K/akt cell survival pathway.

The flow chart in Figure 1 shows that the same key signal transducers are activated by both EGFR and IR, either directly or indirectly. Importantly, although either receptor can stimulate both the Ras/ERK and PI3K/akt pathways, the major routes of activation are different. IR phosphorylates IRS proteins, which are linked to the activation of Ras/ERK pathway through binding to the Grb2–SOS complex (SOS is a guanine nucleotide exchange factor for Ras), whereas EGFR activates the same pathway either by direct binding of Grb2–SOS or by binding and phosphorylation of Shc, which then recruits the Grb2–SOS complex (White, 1998; Sebastian et al., 2006). Likewise, the PI3K/akt pathway is activated by IR via either direct or IRS-mediated recruitment of PI3K, whereas EGFR-mediated PI3K activation occurs mainly via a more intricate route that involves EGFR- and Src-induced phosphorylation of the Grb2-associated binder 1 (GAB1) (Rodrigues et al., 2000; Gu and Neel, 2003; Kiyatkin et al., 2006).

There are several points of crosstalk between the EGFR and IR signaling networks (Figure 1). Following the initial activation of PI3K, the production of phosphatidylinositol-3,4,5-triphosphate (PIP3) in the plasma membrane leads to the membrane recruitment of GAB1 and IRS proteins through their pleckstrin homology (PH) domains. The membrane-recruited and subsequently tyrosine-phosphorylated GAB1 and IRS influence the Ras/ERK and PI3K/akt pathways in multiple ways. They bind p85, the regulatory subunit of PI3K, and alleviate the intrinsic inhibition of PI3K, which further increases PIP3 production, thereby generating a positive feedback and a crosstalk point between EGF and IR (Ogawa et al., 1998; Gu and Neel, 2003). Both GAB1 and IRS can recruit Grb2–SOS, leading to Ras activation (Myers et al., 1994; Lewitzky et al., 2001; Weng et al., 2001), or can bind the GTPase-activating protein RasGAP, which catalyzes Ras deactivation (Montagner et al., 2005). An important crosstalk point emerges because of the binding of SH2 domain-containing tyrosine protein phosphatase 2 (SHP2) to the phosphorylated GAB1 and IRS proteins, which results in both positive and negative regulations of downstream signaling (Noguchi et al., 1994; Yamauchi et al., 1995; Myers et al., 1998; Yart et al., 2001; Asante-Appiah and Kennedy, 2003). Intriguingly, although tyrosine phosphatase SHP2 negatively regulates IRS, PI3K and PI3K/akt signaling (Noguchi et al., 1994; Asante-Appiah and Kennedy, 2003), it positively influences ERK activity, which is partly explained by dephosphorylation of the specific sites involved in RasGAP binding (Cunnick et al., 2002; Agazie and Hayman, 2003; Montagner et al., 2005; Stoker, 2005). Downstream targets of EGFR and IR, such as ERK, GSK3 (glycogen synthase kinase 3) or mTOR (the mammalian target of rapamycin), feedback and phosphorylate GAB1 and IRS on serine/threonine residues, which disable tyrosine phosphorylation at sites engaging PI3K, Grb2, RasGAP or SHP2 (Paz et al., 1997; Gu and Neel, 2003; Johnston et al., 2003). These negative feedback loops generate additional crosstalk points between EGFR and IR.

Although many mechanisms of EGFR–IR crosstalk are well characterized at the molecular level, this knowledge is insufficient to understand cellular responses to EGF plus insulin at the systems level, owing to the multitude of interpathway interactions and feedback loops. This paper brings together experimental studies of combined EGF and insulin signaling with computational modeling of the interactive EGFR and IR networks. We show that, although in HEK293 cells insulin by itself is a poor activator of ERK, it greatly enhances MAPK pathway activation by physiological EGF concentrations. The computational model elucidates the function of feedback loops and crosstalk nodes in combined EGFR and insulin signaling. We demonstrate that synergistic activation of the mitogenic pathway by EGF plus insulin primarily occurs upstream of and at the Ras/Raf level. This potentiation of Ras/ERK response is initiated by insulin-induced PIP3 increase, which leads to subsequent increases in membrane recruitment of Grb2–SOS and SHP2 by GAB1 and IRS. The computational model unveils that insulin makes the mitogenic EGFR signal more robust toward GAB1 knockdown.
Our results may have important ramifications for the identification of therapeutic targets aimed at eliminating insulin-induced amplification of mitogenic and survival signaling stimulated by low growth factor levels in tumor cells.

Results

Building a computational model of the EGF and insulin signaling networks

We have developed a computational model to describe in quantitative terms how cell stimulation by EGF and insulin is linked to the activation of Ras/ERK and PI3K/AKT pathways. The current model stems from our previously developed EGFR network models that were based on *in vitro* and *in vivo* measurements of signaling kinetics. A number of EGFR signaling model predictions were validated in our own studies (Kholodenko et al., 1999; Moehren et al., 2002; Kiyatkin et al., 2006; Birtwistle et al., 2007) and, in addition, tested by other groups (Schoebel et al., 2002; Hatakeyama et al., 2003; Resat et al., 2003; Blinov et al., 2006). This paper extends our previous models to incorporate IR signaling and regulatory processes involved in EGFR–IR crosstalk. However, we do not create a combinatorially complex *in silico* replica of all distinct biochemical species and interactions, which would be impractical (Borisov et al., 2005; Hlavacek et al., 2006; Birtwistle et al., 2007). Instead, we construct a basic, minimal model of the combined EGFR and IR networks. The goal of this model is to provide an insight into the mechanisms of cellular responses to combined EGF and insulin treatment that can account for our data.

The model involves 78 variables for different molecular species, 111 chemical reactions (processes) and more than 200 parameters. A list of reactions, rate equations and parameter values is given in the Supplementary Table S1, and the model SBML file is provided. For many reactions, the parameter values are quantitatively consistent with the previously published values, whose derivation is fully documented in the papers by Kholodenko et al. (1999), Moehren et al. (2002), and Markevich et al. (2004a, b). For additional processes and parameters that describe multi-step processes as single reactions, Supplementary Table S1 cites the corresponding references or indicates that the parameter value was optimized using a training set of data (see Materials and methods). Below, we describe the major signaling processes that are considered and analyzed by this model.

In the model, signal transduction is initiated by ligand (EGF or/and insulin) binding to their cognate receptors. This causes dimerization and autophosphorylation of EGFR, or an allosteric transition and autophosphorylation of the kinase activation loop of the predimerized IR, which leads to activation of the IR kinase and autophosphorylation of its cytoplasmic domain (De Meyts and Whittaker, 2002; Sebastian et al., 2006). The model considers that phosphorylated EGFR can directly bind Shc, Grb2–SOS, PI3K, protein phosphatase(s) and RasGAP (Jones et al., 2006; Sebastian et al., 2006). The membrane recruitment of cytoplasmic SOS is critical for the initiation of the Ras/ERK pathway by both EGFR and IR (Medema et al., 1993; Aronheim et al., 1994; Kholodenko, 2000). Interestingly, the direct recruitment of the Grb2–SOS complex by EGFR was shown to be a less effective route of Ras activation than the EGFR–Shc–Grb2–SOS mediated pathway owing to the corresponding binding affinities (Ravichandran et al., 1995; Kholodenko et al., 1999). At the membrane, SOS catalyzes the transformation of Ras-GDP into active Ras-GTP, whereas RasGAP catalyzes the reverse process of Ras deactivation.

In the model, phosphorylated IR can directly associate with IRS, PI3K, phosphatase(s), and RasGAP (Staub et al., 1994). The IRS family of major IR docking proteins consists of at least six members, IRS1–6; however, the model considers only a single ‘representative’ IRS protein. Importantly in the model, Src is strongly activated by EGFR and more weakly by IR (Schmelzle et al., 2006). Following initial PI3K stimulation and production of PIP3, IRS and GAB1 bind PIP3, and get phosphorylated by IR or EGFR/Src. Phosphorylated IRS and GAB1 recruit cytoplasmic proteins PI3K, Grb2–SOS, SHP2, and RasGAP to the plasma membrane, which results in additional PIP3 production and both activatory and inhibitory regulations of Ras activity (Myers et al., 1994; Ogawa et al., 1998; Gu and Neel, 2003). PIP3 is converted back to phosphatidylinositol-4,5-diphosphate (PIP2) by PTEN (phosphatase and tensin homologue) (Weng et al., 2001). Although both IRS and GAB1 have multiple tyrosine phosphorylation sites, the minimal model represents them by a single, virtual phosphorylation site (Birtwistle et al., 2007). There are some experimental data supporting this assumption and showing that binding of multiple SH2 domain-containing proteins correlates with the overall phosphorylation levels of GAB1 (Figure 4D and E below) and IRS (Goldstein et al., 2000).

Where appropriate, we describe complex yet sequential multi-step processes as a single, semi-mechanistic step. As these condensed processes are sequential, our simplifications allow the reduced model to retain the original network topology. For instance, the activation of Raf by Ras includes a conformational change in Raf caused by binding to Ras-GTP, followed by the dissociation of 14-3-3 protein, dephosphorylation of inhibitory S259 and phosphorylation of activatory S338 sites (Dhillon et al., 2002). In the model, all these processes are considered as a single partial step of Raf activation. The complete Raf activation requires tyrosine phosphorylation by kinases, which are thought to belong to the Src family kinases (SFK) (Wellbrock et al., 2004). In the model, these kinases are linked to Src activity, which is differently stimulated by EGFR or IR. Likewise, in the absence of evidence for a distributed mechanism of ERK kinase (MEK) activation by Raf (Kolch, 2005), we use a simplified, one-step description, whereas a distributed mechanism of ERK activation by MEK and deactivation by MKP3 is described as a two-step process (Markevich et al., 2004a).

The model also incorporates and analyzes complex feedback circuitry of the EGFR and IR networks. For instance, PI3K-dependent positive feedback circuits in the model involve GAB1–PI3K and IRS–PI3K interactions (Rodrigues et al., 2000; Johnston et al., 2003; Mattoon et al., 2004). Activated ERK inhibits SOS (Dong et al., 1996; Fucini et al., 1999), GAB1 (Lehr et al., 2004) and IRS (De Foa and Roth, 1997) by direct phosphorylation. Activated mTOR mediates multiple modes of feedback, including positive feedback to AKT and negative feedback loops to IRS (Gual et al., 2003; Sarbassov et al., 2005). Although AKT-induced inhibitory phosphorylation of Raf...
(Zimmermann and Moelling, 1999; Wellbrock et al., 2004) is included in the model, we assume this inhibition to be weak in HEK293 cells, as no noticeable MEK or ERK activation was detected experimentally, following inhibition of AKT activity (see Supplementary Figure S4). The current model involves many parameters that have no analogs in our previously published models. We used the experimental data that are shown in Figures 2 and 3 (excluding experiments with PI3K

Figure 2  Dynamics of EGF or insulin-induced Ras-GTP, ERK and AKT activation, and GAB1 phosphorylation. The left panels show the time courses calculated in silico and right panels show the corresponding time courses measured in HEK293 cells stimulated with insulin (Ins, 100 nM) or EGF (0.1, 1 or 20 nM) for the indicated time intervals (min). Active GTP-bound Ras was immunoprecipitated (IP) from total cell lysates (TCL) by the agarose-conjugated Ras-binding domain (RBD) of Raf as described in Materials and methods. Proteins from Ras-IP or TCL were separated on LDS-PAGE (4–12%), transferred to nitrocellulose membrane, and immunoblotted (IB) with anti-Ras (A) or anti-phospho-ERK1/2 (T202/Y204), anti-phospho-AKT (S473) or anti-phospho-GAB1 (Y627) antibodies (B–D), respectively. The signal intensities of phosphorylated ERK1/2, AKT, or GAB1 normalized against the appropriate signal of α-tubulin protein level are expressed in arbitrary units (AU). Data shown are the mean of normalized signal intensities ± s.d. from five independent experiments each performed in triplicates.
inhibitor) as a training data set to obtain reasonable fit between the model simulations and data by manually varying the parameter values (see Supplementary Table S1). However, when parameters were fitted, their upper and lower bounds were in agreement with experimental observations for similar reaction types. In addition, reaction rates were always constrained not to be faster than the diffusion limit.

Dose dependence of responses to EGF and insulin

Steady-state plasma concentrations of EGF are reported to be in the range of 0.05–0.2 nM, which is much lower than the concentration range commonly used in the studies on isolated cells (Jansson et al., 1993; Lemos-Gonzalez et al., 2007). For HEK293 cells, which express endogenous EGFR and IR, we...
measured both EGF and insulin dose dependencies. We found that half-maximal ERK activation is attained at about 0.1–0.2 nM EGF (Supplementary Figure S1A). Half-maximal phosphorylation of IR was reached at about 50–100 nM insulin, which was sufficient to saturate ERK activation (Supplementary Figure S1B). To exclude crosstalk at the receptor level, we measured EGFR activation by insulin (and vice versa), and found that neither EGF transactivated IR, nor insulin transactivated EGFR (Supplementary Figure S2A). Although EGF was reported to induce phosphorylation of IRS proteins in A431 cells that express high levels of EGFR (Fujioka et al., 2001), EGF could not induce significant IRS phosphorylation in HEK293 cells (Supplementary Figure S2B), which is consistent with other studies (Kadowaki et al., 1987). By contrast, insulin treatment resulted in potent phosphorylation of IRS1 at Y612 (Supplementary Figure S2B), which is one of the nine putative PI3K binding sites (White, 1998). Robust phosphorylation of these sites by IR correlates with a stronger activation of the PI3K target AKT by insulin than by EGF, although the basal level of AKT phosphorylation in HEK293 cells is significant (Supplementary Figure S2B). However, even at saturating concentrations, insulin is a much weaker activator of ERK than physiological, low EGF (Supplementary Figure S2B), which is also supported by independent observations (Weng et al., 2001).

**Figure 3 Continued.**

**Signaling dynamics of EGF and insulin-induced responses**

To analyze the signaling dynamics, we measured experimentally and simulated in silico the time course of activation of Ras/ERK and PI3K/AKT pathways stimulated by step changes in the EGF and insulin concentrations. First, cells were stimulated with 100 nM insulin or with several EGF concentrations that ranged from low concentrations of 0.1 nM to saturating concentrations of 20 nM (Figure 2). Both the experimental data and simulations showed that the activation
Insulin potentiates MAPK responses to physiological EGF concentrations

Our experimental and simulation results show that co-stimulation by low, physiological EGF (0.1 nM) and insulin synergistically activates the Ras/ERK pathway (Figure 3A–C, quantitative analysis of the blots shown in Figure 3A–C is presented in Supplementary Figure S6A–C). Figure 3E demonstrates that at 1.5 min (left panel) and 15 min (right panel) following co-stimulation with insulin and EGF, the concentration of active ERK is 2.57 ± 0.44 and 1.4 ± 0.1 times greater, respectively, than the sum of the active ERK levels observed with each ligand (this synergistic increase is statistically significant, P<0.01, n=5). Importantly, the synergy effect is not a consequence of additional activation of EGFR and/or IR by co-stimulation with two ligands (Supplementary Figure S2A). Neither can it be explained by activation of insulin-like growth factor receptor-1 (IGF-1R) by insulin. Although 100 nM insulin weakly activates IGF-1 receptor at 1.5 min (Supplementary Figure S2C), 10 nM insulin, which does not transactivate IGF-1R (Supplementary Figure S2C), induces the comparable synergistic effect when combined with 0.1 or 1 nM EGF (Supplementary Figure S2D). When ERK activation is maximal, the synergy becomes insignificant (P>0.05) for any studied EGF dose (Figure 3C and E). Furthermore, the synergistic activation of ERK is less pronounced at higher EGF (1 nM, Figure 3C), and disappears at saturating EGF (20 nM, Supplementary Figure S7).

One group of crosstalk interactions that facilitate the ERK response is located upstream of Ras. These interactions affect multiple Ras activation and deactivation routes involving the adaptor proteins GAB1 and IRS and the protein phosphatase SHP2. In the model, co-stimulation of cells by EGF and insulin increases the amount of PIP3 produced by PI3K (relative to EGF stimulation alone) and further facilitates the GAB1 membrane recruitment and its subsequent tyrosine phosphorylation (Figure 3D, left and middle panels). An increase in membrane-bound phospho-GAB1 is corroborated by our data and it is more appreciable at physiological EGF levels (Supplementary Figure S8B, Figure 3D, right panel, and Supplementary Figure S8A).

An increase in the phosphorylation level of membrane-bound GAB1 promotes Grb2–SOS binding, increasing the SOS concentration in close proximity to Ras (Kholodenko et al., 1999). The SOS concentration in close proximity to Ras (Kholodenko et al., 1999). At the same time, this gain in phospho-GAB1 also increases the amount of RasGAP (Ras deactivator) and the protein phosphatase SHP2 bound to GAB1. In the model, this phosphatase negatively regulates the level of phosphorylated receptors, IRS, and GAB1; yet it has the pronounced positive effect on Ras activation, as we showed using a specific SHP2 inhibitor, NSC-87877 (Supplementary Figure S9A). This positive effect is related to the formation of the GAB1–SHP2 and IRS–SHP2 complexes in close proximity to the plasma membrane and subsequent dephosphorylation of the docking sites on EGFR, IR, GAB1, and IRS, which are involved in RasGAP binding. Thus, SHP2 has an essential role in inhibiting RasGAP-catalyzed Ras deactivation, only if there is a strong activatory signal to Ras arising from EGF co-stimulation, whereas the IRS–SHP2 complexes cannot effectively activate Ras on their own. Simulations predict that the net result of these insulin plus EGF-induced interactions is an increase in positive signaling and decrease in negative signaling to Ras. This amplifies the Ras-GTP level, which is corroborated by the experimental data (Figure 3A).

The second group of crosstalk interactions occurs downstream of Ras and involves processes with positive and negative effects on ERK responses. In the model, at any given Ras-GTP load, the simultaneous exposure to insulin and EGF increases Raf activity relative to insulin alone, owing to EGF-induced stimulation of SFK, which further activates Raf. At the same time, an increase in AKT activity brought about by simultaneous stimulation with insulin and EGF should negatively influence Raf. Yet, the experimental data (Supplementary Figure S4) and simulations suggest that in HEK293 cells the negative influence of AKT on Raf is negligible in comparison with increased Raf activation, owing to increases in Src and Ras activities. Thus, the net effect of insulin-EGF co-stimulation on MEK and ERK activation is positive (Figure 3B and C).
PI3K inhibition suppresses insulin–EGF synergy in Ras/ERK responses

To test the model predictions on the PIP3/GAB1-mediated signaling routes that result in synergistic Ras/MAPK activation by insulin and EGF, the cells were pretreated with the PI3K inhibitor wortmannin (WT). As expected, 100 nM WT prevented AKT phosphorylation in response to insulin and EGF (Figure 3E). By contrast, tyrosine phosphorylation of EGFR, IR and Shc was not affected by PI3K inhibition (Supplementary Figure S10 and Kiyatkin et al., 2006).

Although the model is trained to describe the experimental time-series data for unperturbed cells, simulations of pharmacological interventions, such as inhibition of network nodes and small interfering RNA (siRNA) experiments (see below), were not fitted to the data. Rather, the model predictions are merely compared with the experimental data. The simulations and data suggest that EGF-induced MEK/ERK activation is inhibited by WT due to the disruption of GAB1–PI3K positive feedback. The model predicts that owing to inhibition of the GAB1 and IRS–SHP2 membrane recruitment, WT suppresses synergistic amplification of Ras-GTP/MEK/ERK responses, which is supported by our experimental data (Figure 3A–C and E). The model simulations suggest that although WT disrupts the EGF–insulin synergy, the maximal activation of MEK and ERK remains significant, following 5-min stimulation with EGF or EGF plus insulin. The data presented in Figure 3E (middle panel) validate the model prediction and also show that insulin is unable to enhance MEK and ERK when PI3K is inhibited (Figure 3E, all panels). The agreement between the model predictions and experimental observations substantiates the computational model.

GAB1 complexes with Grb2, SHP2 and PI3K measured by immunoprecipitation analysis

Quantitative data on the dynamics of protein–protein complexes can further validate the model and provide insight into mechanisms of integrative cellular responses to insulin–EGF co-stimulation. As the data (Figure 3D and Supplementary Figure S8) suggest that GAB1 is one of the key mediators of insulin–EGF crosstalk, we determined GAB1 interactions with other network nodes. Although it is not immediately obvious, the simulations predict that following co-stimulation with insulin and EGF, the amount of GAB1-bound PI3K will increase (compared with EGF stimulation) only on a short time scale (Figure 4A, blue dashed lines versus green dotted lines), whereas the post-peak concentration of GAB1-bound PI3K will decrease owing to dephosphorylation of GAB1 phosphotyrosines by GAB1-bound SHP2 and the competitive recruitment of PI3K by phosphorylated IRS. To test this prediction, the binding of PI3K and SHP2 to GAB1 and of PI3K to IRS4 (the major IRS protein in these cells (Fantin et al., 1998)) was measured by immunoprecipitation analysis following stimulation of HEK293 cells with EGF, insulin and their combination. The data confirm that insulin–EGF co-stimulation increased the PI3K–GAB1 complex concentration at 30 s and 1.5 min, whereas the post-peak levels were decreased compared with EGF only stimulation, in agreement with in silico predictions (Figure 4A, right panel). At 3 min of stimulation with insulin and EGF, the data show robust PI3K–IRS binding response and decreased PI3K–GAB1 binding, compared with EGF alone (Figure 4B). The predicted increase in GAB1-bound Grb2 (Figure 4A, solid versus dash–dotted lines) and SHP2 (Figure 4C) correlates with an increase in the total level of

Figure 4  Association of Grb2, PI3K, or SHP2 with GAB1 in response to EGF, insulin, or their combination. (A) The left and middle panels show simulated dynamics of GAB1-containing complexes (scales on the Y axis are different). The right panel shows quantitation of p85–PI3K in GAB1-IP (see Materials and methods). (B) HEK293 cells were stimulated with 100 nM insulin and/or 0.1, 1, and 20 nM EGF for 1.5 min. GAB1-IP was analyzed for total phosphotyrosines (pY20), Grb2, SHP2, or GAB1 proteins. Representative blots are shown in each panel (n=3).
GAB1 tyrosine phosphorylation (Figures 3D and 4E) and is supported by the data shown in Figure 4D for 1.5-min stimulation (quantitation of the blots shown in Figure 4B, D and E is presented in Supplementary Figure S11A–C).

As WT disrupts GAB1–PI3K positive feedback, the total GAB1 phosphorylation level and the concentrations of GAB1-bound Grb2, SHP2, and PI3K decrease dramatically in WT-treated cells (Figure 4E). We conclude that the loss of insulin–EGF synergy caused by WT arises from the loss of the GAB1-mediated membrane recruitment of signaling molecules. The following section presents direct *in silico* and experimental analyses of GAB1 depletion.

**Computational predictions and experimental validation of the GAB1, SHP2, and Src control of ERK responses to EGF and insulin**

We have recently shown that RNA interference (RNAi)-mediated GAB1 depletion can reduce the peak amplitude and decrease the duration of ERK activation (Kiyatkin *et al.*, 2006). These findings are also supported by independent data (Meng *et al.*, 2005). To get further insight into the mechanisms of crosstalk, we simulated the dynamics of ERK responses to EGF versus EGF plus insulin in cells where the GAB1 protein level was held at different levels with respect to control (Figure 5A).

As expected, GAB1 suppression reduces the phospho-ERK level, and its decline is more pronounced for EGF (Figure 5A, left panel) than for EGF plus insulin (Figure 5A, right panel). To test the model, HEK293 cells were transfected with siRNA against human GAB1, resulting in ~75% reduced GAB1 protein level relative to control (non-targeting siRNA-transfected cells) (Figure 5C, quantitation of blots is shown in Supplementary Figure S11D). The phospho-ERK levels measured at 1.5 min following stimulation (Figure 5C) are consistent with the simulations (Figure 5B, left). These results confirm the *in silico* prediction of the larger influence of GAB1 depletion on EGF- rather than on EGF plus insulin-induced ERK phosphorylation.

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**Figure 5** Effects of GAB1 depletion on ERK activation induced by EGF, insulin, or their combination. (A) Computational analysis of ERK activation kinetics in response to 0.1 nM EGF in the absence (left panel) or presence (right panel) of 100 nM insulin at the indicated levels of GAB1 protein. Time courses are shown for 225 nM (control; black—solid line), 150 nM (red—long dash line), 115 nM (dark yellow—short—long—short dash line), 95 nM (dark pink—long—short—short dash line), 75 nM (green—short dash line), 37.5 nM (blue—dash—dot line), 15.0 nM (cyan—dash—dot—dot line), 1.5 nM (gray—long—short dash line) and 0 nM (dark red—dotted line) GAB1 concentrations. (B) Simulated dependences of phospho-ERK level at 1.8 min (left panel) and maximal phospho-ERK level (right panel) on the GAB1 abundance for cells stimulated with 0.1 nM EGF in the presence (red solid line) or absence (blue dashed line) of 100 nM insulin. (C) HEK293 cells transfected with specific siRNA against GAB1 (+) or non-targeting siRNA (−) were stimulated with 100 nM insulin and/or 1 nM EGF for 1.5 min. Immunoblots were analyzed for phosphorylated ERK1/2 (T202/Y204) or GAB1 (Y627). GAB1 protein levels demonstrate the efficacy of GAB1 suppression. GAPDH was used as a loading control. Representative blot (left panel) and the bar graph of respective numerical values (right panel) are shown.
The computational results unveil that insulin endows the mitogenic EGFR pathway with increased robustness towards GAB1 downregulation. The simulations show that for EGF plus insulin stimulation, the peak level of phospho-ERK decreases only slightly with GAB1 depletion, whereas for EGF-induced ERK activation, the peak level decreases significantly (Figure 5B, right panel). The difference between EGF and insulin plus EGF-induced maximal ERK activity is larger in cells with low GAB1 expression than in cells with high GAB1 expression (Figure 5B, right panel), although the absolute value of the peak ERK response is greater for larger GAB1 abundance (Figure 5A). This dependence of the gain in ERK signaling on the GAB1 abundance resembles the behavior of a non-linear amplifier, in which a change in the input does not produce a proportional change in the output. Moreover, the GAB1-mediated amplification of ERK signaling depends on the particular cellular context.

The model predicts that expression of GAB1 PH domain deletion mutant, which is unable to bind PIP3, affects ERK activation in a similar manner as GAB1 knockdown (Supplementary Figure S12A and Figure 5A, left panel). Likewise, co-stimulation with EGFR and insulin makes ERK activation less sensitive to such GAB1 mutation (Supplementary Figure S12A and Figure 5A, right panel). The effects of mutant GAB1 expression or GAB1 suppression differ from the inhibition of PI3K production by wortmannin (Figure 3C and E). When PI3K is present in the membrane, but GAB1 is depleted or mutated, PI3K facilitates the formation of PI3P–IRS–SHP2 complexes, which partially compensate the lack of PI3P–GAB1–Grb2–SOS and PI3P–GAB1–SHP2 complexes by IRS–SHP2-dependent suppression of RasGAP signaling. The simulations show that when IRS–SHP2 interactions are disrupted, synergistic increase in ERK activity by EGF plus insulin co-stimulation can be observed before, but not after the time when ERK reaches its peak activity (Supplementary Figure S12B). Disruption of GAB1–SHP2 interactions substantially decreases the duration of ERK signaling induced by EGF, while the insulin plus EGF synergy is observed (Supplementary Figure S12C). The effects of other perturbations to the system are presented in Supplementary Figures S12D and S12E. The lack of GAB1 phosphorylation by EGFR leads to phospho-ERK changes that are qualitatively similar to the changes induced by the lack of GAB1–PI3K binding (Supplementary Figure S12D), whereas the effects of IRS depletion are qualitatively similar to the effects of disruption of IRS–SHP2 interactions (Supplementary Figure S12E).

The phospho-ERK levels measured in Src-depleted cells (Supplementary Figure S13A) agree reasonably well with the corresponding model predictions (Supplementary Figure S13C). Neither EGFR nor IR autophosphorylation levels were affected by Src downregulation (data not shown). Moreover, we found that RNAi-mediated suppression of Src in HEK293 cells led to a decrease in GAB1 phosphorylation that was more pronounced for EGF than for insulin. Considering a substantial Src contribution to GAB1 phosphorylation under various conditions (Nishida et al., 1999; Bisotto and Fixman, 2001; Chan et al., 2003; Jin et al., 2005), the model explains this observation by the fact that Src is preferably activated by EGFR and not by IR.

At the same time, the model predicts that the influence of SHP2 activity inhibition on ERK phosphorylation will be pronounced after 5 or more minutes following the onset of stimulation (Supplementary Figure S13D), whereas the experimental data show that this effect is already substantial at 1.5 min after stimulation (Supplementary Figure S13B). This shows the limitations of the minimal model, which should be improved in the future studies.

Discussion

Physiological stimuli never act in isolation, and cells in the body are often exposed to insulin and EGF at the same time. Receptors, adaptors, and enzymes involved in insulin and EGF signaling are relatively well-described and form complex networks with intricate interaction circuitry. Downstream of IR and EGFR, these networks significantly overlap. This paper employs a combined experimental and computational approach to address the question of how multiple crosstalk mechanisms in the IR and EGFR networks operate together in a cell-dependent context.

The dynamics of ERK activation depends on a complex interplay of a multitude of regulatory signals and interactions induced by combined insulin and EGF stimulation. Using only qualitative arguments, it is almost impossible to predict the ERK dynamics that results from multiple non-linear interactions and feedback loops, and a testable computational model helps us provide insights into key causative relationships (Kholodenko, 2006). Previous experimental studies indicated that insulin enhances growth-factor-dependent mitogenesis, but the molecular mechanisms remained obscure, as insulin treatment alone fails to induce mitogenesis in most cell types. Using ERK phosphorylation as a marker for mitogenic signaling, we show that insulin leads to the co-localization of mitogenic signaling intermediates at the plasma membrane, whereas EGF-dependent processes contribute to tyrosine phosphorylation of key intermediates. Insulin thereby indirectly boosts signaling events induced by physiological, low EGF doses, despite insulin itself being unable to directly trigger mitogenic signaling. Our results suggest that the regulation of PI3K signaling by non-mitogenic stimuli might be an important means to control the key quantitative aspects of growth-factor-induced MAPK signaling (e.g., signal amplitude and duration).

Our results suggest that co-stimulation with low, physiological EGF doses and insulin endows the mitogenic EGFR pathway with more sustained ERK signaling. Early observations showed that the duration of ERK activation is crucial for cell fate decisions (Marshall, 1995). In these classical experiments, PC12 cells proliferated after transient ERK activation by EGF, but differentiated after more sustained ERK activation by the nerve growth factor. Subsequent work revealed that the duration of ERK activation is sensed by a network of immediate early genes, including the transcription factor c-Fos (Murphy et al., 2002), while the differential timing of ERK signaling is explained by subtle alterations in signaling circuits, induced by distinct ligands (Kholodenko, 2007).

Crosstalk interactions in receptor-induced Ras-MAPK signaling typically occur upstream or at the level of Raf.
Our computational and experimental results also confirm that the major interactions involved in insulin–EGF crosstalk and responsible for amplification of ERK signaling in HEK293 cells operate at the level of protein adaptors, such as GAB1 and IRS, and at the Ras/Raf level. However, in other cellular systems PI3K-induced crosstalk interactions can occur downstream of Ras/Raf. For instance, PIP₃ generated by PI3K enables Rac activation. Subsequently, Rac activates PAK, which in turn stimulates Raf by S338 phosphorylation and MEK by S298 phosphorylation (Chaudhary et al., 2000; Xiang et al., 2002). An alternative pathway of PIP₃-dependent ERK activation involves phosphoinositide-dependent kinase-1 (PDK1), a target of PI3K, which can activate MEK directly (Sato et al., 2004) and ERK indirectly through the activation of certain PKC isoforms (Myers et al., 1996; Schonwasser et al., 1998; Newton, 2003), although it has been reported that PKCs are not involved in EGF-induced ERK activation (Xu et al., 2002; Robin et al., 2004). However, in HEK293 cells, we found that the inhibition of c-Raf by GW-5074 eliminated MEK and ERK activities (Supplementary Figure S14), which supports the conclusion that major insulin–EGF crosstalk mechanisms to amplify ERK signaling are localized upstream of Ras and at the Ras/Raf level.

The model analyzed interactions that involve five key network nodes, the adaptor proteins GAB1 and IRS, PI3K/PIP₃, soluble tyrosine kinase Src and protein tyrosine phosphatase SHP2, which are all localized upstream of MEK/ERK. Together, these nodes contribute to the amplification of ERK signaling in the context of multiple interactions, which involve insulin and EGFs, Grb2–SOS, RasGAP, as well as positive and negative feedback controls from downstream targets of insulin and EGF. The model explains how the network structures, which involve (i) GAB1-mediated insulin and EGF coincidence detection, (ii) coherent feed-forward loop from EGF to Raf via Src, and (iii) multiple positive (PI3K–PIP₃–GAB1–PI3K) and negative (ERK–GAB1, ERK–SOS, mTOR–IRS) feedback loops contribute to the control of insulin plus EGF signaling (a network motif where an initial input signal (A) induces an intermediate input signal (B), and both the initial and intermediate inputs are needed to generate the final output (C), is referred as a coherent feed-forward loop (Mangan et al., 2003)). The simplified scheme shown in Figure 6 illustrates how insulin enhances EGF-induced mitogenesis through two partially redundant and compensating signaling branches via IRS and GAB1. The goal of our computational model is to predict and explain a large number of diverse patterns of signaling dynamics obtained from our own experiments and

**Figure 6** Mechanisms of insulin–EGF signal integration. (A) Weak ERK activation in response to insulin. IR is a strong activator of PI3K signaling, and insulin stimulation induces pronounced adaptor recruitment to the membrane (GAB1 → PIP₃-GAB1; IRS → PIP₃-IRS). The membrane-associated IRS proteins (but not GAB1) are the preferred substrates of the IR kinase (PIP₃-IRS → PIP₃-IRS). However, phosphorylated IRS proteins do not effectively recruit Ras activators, such as Grb2–SOS, implying that insulin is a weak activator of ERK signaling. (B) ERK activation in response to physiological EGF doses. Active EGFR phosphorylates the membrane-associated GAB1 (PIP₃-GAB1 → PIP₃-pGAB1), which in turn efficiently recruits Grb2–SOS and SHP2, and thereby activates Ras-MAPK signaling (for simplicity, the Shc-dependent pathway of EGF-induced MAPK activation is not shown). However, ERK phosphorylation induced by low doses of EGF may remain suboptimal, as the GAB1 recruitment to the membrane is submaximal and highly transient (GAB1 → PIP₃-GAB1). GAB1 phosphorylation and Raf activation are enhanced by EGF-induced Src kinase activity. (C) ERK activation can increase upon insulin and EGF co-stimulation. Synergistic ERK activation predominantly arises from coincidence detection of insulin and EGF stimuli at the level of GAB1; the GAB1 adaptor is massively recruited to the membrane by IR signaling (GAB1 → PIP₃-GAB1) and subsequently phosphorylated by EGFR and active Src (PIP₃-GAB1 → PIP₃-pGAB1). The relative activities of the processes are illustrated by the boldness of the arrows.
literature data. We applied in silico perturbations to our model and, wherever possible, verified the predicted temporal dynamics in vivo. Our strategy is similar to the so-called pattern-oriented modeling, in which a large number of quantitative and also qualitative patterns help us exclude models that are too simple in structure or too uncertain experimentally (Grimm et al., 2005).

We tested the model by using perturbations, including different ligand concentrations, specific inhibitors, and RNAi-mediated suppression of key protein levels. These perturbations corresponded to single-parameter changes, which represent the greatest challenge for any model, because a multitude of temporal patterns have to be described without changes in the rate constants. Our model generated qualitative predictions of the temporal dynamics for diverse cellular responses. In particular, downregulation of GAB1 and Src expression levels using siRNA, or inhibition of PI3K resulted in a significant decrease in synergistic ERK activation by insulin plus EGF. The simulated ERK dynamics in cells with different GAB1 expression (Figure 5A and B) suggest that insulin significantly increases robustness of ERK activation to GAB1 downregulation. The experimental data for GAB1-depleted HEK293 cells demonstrate a significantly larger decrease in the ERK active fraction for EGF than for EGF plus insulin, supporting the modeling results (Figure SC).

Yet, the systems biology approach, while revealing a critical role of GAB1, also points to the fact that this adaptor is just one of several critical players in a complex set of interactions that generate crosstalk between the EGF and IR networks. Despite the fact that RNAi-mediated knockdown of GAB1 expression significantly decreases ERK signaling (Figure SC), the downregulation of additional critical nodes is required to uncouple and completely suppress insulin and EGF-induced Ras/ERK activation. Overall, the analysis presented here demonstrates the feasibility of using computational models to identify critical combinations of therapeutic targets and predict their effects on complex cellular responses to concurrent external cues.

Materials and methods

Antibodies

Rabbit anti-phospho-MEK (S217/S221, anti-phospho-Shc (Y317), anti-IGF-1 receptor β (all from Cell Signaling)), anti-phospho-IR β (Y1162/1163), anti-GRB2 (C-23), anti-GAB1 (H-198), anti-phospho-GAB1 (Y627), anti-SHPT2 (C-18) (all from Santa Cruz Biotechnology), anti-phospho-EGRF (Y1173), anti-phospho-lrs1 (Y612) (BD Biosciences), anti-PI3K-p85 α, anti-GAB1 (CT) (all from Millipore); mouse anti-phospho-ERK1/2 (T202/185 and Y204/187), anti-phospho-AKT1 (S473) (S87F11) (both from Cell Signaling), anti-α-tubulin (DM1A), anti-Src (GD1) (both from Millipore), anti-Ras (BD Biosciences), anti-glyceraldehyde-3-phosphate dehydrogenase (GAPDH) (6C5) (Chemicon) and anti-phosphoestrogens (pY20) (BioLegend). Secondary horse anti-mouse HRP-linked IgG antibodies were from Cell Signaling, HRP-conjugated ImmunoPure goat anti-rabbit IgG antibodies were from Pierce. HRP-conjugated Protein A was from Millipore.

Other reagents

Human recombinant EGF and IGF-1 were from PeproTech, and insulin was from SIGMA. Wortmannin (WT, PI3K inhibitor) (Cell Signaling), NSC-87877 (SHP1/2 inhibitor), AKT-VIII (AKT 1/2/3 inhibitor) (both from Calbiochem) and GW-5074 (Tocris Biosciences) were dissolved in DMSO and used at indicated concentrations. Annealed, desalted GAB1 and c-Src siGENOME SMARTPool siRNAs and siCONTROL Non- Targeting siRNA Pool were from Dharmaco. siRNA stock solution was prepared according to the manufacturer’s instructions and stored at −80 °C. Other chemical reagents were from Fisher Scientific.

Cell culture and transfection

HEK293 (ATCC No. CRL-1573) were cultured in DMEM/F-12 (GIBCO) supplemented with 10% FBS and penicillin-streptomycin solution (Mediatech). The protocol of RNAi-mediated transient gene silencing using synthetic siRNA was described previously (Kiyatkin et al., 2006).

Cell stimulation and protein extraction

For ligand-response studies, the cells were starved from 12 to 16 h in FBS-free medium and stimulated with the indicated ligand concentrations. Preparation of total cell lysates and subcellular fractionation for signaling analysis was performed as described previously (Kiyatkin et al., 2006).

Protein immunoprecipitation

50 μl of recombinant protein A-Sepharose 4B (Sigma-Aldrich) 50% beads slurry was preincubated with the indicated primary antibody (8 μg) while gently mixing for 2 h at room temperature. The beads were incubated with 0.5μg of total lysate protein for 4 h at 4 °C. After the incubation, the beads were washed twice with ice-cold IP buffer (20 mM HEPES (pH 7.4), 150 mM NaCl, 0.1% Triton-X and 10% glycerol) and once with PBS. The proteins were eluted in NuPAGE LDS Sample Buffer (25% 4 × NuPAGE LDS + 10% 10 × NuPAGE Sample Reducing Agent + 65% PBS), heated for 5 min at 75 °C and spun down.

Ras activation assay

Active (GTP-bound) Ras from 500 μg total cell lysates was captured with 30 μl of Raf-1 Ras binding domains (RBD) bound to glutathione-agarose beads (Millipore) for 3 h at 4 °C. Protein complexes were collected by brief centrifugation and washed three times with ice-cold IP buffer supplemented with 10 mM MgCl2. Ras-GTP was released from agarose beads with NuPAGE LDS sample buffer.

Electrophoresis and immunoblotting procedures were conducted according to the methods described in Aksamitiene et al (2007).

Data evaluation

Chemiluminescence signals from immunoreactive bands were detected on a KODAK Image Station 440CF after treatment with a working solution of SuperSignal West Dura Extended Duration Substrate (Pierce Biotechnology) and quantified using KODAK Digital Science software (Kodak Scientific Imaging Systems). To enable comparison, the capture times and number of frames were set equal for each separately exposed membrane, or the whole membranes were exposed simultaneously. Results were plotted and statistically analyzed using SigmaPlot and Microsoft Excel software.

Computational modeling analysis

Numerical integration was performed using Dsolve and PottersWeel. Dsolve (Goryanin et al., 1999) is based on numerical techniques, developed by Khibnik et al (1993). The program files are freely available together with the model SBML file. Ranges of kinetic parameters were constrained based on the literature data and in vivo measurements of the signaling kinetic. The list of the quantitative parameters is presented in Supplementary Table S1, which documents how these parameters were determined. The parameter values marked ‘fitted’ were varied manually within the upper and lower bounds taken
from the experiment to describe a training data set, which includes the experimental time courses in Figures 2 and 3 obtained for an unperturbed HEK293 system. The simulations of the perturbation results, such as inhibition of PI3K, SHP2 and EGFR, and GAB1, IRS or Src depletion, were not fitted to the data. Rather, the model predictions were merely compared with the experimental data.

**Supplementary information**

Supplementary information is available at the Molecular Systems Biology website (www.nature.com/msb).

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**Conflict of interest**

The authors declare that they have no conflict of interest.

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