The global dynamics of gene regulatory networks are known to show robustness to perturbations in the form of intrinsic and extrinsic noise, as well as mutations of individual genes. One molecular mechanism underlying this robustness has been identified as the action of so-called microRNAs that operate via feedforward loops. We present results of a computational study, using the modeling framework of stochastic Boolean networks, which explores the role that such network motifs play in stabilizing global dynamics. The paper introduces a new measure for the stability of stochastic networks. The results show that certain types of feedforward loops do indeed buffer the network against stochastic effects.

The term *canalization* was coined by the geneticist C. Waddington to describe the theory that embryonal development is buffered against genetic and environmental perturbations. It is only recently that a molecular basis for this phenomenon has been suggested. Recent research has highlighted how the intrinsic stochasticity of gene expression can drive changes in phenotypes. Short

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segments of single-stranded RNA, so-called microRNAs (miRNA), represent an
entirely novel agent of gene regulation discovered relatively recently, and have
been proposed to function as canalizing agents that buffer the effects of such
stochasticity in gene expression. According to this theory, when miRNA expres-
sion is perturbed, stochasticity in gene expression can result in transitions to
distinct cellular phenotypes. As miRNAs bind to gene targets they downreg-
ulate translation of target mRNA into protein. Embedded in several different
types of so-called feedforward loops (FFLs), miRNAs help smooth out noise and
generate canalizing effects in gene regulation by overriding the effect of certain
genes on others.

Much experimental work remains to be done in elucidating this concept, and
recent years have seen an explosive growth of publications in this area. There
have also been a number of computational studies focused on canalization. In
this paper, we carry out a computational study of the ability of the feedforward
loop motif to buffer a gene regulatory network against intrinsic noise. This is
done using stochastic Boolean network models as a computational instantiation
of gene regulatory networks. We introduce a measure on networks that captures
its “distance-to-deterministic” characteristics in terms of the stability of their
attractors. For a given network, we successively introduce feedforward loops
and track the resulting change in dynamics. The results show clearly that the
feedforward loop motif buffers the network phenotype, in terms of stability of
attractors, against perturbations from intrinsic noise.

I. Introduction

The term canalization was coined by the geneticist C. Waddington to describe the theory
that embryonal development is buffered against genetic and environmental perturbations. It
is only recently that a molecular basis for this phenomenon has been suggested. Recent re-
search has highlighted how the intrinsic stochasticity of gene expression can drive changes in
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An understanding of canalization in evolutionary biology is important as a cornerstone of natural selection and the emergence of new phenotypes, as well as for the understanding of diseases such as cancer. Transitions to new phenotypes have been implicated as one of the driving forces of tumorigenesis; and, interestingly, significantly altered expression of miRNAs is a feature of several cancers. Much experimental work remains to be done in elucidating this concept, and recent years have seen an explosive growth of publications in this area. There have also been a number of computational studies focused on canalization. Several papers have studied computational models that capture the evolution of canalization in networks and their ability to support significant mutation without change in the phenotype, while others have studied models of stochastic gene expression as the internal source of noise in regulatory networks. A detailed stochastic model of the ability of miRNAs to buffer gene expression noise in a feedforward loop has been proposed, providing evidence that this type of network motif can in fact play a canalizing role. There is evidence that miRNAs do not regulate their target genes directly; rather they act as post-transcriptional regulators by reducing the amount of mRNA and by repressing mRNA translation, e.g., by binding to the 3'-UTR of a mRNA, which prevents this mRNA from being translated.

In this paper, we carry out a computational study of the ability of the feedforward loop motif to buffer a gene regulatory network against intrinsic noise. This is done using stochastic Boolean network models as a computational instantiation of gene regulatory networks. We introduce a measure on networks that captures their “distance-to-deterministic” characteristics in terms of the stability of their attractors. For a given network, we successively
introduce feedforward loops and track the resulting change in dynamics. The results show clearly that the feedforward loop motif buffers the network phenotype, in terms of stability of attractors, against perturbations from intrinsic noise.

II. Modeling framework

A. Stochastic Discrete Dynamical Systems

In this study, the recently developed framework of stochastic discrete dynamical systems (SDDS)\textsuperscript{20} is used to model gene regulatory networks. This framework is an appropriate set-up to model the effect of intrinsic noise on network dynamics. A stochastic discrete dynamical system in the variables $x_1, \ldots, x_n$, which in this paper represent genes, is defined as a collection of $n$ triplets

$$ F = (f_i, p^\uparrow_i, p^\downarrow_i)_{i=1}^n, $$

where

- $f_i : \{0, 1\}^n \rightarrow \{0, 1\}$ is the update function for $x_i$ for all $i = 1, \ldots, n$,
- $p^\uparrow_i \in (0, 1]$ is the activation propensity,
- $p^\downarrow_i \in (0, 1]$ is the degradation propensity.

The stochasticity originates from the propensity parameters $p^\uparrow_i$ and $p^\downarrow_i$, which should be interpreted as follows: If there would be an activation of $x_i$ at the next time step, i.e., $x_i(t) = 0$, and $f_i(x_1(t), \ldots, x_n(t)) = 1$, then $x_i(t + 1) = 1$ with probability $p^\uparrow_i$. The degradation probability $p^\downarrow_i$ is defined similarly.

All variables are synchronously updated and one time step can be interpreted as the average time needed for transcription and translation of the fastest of the genes considered. The propensity parameters for this fastest gene will be set to 1, and the propensity parameters of genes with slower transcription and translation take proportionately lower values. Thus, this framework can be interpreted as introducing a very general stochastic sequential update scheme, which also allows for a variable not to be updated at all at a given step, a generalization of the usual approach in, e.g.,\textsuperscript{21}. 


B. Quantifying Stochasticity

In a deterministic discrete dynamical system, each initial configuration lies in exactly one basin of attraction. This changes when stochasticity is introduced. Now, from one initial configuration, different attractors may be reached. To measure the degree of stochasticity in particular dynamics, we look at every initial configuration and regard its transition probabilities to the different attractors. If each initial configuration only transitions to one attractor, the dynamics are deterministic, whereas lower maximal transition probabilities to attractors lead to proportionately more stochastic dynamics. In our context, the different attractors may be interpreted as different cellular phenotypes, which makes the connection to phenotypic stability discussed in the introduction\textsuperscript{14}. Thus, this computational project focuses on the stability of attractors under intrinsic noise, as it is affected by the introduction of feedforward loops.

Based on this idea, we can define the degree of stochasticity in the dynamics of an SDDS $F = (f_i, p^↑_i, p^↓_i)_{i=1}^n$. Let $\mathcal{A}(F)$ be the set of all attractors of $F$. Then we can calculate the average maximal transition probability to an attractor, where all $2^n$ state space configurations are considered and weighted equally, as follows:

$$
\mu(F) = \frac{1}{2^n} \sum_{x \in \{0,1\}^n} \left( \max_{A \in \mathcal{A}(F)} \mathbb{P}(x \xrightarrow{F} \cdots \xrightarrow{F} A) \right) \in [0, 1].
$$

(2)

When $F$ is a deterministic system, $\mu(F) \equiv 1$ always holds true. In comparison, for a stochastic system with $a$ attractors, values as low as $1/a$ may be obtained; for stochastic systems with a single attractor, $\mu(F) \equiv 1$ because the single attractor is eventually approached from any initial configuration.

Most limit cycles that are attractors in a deterministic system are no longer attractors in an SDDS, because a cycle can be exited with a certain probability. Nevertheless, one particular kind of limit cycle, which consists of $2^k$ elements and in which all but $k$ bits are fixed, remains an attractor even in a SDDS. One such example is a 2-cycle, in which just one bit switches states, e.g., 000 $\leftrightarrow$ 001. Table\textsuperscript{4} shows that such limit cycles occur very rarely by chance, and for computational reasons, we decided to include only steady states and limit cycles of length $2^5$ or less in this study. This restriction does not influence the study since longer limit cycles that remain attractors in the SDDS practically do not occur.
TABLE I. This table shows the average number of steady states and limit cycles that remain attractors in the SDDS framework for different network sizes. We found fewer than thirty such 8-cycles among more than 250,000 networks of different sizes, and no 16-cycles at all. This shows that including only limit cycles of length 16 and less is not restricting the study.

| Network Size | 5     | 15    | 30    | 50    |
|--------------|-------|-------|-------|-------|
| Cycle Length |       |       |       |       |
| 1            | 2.8351| 3.6577| 4.3492| 4.8709|
| 2            | 0.1529| 0.1522| 0.1540| 0.1486|
| 4            | 0.0244| 0.0378| 0.0415| 0.0415|
| 8            | 0.0002| 0     | <0.0001| 0     |
| 16           | 0     | 0     | 0     | 0     |
| Sample Size  | 120000| 42000 | 40000 | 62500 |

The basic procedure underlying the computational study is, for a given SDDS, referred to as the “basic” network, to construct several “extended” networks by successively adding nodes, which we shall refer to as miRNAs, together with one or more feedforward loops involving the new miRNAs in a specific way. We then measure the change in the stochasticity measure described above. Let $F = (f_i, p_i^+, p_i^-)_{i=1}^{n_1}$ be the basic network, and let $F^* = (f_i^*, q_i^+, q_i^-)_{i=1}^{n_2}$ be the extended network, $n_1 \leq n_2$. We hypothesize that the dynamics in the feedforward loop enriched network are less stochastic. To compare the dynamics of two systems with respect to their degree of determinism, we consider their difference $m$ in $\mu$-values

$$m(F, F^*) = \begin{cases} 0 & \text{if } \mu(F^*) = 1 \\ \frac{\mu(F^*) - \mu(F)}{1 - \min(\mu(F), \mu(F^*))} & \text{otherwise.} \end{cases}$$

The denominator scales this difference into the range $[-1, 1]$. Here, $m = 0$ means that both networks have equally stochastic dynamics. If $m$ is positive, the extended network $F^*$ is dynamically less stochastic than the basic network $F$, and a negative value of $m$ suggests the opposite. The magnitude describes the difference in degree of stochasticity. A magnitude of 1 means that one of the networks has the dynamics of a deterministic system, whereas a magnitude of 0.5 suggests that one system is 50% less stochastic than the other.
III. Methods

For each set of input nodes we generated a certain number $N$ of basic Boolean SDDS $F$, introduced miRNAs, in a way that will be specified later in this section, to obtain the extended network $F^*$, and then compared their degree of determinism via $m(F, F^*)$. We considered 4 network sizes $n$: 5, 15, 30, 50 nodes. The corresponding number $N$ is 20000, 7000, 5500, 5500, respectively. The networks generated are random, subject to the following restrictions.

Large-scale studies of *B. subtilis*, *E. coli* and *S. cerevisiae* strongly suggest that the in-degrees of nodes in a transcriptional regulatory network are Poisson distributed with a mean of about two\(^{22}\). Thus, we chose a Poisson distribution with parameter $\gamma = 2$ for the basic network. The only other restriction is that each gene is regulated by at least one other gene, which raises the average in-degree to approximately 2.2. The regulators of each gene are randomly chosen from the set of all $n$ genes in the network, allowing self-regulation.

All propensity parameters for transcription factors and miRNAs are also randomly chosen from $[0, 1]$. For computational reasons, the full interval $[0, 1]$ is not used since a propensity parameter close to 0 might strongly decelerate convergence to attractors, slowing the performance of the simulation. However, 0.2 as the lower limit for the propensity parameters still allows one process to happen up to five times more frequently than another. Each gene is regulated by a certain number of genes, depending on its in-degree, and random Boolean functions that actually depend on all input variables are used as update functions.

After creating the basic network, we generate an extended network in a way reminiscent of post-transcriptional regulation by miRNAs. (Since we do not include a corresponding protein node for each gene node, the analogy is limited). Starting with the basic SDDS, miRNAs are iteratively added to this system by randomly choosing one gene as a transcription factor (TF) that induces transcription of the introduced miRNA. This miRNA, in turn, reduces the mRNA level of its own transcription factor; one example for such coregulation is the interplay between miR-133b and Pitx3 in midbrain dopamine neurons\(^{23}\). A lower TF mRNA level leads to a lower TF protein level, which then affects the regulation of all TF target genes. In the Boolean framework, a gene-specific threshold is used to distinguish between low (0) and high concentration (1). For some target genes, even after the TF mRNA reduction, there might still be enough transcription factor so that the target concentration is on the same side of the
TABLE II. Example of how an additional miRNA is embedded as an input variable in a target
gene’s update function. The light grey rectangle contains the original random update function,
in which \(a_1, \ldots, a_4 \in \{0, 1\}\) can be any Boolean values, with the sole restriction that the update
function must depend on both inputs. Unless both miRNA and transcription factor are present, the
miRNA has no influence since there is no TF mRNA that can be degraded or there is no miRNA
that can catalyze this degradation. Only if both are present (gray rows), can the miRNA reduce
the TF mRNA level to an extent that the target concentration changes because of the miRNA.

| miRNA(t) | TF(t) | other input(t) | target(t+1) |
|---------|-------|----------------|-------------|
| 0       | 0     | 0              | \(a_1\)     |
| 0       | 0     | 1              | \(a_2\)     |
| 0       | 1     | 0              | \(a_3\)     |
| 0       | 1     | 1              | \(a_4\)     |
| 1       | 0     | 0              | \(a_1\)     |
| 1       | 0     | 1              | \(a_2\)     |
| 1       | 1     | 0              | \(a_1\)     |
| 1       | 1     | 1              | \(a_2\)     |

threshold as if no reduction had taken place; for other target genes, the target concentration
might change significantly because of the TF mRNA reduction. Since this reduction is caused
by the miRNA presence, the miRNA becomes a new regulator of the target genes in the latter
case. One input variable in this study, called miRNA strength, describes the probability that
transcription factor-target gene pairs fall into this latter case, i.e., that the TF mRNA level
is significantly reduced, so that the Booleanized target concentration is the same as if no
transcription factor had been present at all. If, for instance, the miRNA strength is 0.5,
any miRNA regulates on average half of its transcription factor’s target genes. However, we
require each miRNA to regulate at least one target gene. This restriction ensures that each
miRNA is part of at least one feedforward loop, consisting of transcription factor, miRNA,
and target gene. Table III depicts an example of how the update function of target genes
regulated by a miRNA is expanded, taking into consideration the mode of action of miRNAs.
Only if miRNA is present and TF mRNA is transcribed, the mRNA reduction takes place.
In this case we see the same output as if no TF mRNA were present.

Because all networks with exactly one attractor already have totally deterministic dy-
namics, in the sense defined earlier, i.e., $\mu \equiv 1$ for those networks, only basic networks with multiple attractors are considered in this study. Those are the interesting networks, in which actual stabilization of the dynamics might be observed. Particularly interesting dynamics occur if at least two attractors possess a relatively large basin of attraction. The network selection process therefore favors networks with multiple large basins of attraction by picking a network only if at least two attractors are found more than once starting from twenty random initial configurations.

If the extended network $F^*$ does not possess multiple attractors, $m(F, F^*) = 1$ by definition. One could argue that the loss of attractors in the extended network is one feature of stabilization through feedforward loops. On the other hand, however, this could be seen as an experimental bias. To consider both views, we use $m$ to define two output measures, $m_1$ and $m_2$, one regarding any extended network and the other considering only those network pairs in which the extended network also possesses multiple attractors with at least two large basins of attraction. For a given set of input variables, we generate $N$ basic and extended networks, and measure

$$m_1 = \frac{1}{N} \sum_{i=1}^{N} \{ m(F, F^*) | \text{basic network } F \text{ and extended network } F^* \text{ have multiple attractors} \},$$

(4)

$$m_2 = \frac{1}{N} \sum_{i=1}^{N} \{ m(F, F^*) | \text{only the basic network } F \text{ is required to have multiple attractors} \}.$$  

(5)

For any set of input variables, we expect $m_2 \geq m_1$ since all network pairs with less than two attractors in the extended network, which are omitted in $m_1$, have $\mu \equiv 1$ and thus a mean value closer to 1. However, we do not want to prefer one or the other measure and thus we report results for both, which have been obtained independently, i.e., a network pair that was used for $m_1$ is not used for $m_2$.

A full analysis of the state space of a SDDS is only possible for small networks, so we used random sampling of initial configurations and an estimate of transition probabilities to attractors to approximate $m(F, F^*)$. We created a set of 100 random initial configurations, which were used in both networks to find the transition probabilities to attractors by updating each configuration 50 times, until an attractor was reached. In a small preliminary
study, we found that these two values yield a good trade-off between accuracy and efficiency.

IV. Results

Overall, we created over 300,000 pairs of basic and feedforward loop enriched networks. The results for networks with sizes ranging from 5 to 50 genes can be seen in Figure 1. None of these networks were required to be strongly connected, and in all of them the introduced miRNAs had full strength, meaning that each miRNA regulates all of its transcription factor’s target genes in a feedforward loop structure. The main result is that both measures, $m_1$ and $m_2$, are indeed positive for all network sizes and numbers of introduced miRNAs, indicating that miRNA-mediated FFLs can actually stabilize networks. It can also be seen that the impact of a single miRNA/FFL decreases when the network becomes larger. This means that larger networks require more miRNAs/FFLs for the same degree of stabilization.

Table III shows the results for networks of size 50. We see that more miRNAs and thus more FFLs stabilize the dynamics. Whereas one miRNA of full strength with $m_1 \approx 4\%$ only has a small impact, five such miRNAs already lead to $m_1 \approx 12\%$, and the introduction of thirty miRNAs of full strength stabilizes the stochastic system quite a lot ($m_1 \approx 0.37\%$). As expected, $m_2$ yields higher values and thirty miRNAs already reduce the stochasticity in the dynamics by more than 50% ($m_2 \approx 0.57\%$). In the case that each miRNA regulates on average only half its transcription factor’s target genes (but at least one), all values are considerably lower; the general behavior does not change, however.

These results raise the question whether a given network can be fully stabilized by introducing a sufficient number of miRNAs. Indeed, under certain conditions this is possible by ensuring the existence of a unique steady state. If an $n$-gene network contains no self-regulating genes, then $n$ miRNAs with full strength, each regulated by another gene, suffice to have fully deterministic network dynamics. Since the miRNA has full strength, it will downregulate any present mRNA, which ensures that only the value $a_1 \in \{0, 1\}$ (compare Table IV) can be taken by the target gene at a steady state. Each gene is regulated by at least one other gene. Hence, each gene and its regulated miRNA can only take the value $a_1$ in its truth table and the existence of a unique steady state is guaranteed. Thus, $\mu \equiv 1$ for such networks, which is equivalent to fully deterministic dynamics in the sense defined in
TABLE III. Comparison of the degree of stochasticity via $m_1$ and $m_2$ for not necessarily strongly connected networks of 50 genes, in which various numbers of miRNAs with full strength (Part a) and with strength 0.5 (Part b) are introduced. Overall, the more miRNA-mediated FFLs are introduced, the less stochastic the network dynamics become.

| Number of miRNAs | 1  | 3  | 5  | 8  | 10 | 15 | 30 |
|------------------|----|----|----|----|----|----|----|
| Average Number of FFLs | 2.401 | 7.298 | 12.36 | 19.79 | 24.77 | 37.27 | 74.50 |
| $m_1$            | 0.0374 | 0.0700 | 0.1083 | 0.1548 | 0.1756 | 0.2609 | 0.4042 |
| $m_2$            | 0.0687 | 0.1444 | 0.2178 | 0.2832 | 0.3203 | 0.4297 | 0.5967 |

| Number of miRNAs | 1  | 3  | 5  | 8  | 10 | 15 | 30 |
|------------------|----|----|----|----|----|----|----|
| Average Number of FFLs | 1.51 | 4.56 | 7.65 | 12.32 | 15.45 | 23.21 | 46.48 |
| $m_1$            | 0.0186 | 0.0523 | 0.0920 | 0.1064 | 0.1356 | 0.1653 | 0.2949 |
| $m_2$            | 0.0441 | 0.1138 | 0.1653 | 0.2052 | 0.2536 | 0.3111 | 0.4682 |

Section II

A. Derrida Values

In this study, we introduce a new measure for the robustness of stochastic networks by quantifying the degree of determinism of network dynamics. Another measure that can be used in the Boolean context was suggested by Derrida in 1986. Pairs of initial configurations of fixed Hamming distance are sampled from the entire state space, and their mean normalized Hamming distance, after being updated using update functions and propensity parameters, is defined as the Derrida value for a given initial Hamming distance. Lower Derrida values reflect more stable dynamics. To take time dependencies into account, we also considered the mean Hamming distance after two and three time steps as has been done earlier. Table displays the percent change in Derrida values starting with a basic 50-gene network and introducing 30 miRNAs. In all cases, the change is negative, i.e., the Derrida...
TABLE IV. If each of a target gene’s transcription factor regulates a miRNA that degrades the transcription factor mRNA, then only the fixed value $a_1 \in \{0,1\}$ can be taken on at a steady state because each transcription factor and its regulated miRNA have to take on the same value, 0 or 1, at a steady state (gray rows). Here, the light gray rectangle contains the original update function, and $a_1, \ldots, a_4$ are any Boolean values such that the update function depends on both inputs.

| miRNA1(t) | miRNA2(t) | TF1(t) | TF2(t) | target(t+1) |
|-----------|-----------|--------|--------|-------------|
| 0         | 0         | 0      | 0      | $a_1$       |
| 0         | 0         | 0      | 1      | $a_2$       |
| 0         | 0         | 1      | 0      | $a_3$       |
| 0         | 0         | 1      | 1      | $a_4$       |
| 0         | 1         | 0      | 0      | $a_1$       |
| 0         | 1         | 1      | 0      | $a_3$       |
| 0         | 1         | 1      | 1      | $a_3$       |
| 0         | 0         | 0      | 0      | $a_1$       |
| 1         | 0         | 0      | 1      | $a_2$       |
| 1         | 0         | 1      | 0      | $a_1$       |
| 1         | 0         | 1      | 1      | $a_2$       |
| 1         | 1         | 0      | 0      | $a_1$       |
| 1         | 1         | 0      | 1      | $a_1$       |
| 1         | 1         | 1      | 0      | $a_1$       |
| 1         | 1         | 1      | 1      | $a_1$       |

values decreased, indicating that the extended network exhibits more stable dynamics than the basic network, which we observed for different network sizes as well. Thus, another commonly used robustness measure also agrees with our hypothesis, which suggests that our findings are independent of the choice of robustness measure.
TABLE V. Derrida values for initial small disturbances of a Hamming distance up to 5 were simulated for a basic 50-gene network and an extended network with an additional 30 miRNAs of full strength. Multiple time steps were taken into account to consider time dependencies. The table shows that the percent change of the Derrida values from the basic to the extended network is always negative, indicating that our findings do not depend on the choice of the robustness measure.

| Hamming Distance | Time Steps |
|------------------|------------|
| 1                | 1          |
|                  | -2.8343    |
|                  | -5.5589    |
|                  | -7.7702    |
|                  | -9.6424    |
|                  | -11.1946   |
| 2                | -3.6709    |
|                  | -5.0620    |
|                  | -6.3319    |
|                  | -7.3764    |
|                  | -8.3597    |
| 3                | -6.4927    |
|                  | -7.4783    |
|                  | -8.3548    |
|                  | -9.0794    |
|                  | -9.8100    |

V. Discussion

We have examined the effect of feedforward loop motifs in stochastic Boolean network models of transcriptional networks, in analogy to the regulatory effects of miRNAs. Our goal was to test the hypothesis that these regulatory motifs have the effect of buffering the network against stochastic effects in the sense that they stabilize the basins of attraction. To this end, we conducted a computational experiment on a large number of randomly generated networks. The networks were modified by introducing additional nodes and feedforward motifs in a way that suggests regulation by miRNAs. To capture the effect on network stability we introduced a new measure of stochasticity of a network suitable for this purpose. Using this measure, as well as the classical measure of Derrida values, we showed that indeed the introduction of miRNAs has the hypothesized buffering effect.

The number of miRNAs that are introduced strongly influences the magnitude of the stabilizing effect, so that one might wonder how many feedforward loops can be expected to be found in actual gene regulatory networks. In a data set from *E. Coli*, among 424 nodes with 519 edges, 40 FFLs have been found. In *S. cerevisiae*, among 685 genes with 1,052 interactions, there are at least 70 FFLs. However, restricting the data to subnetworks, we find other occurrence frequencies of FFLs. A subnetwork of *E. Coli* of 67 nodes with 102 edges containing 42 FFLs was identified (some new FFLs had been found by then),
and in *D. melanogaster*, a subnetwork of 54 nodes and 167 edges contained as many as 157 FFLs. These numbers indicate that the question of how many FFLs are reasonable in a gene network of a certain size seems to depend strongly on the average in-degree of the nodes; whereas even large networks with average in-degree of less than 2 have few FFLs, this number can rise quickly when the network becomes more highly connected, as indicated by the considered network of *D. melanogaster*, with an average in-degree of approximately 3.

Additionally, we looked at the correlations between the number of attractors in both networks and the number of common attractors, where we defined a configuration in both networks to equal if the states of all genes, i.e., the first $n$ bits, coincide. Figure 2 shows the observed correlations, and we notice expected decreasing correlations between all three variables when more miRNAs are introduced. Surprisingly, the number of attractors of the extended network is much more strongly correlated with the number of common attractors than the respective number for the basic network, the cause of which remains to be explored.

This study can be extended in several ways, which we are planning to pursue. To make the study design more realistic, it is useful to introduce additional nodes for proteins, in order to be able to implement more mechanistic details of miRNA regulation. Also, here we do not restrict the regulatory rules to those that correspond to activation and inhibition only, which does not allow the classification of feedforward loops into coherent and incoherent, an important distinction. Also, a more careful study remains to be done on the effect of miRNAs relative to their position in the network and the local network topology into which they are embedded. Finally, another limitation of this work is that only intrinsic noise is being considered as a perturbation. It is important, however, to also take extrinsic noise into account, which requires an extension of the SDDS framework.

VI. Conclusions

This study provides computational evidence that miRNA-mediated feedforward loops have the effect of buffering the network against phenotypic variation due to stochastic effects. Introducing a feedforward loop motif has a local effect on network dynamics that propagates to a generally much smaller global effect on attractor stability. Thus, as the number of feedforward loop motifs increases, the overall stabilizing effect increases as well. In our
study, the number of miRNAs introduced is of a relative order of magnitude that might be expected in an actual transcriptional network. Thus, our computational experimental setup can be used in conjunction with an appropriate experimental system to investigate the effects of individual miRNA actions.

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FIG. 1. $m_1$ and $m_2$ are plotted against the number of introduced miRNAs. Networks are not necessarily strongly connected, and the miRNA has full strength. The size of the considered networks varies from 5 (solid line) to 50 (dotted line). The impact of a single FFL on the dynamics is larger in smaller networks, which suggests that larger networks require more FFLs for the same amount of stabilization.

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FIG. 2. The correlation between the number of attractors in the basic and the extended network and the number of common attractors is compared pairwise. These values have been obtained from not necessarily strongly connected networks of 50 genes and the introduced miRNAs had full strength.