Integrated Systems and Technologies

Functional Roles of Multiple Feedback Loops in Extracellular Signal-Regulated Kinase and Wnt Signaling Pathways That Regulate Epithelial-Mesenchymal Transition

Sung-Young Shin¹, Oliver Rath², Armin Zebisch²,⁴, Sang-Mok Choo³, Walter Kolch⁴, and Kwang-Hyun Cho¹

Abstract

Epithelial-mesenchymal transition (EMT) is a key event in the generation of invasive tumor cells. A hallmark of EMT is the repression of E-cadherin expression, which is regulated by various signal transduction pathways including extracellular signal-regulated kinase (ERK) and Wnt. These pathways are highly interconnected via multiple coupled feedback loops (CFL). As the function of such coupled feedback regulations is difficult to analyze experimentally, we used a systems biology approach where computational models were designed to predict biological effects that result from the complex interplay of CFLs. Using epidermal growth factor (EGF) and Wnt as input and E-cadherin transcriptional regulation as output, we established an ordinary differential equation model of the ERK and Wnt signaling network containing six feedback links and used extensive computer simulations to analyze the effects of these feedback links in isolation and different combinations. The results show that the feedbacks can generate a rich dynamic behavior leading to various dose-response patterns and have a decisive role in determining network responses to EGF and Wnt. In particular, we made two important findings: first, that coupled positive feedback loops composed of phosphorylation of Raf kinase inhibitor RKIP by ERK and transcriptional repression of RKIP by Snail have an essential role in causing a switch-like behavior of E-cadherin expression; and second, that RKIP expression inhibits EMT progression by preventing E-cadherin suppression. Taken together, our findings provide us with a system-level understanding of how RKIP can regulate EMT progression and may explain why RKIP is downregulated in so many metastatic cancer cells. Cancer Res; 70(17); 6715–24. ©2010 AACR.

Introduction

Epithelial-mesenchymal transition (EMT) plays an important role in embryonic development, and the occurrence of EMT during tumor progression allows benign (i.e., noninvasive and nonmetastatic) tumor cells to acquire the capacity to infiltrate surrounding tissue and to ultimately metastasize to distant sites (1–3). During this transition, epithelial cells undergo a morphologic change that is appropriate for migration in the extracellular matrix and settlement in an area involved in organ formation or metastasis. EMT is characterized by loss of cell adhesion, repression of E-cadherin, and increased cell mobility. Many oncogenic pathways activated by growth factors such as epidermal growth factor (EGF), tumor growth factor, and hepatocyte growth factor (HGF), Src, Ras, integrin, Wnt/β-catenin, and Notch can induce EMT (4). In particular, extracellular signal-regulated kinase (ERK) and Wnt/β-catenin signaling pathways have been shown to activate the transcriptional repressors Snail and Slug that suppress E-cadherin expression, leading to the development of EMT (4–6).

Polypeptides including EGF and Wnt secreted in many malignant tumor cells have been reported to be involved in EMT, and the corresponding signal transduction pathways form a highly interconnected network through multiple coupled feedback loops (CFL). Table 1 summarizes all possible positive and negative feedback loops suggested from previous experiments (7–24), which comprise CFLs between the EGF and Wnt signaling pathways. The compilation of feedback loops and topologies of the networks considered here is based mainly on studies done in colorectal cancer and, to a smaller extent, in breast and prostate cancer cells. These observations indicate a highly complex regulation through a variety of feedback loop motifs. They further suggest that feedback loops play key roles in the regulation of E-cadherin and EMT. As little is known about the functional roles of the feedback loops that interconnect the ERK and Wnt signaling pathways.
Our focus was to identify the functional roles of such CFLs. However, it is difficult to experimentally analyze the effects of such CFLs or in fact even combinations of simple feedback regulations and to envisage how the behaviors of such a complex nonlinear system can be understood without using a formal in silico approach. In this article, we used computational modeling to predict the behavior of the system and its responses to perturbations of the CFLs. For this purpose, we integrated experimental data related to ERK and Wnt signaling pathways and constructed a map of the protein interactions and regulations. We note, however, that the constructed network might not be conserved in all cancer cells because of their inherent heterogeneity as well as tumor- and cell type–specific differences. This implies that there could be different network topologies depending on tumors, tumor cell types, and heterogeneity between tumor cells that arise from stochastic processes or interactions with different neighboring cells. To address this problem, we considered different possible network topologies and analyzed their respective dynamic characteristics. In particular, we first developed a system-level signaling network model that contains protein interactions and regulations reported in the ERK and Wnt signaling pathways by integrating available experimental results and using an established basic mathematical model of each pathway. As our model was developed mainly on data from colorectal cancer cell lines, it is most likely applicable to this type of cancer. However, it is generic in the sense that it also will apply to other cancers with a similar network topology and protein expression profiles such as breast and prostate cancers (25). Then, we compared different network topologies by introducing individual or combinations of feedbacks through extensive computational simulations. We found that the coupled positive feedback loops (CPFL), formed by ERK phosphorylation of RKIP and transcriptional repression of RKIP expression by Snail, cooperatively cause a switch-like behavior of E-cadherin expression. Furthermore, we revealed that the RKIP expression inhibits the EMT progression by counteracting the suppression of E-cadherin.

### Materials and Methods

#### Mathematical modeling and model reduction

The ordinary differential equations for the mathematical model of ERK and Wnt signaling pathways including their transcriptional regulations are presented in Supplementary Data A, where the reduction procedure of the developed mathematical model is also described. The numerical integration

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**Table 1. All feedback loops between the EGF and Wnt signaling pathways and key effects of these feedback loops suggested from the simulation study**

<table>
<thead>
<tr>
<th>Type</th>
<th>Possible feedback loops</th>
<th>Related tumors</th>
<th>Experimental observation</th>
<th>Key effects of the feedback loops suggested from this study</th>
</tr>
</thead>
<tbody>
<tr>
<td>PFL</td>
<td>β-Catenin/TCF complex → Ras → ERK → GSK3β → β-catenin/TCF complex</td>
<td>Colon and liver cancer</td>
<td>(7–14)</td>
<td>It increases the sensitivity of Snail and E-cadherin expression to EGF stimulation.</td>
</tr>
<tr>
<td>PFL</td>
<td>ERK → GSK3β → PKCδ → ERK</td>
<td>Colon and prostate cancer</td>
<td>(10, 15)</td>
<td>It induces the switch-like behavior of E-cadherin expression in response to EGF stimulation. It increases a large state change of E-cadherin in response to EGF stimulation.</td>
</tr>
<tr>
<td>PFL</td>
<td>ERK → RKIP → MEK → ERK</td>
<td>Colon cancer</td>
<td>(16–18)</td>
<td>It induces the switch-like behavior of E-cadherin expression. It decreases the state transition interval in the state change of E-cadherin.</td>
</tr>
<tr>
<td>PFL</td>
<td>ERK → Snail → RKIP → MEK → ERK</td>
<td>Prostate cancer</td>
<td>(19)</td>
<td>If one of these feedback loops is blocked, the switch-like behavior disappears.</td>
</tr>
<tr>
<td>PFL</td>
<td>ERK → GSK3β → Snail → RKIP → MEK → ERK</td>
<td>Prostate cancer</td>
<td>(20)</td>
<td></td>
</tr>
<tr>
<td>NFL</td>
<td>ERK → Ras → ERK</td>
<td>Skin cancer</td>
<td>(21)</td>
<td>This feedback loop, together with the PFL mediated by RKIP, increases EC_{50} of E-cadherin expression in response to EGF.</td>
</tr>
<tr>
<td>NFL</td>
<td>β-Catenin/TCF complex → Axin → β-catenin/TCF complex</td>
<td>Colon and liver cancer</td>
<td>(22, 23)</td>
<td>It does not have any significant effect on the cellular responses considered here.</td>
</tr>
</tbody>
</table>

**NOTE:** The arrow denotes activation and the blunted arrow denotes inhibition.

**Abbreviations:** NFL, negative feedback loop; PFL, positive feedback loop.
was performed using Matlab R2009a software. The kinetic parameters were obtained or modified from the previous model (18, 26), and those not available from the literature were estimated through iterative simulations such that they are qualitatively well in accord with the experimental evidence. The parameter estimates used in the numerical simulation are summarized in Supplementary Table S2 of Supplementary Data A.

Quantification of the results and identification of dominant feedback combinations using Karnaugh-map analysis

To quantify the simulation results and to characterize the dose-response curves, we introduced the following three output indexes (Fig. 1): STI, SD, and EC50. STI denotes the stimulus range corresponding to the state transition between 80% (RR80) and 20% (RR20) of the E-cadherin response range (RR), respectively. SD indicates the range of E-cadherin expression level between RR20 and RR80. EC50 denotes the range of stimulus over which the expression of E-cadherin decreases by 50%. Using these three output indexes, we quantified the dose-response curves and summarized them in a table describing the relationships between 32 feedback combinations and output indexes (Fig. 1, step 1; see also Supplementary Tables S1–S3 of Supplementary Data D).

To discretize the continuous values of output indexes, we identified the feedback combinations that have a statistically significant effect on output indexes. It should be noted that the 32 values of each output index are observed to follow almost the normal distribution. We let $Z_{SD}$, $Z_{EC50}$, and $Z_{STI}$ be the normalized probability variables of $X_{SD}$, $X_{EC50}$, and $X_{STI}$, respectively, and selected those with $Z \geq Z_0$ for $Z_{SD}$ and $Z_{EC50}$.
where the statistical significance $P(Z > Z_0) = 0.1$ and $Z < -Z_0$ for $Z_{crit}$ where $P(Z < -Z_0) = 0.1$, and then assigned them with the logical value of "1"; the nonsignificant values were assigned with the logical value of "0" (Fig. 1, step 2). Then, we constructed a truth table where each possible feedback combination is represented by a binary string of length five. Each bit of the string represents an individual feedback link (1 if the corresponding link is connected and 0 if not). The three output indexes are also represented by binary strings of length three (Fig. 1, Truth table). By scanning out each line of the truth table, we could enumerate all possible feedback combinations that have a significant effect on each output index. We then constructed a Boolean function by summing up those with the Boolean operator (+). However, as the resulting function can take a complex form, we then further apply Karnaugh-map (K-map) analysis to obtain a simplified form. This enables us to determine the simplest form of the logic function that represents the relationship between the feedback combinations and the corresponding output index (Fig. 1, steps 3 and 4). For instance, the Boolean function representing the relationship between the feedback combinations and SD was expressed as seven feedback combinations as follows: $F_1F_2F_3F_4$, $F_1F_2F_3F_4$, $F_1F_2F_3F_4$, $F_1F_2F_3F_4$, $F_1F_2F_3F_4$, $F_1F_2F_3F_4$, $F_1F_2F_3F_4$, $F_1F_2F_3F_4$, $F_1F_2F_3F_4$ (Fig. 1, bottom). By applying the K-map analysis method, this Boolean function can be simplified as three feedback combinations: $F_1F_2F_3F_4$, $F_1F_2F_3F_4$, and $F_1F_2F_3F_4$. See Supplementary Data C for further details.

Results

Mathematical model

A diagrammatic molecular interaction map. To develop a mathematical model, we assembled the interactions between ERK and Wnt signaling pathways including their transcriptional regulations in the form of a diagrammatic interaction map (Fig. 2). This map organizes the protein interactions and functional regulations and provides a basis for further development of a mathematical model. The map consists of three modules: the ERK and Wnt signaling pathway modules and the gene transcriptional regulation module (see Supplementary Data A for details). The interaction map shows multiple CFLs that are formed by protein interactions and regulations (Fig. 3A). We have investigated the role of these feedback loops in the EMT progression through the regulation of E-cadherin.

Simulation results

Cellular responses to oncogenic stimulation in the ERK and Wnt signaling pathways are greatly affected by the different feedback combinations. We have reconstructed a complex network model of the ERK and Wnt signal transduction pathways by integrating protein interactions and regulations from the literature. To investigate how diverse cellular responses are produced from this signaling network, we considered various network topologies obtained from the combination of all possible protein interactions and regulations that are likely to play essential roles in the ERK and Wnt signaling pathways by forming coupled feedbacks. Such interactions and regulations include Ras stabilization by the β-catenin/T-cell factor (TCF) complex ($F_1$), inhibition of the physical interaction between Raf-1 and mitogen-activated protein/ERK kinase (MEK) by RKIP ($F_3$), inhibition of the SOS/Grb2 complex by active ERK ($F_5$), inhibition of glycogen synthase kinase 3β (GSK3β) by active ERK ($F_4$), inhibition of protein kinase Cδ (PKCδ) by GSK3β ($F_5$), and expression of Axin by the β-catenin/TCF complex ($F_6$; Fig. 3A). By considering the protein level of RKIP, $F_2$ implicitly encompasses the transcriptional regulation of RKIP by Snail. As each of these interactions and regulations ($F_1$–$F_6$) is a key element within a feedback loop, we call it a feedback link. It should be noted that the feedback link denotes protein interaction or regulation but not the feedback loop itself. By considering all possible combinations of such feedback links (referred to as "feedback combinations"), we have 64 ($2^6 = 64$ possible combinations) network topologies in total (note that feedback links can have two possible states: "connected" and "nonconnected"). We have simulated the cellular responses of each network topology to normalize oncogenic stimuli of EGF and Wnt. The activation of ERK; the expression of Snail, Slug, and E-cadherin; and the formation of the β-catenin/TCF complex were considered as outputs. As in the initial stage of simulation we found that the feedback link $F_6$ does not have a significant effect on the change of those responses (data not shown), we excluded it from further analysis. Thus, we have a total of 32 network topologies ($2^5 = 32$ possible combinations; Fig. 3B). To investigate how the outputs of the signaling networks vary depending on a specific network topology, the 32 network topologies were analyzed using principal component analysis (PCA). Here, PCA was carried out with respect to five outputs (the activation of ERK; the expression of Snail, Slug, and E-cadherin; and the formation of the β-catenin/TCF complex), and then the results were projected onto two dimensions for which 96% of the output variances were accounted (Fig. 3B). The 32 network topologies can be clustered into four groups: group A, which has a dominant effect on both the first principal component (PC1) and the second principal component (PC2) both in a positive way, group B on PC1 in a positive way, group D on PC1 in a negative way, and group C on PC2 in a negative way. Each group has a distinct characteristic of feedback link combinations: networks in group A have both the feedback links $F_1$ and $F_4$ but not $F_5$ networks in group B contain $F_1$ but not $F_3$ and $F_5$, networks in group C have $F_2$ and $F_3$ but not $F_1$ and $F_4$, and networks in group D have $F_6$. Although the degree of group separation may depend on the parameter values used in the numerical simulation, these results suggest that different feedback topologies in the network can cause different cellular responses to the same oncogenic stimulation and that some particular feedback combinations have a profound effect on the output of signaling networks. This result also could potentially explain how biological specificity can be encoded by combinatorial linkage of a limited array of components.

The CFLs mediated by RKIP cause a switch-like behavior of E-cadherin expression. E-cadherin plays crucial and direct roles in EMT at developmental stages as well as during the...
progression of malignant tumor cells (i.e., loss of E-cadherin expression is an important mark of EMT; refs. 1–3). To systematically identify feedback combinations that have a significant effect on E-cadherin expression, we simulated the dose-response characteristics of E-cadherin expression for 32 network topologies as three-dimensional surfaces with respect to two oncogenic stimuli (i.e., EGF and Wnt). We found that E-cadherin expression is remarkably affected by different feedback combinations (Supplementary Fig. S1A and B of Supplementary Data C). To further investigate, we considered the dose-response curves of E-cadherin expression to EGF (and Wnt) stimulation for a fixed level of Wnt (and EGF, respectively) and also the dose-response curves for a gradual increase of both EGF and Wnt at the same level. By introducing three output indexes [i.e., state difference (SD), state transition interval (STI), and effective concentration 50 (EC50)], we quantified the dose-response curves and identified the feedback combinations (i.e., combinations of connected and nonconnected feedback links) that have a significant effect on each output index. To this end, we used K-map analysis (27), which is a graphical method widely used in engineering circuit analysis for simplification of logic equations representing the Boolean logic relationship between input and output (see Fig. 1 and Materials and Methods for detailed procedures and Supplementary Data B for detailed K-map analysis).

The feedback combinations that have a significant influence on three output indexes and, among those, the most commonly involved feedback combinations are summarized in Supplementary Table S1 of Supplementary Data C, where all these were investigated by gradually increasing EGF for five fixed levels of Wnt and vice versa. Supplementary Table S1 of Supplementary Data C also shows the feedback combinations that have a significant effect on the output indexes for simultaneous increase of both EGF and Wnt. In particular, the feedback combination $F_3F_4F_5$ was most commonly observed with respect to SD for all levels of Wnt. The logical expression $F_3F_4F_5$ means all possible combinations containing two feedback links $F_4$ and $F_5$ while not containing $F_3$. This suggests that network topologies containing this feedback combination would have a significant effect on SD for a gradual increase of EGF with any fixed level of Wnt. The three feedback combinations $F_1F_1F_2$, $F_1F_2F_3F_4$, and $F_1F_2F_3F_4F_5$ were most commonly observed with respect to STI, and the two feedback combinations $F_1F_2F_3F_4F_5$ and $F_1F_2F_3F_4F_5$ were most commonly observed with respect to EC50 for all levels of Wnt. These results suggest that network topologies containing these feedback combinations would have a
significant effect on STI and EC50, respectively, for gradual increase of EGF at any fixed level of Wnt. However, there was no commonly observed feedback combination for different levels of EGF with respect to the three output indexes. The commonly observed feedback combinations for gradual increase of both EGF and Wnt were similar to those for gradual increase of EGF. We note, however, that the feedback combinations \( F_4F_5 \) and \( F_5 \) significantly affected STI and EC50, respectively, only for the combined stimulation. Furthermore, the network topologies having a significant effect on SD and STI contained the feedback links \( F_4F_5 \) and \( F_5 \), respectively (Fig. 4A and B), and those affecting

\[ \text{Figure 3.} \] Multiple feedback loops connect the ERK and Wnt signaling pathways. A, the ERK and Wnt signaling pathways are highly interconnected with each other through multiple feedback loops. Lines with a bar end denote negative feedback links, whereas lines with an arrow end denote positive feedback links. The feedback link \( F_1 \) represents Ras stabilization by the \( \beta \)-catenin/TCF complex, \( F_2 \) represents the inhibition of MEK by RKIP, \( F_3 \) represents the inhibition of SOS/Grb2 complex by ERK, \( F_4 \) represents the inhibition of GSK3\( \beta \) by ERK, \( F_5 \) represents the inhibition of PKC\( \delta \) by GSK3\( \beta \), and \( F_6 \) represents the expression of Axin by \( \beta \)-catenin/TCF complex. B, PCA of the network topologies that are determined by different feedback combinations. The 32 network topologies were analyzed using PCA with respect to the five outputs (the activation of ERK; the expression of Snail, Slug, and E-cadherin; and the formation of the \( \beta \)-catenin/TCF complex), and then the results were projected onto two dimensions for which 96% of output variances were accounted. Label of each feedback combination is shown in the right of this panel.
EC_{50} contained the feedback links F_2F_3 (Fig. 4C). Those feedback links form feedback loops with nonindexed links (see the red line of Fig. 4). Taken together, this implies that the feedback loops containing F_2 have two roles in regulating E-cadherin expression: one is to increase EC_{50} and the other is to shorten STI. This suggests the hypothesis that CPFLs mediated by RKIP induce a switch-like behavior of E-cadherin for the gradual increase of EGF irrespective of the levels of Wnt because the increase of EC_{50} and decrease of STI in a dose-response curve characterize the switch-like response. To verify the hypothesis and further investigate the functional roles of the multiple CPFLs included in the signaling network, we distinguished the CFL mediated by RKIP into two PFLs: ERK \rightarrow RKIP \rightarrow MEK \rightarrow ERK (PFL1) and ERK \rightarrow Snail \rightarrow RKIP \rightarrow MEK \rightarrow ERK (PFL2). So, we have four PFLs to analyze (Fig. 5). We removed each feedback loop one by one and simulated the resulting model for a gradual increase of EGF. Specifically, PFL1 is removed by removing the RKIP phosphorylation by ERK, PFL2 by removing the RKIP regulation by Snail, PFL3 by removing the PKC\delta inhibition by GSK3\beta, and PFL4 by removing the Ras expression by \beta-catenin/TCF. Deletion of either PFL1 or PFL2 significantly diminished the switch-like behavior of cellular responses such as E-cadherin expression. In addition, the deletion of PFL3 suggests that this feedback loop also plays an important role in causing a switch-like behavior of the Snail and E-cadherin expression curves. On the other hand, when PFL4 was removed, the Snail and E-cadherin expression curves were shifted to the right but the switch-like shapes were well conserved, which suggests that the cross talk with the Wnt signaling pathway increases the sensitivity of Snail and E-cadherin expressions to EGF stimulation. Taken together, these simulation results suggest that multiple CPFLs cooperatively induce a switch-like behavior of the cellular responses.

RKIP regulation of E-cadherin expression. Most colon cancers have constitutively activated mutations in either or often both of ERK and Wnt signaling pathways (28–31), and the expression level of RKIP was found significantly decreased in various metastatic cancer cells (32–36). This suggests that the basal level of RKIP expression influences the dynamics of a signaling network in different ways depending on whether ERK, Wnt, or both pathways are persistently activated. To probe this, we simulated our model for two different initial conditions of RKIP (high or low) and three different combinations of sustained stimulations (EGF = 1 and Wnt = 1, EGF = 1 and Wnt = 0, or EGF = 0 and Wnt = 1). The simulation results showed that higher RKIP levels delayed the suppression of E-cadherin and the activation of ERK when both the EGF and Wnt signaling pathways (EGF = 1 and Wnt = 1) were persistently stimulated (Supplementary Fig. S2 of Supplementary Data C). On the other hand, the suppression of E-cadherin expression at a higher RKIP level was much slower and less pronounced when only the EGF signaling was active (EGF = 1 and Wnt = 0). This suggests that the cross talk between ERK and Wnt signaling pathways might...
have a synergistic role in suppressing the E-cadherin expression. We note, however, that RKIP is not crucial in the regulation of E-cadherin expression when only the Wnt signaling pathway is persistently stimulated (EGF = 0 and Wnt = 1). In summary, we found that the initial level of RKIP and the different combination of constitutively activated mutations in either or both of ERK and Wnt signaling pathways play an important role in shaping the dynamics of ERK activity and determining the E-cadherin expression.

To further investigate the effect of RKIP on cellular responses, we increased the RKIP expression by changing its production rate and simulated the activation of ERK and the expression of Snail, Slug, and E-cadherin when both EGF and Wnt were constitutively activated. The simulation results show that the activation of ERK (Fig. 5C) and the expression of Snail and Slug (Supplementary Fig. S3A and B of Supplementary Data C) are decreased, and E-cadherin expression is upregulated as the level of RKIP increases (Fig. 5D). We found that all these responses show switch-like behaviors. In other words, when RKIP increment is below or above a certain threshold range (3–4 in this case), phospho-ERK, Snail, Slug, and E-cadherin do not change, but they are dramatically induced (Snail and Slug) or decreased (phospho-ERK and E-cadherin) when RKIP levels cross the narrow threshold range. However, when one of the two PFLs (PFL1 and PFL2 in Fig. 5C and D and Supplementary Fig. S3A and B of Supplementary Data C) mediated by RKIP was blocked, the switch-like behaviors disappeared. This can be explained by the two RKIP-mediated PFLs that cooperatively compensate for the increase of RKIP up to the point where RKIP levels are high enough to sequester Raf completely and thereby block MEK activation and the positive feedback. Figure 5C and D (insets) show how RKIP determines the system dynamics (see also the insets of Supplementary Fig. S3 of Supplementary Data C). When RKIP increment is below the threshold range, there was no prominent change for the activation of ERK and the expression of Snail and Slug because the expression of RKIP is suppressed by Snail and the phospho-RKIP level is decreased by the reduced ERK activity.

Figure 5. CPFLs cooperatively induce a switch-like behavior of cellular responses, and RKIP determines ERK activation and E-cadherin suppression. A, Snail expression in response to normalized EGF simulation. B, E-cadherin expression. "Control" denotes the case when all PFLs are connected, and "No PFL" \((j = 1, \ldots, 4)\) indicates the case when PFL\(j\) is removed. Within a certain range of concentration values of RKIP, a switch-like behavior of cellular responses is observed. This switch-like behavior disappears if one of the CPFLs mediated by RKIP is blocked. C, activation profile of ERK. D, expression profile of E-cadherin. The right inset shows the illustration of four feedback loops through which the ERK signaling pathway is regulated through ERK \(\rightarrow\) RKIP \(\rightarrow\) MEK \(\rightarrow\) ERK (PFL1), ERK \(\rightarrow\) Snail \(\rightarrow\) RKIP \(\rightarrow\) MEK \(\rightarrow\) ERK (PFL2), ERK \(\rightarrow\) GSK3β \(\rightarrow\) PKCδ \(\rightarrow\) Ras \(\rightarrow\) ERK (PFL3), and ERK \(\rightarrow\) GSK3β \(\rightarrow\) Destruction complex \(\rightarrow\) β-catenin \(\rightarrow\) Ras \(\rightarrow\) ERK (PFL4).
Discussion

ERK and Wnt signaling pathways implicated in the EMT process contain multiple CFLs that form a highly interconnected network. However, little is known about the functional role of these feedback loops. In this study, we found that the positive feedback in which activated ERK counteracts the inhibition of PKCδ by GSK3β has a significant role in inducing a large state change of E-cadherin in response to EGF stimulation, whereas the positive feedback mediated by RKIP decreases the state transition interval in the state change of E-cadherin (Fig. 4A and B; Table 1). CFLs composed of the RKIP-mediated PFLs and the negative feedback loop where ERK phosphorylates and inhibits the SOS/Grb2 complex increase EC50 in response to the EGF stimulation (Fig. 4C; Table 1). On the other hand, it turns out that the negative feedback loop where the β-catenin/Tcf complex induces Axin does not have any significant role in the regulation of E-cadherin expression.

In a previous study (18), we found that the PFL formed by the phosphorylation of RKIP by ERK induces a switch-like behavior of ERK and MEK activities. In this study, we have further revealed that such switch-like behaviors are cooperatively caused by CFLs through phosphorylation of RKIP by ERK and transcriptional repression of RKIP by Snail (Fig. 5C and D). If one of these feedback loops is blocked, the switch-like behavior becomes weaker or even disappears.

From extensive in silico simulation of the dose-response characteristics to oncogenic stimulation and the K-map analysis method (Fig. 1), we found that particular feedback combinations had a significant effect on E-cadherin regulation. Some of them were commonly observed for gradual increase of EGF with a fixed level of Wnt, whereas none was observed for a gradual increase of Wnt with a fixed level of EGF (Supplementary Table S1 of Supplementary Data C). This suggests that the dose-response of E-cadherin expression to EGF stimulation is mostly not affected by the cross talk with Wnt but that Wnt stimulation is significantly affected by the cross talk with EGF. This result might raise the interesting possibility that the ERK and Wnt signaling pathways play different roles in inducing EMT, which can also be partially supported by other recent experimental results. For instance, both the ERK and Wnt signaling pathways are activated by the growth factors secreted by malignant tumor cells with epithelial phenotype, and they contribute to the EMT process possibly at an initial phase (6). However, in transformed mesenchymal cells, the ERK signaling pathway is found in the nucleus of the mesenchymal-like cells at the invasive front of colorectal tumors, whereas it is cytosolic in the central epithelial areas of the same tumor (41). These changes were attributed to slightly different microenvironments. Our results suggest that combinatorial changes in the network topology could be the molecular substrate for such dramatic influences of the microenvironment. This flexible topology afforded by different feedback link combinations also would enable the facile transition of cells between epithelial and mesenchymal morphologies as observed in tumors. In summary, our analysis shows that feedback loops formed by combination of feedback links have roles not only in shaping dynamic

Several recent studies showed a statistically significant inverse relationship between RKIP expression and metastasis and overall survival both in animal models and in human cancer patients (32–36, 38–40). For instance, Fu and colleagues (33) showed that highly metastatic C4-2B prostate cancer cells lack RKIP expression and that enforced expression of RKIP decreased cell invasiveness in in vitro assays and suppressed the development of lung metastasis when implanted into mouse prostate without affecting the growth of the primary tumor. In human cancer patients, it was shown that the retention of RKIP expression in colorectal tumors correlates with a low risk of metastatic relapse and improved survival (32). Similar results were obtained by Minoo and colleagues (36), who used colon cancer tissue microarrays to show that the loss of cytoplasmic RKIP is associated with distant metastasis, vascular invasion, and worse survival. Thus, although an important role for RKIP in metastasis suppression especially in colon and prostate cancer is now emerging, the underlying mechanisms of how RKIP expression is altered in tumors and how RKIP counteracts the development of metastasis are less well understood. The regulation of RKIP expression seems to involve at least two mechanisms. One is the silencing of the rkip gene promoter by hypermethylation (38), and the other is the repression of the rkip gene promoter by Snail (19). The molecular function of RKIP as inhibitor of the ERK pathway that inhibits MEK phosphorylation by Raf is well characterized (16, 17). Recent data show that ERK can inactivate RKIP as part of a feedback loop and thereby confer nonlinear dynamics on ERK activation (18). However, thus far, this feedback loop only has been analyzed in isolation. The same is true for most other feedback loops described here.

Our analysis of the combinatorial effects of such feedback loops revealed several new findings. One is that RKIP is an important feedback loop that exerts major control over both STI and EC50 and that RKIP levels are crucial for these effects (Fig. 4B and C). Thus, the CFLs where ERK phosphorylates RKIP and Snail transcriptionally represses the expression of RKIP cause a switch-like behavior of E-cadherin expression (Fig. 5D), where the RKIP expression determines the EMT progression by antagonizing the suppression of E-cadherin. Another result is that subtle changes in the combination of feedback links may substantially change the output of the network. This is interesting in light of the observation that β-catenin is found in the nucleus of the mesenchymal-like cells at the invasive front of colorectal tumors, whereas it is cytosolic in the central epithelial areas of the same tumor (41). These changes were attributed to slightly different microenvironments. Our results suggest that combinatorial changes in the network topology could be the molecular substrate for such dramatic influences of the microenvironment. This flexible topology afforded by different feedback link combinations also would enable the facile transition of cells between epithelial and mesenchymal morphologies as observed in tumors.
response kinetics of networks but also in the flexible specification and diversification of responses.

**Disclosure of Potential Conflicts of Interest**

No potential conflicts of interest were disclosed.

**Acknowledgments**

We thank J.R. Kim, D. Shin, and D.S. Kim for their helpful discussions and suggestions.

**References**


**Grant Support**

National Research Foundation of Korea grants funded by the Korea Government, Ministry of Education, Science and Technology: 2009-008064 and 2010-001192. The W. Kolch lab was supported by Science Foundation Ireland grant 06/CE/B129 and Cancer Research UK. A. Zehnbach was supported by an EMBO long-term fellowship.

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Received 04/29/2010; revised 06/23/2010; accepted 07/15/2010; published OnlineFirst 08/24/2010.
Functional Roles of Multiple Feedback Loops in Extracellular Signal-Regulated Kinase and Wnt Signaling Pathways That Regulate Epithelial-Mesenchymal Transition

Sung-Young Shin, Oliver Rath, Armin Zebisch, et al.

Cancer Res  Published OnlineFirst August 24, 2010.

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